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Peculiar clinicopathological features of immune-mediated necrotizing myopathies

Yves Allenbach\textsuperscript{a,b} and Olivier Benveniste\textsuperscript{a,b}

\textbf{Purpose of review}  
In the past decade, immune-mediated necrotizing myopathies have emerged as a separate entity in the heterogeneous group of autoimmune myopathies. This group is characterized by clinical manifestations restricted to the muscle tissue, and until recently, the definition was based on muscular pathological features.

\textbf{Recent findings}  
It was shown that they are the most severe autoimmune myopathies in term of muscle damages. They have been associated with two myositis-specific antibodies: either anti-signal recognition particle (anti-SRP) or anti-hydroxy-3-methylglutaryl-CoA reductase (anti-HMGCR) antibodies. These two antibodies are now considered as immune-mediated necrotizing myopathy (IMNM) diagnostic criteria. Each antibody delineates a homogenous subgroup of IMNM patients in terms of severity and IMNM without myositis-specific antibodies have a high risk of malignancy. In addition, pathological observations as well as in-vitro experiments suggest the pathogenic role of anti-SRP and anti-HMGCR antibodies.

\textbf{Summary}  
IMNM are muscle-specific autoimmune diseases associated with a severe weakness and a risk poor muscle strength recovery. Anti-SRP and anti-HMGCR antibodies are specifically associated with this condition and are crucial for the diagnosis and the prognosis. The muscle biopsy remains necessary for IMNM diagnosis in absence of myositis-specific antibodies.

\textbf{Keywords}  
autoantibodies, hydroxy-3-methylglutaryl-CoA reductase, myositis, necrotizing myopathies, signal recognition particle

\textbf{INTRODUCTION}  
The spectrum of immune myopathies range from muscle-specific autoimmune diseases to systemic autoimmune diseases. Historically, they were classified into two categories, namely, polymyositis and dermatomyositis, based on the presence or absence of the characteristic dermatomyositis skin rash \cite{1}. Nevertheless, both groups are heterogeneous, as patients may present with extramuscular manifestations and/or different types of myositis-specific autoantibodies \cite{2}.

Initially, in 1986, antisignal recognition particle (anti-SRP) antibody was identified in a subgroup of polymyositis patients \cite{3}. Later, in 2002, it was shown that muscle biopsies from anti-SRP antibody positive (anti-SRP\textsuperscript{+}) patients showed the presence of necrotic muscle fibre without significant muscle inflammation \cite{4}. One year later, a group of immune-mediated necrotizing myopathies (IMNMs) was recognized for the first time as a separate entity. The definition was based on pathological criteria showing predominant muscle fibre necrosis with no or mild muscle infiltrates \cite{5}. On the basis of this definition, a new myositis-specific antibody targeting the hydroxy-3-methylglutaryl-CoA reductase (HMGCR) protein was discovered in a subset of IMNM patients \cite{6,7}.

Now, IMNMs are a homogenous group of severe autoimmune muscle diseases without clinically relevant extramuscular manifestations. Yet, variations can be observed in IMNMs depending on serological status. This is one of the reasons why IMNM diagnosis criteria have been refined recently \cite{8} to distinguish anti-SRP\textsuperscript{+} IMNM, anti-HMGCR\textsuperscript{+} IMNM and antibody negative IMNM.

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**KEY POINTS**

- IMNM are the most severe autoimmune myopathies with a risk of poor muscle recovery.
- In presence of either anti-SRP or anti-HMGCR antibodies, the muscle biopsy is not necessary anymore to the diagnosis.
- For IMNM without myositis-specific antibodies, the muscle biopsy remains necessary for the diagnosis and this group has a high risk of malignancy.
- Pathological finding and in-vitro experiment suggest the pathological role of anti-SRP and anti-HMGCR antibody.

In this review, we aim to describe the clinico-pathological features of INMN patients as well as prognosis, pathophysiology and therapeutic approaches, focusing on the recent advances in the field.

