Vertebral Endplate Changes Are Not Associated with Chronic Low Back Pain among Southern European Subjects: A Case Control Study


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**ORIGINAL RESEARCH**

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**BACKGROUND AND PURPOSE:** Data on the association between vertebral endplate changes and low back pain are contradictory. This study was designed to assess whether this association exists among Southern European subjects.

**MATERIALS AND METHODS:** Patients in this study serving as cases were 35–50 years of age with low back pain lasting >90 days, for whom a lumbar MR imaging had been prescribed. Controls were subjects 35–50 years of age, having a cranial MR imaging for headache with normal findings, and no history of clinically relevant LBP. Two hundred forty cases and 64 controls were recruited consecutively in the radiology services across 6 cities in Spain. Imaging findings and subject characteristics were gathered through previously validated instruments. Radiologists who interpreted MRI were blinded to the subject characteristics. A multivariate logistic regression model was developed to assess the association of vertebral endplate changes with LBP, adjusting for sex, age, body mass index, lifetime exposure to smoking, physical activity, disk degeneration, and the interaction between disk degeneration and vertebral endplate changes.

**RESULTS:** Vertebral endplate changes were found in 80.4% of the cases and in 87.5% of the controls. In the regression model, disk degeneration was the only variable showing a confounding effect. Results showed that after adjusting for disk degeneration, the presence of vertebral endplate changes is associated with the absence of chronic LBP (OR for LBP: 0.31; 95% CI, 0.10–0.95).

**CONCLUSIONS:** In Southern European subjects, vertebral endplate changes are not associated with chronic LBP.

**ABBREVIATIONS:** BMI = body mass index; CI = confidence interval; LBP = low back pain; OR = odds ratio; RMQ = Roland-Morris Questionnaire; VAS = visual analog scale

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**L**BP is defined as pain between the costal margins and the inferior gluteal folds, which may be associated with pain referred down to the leg (leg pain) and is usually accompanied by painful limitation of movement. Diagnosing common LBP implies that the pain is not related to conditions such as fractures, direct trauma, or systemic diseases, such as spondyloarthritis or neoplastic, infectious, vascular, metabolic, or endocrine-related processes.

In certain cases of LBP in which pain radiates down the leg and in which signs and symptoms correlate with radiologic findings, disk protrusion, disk herniation, and spinal stenosis are considered to be plausible causes of pain. However, these cases represent <5%, and the rest are considered to be “nonspecific,” implying that no structural cause is triggering the pain. In “nonspecific” cases, pain is thought to originate in soft tissues.

Vertebral endplate changes are defined as changes in the signal intensity of vertebral endplates and subchondral bone. Three different types have been described. Type I (low T1 and high T2 signals) is thought to reflect an ongoing active degenerative process, type II (high T1 and T2 signals) is believed to correspond to a more stable chronic degeneration, and type III (low T1 and T2 signals) is considered to be a sign of subchondral bone sclerosis.

Vertebral endplate changes, also called “Modic” changes, have been observed in MR images of patients with LBP and have therefore been suspected to be a potential cause of “nonspecific” LBP, especially in chronic cases. “Chronic” cases are defined as those in which LBP lasts for ≥12 weeks.

However, vertebral endplate changes have also been observed in MR imaging performed in the general population and asymptomatic subjects. These more likely to be found among men, especially as the subject gets older and signs of disk degeneration appear.

It is currently debatable whether vertebral endplate changes...
Materials and Methods

This was a case control study conducted in 6 hospitals across 6 cities in Spain. This study protocol was approved by the institutional review boards of the participating hospitals and was registered at the ClinicalTrials.gov data base (identifier NCT00479063).

Subjects

All subjects screened for this study were 35–50 years of age and had been prescribed an MR imaging. Patients had been referred for a lumbar MR imaging for LBP lasting ≥90 days. Controls were patients with headache who had been referred for a cranial MR imaging, which turned out to have normal findings, and who either had no history of LBP or had only experienced 1 episode in their life, which had lasted for <7 days.

Exclusion criteria were ethnicity not Spanish, history of spine surgery, currently pregnant, scoliosis with >15° curvature, vertebral fractures, “red flag” for fractures or potential underlying systemic diseases, and signs suggesting cauda equina syndrome. When this study was designed, no data on the prevalence of vertebral endplate changes among Spanish asymptomatic subjects and patients with chronic LBP were available. Therefore, available data from studies conducted in the Northern European and Anglo-Saxon environments were used to calculate sample size.

