

EXTENDED REPORT

Development of a consensus core dataset in juvenile dermatomyositis for clinical use to inform research

Liza J McCann, ¹ Clarissa A Pilkington, ^{2,3} Adam M Huber, ⁴ Angelo Ravelli, ⁵ Duncan Appelbe, ⁶ Jamie J Kirkham, ⁶ Paula R Williamson, ⁶ Amita Aggarwal, ⁷ Lisa Christopher-Stine,⁸ Tamas Constantin,⁹ Brian M Feldman,¹⁰ Ingrid Lundberg,¹¹ Sue Maillard,² Pernille Mathiesen,¹² Ruth Murphy,¹³ Lauren M Pachman,^{14,15,16} Ann M Reed, ¹⁷ Lisa G Rider, ¹⁸ Annet van Royen-Kerkof, ¹⁹ Ricardo Russo, ²⁰ Stefan Spinty, ²¹ Lucy R Wedderburn, ^{2,3,22,23} Michael W Beresford ^{1,24}

Handling editor Tore K Kvien

► Additional material is published online only. To view please visit the journal online (http://dx.doi.org/10.1136/ annrheumdis-2017-212141).

For numbered affiliations see end of article.

Correspondence to

Dr Liza J McCann, Department of Paediatric Rheumatology, Alder Hey Children's NHS Foundation Trust, Liverpool L14 5AB, UK: liza.mccann@alderhey.nhs.uk

Received 27 July 2017 Revised 23 September 2017 Accepted 1 October 2017 Published Online First 30 October 2017

ABSTRACT

Objectives This study aimed to develop consensus on an internationally agreed dataset for juvenile dermatomyositis (JDM), designed for clinical use, to enhance collaborative research and allow integration of data between centres.

Methods A prototype dataset was developed through a formal process that included analysing items within existing databases of patients with idiopathic inflammatory myopathies. This template was used to aid a structured multistage consensus process. Exploiting Delphi methodology, two web-based questionnaires were distributed to healthcare professionals caring for patients with JDM identified through email distribution lists of international paediatric rheumatology and myositis research groups. A separate questionnaire was sent to parents of children with JDM and patients with JDM, identified through established research networks and patient support groups. The results of these parallel processes informed a face-to-face nominal group consensus meeting of international myositis experts, tasked with defining the content of the dataset. This developed dataset was tested in routine clinical practice before review and finalisation.

Results A dataset containing 123 items was formulated with an accompanying glossary. Demographic and diagnostic data are contained within form A collected at baseline visit only, disease activity measures are included within form B collected at every visit and disease damage items within form C collected at baseline and annual visits thereafter.

Conclusions Through a robust international process. a consensus dataset for JDM has been formulated that can capture disease activity and damage over time. This dataset can be incorporated into national and international collaborative efforts, including existing clinical research databases.

INTRODUCTION



To cite: McCann LJ, Pilkington CA, Huber AM, et al. Ann Rheum Dis 2018;77:241-250.

Juvenile dermatomyositis (JDM) is associated with significant morbidity and mortality. 1-3 To better understand this rare disease,4 international collaboration is essential. This is feasible with the development of national and international electronic web-based registries and biorepositories.⁵ ⁶ For good clinical care and to aid comparison of data

between groups, it is crucial to have a common dataset that clinicians and researchers collect in a standardised way, with items clearly defined. The International Myositis and Clinical Studies (IMACS) Group^{7–9} and Paediatric Rheumatology International Trials Organisation (PRINTO)^{10–12} IDM core sets were developed predominantly for research studies. Existing myositis registries include partially overlapping but different dataset items, making comparison between groups challenging. 13 This study aimed to define optimal items from existing datasets that would be useful to collect in routine practice, within accessible disease-specific registries, that, when measured over time, would help capture disease outcome/treatment response. which would facilitate both patient care and translational research.

METHODS

The study protocol and background work have been published. 13 14 The study is registered on the Core Outcome Measures in Effectiveness Trials initiative database. 15 The Core Outcome Set—STAndards for Reporting standards for reporting were followed.¹⁶ The study overview is shown in figure 1.

Background work

A steering committee (SC) developed a prototype dataset by scrutinising all items within existing international databases of juvenile-onset myositis (JM) and adult-onset myositis, 1 17-19 informed by a literature search and detailed analysis of the UK Juvenile Dermatomyositis Cohort Biomarker Study and Repository (JDCBS). 13 19 Leading representatives of each partner organisation 9 12 17 20 21 detailed in the study protocol ¹⁴ approved the template/ provisional dataset.

Stakeholder groups

This study design aimed to employ representation from healthcare professionals with experience in myositis working as physicians, allied health professionals or clinical scientists in paediatric or adult medicine within rheumatology, neurology or dermatology¹⁴ and consumers (patients with JM and their parents or carers).



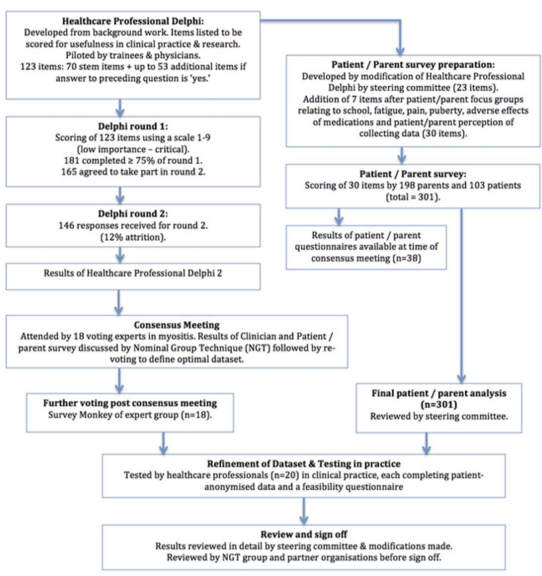


Figure 1 Flow chart showing study overview.

Healthcare professional Delphi process

A two-stage Delphi process was undertaken. 14 Items contained within the prototype dataset were listed and further modified by the SC to ensure clarity. The items were formatted into a custom-made electronic questionnaire, piloted before distribution. After modifications, the Delphi template included 70 items with an additional 53 conditional on previous response (detailed in online supplementary table S1). Participation was invited via membership lists of IMACS, Childhood Arthritis and Rheumatology Research Alliance (CARRA), Juvenile Dermatomyositis Research Group (JDRG) UK and Ireland, Paediatric Rheumatology European Society (PReS) JDM working party and PRINTO Centre Directors. These are representative of international paediatric rheumatology and myositis specialty groups, capturing opinion of clinicians, scientists and allied health professionals. The estimated membership of these groups totals more than 1000. However, the majority of members belong to more than one organisation and membership lists include retired/non-active members or specialists working in adult-onset myositis potentially less inclined to answer a paediatric-specific survey.¹⁴ Participants were asked to rate the importance of each item for clinical practice and separately for value in research,

using a scale of 1–9: 1–3 (of low importance), 4–6 (important but not critical) and 7–9 (critical). An option of 'unable to score' was given and free text comments were allowed. Delphi 2 was sent to participants who scored 75% or more of the items in round 1 of the Delphi. Each participant was asked to re-score each item, having been shown the distribution of scores for the group as a whole and their own score.

