Introduction

Mediastinal masses are mostly found in the anterior compartment and include several pathological entities. Tumors of the anterior mediastinum are more likely to be malignant, and more frequently comprise thymoma, lymphoma and germ cell tumors (1).

Clinical presentation of a mediastinal mass may be related to the compression or invasion of the adjacent structures leading, in some cases, to life-threatening emergencies. In other patients, these tumors may be heralded by non-local symptoms caused by paraneoplastic neurological diseases (PND) caused by an autoimmune response to antigens co-expressed by the cancer and the nervous system. PND may affect multiple levels of the nervous system from the cerebral cortex to the neuromuscular junction and the muscle. Early diagnosis of PND can be critical for the detection of a new or recurrent occult tumor. Neural-specific autoantibodies serve as diagnostic markers for PND and in some cases may predict the presence of a specific type of cancer. This review focuses on the clinical and immunological characteristics of the PND associated with the most frequent mediastinal cancers, namely thymoma, lymphoma and germ cell tumors.

The role of thymoma in autoimmunity

Immunological tolerance can be defined as the state of unresponsiveness to a self or non-self antigen. The thymus is the central organ of immunologic tolerance. Progenitor T cells migrate from the bone marrow to the thymus to mature and differentiate through interactions with cortical and medullary thymic epithelial cells (TECs). In this tightly regulated process T cells undergo positive and negative selection, and the cells with high reactivity to self-antigens...
(potential cause of autoimmunity) are eliminated (3). Thymomas originate from TECs and are distinct into histological subtypes according to the W.H.O. classification published in 1999 (4) and modified in 2004 (5).

The pathogenic mechanisms of thymoma-associated autoimmunity have been investigated in relation to myasthenia gravis (MG). Several abnormalities have been described in the molecular biology and immunocytochemistry of thymoma that can alter T-cell development. In MG-associated thymoma, neoplastic epithelial cells still express epitopes of the acetylcholine receptor (AChR) subunits and titin, but also show a decreased expression of the major histocompatibility complex (MHC) class II molecules, which are required for positive selection of T lymphocytes. Most thymoma neoplastic cells also fail to express the Autoimmune Regulator (AIRE) transcription factor, which is critical for T cell negative selection. Moreover, it has been shown that the production of regulatory T-lymphocytes, which suppress the immune response toward self antigens are diminished in thymomas (6).

PND associated with thymoma

The neurologists’ interest in thymoma was initially stirred by its association with MG. How this rare, slow-growing tumor came to be involved in the pathogenesis of a prototypical autoimmune disease has long been a matter of debate. Later, it became evident that thymoma was central to a broader disease spectrum and what we learnt from paraneoplastic MG paved the way to the understanding of the mechanisms of PND of the peripheral and central nervous system (CNS). All these conditions, as well as non-neurological thymoma-associated PDs, can also occur as idiopathic non-paraneoplastic diseases.

Disorders of the neuro-muscular transmission

Myasthenia Gravis

MG is the most common thymoma-associated disease, as up to 40% of individuals with thymoma develop MG, while 15–20% of MG patients are diagnosed with thymoma.

MG is a disorder of the neuromuscular junction, caused by antibodies (Abs) against extracellular determinants of postsynaptic proteins. The antibody attack results, through different mechanisms, in morphological and functional alterations leading, in turn, to the impairment of the neuromuscular transmission.

The clinical hallmark of MG is fluctuating muscle weakness, worsened by exertion and relieved by rest. The more common involvement of certain muscle groups (extrinsic ocular, facial, oropharyngeal, neck, limb proximal muscles) accounts for a typical clinical picture in most cases. On the other hand, weakness severity shows remarkable variability, from purely ocular symptoms to severe generalized disease. Weakness of respiratory muscles can lead to respiratory failure (so-called myasthenic crisis) requiring assisted ventilation (7). Around 90% of MG patients have serum Abs to the AChR. These Abs induce severe alterations of the post-synaptic membrane through complement activation and increased AChR degradation, and, to a lesser extent, by interfering with the ACh binding site (8). MG with AChR Abs (AChR-MG) is frequently associated with thymus alterations, such as follicular hyperplasia and thymoma, both playing a role in the disease pathogenesis (9).

