The effect of calpain 3 deficiency on the pattern of muscle degeneration in the earliest stages of LGMD2A

M Vainzof, F de Paula, A M Tsanaclis, M Zatz

Limb girdle muscular dystrophy type 2A (LGMD2A) is caused by mutations in the calpain 3 gene. In a large family affected by LGMD2A with four severely affected members, three additional asymptomatic relatives had very high serum creatine kinase concentrations. All were homozygous for the R110X mutation and showed a total absence of calpain 3 in the muscle. Histological analysis in these three rare preclinical cases showed a consistent but unusual pattern, with isolated fascicles of degenerating fibres in an almost normal muscle. This pattern was also seen in one patient with early stage LGMD2A who had a P82L missense mutation and a partial deficiency of calpain 3 in the muscle, but was not seen in early stage patients affected by other forms of LGMD. These findings suggest that a peculiar pattern of focal degeneration occurs in calpainopathy, independently of the type of mutation or the amount of calpain 3 in the muscle.

CASE REPORT

Among more than 250 patients with LGMD submitted to muscle histological and protein analysis we identified one highly inbred family affected by LGMD2A in which there were four severely affected adult members and three still asymptomatic young individuals with high serum creatine kinase concentrations (increased 10–12-fold; family 22 in Passos-Bueno and colleagues'). All affected individuals and all those with very high CK concentrations had a homozygous R110X mutation in the calpain 3 gene. Muscle protein analysis showed a total absence of calpain 3 in muscle (fig 1D).

A muscle biopsy from one affected relative showed typical dystrophic features (fig 1A). In the three preclinical patients, muscle histology revealed an almost normal histological pattern. However, an isolated fascicle of degenerating fibres was seen. Acid phosphatase staining was positive and there was patchy labelling for dystrophin and the sarcoglycan proteins, and an increased staining for utrophin in the plasma membrane from these degenerated fibres (fig 1B).

This pattern of focal degeneration was subsequently seen in one additional young patient with sporadic LGMD2A in the initial stages of the disease (fig 1C). She was identified through the deficiency of calpain 3 in the muscle, and DNA analysis detected a homozygous missense P82L mutation in the calpain 3 gene.

This histological pattern was not seen in other patients affected by muscular dystrophy who were studied by our group, or in preclinical or early stage cases of Xp21 muscular dystrophy, dystrophinopathies, sarcoglycanopathies, or telethoninopathy, and suggests that a different mechanism for the onset of degeneration may occur in patients with primary calpain 3 deficiency.

DISCUSSION

Calpain 3 is a proteolytic enzyme of the calpain superfamily whose specific role is not known. The protein is composed of 821 amino acids, organised into four domains, which include...

Abbreviations: AD, autosomal dominant; AR, autosomal recessive; LGMD, limb girdle muscular dystrophy; NS, N-terminus domain I; SG, sarcoglycan...
a cysteine proteolytic domain (II) and a calcium binding domain (IV). There are also three short specific inserted sequences located at the N-terminus domain I (NS), in the protease domain II (IS1), and between domains III and IV (IS2). In the IS2 region there is a titin binding site and a nuclear localisation signal.10

“Our results showed that the pattern of focal degeneration was not related to the amount of calpain 3 in the muscle because in the first family the deficiency was total, whereas in the second one it was only partial.”

Various intracellular enzymes and transcription factors, in addition to cytoskeletal proteins, are processed by calpain, resulting in modification of their structures and activities. This suggests that calpain may play an important role in intracellular signal transduction.11 Complex proteolysis, promotion, and suppression of the network involving calpain 3 protein has been suggested in the response to muscle wasting. Thus, substrate processing by calpain 3 is essential for the proper function or maintenance of skeletal muscle.12 In this regard, preservation of the proteolytic domain of calpain 3 would be of utmost importance.

