ISSLS prize in clinical science 2022: accelerated disc degeneration after pubertal growth spurt differentiates adults with low back pain from their asymptomatic peers

Anni Aavikko1 · Martina Lohman2 · Leena Ristolainen3 · Hannu Kautiainen4 · Kalevi Österman3 · Dietrich Schlenzka3 · Teija Lund1

Received: 15 October 2021 / Revised: 20 January 2022 / Accepted: 12 March 2022 / Published online: 25 March 2022
© The Author(s) 2022

Abstract
Purpose In this prospective observational cohort study, the development of lumbar intervertebral discs (LIVD) on magnetic resonance imaging (MRI) was investigated from childhood to adulthood with emphasis on the possible association of disc degeneration (DD) to low back pain (LBP).

Methods In 2021, 89 subjects who were enrolled in 1994 in a longitudinal study with lumbar spine MRI at ages 8, 11 and 18 were invited to participate in a long-term follow-up comprising a clinical examination, selected patient-reported outcome measures and a lumbar spine MRI. We assessed all MRIs (three lowest LIVDs) with the Pfirrmann summary score, and the ratio of signal intensity of nucleus pulposus to signal intensity of cerebrospinal fluid (SINDL). We further analyzed whether disc changes at any age were associated with self-reported LBP at age 34.

Results Of the 48 subjects in the follow-up, 35 reported LBP at age 34. The Pfirrmann summary score significantly increased with age (p < 0.001). Subjects reporting LBP at age 34 demonstrated statistically significantly higher summary scores at age 18 and 34 compared to asymptomatic subjects (p = 0.004 at age 18, and p = 0.039 at age 34). SINDL significantly decreased with age (p < 0.001 for all levels separately), but no significant differences between subjects with or without LBP at age 34 were noticed.

Conclusion Subjects with LBP at age 34 had more widespread or severe DD already at age 18 compared to those without LBP.

Keywords Low back pain · Disc degeneration · Long-term · Lumbar spine · MRI

Introduction

Low back pain (LBP) constitutes one of the most common global causes of disability regardless of age [1]. Prevalence of LBP has been reported to reach adult levels by the end of puberty [2], and some studies have identified childhood LBP as a significant risk factor for LBP in adulthood [3].

Despite extensive research, the etiology of LBP remains unknown resulting in unpredictable treatment outcomes. Imaging findings of disc degeneration (DD) become more prevalent with age and are frequently seen in asymptomatic individuals [4]. Findings of DD on magnetic resonance imaging (MRI) in early adulthood seem to predispose the individual for a more rapid progression of degeneration [5]. A systematic review found strong evidence for the association of disc herniation on MRI with progression of DD, while evidence for the extent of disc degeneration or the presence of Modic 1 changes was insufficient [6].

MRI findings such as disc degeneration, disc protrusion, bulge, and extrusion have been associated with LBP [7]; the association of high intensity zones (HIZ) or Modic changes has proven inconsistent [7–9]. Moreover,
the value of MRI findings in predicting future LBP seems low [10]. In a 30-year follow-up study of young males with LBP, degenerated lumbar intervertebral discs (LIVD) in early adulthood demonstrated progressive degeneration over time but did not predict future pain or disability [11].

Our present understanding of the natural history of the LIVD is insufficient to differentiate age-related changes from possible pathologic findings related to LBP. The few long-term studies with MRI published to date lack clinical information [5], report on specific populations [11, 12] or older subjects [13]. In the present study, our objective was twofold: first, to describe the natural history of LIVD from childhood to adulthood; and second, to investigate whether findings of DD were associated with the clinical symptom of LBP.

**Methods**

**Study design and subject recruitment (Fig. 1)**

In 1994, we randomly recruited healthy 8-year-old school children from six primary schools in the urban capital area of Helsinki to investigate the normal growth of the lumbar spine from childhood to early adulthood. The subjects (n = 94 in the beginning of the study) were examined at the age of 8, 11, and 18. In January 2021, an invitation letter for a follow-up examination was sent to all those subjects who were enrolled in 1994, and whose contact details were available from the national population data service agency (n = 89). Subjects who did not respond to the first invitation letter were contacted anew via mail.

All four investigations included a semi-structured interview, a clinical examination and a lumbar spine MRI. The final examination in 2021 additionally included selected patient-reported outcome measures (PROM).

**Semi-structured interview**

The semi-structured interview included the following information: physical demands of work, sports activities and smoking habits; history of LBP without specific trauma (last week/last month/last year/earlier); usage of pain medication; and history of contacting a healthcare provider due to LBP.

