ABSTRACT

Introduction  Idiopathic inflammatory myopathies (IIMs) are rare diseases characterised by non-suppurative inflammation of skeletal muscles and muscle weakness. Additionally, IIM is associated with a reduced quality of life. Strength training is known to promote muscle hypertrophy and increase muscle strength and physical performance in healthy young and old adults. In contrast, only a few studies have examined the effects of high intensity strength training in patients with IIM and none using a randomised controlled trial (RCT) set-up. Thus, the purpose of this study is to investigate the effects of high-intensity strength training in patients affected by the IIM subsets polymyositis (PM), dermatomyositis (DM) and immune-mediated necrotising myopathy (IMNM) using an RCT study design.

Methods and analysis  60 patients with PM, DM or IMNM will be included and randomised into (1) high-intensity strength training or (2) Care-as-Usual. The intervention period is 16 weeks comprising two whole-body strength exercise sessions per week. The primary outcome parameter will be the changes from pre training to post training in the Physical Component Summary measure in the Short Form 36 health questionnaire. Secondary outcome measures will include maximal lower limb muscle strength, skeletal muscle mass, functional capacity, disease status (International Myositis Assessment and Clinical Studies Group core set measures) and questionnaires assessing physical activity levels and cardiovascular comorbidities. Furthermore, blood samples and muscle biopsies will be collected for subsequent analyses.

Ethiscs and dissemination  The study complies with the Helsinki Declaration II and is approved by The Danish Data Protection Agency (P-2020–553). The study is approved by The Danish National Committee on Health Research Ethics (H-20030409). The findings of this trial will be submitted to relevant peer-reviewed journals. Abstracts will be submitted to international conferences.

Trial registration number  NCT04486261.

INTRODUCTION

Idiopathic inflammatory myopathies (IIMs), collectively termed myositis, is a heterogeneous group of rare diseases that share features as non-suppurative inflammation of skeletal muscles and muscle weakness. The main subsets of IIMs consists of polymyositis (PM), dermatomyositis (DM), sporadic inclusion body myositis (sIBM) and immune-mediated necrotising myopathy (IMNM).

Patients with myositis generally respond well to prednisolone and other anti-inflammatory drugs, with sIBM as an exception. Even though anti-inflammatory drugs in general reduce disease activity in IIM, patients typically do not fully regain their pre-disease muscle function. In addition, patients with IIM have reduced quality of life. Patients with a disease duration of roughly 7 years scored ~50% lower within the physical domain of the Short Form 36 (quality of life) questionnaire compared with healthy age-matched adults. In a recent OMERACT (Outcome Measures in Rheumatology) survey on patient-reported outcome measures, patients with myositis listed ‘muscle symptoms’,
‘fatigue’ and ‘levels of physical activity’ as key challenging aspects of daily living. Therefore, it is paramount to address these aspects and improve physical function to increase quality of life for patients with myositis.

The effect of physical exercise has been investigated in patients with PM and DM in a few smaller non-randomised studies and in general, physical exercise was shown to be a safe and effective therapeutic tool to improve physical function and activities of daily living.

Two randomised controlled trials (RCT) concerning endurance training have been conducted in patients with PM and DM and and aerobic exercise in combination with resistive endurance training led to improvements in general health, exercise performance and aerobic capacity, respectively, in patients with PM, DM and IMNM. The effect of high intensity strength training has only been investigated in limited number of patients with myositis (excluding sIBM) with promising results in terms of increased muscle strength, reduced disease activity and reduced signs of physical impairment, along with no signs of increased inflammation within the trained muscles. However, none of these studies included assessments of quality of life and despite the promising results, a strict RCT study design is currently lacking.

Two training studies performed comprehensive immunohistochemistry analysis, with a focus on inflammation and none of the studies observed signs of increased inflammation following the training interventions. Nonetheless, the effect of high-intensity strength training at the myocellular level has never been investigated in patients with PM, DM and IMNM.

The aims of the present study, therefore, are to use a RCT study design to investigate the effects of high-intensity strength training on (1) quality of life, (2) muscle strength, muscle mass, physical function and disease activity in patients with IIM compared with patients with control (IIM) receiving Care-as-Usual and (3) additionally, explorative outcomes as the underlying myocellular adaptations will be examined by repeated muscle biopsy sampling.