Sample size was established at 236 cases and 59 controls (total, 295 subjects) on the basis of the following assumptions: prevalence of vertebral endplate changes at 40% among patients with LBP and 19% among controls, α error of .05, statistical power of 90%, and a ratio of cases per controls of 4:1.

Procedure

All the subjects who complied with inclusion criteria and signed the informed consent for participating in the study were included. Neither subjects nor physicians received compensation for participating in this study.

Controls were invited to participate once the recruiting radiologist had established that the cranial MR imaging had normal findings. For controls, participating in the study implied undergoing a lumbar MR imaging. This study did not require any further modifications to patient clinical management.

All the subjects underwent a lumbar MR imaging. Once the images had been obtained, auxiliary personnel provided the subjects with a self-administered questionnaire to gather demographic and clinical data and assigned a code number to each one. At the analysis phase, this code made it possible to match each subject’s clinical data and MR imaging.

Each subject completed a self-administered questionnaire. Once completed, the auxiliary personnel added the code assigned to the subject and sent the questionnaire to a central coordination office. At the coordination office, data were introduced into a data base by 2 administrative assistants separately, who double-checked that the data introduced coincided with the questionnaires.

The auxiliary personnel sent the research coordinator a list indicating the code number assigned to each subject. For ethical reasons and following Spanish law on patient privacy and management of clinical data for research purposes, the research coordinator (F.M.K.) substituted another name on each image by the code assigned to the subject. Once subject identities had been masked on the images, he distributed these among participating radiologists, who read them at their own institutions.

At the beginning of the study, the research coordinator (F.M.K.) had established with the auxiliary personnel a key, within the patient’s code, making it possible to distinguish cases from controls. The key was communicated to the statisticians when they started data analysis. The key remained unknown for the rest of the research staff, including radiologists. The research coordinator did not interpret any images.

This procedure ensured the following: 1) Radiologists were unaware of whether the images they were reading corresponded to a “case” or a “control,” 2) they were blinded to subject demographic and clinical characteristics, 3) the only researchers who were aware of the code identifying each as a “case” or a “control” (the research coordinator and the statisticians) did not read the images, and 4) only the auxiliary personnel and the statisticians who conducted the analysis had access to subject demographic and clinical data.

Clinical Assessment

The self-administered questionnaire gathering demographic and clinical data included age (date of birth), sex, height (centimeters), weight (kilograms), lifetime smoking exposure (pack-years smoked), duration of pain (asking separately whether pain had lasted for ≥90 days and the date of appearance), degree of physical activity, current pain severity, and current degree of disability. Physical activity was assessed through a previously validated questionnaire. Pain intensity was measured with a 10-cm VAS and LBP-related disability, with the validated version of the RMQ. Values, ranging from better to worse, are 0–10 for VAS and 0–24 for RMQ.

Subjects completed all the self-administered questionnaires by themselves, in private. The only instructions they received were those included in the validated instruments measuring physical activity, pain intensity, and disability. They received no help or additional instructions from health care personnel, research staff, or other third parties.

MR Imaging Evaluation

Eleven practicing general radiologists, working in 6 general hospitals located in 5 different geographic regions in Spain, participated in this study. Their postresidency experience as radiologists ranged from 12 to 18 years, and their experience interpreting spine imaging ranged between 7 and 12 years. They were trained in different institutions without fellowships. Their interobserver agreement when reporting
vertebral endplate changes had previously been shown to be better than that for any other radiologic finding.19–21

The methods for MR imaging evaluation used in this study were consistent with those used in studies focusing on radiologist agreement when interpreting lumbar MR imaging and specifically vertebral endplate changes.19–22 All examinations were performed on six 1.5T MR imaging systems with a 6-channel phased array spine coil. The following sequences were used without fat suppression:

