An Assessment of Morphological and Pathological Changes in Paravertebral Muscle Degeneration using Imaging and Histological Analysis

Ding-Chao Zhu
Yuying Children's Hospital of Wenzhou Medical College: Wenzhou Medical University Second Affiliated Hospital

Jia-Hao Lin
Yuying Children's Hospital of Wenzhou Medical College: Wenzhou Medical University Second Affiliated Hospital

Jia-Jing Xu
The second affiliated hospital and Yuying children's hospital of Wenzhou medical university

Qiang Guo
Yuying Children's Hospital of Wenzhou Medical College: Wenzhou Medical University Second Affiliated Hospital

Yi-Han Wang
The second affiliate hospital and Yuying Children's hospital of Wenzhou medical university

Chao Jiang
The second affiliated hospital and Yuying children's hospital of Wenzhou medical university

Hui-Gen Lu
The second hospital of Jiaxing

Yao-Sen Wu (✉️ wuyaosen@wmu.edu.cn)
Yuying Children's Hospital of Wenzhou Medical College: Wenzhou Medical University Second Affiliated Hospital  https://orcid.org/0000-0002-3844-2950

Research article

Keywords: low back pain, paravertebral muscle, magnetic resonance imaging, fatty degeneration, inflammatory.

DOI: https://doi.org/10.21203/rs.3.rs-482090/v1

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Abstract

Objective: The high signal of paravertebral muscle (PVM) on T2WI is usually considered to be fatty degeneration. However, it is difficult to distinguish inflammatory edema from fatty degeneration on T2WI. The purpose of this study was to identify different types of PVM high signal in patients with low back pain (LBP) through MRI and histology.

Methods: Seventy patients with LBP underwent MRI. The signal change of multifidus both on T2WI and fat suppression image (FSI) was quantified by Image J. Furthermore, 25 of the 70 patients underwent surgery for degenerative lumbar disease and their multifidus were obtained during the operation. Histological analysis of the samples was performed by HE staining.

Result: Three types of PVM signal changes were identified from the MRI. Type 1 (n=36) indicated fatty degeneration characterized by a high signal on T2WI and low signal on FSI. High signal on both T2WI and FSI, signifying type 2 meant inflammatory edema (n=9). Type 3 (n=25) showed high signal on T2WI and partial signal suppression on FSI, which meant a combination of fatty degeneration and inflammatory edema. Histological results were consistent with MRI. Among the 25 patients who underwent surgery, type 1 (n=14) showed adipocytes infiltration, type 2 (n=3) showed inflammatory cells infiltration and type 3 (n=8) showed adipocytes and inflammatory cells infiltration.

Conclusion: From our results, there are three types of pathological changes in patients with PVM degeneration, which may help to decide on targeted treatments for LBP.

Introduction

Low back pain (LBP) is a complex and multifactorial disorder commonly found in middle-aged and elderly individuals [1]. Previous studies have demonstrated that a number of lesions, such as nerve root compression, disc degeneration, modic change, facet joint osteoarthritis, and spinal stenosis are all associated with LBP [2–4]. However, the main source and related mechanism of LBP are currently unknown. Paravertebral muscle (PVM) degeneration has attracted more attention from researchers. A number of studies have shown the relationship between LBP and PVM degeneration. In the process of PVM degeneration, normal muscle fiber morphology changes and is replaced by adipose tissue resulting in a decrease in the stability of the vertebral column [5]. This may be one of the important reasons behind LBP. Additionally, Parkkola et al. [6] found increased fatty infiltration (FI) and reduced cross-sectional area (CSA) of PVM in patients with LBP compared with the control group. Storheim et al. [7] found that patients with less fatty infiltration of PVM had better results after the treatment of LBP. However, the pathological process and specific mechanism of PVM from normal to degeneration are still unclear.