In anti-AChR negative patients, Abs to other synaptic proteins (such as the muscle-specific tyrosine kinase receptor, MuSK, and the low-density lipoprotein receptor-related protein 4, LRP4) can be detected and identify distinct disease subtypes (10). The MuSK-LRP4-Agrin complex is essential for the neuromuscular junction formation and its maintenance in adult life. MuSK, activated by the nerve secreted agrin through its co-receptor LRP4, promotes AChR clustering (11), and LRP4 delivers retrograde signals contributing to presynaptic differentiation (12). Five to seven percent of anti-AChR negative patients have Abs to MuSK, serum Abs to LRP4 can be found in a low proportion of anti-AChR and -MuSK negative cases and, more recently, serum immunoglobulin G (IgG) to agrin have been reported (10). In addition, a few patients with clinical and electrophysiological signs of MG do not have detectable Abs (seronegative MG). Currently, there are no data supporting a pathogenic link with the thymus for all these forms of MG.

Despite isolated cases of thymoma have been described in MuSK-MG (13), LRP4-MG (14) and in the seronegative disease (15), the association with thymoma is essentially restricted to AChR-MG.

Clinical and pathogenic aspects of thymoma-associated AChR-MG

In the great majority of patients, the onset of MG leads to thymoma detection, as a study of the mediastinum by computed tomography scan is routinely performed upon the neurological diagnosis. More rarely, MG presentation
occurs months, and even years, after thymoma treatment and can be a sign of tumor recurrence.

In MG patients, the risk of having a thymoma is low in the first two decades and after 70 years of age; it increases in between, with a peak incidence in the fourth through the sixth decade. Figure 1 shows the age at thymoma diagnosis in 256 MG patients seen in our institution between 1980 and 2016. As shown in the figure, there was no gender bias in our population. This finding has been reported in other thymoma-associated MG cohorts (16,17).

Thymoma-associated MG is generally severe, with a high rate of life-threatening symptoms, and patients tend to remain dependent on immunosuppressive treatment. A broad range of other neurological and non-neurological PDs can occur in these cases, both before and after tumor treatment. The onset of any of these disorders as well as a serious worsening of MG symptoms may predict a tumor recurrence and should prompt specific imaging studies.

Patients with AChR-MG often have serum Abs to the giant muscle protein titin and the ryanodine receptor (RyR). These Abs, formerly called anti-striational Abs, are strongly associated with MG-related thymoma (anti-titin are positive in 95% and anti-RyR Abs in 70% of these patients), are detected also in nearly 50% of late-onset non-thymoma cases, while are very uncommon in early-onset MG (18,19). As anti-striational Abs target intracellular antigens their pathogenicity is uncertain, though they are valuable markers of thymoma, at least in young MG patients (20).

Kv1.4 Abs that target the alpha subunit of the muscle voltage-gated potassium channel (VGKC) were reported in 12–15% of Japanese MG patients, in association with thymoma, severe MG and myocarditis with arrhythmias (21,22). Interestingly, these findings were not confirmed in Caucasian patients (23).

Thymoma-associated MG is thought to be caused by defects of central tolerance related to the neoplastic thymus microenvironment. Similar mechanisms may be operating in other PDs. As mentioned before, in the normal thymus, developing T lymphocytes go through sequential maturation stages from CD4⁺CD8⁺ double positive (DP) cells in the thymic cortex, to CD4⁺CD8⁻ and CD4⁻CD8⁺ single positive (SP) T cells in the medulla. Positive and negative selection processes secure that (mostly) self-tolerant SP T cells are released into the periphery. Stromal cells, such as TECs and dendritic cells are crucial to the shaping of T lymphocyte repertoire (24).

Thymoma subtypes differ in their thymopoietic capacity (22). AB and B thymomas are able to support thymocyte development and harbor a high number of DP cortical T cells. Conversely, thymopoietic function is very low in A thymoma and nearly absent in thymic carcinoma (16).

