Prevalence and clinical features of lumbar zygapophyial joint pain: a study in an Australian population with chronic low back pain

Anthony C Schwarzer, Shih-chang Wang, Nikolai Bogduk, Patrick J McNaught, Rodger Laurent

Abstract

Objectives—To determine the prevalence of pain arising from the zygapophyial joint in patients with chronic low back pain and to determine whether any clinical features could distinguish patients with and without such pain.

Methods—Sixty-three patients with chronic low back pain were studied prospectively. All patients underwent a detailed history and physical examination as well as a series of intra-articular zygapophyial joint injections of 0-5% bupivacaine starting at the symptomatic level to a maximum of three levels or until the pain was abolished. They also received injections of normal saline into paraspinal muscles to act as controls.

Results—All patients proceeded with the injections. Twenty (32%; 95% confidence interval (CI) 20 to 44%) obtained greater than 50% relief of their pain following the administration of saline. Fifty-seven patients completed the study; 23 of them (40%; 95% CI 27 to 53%) failed to obtain relief following the injection of saline but obtained relief following one or more intra-articular injections of local anaesthetic. None of the historical features or clinical tests could discriminate those patients with and those without zygapophyial joint pain.

Conclusion—Pain originating from the zygapophyial joint is not uncommon, but this study failed to find any clinical predictors in patients with such pain.


There has been considerable debate as to the role of the zygapophyial joint in chronic low back pain. Whereas some have considered this joint to be a significant source of pain, others have supported opposite views. Givingey first advanced the notion that this joint could be the source of a set of symptoms and signs in the low back and coined the term ‘facet syndrome’, and early pathomorphological studies supported this idea. Anatomical studies established that the lumbar zygapophyial joints were richly innervated by nociceptive fibres which provided the anatomical substrate for pain from these joints. The landmark study by Mooney and Robertson provided important evidence for the existence of zygapophyial joint pain. By injecting the zygapophyial joints of normal volunteers under fluoroscopic guidance, they were able to induce both back and referred leg pain; then by injecting the same joints with local anaesthetic, they could abolish the pain. Using similar techniques, a number of authors have been able to abolish pain in 8–94% of patients suffering from low back pain. Further studies demonstrated a range of pathological changes which could affect the zygapophyial joint including osteoarthrosis, and even occult fractures.

A major problem has been the inability to diagnose reliably the pain arising from the zygapophyial joint, using either clinical tests or conventional imaging techniques. Two small studies reported sets of clinical signs predictive of zygapophyial joint pain, but their predictive value was not borne out in larger studies. Plain radiographs and computed tomography are capable of showing morphological changes, but these occur as frequently in asymptomatic individuals as in symptomatic individuals.

The only reliable method for making a diagnosis of zygapophyial joint pain is the injection of local anaesthetic into the putatively painful joint or around the nerves which innervate it. These diagnostic blocks are based on the precept that if a structure in the low back is the source of pain, anesthetising that structure should abolish the pain for a period of time commensurate with the duration of action of the local anaesthetic used. If the pain is from a site other than the zygapophyial joint, such a procedure would fail to abolish the pain.

One possible confounder, however, is the false positive response: injection of local anaesthetic may bring about relief of pain not through the pharmacological action of the drug but through some other effect beyond the immediate control of the treating physician. In order to counter this effect, the injection of an inactive substance can be used as a control. If the pain is not relieved by such a control injection but is relieved by the local anaesthetic, this provides strong evidence that the source of pain is indeed the putative joint.

In Australia, patients with back pain are commonly seen by rheumatologists. Of relevance to them is the prevalence of zygapophyial joint pain and how the diagnosis should
be made. Because of the different social and medicolegal context of back pain, prevalence figures from the United Kingdom or North America do not necessarily apply to Australia. There has been one previous Australian study, but neither is nor any other study of lumbar zygapophysial joint pain used placebo controls.

The present study sought to assess the prevalence of pain arising from the lumbar zygapophysial joints and to determine if there are clinical signs that reliably discriminate patients whose pain arises from these joints from patients whose pain arises from some other source. The null hypothesis addressed was that lumbar zygapophysial joint pain is uncommon in an Australian population suffering from low back pain and cannot be predicted clinically using conventional clinical tests.

