Muscle MRI has become a very useful tool in the diagnosis and follow-up of patients with muscle dystrophies. Muscle MRI provides us about many aspects of the structure and function of skeletal muscles, such as the presence of oedema or fatty infiltration. In the last years many reports have described the particular muscles that are involved in these muscle disease. This knowledge can facilitate the diagnosis in many cases. In the present paper we review the main changes observed in muscle MRI of patients with muscle dystrophies.

Key words: Limb-girdle muscular dystrophies; muscle MRI

Introduction

Muscle MRI has become a useful tool for diagnosis and follow-up of patients with limb-girdle muscle dystrophies (LGMD). Muscle MRI provides information on different aspects of muscle structure and function of muscles of the body. The sequences available allow to identify fatty infiltration (T1 or 3-point Dixon sequences) or oedema (STIR or T2) in the muscles (1). Therefore, muscle MRI is useful to select a muscle for the biopsy in patients without clear muscle weakness or in patients with a severe degree of muscle atrophy, in which muscle tissue can be completely substituted by fat. Moreover, because MRI does not use radiation, it is a good technique to follow-up the progression of the disease in patients (2). There is a growing number of evidences demonstrating that the degree of muscle fatty infiltration observed in muscle MRI correlates with the muscle strength and functional status of patients with LGMD (2-4). For this reason, muscle MRI is being progressively included as a primary endpoint in clinical trials of patients with muscle dystrophy (5, 6).

It is well known that every single muscle dystrophy induces fatty infiltration of a particular group of muscles. This fact allows investigators to design diagnosis algorithms based on the MRI findings (7). However the number of reports describing the changes observed in every muscle disease has grown in the last years and a review of the main findings observed in every disease is needed. Our aim is to describe our protocol to perform muscle MRI in patients with muscle dystrophy and to summarize the main findings of MRI studies in LGMD.

Methods

Muscle MRI protocol

Good quality Muscle MRI studies can be conducted in a 1.5 Tesla MRI scanner. In our center, we are equipped with a Philips Achieva XR Magnetic Resonance System (Phillips, Eindhoven, Netherlands). Our MRI system has a moveable tabletop that allows performing whole body studies in a short period of time (less than 30 minutes) without relocating the patient. Our standard protocol includes T1 and STIR sequences. T1 weighted spin-echo sequence obtains axial and coronal images using the following parameters: repetition time: 300 ms, echo-time: 10 ms and thickness 10 mm. Short-time inversion recovery (STIR) sequences obtains coronal images using the following parameters: repetition time 2,500-3,500 ms, echo time 60 ms, inversion time 150 ms in 10 mm slices.

We quantify the degree of muscle fatty infiltration using the Mercuri scale modified by Fischer et al in 2008 that has been used to analyse both muscle MRI and CT scans (8) (Fig. 1):

- **Normal muscle appearance**: 0 points.
- **Mild involvement**: traces of increased signal intensity on the T1-weighted MR sequences: 1 point.
- **Moderate involvement**: increased T1-weighted signal

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intensity with beginning confluence in less than 50% of the muscle: 2 points.
- **Severe involvement**: increased T1-weighted signal intensity with beginning confluence in more than 50% of the muscle: 3 points.
- **End-stage appearance**: entire muscle replaced by increased density of connective tissue and fat: 4 points.

Our protocol to analyze a whole body muscle MRI is the following: first we quantify the degree of fatty infiltration of the muscles of all body. Then, we fill out a table containing the name of the muscles with the value of the quantification. In this way, we are able to easily identify which are the most and the less involved muscles. Finally, we compare the results with the patterns already published in the literature to identify a probable diagnostic.

We use 3-point Dixon sequences for investigation purposes only, for example in natural history studies or in clinical trials. 3-point Dixon studies require a specific software to quantify the amount of fat and water per pixel in every muscle (9). Although the process of analysis of the image is longer than with T1 sequence, the results are more reliable and can be compared from one center to the other.

We use CT scan to study patients in which muscle MRI is contraindicated, for example in patients with a metallic prosthesis, a pacemaker or with claustrophobia. In these cases, whole body CT scan is performed using the following parameters: 140 kV and 120-350 mA. Section thickness is of 1.25 mm, with a section interval of 0.6 mm. The axial images are 5 mm thick, with an increment of 5 mm.

