The Czech version of the Juvenile Arthritis Multidimensional Assessment Report (JAMAR)

Pavla Doležalová1 · Marek Bohm1 · Lenka Linková1 · Melánia Safrídová1 · Dana Němcová1 · Tomáš Dallos2 · Alessandro Consolaro3,4 · Francesca Bovis3 · Nicolino Ruperto3 · For the Paediatric Rheumatology International Trials Organisation (PRINTO)

Received: 22 December 2017 / Accepted: 11 January 2018 © The Author(s) 2018. This article is an open access publication

Abstract
The Juvenile Arthritis Multidimensional Assessment Report (JAMAR) is a new parent/patient-reported outcome measure that enables a thorough assessment of the disease status in children with juvenile idiopathic arthritis (JIA). We report results of the cross-cultural adaptation and validation of the parent and patient versions of the JAMAR in the Czech language. The reading comprehension of the questionnaire was tested in ten JIA parents and patients. The participating centre was asked to collect demographic and clinical data along the JAMAR questionnaire in 100 consecutive JIA patients or all consecutive patients seen in a 6-month period and to administer the JAMAR to 100 healthy children and their parents. The statistical validation phase explored descriptive statistics and the psychometric issues of the JAMAR: the three Likert assumptions, floor/ceiling effects, internal consistency, Cronbach’s alpha, interscale correlations, test–retest reliability, and construct validity (convergent and discriminant validity). A total of 103 JIA patients (5.8% systemic, 35.9% oligoarticular, 37.9% RF-negative polyarthritis, 20.4% other categories) and 100 healthy children, were enrolled. The JAMAR components discriminated well healthy subjects from JIA patients. Notably, there was no significant difference between healthy subjects and their affected peers in the school-related problems variable and in the Psychosocial Health of the Paediatric Rheumatology Quality of Life scale. All JAMAR components revealed good psychometric performances. In conclusion, the Czech version of the JAMAR is a valid tool for the assessment of children with JIA and is suitable for use both in routine clinical practice and clinical research.

Keywords Juvenile idiopathic arthritis · Disease status · Functional ability · Health-related quality of life · JAMAR

Introduction
The aim of the present study was to cross-culturally adapt and validate the Czech parent, child/adult version of the Juvenile Arthritis Multidimensional Assessment Report (JAMAR) [1] in patients with juvenile idiopathic arthritis (JIA). The JAMAR assesses the most relevant parent/patient-reported outcomes in JIA, including overall well-being, functional status, health-related quality of life (HRQoL), pain, morning stiffness, disease activity/status/course, articular and extra-articular involvement, drug-related side effects/compliance and satisfaction with illness outcome.

This project was part of a larger multinational study conducted by the Paediatric Rheumatology International Trials Organisation (PRINTO) [2] aimed to evaluate the...
We report herein results of the cross-cultural adaptation and validation of the parent and patient versions of the JAMAR in the Czech language.

Materials and methods

The methodology employed has been described in detail in the introductory paper of the supplement [4]. In brief, it was a cross-sectional study of JIA children, classified according to the ILAR criteria [5, 6] and enrolled from September 2011 to December 2012. Children were recruited after Ethics Committee approval and consent from at least one parent.

The JAMAR

The JAMAR (1) includes the following 15 sections:

1. Assessment of physical function (PF) using 15 items in which the ability of the child to perform each task is scored as follows: 0 = without difficulty, 1 = with some difficulty, 2 = with much difficulty, 3 = unable to do and not applicable if it was not possible to answer the question or the patient was unable to perform the task due to their young age or to reasons other than JIA. The total PF score ranges from 0 to 45 and has 3 components: PF-lower limbs (PF-LL); PF-hand and wrist (PF-HW) and PF-upper segment (PF-US) each scoring from 0 to 15 [7]. Higher scores indicating higher degree of disability [8–10].