Patient and parent survey

The healthcare professionals' survey was modified into separate parent and patient questionnaires as per protocol, ¹⁴ formatted for computer or paper format completion. The questionnaires and age-appropriate information leaflets were reviewed by patient and public involvement coordinators and by parent/ young people's focus groups. ¹⁴ The focus groups also reviewed patient/parent-reported outcome measures (PROMs) used for JDM and other rheumatology conditions, ^{22–27} and opinions were summarised (online supplementary table S2). Thirty items were included in patient/parent questionnaires; 23 from adaptation of the Delphi (combining or simplifying items from the healthcare professional questionnaire and selecting items

Table 1 Definition of consensus for each stage of the study (defined a priori)

Consensus		Definition of consensus				
classification	Description	Healthcare professionals' Delphi	Patient/parent survey	Consensus meeting*		
Consensus in	Consensus that outcome should be included in core set	≥70% of participants scoring '7–9' 'critical for decision-making'	≥70% of participants scoring 'really important'	≥80% of participants voting for inclusion in core outcome set		
Consensus out	Consensus that outcome should not be included in the core outcome set	≥70% of participants scoring '1–3' 'low importance'	≥70% of participants scoring 'not that important'	<80% of participants voting for inclusion in core outcome set		
Equivocal	Uncertainty about importance of outcome	All other responses	All other responses	Further discussion by NGT and revoting allowed		

^{*}More stringent consensus cut-off for consensus meeting. NGT, nominal group technique.

particularly relevant to patients/parents), 2 additional questions added by the SC to determine patient/parent perspectives on collecting and storing information, plus 5 questions suggested by patients/parents within focus groups (online supplementary table S1). The scoring system was simplified into three categories of 'not that important', 'important' and 'really important'. An option of 'unable to score' was given and free text comments were allowed. Participation was open to any patient with JM child/adult, or any parent/carer of a child with IM. Patients with adult-onset myositis (onset ≥18 years) were excluded. Information leaflets and questionnaires were in English only; translators could be used if available. Patients/parents were signposted to the study via email distribution lists/websites of North American and UK patient support groups (Cure JM and Myositis UK), 28 29 the lead of the JDRG patient/parent groups and JDRG coordinator.²⁰ In addition, following site-specific ethics approval, UK centres participating in the JDCBS^{19 30} and a Netherlands site invited patients/parents to participate.

Data analysis

For each item, the number and percentage of participants who scored the item and the distribution of scores (grades 1–9) were summarised for each stakeholder group. Consensus definitions were applied as 'consensus in' versus 'equivocal' or 'consensus out' according to predefined consensus definitions (table 1).

Consensus meeting

Eighteen voting delegates were invited to a 2-day consensus meeting, led by a non-voting facilitator (MWB). International representatives were experts in myositis from paediatric rheumatology/myositis groups and professionals who care for patients with myositis including neurologists, dermatologists, adult rheumatologists and physiotherapists. Prior to the meeting, delegates were sent a summary of results to review. During the consensus meeting, Delphi 2 results and patient/parent results were presented for each item—as shown in online supplementary figure 1. Items achieving 'consensus in' within the Delphi and patient/parent questionnaires were voted on immediately. Those not achieving 'consensus in' were discussed by nominal group technique. Consensus was defined a priori as ≥80% (table 1). Discussion and re-voting allowed refinement of items or associated definitions. The process continued until consensus was reached or until it was clear that consensus would not be reached.

Testing in practice

The proposed dataset was formatted into three sections (forms A, B and C) and tested in clinical practice. Members of the expert group were asked to test the dataset themselves and/ or delegate a member of their department unfamiliar with the

dataset. Clinicians completed patient-anonymised data on one to two patients under their care and a feasibility questionnaire (online supplementary table S3). Feedback was considered by the SC and refinements made. The dataset was sent to the expert group, including representatives of partner organisations (IMACS, CARRA, PRINTO, PReS JDM working group, JDRG, Euromyositis) for comment.

RESULTS

Two hundred and sixty-two healthcare professionals accessed the system (26% of the estimated total membership of specialty groups). 181/262 (69%) completed ≥75% of Delphi 1 (June-September 2014). One hundred and sixty-five agreed to take part in Delphi 2 (November 2014-January 2015); from these, 146 replies were received (12% attrition). One hundred and seventy-two participants provided full demographic data in round 1 showing that survey responses were received from Europe (44%), North America (34%), Latin America (12%), Asia (6%), Australia/Oceania (0.5%), Middle East (3%) and Africa (0.5%). Respondents primarily were paediatric or adult rheumatologists (85%) or had an interest in rheumatology (8%), but also included clinical academics (specialty not defined, 4%), dermatologists (0.5%), neurologists (0.5%), physiotherapists (1%) or other professionals (1%). The majority of respondents had substantial experience in the specialty (74% with ≥10 years of experience) and worked within paediatrics/mainly paediatrics (82.5% vs 17.5% of respondents working with adults). Responses were summarised as percentages of participants ranking items as critical for decision-making (score 7-9) for each item (clinical/ research), shown in online supplementary table S1. Availability of investigations to clinicians within clinical practice was also summarised from responses received in Delphi 1 (online supplementary table S4 and online supplementary figure S1).

Patient/parent surveys

In total, 301 surveys were completed (198 from parents, 103 patients). To allow time for sufficient data capture for parent/patient questionnaires, data collection continued after the consensus meeting. At the consensus meeting, data were available from 16 completed patient surveys and 22 parent surveys. Decisions made at the consensus meeting with 38 responses still held true in the final analysis of 301 replies. Responses were received from Europe (53%), North America (44%) and other continents (3%). Patients completing the questionnaire were a median of 15 years of age (IQR 12–17). Parents completed questionnaires for children who had a median age of 11 years (IQR 7–15). Overall, there was good agreement between patient/parent surveys and the healthcare professionals' Delphi and items agreed at the consensus meeting (online supplementary table S1). Key exceptions are summarised in table 2.