**Muscle MRI in dystrophinopathies**

Mutations in the dystrophin gene produce different phenotypes such as Duchenne muscle dystrophy and Becker muscle dystrophy. The MRI pattern of muscle atrophy has been well investigated both in patients with Duchenne and Becker muscle disease (4, 10, 11). In both cases the pattern is very similar. Studies have been focused on pelvic, thigh and leg muscles (Fig. 2). In general muscle atrophy is symmetric. The muscles more commonly involved are:

**a. Pelvis**: *Gluteus maximus* and *medius* are involved from the onset of the disease and become progressively atrophied until late stages of the disease when all the glutei muscles are atrophic. In this later stage *psosas* and *iliacus* muscles could also become atrophic. *Obturator internus* and *externus* are rarely involved.

**b. Thighs**: Muscles from the anterior and posterior compartment of the thigh are involved in most of the patients, including *adductor major, biceps, semi-
membranosus, semitendinosus and the vasti. In contrast sartorius, gracilis and adductor longus are not involved until later stages of the disease.

c. Legs: There is a common involvement of both gastrocnemius, soleus and peroneus muscles. In contrast tibialis posterior is not commonly involved through all the progression of the disease.

Keys for the diagnosis

In more than 90% of the patients there is a moderate to severe atrophy of gluteus maximus, medius, semimembranosus and the vasti (4).

Mild involvement or not involvement of psoas, iliacus, sartorius, gracilis and adductor longus. To find a normal adductor longus among completely atrophic thigh muscles is a good clue for the diagnosis.

There is only one study that has analysed muscles of upper limbs in patients with Duchenne muscle dystrophy (12), showing a preferential involvement of triceps, biceps, teres major and periscapular muscles including supraspinatus, infraspinatus and subscapularis. In contrast, deltoid is usually not involved.

The pattern of muscle atrophy of symptomatic carriers of a mutation in the dystrophin gene is very similar to the one found in Becker muscle dystrophy (13). As we have just mentioned, a normal adductor longus is a clue for the diagnosis. Moreover, many of the symptomatic carriers have clear asymmetric changes that, when present, can easily guide the diagnosis.

Muscle MRI in autosomal dominant limb girdle muscle dystrophies (AD-LGMDs)

1. LGMD1A or Myotilinopathy

Patients with mutations in the MYOT gene share a common pattern of muscle fatty infiltration on MRI studies with other myofibrillar myopathies, such as the ones

![Figure 3](image-url). Muscle MRI of two patients with mutations in the MYOT gene. A). Patient with mild weakness: muscle MRI shows mild involvement of gluteus minimus, biceps (Bi), tibialis anterior (TA) and tibialis posterior (TP). B) Patient with severe weakness: muscle MRI shows involvement of all glutei muscles, although gluteus maximus is less involved than gluteus minimus or medius. In the thighs, semitendinosus (ST) is not involved. In contrast, there is a clear involvement of the posterior muscles of the thighs. Tibialis posterior (TP) is more atrophic than peroneus muscle (Per).
produced by mutation in the ZASP and the FLNC gene [8, 14, 15] (Figure 3). The muscles involved are:
a. **Pelvis**: Although there are not enough data to establish a common pattern, in our experience gluteus maximus is less involved than gluteus medius and minimus.
b. **Thighs**: In general, adductor major, biceps and semimembranosus are involved in all patients. Vasti muscles and sartorius can be also atrophic in many cases. In contrast semitendinosus is not involved until late stages of the disease.
c. **Legs**: There is a common involvement of gastrocnemius medialis, soleus, tibialis anterior and posterior. Peroneus muscles is not involved until late stages of the disease.

**Keys for the diagnosis**

To find a spared semitendinosus muscle associated to a severe atrophy of biceps and semimembranosus is a clue for the diagnosis.

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Atrophy of tibialis posterior in a higher degree than peroneus muscle is also useful to identify this disease.

