Neurological disorders associated with glutamic acid decarboxylase antibodies: a Brazilian series

Distúrbios neurológicos associadas a anticorpos anti-GAD (descarboxilase do ácido glutâmico): uma série de casos brasileira

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ABSTRACT
Neurological disorders associated with glutamic acid decarboxylase (GAD) antibodies are rare pleomorphic diseases of uncertain cause, of which stiff-person syndrome (SPS) is the best-known. Here, we described nine consecutive cases of neurological disorders associated with anti-GAD, including nine patients with SPS and three cases with cerebellar ataxia. Additionally, four had hypothyroidism, three epilepsy, two diabetes mellitus and two axial myoclonus.

Key words: glutamic acid decarboxylase antibodies, anti-GAD, neurological disorders, stiff-person syndrome, cerebellar ataxia.

RESUMO
Distúrbios neurológicos associados com anticorpos anti-GAD são doenças pleomórficas, raras, de causa incerta, das quais a rigidez muscular espasmódica (SPR) é a mais conhecida. Neste estudo, descrevemos nove casos consecutivos de distúrbios neurológicos associados com a presença de anticorpos anti-GAD, incluindo nove pacientes com SPR e três casos com ataxia cerebelar. Adicionalmente, foram encontrados quatro casos com hipotireoidismo, três com epilepsia, dois com diabetes mellitus e dois casos com mioclonia axial.

Palavras-Chave: anticorpos anti-descarboxilase do ácido glutâmico, anti-GAD, distúrbios neurológicos, rigidez muscular espasmódica, ataxia cerebelar.

Neurological disorders associated with glutamic acid decarboxylase (GAD) antibodies include stiff-person syndrome (SPS) and its variants — stiff trunk syndrome, stiff limbs syndrome, progressive encephalomyelitis with rigidity, SPS-plus and paraneoplastic SPS — as well as epilepsy and ataxia¹⁻⁸. These syndromes are often found in association with other autoimmune disorders, such as diabetes mellitus and autoimmune thyroid disease. These autoimmune disorders, which often occur in association with SPS, are caused by glutamic acid decarboxylase (GAD) and amphiphysin (presynaptic antigens) and GABA(A) receptor-associated protein (GABARAP) and gephyrin (postsynaptic antigens). The autoantibodies anti-amphiphysin and anti-gephyrin have been linked to paraneoplastic SPS⁹⁻¹⁵.

The pathogenetic role of autoantibodies in SPS is unclear. However, all SPS autoantigens identified to date are synaptic proteins involved in inhibitory synaptic transmission, such as GAD and amphiphysin (presynaptic antigens) and GABA(A) receptor-associated protein (GABARAP) and gephyrin (postsynaptic antigens). The autoantibodies anti-amphiphysin and anti-gephyrin have been linked to paraneoplastic SPS⁹⁻¹⁵.

The objective of this study was to present the clinical and laboratory findings of 12 patients with neurological disorders associated with anti-GAD antibodies.

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METHODS

Twelve consecutive patients with neurological disorders associated with anti-GAD antibodies were evaluated over a 17-year period at the Movement Disorders Unit in the Neurology Service, Hospital de Clínicas, Universidade Federal do Paraná, Pontifícia Universidade Católica do Paraná (Curitiba, PR, Brazil) and Medical School of UNIOESTE (Cascavel, Paraná, Brazil). Extensive workups, including neuroimaging, cerebrospinal fluid (CSF) analysis, electroencephalography (EEG) tracing, electroneuromyography (ENMG) and immunological tests (serum anti-GAD, antithyroid, antinuclear and antiparietal cell antibodies), were carried out.

The Dalakas criteria\textsuperscript{16} for the diagnosis of typical SPS were used. SPS variants were defined in accordance with Baker et al.\textsuperscript{5} as stiff trunk syndrome, stiff limb syndrome, progressive encephalomyelitis with rigidity and paraneoplastic SPS.

Seven patients were females. Mean age of diagnosis was 47.25 years (11 to 58 years). Two of the patients have been previously described\textsuperscript{17,18}.