Table 2 Key differences between opinions of patients/parents and healthcare professionals

Item	Patients' opinion	Parents' opinion	Healthcare professionals' opinion	Outcome from consensus meeting	Comments/reasons for retaining in dataset	
Raynaud's phenomenon	Equivocal	Equivocal	Consensus in	Consensus in	Important for overlap phenotypes especially myositis—scleroderma	
Use of an age-appropriate patient/ parent measure of function	Equivocal	Equivocal	Consensus in	Consensus in	Retained (with the option of using alternative tools to allow for country-specific requirements) since	
Use of an age-appropriate patient/ parent measure of quality of life	Equivocal	Equivocal	Consensus in	Consensus in	these are standard outcome measures for research in JDM	
Parent/patient global assessment VAS	Equivocal	Equivocal	Consensus in	Consensus in		
Physician global assessment VAS	Equivocal	Equivocal	Consensus in	Consensus in		
Fatigue due to myositis (within PROM)	Equivocal	Consensus in	Consensus in	Consensus in—as part of a PROM	Quantifiable outcome measure	
Questions related to physiotherapy	Equivocal	Equivocal	Consensus in	Consensus in	Increasingly a defined therapeutic intervention; omitting would be akin to not asking about medicines	
Pubertal assessment	Equivocal	Equivocal	(Not asked)*	Consensus in	Important outcomes of disease activity/damage/	
Height of patient	Equivocal	Consensus in	Consensus in	Consensus in	adverse effects of medication	
Weight of patient	Equivocal	Consensus in	Consensus in	Consensus in		
Items related to major organ involvement—cardiac/pulmonary/gastrointestinal	Equivocal	Consensus in	Consensus in	Consensus in	Important implications for disease severity, treatment and prognosis	
Specific questions about pain	Consensus in	Consensus in	(Not asked)	Consensus out	Thought to be part of standard care (questions that	
Specific questions about medicines	Consensus in	Consensus in	(Not asked)	Consensus out	would be asked by a clinician in a clinic consultation)	
Irritability due to JDM	Equivocal	Consensus in	(Not asked)	Consensus out	Too non-specific and variable interpretation in different countries	

^{*}Added to patient/parent questionnaire after discussion in patient/parent focus groups.

JDM, juvenile dermatomyositis; PROM, patient/parent-reported outcome measure; VAS, Visual Analogue Scale.

Consensus meeting and output

All invited experts (n=18) attended the consensus meeting (Liverpool, March 2015), representing Europe (n=10), North America (n=6), Latin America (n=1) and Asia (n=1). Specialties included paediatric rheumatology (n=13), adult rheumatology (n=2), paediatric dermatology (n=1), paediatric neurology (n=1) and physiotherapy (n=1). Parents/patients were not included. Output from the consensus meeting is shown in online supplementary table S1. A set of recommendations for first visit, for each visit and for annual assessment was made. Refinement took place following the consensus meeting via three rounds of SurveyMonkey, principally to better define myositis overlap features and disease damage items (shown in online

supplementary table S1), with the same members of the expert group (100% response rate).

Testing the dataset in practice

Glossaries of definitions/instructions to aid completion, along with muscle strength-testing sheets, were formulated into appendices, approved by the SC. Twenty clinicians tested the dataset (October 2016–April 2017); eight were present at the consensus meeting, three had completed the Delphi and nine were new to the dataset. Time taken to complete the dataset in clinical practice ranged from 5 to 45 min (median time 15 min).

Table 3 Summary of items included in the JDM optimal dataset, form A (completed at first/baseline visit only)				
Section heading	ltem:	s	Additional items conditional on previous response (summary)	
Personal factors/	1	Date of birth (year and month of birth±day of birth)		
demographics	2	Sex of patient		
Diagnostic factors	3	Date (year and month) of first symptom of myositis		
	4	Date (year and month) of diagnosis of JDM		
	5	At the time of diagnosis did the patient have proximal muscle weakness?		
	6	At the time of diagnosis did the patient have typical skin features of JDM (Gottron's/heliotrope)?		
	7	Was an MRI scan done at diagnosis?	Choice of options for MRI result (four options)	
	8	Was a muscle biopsy done at diagnosis?	Choice of options for biopsy result (four options plus total biopsy score if available)	
	9	Were myositis-specific antibodies tested at diagnosis?	If positive, asked to select all that apply (eight options)	
	10	Were myositis-associated antibodies tested at diagnosis?	If positive, asked to select all that apply (nine options)	
Treatments received prior to	11	Did this patient receive systemic glucocorticoid prior to diagnosis of JDM?	If yes, asked to select all that apply (three options)	
diagnosis of JDM	12	Did this patient receive any synthetic or biologic disease modifying anti- rheumatic drug prior to the diagnosis of JDM?	If yes, asked to select all that apply (13 options)	
JDM, juvenile dermatomyosi	itis.			

 Table 4
 Summary of items included in the JDM optimal dataset, form B (completed at every visit representing status of the patient at the current time point)

Section heading	Item	ıs	Additional items conditional on previous response (summary)
Growth	1	Height of patient (in centimetres)	
	2	Weight of patient (in kilograms)	
Muscular involvement	3	Presence of symmetrical proximal muscle weakness	
	4	Childhood Myositis Assessment Scale score	State score (out of 52)
	5	Manual Muscle Testing score	State score (out of 80)
	6	VAS score for global muscle disease activity	If measured, mark score on 10 cm line*
Skeletal involvement	7	Arthritis due to myositis	
	8	Joint contractures due to myositis	
	9	VAS score for global skeletal disease activity	If measured, mark score on 10 cm line*
Cutaneous involvement	10	Gottron's papules or Gottron's sign	
	11	Heliotrope rash	
	12	Periungual capillary loop changes (plus measure of capillary density if available)	
	13	Malar or facial erythema	
	14	Linear extensor erythema	
	15	'V' sign	
	16	Shawl sign	
	17	Non sun-exposed erythema	
	18	Extensive cutaneous erythema, which may include erythroderma	
	19	Livedo reticularis	
	20	Cutaneous ulceration	
	21	Mucus membrane lesions	
	22	Mechanic's hands	
	23	Cuticular overgrowth	
	24	Subcutaneous oedema	
	25	Panniculitis	
	26	Alopecia (non-scarring)	
	27	Calcinosis (with active disease)	
	28	VAS score for global cutaneous disease activity	If measured, mark score on 10 cm line*
Features suggestive of	29	Does this patient have a myositis overlap condition?	If yes, asked to select all that apply (four option
myositis overlap	30	Raynaud's phenomenon	
	31	Sclerodactyly	
Gastrointestinal involvement	32	Dysphagia due to myositis	
	33	Abdominal pain due to myositis	
	34	Gastrointestinal ulceration due to myositis	
	35	VAS score for global gastrointestinal disease activity	If measured, mark score on 10 cm line*
Pulmonary involvement	36	Pulmonary involvement/respiratory muscle weakness or interstitial lung disease due to myositis	
	37	Dysphonia due to myositis	
	38	VAS score for global pulmonary disease activity	If measured, mark score on 10 cm line*
Cardiovascular involvement	39	Cardiovascular involvement due to myositis	
	40	BP recording	State systolic and diastolic measurement
	41	BP elevated suggesting hypertension (for age of patient)	
	42	VAS score for global cardiovascular disease activity	If measured, mark score on 10 cm line*
Constitutional features	43	Fever (>38°C) due to myositis	
	44	Weight loss (>5%) due to myositis	
	45	Fatigue due to myositis	
	46	VAS score for global constitutional disease activity	If measured, mark score on 10 cm line*
Global disease assessment	47	Physician VAS score of global disease activity	If measured, mark score on 10 cm line*
by clinician	48	Physician VAS score of extramuscular disease activity	If measured, mark score on 10 cm line*
Global disease assessment by patient/parent	49	Patient/parentVAS score for global disease activity	If measured, mark score on 10 cm line* and star who completed (four options)
	50	Patient/parentVAS score for pain	If measured, mark score on 10 cm line*
PROM	51	Use of an age-appropriate PROM of function	Asked to state PROM used and score
	52	Use of an age-appropriate patient/parent-reported measure of quality of life	Asked to state PROM used and score