- A localizer sequence of 5 images, 30 ms/10 ms/50° (TR/TE/flip angle) consisting of 2 coronal and 3 sagittal images in orthogonal planes
- Sagittal T1-weighted spin-echo, 440–550 ms/14–20 ms (TR/TE), 156–307 × 192–512 matrix, 270-mm FOV, 11 sections of 4-mm thickness, 2 acquisitions
- Sagittal T2-weighted turbo spin-echo 2896–3300 ms/102–120 ms (TR/TE), 156–307 × 192–512 matrix, 270-mm FOV, 11 sections of 4-mm thickness, 2 acquisitions, echo-train length of 12
- Axial T2-weighted turbo spin-echo 2896–3040 ms/103–120 ms (TR/TE), 156–307 × 192–512 matrix, 270-mm FOV, 15 sections of 5-mm thickness, 3 acquisitions, echo-train length of 5. Sections were placed in the plane of the 5 lower disks

Note: TE eff indicates effective echo-time.

Table 1: MR imaging protocol

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Results

Ten radiologists from the 6 participating hospitals screened 64 subjects without LBP and 287 patients with LBP with a lumbar MR imaging. Among the latter, 47 were excluded for pain lasting <90 days. Therefore, 304 subjects were included, 240 cases and 64 controls.

Among the 240 patients with LBP who were included, the median (P25, P75) age was 43 (38–47) years, and 132 (55.0%) were women. Among the 64 subjects without LBP, the median age was 45 (41–47) years, and 29 (45.3%) were women. Table 2 details sample characteristics. There were some missing values, due to questions patients left unanswered in the questionnaires.

The question left unanswered by more subjects was the one requesting the date of appearance of pain; among the 240 patients with chronic LBP, 55 left it unanswered. Because it was hypothesized that differences between patients who did and did not answer this question might exist, their characteristics were compared (On-line Table 1) and a sensitivity analysis was performed, in which these 55 cases were excluded. Characteristics of patients who answered and did not answer this question were very similar (On-line Table 1). Therefore, according to the parsimony principle, only results from the analysis including all the recruited subjects are shown (Table 3).

At least 1 vertebral endplate change (of any type and location) was found in 230 (76%) cases versus 46 (71%) controls. The difference was statistically significant (p = 0.007). When excluding patients with any vertebral endplate change, the association was even stronger (p = 0.001).

Absolute and relative frequencies were used to describe categorical variables. For continuous ones, medians and interquartile range or mean and SD were used, depending on whether data were normally distributed.

The χ² test was used to compare the proportion of subjects showing vertebral endplate changes between cases and controls. To control for potential confounders, we performed a multivariate logistic regression to estimate the association between vertebral endplate changes and chronic LBP. Based on previous studies, the variables, which were included in the maximal model, were age (years), sex, BMI (kilogram per square meter), lifetime exposure to smoking (pack-years smoked), exposure to high-intensity physical activity, severe disk degeneration, and the interaction between disk degeneration and vertebral endplate changes.6,12,15,24

We used a backward strategy; those variables that changed ≥10% of the effect size when they were eliminated from the model were considered as confounders. The collinearity of the maximal models was evaluated by using the criteria proposed by Belsley.25 and the significance level was set at 5%.

The STATA (Version 11.0; StataCorp, College Station, Texas) program was used for analysis.
The results from sensitivity analysis, in which only the 185 patients with LBP (240) were included, were very similar to those excluded (On-line Table 1). Third, differences in results across studies may also derive from differences in sample characteristics, study design, and methodologies.3,7,13,29-31 For instance, results may be influenced by differences in disk degeneration, genetic profile, or other unknown confounders. For instance, in some studies, the exclusion of 55 cases for the sensitivity analysis sufficed for the association of vertebral endplate changes and absence of LBP to cease to be significant, despite the fact that the characteristics of the subjects included were very similar to those excluded (On-line Table 1). Third, differences in results across studies may also derive from differences in sample characteristics, study design, and methodologies.3,7,13,29-31 For instance, results may be influenced by whether there were the following conditions: 1) Asymptomatic subjects were included, 2) imaging reading was blinded to patient clinical characteristics, 3) reporting followed a previously validated nomenclature and protocol, 4) images were applied—eg, age, sex, weight, height, patients who showed any type of vertebral endplate change or severe disk degeneration, and so forth).

