Seronegative Myasthenia Gravis and a Biermer’s Anemia: A Rare Association

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Abstract
We report a rare association of seronegative myasthenia gravis and a Biermer’s anemia (or pernicious anemia). A Senegalese patient of 31 years has been followed for a vitamin B12 deficiency anemia, 12 months before his hospitalization in our department. She has been admitted for an intense and invalidating fatigability in spite of the correction of anemia, associated to a right ptosis. This clinical picture has electively been improved to the prostigmine test. The electromyography had revealed a compatible decrement with a diagnosis of myasthenia. The positivity of the antibodies anti gastric parietal cells and the twice negativity of the antibody against acetylcholine receptor (AChR) and muscle-specific kinase (MuSK) had permitted to deduct a diagnosis of seronegative myasthenia and Biermer’s anemia. The evolution was favorable under substitutive B12 vitamin therapy associated to corticotherapy and azathioprine. We insist on the research and the early treatment of a myasthenia, in a context of Biermer’s anemia, before suggestive clinical signs in spite of the negativity of the anti-Rach antibodies and anti-Musk.

Keywords
Myasthenia Gravis, Pernicious Anemia, Seronegative

1. Introduction
The Biermer’s anemia is a gastritis auto-immune atrophic characterized by the absence of gastric intrinsic factor secretion caused by a malabsorption of the B12 vitamin at the ileal level [1].

Myasthenia gravis is an auto-immune illness affecting the neuromuscular junction, usually associated to the presence of antibodies against acetylcholine receptor (AChR) and muscle-specific kinase (MuSK).
In 20% of the cases, these antibodies are not however detectable, the myasthenia gravis is said negative [2].

The Myasthenia gravis association and Biermer’s anemia is rare. We report the first Senegalese observation of an association of negative Myasthenia and Biermer’s anemia.

2. Observation

This was about a patient of 31 years, accountant by trade who has been followed 12 months before her admission in our department for a megaloblastic anemia labeled Biermer’s anemia under B12 vitamin therapy. She was hospitalized in our institution on February 05th, 2015 for an intense and invalidating fatigability in the effort so far attributed to anemia. To the questioning, she reported a generalized fatigability associated to a tendency to the closing of her right eye especially occurring at the end of the day.

Her general state was maintained, the mucous membranes were colored, and the admission constants were blood pressure 120/70 mmHg, cardiac frequency was 84 beatings per minute, respiratory frequency 18 cycles per minute, temperature 37.2°C.

The admission exam revealed a palmoplantar melanoderma associated to a unilateral ptosis and a difficulty to horizontally maintain her arms for a duration over 30 seconds. The rest of the clinic examination was normal. The prostigmine test has been done right after with a significant improvement of the clinical symptomatology.

At the paraclinical examination the hemogram shows a hemoglobin level of 12 g/dl (normal < 11 g/dl) normochromic, normocytic, the level of platelets of 369,000/mm³ (normal: 150,000 - 450,000) and the leukocytes level of 8140 cells/mm³ (normal: 4000 - 10,000). There was no inflammatory syndrome with a sedimentation rate at the first hour to 25 mm (normal < 15 mm) and a C-reactive protein of 5 mg/l (normal < 6 mg/l). The electrophoresis of the serum proteins was without particularities, the ferritinemny was 50.97 ng/ml (normal: 12 - 150 ng/ml). Elsewhere the dosage of the TSH us showed a rate of 1.47 mUI/L (normal between 0.27 and 4.2), the antibodies against-ENA, against-CCP2 and anti-TPO test was negative. The antibody against-parietal cells were negative and the antibody against intrinsic factors test was positive. The dosage of the auto-antibodies against Musk was 0.03 (normal lower to 0.05) and the dosage of the antibodies against AChR was twice lower than 0.20 UI. The duodenal-gastro-oesophageal fibroscopy revealed an atrophic gastritis of the fund us and the histology revealed an intestinal metaplasia. The electromyography identified a decrement of 5.5% on the pick 5 and 17, 1% on the pick 9 at the orbicular eyelid, a decrement of 30% on the picks 1 and 2 at the abductor thumb (Figure 1).