METHODS AND ANALYSIS

The current RCT is registered at ClinicalTrials.gov and any changes to the protocol will be implemented here.

Study design

The study is a two-armed RCT. Sixty patients diagnosed with IIM (PM, DM and IMNM) will be included in a 16-week training intervention study (figure 1). Patients will be allocated randomly in the two subject groups in a 1:1 ratio, with stratification of IIM subgroups to ensure even distribution between intervention arms (training vs no training). The randomisation code will be generated by a biomedical laboratory technician with no further relation to the study, using an online tool (Research Randomizer, www.randomizer.org). The study has conformed to the Standard Protocol Items: Recommendations for Interventional Trials guidelines for constructing a clinical trial.

Outside the scope of the RCT, we intend to perform a 1-year follow-up measurement, which would include the same outcome variables as pre-intervention and post-intervention, with the exception of muscle biopsies.

Blinding

The two physiotherapists who will be conducting the pre and post testing of physical function, maximal lower limb muscle power and dual-energy X-ray absorptiometry (DEXA) scans will be blinded to participants’ group allocation. The physician assessing IIM-specific disease measures (disease damage and activity, etc), as well as the statistician performing all statistical analysis will also be group allocation blinded. The patients and the lead investigator in charge of the supervised training cannot be blinded for group allocation.

Patients

Only patients with an affiliation to Copenhagen University Hospital, Rigshospitalet, will be assessed for eligibility in the study. Eligible patients will be identified through the electronic patient record system at Rigshospitalet by an extraction on diagnosis codes (M33.1—‘Other dermatomyositis’, M33.2—’Polymyositis',
The weight will be progressively adjusted across successive
load corresponding to 10 RM to failure, which will be
will consist of three sets of each exercise using a training
sessions at an intensity of 15 RM. At week three each session
each exercise will be per
loads will be estimated based on maximal test (five repeti-
protocol will consist of two exercise sessions per week and
months prior to inclusion in the study.
IIM diagnosis established at
Prednisolone ≤5mg/day and stable dosage of
immunosuppressive treatment for at least 1 month prior to
inclusion in the study.
EULAR/ACR, European League Against Rheumatism/American
College of Rheumatology; IIMs, idiopathic inflammatory
myopathies.

M33.9—‘Dermatopolymyositis unspecified’ and G72.49—
‘Other inflammatory and immune myopathies, not else-
where classified’). The study intends to include stable
patients with IIM only. Specific inclusion and exclusion
criteria are listed in table 1. Patients deemed eligible by
the leading physician will receive invitation by letter or
e-mail.

Patient and public involvement
To strengthen the study in general and the strength
training protocol, five patients with myositis were recruited
for an advisory board to advise the research group in
matters relevant for the patients and their role in the
research study. The advisory board will persist through
the entirety of the study and asked to give feedback on
all matters relevant for the patients. Patients were chosen
based on age, disease length, sex and general background
to make sure the advisory board was as diverse as possible.

Intervention protocol
High-intensity strength training
The high-intensity (ie, heavy load) strength training
protocol will consist of two exercise sessions per week and
all training sessions will be supervised. The initial training
loads will be estimated based on maximal test (five repeti-
tions maximum (RM)) prior to the first training session.
The first 2 weeks will be familiarisation period, where
each exercise will be performed in three sets of 10 repeti-
tions at an intensity of 15 RM. At week three each session
will consist of three sets of each exercise using a training
load corresponding to 10 RM to failure, which will be
kept for the remaining part of the training intervention.
The weight will be progressively adjusted across successive
exercise sessions when participants are able to complete
two extra repetitions (ie, 12 repetitions) in the last set of
the respective exercise. The training protocol will be a
whole-body training protocol and consist of five exercises
using machines: horizontal bench press, horizontal leg
press, seated rows, weighted knee extension and seated
biceps curls.
The respective muscles will be working for approxi-
ately 45 s per set, with intermittent pauses of 90 s.27 28
Borg scale (6–20) will be used for assessing perceived
exertion during and following the exercise sessions (aim
following session: 16–18). All training loads for each
training session will be recorded in an individual training
diary for each patient (eg, exercise adherence, training
load and adverse events).

