Dear Sir,

Stiff man syndrome is a rare neurological pathology first described in 1956 by Moersch and Woltman [1]. Four forms of the illness have been identified: stiff person syndrome, which is characterized by contractions and axial muscle rigidity; progressive encephalomyelitis with rigidity, characterized by diffuse rigidity and cognitive disorders; jerking stiff man syndrome, characterized by limb myoclonus and respiratory disorders; and stiff leg syndrome (SLS), which is characterized by focal rigidity and is different in various ways [2]. Anti-GAD antibodies are positive in only 15% of cases of SLS. Contractions and rigidity affect only one or two limbs. A paraneoplastic etiology is rarely associated with it. Medullary and cerebral MRI do not usually show any abnormalities [3].

We report the case of a patient suffering from SLS with anti-amphiphysin antibodies revealing breast cancer, with paraneoplastic transverse myelitis (TM) images appearing later in MRI with no associated neurological deficit.

**Case Report**

Mrs. H., aged 65, presented to the Neurology Department in July 2009 with painful stiffness in the lower limbs and difficulty in walking. The symptoms had first appeared 2 weeks earlier. Her medical history included type 2 diabetes treated with metformin and a thyroid goiter treated with surgery and levothyroxine.

On clinical examination, she presented permanent hypertonia in the lower limb and bilateral dystonia of the extensor hallucis longus muscles. Intermittent spasms were associated with hypertonia. The spasms lasted several seconds and were spontaneous, painful, and made worse by anxiety and cutaneous stimulation. The tibialis anterior muscles were hypertrophied.

Electroneuromyography (ENMG) carried out under diazepam was normal. Cerebral and medullary MRI (fig. 1) were normal. A PET scan showed an anomaly on the right breast. A mammography, breast ultrasound and biopsy confirmed the diagnosis of infiltrating canal carcinoma of the breast.

The standard biological assessment was normal. Cerebrospinal fluid (CSF) tests showed pleocytosis (14 white blood cells), a high protein level (0.81 g/l) and intrathecal IgG synthesis with monoclonal bands. Other biological examinations were done: Lyme disease, HTLV (1 and 2) and tetanus serology were negative. Tests for antiphospholipid, antinuclear and anti-GAD antibodies were negative. No antineural antibodies (Hu, Yo, Ri, CV2, Ma1, Ma2) were found, but anti-amphiphysin antibodies were found in blood and CSF (qualitative measurement).

Intravenous corticotherapy (1 g/day for 5 days), associated with diazepam (5 mg 3 times per day) and baclofene (10 mg 3 times per day), was introduced. Spasms receded after 3 days, and the patient was able to walk again.

Evolution was marked by frequent relapses with profuse sweating without hyperthermia, treated by corticotherapy, diazepam and baclofene. Another ENMG showed simultaneous activity in the agonist and antagonist muscles and continuous activity in anterior tibialis muscles with intermittent spasms triggered after cutaneous stimulation.

Spasms in the legs exacerbated by emotion and cutaneous stimulation, hypertonia of the lower limbs, adrenergic signs, ENMG data, amphiphysin antibodies and pharmacosensitivity led to a diagnosis of SLS. Cancer was treated by surgery, radiotherapy and chemotherapy. Symptoms improved with the introduction of chemotherapy. After the fifth round, SLS worsened with no associated neurological deficit (pyramidal, sensitive or sphincteric).

Six months after disease onset, the patient complained of leg pain; therefore, an...
other medullary MRI was carried out, which showed an unexpected TM with a T2 hypersignal from level Th6 to Th10 (fig. 1). Anti-NMO antibodies were negative. As there was no worsening of SLS and no associated neurological deficit, TM was considered as an incidental finding and no specific treatment was administrated.

One year after disease onset, the patient could walk with two sticks and no longer presented with spasms; however, she still suffered from stiff lower limbs with no pyramidal syndrome. Her only treatment was tizanidine. No further medullary abnormalities were found in the medullary MRI carried out 3 months later.

**Discussion**

Our patient had two paraneoplastic neurological presentations caused by anti-amphiphysin antibodies: firstly SLS, and secondly by extensive TM in imaging. The only sign of TM was nonspecific leg pain with no sign of associated medullary suffering and no worsening of SLS. To our knowledge, this is the first case of SLS and TM to be described in the literature. We suggest that a common medullary physiopathology may occur in these two pathologies.

The literature [3–19], including this case report, provides details of 31 patients: 23 women and 8 men (sex ratio: 1/3), with an average age of 49 (28–71), suffering from SLS. The search for anti-GAD antibodies was positive in 13 patients (42%). However, the search for anti-amphiphysin antibodies was carried out in only 10 patients and was positive in only 2. Our patient was the only one to have pleocytosis. In 1 of 27 patients, medullary MRI showed a dispersed T2 hypersignal [3]. Our patient was the only one to undergo another MRI showing extensive TM 6 months after symptom onset. Virani et al. [20] reported a case of extensive paraneoplastic TM with anti-amphiphysin antibodies in breast cancer with stiffness and spasms, as well as sensory disorders. Holmøy et al. [21] described a case of SLS with an anatomopathological analysis of the spinal cord showing infiltration of CD8+ T cytotoxic lymphocytes, chromatolysis, vacuolization of anterior horn cells and gliosis.

The largest series of myelitis with anti-amphiphysin antibodies reported 17 cases in 63 patients with anti-amphiphysin autoimmunity [22]. Presentations were generally subacute with predominantly motor involvement. CSF tests were usually mildly pleocytic with an increased protein level. Seven patients underwent a medullary MRI examination. Five had T2-weighted signal abnormality in the spinal cord and two had normal imaging but clinical symptoms of myelitis. The authors concluded that patients with isolated anti-amphiphysin antibodies are more likely to be women with breast cancer and to have stiff man syndrome or myelopathies compared to patients with anti-amphiphysin antibodies coexisting with autoantibodies [22]. Pittock and Lucchinetti [23] reported that patients presenting with paraneoplastic longitudinally extensive TM are often anti-amphiphysin positive.

The role of anti-amphiphysin antibodies is not fully understood, but injection of

![Fig. 1. Medullar MRI a initially normal and b 6 months after disease onset showing extensive transverse myelitis with T2 hypersignal from level Th6 to Th10.](image-url)
anti-amphiphysin antibodies causes stiff person syndrome in murine models [24]. Medullary inflammatory reaction could be caused by these antibodies or by neurone hyperactivity leading to excitotoxicity.

This overlap between SLS and paraneoplastic TM with anti-amphiphysin antibodies suggests a common physiopathology of anti-amphiphysin antibodies in stiff man syndrome, SLS and TM. In our case, SLS and TM are probably epiphenomena caused by diffuse spinal involvement induced by anti-amphiphysin autoimmunity. SLS and TM are both medullar pathologies, suggesting that anti-amphiphysin antibodies target epitopes that are highly expressed in the spinal cord. These antibodies may induce two physiopathological mechanisms: (1) functional abnormalities, leading to motor hyperexcitability resulting in SLS; and (2) toxic lesions, causing TM. Both mechanisms inducing SLS and TM are associated in our case report.

Acknowledgement

The authors wish to thank Frances Sheppard of the Clinical Investigation Center of Besançon (INSERM CIT 808) for correcting and improving the English in the manuscript.

Disclosure Statement

The authors have nothing to disclose.

References