**Patient-reported outcome measures**

The participants filled in the following standardized PROMs before the clinical examination: International Physical Activity Questionnaire-Short Form (IPAQ-SF) to assess their physical activity level [14], the EQ-5D-5L for quality of life [15], Oswestry Disability Index (ODI) as a symptom-specific measure [16] and Numeric Rating Scale (NRS, 11-point scale) for pain intensity during the past week including back, neck, arm and leg pain [17].

**Clinical examination**

The subjects’ weight was measured with a balance-beam scale; height was self-reported. Body Mass Index (BMI) was calculated using the standard formula for adults (weight in kilograms divided by the square of height in meters). In addition, a standard clinical examination was performed.

**MRI investigation**

The MRI investigations at the ages of 8 and 11 were performed with a high-field 1.0 T MRI scanner using a dedicated spine coil, while the subsequent investigations were obtained with a high-field 1.5 T MRI scanner. All MRI investigations were performed before 10 am to prevent possible diurnal variations in the signal intensity (SI). Specifically, the MRIs in 2021 were performed using a 1.5 T Canon Vantage Orian scanner (Canon Medical Systems Corporation, Otawara, Japan), a dedicated spine coil and the following four sequences:

- T2 sagittal sequence (TR 4950 ms, TE 120 ms, FOV 300, slice thickness 3.5 mm, slice interval 0.3, acq2).
- T1 sagittal sequence (TR 500 ms, TE 9 ms, FOV 300, slice thickness 3.5 mm, slice interval 0.3, acq1).
- STIR sagittal sequence (TR 4600 ms, TE 50 ms, TI 125, FOV 300, slice thickness 3.5 mm, slice interval 0.3, acq1).
- T2 axial sequence with imaging planes in the direction of the three lowest discs 1 + 1 + 1 (TR 3150 ms, TE 120 ms, FOV 220, slice thickness 3 mm, slice interval 0.3, acq1).

The morphology of the three lowest LIVDs on sagittal T2-weighted images was assessed both qualitatively and quantitatively. For the qualitative visual evaluation, the MRI images including the historical ones were assessed using the Pfirrmann classification [18]. A musculoskeletal radiologist (second author) and a spine surgeon (last author) independently graded the discs without any knowledge of the subject’s clinical characteristics. The inter-rater reliability (IRR, kappa) between the two evaluators for the three lowest intervertebral discs ranged from 0.73 to 0.84 indicating substantial agreement [19]. In case of discrepancy, we used the assessment of the third evaluator (fifth author) for consensus. The individual grades (1–5) of the three LIVDs were added up for a Pfirrmann summary score (range 3–15).
The visual analysis included assessment of any possible HIZ or Modic changes, as well as disc protrusions and herniations.

For the computerized quantitative evaluation of the SI of the three lowest LIVDs, an ellipsoid region of interest (ROI) was digitally marked in midline on T2-weighted sagittal images from each nucleus pulposus. As an internal reference, the SI of the adjacent cerebrospinal fluid (CSF, lat. Liquor cerebrospinalis) was used resulting in a disc to CSF-SI ratio (SINDL) [21].

This manuscript follows the STROBE guidelines for reporting observational studies when applicable.

**Data analysis**

Data are presented as means with standard deviation (SD), median with interquartile range (IQR) or as counts with percentages. Statistical comparison between the groups was performed by t-test, Mann–Whitney test or Chi-square test, when appropriate. In the case of violation of the assumptions (e.g., non-normality) for continuous variables, a bootstrap-type method or Monte Carlo p-values (small number of observations) for categorical variables were used. Repeated measures in Pfirrmann summary score and SINDL between the groups were analyzed using mixed-effects models, with an unstructured covariance structure (Kenward–Roger
method to calculate the degrees of freedom). Fixed effects included group, time and group × time interactions. The repeated measurements were taken at different age, including 8, 11, 18, and 34 years. Mixed models allowed analysis of unbalanced datasets without imputation; therefore, we analyzed all available data with the full analysis set. The normality of variables was evaluated graphically and by using the Shapiro–Wilk W test. The Stata 17.0, StataCorp LP (College Station, TX, USA) statistical package was used for the analysis.