2. **LGMD1B or Laminopathy**

There are many publications describing the pattern of muscle involvement in patients with mutations in the LMNA gene (16-18)). The studies already published in adult patients have been focused on muscles of the lower limbs. The pattern seems to be homogeneous among patients and is characterized by a symmetric involvement of posterior muscles of the thighs and legs associated to a severe involvement of the paravertebral muscles (Figure 4). We have observed that this pattern is not different of the muscle involvement observed in patients with mutations in the STA gene (19). The muscles preferentially involved are:

a. **Trunk muscles**: Paraspinal muscles become involved from the onset of the disease, especially multifidus and longissimus. Different to other muscle diseases, such as Pompe disease, abdominal muscles are not commonly involved during the progression of the disease.

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**Figure 4.** Muscle MRI of two patients with mutations in the LMNA gene. A) Patient with mild weakness. In this patient we observed fatty infiltration in the gluteus medius, semimembranosus (SM) and gastrocnemius medialis (GMed) muscles. B) Patient with severe weakness: fatty infiltration was observed in gluteus minimus and medius. All the muscles of the thighs were involved, except rectus femoris (RF), adductor longus (AL), sartorius (Sar) and gracillis (Gra) that were not atrophic.
b. Pelvis: *Gluteus minimus* and *medius* are involved in most of the patients, and in all the cases analysed by us these two muscles have a more severe degree of muscle atrophy than *Gluteus maximus*. *Psoas* muscle is rarely involved in these patients.

c. Thighs: The posterior muscles of the thighs, especially *semitendinosus*, the long head of *biceps* and the *adductor major* muscles are involved from the onset of the disease. In later stages *semitendinosus* and the short head of the *biceps* become also atrophic. *Vastus* muscles are involved in most of the patients: *vastus intermedius* is usually the first *vasti* muscle involved, followed by *vastus lateralis* and *medialis*. We have observed that the lateral part of the *vastus lateralis* is not involved until very late stages of the disease. *Rectus femoris* is normally not involved and becomes hypertrophic in many patients, as well as *sartorius* and *gracilis* muscles.

d. Legs: The posterior compartment of the legs is involved in all patients. *Medial gastrocnemius* is severely involved in most of the patients, associated in many cases to atrophy of *soleus* and *gastrocnemius lateralis*. The muscles of the anterolateral compartment are rarely involved during the progression of the disease.

**Keys for the diagnosis**

The combination of atrophy involving the anterior and posterior compartment of the thighs, associated to a severe atrophy of the posterior muscles of the legs and hypertrophy of *rectus femoris* is a very suggestive pattern of involvement in patients with mutations in the *LMNA* gene.

### 3. LGMD1C or Caveolinopathy

There is not enough information published of patients with caveolinopathy to establish a pattern of muscle involvement. We have studied patients with rippling muscle disease or isolated hyperckemia and the muscle MRI is normal in these cases.

### 6. LGMD-1E or Desminopathy

Patients with mutation in the *DES* gene develop a myofibrillar myopathy. The radiological features are very similar to those found in patients with mutation in the *CRYAB* gene [8]. There are many reports describing the common features in these patients, although the studies have focused mainly on thighs and legs area (Fig. 5) (14).

a. Thighs: Fatty infiltration involving the *semitendinosus*, the *gracilis* and the *sartorius* muscles occurs from the onset of the disease. This pattern is very suggestive of desminopathy and is not commonly observed in other diseases. Latter on, the *vasti* can also be involved.

b. Legs: *Peroneus* is the most commonly involved muscle in these patients, associated commonly to atrophy of the posterior muscles of the legs.

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**Figure 5.** Muscle MRI of a patient with mutation is the *DES* gene. Muscle MRI shows a preferential involvement of *semitendinosus* (ST), *sartorius* (Sar) and *gracillaris* (Gra) muscles, while in contrast *semitendinosus* (SM) and *biceps* (Bi) are not involved. In the legs, there is an involvement of *tibialis anterior* (TA) and peroneus muscles (Per). Courtesy of Dr. Giorgio Tasca.
Muscle MRI in autosomal recessive limb girdle muscle dystrophies (AR-LGMDS)