Continued

Table 4 Continued		
Section heading	Items	Additional items conditional on previous response (summary)
Investigations	53 Elevation of any muscle enzyme (including CPK, LDH, aldolase, AST/SGOT, ALT/SGPT) above normal range	If elevated, asked to select which apply (five options)
Specimens available	Has this patient had specimens taken that may be available for specific research projects? This may include DNA, serum, biomarkers, biopsy tissue or other material	If answer is 'yes', asked to select which apply (three options)
Treatment	55 Is the patient on treatment (now or since last visit)?	Asked to select all that apply (16 options) and to state dose, route and frequency for each medication
	Is the patient doing a regular exercise routine prescribed by a healthcare professional aimed at improving/maintaining:(A) range of movement?(B) muscle strength?	

^{*0} is inactive or lowest score and 10 is most active or highest score on 10 cm VAS scores.

ALT, alanine transaminase; AST, aspartate aminotransferase; BP, blood pressure; CPK, creatine phosphokinase; JDM, juvenile dermatomyositis; LDH, lactate dehydrogenase; PROM, patient/parent-reported outcome measure; SGOT, serum glutamic oxaloacetic Transaminase; SGPT, serum glutamic-pyruvic transaminase; VAS, Visual Analogue Scale.

In addition, 15/20 (75%) found the dataset helpful in practice. Feedback was reviewed in detail by the SC and refinements made.

Completed optimal dataset

The resulting optimal dataset is summarised within tables 3–5 representing three forms. They consist of 123 items: 12 (plus 6 items conditional on responses to the initial 12) within form A, to be completed at first/baseline data entry only; 56 (plus 20 conditional on responses to the 56) within form B, to be completed at every clinic visit representing status of the patient at the current time point; and 55 (plus 15 conditional on responses to the 55) within form C, to be completed at baseline and then annually to capture disease damage. The complete dataset with glossary of definitions and muscle strength-testing sheets can be found in the website of University of Liverpool (http://ctrc.liv.ac.uk/ JDM/) and online supplementary table S5.

DISCUSSION

An internationally agreed JDM dataset has been designed for use within a clinical setting, with the potential to significantly enhance research collaboration and allow effective communication between groups. The accompanying glossary of definitions may be particularly helpful to those in training or physicians less familiar with JDM and for standardisation of the information. Key items are included within the dataset that allow documentation of disease activity and damage with the ability to measure change over time. If adopted widely, the dataset could enable analysis of the largest possible number of patients with JDM to improve disease understanding. It is anticipated that further ratification of the dataset will take place when incorporated into existing registries and national/international collaborative research efforts. It is acknowledged that updates may be needed in the future to incorporate advances in JDM.

When tested in practice by a small number of clinicians, the forms took between 5 and 45 min to complete. The wide range is likely to be due to some respondents interpreting this question as time taken to complete the actual forms, while others may have documented time taken to complete all the tasks within the forms, including clinical examination. It is likely that completion time will be reduced as clinicians become familiar with the questions over time and employment of electronic data entry systems. The dataset does not encompass every aspect of a clinic consultation. Other factors such as adverse effects to medication

or details of pain (ranked important by patients/parents) should be covered as part of standard care.

This study has benefited from the enormous contribution of patients and parents. It is interesting that patients do not necessarily perceive items such as shortness of breath, chest pain and abdominal symptoms as important in JDM whereas for clinicians, major organ involvement has important implications for prognosis and treatment choices. 31-37 Likewise, growth and pubertal parameters were rated less important by patients/ parents but retained due to impact of active disease and corticosteroid treatment on growth. 38 39 Self-assessment is allowable to make pubertal assessment more acceptable to patients.⁴⁰ Notable discrepancies in healthcare professional and patient/ parent opinion included the use of PROMs capturing function and health-related quality of life (HRQOL). The benefits and limitations of individual tools have been described.²² ²⁷ Within this study, comments from patient/parent surveys and focus groups suggested a dislike of 0-10 cm scales used in VAS measurements (data not shown). It is possible that a pain/general VAS is not adequate to capture the complexity of pain or overall feelings for a patient, particularly due to the variability of the disease. Despite this caveat, clinicians recognise the need to have outcome-driven data that include measures of activity, participation, pain and HRQOL.²⁷ Patients with JDM have been found to have significant impairment in their HRQOL compared with healthy peers. 41 PROMs used within the IMACS and PRINTO core sets, including the Childhood Health Assessment Questionnaire and Child Health Questionnaire, are not designed specifically for JDM but have been evaluated and endorsed for use in juvenile myositis.²² The Juvenile Dermatomyositis Multidimensional Assessment Report (JDMAR) is a multifunctional tool that includes function, quality of life, fatigue and adverse effects of medications that has been specifically developed for JDM.²³ It is currently undergoing further validation. Fatigue, rated as important by parents in this work, is included within the JDMAR. During the consensus meeting, it was not possible to define a single agreed PROM for function (activity) or HRQOL (participation) despite taking into consideration results of the healthcare professionals' Delphi, patient/parent surveys and feedback from patients within a UK focus group (online supplementary table S2). The difficulty of PROMs being internationally accepted was discussed and noted. Specifically, items within tools developed in Europe/North America may not be relevant in economically less developed countries. It was agreed that the dataset would include a recommendation to use 'an age-appropriate patient/

 Table 5
 Summary of items included in the JDM optimal dataset, form C (completed at baseline visit and then annual visits only)

