POLYMYOSITIS

Clinical investigation in two sisters

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ABSTRACT - We present an investigation of a case of polymyositis affecting two sisters of one same parenthood. Their cases have been documented for almost two decades, being investigated by means of a diagnostic protocol which combined clinical findings as well as laboratorial, histopathological and image tests. In both cases, clinical manifestations started in childhood, without signs of involvement of the central and peripheral nervous system. Both patients proved to respond to a therapeutics based on corticosteroids. The degree of relatedness between their parents corroborate the notion that genetic factors may contribute to the development of the disease.

KEY WORDS: polymyositis, inflammatory myopathy.

Idiopathic myopathies are a heterogeneous group of acquired muscle diseases which affect 1 in 100,000 individuals1. Polymyositis (PM) comprises an inflammatory myopathy that results from an abnormal immune response to skeletal muscle fibers and associated tissues. Activation of auto-reactive T-lymphocytic lymphocytes and molecular mimicry are probable mechanisms underlining this pathology2. Moreover, the majority of patients diagnosed with PM display high levels of antibodies to self-antigens, such as cell membrane components and nuclear proteins3-4. Current diagnosis of PM is assessed by means of compatible clinical findings (proximal muscle weakness) and measurements of serum muscle enzymes, electromyography and muscle biopsy5. Even though causes for PM remain undetermined, an increasing awareness that genetic factors are implicated has evolved from different studies6-7. In spite of that, the occurrence of more than one case within the same family still consists in a rare event6,8,8.

This study has the purpose to describe anamnestic and testing procedures which resulted in the diagnosis of two patients with PM within the same parenthood, and to report treatment and evolution of both patients documented over almost two decades.

CASES

Patient 1. G.R.V.C., 26 years old female Caucasian, daughter of parents related to the third degree, was born presenting Apgar 10. The very first symptoms of mobility disorder arose at the age of 5 years old as complaints of strong muscular pains in both lower limbs after extenuating physical exercises. At the age of nine, the patient developed the habit of walking on her toe tips, what prompted medical attention. Preliminary analysis revealed general hipotony, predominantly on the pelvic and scapular muscles. At this time, the following laboratorial tests...
were performed: complete hemogram (normal); summary of urine (normal); oxalacetic transaminase 145 UI/l max RV 30 UI/l; pyruvate transaminase 186 UI/l (max RV 37 UI/l); lactate dehydrogenase 419 UI/l (max RV 240 UI/l); creatine kinase (CK) 2,600 UI/l (max RV 70 UI/l); aldolase 26.3 UI/l (max RV 7.6 UI/l). X-ray analysis of the lumbar and the sacrum-coccyx regions of the vertebral column yielded normal results.

Studies on nerve-motor conduction revealed no alteration on either latency or amplitude of the action potentials of the nerves studied (median, ulnar and fibular), being the speed of conduction preserved as well. Studies on sensitive conduction of the same nerves have also failed to demonstrate alterations. Nonetheless, quantitative electromyography yielded motor units of greatly reduced length and amplitude, being polyphasic in their majority, mainly on the biceps, deltoid and quadriceps muscles in spite of no signs of denervation. Tomographic analysis failed to demonstrate any intracranial expansive processes either above or below the tentorium as well as absence of any pathological calcifications, attesting a cranium-encephalic status within the parameters of normality. Additional tomographic analysis revealed a similarly healthy vertebral condition, characterized by canal with dimensions and configuration compatible with normality, lacking pathological compression upon the dural sac. All findings above are consistent with a myopathic condition of idiopathic etiology.

Biopsies of biceps and quadriceps were performed under local anesthesia, being samples fixed before histological processing. Both tissues revealed a degenerative process characterized by disorganization of striated muscle fibers with various degrees of atrophy, displaying multiple endomysial infiltrates by polymorphonuclear cells. In some areas, intensive fagocytosis of muscle fibers by inflammatory mononuclear cells could be observed, with signs of bundle regeneration and endomysial fibrosis. Such findings are compatible with a myopathic condition of acute and chronic polymyositis.

Patient 2. M.R.V.C., 23 years old, female, Caucasian, sister of G.R.V.C., was born presenting Apgar 8. The first symptom of mobility disorder arose at the age of nine, by walking with her left foot in valgus. Having in mind her sister’s condition, myopathy was taken under consideration. At that time, the following laboratorial tests were performed: complete hemogram with eosinophilia at 9% (max RV 3%); oxalacetic transaminase 24.5 UI/l; pyruvate transaminase 29 UI/l; lactate dehydrogenase 273 UI/l; creatine kinase 1,580 UI/l; aldolase 19.7 UI/l. Max RV were as cited previously. X-ray analysis of hands and feet revealed osseous age within the normality. Similarly to her sister’s case, studies of nerve-motor conduction have not shown major alterations of either latency or amplitude of the action potentials of the nerves investigated. Accordingly, electromyographic examination confirmed motor units of reduced length and amplitude, largely polyphasic and restricted to the proximal muscles. Biopsies of these muscle revealed an interstitial microscopic aspect similar to the previous case, with several degenerative inflammatory areas and bundles being regenerated. On what concerned its intensity, the degenerative process that was observed in these samples exceeded the severity of the preceding case, in spite of the lower levels of muscular enzymes. The accentuated disappearance of the striated aspect and the formation of large vacuoles in the muscle cells could translate this increased myopathy. Both biochemical evidence and histopathological findings attest an intense myopathic process, greater in severity than the preceding case, and consistent with acute polymyositis.

