Clinical imaging in genetic neuromuscular disorders

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Role of radiologic imaging in genetic and acquired neuromuscular disorders

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Abstract

Great technological and clinical progress have been made in the last two decades in identifying genetic defects of several neuromuscular diseases, as Spinal Muscular Atrophy, genetic muscular dystrophies and other genetic myopathies. The diagnosis is usually challenging, due to great variability in genetic abnormalities and clinical phenotypes and the poor specificity of complementary analyses, i.e., serum creatine kinase (CK) and electrophysiology. Muscle biopsy represents the gold standard for the diagnosis of genetic neuromuscular diseases, but clinical imaging of muscle tissue is an important diagnostic tool to identify and quantifies muscle damage. Radiologic imaging is, indeed, increasingly used as a diagnostic tool to describe patterns and the extent of muscle involvement, thanks to modern techniques that enable to define the definition of degrees of muscle atrophy and changes in connective tissue. They usually grade the severity of the disease process with greater accuracy than clinical scores. Clinical imaging is more than complementary to perform muscle biopsy, especially as ultrasound scans are often mandatory to identify the muscle to be biopsied. We will herein provide detailed examples of the radiologic methods that can be used in genetic and acquired neuromuscular disorders, stressing pros and cons.

Key Words: Muscle Imaging, MRI, CT, genetic muscle disorders, myopathies, dystrophies

Great technological and clinical progresses have been made in the last two decades in identifying genetic defects of several neuromuscular diseases, as Spinal Muscular Atrophy, genetic muscle dystrophies and other genetic myopathies. However, the diagnosis is usually challenging, due to great variability in genetic abnormalities and clinical phenotypes, the complexity of the molecular genetic approaches and the poor specificity of complementary analyses, i.e. blood CK, electrophysiology, and others. Although muscle biopsy represents the gold standard for the diagnosis of genetic neuromuscular diseases, clinical imaging of muscle tissue is an important diagnostic tool for the identification and quantification of muscle changes. Radiologic imaging is, indeed, increasingly used as a diagnostic tool to describe patterns and the extent of muscle involvement, thanks to modern techniques that enable the definition of degrees of muscle atrophy and changes in connective tissue. They usually grade the severity of the disease process with greater accuracy than clinical scores. Further, clinical imaging is more than complementary to perform muscle biopsy, especially as ultrasound scans are often mandatory to identify the muscle to be biopsied. We will herein provide detailed examples of the several radiologic methods that can be used, stressing for each one their respective pros and cons.

Computer Tomography

Computer Tomography (CT) has been widely used to assess the presence and extent of changes in skeletal muscle of patients with congenital neuromuscular diseases. In particular, the use of CT has made it possible to detect the selective involvement of muscle groups; moreover, CT has been useful in establishing disease progression and identifying asymptomatic patients. The myopathic patients usually undergo gradual loss of muscle strength, which doesn't affect every muscle group to the same degree. Selective involvement of specific muscle groups and savings of others has been confirmed by different studies using CT. Fatty replacement of skeletal muscle tissue is seen as an area with a clear decrease in density, therefore affected muscles are less identifiable as a consequence of the loss of contrast with the subcutaneous adipose tissue. A CT scan is usually more accessible than magnetic resonance (RM), and it allows for the quick and correct evaluation of muscle changes, especially fatty degeneration, by assessing...
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Muscle density and morphology. Examination is not operator-dependent and allows for the evaluation of the deepest muscles. However, CT scanning needs high doses of ionizing radiation, and has therefore it has been almost entirely replaced by ultrasound (US) and magnetic resonance imaging (MRI). Another drawback are the poor contrasts of the soft tissues, which makes identification of inflammatory changes difficult or impossible; as rule of thumb, muscle edema appears before the muscle degeneration. Furthermore, though TC provides objective information that is easy to be duplicated by different observers and different apparatuses, some muscle artifacts may appear in muscle adjacent to cortical bone tissue.5

Ultrasound
Ultrasound is a well-known validated imaging technique for the evaluation of suspected muscle diseases. It is especially useful with children, due to the lack of ionizing radiations.11,12 It is typically used to identify muscle thickness and to evaluate echogenicity changes, and can identify both atrophic changes and fatty degeneration, which is a typical feature of neuromuscular dystrophic diseases, like Duchenne Muscle Dystrophy. Its applications range from initial assessment, ultrasound-guided biopsy and follow-up of the patients (Figs. 1 and 2). Guiding biopsy with US, allows for the targeting the more affected muscles.12-15 On top of being low-cost and largely available, ultrasound scans have good resolutions, up to 1 mm, which allows for the identification of tertiary muscular bundles.16 US also enables dynamic analysis of muscle contraction by short videos to detect muscle contraction and/or pathologic fasciculations.16 Unfortunately, it also has severe disadvantages: its application is usually limited to superficial muscle groups, since it is not sufficiently effective in deep