2. Rating of the intensity of the patient’s pain on a 21-numbered circle Visual Analogue Scale (VAS) [11].

3. Assessment of the presence of joint pain or swelling (present/absent for each joint).

4. Assessment of morning stiffness (present/absent).

5. Assessment of extra-articular symptoms (fever and rash) (present/absent).

6. Rating of the level of disease activity on a 21-circle VAS.

7. Rating of disease status at the time of the visit (categorical scale).

8. Rating of disease course from previous visit (categorical scale).

9. Checklist of the medications the patient is taking (list of choices).

10. Checklist of side effects of medications.

11. Report of difficulties with medication administration (list of items).

12. Report of school/university/work problems caused by the disease (list of items).

13. Assessment of HRQoL, through the Physical Health (PhH), and Psychosocial Health (PsH) subscales (five items each) and a total score. The four-point Likert response, referring to the prior month, are ‘never’ (score = 0), ‘sometimes’ (score = 1), ‘most of the time’ (score = 2) and ‘all the time’ (score = 3). A ‘not assessable’ column was included in the parent version of the questionnaire to designate questions that cannot be answered because of developmental immaturity. The total HRQoL score ranges from 0 to 30, with higher scores indicating worse HRQoL. A separate score for PhH and PsH (range 0–15) can be calculated [12–14].

14. Rating of the patient’s overall well-being on a 21-numbered circle VAS.

15. A question about satisfaction with the outcome of the illness (yes/no) [15].

The process of cross-cultural adaptation was conducted according to international guidelines with 2–3 forward and backward translations. In those countries for which the translation of JAMAR had been already cross-cultural adapted in a similar language (i.e. Spanish in South American countries), only the probe technique was performed. Reading comprehension and understanding of the translated questionnaires were tested in a probe sample of ten JIA parents and ten patients.

The participating centre was asked to collect demographic and clinical data along the JAMAR questionnaire in 100 consecutive JIA patients or all consecutive patients seen in a 6-month period and to administer the JAMAR to 100 healthy children and their parents. The statistical validation phase explored the descriptive statistics and the psychometric issues [16]. In particular, we evaluated the following validity components: the first Likert assumption [mean and standard deviation (SD) equivalence]; the second Likert assumption or equal item–scale correlations (Pearson \( r \)); third Likert assumption (item internal consistency or linearity for which each item of a scale should be linearly related to the total score); second Likert assumption (item internal consistency or linearity for which each item of a scale should be linearly related to the total score that is 90% of the items should have Pearson \( r \geq 0.4 \)); floor/ceiling effects (frequency of items at lower and higher extremes of the scales, respectively); internal consistency, measured by the Cronbach’s alpha, interscale correlation.
(the correlation between two scales should be lower than their reliability coefficients, as measured by Cronbach’s alpha); test–retest reliability or intra-class correlation coefficient (reproducibility of the JAMAR repeated after 1 or 2 weeks); and construct validity in its two components: the convergent or external validity which examines the correlation of the JAMAR subscales with the six JIA core set variables, with the addition of the parent assessment of disease activity and pain by the Spearman’s correlation coefficients ($r$) [17] and the discriminant validity, which assesses whether the JAMAR discriminates between the different JIA categories and healthy children [18].

Quantitative data were reported as medians with 1st and 3rd quartiles and categorical data as absolute frequencies and percentages.

The complete Czech parent and patient versions of the JAMAR are available upon request to PRINTO.

Results

Cross-cultural adaptation

The Czech JAMAR was fully cross-culturally adapted from the standard English version with two forward and two backward translations with a concordance for 113/123 translations lines (91.9%) for the parent version and 119/120 lines (99%) for the child version.

In the probe technique analysis, all 123 lines were understood by at least 80% of the parents (median 100%; range 90–100%). All the 120 lines of the patient version of the JAMAR were understood by at least 80% of the children (median 100%; range 100–100%). The texts of the parent JAMAR and of the child JAMAR were unmodified after the probe technique.

Demographic and clinical characteristics of the subjects

A total of 103 JIA patients and 100 healthy children (total of 203 subjects), were enrolled at the paediatric rheumatology centre.

In the 103 JIA subjects, the JIA categories were 5.8% with systemic arthritis, 35.9% with oligoarthritis, 37.9% with RF-negative polyarthritis, 1.0% with RF-positive polyarthritis, 2.9% with psoriatic arthritis, 12.6% with enthesitis-related arthritis and 3.9% with undifferentiated arthritis (Table 1).

A total of 202/203 (99.5%) subjects had the parent version of the JAMAR completed by a parent (103 from parents of JIA patients and 99 from parents of healthy children). The JAMAR was completed by 178/202 (88.1%) mothers and 24/202 (11.9%) fathers. The child version of the JAMAR was completed by 163/203 (80.3%) children aged five or older. Also patients younger than 7 years, capable to assess their personal condition and able to read and write, were asked to fill in the patient version of the questionnaire.

Discriminant validity

The JAMAR results are presented in Table 1, including the scores [median (1st–3rd quartile)] obtained for the PF, the PhH, the PsH subscales and total score of the HRQoL scales. The JAMAR components discriminated well between healthy subjects and JIA patients.