RESULTS

Among a total of nine SPS cases, four presented the classic form and five the variants (two cases of stiff lower limbs, two of stiff one-limb and one of stiff three-limbs). Additionally, we found three cases of cerebellar ataxia. No cases of progressive encephalomyelitis with rigidity or paraneoplastic SPS were found. With one exception, all patients had elevated serum anti-GAD titers, ranging from 20 to 800 U/mL (normal \(\leq 1\)).

ENMG results were compatible with a diagnosis of SPS in six cases. Neuroimaging and EEG tracings were normal in all the patients. CSF analysis revealed nonspecific changes (mild pleocytosis) in two cases.

Three patients suffered from epileptic seizures (one with the classic form of SPS, the other with stiff lower limb variant and the last with cerebellar ataxia). Two patients (one with the classic form and the other with stiff lower limb variant) had segmental axial myoclonus.

A total of five patients had clinical and laboratory signs of autoimmune endocrinopathy (four patients with Hashimoto’s thyroiditis, two with diabetes mellitus).

The patients with SPS were treated with diazepam (15 to 30 mg/day), baclofen (30 to 60 mg/day) and intravenous immunoglobulin (IVIg, 0.4 g/kg/day for 5 days, followed by single monthly doses). Among the nine SPS patients, seven showed significant clinical improvement and two (with stiff one-limb) were partially responsive. The patients with cerebellar ataxia were treated with IVIg and responded well to this therapy. The Table summarizes the results for all the patients.

Two illustrative cases of neurological disorders associated with anti-GAD antibodies are described below.

ILLUSTRATIVE CASES

CASE 1

A 50-year-old female patient with four years history of progressively worsening signs and symptoms, consisting lumbar muscles contractures, pain and difficulty walking, sought medical attention in her hometown, initially with suspected postural lumbago and later with a possible diagnosis of ankylosing spondylitis. As her condition progressed, increased contracture and rigidity of the paravertebral muscles was observed (mainly in the lumbar region) with hyperlordosis (Fig 1). Her condition fluctuated, alternating between periods when the muscular contracture and rigidity worsened and she experienced pain and difficulty walking, with frequent falls, and periods when there was a partial improvement. She had a positive family history of breast cancer. She had also been treated for breast cancer, and there was no evidence of a relapse. On her latest assessment, in addition to muscular rigidity and bilateral contractures in the lumbar muscles, the patient presented ataxia with increase in the anteroposterior and lateral sway and wide based gait. She had been treated with baclofen (60 mg/day) and physical therapy, but with no significant improvement. On physical examination, she had a normal abdominal musculature with increased muscular atrophy in the lower limbs and foot deformities. Her mental status was normal and she had no signs of cognitive impairment. She had not received any immunomodulatory agents. On her last laboratory evaluation, she had increased serum anti-GAD titers (800 U/mL), with normal thyroid function tests (TSH, FT3, FT4). Her condition has been fluctuating over the last six months.

Table. Neurological disorders associated with anti-GAD antibodies.

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<td>3</td>
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<td>M/F</td>
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<td>57/58</td>
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<td>-</td>
<td>-</td>
<td>NC</td>
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<tr>
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Synd.: syndrome; M: male, F: female; EMG: electromyography; CSF: cerebrospinal fluid; EEG: electroencephalogram; Ab: antibodies; HT: Hashimoto’s thyroiditis; NC: not carried out.
The paravertebral region, particularly the lumbar region and lower limbs, she sometimes has sudden, jerky, movements and falls to the ground. She reports increasing difficulty walking because of severe muscular rigidity and is taking baclofen (40 mg/day) and diazepam (30 mg/day).