Section heading	Items		Additional items conditional on previous response (summary)
Muscular damage items	1	Muscle atrophy (clinical)	
	2	Muscle weakness not attributable to active muscle disease	
	3	Muscle dysfunction: decrease in aerobic exercise capacity	
	4	VAS for global muscle disease damage	Mark score on 10 cm line*
Skeletal damage items	5	Joint contractures (due to myositis)	
	6	Osteoporosis with fracture or vertebral collapse (excluding avascular necrosis)	
	7	Avascular necrosis	
	8	Deforming arthropathy	
	9	VAS for global skeletal disease damage	Mark score on 10 cm line*
Cutaneous damage items	10	Calcinosis (persistent)	
	11	Alopecia (scarring)	
	12	Cutaneous scarring or atrophy (depressed scar or cutaneous atrophy)	
	13	Poikiloderma	
	14	Lipoatrophy/lipodystrophy	
	15	VAS for global cutaneous disease damage	Mark score on 10 cm line*
Gastrointestinal damage items	16	Dysphagia (persistent)	
-	17	Gastrointestinal dysmotility, constipation, diarrhoea or abdominal pain (persistent)	
	18	Infarction or resection of bowel or other gastrointestinal organs	
	19	VAS for global gastrointestinal disease damage	Mark score on 10 cm line*
Pulmonary damage items	20	Dysphonia (persistent)	
,	21	Impaired lung function due to respiratory muscle damage	
	22	Pulmonary fibrosis	
	23	Pulmonary hypertension	
	24	VAS for global pulmonary disease damage	Mark score on 10 cm line*
Cardiovascular damage items	25	Hypertension requiring treatment for >6 months	Mark score on To chi line
caratovascalar admage rems	26	Ventricular dysfunction or cardiomyopathy	
	27	Assessed in adults (>18 years of age) only: angina or coronary artery bypass	
	28	Assessed in adults (>18 years of age) only: myocardial infarction	
	29	VAS for global cardiovascular damage	Mark score on 10 cm line*
Peripheral vascular damage items	30	Tissue or pulp loss	Wark score on To chi line
reliplieral vasculal dalliage itellis	31	Digit loss or limb loss or resection	
	32	Venous or arterial thrombosis with swelling, ulceration or venous stasis	
	33	Assessed in adults (>18 years of age) only: claudication	
	34	VAS for global peripheral vascular disease damage	Mark score on 10 cm line*
Duboutal status of mations			Tanner score (1–5)
Pubertal status of patient	35	Pubertal assessment completed by physician or by patient (self-assessment)	Tanner score (1–5)
Endocrine damage items	36	Growth failure	
	37	Delay in development of secondary sexual characteristics (>2 SD beyond mean for age)	
	38	Hirsutism or hypertrichosis	
	39	Irregular menses	
	40	Primary or secondary amenorrhoea	
	41	Diabetes mellitus	
	42	In adults (>18 years of age): infertility—male or female	
	43	In adults (>18 years of age): sexual dysfunction	
	44	VAS for global endocrine disease damage	Mark score on 10 cm line*
Ocular damage items	45	Cataract resulting in visual loss	
	46	Visual loss, other, not secondary to cataract	
	47	VAS for global ocular disease damage	Mark score on 10 cm line*
Infection damage items	48	Chronic infection	
	49	Multiple infections	
	50	VAS for global infection damage	Mark score on 10 cm line*
Malignancy	51	Presence of malignancy	
	52	VAS for malignancy (complications)	Mark score on 10 cm line*
Other damage	53	Death	Include cause and date of death
	54	VAS for any other damage	Mark score on 10 cm line* and add detail of other damage
Global disease assessment damage	55	Physician VAS of global disease damage	Mark score on 10 cm line*

^{*0} is inactive or lowest score and 10 is most active or highest score on 10 cm VAS scores.

JDM, juvenile dermatomyositis; VAS, Visual Analogue Scale.

parent-reported outcome of function' and 'an age-appropriate patient/parent-reported measure of quality of life'. More work is needed to make PROMs acceptable to patients/parents and applicable to their disease. 42 43

This study is limited by the fact that patient/parent questionnaires were available in English only, reducing the number of countries that could contribute; hence, there is low patient participation outside of Europe and the USA. Complete data from patient/parent surveys were not available at time of the consensus meeting. However, reanalysis of outcomes after the close of the patient/parent survey showed that decisions made at the consensus meeting still held. Initial response rate to Delphi 1 was low (estimated at 26% of potential specialty group membership). However, not all members of the respective organisations contacted would be expected to answer a paediatric-specific survey as described previously. Response rates and attrition between Delphi 1 and 2 were as expected from paediatric rheumatology studies with similar methodology. 44-46 Despite inclusion of neurology and dermatology experts in the consensus meeting, the participants of this study were primarily rheumatologists.

Considerable discussion took place during the consensus meeting regarding the assessment of cutaneous disease in myositis. There are many tools available, 22 but no single tool has been universally accepted. It can be difficult to define skin activity versus damage, particularly without a skin biopsy. After voting on individual skin items and comparing two tools endorsed in JDM, the abbreviated Cutaneous Assessment Tool (aCAT) and Disease Activity Score (DAS) skin score, ²² agreement was reached to use items within the aCAT as disaggregated skin manifestations. These items are recognised to reflect cutaneous lesions associated with disease activity and damage in juvenile and adult myositis.²² Within the item 'periungual capillary loop changes', 'measure of nailfold capillary density if available' was added in recognition of nailfold density relating to prognosis. 47 48 A direct comparison of all available skin tools was outside the remit of this study. Recent published work evaluating the Cutaneous Dermatomyositis Disease Area and Severity Index (CDASI) and the Cutaneous Assessment Tool Binary Method (CAT-BM) in JDM confirms the reliability of both tools when used by paediatric dermatologists or rheumatologists.⁴⁹

The consensus-driven dataset developed in this study, like IMACS and PRINTO core sets, includes physician and patient/ parent global activity, each of which is included in recently defined response criteria for minimal, moderate and major improvement in JDM.8 IMACS measures muscle strength using Manual Muscle Testing, whereas CMAS is used within the PRINTO core set. Both were retained in the consensus dataset. Both tools have been found to have very good inter-rater reliability (when summary scores are used)²² and either is allowed in the recently defined American College of Rheumatology/ European League Against Rheumatism-approved response criteria. The overlap between the IMACS/PRINTO core sets and items contained within the consensus dataset is unsurprising as all core sets aim to capture and measure disease activity and damage over time. A key difference is that the consensus dataset does not use specific tools to record disease activity, such as the Myositis Disease Activity Assessment Tool or the DAS, but rather uses disaggregated items, each of which has been evaluated by a multistage consensus-driven process that considered value for both clinical use and research. The dataset was developed with a key aim for it to be incorporated into existing registries, allowing comparison of data between groups. The already available web-based Euromyositis registry, www.euromyositis.eu, is

free to use in clinical practice and for research and includes a JDM proforma, which will be modified where needed to include items in this new dataset. Likewise, at the time of writing, the CARRA Registry is in the final stages of adding JDM (https:// carragroup.org/) and will include the items contained in this consensus dataset. The JDCBS (https://www.juveniledermatomy ositis.org.uk/) aims to incorporate this dataset as far as possible.