**MUSCLE CLINICAL PICTURE**

IMNMs represent 20–30% of patients suffering from auto-immune myositis [9,10*,11], and the overall prevalence of myositis is 14 out of 10 000 [12]. It can occur at any age, ranging from juvenile cases to elderly patients [13,14]. Most of the cases occur between 40 and 60 years of age, and women are more frequently affected [7,9,10*,13]. Anti-HMGCR+ IMNMs may be triggered by a statin exposure in one-half to two-thirds of the cases [7,13]. The statin exposure in IMNM patients varies depending on the age of onset; in patients older than 50 years, a statin exposure is observed in greater than 90% of the cases [7]. Interestingly, Dr. A. Mammen suggests that statins are also present in some foods and dietary supplements (oyster mushroom, red yeast rice or pu-erh tea), and this may trigger the disease, especially in Asian people in whom statin drug exposure in anti-IMNM patients is low [15].

At presentation, patients usually complain of muscle signs. Extramuscle manifestations are uncommon. There are two clinical phenotypes of the disease: patients with a subacute onset and patients with a slowly progressive disease.

Patients with a subacute onset are the most common and account for greater than two-thirds of the cases [13,14]. Typically, patients complain from a rapidly progressive muscle deficit occurring within weeks or months (less than 6 months) before the first hospital visit; they generally visit the Rheumatology department. In addition, myalgia is regularly reported (30–50%) in IMNMs [10*,13,16]. One-quarter to one-third of anti-SRP+ or anti-HMGCR+ patients suffer from a slowly progressive onset of disease, greater than 6–12 months [13,14]. These later patients may present as dystrophic patients and are seen in Neurology/Myology departments.

Clinical examination shows a severe proximal, bilateral and symmetric muscle weakness. The hip flexor muscles are the most severely affected compared with other muscle groups in anti-SRP+ and anti-HMGCR+ patients [10*,17*]. Compared with other autoimmune myopathies, INMN patients more frequently harbour severe muscle weakness [16,18*]. Nevertheless, there are some variations between anti-SRP+ and anti-HMGCR+ patients. Anti-SRP+ patients suffer from a more severe muscle weakness than anti-HMGCR patients [9,18*].

With a slowly progressive onset, muscle strength can be clinically considered normal (isolated increased creatine kinase level) for years, but more frequently, in addition to the muscle weakness, a muscle atrophy with some cases of winged scapula is observed [14], mimicking a limb girdle muscle dystrophy.

Life-threatening complications must always be considered. Swallowing troubles are frequent in IMMN patients, especially in anti-SRP+ patients where they occur in more than half of cases [9].

In addition, dyspnoea related to a weakness of the respiratory muscles may be observed. Usually, it occurs in very severely affected patients in whom exercise dyspnoea is difficult to demonstrate. The patients may complain of morning signs related to hypercapnia. This lung insufficiency can be screened for by nocturnal polygraphia, morning arterial gas measurements and pulmonary functional testing, ideally performed in both the sitting and supine position. Indeed, a more than 20% decrease in vital capacity measured in the supine position versus the sitting position may be helpful in detecting severe or predominantly diaphragmatic weakness [19].

According to myopathological findings, the mean creatine kinase level in IMMN is higher than that observed in other myositis cases [9,18*,20]. Higher creatine kinase levels are also measured in anti-SRP+ patients than anti-HMGCR+ patients [9,18*].