The patient underwent extensive additional tests, including MR neuroimaging of the skull and cervical/thoracic and lumbosacral spine, which revealed L4-L5 discopathy. The following tests were unremarkable: CSF analysis, complete blood count, fasting blood sugar, lipid profile, creatinine, TSH, T3, T4F, TGO, TGP, amylase, VDRL, electrolytes, mammography, chest X-ray, abdominal echography, whole-body PET-CT, antinuclear antibody (ANA), protein electrophoresis, muscle enzymes, erythrocyte sedimentation rate (ESR) and C-reactive protein (CRP). Bone densitometry showed osteopenia. ENMG showed the presence of continuous muscular activity (Fig 2). The patient is receiving IVIg therapy with a progressive improvement in her clinical picture.

**CASE 2**

A 52-year-old female patient who presented initially with diplopia had difficulty walking and had progressively worsening imbalance, which deteriorated markedly over 20 days. She had an ophthalmic assessment, which proved inconclusive, and was referred for neurological assessment in her hometown. This revealed gait ataxia, difficulty with tandem gait and the presence of horizontal gaze-evoked nystagmus, as well as a complaint of diplopia. The patient was receiving thyroid hormone therapy (100 μg/day). Further investigation revealed normal routine blood tests, except for ESR (60). Brain neuroimaging (CT and MRI) was normal. CSF analysis revealed the presence of 16 cells (100% lymphomononuclear), but no other abnormalities. Over time the patient complained of nausea and vomiting, and used an antiemetic (metoclopramide), dimenhydrinate and clonazepam. Viral cerebellitis was suspected. She later presented with a clinical picture of urinary infection, which subsequently improved with norfloxacin. The clinical picture of ataxia remained unchanged, and 30 days later the patient had a generalized tonic-clonic seizure while asleep, which was repeated the following day. The patient was hospitalized and phenytoin was administered intravenous (i.v.). Neurological examination revealed cerebellar ataxia with nystagmus and diplopia; a new brain MRI was normal. Phenytoin was replaced by lamotrigine in a daily dose of 200 mg, and methylprednisolone pulse therapy (1 g/day for 5 days) was started. At the end of the five days, the methylprednisolone was substituted by prednisone (40 mg/day). New CSF tests were unremarkable, and a further detailed investigation, including tests for vasculitis and screening for occult neoplasms, infections (serologic tests to detect antibodies against HIV, Epstein-Barr, varicella-zoster and herpes simplex viruses) and Hu, Ri and Yo antibodies, was normal. EEG tracing was also unremarkable.

Because the patient’s clinical picture deteriorated, she was referred for specialized assessment. She continued to complain of dizziness, imbalance and diplopia, and neurological examination revealed marked gait ataxia (she could not perform tandem gait unsupported) with horizontal and vertical gaze-evoked nystagmus, as well as slight dysarthria.

As autoimmune cerebellar ataxia was suspected, further investigations were again carried out, including routine blood chemistry, VDRL, complete blood count, coagulogram, ESR, liver function, thyroid tests, anti-DNA tests, a new MRI and a new CSF examination, as well as measurement of antigliadin, antithyroglobulin and antiperoxidase antibodies, all of which were unremarkable, except for the anti-GAD antibody test (39 IU/mL; normal value <1.0 IU/mL).

As the diagnosis was cerebellar ataxia and epileptic seizures associated with anti-GAD antibodies, the patient underwent IVIg treatment (0.4 g/kg/day) for five days followed by a single monthly i.v. dose (for 12 months) and, subsequently,

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**Fig 1.** Electromyography of patient 1, with continuous motor unit activity in paravertebral muscles.

**Fig 2.** Patient 1 with stiff-person syndrome and contracture and rigidity of the paravertebral muscles (reproduced with patient’s authorization).
azathioprine p.o. (2 mg/kg/day). There was a gradual improvement in the clinical picture and, one year after the start of the specific treatment, neurological examination showed only mild gait ataxia (more apparent in tandem gait) and the presence of mild horizontal nystagmus in both extreme gazes. Lamotrigine (200 mg/day) p.o. and clonazepam (1.0 mg/night) were continued; complete blood count and liver function tests were normal and azathioprine was discontinued.