Patients and methods

Patients

The study population consisted of patients with low back pain referred by rheumatologists at the Royal North Shore Hospital, a major Teaching Hospital in Sydney, between January 1990 and October 1991. The cause of their back pain was not evident from non-invasive diagnostic techniques. Patients were usually referred for the study if they failed to respond to conservative therapy; they were not referred if they were suspected of having anything other than a mechanical cause for their low back pain. Patients were excluded if they were under the age of 18 or over the age of 80 years, if there was a history of previous spinal surgery, if they had a neurological deficit, or if they had a malignancy, spinal infection or inflammatory spinal disorder such as ankyllosing spondylitis. Patients included in the study were restricted to categories 1, 2 or 3 in the classification used by the Quebec Task Force for activity related spinal disorders as recommended by Deyo. None of the patients was receiving 'worker's compensation'. The intention was to study patients with chronic low back pain who had visited a rheumatologist based at a tertiary referral centre. Although such a group represents a small subset of patients in the community with low back pain, these patients are the most difficult to treat and are the most costly to manage.

Seventy one patients were referred, of whom eight failed to proceed: two patients declined to proceed when they were told about the nature of the investigations; two patients were going to leave the country during the investigations and therefore could not proceed; one patient was diagnosed with a carcinoma of the colon and did not commence the study; and three patients experienced too little pain for them to want to proceed with any investigation. Sixty three patients were therefore admitted to the study.

The median age of the population was 59 years (interquartile range 51.0 to 68.0) and the female: male ratio was 3:1. The median duration of low back pain was seven years (interquartile range two to 20) and the median duration of the current episode of low back pain was one year (interquartile range six months to two years). The study was approved by the Ethics Committee at Royal North Shore Hospital and informed consent was obtained from all participants.

Assessments

A detailed history was obtained from all patients and a physical examination was performed by the principal investigator. Aspects considered important were duration and severity of back pain, precipitants, exacerbating and relieving features, site of pain, sites of referred pain, associated symptoms, influence of pain on lifestyle, treatment both past and present, past history, social history and employment history. All patients were requested to document their average level of pain over the previous 24 hours and to rate their average day pain, night pain and pain on movement, using 10 cm visual analogue scales (VAS). The left side of the scales showed 'no pain' and the right side showed 'worst possible pain'; all measurements were in millimetres measured from the left hand margin. Patients were also requested to complete several psychometric questionnaires: the McGill-Melzack Pain Questionnaire (MPQ); the State-Trait Anxiety Inventory; the Beck Depression Inventory (BDI).

The physical examination included a variety of measures that might be used in conventional medical practice in the assessment of patients with low back pain: range of lumbar spinal movement between L1 and S1, noting movements that induce pain; flexion (measured according to the modified Schober's index, using a spirit goniometer positioned 5 cm above the dimples of Venus, and using the extent of distraction over the distance from the level of the C7 spinous process to the level of the iliac crest); extension measured using a spirit goniometer; lateral flexion (measured as finger to floor distance using an erect rule, and also using a spirit goniometer held against the arm of patients to the side of lateral flexion); sites of spinal and paraspinal tenderness; extent of straight leg raising using a spirit goniometer held against the lower thigh just above the patella; combinations of rotation and extension with the patient standing and assessing whether these manoeuvres are painful; a complete rheumatological and neurological examination. In addition, blood was taken for a full blood count, erythrocyte sedimentation rate and biochemical profile.

A study was performed to assess intraobserver and interobserver agreement using the following tests: modified Schober's index; distraction of C7 to iliac crest distance with forward flexion; forward flexion using a spirit goniometer; extension using a spirit goniometer; lateral flexion to the right and left using finger to floor distance and spirit goniometer; extent of straight leg raising using a spirit goniometer; rotation to the right combined with extension to the left and
rotation to the left combined with extension to the right assessing whether pain was induced.

The examinations were performed by the principal investigator and a co-investigator (RL) on 10 patients. Each patient was assessed by each examiner at times 0, 15 minutes, and 30 minutes.