In summary, the JAMAR revealed that JIA patients had a greater level of disability and pain, as well as a lower HRQoL than their healthy peers.

Psychometric issues

The main psychometric properties of both parent and child versions of the JAMAR are reported in Table 2. The following results section refers mainly to the parent’s version findings, unless otherwise specified.

Descriptive statistics (first Likert assumption)

There were no missing results for all JAMAR items, since data were collected through a web-based system that did not allow skipping answers and input of null values. The response pattern for both PF and HRQoL was positively skewed toward normal functional ability and normal HRQoL. All response choices were used for the different HRQoL items except for items 1, 6, 7 and 8, whereas a reduced number of response choices was used for all the PF items except for items 1, 3, 4 and 10.

The mean and SD of the items within a scale were roughly equivalent for the PF and for the HRQoL items, except for HRQoL item 8 (data not shown). The median number of items marked as not applicable was 0% (0–0%) for the PF and 1% (1–3%) for the HRQoL.

Floor and ceiling effect

The median floor effect was 94.2% (88.3–96.1%) for the PF items, 70.9% (52.4–74.8%) for the HRQoL-PhH items, and 71.8% (67.0–75.7%) for the HRQoL-PsH items. The median ceiling effect was 0% (0–1%) for the PF items, 3.9% (1.9–3.9%) for the HRQoL-PhH items, and 0% (0–1%) for the HRQoL-PsH items. The median floor effect was 53.4% for the pain VAS, 47.6% for the disease activity VAS and 44.7% for the well-being VAS. The median ceiling effect was 0% for the pain VAS, 0% for the disease activity VAS and 0% for the well-being VAS.
Table 1: Descriptive statistics (medians, 1st–3rd quartiles or absolute frequencies and %) for the 103 JIA patients

| Category                        | Female (N=6) | Oligoarthritis (N=37) | RF− polyarthrits (N=39) | RF+ polyarthrits (N=1) | Psoriatic arthritis (N=3) | Enthesitis related-arthritis (N=13) | Undifferentiated arthritis (N=4) | All JIA patients (N=103) | Healthy (N=100) |
|--------------------------------|--------------|------------------------|-------------------------|------------------------|---------------------------|-----------------------------------|----------------------------------|--------------------------|------------------|
| **Age at visit**               | 13.3 (11.5–14.3) | 10.3 (7.3–14.4) | 9.6 (6.1–14.9) | 18.9 (18.9–18.9) | 12.9 (5–13.1) | 13.5 (12.3–15.7) | 12.5 (11.6–13.6) | 11.6 (7.5–14.5) | 10.4 (9.1–13.3) |
| **Age at onset**               | 9.8 (8–10) | 3.1 (2–8.2) | 4.6 (2.5–7.6) | 15.7 (15.7–15.7) | 6.4 (3–12.5) | 9 (8.4–10.4) | 8.4 (7.4–9.3) | 5.8 (2.8–8.8)** | 4.5 (2–6.8) |
| **Systemic**                   | (N=6) | 24 (64.9%) | 29 (74.4%) | 1 (100%) | 1 (33.3%) | 2 (15.4%) | 3 (75%) | 61 (59.2%) | 58 (58%) |
| **Oligoarthritis**            | (N=37) | 10.3 | 9.6 (6.1–14.9) | 18.9 (18.9–18.9) | 12.9 (5–13.1) | 13.5 (12.3–15.7) | 12.5 (11.6–13.6) | 11.6 (7.5–14.5) | 10.4 (9.1–13.3) |
| **RF− polyarthrits**          | (N=39) | 3.1 (3.1–3.1) | 2 (0.4–6.7) | 4.3 (3.1–4.9) | 4 (3.2–5.3) | 20.5 (18–23) | 9 (5–15) | 20.5 (18–23) | 9 (5–15) |
| **RF+ polyarthrits**          | (N=1) | – | 10 (6–14) | 8 (7–10) | 10 (6–14) | 8 (7–10) | 20.5 (18–23) | 9 (5–15) | 20.5 (18–23) | 9 (5–15) |
| **Psoriatic arthritis**       | (N=3) | 12.9 (5–13.1) | 13.5 (12.3–15.7) | 12.5 (11.6–13.6) | 11.6 (7.5–14.5) | 10.4 (9.1–13.3) | 10.4 (9.1–13.3) | 10.4 (9.1–13.3) | 10.4 (9.1–13.3) |
| **Enthesitis related-arthritis** | (N=13) | – | 10 (6–14) | 8 (7–10) | 10 (6–14) | 8 (7–10) | 20.5 (18–23) | 9 (5–15) | 20.5 (18–23) | 9 (5–15) |
| **Undifferentiated arthritis** | (N=4) | – | 10 (6–14) | 8 (7–10) | 10 (6–14) | 8 (7–10) | 20.5 (18–23) | 9 (5–15) | 20.5 (18–23) | 9 (5–15) |

Data related to the JAMAR refers to the 103 JIA patients and to the 99 healthy subjects for whom the questionnaire has been completed by the parents.