Our results showed that the pattern of focal degeneration was not related to the amount of calpain 3 in the muscle because in the first family the deficiency was total, whereas in the second one it was only partial (fig 1D). In addition, it is not related to a specific mutation, because it was seen as the consequence of two different mutations. The R110X mutation is located in the IIa protease domain of calpain 3, whereas the P82L mutation is located between the N-terminal first NS specific insertion sequence and the IIa protease domain. It is important to note that both mutations are at the N-terminal region of the protein, and possibly affect its proteolytic domain function.

It has also been suggested that calpain 3 deficiency may cause myonuclear apoptosis and a profound perturbation of the 1kb8–nuclear factor-κB pathway in the earliest stages of the disease. Apoptotic myonuclei would be distributed in clusters.13 This would be consistent with our finding of clustered degenerated fibres grouped in possible clones.

Figure 1 Histological, histochemical, and immunohistochemical analysis of muscle biopsies from: (A) one adult patient affected by LGMD2A who was from the family with the R110X mutation, and who showed a typical dystrophic pattern; (B) one of the asymptomatic members from this family, who showed focal degeneration in the haematoxylin and eosin (HE) and the acid phosphatase reactions. The same bundle of fibres shows a patchy pattern of labelling for dystrophin and sarcoglycans, and strong sarcolemmal labelling for utrophin; (C) the patient with early stage LGMD2A who had the P82L mutation, showing focal degeneration; (D) multiplex western blot analysis for dystrophin (DYS) and calpain 3 (CALP3; using the antibody used by Anderson and colleagues1), showing the presence of a 427 kDa dystrophin band and the 94 kDa band of calpain 3 in normal controls (N), the absence of the 94 kDa calpain 3 band in patient B (preclinical), and partial deficiency in patient C. MYOS, myosin content in the Ponceau prestained blot for the evaluation of the amount of loaded muscle proteins.

Take home messages

• Histological analysis of muscle in preclinical and early stage cases of limb girdle muscular dystrophy (LGMD) 2A showed a consistent but unusual pattern, with isolated fascicles of degenerating fibres in an almost normal muscle.
• This pattern was not seen in early stage patients affected by other forms of LGMD.
• A peculiar pattern of focal degeneration appears to occur in calpainopathy, independently of the type of mutation or the amount of calpain 3 in the muscle.

In conclusion, histological alterations in rare preclinical and early stage LGMD2A muscle biopsies suggest that in calpainopathy a peculiar pattern of focal degeneration occurs in the earliest stages of the disease.

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Authors’ affiliations

M Vainzof, F de Paula, M Zatz, Human Genome Research Centre, Department of Biology, IBUSP, University of São Paulo, São Paulo, Sao Paulo - CEP. 05508–900, SP Brazil
A M Tsanaclis, Department of Pathology, FMUSP, University of São Paulo

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REFERENCES


Teaser

Pathology’s top ten one liners—and what they really mean

1 Enucleated specimen of right eye, inadequate for opinion: excise the left eye, too.

2 Small round cell tumour, advised immunohistochemistry for a definite diagnosis: I don’t know what the hell it is.

3 Compatible with lichen planus: doesn’t look like it. But if you insist, I don’t resist.

4 Florid reactive hyperplasia, lymph node; advised close clinical follow up: boss, wait till it turns into a full blown lymphoma, then I’ll type it.

5 Borderline serous cystadenoma, ovary, with focal microinvasion: phew, this’ll save my skin, if the patient throws a met 10 years later!

6 Early ill formed epithelioid granulomas with occasional acid fast bacilli: I have an excellent imagination!

7 Special stains for fungi, bacteria and parasites are not contributory: I didn’t look hard enough.

8 Metastatic poorly differentiated neoplasm, cerebellum, with possibilities of carcinoma, sarcoma, melanoma, lymphoma . . .: looking for the primary is your job; anyway, how does it matter now?

9 Appendix showing lymphoid hyperplasia: you knocked off a perfectly normal one.

10 Poorly preserved biopsies from multiple sites, unsuitable for definite opinion: only a necropsy can solve the issue!