Care-as-Usual
Care-as-Usual is defined as maintaining the level of phys-
ical activity at the same level as prior to initiation of the
study. Furthermore, the usual medical treatment related
to the myositis disease will be maintained throughout the
timeline of the study for both groups.

Outcome variables
All study outcome measures are presented in table 2.
All outcome measures will be measured at baseline and
following the 16-week intervention period. Demographi-
cal information (age, gender, disease duration and
time from first symptoms) will be drawn from clinical
records from the electronic patient record system at
Rigshospitalet.

Primary outcome
The primary outcome variable will be the change in the
Physical Component Summary (PCS) measure from
baseline to 16 weeks assessed by the Short Form-36 health
questionnaire (SF-36), with scores ranging from 0 (worst)
to 100 (best).32 The SF-36 is proposed by The Interna-
tional Myositis Outcome Assessment Collaborative Study
Group (IMACS) as the preferred quality of life assess-
ment tool.32

Secondary outcomes
Assessment of physical function and maximal muscle strength
Physical function will be tested using Functional Index
3, 35 30 s chair rise,34 timed up-and-go35 and 2-min walk
testing.36 Leg power will be measured by power rig.37
Handgrip strength will be measured38 and lastly a test
for static balance with three feet positions (feet together,
semi tandem and full tandem) will be performed.39

Body composition
Body composition as well as whole-body, appendicular
(arms and legs) and lower-limb lean mass will be evaluated
by DEXA. Bioimpedance measures will also be collected.

Disease activity and damage
Several outcome measures proposed by the IMACS to
evaluate disease activity and disease damage in patients
with IIM will be obtained.2 Patient and Physician Global

| Inclusion criteria                                                                 | Exclusion criteria                                                                 |
|-----------------------------------------------------------------------------------|------------------------------------------------------------------------------------|
| Patients, age ≥18 years old, fulfilling the criteria for IIMs by EULAR/ACR.42 43 | Patients with sporadic inclusion body myositis and overlap myositis (myositis combined with another autoimmune rheumatic diseases), except Sjögren Syndrome. |
| Prednisolone ≤5mg/day and stable dosage of immunosuppressive treatment for at least 1 month prior to inclusion in the study. | Comorbidity preventing resistance training (eg, severe heart/lung disease, uncontrolled hypertension (systolic >160 mm Hg and/or diastolic >100 mm Hg), severe knee/hip arthritis). |
| IIM diagnosis established at least 6 months prior to inclusion in the study. | Alcohol and/or drug abuse. Defined by the guidelines issued by The Danish Health Authority. |

Table 1 Inclusion and exclusion criteria

EULAR/ACR, European League Against Rheumatism/American College of Rheumatology; IIMs, idiopathic inflammatory myopathies.
Assessment of Disease Activity and Extramuscular Global Assessment will be evaluated using a Visual Analogue Scale (VAS, 100 mm). The Manual Muscle Test will be used to determine muscle strength in eight predefined muscles. Muscle strength is graded from 0 (zero; no contraction felt in the muscle) to 10 (normal; holds test position against strong pressure). Perceived physical function is reported by the patients, using the Health Assessment Questionnaire. Plasma creatine kinase (CK) will be measured by blood sampling. Patient and Physician Global Assessment of Disease Damage will be evaluated using a VAS (100 mm).

**Questionnaires**

The International Physical Activity Questionnaire - long (IPAQ-long) concerning the level of self-reported physical activity and a questionnaire concerning medical conditions, current medication, heart symptoms, smoking habits and so on, will be filled out by all study participants. The Mental Health Component Summary measure from the SF-36 will also be recorded.

**Cardiovascular co-morbidities**

Traditional cardiovascular risk factor will be measures, including body mass index, systolic and diastolic blood pressure, plasma lipid profile (low-density lipoprotein, high-density lipoprotein, triglycerides and total cholesterol) and glycated haemoglobin (HbA1c). In addition, troponins, N-terminal pro b-type Natriuretic Peptide (NT-proBNP) and ECG will be measured.

**Explorative outcomes**

**Muscle biopsy**

Biopsy samples will be acquired ad modum conchotome vastus lateralis (~100 mg). Immunohistochemistry will be used to analyse myofiber cross-sectional area, fibre type composition, B-lymphocytes and T-lymphocytes, macrophages, satellite cells and myonuclei.