Research priorities defined during the consensus meeting included the need to further develop skin assessment tools that are practical within a busy clinical setting, develop an abbreviated muscle assessment tool that removes redundant items from a combined Childhood Myositis Assessment Scale and Manual Muscle Testing and to further develop PROMs so that they are applicable to JDM and acceptable to patients.

CONCLUSION

Through a robust international consensus process, a consensus dataset for JDM has been formulated that can capture disease activity and damage over time. This dataset can be incorporated into national and international collaborative research efforts, including existing clinical research databases and used routinely while evaluating patients with JDM.

Author affiliations

¹Department of Paediatric Rheumatology, Alder Hey Children's NHS Foundation Trust, Liverpool UK

²Department of Paediatric Rheumatology, Great Ormond Street Hospital for Children NHS Foundation Trust, London, UK

³Arthritis Research UK Centre for Adolescent Rheumatology, University College London, University College London Hospital, London, UK

⁴Division of Pediatric Rheumatology, IWK Health Centre and Dalhousie University, Halifax, Nova Scotia, Canada

⁵Division of Rheumatology, Università degli Studi di Genova and Istituto Giannina Gaslini Pediatria II-Reumatologia, Genoa, Italy

⁶Department of Biostatistics, MRC North West Hub for Trials Methodology Research, University of Liverpool, Liverpool, UK

Department of Clinical Immunology, Sanjay Gandhi Postgraduate Institute of Medical Sciences, Lucknow, India

⁸Johns Hopkins Myositis Center, Johns Hopkins University, Baltimore, Maryland, USA ⁹2nd Department of Paediatrics, Semmelweis University, Budapest, Hungary

¹⁰Department of Pediatrics, The Hospital for Sick Children and University of Toronto, Toronto, Canada

¹¹Department of Medicine, Rheumatology Unit, Karolinska Institutet, Karolinska

University Hospital, Stockholm, Sweden

¹²Department of Paediatrics and Adolescent Medicine, Naestved Hospital, Region Zeeland, Naestved, Denmark

¹³Department of Dermatology, Royal Hallamshire Hospital, Sheffield, UK

¹⁴Department of Pediatrics, Feinberg School of Medicine, Northwestern University, Chicago, Illinois, USA

¹⁵Division of Rheumatology, Ann and Robert H. Lurie Children's Hospital of Chicago, Chicago, Illinois, USA

Centre for Clinical Immunology, The Stanley Manne Children's Research Centre, Chicago, Illinois, USA

⁷Department of Paediatrics, Duke University, Durham, North Carolina, USA

¹⁸Environmental Autoimmunity Group, Clinical Research Branch, National Institute of Environmental Health Sciences, Bethesda, Maryland, USA

¹⁹Department of Paediatric Immunology and Rheumatology, Wilhelmina Children's Hospital University Medical Centre, Utrecht, The Netherlands

²⁰Department of Paediatric Immunology and Rheumatology, Paediatric Hospital Dr. Juan P. Garrahan, Buenos Aires, Argentina

²¹Department of Paediatric Neurology, Alder Hey Children's NHS Foundation Trust, Liverpool, UK

²²Infection, Immunology, and Rheumatology Section, UCL GOS Institute of Child Health, University College London, London, UK

²³NIHR-Biomedical Research Centre at GOSH, London, UK

²⁴Department of Women's and Children's Health, Institute of Translational Medicine, University of Liverpool, Liverpool, UK

Acknowledgements We would like to acknowledge the Euromyositis Research Steering Committee for their collaboration and impetus for this work, in particular Professor Ingrid Lundberg, Dr Hector Chinoy, Professor Jiri Vencovsky and Niels Steen Krogh. We acknowledge the IMACS, CARRA, PRINTO and PReS working groups for their support and collaboration. We would like to acknowledge the IMACS

membership who contributed to this work and the IMACS Scientific Committee for guidance, particularly Dr Rohit Aggarwal, Dr Frederick Miller and Dr Dana Ascherman. We would like to acknowledge the PRINTO Directors who contributed to this work and particularly acknowledge Dr Nicola Ruperto, representing PRINTO. We acknowledge the support of OMERACT and in particular, would like to thank Professor Maarten Boers for his advisory role. We would like to thank the UK Trainees Group and individuals selected by the steering committee for piloting the questionnaire before distribution. We would like to acknowledge our collaborators within the COMET group, in particular, Heather Bagley, Patient and Public Involvement Coordinator. We would also like to acknowledge the help and advice of Professor Bridget Young, Olivia Lloyd and Helen Hanson and the Clinical Studies Group consumer representatives and BSPAR Parent Group, particularly Sharon Douglas. We would like to thank young people from NIHR Young Person's Advisory Group and the JDM Young Person's Group, for their advice on patient questionnaires and information leaflets and Hema Chaplin. Patient and Public Involvement and Engagement Lead for the ARUK Centre for Adolescent Rheumatology for facilitating these groups. We acknowledge the support and collaboration of patient/parent support groups, Cure JM and Myositis UK. In particular, we would like to thank these organisations for promoting the patient and parent questionnaires via their websites/email lists. We would like to thank Katie Arnold, Lawrence Brown, Kath Forrest and Karen Barnes for their administrative support for this work. We would also like to thank Eve Smith and Nic Harman for note keeping during the consensus meeting. We would like to thank the following people for testing the dataset in practice: Bianca Lang, Silvia Rosina, Heinrike Schmeling, Krystyna Ediger, Olcay Jones, Latika Gupta, Maria Martha Katsica, Kiran Nistala, Parichat Khaosut, Ceri Turnbull, Joyce Davidson and Megan Curran.

Collaborators International Myositis Assessment and Clinical Studies group (IMACS), Paediatric Rheumatology INternational Trials Organisation (PRINTO at www.printo.it), Juvenile Dermatomyositis Research Group (JDRG) UK and Ireland, Childhood Arthritis and Rheumatology Research Alliance (CARRA), Paediatric Rheumatology European Society (PReS) JDM working group, Euromyositis Steering Committee, Core Outcome Measures in Effectiveness Trials (COMET) and Outcome Measures in Rheumatology (OMERACT) groups. A full list of contributors to the JDCBS can be found on the JDRG website (http://www.juveniledermatomyositis.org. uk).