**DISCUSSION**

Classical SPS is a rare neurological disorder characterized by severe and incapacitating axial and proximal limb rigidity and stiffness due to continuous motor unit activity4-8. This unusual syndrome was first described by Moersch and Woltman in 1956, who defined it as “progressive fluctuating muscular rigidity and spasm” (“Stiff-man” syndrome) after reporting one case and studying 13 others19. Rigidity is often exacerbated by anxiety, sudden movements or external stimuli, causing intermittent pain-ful spasms and frequently leading to skeletal deformity. Exaggerated lumbar lordosis and pain are almost universal findings, and diagnosis is supported by a finding of anti-GAD antibodies in the blood or CSF2. SPS frequently presents with psychiatric symptoms, such as anxiety, phobia and depression, and psychogenic movement disorder is a common misdiagnosis.

In our series of patients with neurological disorders associated with anti-GAD antibodies, we found nine patients with SPS and its variants. Four had a classical clinical presentation of SPS, and the other five presented with SPS variants (two with stiff lower limbs, two with stiff one-limb and one with stiff three-limbs). ENMG examination was diagnostic in all the patients with classical SPS and stiff lower limbs syndrome; however, in the patients with stiff three-limbs or stiff one-limb syndrome, it was not diagnostic, and the diagnosis was confirmed after high serum anti-GAD antibodies had been detected. Autoimmune endocrinopathies, particularly diabetes mellitus and hypothyroidism (Hashimoto’s thyroiditis), were found in some of the patients. Two patients suffered from epileptic seizures and segmental myoclonus affecting the trunk.

Clinically, stiff-person syndrome presents as neurological paraneoplastic syndromes in about 5% of cases1,20,21. In such cases, paraneoplastic stiff-person syndrome is usually associated with amphiphysin or anti-gephyrin antibodies20. While a few cases of paraneoplastic SPS have been described in anti-GAD-antibody-positive patients with thymoma and other solid tumors, a finding of anti-GAD antibodies in patients with other classical paraneoplastic neurological syndromes is rare10,20.

Progressive encephalomyelitis is a rare form of SPS characterized by brainstem, extrapyramidal and spinal cord dysfunction with acute onset of painful rigidity and muscle spasm in association with nystagmus, opsoclonus, ophthalmoparesis, deafness, dysarthria, dysphagia and autonomic dysfunction21,22. In our series, there were no cases of paraneoplastic SPS or progressive encephalomyelitis with rigidity.

Ataxia associated with anti-GAD antibodies is a very rare condition and is considered to be a component of a polyglan-dal autoimmune syndrome in patients with circulating anti-GAD antibodies (10%)21. Most patients with ataxia and anti-GAD antibodies are women and have insulin-dependent diabetes mellitus. In general, there is a slowly progressive pancerebellar syndrome, and cerebellar atrophy is sometimes found in MRI examinations4,22,23. Three of our patients had acquired secondary cerebellar ataxia associated with anti-GAD antibodies, and two, whose CSF analysis revealed mild pleocytosis, had Hashimoto’s thyroiditis. One patient in this group had diabetes mellitus and epileptic crises without myoclonus. There was no difference in anti-GAD levels between this group of cerebellar ataxia patients and the SPS group.

The prevalence of neurological disorders associated with anti-GAD antibodies in epileptic patients is low (less than 10%)25,26. In this series, we found two patients who suffered from epileptic seizures (one from the SPS group and the other from the cerebellar ataxia group). Some authors recommend actively searching for anti-GAD antibodies in patients with atypical refractory epilepsy either on its own or in association with diabetes mellitus or cerebellar ataxia25,26.

The heterogeneity of the neurological presentation in patients with neurological syndromes associated with anti-GAD antibodies is probably related to the widespread distribution of GABAergic neurons in the central nervous system as these cells have a large amount of GAD22.

In conclusion, in this series of neurological disorders associated with anti-GAD antibodies, we found nine patients with stiff-person syndrome (the classical form and its variants) and three with cerebellar ataxia. Four had hypothyroidism, three epilepsy, two diabetes mellitus and two had segmental myoclonus affecting the trunk.

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References