INJECTIONS OF ZYGAPOPHYSIAL JOINTS

The criterion standard adopted for the diagnosis of zygapophysial joint pain was the response to local anaesthetic blocks of these joints. The procedure was performed using an image intensifier. Under single blind conditions, patients underwent a placebo injection followed by a series of intra-articular zygapophysial joint injections. The placebo injection was always the first injection; subsequently, on separate occasions, intra-articular injections were performed at L5–S1, L4–5, and L3–4, in that order. The injections were one week apart and the effect of each procedure was assessed for eight hours. Injections were performed sequentially until the patient became pain free or until all three levels had been injected. If a patient obtained less than 50% relief of pain at one level, that joint was not reinjected but the next joint was injected on the subsequent occasion. When it was suspected that the pain may have arisen from a level higher than L3–4, higher joints were injected; this was the case in three patients. Although patients consented to undergo placebo injections as part of the study, the order of injections was not specified and, therefore, patients were unaware of when they were receiving placebo and active injections.

Placebo injections were performed first in order to avoid any carry over effect from injection of an active agent and in order to avoid conditioning. The literature indicates that placebos become more effective when there is prior conditioning with non-placebos. Performing the placebo injection first avoids this risk of amplifying the placebo response rate.

Before each and every procedure, up to 5 ml of 1% lignocaine (Xylocaine 1%<sup>R</sup>, Astra) was infiltrated subcutaneously to provide adequate surface anaesthesia. Injections were either unilateral or bilateral, depending on whether the pain was unilateral or bilateral. Both placebo and active injections were performed with a 25 gauge, 9 cm spinal needle (Spinocan<sup>R</sup>, B Braun Melsungen AG, Germany). The placebo injections consisted of 0·5 ml of normal saline injected into muscle superficial to the zygapophysial joints in the region of the patients' pain and not deeper than 40 mm from the skin. Placebo injections were performed in this way in order not to interfere with the zygapophysial joint. Active injections consisted of 0·1–0·3 ml of contrast medium (meglumine iothalamate, Conray 280<sup>R</sup>, May and Baker) to confirm intra-articular placement (figure), followed by 1·5 ml of 0·5% bupivacaine hydrochloride (Marcain<sup>R</sup>, Astra). Smaller volumes were used when joints would not accept the full 1·5 ml.

**Assessment of Pain Relief**

Patients were assessed after both placebo and active injections using a series of 10 cm VASs which were completed by the patient before the procedure, 30 minutes after the procedure and then hourly for eight hours. Following each injection patients were instructed to carry on with their usual daily activities in addition to completing their VAS. Two major types of response were recognised: positive or negative. A positive response to placebo or intra-articular injections was defined as a 50% or greater reduction in pain maintained for at least three hours, consistent with the expected duration of action of bupivacaine. Amongst patients who exhibited a positive response, patients were identified in whom there was greater than 90% relief of pain or complete relief of pain for a duration of at least three hours. Patients exhibiting positive responses to the placebo injection subsequently also underwent intra-articular injections of bupivacaine if and once their usual pain returned.

The diagnosis of zygapophysial joint pain was made if the patient exhibited a positive response to zygapophysial joint injections at one or more levels but failed to respond positively to the placebo control injection.

**Statistical Analysis**

All results of history, examination and investigations were recorded on an SPSS database<sup>60</sup> using an IBM personal computer.

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Statistical analysis was performed using BMDFit and the SAS statistical program. For interobserver agreement, the kappa statistic was used for categorical variables and the intraclass correlation coefficient was used for continuous variables. The intraclass correlation coefficient was calculated using estimates of mean squares from a one way analysis of variance and p values were based on the F test.

The information obtained on history, physical examination and psychometric assessment was compared in patients with a diagnosis of zygapophysial joint pain and those who did not have zygapophysial joint pain. The normality of quantitative variables was assessed using the Shapiro-Wilk test. As quantitative values followed a non-parametric distribution, the Wilcoxon rank sum test was used to compare the median scores for each parameter in responders and non-responders to zygapophysial joint injections. The results were expressed as the means of the rank sums in each group, with p values. When examining the association between two discrete variables, the Pearson Chi Squared test was used. The Bonferoni method was used to take account of multiple tests. A result was considered significant if p was less than 0.01.

Results
In all, 316 injections were performed at 218 levels. Injections were performed bilaterally at 98 segmental levels (45%), while 67 injections (31%) were right sided and 53 (24%) were left sided. Sixty three patients underwent placebo injections and intra-articular injections were performed at the following levels and at the following frequencies: 57 at L5–S1, 49 at L4–5, 46 at L3–4, two at L2–3 and one at L1–2.