1. LGMD2A or calpainopathy

There is an important variability in the patterns described in patients with mutations in the CAPN3 gene (21-23). However, there are some common traits that can be found in most of the patients (Figure 6). The common muscles involved in this disease are:

a. **Trunk and pelvis area**: Paravertebral muscles are involved in most of the patients from the onset of the disease. In contrast, abdominal muscles are rarely involved and they are not affected until late stages. The glutei muscles are also involved from the onset, especially gluteus medius and minimus, while gluteus maximus tend to be involved in more advanced patients (24). Psoas muscles is not commonly involved until late stages.

b. **Thighs**: Adductor major and longus and semimembranosus are the first muscles involved. Later on the progression of the disease, the rest of the muscles of the posterior compartment of the thigh become involved. In many patients there is a clear difference between the anterior muscles (nearly spared) and posterior muscles (atrophic) of the thighs. However, involvement of the anterior compartment is not uncommon, and is frequently observed in medium and advanced patients. The vasti muscles are involved: in a first step the vastus intermedius and then the vastus medialis and lateralis. Rectus femoris can also be involved in many patients. In general, sartorius and gracillis are not involved until very late stages of the disease.

c. **Legs**: One of the clues for the diagnosis is the common involvement of gastrocnemius medialis and soleus from the onset of the disease. However, as the disease advances, atrophy involves the gastrocnemius lateralis, the peroneus and the tibialis anterior.

![Figure 6](image-url)
Muscle MRI in muscular dystrophies

**Tibialis posterior** is not commonly involved until very late stages of the disease.

**Keys for the diagnosis**

a. Predominant involvement of posterior muscles of the thighs associated to atrophy of *gastrocnemius medialis* and *soleus*.

b. The lateral part of the *vastus lateralis* is not involved until late stages of the disease. Different to other diseases, *rectus femoris* is also involved in many patients.

2. **LGMD2B or disferlinopathy**

The reports published on the radiological features of patients with mutation in the DYSF gene have mainly been focused in the study of the lower limbs (Fig. 7). The two most common phenotypes associated to mutations in the DYSF gene, the limb girdle muscle dystrophy and the distal posterior myopathy (Miyoshi myopathy) are not differentiable using MRI (2, 25). Common muscles involved are:

- **Pelvis:** *Glutei minimus* is commonly involved in dysferlin patients. However, the *glutei medius* and *maximus* are also involved later on the progression of the disease. Psoas muscles may be not involved until late stages of the disease.
- **Thighs:** Adductor muscles are involved from the onset of the disease. The first change is an hyperintense signal in STIR sequences, reflecting the presence of oedema before fatty muscle infiltration is observed in T1 sequences (2). *Vastus lateralis*, *medialis* and *intermedius* are involved early in the progression of the disease. Later on, posterior muscles of the thighs, especially the *semitendinosus* and the *semimembranosus* become atrophic. *Biceps* short head is less affected in many patients, until late stages of the disease when all muscles of the thighs, except *sartorius* and *gracillis* become atrophic.
- **Legs:** An hyperintense signal in STIR sequences involving the *gastrocnemius medialis* is commonly ob-

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*Figure 7.* Progression of muscle atrophy in patients with mutations in the DYSF gene. There is a mild involvement of the adductor major (AM) and gastrocnemius medialis (GMed) in the first stages of the disease (A). In a second stage (B), atrophy involves also the vastus lateralis (VL) and the soleus (Sol). Then (C), fatty infiltration progresses and involves the rest of the vasti muscles and the posterior muscles of the thighs, but sartorius (sar) and gracillis (gra) are commonly not involved. In the legs, the vastus lateralis and peroneus become atrophic. In the most advanced stages (D), all the muscles are infiltrated by fat, but in some patients tibialis posterior (TP) may be not involved.
served at the onset of the diseases. Atrophy involves this muscle first and then progress affecting gastrocnemius lateralis and soleus. In a more advanced stage antero-lateral muscles are also atrophied. Tibialis posterior is not involved until late stages.

**Clues for the diagnosis**

STIR hyperintensities in adductor and gastrocnemius medialis muscles at the onset of the disease (26, 27).

Involvement of the vasti muscles early in the progression of the disease associated to atrophy of adductor muscles and gastrocnemius medialis.