**DIAGNOSTIC TOOLS**

**Myositis antibody-detection**

In the case of autoimmune myositis, screening for myositis auto-antibodies is crucial, as by definition,
it is a key diagnostic tool and separates a homogeneous group of patients in terms of phenotype and prognosis [2]. In IMNM, anti-SRP and anti-HMGCR are considered (for the first time in autoimmune myopathies) a key diagnostic criterion [2]. In the case of IMNM, the clinician must test for anti-SRP as well as anti-HMGCR [21,22]. Screening for anti-nuclear antibodies was not sensitive enough to detect anti-SRP and anti-HMGCR antibodies using a screening test in HEp-2 cells [21,22]. SRP and HMGCR are intracytoplasmic proteins related to the endoplasmic reticulum, and indirect immunofluorescence may be negative (Fig. 1). SRP is key in the delivery of newly synthesized proteins [23], and HMGCR is a key enzyme in cholesterol biosynthesis [24]. There are different commercial kits for myositis-specific auto-antibody detection that have a limited risk of false-positive or false-negative results [2,11,21,22]. Immunoprecipitation remains the gold standard of auto-antibody detection, but it is not routinely feasible. For these reasons, the results of the tests must be challenged with the clinico-pathological picture, and a retest can be performed using different methods if doubts exist.

Muscle biopsy

In the case of a typical clinicobiological picture of IMNMs with a positive detection of either anti-SRP or anti-HMGCR antibody, a muscle biopsy is not necessary to diagnose anti-SRP+ or anti-HMGCR+ IMNM [8*]. In other cases, a muscle biopsy must be performed.

The detection of the elementary lesions (muscle fibre, vascular, connective domains) and their distribution is important to recognize IMNMs (Fig. 2) [25]. Initially, IMNMs were defined on the basis of pathological criteria, including predominant muscle necrosis, whereas inflammatory infiltrates were mild or absent [5]. It has been shown that the percentage of necrotic muscle fibres is low in IMNMs (3.2% in anti-SRP+ patients vs. 1.8% in anti-HMGCR patients, P < 0.05) [18*]. The necrotic muscle fibres are randomly distributed, whereas they are confined in the perifascicular area in antisynthetase syndrome, another condition wherein significant muscle fibre necrosis is observed [18*]. However, indirect signs of necrosis, such as regenerating muscle fibres, are much more frequent [18*].

According to the first definition of IMNM, the inflammatory infiltrates are mild and mainly composed of macrophages [9,18*,26]. Nevertheless, significant inflammatory infiltrates with cell densities reaching the same range as those measured in other myositis are observed in one-quarter of IMNM cases (Fig. 2) [18*]. A subgroup of IMNM patients may have diffuse sarcolemmal major histocompatibility complex (MHC)–I positive staining. Together, those data showed that a subset of INMN patients have significant signs of muscle inflammation that are correlated with the percentage of necrotic muscle fibres [18*]. Thus, significant muscle inflammation is not considered an exclusion criterion in the new antibody-positive IMNM definition [8*].

The pathological criteria for auto-antibody negative IMNM are the presence of necrotic fibres and regenerative fibres (with different stages) with a scattered distribution and the presence of macrophage-predominant, paucilymphocytic infiltrates. Additional consistent features are sarcolemmal...
MHC class I expression (seen on nonnecrotic/nonregenerating fibres), sarcolemmal complement deposition, endomysial fibrosis and proliferation, and enlarged capillaries [8*].

EXTRASKELETAL MUSCLE FEATURES

Cardiac muscle features
Conduction abnormalities, rhythm disorders and left ventricular dysfunction have been reported in 2–40% of anti-SRP cases [14,16,20,27,28]. If clear myocarditis has been reported in anti-SRP+ patients, the significance and the specificity of all of the above-mentioned cardiac manifestations remain to be established. Nevertheless, in our opinion, a systematic cardiac screening is necessary for anti-SRP patients. To date, cardiac involvement has not been clearly associated with anti-HMGCR patients.

Extramuscular autoimmune features
IMNM may be considered a pure muscle autoimmune disease. Specifically, in anti-HMGCR no auto-immune extra-muscle involvement has been
reported. Anti-SRP+ patients do not suffer frequently from extramuscular manifestations, even though arthralgia (0–39%) [14,20] and Raynaud’s phenomenon (0–26%) [20,29] have been reported. Anti-SRP IMNM may also result in an interstitial lung disease that is detected when a computed tomography (CT)-scan is performed (0–22%) [20,28]. Again, this finding is not clinically significant and is not associated with pulmonary function test abnormalities.