![Fig 1. Normal appearance of the tight on ultrasound, computed tomography and magnetic resonance scans at the level of the middle third: in the ultrasound images (axial and longitudinal scans), the normal muscles are characterized by heterogeneous ecostructure and are surrounded by slightly hyperechoic fat; in the CT image, the normal muscle is slightly iperdense (10 to 40 HU), while fat tissue is ipodense (-100 to -50 HU); in the MR image (T1-weighted), the normal muscular appearance is characterized by a homogeneous isointense ("grayish") signal intensity. The subcutaneous soft tissues, the intermuscular septa, the intramuscular aponeurosis, and the cancellous bone as well, have a brighter hyperintense signal intensity due to the presence of fat; the femoral cortex is hypointense, the typical signal intensity of cortical bone.](image1)

![Fig 2. Fatty replacement of the anterolateral compartment muscles of the leg following common fibular nerve injury: ultrasound and magnetic resonance axial images at the level of the middle third of the tibialis anterior belly. The right anterolateral compartment muscles are denervated, so, compared to the contralateral ones, these muscles are atrophic with increase of intramuscular fat: in the ultrasound images, there is a marked thinning of the tibialis anterior muscle with increased echogenicity and loss of physiological heterogeneity of the ecostructure; in the T1-weighted MR image, these muscles are brighter.](image2)
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Fig.3  Fatty degeneration: axial T1-weighted images of the thigh showing, on the left, the normal muscular appearance on MR, characterized by a homogeneous grayish ("isointense") signal intensity. On the right, there is a severe and diffuse muscular fatty degeneration: these muscles are hyperintense compared to the previous ones because of the increase of intramuscular fat.

Fig.4  Intramuscular edema: axial STIR images of the thigh showing, on the left, normal muscle signal intensity. In this sequence, the signal coming from the fat is suppressed, so the fat-containing structures are hypointense; on the right, the muscles are hyperintense, particularly the vastus medialis muscle due to the increase of water typical of inflammatory changes.

Magnetic Resonance

Due to drawbacks of CT and US, MR is increasingly used in the evaluation of patients with genetic neuromuscular diseases. The technique provides excellent soft tissue characterization, so it perfectly evaluates shape, volume and morphological features of normal skeletal muscle. Specifically, this technique easily allows for determination of fatty degeneration in late-stage muscle dystrophy, aside from recognition of decreases in muscle volume. Thanks to this optimal soft tissue characterization, MRI easily distinguishes the different muscles in the same muscle group, which is important in the identification of a specific pattern of involvement. Using qualitative grading, MRI can assess the degree of fat replacement of muscles and thus follow disease progression. Fatty degeneration is usually evaluated in the T1-weighted images, where it appears as area of signal hyperintensity (Fig 3).

T2-weighted images with fat signal suppression and STIR sequences identify muscle edema, which appear as a diffuse signal hyperintensity of the involved muscles (Fig. 4). This is useful in detecting early signs of neuromuscular diseases, because inflammatory changes are often the first manifestation of the muscle disease, much earlier than fat and scar replacement of the muscle tissue. Recent studies suggest that the use of intravenous administration of contrast medium would provide better details regarding the detection and extent of muscle involvement, although this leads to a substantial increase of scan time. MRI has higher sensitivity in detecting muscle dystrophic disorders and a good inter-observer and intra-observer agreements.

Therefore, MRI is preferred to CT and US in the evaluation of genetic dystrophic neuromuscular disorders, and also because MRI doesn't expose the patients to ionizing radiation and, thus, can be safely used with young patients. In these cases, RM protocol include Turbo-spin-echo T1-weighted, in T2-weighted and in T2 with fat suppression and, eventually STIR (short-tau inversion recovery) sequences. MR images are usually acquired in the axial plane with slice thickness of 4 mm. When recommended, it is also possible to obtain images according to other anatomical planes (coronal, sagittal); multiplanar scans, of course, provide more anatomical details. The typical MR protocol applied in this type of study encompasses T1-weighted scans of the pelvic girdle and lower limbs; there is still little experience in describing neuromuscular diseases by analyzing scans of to chest and upper limbs or by whole-body MRI. Whole-body MRI can be useful in the assessment of Congenital and Metabolic Myopathies, which typically show muscular involvement of regions other than limbs, without using great doses of ionizing radiation. Myopathic patients usually undergo a partial and gradual loss of muscle strength, but this loss doesn't affect all muscle groups with the same degree of involvement. Selective involvement of certain muscles and and saving of others was confirmed by studies using imaging techniques, in particular CT and MR. For this reason, it is important to evaluate not only the different muscular compartments, but every single muscle in them, in order to detect specific patterns that can narrow the differential diagnosis and help the search for specific genetic defects. For example, some authors have demonstrated specific patterns of involvement in the Central Cores Myopathies, dysferlinopathy (LGMD-2B), hyaline body myopathy (HBM), tubular aggregate myopathy (TAM) and myotonic dystrophy (MD) (Fig. 5) while others...
claim that, by MRI, specific patterns of involvement are powerful predictors in congenital myopathies due to ryanodine receptor type 1 gene mutations. Due to the lack of ionizing radiation, MRI is preferred to CT in young patients and in the assessment of asymptomatic patients with relatives affected by genetic disorders. There are experimental techniques, both applied to CT and MR, that can quantify the degree of muscle degeneration, evaluate the relationship between muscle fibers and fibrous tissue or measure the amount of intramuscular fat tissue. These techniques can be useful in the evaluation of therapies, but use experimental software and/or MRI sequences that are not easily available and difficult to apply to the clinical practice.