**Statistical considerations**

The calculation of the number of subjects is based on the PCS values from the SF-36 questionnaire in patients with PM and DM reported by Poulsen et al. Reported PCS values were 36.5 with a SD for 9.5. The current study is a superiority trial and intends to demonstrate a significant change with the training intervention protocol of at least 20% with a statistical significance level of 0.05 and a
statistical power of 80% while anticipating a dropout rate of 10%. Based on sample size calculations (www.scaledenvelope.com/power/continuoussuperiority) based on the above-mentioned values and dropout rate, a total of 60 patients was estimated to be recruited.

**Statistical analysis**
The primary outcome variable is the change in PCS using the SF-36 questionnaire from baseline to post 16 weeks of intervention. The statistical analysis of the primary outcome will be conducted using an ‘intention-to-treat’ approach. For the primary outcome, an independent t-test will be conducted to determine the difference in change between the two groups. Likewise, independent t-tests will be conducted to determine all other outcomes measured throughout the study. In addition, a ‘per protocol’ approach also will be employed. The criteria for being included in the ‘per protocol’ analysis is having participated in at least two-third of all exercise sessions.

**ETHICAL ASPECTS AND DISSEMINATION**

**Ethical considerations**
The study will stay true to the Helsinki declaration II and is approved by The Danish National Committee on Health Research Ethics (H-20030409). Further, the study is approved by The Danish Data Protection Agency (P-2020–553) and all data accumulated will be handled confidentially and under secrecy in accordance with the guidelines and approval conditions of The Danish Data Protection Agency. Written and verbal informed consent will be collected from all patients prior to participation in the study, according to Danish law (see online supplemental file 1 for patient consent form). In publication, there will not be any information that could identify any of patients partaking in the project.

There is no commercial interest at stake within the project. Possible ‘conflict of interests’ will be uncovered from the start of the intervention.

**Dissemination policy**
The results of the current RCT will be published in peer-reviewed journals. Abstracts will be submitted for poster presentations at international conferences (eg, American College of Rheumatology). Authorship is granted to authors who provide essential contributions to the creation of the final publications. Both contributions via writing and/or assisting in conducting the clinical trial are accepted.

**Author affiliations**

1Department of Pathology, Department of Clinical Research, University of Southern Denmark, Odense, Denmark
2Research Unit of Clinical Biomechanics, Department of Sports Science and Clinical Biomechanics, University of Southern Denmark, Odense, Denmark
3Department of Rheumatology, Odense University Hospital, Odense, Denmark
4Geriatric Research Unit, Department of Medicine, Copenhagen University Hospital, Copenhagen, Denmark
5Geriatric Research Unit, Department of Medicine, Copenhagen University Hospital, Copenhagen, Denmark
6Research Unit of Clinical Biomechanics, Department of Sports Science and Clinical Biomechanics, University of Southern Denmark, Odense, Denmark
7Department of Rheumatology, Odense University Hospital, Odense, Denmark

**Acknowledgements** We thank Professor Ingrid Lundberg (Karolinska Institute), Associate Professor Helene Alexanderson (Karolinska Institute), PhD Anders Jørgensen and the patient advisory board for their valuable input to the design of the current randomised controlled trial protocol. We used the Standard Protocol Items: Recommendations for Interventional Trials checklist when writing this report.

**Contributors** LPD is the principal investigator on the current trial. KYJ is the coordinator of the trial and has drafted the manuscript. PA, CS and HDS are co-coordinators of the trial and supply academical depth and experience. EB provided statistical expertise. JLN provided insight and expertise concerning clinical trials. All authors took part in the study design and assisted with the project funding. All authors have participated in the design of the trial and assisted with the draft of the manuscript and read and approved the final manuscript.

**Funding** This work was supported by The Danish Rheumatism Association—grant number R185–6606; The AP Møller Foundation—grant number 20-L-0001; Copenhagen University Hospital, Rigshospitalet—grant number N/A.