Fifty seven patients completed the study. Five patients failed to proceed to the intra-articular injections and one patient left the study after receiving the first injection. Of the five patients who failed to proceed, four were free of pain for a prolonged period after the placebo injection and therefore did not require further injections, and one developed a generalised skin rash, which was unrelated to any of the procedures. By the time the rash resolved, the patient’s pain had largely subsided and therefore blocks of the zygapophysial joints were no longer required. The patient who left the study experienced features of L5 motor and sensory loss after injection of the L5–S1 zygapophysial joint. This was believed to be caused by extracapsular spread of local anaesthetic and the neurological symptoms resolved completely after six hours. The analgesic response could not be assessed and it was recorded as a failed block. The patient subsequently withdrew from the study. No other patient suffered anaesthesia to a spinal nerve.

Twenty of the 63 patients (32%; 95% confidence interval (CI) 20 to 44%) who underwent placebo injections obtained a greater than 50% reduction in pain maintained for at least three hours. A diagnosis of zygapophysial joint pain was made in 23 of 57 patients (40%; 95% CI 27 to 53%). When those patients who could not complete the study were included in the analysis a diagnosis of zygapophysial joint pain could be made in 37% (95% CI 25 to 49%). Eighteen patients obtained 90% or greater relief of pain; seven of these obtained complete relief of pain. If a positive response to injection is defined as abolition of 90% of the original pain, the prevalence of zygapophysial joint pain was 32% (95% CI 20 to 44%); if a positive response to injection is defined as total abolition of pain, this figure reduces to 12% (95% CI 3 to 21%) of those who completed the study or 11% (95% CI 3 to 19%) of those admitted to the study.

Eighteen patients obtained relief at only one level. Five other patients obtained relief at more than one level after single injections on separate occasions; three had relief at two levels and two at all three levels. When the joints considered as sources of pain were those in which injection gave the greatest relief for each patient, L5–S1 (eight) and L4–5 (eight) were the most common levels, followed by L3–4 (six) and L2–3 (one).

Assessment of interobserver agreement revealed that measurements of forward flexion were reliable (intraclass correlation coefficients of 0.81–0.94 for the Schober’s test and 0.73–0.93 using the inclinometric method), but measurements of extension were variably reliable (0.18–0.83). For lateral flexion, the inclinometric measurements performed slightly better than measurements using the vertical rule. There was greater interobserver agreement for range of lateral flexion than finger to floor distance. Intraclass correlation coefficients ranged from 0.80 to 0.98 for the inclinometric method and from 0.73 to 0.86 when range of movement in centimetres was considered. There was moderate interobserver agreement using the inclinometric method for straight leg raising; intraclass correlation coefficients ranged from 0.63 to 0.86. For rotation combined with extension, kappa scores ranged from 0.35 to 0.80, indicating fair to substantial agreement. Intraobserver agreement was best for tests of forward flexion and lateral flexion and least reliable for tests that measured extension.

There was no statistically significant difference in the demographic features, levels of pain and findings on history and physical examination between patients with and without pain originating from the zygapophysial joint (tables 1–4). Levels of anxiety and depression were not significantly different.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Comparison of demographic features in patients with and without a diagnosis of zygapophysial joint pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feature</td>
<td>Z joint positive</td>
</tr>
<tr>
<td>---------</td>
<td>------------------</td>
</tr>
<tr>
<td>Age (years)</td>
<td>61</td>
</tr>
<tr>
<td>Duration of low back pain (months)</td>
<td>132</td>
</tr>
<tr>
<td>Duration of current episode of low back pain (months)</td>
<td>9</td>
</tr>
</tbody>
</table>

*Derived from Wilcoxon rank sum test.

Z joint = zygapophysial joint; median values given.
Table 2  Level of pain: comparison between patients with and without zygapophysial joint pain

<table>
<thead>
<tr>
<th>Feature</th>
<th>Z joint positive</th>
<th>Z joint negative</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>VAS (mm)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Day pain</td>
<td>47</td>
<td>22</td>
<td>0.47</td>
</tr>
<tr>
<td>Night pain</td>
<td>22</td>
<td>22</td>
<td>0.98</td>
</tr>
<tr>
<td>Pain on movement</td>
<td>50</td>
<td>70</td>
<td>0.23</td>
</tr>
<tr>
<td>PPI</td>
<td>2</td>
<td>2</td>
<td>0.40</td>
</tr>
<tr>
<td>PRI (T)</td>
<td>21</td>
<td>21</td>
<td>0.84</td>
</tr>
</tbody>
</table>

Wilcoxon rank sum test. \( Z \) joint = zygapophysial joint; median values. VAS = visual analogue scale; PPI = present pain intensity, derived from the McGill Pain Questionnaire; PRI (T) = total of the rank values of the words in the McGill Pain Questionnaire.