JAMAR Juvenile Arthritis Multidimensional Assessment Report, ESR erythrocyte sedimentation rate, MD medical doctor, VAS Visual Analogue Scale (score 0–10; 0 = no activity; 10 = maximum activity), LOM limitation of motion, ANA anti-nuclear antibodies, PF physical function (total score ranges from 0 to 45), HRQol health-related quality of life (total score ranges from 0 to 30), PhH physical health (total score ranges from 0 to 15), PsH psychosocial health (total score ranges from 0 to 15)

*p values refer to the comparison of the different JIA categories or to JIA versus healthy. *p < 0.05, **p < 0.001, #p < 0.0001
Equal item–scale correlations (second Likert assumption)

Pearson’s item–scale correlations corrected for overlap were roughly equivalent for items within a scale for 87% of the PF items, with the exception of PF items 14 and 15, for 100% of the HRQoL items and for 100% of the HRQoL items.

Items internal consistency (third Likert assumption)

Pearson’s item–scale correlations were $\geq 0.4$ for 87% of items of the PF (except for PF items 14 and 15) and for 100% of items of the HRQoL.
Cronbach’s alpha internal consistency

Cronbach’s alpha was 0.87 for PF-LL, 0.93 for PF-HW, 0.77 for PF-US. Cronbach’s alpha was 0.91 for HRQoL-PhH and 0.81 for HRQoL-PsH.

Interscale correlation

The Pearson’s correlation of each item of the PF and the HRQoL with all items included in the remaining scales of the questionnaires was lower than the Cronbach’s alpha.

Test–retest reliability

Reliability was assessed in 20 JIA patients, by re-administering both versions (parent and child) of the JAMAR after a median of 2 days (2–2 days). The intraclass correlation coefficients (ICC) for the PF total score showed an almost perfect reproducibility (ICC 0.98). The ICC for the HRQoL-PhH and HRQoL-PsH scores showed an almost perfect reproducibility (ICC 0.89 and 0.86, respectively).

Convergent validity

The Spearman’s correlation of the PF total score with the JIA core set of outcome variables ranged from 0.4 to 0.7 (median 0.5). The PF total score best correlation was observed with the parent assessment of pain ($r=0.7$, $p<0.001$). For the HRQoL, the median correlation of the PsH with the JIA core set of outcome variables ranged from 0.2 to 0.7 (median 0.4), whereas for the PsH ranged from −0.1 to 0.2 (median 0.04). The PhH showed the best correlation with the parent’s assessment of pain ($r=0.7$, $p<0.001$) and the PsH with the parent global assessment of well-being ($r=0.45$, $p<0.001$). The median correlations between the pain VAS, the well-being VAS, and the disease activity VAS and the physician-centered and laboratory measures were 0.4 (0.2–0.5), 0.2 (0.04–0.3), 0.4 (0.2–0.5), respectively.

Discussion

In this study, the Czech version of the JAMAR was cross-culturally adapted from the original standard English version with two forward and two backward translations. According to the results of the validation analysis, the Czech parent and patient versions of the JAMAR possess satisfactory psychometric properties. The disease-specific components of the questionnaire discriminated well between patients with JIA and healthy controls. Notably, there is no significant difference between the healthy subjects and their affected peers in the psychosocial quality of life and school-related problems variables. This finding indicates that children with JIA adapt well to the consequences of JIA.

Psychometric evaluation was good for all domains with few exceptions: two PF items (“bend head back” and “bite a sandwich or an apple”) showing a lower items internal consistency. However the overall internal consistency was good for all the domains. In the external validity evaluation, the Spearman’s correlations of the PF and HRQoL scores with JIA core set parameters ranged from very weak to moderate.