**Competing interests** None declared.

**Patient consent for publication** Not required.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Supplemental material** This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

**Open access** This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

**ORCID ID**
Kasper Yde Jensen http://orcid.org/0000-0003-4232-6079

**REFERENCES**

1 Wiesinger GF, Quittan M, Nuhr M, et al. Aerobic capacity in adult dermatomyositis/polymyositis patients and healthy controls. Arch Phys Med Rehabil 2000;81:1–5.
2 Rider LG, Giannini EH, Harris-Love M, et al. Defining clinical improvement in adult and juvenile myositis. J Rheumatol 2003;30:603–17.
3 Mammen AL. Necrotizing myopathies: beyond statins. Curr Opin Rheumatol 2014;26:679–83.
4 Basharat P, Christopher-Stine L. Immune-Mediated necrotizing myopathy: update on diagnosis and management. Curr Rheumatol Rep 2015;17:72.
5 Dalakas MC. Inflammatory muscle diseases. N Engl J Med 2015;372:1734–47.
6 Carroll MB, Newkirk MR, Surnmer NS. Necrotizing autoimmune myopathy: a unique subset of idiopathic inflammatory myopathy. J Clin Rheumatol 2016;22:376–80.
7 Selva-O’Callaghan A, Pinal-Fernandez I, Trallero-Araguas E, et al. Classification and management of adult inflammatory myopathies. Lancet Neurol 2018;17:816–28.
8 Zong M, Lundberg IE. Pathogenesis, classification and treatment of inflammatory myopathies. Nat Rev Rheumatol 2011;7:297–306.
9 Gordon PA, Winer JB, Hoogendijk JE, et al. Immunosuppressant and immunomodulatory treatment for dermatomyositis and polymyositis. Cochrane Database Syst Rev 2012;2012:CD003643.
10 Askanas V, Engel WK. Inclusion-body myositis: a myodegenerative conformational disorder associated with Abeta, protein misfolding, and proteasome inhibition. Neurology 2006;66:539–48.
Open access