This aspect was compatible with a myasthenia. The thoracic tomodensitometry was unremarkable. The diagnosis of a seronegative myasthenia gravis has been put association of a Biermer anemia.
Figure 1. Results compatible with myasthenia gravis. (a) Large deltoid; (b) Orbicularis of the eyelid; (c) Short abductor of the thumb.

Therapeutically, the B12 vitamin therapy treatment has been followed at the rate of 1000 UI/month and the administered treatment was the azathioprine, 100 mg/day associated with a corticotherapy of 10 mg/day that permitted a significant improvement of the symptoms. The treatment of anticholinesterase only had not entailed a favorable evolution of the symptoms. The evolution is until this day stable with a normal performance of daily activities and a receding of more than 2 years of follow-up.

3. Discussion

We reported an association of seronegative myasthenia gravis and Biermer’s anemia.

The myasthenia is defined as an auto-immune illness concerning the neuromuscular junction and its impact varies between 1.5 and 10 cases for 1 million inhabitants [3] [4]. In 20% of the cases it is seronegative, characterized by the negativity of the anti-RACH and anti-MUSK Ac of the techniques of routine detection [5]. We did not find any association with seronegative myasthenia and Biermer’s disease in the literature. In our observation, the diagnosis of negative myasthenia has been put on the basis of the myasthenia diagnosis criteria (existence of a muscular weakness fluctuating spontaneously or under the effect of the effort, associated with a clear answer to the test of the anticholinesterase of fast action, either to electrophysiological signs of post-synaptic block [6] [7]. associated to the negativity of the anti-MUSK antibodies and the anti-receptors antibodies of the acetylcholine twice.

The seronegative myasthenia gravis would have a superimposable clinical and epidemiological presentation with immunopositive forms. However in some sets, it has been reported a more advanced age of onset and a less clear feminine predominance. The resemblance on the clinic plan is globally found in the literature even though a subgroup of negative myasthenia presents some predominantes
oculo-bulbar demonstrations [8].

The patient was relatively young and presented a generalized shape. The seriousness of the myasthenia is globally similar be the myasthenia is positive or negatives [9].

The myasthenia can sometimes associate to other auto-immune illnesses, the thyroids dysfunction remains the first reasons of association with 5% to 10% of the cases [10], and the dosage of the TSHUs was at a normal rate in our patient. The myasthenia can associate to numerous diseases, the association to Biermer’s anemia that can be defined as an auto-immune gastritis at the origin of a B12 vitamins deficiency responsible for a macrocytic anemia and less frequently of neurological demonstrations [11]. The association of Biermer’s anemia and myasthenia can associate to numerous diseases, the association to Biermer’s anemia that can be defined as an auto-immune gastritis at the origin of a B12 vitamins deficiency responsible for a macrocytic anemia and less frequently of neurological demonstrations [11]. The association of Biermer’s anemia and myasthenia is very little described and we did not find an association of seronegative myasthenia gravis and Biermer’s anemia in the literature. Some sets in the literature valued the association and described it without specifying the immunological characters of Biermer’s anemia [12]. However isolated cases of associations exist. In Goulon et al.’s set [2], as well as Fraisse et al. [13], it has been respectively noted proportions of 3/84 patients having developed some antibodies against-intrinsic factors and no patient developed an authentic Biermer’s anemia and 2/45. The Biermer’s precedes most often myasthenia [13], which was the case of our patient. In a context Biermer anemia, as in our observation the intervening of a fatigability to the effort associated to a ptosis must evoke an association to Myasthenia.

The negativity of the antibodies doesn’t exclude the diagnosis. This negativity would be due to an imperfect sensitivity of the usual techniques of antibodies against-AChR and MuSK tests and/or the controlled presence of antibodies aimed at unidentified antigenic targets [5].

Therapeutically, we noted a good regression of the symptomatology of the Biermer anemia under B12 vitamin therapy alone even though some authors reported a use of the corticotherapy for an improvement of hematological manifestations [14]. As for the myasthenia it has not only been improved under azathioprine associated to the corticotherapy after an absence of improvement under corticotherapy alone, which was in accordance with the therapeutic algorithm of myasthenia.

4. Conclusion

The seronegative myasthenia-Biermer’s anemia association has not been described in literature. The seronegative character of myasthenia doesn’t exclude its association to other auto-immune diseases.

References

B. Djiba et al.