On the contrary, it must be mentioned that some overlapping myositis may have myopathological features similar to IMNM, with diffuse randomly distributed necrotic/regenerating muscle fibres. This is the case in myositis-associated scleroderma [30].

**Malignancy in immune-mediated necrotizing myopathies**

An increased risk of malignancy in autoimmune myopathies is mainly observed in dermatomyositis, but it has also been reported in ‘polymyositis’ [31]. On the basis of this observation and previous case reports describing IMNM and cancer, the risk of malignancy in IMNM patients was compared with that expected in a sex and age-matched population. It was shown that patients with antibody-negative IMNM have a higher risk of malignancy, whereas the risk was similar for anti-SRP patients [32]. A mild increase was observed in two studies of anti-HMGCR+ patients [32*,33], but this was not observed by others [17*]. In the case of cancer association, malignancy occurs within 3 years before or after the diagnosis and in most cases within 1 year of diagnosis [32*]. Cancer affects mainly patients older than 50 years. No specific type of cancer is observed in IMNM, and patients with malignancy have a lower survival rate [32*].

**IMMUNE-MEDIATED NECROTIZING MYOPATHY PROGNOSIS**

Patients suffering from autoimmune myopathies have a higher mortality rate than the general population [34]. The main causes of death in autoimmune myopathies are malignancy and diseases of the respiratory and circulatory systems [34]. All of these life-threatening complications, except severe interstitial lung disease, may occur in IMNM patients.

One-quarter of IMNM patients suffer from difficulties in their daily living, graded as modified Rankin Scale scores of 3–5 [9]. Only 50% of anti-SRP+ patients reached near-full or full strength after 4 years of treatment, and a younger age at onset is associated with more severe weakness [10*]. Only 44% of anti-HMGCR+ patients reached full strength with immunosuppressive therapy, and younger patients had more severe disease and a worse prognosis than older patients [17*]. Of note, statin exposure was not independently associated with the rate of muscle strength improvement [17*].

These observations are in line with the presence of significant muscle damage in IMNM. Compared with the other myositis, IMNM was characterized by a higher proportion of thigh muscles with oedema, atrophy and fatty replacement (Fig. 3) [35**]. According to the clinicobiological observations, among IMNM, anti-SRP+ patients had more atrophy and fatty replacement than anti-HMGCR+ patients [35**].

Together, the data show that IMNMs are the most severe group of myositis with a poor outcome in terms of muscle strength compared with dermatomyositis or the antisynthetase syndrome.

**PATHOPHYSIOLOGY**

The pathophysiology of IMNM is not fully understood. Some human leukocyte antigen (HLA) class I and II antigens have been associated with IMNM [36–38]. Statin exposure is another factor associated with anti-HMGCR+ IMNM onset, suggesting that in genetically predisposed patients, the drug may trigger the disease. It has been shown that HMGCR expression is upregulated in regenerating myofibers in anti-HMGCR IMNM patients [7]. Similarly, HMGCR and SRP proteins were detected at the sarcolemmal level in IMNM patients [18*]. This observation and the presence of membrane attack complexes at the surface of nonnecrotic muscle fibres suggest a pathogenic role of the auto-antibodies. This hypothesis is reinforced by the observation of IgG and C1q at the sarcolemmal level in IMNM patients, suggesting classical pathway activation (antibody-dependant) of complement [18*]. The titre of the auto-antibodies correlates with the disease activity of anti-SRP+ and anti-HMGCR+ myositis [13,17*,22]. In addition, it has been shown in vitro that anti-SRP and anti-HMGCR induce muscle fibre atrophy and impair muscle regeneration [39**]. Together, these observations suggest a pathogenic role of auto-antibodies in IMNM.