Contributors LJM has led all parts of the study including background work, preparation of the protocol, ethics submissions, content of surveys, planning of consensus meeting, testing the dataset and writing the manuscript. CAP, AMH, AR and LRW, as members of the steering committee, have provided intellectual input and practical help into all parts of the study including background work, protocol development, Delphi survey, planning of the consensus meeting, reviewing results, refining the dataset and preparing the manuscript. AMH, AR and CAP also tested the dataset in practice. DA developed the bespoke Delphi system and provided IT support for the study including data analysis. JJK participated in the study design. was responsible for testing the Delphi system, performed the statistical analysis, helped prepare for the consensus meeting, and was involved in reviewing results and preparing the manuscript. PRW has provided expert advice on study design and Delphi methodology and analysis. AA, LC-S, TC, BMF, IL, SM, PM, RM, LMP, AMR, LGR, AvRK, RR and SS attended the consensus meeting and had intellectual input into the study. In addition, AA, AvRK, LMP, LGR and RR tested the dataset in practice. MWB has been responsible for intellectual and financial overview of the study, input into the protocol development and as a member of the steering committee has provided intellectual input into the Delphi survey and consensus meeting, reviewing results, facilitating the consensus meeting, refining the dataset and preparing the

Funding This work was supported by Arthritis Research UK (grant number 20417); January 2014–July 2017. The UK JDM Cohort and Biomarker study has been supported by generous grants from the Wellcome Trust UK (085860), Action Medical Research UK (SP4252), The Myositis Support Group UK, Arthritis Research UK (14518), The Henry Smith Charity. LRW's work is supported in part by Great Ormond Street Children's Charity, the GOSH/ICH NIHR funded Biomedical Research Centre (BRC) and Arthritis Research UK. The JDM Cohort study is adopted onto the Comprehensive Research Network through the Medicines for Children Research Network (www.mcrn.org.uk) and is supported by the GOSH/ICH Biomedical Research Centre. LGR was supported in part by the Intramural Research Program of the National Institute of Environmental Health Sciences National Institutes of Health.

Competing interests IEL has research grants from Astra Zeneca and Bristol Myers Squibb and has served on advisory board for Bristol Myers Squibb.

Patient consent Obtained.

Ethics approval Ethical approval for the study was obtained from the IPHS Research Ethics Committee (reference IPHS-1314-321). The NHS Research Ethics Committee approved final questionnaires, consent/assent forms and patient/parent information leaflets (NRES Committee East Midlands, reference 14/EM/1259; IRAS project ID 160667).

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement All data for this study are held securely within the University of Liverpool. There are no unpublished data openly available. A URL has been established at Liverpool University to hold the published dataset forms as described within the manuscript.

Open Access This is an Open Access article distributed in accordance with the terms of the Creative Commons Attribution (CC BY 4.0) license, which permits others to distribute, remix, adapt and build upon this work, for commercial use, provided the original work is properly cited. See: http://creativecommons.org/licenses/by/4.0/

© Article author(s) (or their employer(s) unless otherwise stated in the text of the article) 2018. All rights reserved. No commercial use is permitted unless otherwise expressly granted.

REFERENCES

- 1 Ravelli A, Trail L, Ferrari C, et al. Long-term outcome and prognostic factors of juvenile dermatomyositis: a multinational, multicenter study of 490 patients. Arthritis Care Res 2010:62:63—72
- 2 Mathiesen PR, Zak M, Herlin T, et al. Clinical features and outcome in a Danish cohort of juvenile dermatomyositis patients. Clin Exp Rheumatol 2010;28:782–9.
- 3 Sanner H, Kirkhus E, Merckoll E, et al. Long-term muscular outcome and predisposing and prognostic factors in juvenile dermatomyositis: a case—control study. Arthritis Care Res 2010:62:1103—11.
- 4 Meyer A, Meyer N, Schaeffer M, et al. Incidence and prevalence of inflammatory myopathies: a systematic review. Rheumatology 2015;54:50–63.
- 5 Lundberg IE, Svensson J. Registries in idiopathic inflammatory myopathies. Curr Opin Rheumatol 2013:25:729–34.
- 6 Rider LG, Dankó K, Miller FW. Myositis registries and biorepositories: powerful tools to advance clinical, epidemiologic and pathogenic research. Curr Opin Rheumatol 2014:26:724–41
- 7 Rider LG, Giannini EH, Harris-Love M, et al. Defining clinical improvement in adult and juvenile myositis. J Rheumatol 2003;30:603–17.
- 8 Rider LG, Aggarwal R, Pistorio A, et al. 2016 American College of Rheumatology/ European League Against Rheumatism Criteria for Minimal, Moderate, and Major Clinical Response in Juvenile Dermatomyositis: an International Myositis Assessment and Clinical Studies Group/Paediatric Rheumatology International Trials Organisation Collaborative Initiative. Ann Rheum Dis 2017;76:782–91.
- 9 National Institutes of Environmental Health and Science. International Myositis Assessment & Clinical Studies Group (IMACS). https://www.niehs.nih.gov/research/ resources/imacs/index.cfm (accessed 8 Apr 2017).
- 10 Ruperto N, Ravelli A, Murray KJ, et al. Preliminary core sets of measures for disease activity and damage assessment in juvenile systemic lupus erythematosus and juvenile dermatomyositis. Rheumatology 2003;42:1452–9.
- 11 Ruperto N, Pistorio A, Ravelli A, et al. The Paediatric Rheumatology International Trials Organisation provisional criteria for the evaluation of response to therapy in juvenile dermatomyositis. Arthritis Care Res 2010;62:1533–41.
- 12 PRINTO. Paediatric Rheumatology INternational Trials Organization. https://www.printo.it/ (accessed 8 Apr 2017).
- 13 McCann LJ, Arnold K, Pilkington CA, et al. Developing a provisional, international minimal dataset for juvenile dermatomyositis: for use in clinical practice to inform research. Pediatr Rheumatol Online J 2014;12:31.
- 14 McCann LJ, Kirkham JJ, Wedderburn LR, et al. Development of an internationally agreed minimal dataset for juvenile dermatomyositis (JDM) for clinical and research use. *Trials* 2015;16:268.
- 15 Core Outcome Measures in Effectiveness Trials Initiative. Development of an internationally agreed minimal dataset for juvenile dermatomyositis (JDM); a rare but severe, potentially life-threatening childhood rheumatological condition. http://www.comet-initiative.org/studies/details/389?result=true (accessed 8 Apr 2017).
- 16 Kirkham JJ, Gorst S, Altman DG, et al. Core outcome set—STAndards for reporting: the COS-STAR statement. PLoS Med 2016;13:e1002148.
- 17 CARRA. Childhood Arthritis and Rheumatology Research Alliance. https://carragroup. org/ (accessed 8 Apr 2017).
- 18 Euromyositis. International collaboration research and treatment database for myositis specialists. https://euromyositis.eu/ (accessed 8 Apr 2017).
- Martin N, Krol P, Smith S, et al. Juvenile Dermatomyositis Research Group. A national registry for juvenile dermatomyositis and other paediatric idiopathic inflammatory myopathies: 10 years' experience; the Juvenile Dermatomyositis National (UK and Ireland) Cohort Biomarker Study and Repository for Idiopathic Inflammatory Myopathies. Rheumatology 2011;50:137–45.
- 20 JDRG. Juvenile Dermatomyositis Research Group. https://www.juveniledermatomyosit is.org.uk/ (accessed 8 Apr 2017).
- 21 PReS. Paediatric Rheumatology European Society. http://www.pres.eu/ (accessed 8 Apr 2017).
- 22 Rider L, Werth V, Huber A, et al. Measures for adult and juvenile dermatomyositis, polymyositis, and inclusion body myositis. Arthritis Care Red 2013;63:118–57.