Vertebral endplate changes (any type)a,b,c,d 240 (80.4) 64 (87.5) 6

Low back pain severity (VAS) (range) b 229 (4) 52 (0–0)

Highly intense physical activity (median) c 199 (25.6) 48 (9.188)

Only type I 10 (4.2) 1 (1.6)

Types I and II 38 (15.8) 14 (21.9)

Only type II 120 (50.0) 40 (62.5)

Types II and III 11 (4.6) 0 (0.0)

Only type III 1 (0.4) 1 (1.6)

No. of vertebral endplate changes per patienta 240 2 (1, 3) 64 3 (2, 4)

Severe disk degenerationa 240 158 (65.8) 64 30 (46.9)

b Adjusted by disk degeneration.

c No. (%).

d P value for the comparison of the proportion of vertebral endplate changes among subjects without LBP, compared with those with LBP: .191.

LBP. In fact, vertebral endplate changes were more common among controls (Table 2) and were associated with absence of chronic LBP (Table 3).

Several factors may explain the inconsistencies between results from previous studies assessing the association between the existence of vertebral endplate changes and LBP. First, given that genetic differences influence disk degeneration,27,28 the existence of vertebral endplate changes and LBP. First, given that genetic differences influence disk degeneration,27,28 which is the factor most strongly associated with vertebral endplate changes,12 genetic differences may also influence the prevalence and clinical relevance of such vertebral changes and might explain variations across studies conducted with different populations. In fact, the only previous study reporting the prevalence of vertebral endplate changes among Southern European (Spanish) patients with LBP found it to be 81%, with a 95% CI, 77%–85%.12 This prevalence is higher than those reported in other geographic settings,4,6,10,13 but virtually identical to the one found in the current study (Table 2).

Second, if vertebral endplate changes are actually not associated with LBP, as this study suggests, inconsistencies could also be explained by fortuitous differences between the characteristics of each study sample, such as age, disk degeneration, genetic profile, or other unknown confounders. For instance, in this study, the exclusion of 55 cases for the sensitivity analysis sufficed for the association of vertebral endplate changes and absence of LBP to cease to be significant, despite the fact that the characteristics of the subjects included were very similar to those excluded (On-line Table 1). Third, differences in results across studies may also derive from differences in sample characteristics, study design, and methodologies.3,7,13,29-31 For instance, results may be influenced by whether there were the following conditions: 1) Asymptomatic subjects were included, 2) imaging reading was blinded to patient clinical characteristics, 3) reporting followed a previously validated nomenclature and protocol, 4) images were
interpreted by several radiologists, 5) radiologists were blinded to other radiologists’ interpretations, 6) the interobserver agreement of participating radiologists had been previously assessed and shown to be acceptable, and 7) potential confounders were accounted for at the analysis phase.

Type II is the most prevalent vertebral endplate change, and the one that most subjects in this study had, either isolated or in combination with type I. This study was designed to assess the association between LBP and any type of vertebral endplate change. Assessing its association with any particular characteristic (eg, type or affected vertebral volume) would have required a different design, with different inclusion criteria, comparison groups, and sample size. Probably because of the 4:1 ratio between cases and controls, some uncommon combinations were more prevalent among the former (eg, isolated type I or combinations of types I/III, II/III, and I/II/III). However, this possibility is not likely to alter the conclusion of this study because a recent systematic review found no differences in the association between LBP and any particular type of vertebral endplate change.

It is unlikely that a “causal,” “predisposing,” or “triggering” factor will not show an “association” with its corresponding condition. However, because the current study was cross-sectional and had a case control design, it might be argued that it does not rule out the possibility of vertebral endplate changes influencing the prognosis of LBP. This is not likely because among asymptomatic subjects, presenting with a type I vertebral endplate change does not increase the risk of developing LBP; vertebral endplate changes do not influence the clinical course of back pain, and they are not a prognostic factor for recovery. Nevertheless, further prospective studies might assess whether subjects with vertebral endplate changes show different responsiveness to certain specific forms of treatment, such as exercise or some types of surgery.

The characteristics and representativeness of cases and controls are essential in any case control study. Age influences the prevalence of vertebral endplate changes, as well as LBP. Therefore, subjects had to be between 35 and 50 years of age to ensure the comparability of age across controls and cases. In this study, controls were subjects without LBP and without a meaningful history of LBP. For ethical and operational reasons, they were recruited among subjects with normal findings on cranial MR imaging, which had been requested for an unrelated condition (headache). Because the lifetime prevalence of LBP is >70%, at the design phase, it was considered unrealistic to exclude all the subjects reporting having had a single short (<7 days) LBP episode and they were accepted as controls. This does not contradict the conclusion of this study; if vertebral endplate changes were to be associated with a single short episode of LBP in patients’ lifetimes, they could be considered as clinically irrelevant.