Table 3  Comparison of examination features between patients with and without zygapophysial joint pain

<table>
<thead>
<tr>
<th>Feature</th>
<th>Z joint positive</th>
<th>Z joint negative</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mod. Schober’s index (cm)</td>
<td>5 1</td>
<td>5 15</td>
<td>0.83</td>
</tr>
<tr>
<td>Flexion (deg)</td>
<td>93</td>
<td>84-5</td>
<td>0.42</td>
</tr>
<tr>
<td>Extension (deg)</td>
<td>17</td>
<td>19-5</td>
<td>0.47</td>
</tr>
<tr>
<td>Lateral flexion (cm)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Left</td>
<td>15-7</td>
<td>14-0</td>
<td>0.64</td>
</tr>
<tr>
<td>Right</td>
<td>14-7</td>
<td>15-5</td>
<td>0.64</td>
</tr>
<tr>
<td>SLR right (deg)</td>
<td>93</td>
<td>85</td>
<td>0.38</td>
</tr>
<tr>
<td>SLR left (deg)</td>
<td>90</td>
<td>84</td>
<td>0.40</td>
</tr>
</tbody>
</table>

Wilcoxon rank sum test. SLR = Straight leg raising; \( Z \) joint = zygapophysial joint; median values.

Table 4  Comparison of history and examination features between patients with and without zygapophysial joint pain

<table>
<thead>
<tr>
<th>Historical or examination feature</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leg pain</td>
<td>0.10</td>
</tr>
<tr>
<td>Dysaesthesia in legs</td>
<td>0.48</td>
</tr>
<tr>
<td>Pain on flexion</td>
<td>0.78</td>
</tr>
<tr>
<td>Pain on extension</td>
<td>0.46</td>
</tr>
<tr>
<td>Pain on right rotation and left extension</td>
<td>0.42</td>
</tr>
<tr>
<td>Pain on left rotation and right extension</td>
<td>0.39</td>
</tr>
</tbody>
</table>

\( \dagger \) \( t \) test.

Table 5  Comparison of scores for the Beck Depression Index (BDI) and the State-Trait Anxiety Index (STAI) in patients with zygapophysial joint pain and patients with other sources of pain

<table>
<thead>
<tr>
<th>Result of psychometric test</th>
<th>Z joint positive</th>
<th>Z joint negative</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>BDI</td>
<td>8</td>
<td>8</td>
<td>0.92</td>
</tr>
<tr>
<td>STAI (state)</td>
<td>44</td>
<td>44-3</td>
<td>0.88</td>
</tr>
<tr>
<td>STAI (trait)</td>
<td>46</td>
<td>45-5</td>
<td>0.87</td>
</tr>
</tbody>
</table>

Wilcoxon rank sum test. \( Z \) joint = zygapophysial joint; median values.

The diagnosis of zygapophysial joint pain. Failure to consider the placebo response may be one reason why several previous estimates of the prevalence of zygapophysial joint pain were so high.\( ^{1-6} \)

The present study addressed the issue of a single zygapophysial joint being the main source of unilateral low back pain or, in the case of bilateral pain, a pair of joints. It expressly did not study the simultaneous contribution of joints from more than one level. It may well be that the prevalence estimate of 40% for zygapophysial joint pain was an underestimate and, had several levels been injected on one occasion, more patients might have been granted the diagnosis. The contribution of multiple joints could form the subject of future studies. Interestingly, five of the 23 patients with a diagnosis of zygapophysial joint pain experienced a greater than 50% diminution in pain after the injection of different joints on separate occasions. No adequate explanation can be provided other than the possibility that these patients obtained placebo responses following some injections but did not experience a placebo response on the day of the saline injection. The majority of the 23 patients (78%) experienced relief at one level only.