A purpose-developed false-color method of muscle imaging was designed and implemented to analyze macroscopic and microscopic structural changes of human skeletal muscle based on processing techniques of medical CT scans, to follow-up in patients suffering with complete permanent denervation of leg muscles (complete Conus and Cauda Equina syndrome) the progression of muscle atrophy to degeneration and the extent of muscle recovery after experimental therapy. Changes in tissue composition within the muscle were visualized by associating different colors to Hounsfield unit values of normal or atrophic muscle, fat and connective loose and fibrous tissue. The results of this analysis were presented as the percentage of different tissues (muscle, loose and fibrous connective tissue, and fat) in cross-sections and in the total volume of muscle on the three-dimensional reconstructions. Authors stress that the advantages of these approaches outweigh the low risk related to irradiation, in particular during follow-up of supervised trials, adding quantitative evidence to clinical assessments.

As mentioned before, in clinical practice it is possible to use rating scales in order to semi-quantitatively assess the degree of muscle degeneration. This type of scaling evaluates the extension of fatty degeneration. Its use is relatively quick and has a good inter-observer and intra-observer agreements.
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One of the most used is a 5-point scale where the signal intensity, evaluated in MRI T1-weighted sequence, is scored using a modification of the scale reported by Lamminen (Table 1 and Figure 7)⁶,³⁴ In the early stages of disease or in mild cases, an MR scan may be normal or slightly pathological, with a selective involvement of a few muscles. In more advanced myopathies, the pattern of involvement is less evident because more muscles are involved; but, spared muscles or less affected ones, which identify the so-called "negative pattern", can be useful diagnostic clues.¹¹ For diagnostic purposes, a quantitative description of the muscle involvement is not necessary, as the degree of fatty degeneration correlates with the severity and duration of the disease. The articles describing the pattern of involvement with the MRI, in different types of myopathy and dystrophy, have a low number of patients so far, and little is known about the spectrum of disease, which is only detectable by studies with higher number of patients. Furthermore, when MRI indications of the disease can be identified, the degree of disease progression assessed by MRI, whether sex, ethnicity, exercising can influence the progression and whether the pattern varies with the type of mutation in a specific gene are not fully understood.

Knowing these points would help with diagnoses and would give useful information about the pathophysiology and therapy of neuromuscular diseases.²² MRI is increasingly used in the evaluation of patients with genetic neuromuscular diseases. Thanks to its high sensitivity in the detection of muscular abnormalities and a good inter-observer and intra-observer agreements, MRI is preferred to CT and US.⁶ In these cases, semi-quantitative rating scales allow for the objective assessment of muscle degeneration and therefore MRI provides an alternative or supportive analysis in follow up aside from clinical and laboratory tests. In patients suspected to suffer or suffering genetic neuromuscular disorders, MRI is, thus, useful for: 1. identifying asymptomatic patients, but with RM signs of muscle involvement (e.g. family members of asymptomatic patients or patients susceptible to malignant hyperthermia); 2. Staging the extent of muscle damage; 3. guiding muscle biopsy, which allows for the selection of the more diseased muscle or muscle group; 4. selecting therapeutic treatments and evaluating new therapies; 5. monitoring disease progression and following-up with patients.

In conclusion, radiologic imaging is increasingly playing a relevant role in neuromuscular disorders: in

### Table 1. Modified Lamminen rating scale.

| SCORE | DESCRIPTION |
|-------|-------------|
| 0     | Normal      |
| 1     | Mild with only traces of increased signal intensity |
| 2     | Moderate with increased signal in less than 50% of affected muscle |
| 3     | Severe with increased signal intensity in more than 50% of affected muscle |
| 4     | Entire muscle replaced by abnormal signal |

Fig 7. Modified Lamminen grading scale. Axial T1-weighted images of the thigh showing different grade of fatty infiltration according to the grading scale.
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particular, those of known or unknown genetic etiology. In spite of the dominant role of Magnetic Resonance Imaging, other approaches maintain important roles: in particular, Ultrasound with reagards to in guiding muscle biopsy. Other promising applications are described in details in other chapters of this Ejtm Special: News of Clinical Muscle Imaging.

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