Results

Fifty-one of the 89 invited subjects consented to this follow-up. The final study group comprised 48 subjects (24 males and 24 females). Two subjects only wanted to attend the clinical examination; one additional MRI was cancelled due to pregnancy. Mean age of the subjects at the time of the final investigation was 34.2 years (SD 0.6). LBP without specific trauma was reported by 35 subjects (73%). A statistically significant difference in the use of pain medication (n = 17 vs n = 0, p < 0.001) and the ODI (25.9 vs 23.3, p = 0.045) was seen between subjects with or without LBP, respectively. Table 1 shows more detailed description of the study population.

Fourteen subjects had contacted a healthcare provider due to their LBP symptom. None of the subjects had a history of spine surgery. Of the 19 subjects who reported LBP at the age of 18, 16 still reported low back pain at the age of 34; of the 29 subjects who did not report LBP at the age of 18, 10 remained asymptomatic at the age of 34 with the remaining 19 reporting new-onset LBP.

Pfirrmann summary score and its association with LBP

The Pfirrmann summary score significantly increased with age (p < 0.001) with the mean score increasing by 3.7 (95% CI: 3.3 to 4.1) during the study period from the initial 3.8 (SD 0.8) at the age of 8. A statistically significant difference was seen in the Pfirrmann summary score at the age of 18 and 34 among those subjects who reported LBP at the age of 34 compared to those who did not report LBP at the age of 34 (p = 0.004 at age 18, p = 0.039 at age 34), Fig. 2.

Table 1  Characteristics of 48 subjects according to LBP at the age of 34

|                          | LBP | p value |
|--------------------------|-----|---------|
|                          | No  | Yes     |
|                          | N=13| N=35    |
| Female, n (%)            |     |         |
| 5 (38)                   | 19 (54) | 0.33   |
| Age, mean (SD)           | 34.2 (0.6) | 34.2 (0.6) | 0.71 |
| Smoking                  |     |         |
| Never                    | 10 (77) | 16 (46) | 0.36 |
| Quit                     | 0 (0) | 12 (34) |
| Smoking                  |     |         |
| Never                    | 3 (23) | 7 (20)  |
| BMI, mean (SD)           | 27.0 (8.0) | 26.7 (4.1) | 0.86 |
| Work/physical demand, mean (SD) | 9.4 (4.9) | 11.1 (5.7) | 0.38 |
| Pain medicine, n (%)     |     |         |
| No                       | 13 (100) | 18 (51) | 0.003 |
| Occassionally            | 0 (0) | 14 (40) |
| Regulary                 | 0 (0) | 3 (9)   |
| IPAQ (MET minperWK), median (IRQ) | 3154 (1028, 9375) | 2438 (1158, 5337) | 0.62 |
| EQ-5D-5L, mean (SD)      | 0.938 (0.113) | 0.873 (0.122) | 0.12 |
| EQ-VAS, mean (SD)        | 82 (11) | 79 (15) | 0.53 |
| ODI, mean (SD)           | 23.3 (2.1) | 25.9 (4.2) | 0.045 |
| Pain global, NRS, mean (SD) | 2.00 (2.45) | 2.94 (2.13) | 0.23 |
| Low back                 | 1.25 (1.71) | 1.54 (1.65) | 0.70 |
| Upper back               | 1.50 (2.24) | 0.97 (1.32) | 0.38 |
| Neck                     | 1.08 (2.39) | 1.94 (2.22) | 0.30 |
| Arm                      | 0.83 (1.59) | 0.89 (1.68) | 0.95 |
| Leg                      | 0.50 (1.17) | 1.09 (1.60) | 0.29 |
Visual assessment of disc changes and their association with LBP

Table 2 illustrates the prevalence of Modic changes, HIZ and disc protrusions in our study population at the age of 34. No statistically significant differences were noticed between subjects with or without LBP.

SINDL and its association with LBP

When assessing SINDL for the three lowest LIVDs separately, a statistically significant decrease with age was noticed at all disc levels ($p < 0.001$ for all levels). No significant differences in SINDL emerged at any of the disc levels between subjects with or without LBP at the age of 34, Fig. 3a–c.