- 23 Varnier GC, Ferrari C, Consolaro A, et al. PReS-FINAL-2012: introducing a new approach to clinical care of juvenile dermatomyositis: the juvenile dermatomyositis multidimensional assessment report. Pediatric Rheumatol 2013;11:P25.
- 24 Hullmann SE, Ryan JL, Ramsey RR, et al. Measures of general pediatric quality of life: Child Health Questionnaire (CHQ), DISABKIDS Chronic Generic Measure (DCGM), KINDL-R, Pediatric Quality of Life Inventory (PedsQL) 4.0 Generic Core Scales, and Quality of My Life Questionnaire (QoML). Arthritis Care Res 2011;63:S420–30.
- 25 van Agt HM, Essink-Bot ML, Krabbe PF, et al. Test–retest reliability of health state valuations collected with the EuroQol questionnaire. Soc Sci Med 1994;39:1537–44.
- 26 Brazier J, Jones N, Kind P. Testing the validity of the Euroqol and comparing it with the SF-36 health survey questionnaire. Qual Life Res 1993;2:169–80.
- 27 Luca NJ, Feldman BM. Health outcomes of pediatric rheumatic diseases. Best Pract Res Clin Rheumatol 2014;28:331–50.
- 28 Cure JM Foundation. http://www.curejm.org/ (accessed 8 Apr 2017).
- 29 Myositis UK. http://www.myositis.org.uk/ (accessed 8 Apr 2017).
- 30 McCann LJ, Juggins AD, Maillard SM, et al. The Juvenile Dermatomyositis National Registry and Repository (UK and Ireland)—clinical characteristics of children recruited within the first 5 yr. Rheumatology 2006;45:1255–60.
- 31 Pilkington CA, Wedderburn LR. Paediatric idiopathic inflammatory muscle disease: recognition and management. *Drugs* 2005;65:1355–65.
- 32 Ramanan AV Feldman BM Clinical features and outcomes of juvenile dermatomyositis and other childhood onset myositis syndromes *Rheum Dis Clin North Am* 2002 2883357
- 33 Feldman BM, Rider LG, Reed AM, et al. Juvenile dermatomyositis and other idiopathic inflammatory myopathies of childhood. Lancet 2008;371:2201–12.
- 34 Shah M, Mamyrova G, Targoff IN, et al. The clinical phenotypes of the juvenile idiopathic inflammatory myopathies. Medicine 2013;92:25–41.
- 35 Robinson AB, Reed AM. Clinical features, pathogenesis and treatment of juvenile and adult dermatomyositis. Nat Rev Rheumatol 2011;7:664–75.
- 36 Schwartz T, Diederichsen LP, Lundberg IE, et al. Cardiac involvement in adult and juvenile idiopathic inflammatory myopathies. RMD Open 2016;2:e000291.
- 37 Singh S, Suri D, Aulakh R, et al. Mortality in children with juvenile dermatomyositis: two decades of experience from a single tertiary care centre in North India. Clin Rheumatol 2014;33:1675–9.

- 38 Huber AM, Lang B, LeBlanc CM, et al. Medium- and long-term functional outcomes in a multicenter cohort of children with juvenile dermatomyositis. Arthritis Rheum 2000;43:541–9.
- 39 Sanner H, Gran JT, Sjaastad I, et al. Cumulative organ damage and prognostic factors in juvenile dermatomyositis: a cross-sectional study median 16.8 years after symptom onset. Rheumatology 2009;48:1541–7.
- 40 Rasmussen AR, Wohlfahrt-Veje C, Tefre de Renzy-Martin K, et al. Validity of selfassessment of pubertal maturation. Pediatrics 2015;135:86–93.
- 41 Apaz MT, Saad-Magalhães C, Pistorio A, et al. Health-related quality of life of patients with juvenile dermatomyositis: results from the Pediatric Rheumatology International Trials Organisation multinational quality of life cohort study. Arthritis Rheum 2009;61:509–17.
- 42 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;40:1262–73.
- 43 Feldman BM, Grundland B, McCullough L, et al. Distinction of quality of life, health related quality of life, and health status in children referred for rheumatologic care. J Rheumatol 2000;27:226–33.
- 44 Brown VE, Pilkington CA, Feldman BM, et al. An international consensus survey of the diagnostic criteria for juvenile dermatomyositis (JDM). Rheumatology 2006;45:990–3.
- 45 Brunner HI, Mina R, Pilkington C, et al. Preliminary criteria for global flares in childhood-onset systemic lupus erythematosus. Arthritis Care Res 2011;63:1213–23.
- 46 Ruperto N, Hanrahan LM, Alarcón GS, et al. International consensus for a definition of disease flare in lupus. Lupus 2011;20:453–62.
- 47 Christen-Zaech S, Seshadri R, Sundberg J, et al. Persistent association of nailfold capillaroscopy changes and skin involvement over thirty-six months with duration of untreated disease in patients with juvenile dermatomyositis. Arthritis Rheum 2008;58:571–6.
- 48 Schmeling H, Stephens S, Goia C, et al. Nailfold capillary density is importantly associated over time with muscle and skin disease activity in juvenile dermatomyositis. Rheumatology 2011;50:885–93.
- 49 Tiao J, Feng R, Berger EM, et al. Evaluation of the reliability of the cutaneous dermatomyositis disease area and severity index and the cutaneous assessment tool-binary method in juvenile dermatomyositis among paediatric dermatologists, rheumatologists and neurologists. Br J Dermatol 2017.