**TREATMENT**

IMNM treatment remains challenging. There has been no randomized clinical trial specifically designed for IMNM patients. Retrospective studies have shown that anti-SRP and anti-HMGCR patients have a long disease duration with frequent relapses when using several treatments, including
corticosteroids and immunosuppressants [9,10*, 13,17*]. It was shown that rituximab administration was efficacious in 75% of cases [10*,40], leading to the consideration of its use as a first-line treatment in anti-SRP patients [8*]. In anti-HMGCR+ patients, intravenous immunoglobulin has been efficacious as a monotherapy [41], showing its importance in the treatment strategy for those patients [8*].

The first-line treatment strategy must be adapted to each patient’s condition and the severity of the muscle disease. The recommendations of the 224th ENMC are shown in Fig. 4.

Disease assessment is challenging in autoimmune myositis. The ACR/EULAR recommendations have been updated to capture a significant improvement in the domains (e.g. muscle, skin, lung or joints) that could be affected in myositis patients [42]. In anti-SRP+ and anti-HMGCR+ IMNM, it was shown that the creatine kinase level is associated with disease activity [13,17*], as the creatine kinase level mirrors the percentage of necrotic muscle fibres [18*]. Regarding the severe muscle damage observed in IMNM patients [35**], it seems logical to target normal creatine kinase levels to treat IMNM and determine disease remission.

In addition, a slight increase in creatine kinase levels may be a sign of important and persistent disease activity in severely affected patients, as a normal creatine kinase level is correlated with muscle mass. Indeed, IMNM patients with important clinically defined sarcopenia and low creatinine blood levels may have subnormal creatine kinase levels, whereas persistent disease activity is determined by muscle oedema using MRI.

**FIGURE 3.** Immune-mediated necrotizing myopathy whole-body MRI. Whole-body MRI (T1) of a severely affected IMNM patient shows important muscle damages in the lower limbs. The important fatty replacement (T1 hypersignal) is observed in the gluteus muscles as well as in the thigh muscles, whereas the muscles of the lower limbs are spared.
MRI of the muscle is an important tool to monitor IMNM patients, as MRI permits the measurement of disease activity and also muscle damage. MRI is sometimes used to avoid treatment escalation in patients with definitive fixed muscle damage with fatty replacement.

To conclude, IMNM is the most severe autoimmune muscular disease with regard to patient
impairment and the risk of muscle damage. IMNM can be anti-SRP+, anti-HMGCR+ or auto-antibody negative. This serological distinction allows the determination of a more definitive phenotype in the patient (more severe in anti-SRP patients), the determination of potential cancer association (mainly in antibody-negative IMNM) and the development of treatment strategies.

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Conflicts of interest
There are no conflicts of interest.

REFERENCES AND RECOMMENDED READING

Papers of particular interest, published within the annual period of review, have been highlighted as:
◆ of special interest
◆◆ of outstanding interest

8. In this conference, an international consensus was reached to refine IMNM criteria. For the first time, IMNM definition is based on serological criteria.
11. In this article, the authors described for the first time the precise muscle phenotype of anti-SRP+ patients. For the first time, they have clearly demonstrated the poor muscle outcome.

In this article, the authors described for the first time the precise muscle phenotype of anti-HMGCR+ patients. For the first time, they have clearly demonstrated the poor muscle outcome, stratifying the outcome based on the age of onset.


In this article, the authors described myopathological changes IMNM with a quantitative approach. This study shows IgG and complement deposits at the sarcocellular level and suggest a pathogenic role of auto-antibody in muscle fibre necrosis.

33. In this study, the authors have compared the prevalence of malignancy in the three IMNM subtypes (anti-SRP+, anti-HMGCR+ and seronegative) to the aged and sex-matched population. They have showed an important increased risk in anti-body-negative IMNM, whereas the risk is mild in anti HMGCR+ patients.
37. This study has reported for the first time the muscle damages occurring in a large cohort of IMNM compared with others myositis. According to the poor outcomes of IMNM, the authors showed the important fatty replacement in IMNM patients compared with the other group of IMNM.

In this study, the authors test in vitro the pathogenic role of the anti-SRP and anti-HMGCR antibodies. For the first time, the authors showed that they can induce muscle atrophy and impair muscle regeneration.