Table 2 Prevalence of Modic, HIZ, and disc protrusions in the study population at the age of 34

| LBP   | Modic L3-L5, n (%) | HIZ L3-L5, n (%) | Protrusion L3-L5, n (%) | p value |
|-------|--------------------|------------------|-------------------------|---------|
| No    | $N = 13$           | $N = 35$         |                         |         |
| 1     | 12 (92)            | 26 (74)          |                         | 0.29    |
| 2     | 1 (8)              | 3 (9)            |                         |         |
|       | 0 (0)              | 6 (17)           |                         |         |
| Yes   |                    |                  |                         |         |

Fig. 2 The mean Pfirrmann summary score during the study period for all subjects and according to LBP at the age of 34. Error bars are for 95% confidence intervals

Fig. 3 The mean SINDL for the three lowest LIVDs (a L3/L4, b L4/L5, c L5/S1) separately during the study period for all subjects and according to LBP at the age of 34. Error bars are for 95% confidence intervals
Discussion

Our objective was to describe the natural development of LIVDs (L3/4, L4/5, L5/S1) from childhood to adulthood in a population randomly selected from healthy school children in 1994. We further wanted to investigate whether disc changes on MRI associated with LBP. Our main finding was that subjects who reported LBP at the age of 34 had statistically significantly higher Pfirrmann summary scores already at the age of 18. The difference remained statistically significant at the age of 34, although it was not as remarkable as earlier due to progression of degenerative changes in the asymptomatic subjects alike. In the clinical setting, this finding suggests that either more severe DD in single LIVD or more extensive DD in multiple LIVDs after pubertal growth spurt may be associated with LBP in adulthood.

When the SI of the three lowest LIVDs relative to the SI of the adjacent CSF (SINDL) was calculated separately for each level, no statistically significant differences were noticed between subjects with or without LBP at the age of 34. This may indicate that changes detected in a single LIVD are not as consequential as more widespread DD. The Pfirrmann summary score introduced by Määttä et al. [20] enabled us to analyze the disc morphology of the lower lumbar spine in its entirety for a more comprehensive understanding.

Some evidence suggests that LBP in childhood or adolescence may predict LBP in adulthood [3]. While most of our subjects with LBP at the age of 18 reported continued LBP at the age of 34, many subjects who were asymptomatic at the age of 18 also developed new LBP by the age of 34. Thus, our results do not allow for any conclusions on the predictive value of early-onset LBP per se for LBP symptoms in adulthood.

Sääksjärvi et al. [11] recently published a 30-year follow-up on 26 male subjects who were initially examined by MRI at the age of 20 due to LBP severe enough to exclude them from military service. In their study population, LIVDs with even slightly decreased SI at baseline were more likely to have severely decreased SI at follow-up compared to healthy discs. However, contrary to our results, they did not notice any significant association between the severity of DD at baseline and current pain or disability. Their study population was different from ours in that all their subjects had LBP at baseline which might reflect on their results. Further, although they used a relative measure of disc SI, their reference was the least degenerated lumbar disc with the highest SI; this might have affected their results especially in older subjects. In our study, we used the adjacent CSF which has proven useful as a SI reference [22].

Our study has several limitations. The concept of self-reported LBP should be taken with certain reservations. It is noteworthy that also our asymptomatic population reported some LBP over the past week in NRS. Subjects who reported LBP nevertheless used significantly more pain medication either regularly or occasionally suggesting that their symptoms were more bothersome. The difference between the groups in ODI was statistically significant but did not reach clinical significance. Further, in our inquiry of life-time prevalence of LBP, memory decay might affect the results, although this is less likely with more severe or recurrent symptoms. It is also possible that due to loss of follow-up our study population was biased toward subjects with current or previous history of LBP. Nineteen of our 48 subjects (40%) reported LBP at the age of 18 indicating that majority of subjects of this long-term follow-up did not experience LBP during childhood or adolescence.

For the visual assessment of LIVDs, we used the Pfirrmann classification. The IRR between the two primary evaluators was substantial and comparable to previous literature [23]. The assessment of the third evaluator was used for consensus. The quantitative disc signal measurement (SI) from mid-sagittal MRI has proven a reliable measure of DD [24]. Our study spanned over three decades covering the evolution of MRI technology. The relative measure between the SI of the disc and the adjacent CSF (SINDL) allowed us to compare LIVDs within a single individual and between individuals over time. We only assessed the three lowest LIVDs, as in our young study population, we did not expect significant DD in the upper lumbar spine.

The power of our study is probably not enough to detect some possible associations between disc morphology and LBP, e.g., Modic 1 changes were present only in four subjects. On the other hand, the association between the Pfirrmann summary score at the age of 18 and 34 and LBP at the age of 34 came across as statistically significant. Despite loss to follow-up, to our knowledge, this is the first longitudinal study describing the natural history of LIVDs on MRI from childhood to adulthood with concurrent data on the clinical symptom of LBP.