3. Sarcoglycanopathies (LGMD 2C-F)

The radiologic characteristics of the patients with mutations in the sarcoglycan genes are not fully established. There are only some case reports published (24). In our experience, the pattern seems to be homogeneous among patients (Tasca et al, in preparation). There is a severe involvement of all glutei muscles and the psoas in most of the patients reported. In the legs, muscles of the anterior and posterior compartment are also severely affected from the onset without involvement of sartorius and gracillis. Different to other muscle dystrophies, muscles of the legs are not involved until late stages of the disease (Figure 8).

4. LGMD-2I

LGMD-2I is produced by mutations in the FKRP gene. Muscle MRI features have been widely reported (24, 28), and the pattern seems to be very similar to patients with mutation in the CAPN3 gene.

a. **Pelvis**: Glutei muscles are involved from the onset of the disease, being severely involved in patients in an intermediate stage of the disease. Psoas and iliacus muscles are not involved until late stages of the disease.

b. **Thighs**: The posterior compartment of the legs is involved from the onset of the disease. Biceps, semitendinosus and semimembranosus are the most severely involved muscles in these patients. Vasti muscles are progressively involved in LGMD-2I patients. Vastus intermedius is the most common vasti muscle involved, associated to a variable degree of involvement of vastus lateralis and medialis. Rectus femoris can be involved in many cases, associate to atrophy of the adductor muscle. In later stages all muscles can be involved, being sartorius and gracillis the less atrophic muscles.

c. **Legs**: Both gastrocnemius and soleus are involved from the onset of the disease. In later stages, peroneus and tibialis anterior become also involved, although they are commonly less atrophic than the posterior muscles of the legs.

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**Figure 8.** Muscle MRI of patients with sarcoglycanopathy and titinopathy. A: 35 years old patient with LGMD-2D: there is a severe involvement of pelvic muscles (including glutei and psoas muscles) and thigh muscles. Sartorius (sar) and gracillis (gra) are not involved until advanced stages of the diseases. In contrast leg muscles are not involved. B: Muscle CT of a 62 years old patient with distal titinopathy: preferential involvement of the tibialis anterior (TA) and the posterior muscles of the thighs is observed. In contrast, the pelvis muscles are not involved.
Clues for the diagnosis

Severe involvement of glutei muscles associated to atrophy of the posterior muscles of the thighs and legs. 

*Vastus intermedius* is more involved than *lateralis* and *medialis* in many cases. In general atrophy affects first *vastus medialis* than *lateralis*.

5. LGMD2J or Titinopathy

Patients with the classical titin phenotype in which there is a preferential involvement of the *tibialis anterior*, have a very characteristic muscle MRI (7). It is very common to find an isolated atrophy of the *tibialis anterior* muscle, that can be the only muscle affected during many years. Later on the progression of the disease, atrophy can also affect *soleus* muscles, the posterior compartment of the thighs and the *rectus femoris* (Fig. 8).

In recent years, many different phenotypes of muscle disease have been associated to mutations in the *TTN* gene, including congenital myopathies or patients with proximal weakness of the lower and upper extremities. There is not a clear muscle MRI pattern in these cases.

However, patients with the hereditary myopathy and early respiratory failure phenotype (HMERF), have a common pattern of muscle MRI involvement. It has been reported that *iliacus, psoas, obturator, semitendinosus, sartorius* and *gracillis* are commonly involved in these patients.

6. LGMD2L

LGMD 2L is produced by mutations in the *ANO5* gene. Although this disease has been recently described, the muscle MRI pattern is very well known and seems to be homogenous among patients (29, 30) (Fig. 9). In these cases, muscle MRI features are very similar to those de-

![Figure 9](image-url)

*Figure 9.* Muscle MRI of a patient with mutations in the *ANO5* gene. A: Muscle MRI of a 53 years old patient with mild involvement of the lower limbs. Axial images showed no involvement of pelvic muscles, but in contrast there was a clear asymmetric involvement of vastus lateralis (VL) and posterior muscles of the thighs. In the legs, muscles of the posterior compartment were also involved, including soleus (Sol) and gastrocnemius medialis. B and C: hypointensities in STIR sequences were observed in the posterior muscles of the thighs and in the vastus medialis muscle (arrows).
scribed in patients with mutations in the DYSF gene (31). The common muscles involved are:

a. Pelvis: Glutei, psoas and iliacus muscles are commonly not involved until late stages of the disease.