The patients included in this study were those seen by rheumatologists practising at a tertiary referral hospital. The prevalence of 40% therefore pertains only to such a population and no inferences can be made as to the wider prevalence of zygapophysial joint pain. These figures are greater than those obtained in a recent study of North American patients with chronic low back pain in which the prevalence of zygapophysial joint pain was 15%.\( ^{14} \) Such discrepancies may be a result of the very different study populations. For example, in the present Australian study, patients were predominantly female, the median age was 59 years and none was receiving worker’s compensation. By contrast, in the American study patients were mainly male, the median age was 38 years and 75% were receiving some compensation.\( ^{16} \)

One potential criticism of the present study is the problem of referral bias. However, referral bias is not an issue: the present study was designed expressly to study the type of patients seen by rheumatologists. In principle it is possible that, because all the rheumatologists knew the focus of the study, there may have been a tendency to refer patients with zygapophysial joint pain as opposed to patients with other diagnoses. However, as borne out by the present study, the diagnosis of lumbar zygapophysial joint pain cannot be made on history and clinical examination. Therefore, the participating clinicians could not have preferentially referred patients with the proband condition. At best they could have selected only patients with idiopathic low back pain once patients with overt radiculopathy, neurological signs or neoplastic, metabolic or inflammatory disorders had been excluded. It is for that type of patient that the present prevalence figure pertains.
In conventional practice, recall bias and observer bias can affect the validity of diagnostic decisions. These were eliminated in the present study by requiring every patient to complete a series of VASs for eight hours after each and every injection while undertaking activities that would customarily bring on their pain. The written record circumscribed any problem the patient might have in remembering how much relief they obtained, and provided the observer with an objective measure of the degree of relief and its duration.

The prevalence estimate of 40% was that obtained when the definition of a positive block was a greater than 50% relief of pain. This threshold of 50% was based on an a priori decision made because it was considered that sources other than the zygapophyseal joint may also be responsible for pain in these patients. However, 32% of patients obtained 90% relief and in these patients other sources of pain would have had an extremely minor role. The 95% confidence limits of this latter estimate establish that the prevalence of lumbar zygapophyseal joint pain would be at least 20%, but could be as high as 44%. When the criterion for zygapophyseal joint pain is the complete loss of pain, the proportion diminishes to 12%. This figure is comparable to those from earlier large studies.15 17

One limitation of the present study is the possibility that some patients responding to placebo injections may in fact have been patients with true zygapophyseal joint pain. A positive placebo response does not exclude real pathology. Thus the true prevalence of lumbar zygapophyseal joint pain may, in reality, be greater than indicated in the present study. However, if patients respond to the placebo injection a diagnosis of zygapophyseal joint pain cannot be sustained unless further placebo injections are performed. Such a study could not be justified on ethical grounds.

In the present study the null hypothesis that zygapophyseal joint pain is an uncommon condition was refuted. A condition with a prevalence even as low as 20% cannot be considered uncommon. However, the study failed to refute the null hypothesis that there are no clinical features that reliably discriminate between patients with pain of zygapophyseal joint origin and pain of other sources. None of the clinical signs tested was found to be a useful discriminator of zygapophyseal joint pain. This study confirms the results of Schwarzer et al.16 Jackson et al.17 and Revel et al.,20 but contradicts the findings of Fairbank et al.2 and Helbig and Lee.12 The last two studies, however, used single uncontrolled blocks, which is likely to be the basis for the difference in results.

Whereas the results of the present study may be disappointing to those intent on diagnosing lumbar zygapophyseal joint pain by clinical examination,4 12 they do dispute the nihilism directed towards this entity in some quarters.18 21 Under stringent, controlled conditions, lumbar zygapophyseal joint pain was found to have a substantial prevalence, at least

in an Australian population of patients with low back pain. Why, some in other countries, fail to encounter this same prevalence might be explained by differences in their patients, their referral patterns and their selection bias. However, in the face of the results of the present study, failure to diagnose lumbar zygapophyseal joint pain in some quarters is not evidence that the condition does not exist.

The present study does not, however, endorse the gratuitous application of the diagnosis. The diagnosis of lumbar zygapophyseal joint pain can only, and should only, be made on the basis of controlled diagnostic blocks in each and every patient. Without such measures, false positive responses exaggerate the prevalence of this condition.

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