In conclusion, subjects who reported LBP at the age of 34 presented with more severe or widespread DD already at the age of 18 compared with asymptomatic subjects; the difference was still present at the age of 34. Our finding is clinically important considering the high prevalence of both LBP and MRI findings in adolescents [25], and might offer some cue toward better understanding “normal” age-related versus “pathologic” DD.

Acknowledgements The authors would like to thank all the participants of this long-term follow-up study. The authors gratefully acknowledge the contribution of Tommi Jauhiainen, MD, in performing the MRI examinations.
Author contributions Conceptualisation: Dietrich Schlenzka, Kalevi Österman, Teija Lund. Data curation: Leena Ristolainen. Formal analysis: Martina Lohman, Dietrich Schlenzka, Hannu Kautiainen, Leena Ristolainen, Teija Lund. Funding acquisition: Kalevi Österman, Teija Lund. Investigation: Anni Aavikko. Methodology: Martina Lohman, Dietrich Schlenzka, Teija Lund. Project administration: Leena Ristolainen. Resources: Leena Ristolainen. Writing—original draft: Anni Aavikko. Writing—review and editing: all the authors. All authors read and approved the final manuscript.

Funding Open Access funding provided by University of Helsinki including Helsinki University Central Hospital. This work was supported by Research Institute Orton, Orton Orthopaedic Hospital, Helsinki, Finland, and Helsinki University Hospital, Helsinki, Finland, through grants by the Ministry of Social Affairs and Health in Finland, and the Siviä Kosti Foundation, Helsinki, Finland.

Data availability The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Declarations

Conflicts of interest All authors declare that they have no conflict of interest.

Consent to participate All participants gave their written informed consent prior to their involvement in the study.

Ethics approval This study was conducted according to the Declaration of Helsinki for research on human participants. The study protocol was approved by the Institutional Review Board, and ethical approval was granted by the Ethics Committee of Helsinki University Hospital on December 9, 2020 (protocol code 3251/2020).

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References

1. GBD (2019) Diseases and Injuries Collaborators (2020) Global burden of 369 diseases and injuries in 204 countries and territories, 1990–2019: a systematic analysis for the Global Burden of Disease Study 2019. Lancet 396:1204–1222. https://doi.org/10.1016/S0140-6736(20)30925-9
2. Jeffries LJ, Millanese SF, Grimmer-Somers KA (2007) Epidemiology of adolescent spinal pain: a systematic overview of the research literature. Spine 32:2630–2637. https://doi.org/10.1097/BRS.0b013e318158d70B
3. Hestbaek L, Leboeuf-Yde C, Ohm Kyvik K, Manniche C (2006) The course of low back pain from adolescence to adulthood: eight-year follow-up of 9600 twins. Spine 31:468–472. https://doi.org/10.1097/01.BRS.0000199958.04073.D9
4. Brinjikji W, Luetmer PH, Comstock B, Bresnahan BW, Chen LE, Deyo RA, Halabi S, Turner JA, Avins AL, James K, Wald JT, Kallmes DF, Jarvik JG (2015) Systematic literature review of imaging features of spinal degeneration in asymptomatic populations. Am J Neuroradiol 36:811–816. https://doi.org/10.3174/ajnr.A4173
5. Makino H, Kawaguchi Y, Seki S, Nakano M, Yasuda T, Suzuki K, Ikegawa S, Kimura T (2017) Lumbar disc degeneration progression in young women in their 20’s: a prospective ten-year follow up. J Orthop Sci 22:635–640. https://doi.org/10.1016/j.jos.2017.03.015
6. Rustenburg CME, Faraj SSA, Ket JCF, Emanuel KS, Smit TH (2019) Prognostic factors in the progression of intervertebral disc degeneration: which patients should be targeted with regenerative therapies? JOR Spine 2:e1063. https://doi.org/10.1002/jssp.2.1063
7. Brinjikji W, Diehn FE, Jarvik JG, Carr CM, Kallmes DF, Murad MH, Luetmer PH (2015) MRI findings of disc degeneration are more prevalent in adults with low back pain than in asymptomatic controls: a systematic review and meta-analysis. Am J Neuroradiol 36:2394–2399. https://doi.org/10.3174/ajnr.A4498
8. Teraguchi M, Yim R, Cheung JP-Y, Samartzis D (2018) The association of high-intensity zones on MRI and low back pain: a systematic review. Scoliosis Spinal Disord 13:22. https://doi.org/10.1186/s13013-018-0168-9
9. Herlin C, Kjaer P, Espeland A, Skouen JS, Leboeuf-Yde C, Karpinnen J, Niinimäki J, Solgaard Soerenstor J, Storheim K, Jensen TS (2018) Modic changes: their associations with low back pain and activity limitation—a systematic literature review and meta-analysis. PLoS ONE 13:e0200677. https://doi.org/10.1371/journal.pone.0200677
10. Steffens D, Hancock MJ, Maher CG, Williams C, Jensen TS, Latimer J (2014) Does magnetic resonance imaging predict future low back pain? A systematic review. Eur J Pain 18:755–765. https://doi.org/10.1002/j.1532-2149.2013.00427.x
11. Sääksjärvi S, Kerttula L, Luoma K, Paaajanen H, Waris E (2020) Disc degeneration of young low back pain patients. A prospective 30-year follow-up MRI study. Spine 45:1341–1347. https://doi.org/10.1097/BRS.0000000000003548
12. Kjaer P, Tunset A, Boyle E, Jensen TS (2016) Progression of lumbar disc herniations over an eight-year period in a group of adult Danes from the general population: a longitudinal MRI study using quantitative measures. BMC Musculoskelet Disord 17:26. https://doi.org/10.1186/s12891-016-0865-6
13. Videman T, Battie MC, Gibbons LE, Gill K (2017) A new quantitative measure of disc degeneration. Spine J 17:746–753. https://doi.org/10.1016/j.spine.2017.02.002
14. Lee PH, Macfarlane DJ, Lam TH, Stewart SM (2011) Validity of the international physical activity questionnaire short form (IPAQ-SF): a systematic review. Int J Behav Nutr Phys Act 8:115. https://doi.org/10.1186/1479-5868-8-115
15. EQ-5D User Guides – EQ-5D, https://euroqol.org/publications/user-guides/, Accessed June 9, 2021
16. Pekkanen L, Kautiainen H, Ylilnen J, Säo P, Häkkinen A (2011) Reliability and validity study of the Finnish version 2.0 of the Oswestry disability index. Spine 36:332–338. https://doi.org/10.1097/BRS.0b013e3181cdd702
17. Farrar JT, Young JP Jr, LaMoreaux L, Poole RM (2001) Clinical importance of changes in chronic pain intensity measured on an 11-point numerical pain rating scale. Pain 94:149–158. https://doi.org/10.1016/S0304-3959(01)00349-9
18. Pfirrmann CWA, Metzdorf A, Zanetti M, Hodler J, Boos N (2001) Magnetic resonance classification of lumbar intervertebral disc