Figure 2 shows thymoma histological subtypes in our population. All thymoma specimens were reviewed over the time by the local pathologists and classified according to the WHO system (4,5).

B thymomas are the most common subtypes related to MG (see Figure 2), and both intratumorous T lymphocyte
maturation and export of mature T cells appear to be prerequisites for paraneoplastic MG development (16). Although neoplastic TECs express single epitopes of AChR subunits and the protein titin, thymoma tissue has a low expression of MHC class II molecules and AIRE gene, both required for an effective thymocyte selection (17). Defective medullary function in “cortical” B thymomas (that have sparse medullary areas) can be responsible for the export of autoreactive CD4+ SP T cells and for a reduced production of T regulatory cells. Such unbalance appears to be a distinctive feature of MG-associated thymomas (17).

Protein tyrosine phosphatase non-receptor 22 (PTPN22) and cytotoxic T lymphocyte associated antigen 4 (CTLA-4) are both inhibitors of T lymphocyte activation. Variants of these genes were reported in association with different autoimmune diseases, including MG (25,26). CTLA-4+49A>G (27,28) and two different PTPN22 genotypes (28,29) were found to be associated with MG and thymoma. In addition, peripheral and intra-tumorous T cells from MG thymoma patients had higher expression levels of the anti-apoptotic factor cellular FLICE-like inhibitory protein (c-FLIP) than non-MG thymomas (30). These results confirm the role of activated T cells in the pathogenesis of thymoma-associated autoimmune diseases.

Few patients with MG may have a thymolipoma, a rare benign tumor of the thymus containing mature adipose tissue and thymic remnants. In these cases, MG is generally anti-AChR positive, and its severity and outcome do not seem to differ from those of the thymoma associated disease (31,32).

**Lambert-Eaton myasthenic syndrome (LEMS)**

LEMS is a rare disorder of the neuromuscular and autonomic transmission, in which Abs to the voltage-gated calcium channel (VGCC) at the nerve terminal markedly reduce acetylcholine release. These patients complain of muscle weakness typically prevalent on leg proximal muscles, loss of tendon reflexes, and autonomic disturbances (33). Paraneoplastic LEMS is generally related to small-cell lung cancer (33), but the association with thymic tumors has rarely been reported (34,35). In these patients, oncologic treatment resulted in marked clinical benefit.

**Disorders of peripheral and CNS**

Acquired neuromyotonia (aNMT) and the Morvan’s syndrome (MoS) are well characterized disorders due to peripheral nerve hyperexcitability. These conditions, when paraneoplastic, are generally associated with thymoma. The earliest evidence for NMT to be an antibody-mediated disease derived from studies of passive transfer of disease-related electrophysiological abnormalities to mice with patients’ IgG. Then, NMT and MoS have been recognized as autoimmune channelopathies since the identification of autoAbs binding to the VGKCs (36).

These Abs were originally detected with a radioimmunoassay in which VGKCs solubilized from mammalian brain membranes and labelled with iodinated-dendrotoxin were precipitated by patients’ IgG. The Abs were believed to bind directly to alpha dendrotoxin-bound VGKC subunits (Kv 1.1, 1.2 and 1.6). However,
it has recently been shown that patients' serum IgG most frequently bind to the extracellular domains of leucine rich glioma inactivated protein 1 (LGI1) and contactin-associated protein-like 2 (CASPR2), two proteins complexed with VGKCs (37,38). Indeed, CASPR2 Abs are more commonly detected in patients with NMT and MoS associated with thymoma (39), while LGI1 Abs are usually found in patients with non-paraneoplastic autoimmune encephalitis (38). In addition, Abs to the Netrin-1 receptors (DCC and UNC5A) have lately been reported in NMT or MoS occurring in patients with thymoma-associated MG (40). The expression of DCC, UNC5A, and Caspr2 proteins was demonstrated both in thymoma samples and normal thymus (40). The pathogenic role of the Netrin-1 receptor Abs has not been investigated.