The results obtained for the parent version of the JAMAR are very similar to those obtained for the child version, which suggests that children are equally reliable proxy reporters of their disease and health status as their parents. The JAMAR is aimed to evaluate the side effects of medications and school attendance, which are other dimensions of daily life that were not previously considered by other HRQoL tools. This may provide useful information for intervention and follow-up in health care.

In conclusion, the Czech version of the JAMAR was found to have satisfactory psychometric properties and it is, thus, a reliable and valid tool for the multidimensional assessment of children with JIA.

Acknowledgements We thank all families who participated in the project and the team that prepared and reviewed the forward and backward translations. We thank the staff of the PRINTO International Coordinating Centre in Genoa (Italy) and in particular Marco Garrone for the overall coordination of the translation process, Silvia Scala and Elisa Patrone for data collection and quality assurance, Luca Villa, Giuseppe Silvestri and Mariangela Rinaldi for the database development and management and the remaining PRINTO team for data entry. The principal investigator of the study was Prof. Angelo Ravelli, MD. The scientific coordinator and study methodologist was Nicolina Ruperto, MD, MPH. The project coordinators were Alessandro Consolaro, MD, PhD, Francesca Bovis, BsA. We also thank Prof. Alberto Martini, PRINTO Chairman. Funding was provided by the Istituto G. Gaslini, Genoa (Italy). The local Czech team was supported by the Grant IGA MZ ČR NT14149-3/2013. Permission for use of JAMAR and its translations must be obtained in writing from PRINTO, Genoa, Italy. All JAMAR-related inquiries should be directed to at printo@gaslini.org. Permission for use of CHAQ and CHQ derived-material is granted through the scientific cooperation of the copyright holder ICORE of Woodside CA and HealthActCHQ Inc. of Boston, Massachusetts USA. All CHQ-related inquiries should be directed to licensing@healthactchq.com. All CHAQ-related inquiries should be directed to gsingh@stanford.edu.

Funding This study was funded and coordinated by Istituto Giannina Gaslini, Genoa, Italy.

Compliance with ethical standards

Conflict of interest Dr. Bohm, Dr. Saifridová, Dr. Němcová, Dr. Linková and Dr. Doležalová received Grant no. IGA MZ ČR NT 14149-3/2013 and funding support from Istituto Giannina Gaslini, Genoa, Italy, for the translation phase and data collection performed at their site, within the EPOCA project. Dr. Doležalová reports also personal fees and non-financial support from Pfizer, Abbvie, Roche, MEDAC, personal fees from Novartis, non-financial support from SOBI out-
side the submitted work. Dr. Ruperto has received Grants from BMS, Hoffman-La Roche, Janssen, Novartis, Pfizer, Sobi, during the con-
duct of the study and personal fees and speaker honorarium from Ab-
bvie, Ablynx, Amgen, AstraZeneca, Baxalta Biosimilars, Biogen Idec, 
Boehringer, Bristol Myers Squibb, Celgene, Eli-Lilly, EMD Serono, 
Gilead Sciences, Janssen, Medimmune, Novartis, Pfizer, Rpharm, 
Roche, Sanofi, Servier and Takeda. Dr. Consolaro, Dr. Bovis and Dr. 
Dallos have nothing to disclose.

Ethical approval All procedures performed in studies involving human 
participants were in accordance with the ethical standards of the insti-
tutional and/or national research committee and with the 1964 Helsinki 
Declaration and its later amendments or comparable ethical standards.

Informed consent Informed consent was obtained from all individual 
participants included in the study as per the requirement of the local 
ethical committee.

Open Access This article is distributed under the terms of the Creative 
Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