© Springer
19. Landis JR, Koch GG (1977) The measurement of observer agreement for categorical data. Biometrics 33:159–174

20. Määttä JH, Karppinen JI, Luk KDK, Cheung KMC, Samartzis D (2015) Phenotype profiling of Modic changes of the lumbar spine and its association with other MRI phenotypes: a large-scale population-based study. Spine J 15:1933–1942. https://doi.org/10.1016/j.spinee.2015.06.056

21. Luoma K, Vehmas T, Riihimäki H, Raininko R (2001) Disc height and signal intensity of the nucleus pulposus on magnetic resonance imaging as indicators of lumbar disc degeneration. Spine 26:680–686. https://doi.org/10.1097/00007632-200103150-00026

22. Luoma EK, Raininko R, Nummi PI, Luukkonen R, Manninen HJ, Riihimäki HA (1997) Suitability of cerebrospinal fluid as a signal-intensity reference on MRI: evaluation of signal-intensity variations in the lumbosacral dural sac. Neuroradiology 39:728–732. https://doi.org/10.1007/s002340050496

23. Urrutia J, Besa P, Campos M, Cikutovic P, Cabezon M, Molina M, Cruz JP (2016) The Pfirrmann classification of lumbar intervertebral disc degeneration: an independent inter- and intra-observer agreement assessment. Eur Spine J 25:2728–2733. https://doi.org/10.1007/s00586-016-4438-z

24. Hu X, Chen M, Pan J, Liang L, Wang Y (2018) Is it appropriate to measure age-related lumbar disc degeneration on the mid-sagittal MR image? A quantitative image study. Eur Spine J 27:1073–1081. https://doi.org/10.1007/s00586-017-5357-3

25. van den Heuvel MM, Oei EHG, Bierma-Zeinstra SMA, van Middelkoop M (2020) The prevalence of abnormalities in the pediatric spine on MRI. Syst Rev Meta-analysis Spine 45:E1185–E1196. https://doi.org/10.1097/BRS.0000000000003527

Publisher’s Note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.