Acquired neuromyotonia
aNMT is characterized by spontaneous muscle activity due to continuous nerve depolarization. Clinical presentation includes myokymia (diffuse muscle twitching), fasciculations and cramps; hyperhidrosis is common and patients with long-standing disease can develop muscle hypertrophy (41). Electromyography shows spontaneous motor unit discharges as doublet, triplet, multiplet bursts (myokymic discharges), and, less commonly, longer bursts with high intraburst frequency (neuromyotonic discharges) (42). In up to 20% of cases aNMT is associated with signs of CNS involvement, such as mood changes and sleep disturbance. aNMT generally occur in association with MG in patients with thymoma. Importantly, both the onset and worsening of aNMT can herald a tumor recurrence (43).

NMT symptoms usually improve with sodium channel-blockings as carbamazepine, phenytoin or lamotrigine. In patients with severe symptoms plasma-exchange and immunosuppression with corticosteroids and azathioprine can be required (44). When associated with thymoma, oncologic treatment can significantly improve the neurological diseases.

Morvan’s syndrome
In patients with MoS, NMT is associated with encephalopathy, sleep disorder with prolonged periods of insomnia (agrypnia), dysautonomia (hyperhidrosis, sphincter dysfunction), and/or pain. CASPR2 Abs are most frequently detected in these patients, often in association with LGI1 Abs. A thymoma can be detected in up to 50% of patients with CASPR2 Abs, generally in association with MG (45).

Both peripheral and central symptoms respond to plasma-exchange and immunosuppressive treatment, and may improve further with thymoma treatment.

Encephalitis
Paraneoplastic encephalitis usually presents with subacute (days to weeks) memory impairment associated with seizures and mood or behavioral disturbances (46).

Some patients develop hallucinations, delusions, and bizarre behavior at the onset of their illness and may initially be referred to a psychiatrist. Seizures and alterations in the level of consciousness ensue in most cases. Brain MRI typically shows hyperintense lesions of the medial temporal lobes on T2-weighted images, but can otherwise be negative. Cerebrospinal fluid (CSF) analysis may detect increased protein content or moderate lymphocytic pleocytosis, and in a minority of patients may show the presence of oligoclonal bands. Thymoma-associated autoimmune encephalitis is generally associated with pathogenic autoAbs binding to membrane neural antigens and often responds favorably to immunotherapy and anti-neoplastic therapy. The presence of Abs to intracellular antigens is rarer.

Serum IgG specific for the extracellular domain of the VGKC-complex proteins LGI1 and CASPR2 are the most commonly Abs detected in patients with thymoma-associated encephalitis. In particular, while LGI1 Abs are mostly detected in non-paraneoplastic encephalitis, patients with thymoma more often have Abs binding to CASPR2 or to both LGI1 and CASPR2 (Figure 3). In a recent analysis performed in 256 patients with LGI1-IgG and/or CASPR2-IgG a thymoma was detected in 33% of patients with both LGI1 and CASPR2-Abs, in 5% of patients with isolated CASPR2- and in less than 1% of patients with isolated LGI1 Abs (47).

Other neuronal autoAbs binding to neuronal membrane proteins detected in patients with autoimmune encephalitis and thymoma are specific to AMPA (AMPA-R) and GABA (GABA-A) receptors. Encephalitis associated with AMPAR-IgG usually presents like classical limbic encephalitis with memory disturbances, alteration of mood and seizures. In up to 40% of patients with AMPAR-IgG a diffuse encephalopathy can be observed, while a minority of patients can develop a different syndrome, such as rapidly progressive dementia or psychosis. Patients with GABAAR-IgG usually develop seizures refractory to standard antiepileptic therapy (48). MRI may reveal multifocal cortical and juxta-cortical hyperintense lesions on T2-weighted
images. Most patients respond to immunotherapy. Thymoma has been detected in around 30% of adult patients with GABAAR Abs (49).