11 Benveniste O, Guiguet M, Freebody J, et al. Long-term observational study of sporadic inclusion body myositis. Brain 2011;134:3716–84.
12 Dalakas MC. Pathogenesis and therapies of immune-mediated myopathies. Autoimmun Rev 2012;11:203–6.
13 Dalakas MC, Koffman B, Fuji M, et al. A controlled study of intravenous immunoglobulin combined with prednisone in the treatment of IBM. Neurology 2001;56:323–7.
14 Lawson Mahowald M. The benefits and limitations of a physical training program in patients with inflammatory myositis. Curr Rheumatol Rep 2001;3:317–24.
15 Regardt M, Wein Henriksson E, Alexanderson H, et al. Patients with polymyositis or dermatomyositis have reduced grip force and health-related quality of life in comparison with reference values: an observational study. Rheumatology 2011;50:578–85.
16 Poulsen KB, Alexanderson H, Daigard C, et al. Quality of life correlates with muscle strength in patients with dermatomyositis. Clin Rheumatol 2017;36:2289–95.
17 Regardt M, Mecoli CA, Park JK, et al. OMERACT 2018 modified patient-reported outcome domain core set in the life impact area for adult idiopathic inflammatory myopathies. J Rheumatol 2019;46:1351–4.
18 Alexanderson H, Stenström CH, Lundberg I. Safety of a home exercise programme in patients with polymyositis and dermatomyositis: a pilot study. Rheumatology 1999;38:608–11.
19 Alexanderson H, Stenström CH, Jensen G, et al. The safety of a resistive home exercise program in patients with recent onset active polymyositis or dermatomyositis. Scand J Rheumatol 2000;29:295–301.
20 Varjú C, Pethő E, Kutás R, et al. The effect of physical exercise following acute disease exacerbation in patients with dermatomyositis. J Rheumatol 1993;20:1340–4.
21 Escalante A, Miller L, Beadmore TD. Resistive exercise in the rehabilitation of polymyositis/dermatomyositis. J Rheumatol 1993;20:1340–4.
22 Bertolucci F, Neri R, Dalise S, et al. Abnormal lactate levels in patients with polymyositis and dermatomyositis: the benefits of a specific rehabilitative program. Eur J Phys Rehabil Med 2014;50:161–9.
23 Wiesinger GF, Quittan M, Aringer M, et al. Improvement of physical fitness and muscle strength in polymyositis/dermatomyositis patients by a training programme. Br J Rheumatol 1998;37:196–200.
24 Alemo Munters L, Dasmaltchi M, Katz A, et al. Improved exercise performance and increased aerobic capacity after endurance training of patients with stable polymyositis and dermatomyositis. Arthritis Res Ther 2013;15:R83.
25 Alemo Munters L, Dasmaltchi M, Andgren V, et al. Improvement in health and possible reduction in disease activity using endurance exercise in patients with established polymyositis and dermatomyositis: a multicenter randomized controlled trial with a 1-year open extension followup. Arthritis Care Res 2013;65:1959–68.
26 Alexanderson H, Munters LA, Dasmaltchi M, et al. Resistive home exercise in patients with recent-onset polymyositis and dermatomyositis -- a randomized controlled single-blinded study with a 2-year followup. J Rheumatol 2014;41:1124–32.
27 Alexanderson H, Lundberg IE. Disease-specific quality indicators, outcome measures and guidelines in polymyositis and dermatomyositis. Clin Exp Rheumatol 2007;25:153–8.
28 Nader GA, Dasmaltchi M, Alexanderson H, et al. A longitudinal, integrated, clinical, histological and mRNA profiling study of resistance exercise in myositis. Mol Med 2010;16:455–64.
29 de Souza JM, de Oliveira DS, Perin LA, et al. Feasibility, safety and efficacy of exercise training in immune-mediated necrotising myopathies: a quasi-experimental prospective study. Clin Exp Rheumatol 2014;32:235–41.
30 Munters LA, Loelll I, Ossipova E, et al. Endurance exercise improves molecular pathways of aerobic metabolism in patients with myositis. Arthritis Rheumatol 2016;68:1738–50.
31 Chan A-W, Tetzlaff JM, Altman DG, et al. SPIRIT 2013 statement: defining standards for protocol items for clinical trials. Ann Intern Med 2013;158:200–7.
32 Miller FW, Rider LG, Chung YL, et al. Proposed preliminary core set measures for disease outcome assessment in adult and juvenile idiopathic inflammatory myopathies. Rheumatology 2001;40:1282–73.
33 Ernste FC, Chong C, Crowson CS, et al. Functional Index-3: a valid and reliable functional outcome assessment measure in patients with dermatomyositis and polymyositis. J Rheumatol 2021;48:94–100.
34 Rikli RE, Jones CJ. Development and validation of criterion-referenced clinically relevant fitness standards for maintaining physical independence in later years. Gerontology 2013;53:255–67.
35 Poddiaud D, Richardson S. The timed “Up & Go”: a test of basic functional mobility for frail elderly persons. J Am Geriatr Soc 1991;39:142–6.
36 Alexanderson H, Bromlan L, Tölbäck A, et al. Functional index-2: validity and reliability of a disease-specific measure of impairment in patients with polymyositis and dermatomyositis. Arthritis Rheum 2006;55:114–22.
37 Hardy R, Cooper R, Shah I, et al. Is chair rise performance a useful measure of leg power? Aging Clin Exp Res 2010;22:412–8.
38 Nordenskiöld UM, Grimby G. Grip force in patients with rheumatic arthritis and fibromyalgia and in healthy subjects. A study with the Griptit instrument. Scand J Rheumatol 1993;22:14–19.
39 Puthoff ML. Outcome measures in cardiopulmonary physical therapy: short physical performance battery. Cardiopulm Phys Ther J 2008;19:17–22.
40 Dorph C, Nennesmo I, Lundberg IE. Percutaneous conchotome muscle biopsy: A useful diagnostic and assessment tool. J Rheumatol 2001;28:1591–9.
41 Jensen KY, Jacobsen M, Schroder HD, et al. The immune system in sporadic inclusion body myositis patients is not compromised by blood-flow restricted exercise training. Arthritis Res Ther 2019;21:293.
42 Bottai M, Tijhjnlund A, Santoni G, et al. EULAR/ACR classification criteria for adult and juvenile idiopathic inflammatory myopathies and their major subgroups: a methodology report. RMD Open 2017;3:e000507.
43 Lundberg IE, Tijhjnlund A, Bottai M, et al. 2017 European League against Rheumatism/American College of rheumatology classification criteria for adult and juvenile idiopathic inflammatory myopathies and their major subgroups. Arthritis Rheumatol 2017;69:2271–82.