1. Filocamo G, Consolaro A, Schiappapietra B, Dalpra S, Lattanzi 
B, Magni-Manzoni S et al (2011) A new approach to clinical care of 
juvenile idiopathic arthritis: the Juvenile Arthritis Multidimen-
sional Assessment Report. J Rheumatol 38(5):938–953
2. Ruperto N, Martini A (2011) Networking in paediatrics: the example 
of the Paediatric Rheumatology International Trials Organisa-
tion (PRINTO). Arch Dis Child 96(6):596–601
3. Consolaro A, Ruperto N, Filocamo G, Lanni S, Braccioli G, 
Garrone M et al (2012) Seeking insights into the EPidemiology, 
treatment and Outcome of Childhood Arthritis through a multi-
national collaborative effort: Introduction of the EPOCA study. 
Pediatr Rheumatol Online J 10(1):39
4. Bovis F, Consolaro A, Pistorio A, Garrone M, Scala S, Patrone 
E et al (2018) Cross-cultural adaptation and psychometric evalua-
tion of the Juvenile Arthritis Multidimensional Assessment 
Report (JAMAR) in 54 languages across 52 countries: review of 
the general methodology. Rheumatol Int. https://doi.org/10.1007/ 
s00296-018-3944-1 (in this issue)
5. Petty RE, Southwood TR, Baum J, Bhattay E, Glass DN, Man-
ners P et al (1998) Revision of the proposed classification crite-
rion for juvenile idiopathic arthritis: Durban, 1997. J Rheumatol 
25(10):1991–1994
6. Petty RE, Southwood TR, Manners P, Baum J, Glass DN, Gold-
enberg J et al (2004) International League of Associations for 
Rheumatology classification of juvenile idiopathic arthritis: sec-
ond revision, Edmonton, 2001. J Rheumatol 31(2):390–392
7. Filocamo G, Sztajnbok F, Cespedes-Cruz A, Magni-Manzoni S, 
Pistorio A, Viola S et al (2007) Development and validation of a 
new short and simple measure of physical function for juvenile 
idiopathic arthritis. Arthritis Rheumatol 57(6):913–920
8. Lovell DJ, Howe S, Shear E, Hartner S, McGirr G, Schulte M et al 
(1989) Development of a disability measurement tool for juve-
nue idiopathic arthritis. The juvenile arthritis functional assess-
ment scale. Arthritis Rheumatol 32:1390–1395
9. Howe S, Levinson J, Shear E, Hartner S, McGirr G, Schulte M 
et al (1991) Development of a disability measurement tool for 
juvenile rheumatoid arthritis. The juvenile arthritis functional 
assessment report for children and their parents. Arthritis Rheu-
matol 34:873–880
10. Singh G, Athreya BH, Fries JF, Goldsmith DP (1994) Measure-
ment of health status in children with juvenile rheumatoid arthri-
tis. Arthritis Rheumatol 37:1761–1769
11. Filocamo G, Davi S, Pistorio A, Bertamino M, Ruperto N, Lat-
tanzi B et al (2010) Evaluation of 21-numbered circle and 10-cen-
timeter horizontal line visual analog scales for physician and par-
ent subjective ratings in juvenile idiopathic arthritis. J Rheumatol 
37(7):1534–1541
12. Duffy CM, Arsenault L, Duffy KN, Paquin JD, Strawczynski H 
(1997) The Juvenile Arthritis Quality of Life Questionnaire—
development of a new responsive index for juvenile rheuma-
toid arthritis and juvenile spondyloarthritides. J Rheumatol 
24(4):738–746
13. Varni JW, Seid M, Knight TS, Burwinkle T, Brown J, Szer IS 
(2002) The PedsQL (TM) in pediatric rheumatology—reliability, 
validity, and responsiveness of the pediatric quality of life inven-
tory (TM) generic core scales and rheumatology module. Arthritis 
Rheumatol 46(3):714–725
14. Landgraf JM, Abetz L, Ware JE (1996) The CHQ user’s manual, 
1st edn. The Health Institute, New England Medical Center, 
Boston
15. Filocamo G, Consolaro A, Schiappapietra B, Ruperto N, Pistorio 
A, Solari N et al (2012) Parent and child acceptable symptom state 
in juvenile idiopathic arthritis. J Rheumatol 39(4):856–863
16. Nunnally JC (1978) Psychometric theory, 2nd edn. McGraw-Hill, 
New York
17. Giannini EH, Ruperto N, Ravelli A, Lovell DJ, Felson DT, Mar-
tini A (1997) Preliminary definition of improvement in juvenile 
arthritis. Arthritis Rheumatol 40(7):1202–1209
18. Ware JE Jr, Harris WJ, Gandek B, Rogers BW, Reese PR (1997) 
MAP-R for Windows: multitrait/multi-item analysis program— 
revised user’s guide. Version 1.0 ed. Health Assessment Lab, 
Boston

Affiliations

Pavla Doležalová1 · Marek Bohm1 · Lenka Linková1 · Melánia Saifridová1 · Dana Němcová1 · Tomáš Dallos2 · 
Alessandro Consolaro3,4 · Francesca Bovis3 · Nicolino Ruperto3 · For the Paediatric Rheumatology International 
Trials Organisation (PRINTO)

Marek Bohm 
Bohm.Marek@seznam.cz

Lenka Linková 
linkovalenka@seznam.cz

Melánia Saifridová 
melkal@seznam.cz

Dana Němcová 
nemcova.dana@vfn.cz