In both cases, initial therapeutics consisted on oral administration of 60 mg of the corticosteroid prednisone per day, which resulted in marked decrease of the serum enzyme levels within the first month (Fig 1). Dosage was gradually reduced according to the therapeutic response. Nowadays, both patients present stable clinic conditions which allow regular daily activities: walking without assistance, absence of dysphasia and minor difficulty in shifting from the seated to the ortostatic position. Both patients attend to physiotherapy sessions twice a week in order to strengthen the pelvic and scapular musculature. M.R.V.C employs a maintenance dose of 5 mg of corticosteroids in alternate days. G.R.V.C. sustains her medication suspended since October/2000, following a 16 years period of stable condition under administration of a similar maintenance scheme.

DISCUSSION

The first reports of polymyositis were based on findings of anamnesis. These studies used to generate inaccurate diagnosis since inflammatory myopathies share clinical symptoms such as mialgia, proximal muscle weakness and electromyographic alterations with several other idiopathic forms of myopathy. The development of diagnostic protocols that couple laboratorial, histopathological and image procedures has greatly facilitated the precise identification of myopathies with inflammatory etiology, what in turn allows hiring an effective therapeutics. In this study, an investigation based on clinical, biochemical and images findings was successfully hired to diagnose polymyositis in two non-twin sisters from the same parenthood.

In both cases, the first clinical signs were observed within childhood. In accordance to a typical myopathic condition, levels of serum muscular enzymes (CK and aldolase) were significantly elevated in preliminary tests. Electromyographic abnormalities confirmed a myopathic status which was not correlated to any detectable cranium-encephalic or neurological disturbances, whose normality was attested by tomographic as well as nerve conduction analysis.
In addition, histopathological analysis of proximal muscle biopsies of each patient revealed an intense degenerative process characterized by marked inflammatory infiltrates, showing a clinical condition compatible with PM. Nonetheless, symptoms usually associated to inflammatory myopathies such as fever, dysphasia and arthralgia\textsuperscript{12,13} have not been observed. Exclusion of dermatomyositis (DM) was possible based on the absence of cutaneous involvement in either patients. Both cases proved to respond to a therapeutic of corticosteroids, noticed the great reduction of the serum muscle enzymes within the first month of treatment (Fig 1). In spite of the long-term administration of steroids, periodic osseous densitometry revealed no major pathological alterations of their bone constitution.

Despite the fact that the etiology of PM remains undetermined, different factors have been correlated...
with the origin of inflammatory myopathies. Some studies implicate viral and bacterial infections as possible triggering factor of the pathogenic process\textsuperscript{9,15,16}. Others enroll immune disturbances and genetic profiles as causative determinants for the onset of PM and DM. In the latter, disruption of immunological tolerance is considered a factor that probably accounts to the development of muscular lesions, since such lesions may consist in a pathological outcome of the autoantibodies found in a significant number of patients with PM\textsuperscript{6,17}. In addition, the augmented frequency of the HLA-B8 and DR3 haplotype among patients diagnosed with this pathology suggests that a genetic profile may predispose to the onset\textsuperscript{18}. In the present study, both patients are sisters from one same parenthood, where the parents are cousins in first degree. It is noteworthy that related individual share sets of haplotype and that, for this reason, consanguine unions tend to predispose their descendants to the onset of recessive and multifactorial phenotypes. Thus, even though a definitive relationship between immunogenetic factors and polymyositis has not been established, we consider that the cases reported in this study corroborate such an association.

The comprehension that the immune system works as an agent that ensures the equilibrium within the organism implies that its disturbance allows or generates unbalance. In autoimmune diseases, it is desirable to apply medicinal treatments that contribute to the recovery from the unbalanced state. In the cases above, treatment with corticosteroid proved to be appropriate for such purpose, considering that patient G.R.V.C. remains with a stable condition despite of having her medication suspended for one year. Nonetheless, it is important to emphasize the need for an early and accurate diagnosis of inflammatory myopathies, so that an appropriate therapeutics may take place around the dawn of the pathological process to allow satisfying outcomes, similar to those described in this study.

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**REFERENCES**