b. Thighs: Adductor muscles and posterior muscles of the thighs, especially the semitendinosus (32), are involved from the onset of the disease. As the weakness progresses there is an involvement of the vastus lateralis, medialis and intermedius. As in dysferlinopathy patients, sartorius, gracilis and rectus femoris are not involved until late stages. STIR hyperintensities are commonly observed in many muscles before atrophy is clearly detected in T1 sequences. The presence of asymmetries in the degree of muscle atrophy is very common among patients.

c. Legs: Soleus and gastrocnemius medialis are commonly involved from the onset of the disease, followed by atrophy of the gastrocnemius lateralis, peroneus and tibialis anterior muscles.

**Clues for the diagnosis**

Asymmetric involvement of the thighs muscles, especially in the degree of involvement of the vasti muscles. Glutei muscles are less involved than muscles of the thighs and legs until very late in the disease.

Increase in the STIR signal is observed in many cases from the onset of the disease.

**Muscle MRI in other hereditary myopathies**

1. Adult onset Pompe disease

Pompe disease, also known as glycogenosis type II, can manifest as slowly progressive weakness involving axial and proximal muscles of the limbs. Pompe patients are in many cases misdiagnosed as patients with muscle dystrophies. Muscle MRI is very useful to identify these patients, as the pattern has been widely described and seems to be very characteristic (Fig. 10). There is a preferential involvement of tongue from the onset of the disease, also in patients without clear clinical involvement of this muscle. Perisacral muscles are commonly involved, especially the subscapularis muscle that is atrophic in many cases (33). Paravertebral muscles are affected early in the progression of the disease, especially the multifidus and longissimus muscle (34). Atrophy of the abdominal muscles runs parallel to progression of the disease. Patients may develop a severe atrophy of rectus abdominis, obliquus internus and externus and transversus abdominalis. Glutei muscles are also involved in all patients. In general these muscles are more involved than psoas-iliacus muscles. The involvement of the adductor muscles occurs early in the progression of the disease followed by involvement of posterior muscles of the thighs, vastus intermedius and finally vastus lateralis and medialis (35). In most of the cases, there is not involvement of the muscles of the legs, than can be normal also in wheel-chair bound patients.

2. Myopathies associated to mutations in the VCP gene

Mutations in the VCP gene have been associated to a characteristic triad of diseases: myopathy with proximal and/or distal involvement in the upper and lower limbs, Paget disease and fronto-temporal dementia (36). Although the muscle MRI pattern has not been published, in our experience atrophy involves muscles of the pelvis, thighs and legs. The most characteristic finding in this disease is that fatty infiltration is not homogeneous and in many cases shows a patchy distribution among the muscles (Fig. 11).

3. Oculo-pharingeal muscle dystrophy

OPMD disease is characterized by the triad of ptosis, dysphagia and proximal weakness of the lower and upper limbs. Although the symptoms are very suggestive, sometimes it can be difficult to differentiate these patients from patients with mitochondrial myopathy. In our experience, muscle MRI is useful in this case. Muscle MRI of the pelvis shows a progressive involvement of glutei muscles, specially glutei minimus. Psoas is also involved early in the progression of the disease. In the thighs there is a preferential involvement of posterior muscles of the thighs and of the adductor muscles. In the legs, soleus and peroneus muscles are commonly involved early. In fact, soleus is commonly more atrophic than gastrocnemius muscles also in advanced patients (37) (Fig. 11).

4. Collagen VI related myopathies.

Although mutations in one of the three COLVI genes have been commonly related with congenital muscle dystrophies, this disease can also present during the adulthood. Muscle MRI is highly specific and has been described elsewhere. The main finding is a muscle atrophy involving the periphery of the vastus lateralis and medialis, while the center of the muscle is not involved (38). This pattern has been described as concentric atrophy (Fig. 11). A similar pattern can also be observed in the gastrocnemius medialis, where there is a ring of atrophy between the gastrocnemius medialis and soleus muscle. Muscle atrophy can also involve other muscles, such as the rectus femoris muscle, the posterior compartment of the thighs, the paravertebral muscles and the upper limbs muscles. In this latter case, a band of muscle atrophy is commonly observed in the triceps and in the subscapu-
Muscle MRI in muscular dystrophies

laries. It has been suggested than in those patients with unknown mutations showing this characteristic pattern, genetic tests can be performed without histologic demonstration of collagen VI deficiency.