AutoAbs binding to the collapsin response mediator protein 5 (CRMP5), also known as CV2, an intracellular protein involved in the regulation of dendritic development and synaptic plasticity, can be observed in patients with thymoma and encephalitis. These patients can present with limbic encephalitis or chorea and immunotherapy is often beneficial (50,51).

**Inflammatory muscle diseases**

Different autoimmune muscle diseases, such as dermatomyositis (52), granulomatous myositis (53) and polymyositis (54) have been reported in association with thymic tumors, with and without MG. All these disorders manifest with muscle weakness, myalgia, increased serum levels of muscle enzymes and abnormal hyperintense on STIR MRI. Cardiac muscle may be involved with heart failure and arrhythmias (54). In these cases, a role for anti-striational Abs has been suggested (21) but it remains uncertain owing to the lack of systematic studies.

**PND associated with lymphoma**

PND are a rare accompaniment of Hodgkin lymphoma (HL) and non-HLs (NHLs) (55). Paraneoplastic cerebellar degeneration (PCD) is the most frequent PND in patients with HL (56). PCD represents a classical PND and occurs most frequently in patients with ovarian, breast and small cell lung cancer or HL, while only few cases of PCD associated with NHL have been reported to date (57). PCD patients usually develop dizziness and vertigo that rapidly progress to severe ataxia, dysarthria, diplopia, and nystagmus. CSF examination can reveal increased protein concentration or mild pleocytosis, and the MRI studies are initially normal while cerebellar atrophy can be observed later in the course of the disease. The symptoms of PCD may precede the diagnosis of HL in up to 80% of patients (58). The most common autoAb detected in patients with PCD and HL are Tr-IgG that bind to the Delta and Notch-like epidermal growth factor-related receptor (DNER) (59), a membrane protein highly expressed in the dendrites of...
Purkinje cells that is critical for the development of the cerebellum.

IgG specific for the metabotropic glutamate receptor 1 (mGluR1) were originally identified in two patients with PCD and HL. To date 16 patients with mGluR1 Abs have been described and 6 of them had a history of hematologic malignancy (HL, 2; NHL, 1; T-cell cutaneous lymphoma, 2; acute lymphocytic leukemia, 1) (60-63).

**Paraneoplastic neurological syndromes associated with mediastinal germ cell tumors**

Mediastinal germ cell cancers are rare and heterogeneous. In a retrospective multicenter study performed in France to establish the frequency and the prognosis of primary mediastinal germ cell tumors, non-seminomatous germ cell tumors (NSGT) were detected in more than 60% of the patients, followed by seminomas observed in 32%, while thoracic teratomas represented the most rare cancer types (64). Few patients with paraneoplastic encephalitis associated with Abs binding to Ma2 or to NMDA receptor (NMDAR) and mediastinal germ cell tumor have been reported to date.

Encephalitis with anti-Ma2 Abs may be associated with seminoma or NSGT. Patients with Ma2-IgG usually develop a limbic encephalitis associated with symptoms due to diencephalic involvement (e.g., hyperthermia, narcolepsy and syndrome of inappropriate antidiuresis) or brainstem dysfunction while some patients can present with symptoms of brainstem dysfunction (e.g., ataxia, diplopia, dysphagia, dysarthria) (65).

Anti-NMDAR encephalitis has been rarely reported in patients, mainly males, with thoracic teratoma. This form of autoimmune encephalitis has characteristic clinical features. Patients develop behavioral disturbances and then seizures often refractory to standard anti-epileptic therapy and movement disorders (oro-buccal dyskinesias and dystonic postures) (66). Coma, central hypoventilation and dysautonomia can occur in the course of the disease leading to patient admission to ICU or even to patient death if appropriate treatment is not performed. There are no standardized guidelines for the treatment of patients with anti-NMDAR encephalitis. Plasma exchange, intravenous immunoglobulin and high dose intravenous steroids associated with cancer removal can result in the improvement of the neurological deficits (67). Rituximab or cyclophosphamide can be efficacious treatments for patients not responding to first-line therapies.

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**Footnote**

*Conflicts of Interest:* The authors have no conflicts of interest to declare.

**References**