“Cases” were not defined as “patients with LBP” but specifically as “chronic patients in whom LBP was serious enough to prescribe a lumbar MR imaging.” These are likely to be patients the clinicians perceive as more serious or those in whom LBP does not improve despite treatment. Chronic patients cause the major clinical, social, and economic burden deriving from LBP, and represent the population in which vertebral endplate changes were more likely to play a causal role. Subjects were recruited consecutively, and all those who complied with inclusion criteria were included. Moreover, the prevalence found among Spanish patients with chronic LBP in this study is very similar to the one from the only previous study conducted in this geographic setting. All these suggest that the generalizability of the results from the current study to middle-aged Spanish patients with chronic LBP is not a major concern.

Results from the current study further support the poor correlation between clinical symptoms and radiologic findings in patients with LBP and the notion that imaging is only appropriate in a specific subset of patients with LBP in whom results may actually change clinical management.

A growing body of evidence questions the effectiveness of spinal fusion and suggests that complex instrumented fusion may cause unnecessary harm to patients. Some surgical societies continue to recommend such a surgery on the basis of the existence of vertebral endplate changes and other radiologic findings. Therefore, unless future high-quality studies show that vertebral endplate changes are indeed clinically relevant, it may be sound to consider changing the way radiologists report findings on lumbar MR imaging, as was the case for plain radiography of the spine, moving away from describing anatomic features to focusing on findings that are more likely to be clinically relevant, such as nerve compression. In the case of vertebral endplate changes, refraining from including this finding in radiologic reports or mentioning it as a finding that is associated with disk degeneration but is likely to be clinically irrelevant may be a way of protecting patients from unnecessarily aggressive forms of treatment or overtreatment.

This study has some weaknesses. Fifty-five (22.9%) of the 240 LBP patients recruited as “cases” did not answer the question on the date when pain appeared. Therefore, although they were chronic (ie, their pain had lasted for >90 days), the exact duration of their pain could not be calculated. It might be argued that these 55 patients are likely to correspond to those who had LBP longer so that it was more difficult for them to recall exactly when the pain appeared. It might also be hypothesized that the 185 patients who provided the date of appearance of pain had an incident to which they attributed pain, even if it was not direct trauma, and did not report retrospective pain. Moreover, the prevalence found among Spanish patients with chronic LBP in this study is very similar to the one from the only previous study conducted in this geographic setting. All these suggest that the generalizability of the results from the current study to middle-aged Spanish patients with chronic LBP is not a major concern.

The reliability of the assessment of lumbar MR imaging is poor for spinal stenosis; fair for osteophytes; and moderate for vertebral endplate changes, Schmorl nodes, disk degeneration, annular tears, and disk contour. However, this limitation is inherent to the use of lumbar MR imaging and does not challenge the clinical applicability of these results.
This study also has some strengths: The definition of “case” corresponded to the type of patients with LBP in whom vertebral endplate changes were more prevalent and more likely to be clinically relevant; cases and controls were recruited in the same setting; the agreement in the reporting of vertebral endplate changes among radiologists interpreting MRI had been shown to be satisfactory by using the methods and imaging protocol implemented in this study; radiologists were blinded to subject characteristics (including whether they were “cases” or “controls”); imaging findings were reported through a previously validated classification; and results were adjusted for potential confounders.

Conclusions

These results suggest that in Spanish subjects, the existence of vertebral endplate changes in the lumbar spine is not associated with chronic low back pain.

References

9.  Kanayama M, Togawa D, Takahashi C, et al. Corresponded to the type of patients with LBP in whom vertebral endplate changes were more prevalent and more likely to be clinically relevant; cases and controls were recruited in the same setting; the agreement in the reporting of vertebral endplate changes among radiologists interpreting MRI had been shown to be satisfactory by using the methods and imaging protocol implemented in this study; radiologists were blinded to subject characteristics (including whether they were “cases” or “controls”); imaging findings were reported through a previously validated classification; and results were adjusted for potential confounders.12-15