5. Facio-scapulo-humeral muscle dystrophy (FSHD)
The clinical diagnosis of FSHD is easy to achieve based only in clinical symptoms in most of the patients. However, muscle MRI studies can be helpful in un-

Figure 10. Muscle MRI of patients with Pompe disease. A: Whole body muscle MRI of a 35 years old patient with mild muscle involvement showing atrophy of tongue, paravertebral and abdominal muscles, gluteus medius (Glu.Me) and posterior muscles of the thighs. B: Muscle MRI of a 43 years old patients with moderate weakness showing involvement of tongue, subscapularis (SSc), paravertebral and abdominal muscles, gluteus medius (Glu.Me) and gluteus minimus and adductor major (AM). C: Muscle MRI of a 44 years old patients with severe weakness showing involvement of tongue, subscapularis (SSc), paravertebral, abdominal, all glutei muscles and anterior and posterior muscles of the thighs.
clear cases. Although FSHD is one of the most common muscle diseases, there are not enough studies on which patterns are commonly observed in this disease. Tasca and collaborators studied MRI muscle involvement of the upper limbs and reported a preferential involvement of periscapular muscles including trapezius, serratus anterior, latissimus dorsi, pectoralis major and rhomboids (39). In contrast, supraspinatus, infraspinatus and subscapularis were not commonly involved in this disease. The frequency of asymmetric involvement was very high. The studies focused on lower limbs have shown a predominant involvement of Tibialis anterior in most of the cases (40). Other muscles can also be atrophic such as the vastii, the posterior compartment of the thighs and both gastrocnemius. Asymmetric changes were also very common in the lower limbs, and this is probably the most important clue for the radiologic diagnosis of the disease (Fig. 12).

Conclusions

Muscle MRI has become a useful tool in the diagnosis of patients with muscle dystrophies. The standardization of the study protocols has allowed obtaining information from many patients with different muscle diseases. But still, most of the studies published are performed in relatively small cohorts of patients and are in general transversal. More information regarding progression of the atrophy during the natural history of the disease is needed. Collaboration between different study groups is necessary to increase the number of patients analyzed and to include patients in different clinical stages (from presymptomatic to advanced patients). In this sense the effort performed by the European Commission creating an E-COST action centered in the application of muscle MRI to the study of muscle diseases is remarkable (http://mymri.eu). Patients associations have also shown interest in radiological studies. This is the case of the Jain Foundation that sponsors a clinical-radiological study to describe the natural history of patients with LGMD-2B that has already recruited 196 patients all over the world (http://www.jain-foundation.org). Although the diagnosis of muscle dystrophies still continues to be based on a careful clinical history, a detailed physical examination and a complete study of the muscle biopsy, information on the muscle MRI pattern could be helpful for the genetic diagnosis of these diseases. Along the same lines, a high level of expertise is needed to identify muscle MRI patterns, that in many cases are not patognomonic of a single disease (41). In this sense, the creation of training schools will improve the level of knowledge among radiologists and neurologists all over the world, making muscle MRI studies more useful for clinical practice.

Figure 11. Muscle MRI of patients with collagen VI related myopathy, HIBM PD and OPMD muscle dystrophies. A: Muscle MRI of a 20 years old patient with mutations in the COL6A3 gene showing concentric atrophy of the vastus lateralis (double arrow) and a band of atrophy between gastrocnemius medialis and soleus (single arrow). B: Muscle MRI of a 56 years old woman with a mutation in the VCP gene producing patchy atrophy of the glutei, posterior muscles of the thighs and both gastrocnemius. C: Muscle MRI of a 72 years old patient with OPMD showing preferential involvement of posterior muscles of the thighs and the soleus muscle.
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References


Figure 12. Muscle MRI of patients with Facio-Scapulo-Humeral muscle dystrophy. Muscle MRI of different patients with FSH muscle dystrophy showing asymmetries in the thighs and legs studies.