At present, the measurement of PVM morphology has been an effective method for reflecting FI and has been used for investigations into the etiology of LBP [8]. Magnetic resonance imaging (MRI) as a reliable measurement method, can clearly identify the characteristics of different groups of muscle and the difference in signal intensity between fat and muscle. Previous studies were mainly carried out on the T2-
weighted image (T2WI). However, fat and liquid have a relatively longer T2-relaxation time and higher signal intensity on T2WI than other soft tissue such as normal skeletal muscle. Therefore, there is a possibility that fatty degeneration of PVM may be combined with inflammatory edema. In this study, we suspected that there might be different types of PVM pathological changes and that the high signal of PVM on T2WI might not only be fatty degeneration, but also inflammation edema. Studying PVM degeneration on both T2WI and FSI could easily distinguish the signal intensity between fat and edema.

A biopsy can help us understand the microstructure of PVM degeneration in patients with LBP. Bahar Shahidi et al. took multifidus muscle biopsy on 22 patients undergoing surgical treatment for the degenerative lumbar disease. They found high levels of muscle degeneration and inflammation and decreased vascularity [9]. Subsequently, Bahar Shahidi et al. found increased fibrogenic gene expression in the multifidus muscle of patients with chronic compared to acute LBP [10]. However, these studies employed a simple histological observation and genetic test but did not combine the histological changes in multifidus with imaging. Furthermore, although there have been previous studies about the imaging and histological changes of damaged PVM after lumbar surgery [11], studies combining imaging and histological changes of non-surgical degenerative PVM are still rare. Therefore, in the present study, we mainly studied PVM degeneration in the natural process at a microscopic and macroscopic level using MRI and histological analyses.

This study found three types of pathology in the process of PVM degeneration, and each corresponded to different imaging. Our study provides a basis for the personalized treatment of PVM-induced LBP.

**Materials And Methods**

**Patients**

A total of 70 patients with LBP who consecutively came to the Second Affiliated Hospital of Wenzhou Medical University were included in the study. Only patients with LBP, no prior spine surgery, no systemic inflammatory disease, no acute trauma, neoplasm, or infection were included in this study. The following data were recorded for each LBP patient: sex, age, body mass index (BMI) and symptoms duration, etc. This study were approved by the Second Affiliated Hospital of Wenzhou Medical University Ethics Committee and followed the guidelines of the Helsinki Declaration. All participants signed written informed consent before the experiments.

**Imaging Evaluation**

All patients underwent MRI examinations. The MRI system was a 1.5 Tesla Imaging System (GE Health care Milwaukee, USA). Images were analyzed with Image J (U.S. National Institutes of Health) and stored on the computer. Two spine surgeons with more than 10 years of work experience completed the assessment. The signal of the multifidus was quantified using the following steps on T2WI and FSI. The scale pixel was first set, and each image converted to a grayscale 8-bit image. Image J was then used to
outline CSA of the multifidus. The signal intensity was then quantified using a threshold technique. The high signal area in the 8-bit image was colored in red using the threshold tool of the program (Fig. 1).

Pathological Examination

A total of 25 underwent surgery for degenerative lumbar spine pathology. During their operation, multifidus about 1cm by 1cm were cut from diseased segment. Multifidus was fixed, dehydrated, impregnated, embedded in paraffin to form wax block and sectioned for haematoxylin and eosin (HE) staining. Histological analysis was made by one experienced pathologist, blinded to the MRI finding and objective of this study.

Statistical Analysis

Statistical analysis was performed using SPSS version 24.0 (SPSS Inc, Chicago, USA). All continuous data were described as mean ± standard deviation. An analysis of variance was carried out to detect differences in age and BMI among the different types of PVM degeneration. The Kruskal-Wallis test was performed to detect differences in disease duration among the different types. The gender in different types was analyzed using the Chi-square test. P values less than 0.05 were considered statistically significant.

Result

MRI Evaluation of Multifidus

The demographic data of the 70 patients are presented in Table 1. Three different types of PVM high signals were identified from the MRI examination. A total of 36 patients (51.42%, 57.33 ± 7.24 years) showed high signal on T2WI and low signal on FSI, which indicated fatty degeneration (Type 1, Fig. 2A.B). Nine patients (12.86%, 26.00 ± 5.12 years) showed high signal on both T2WI and FSI, which signified inflammatory edema (Type 2, Fig. 2C.D). High signal on T2WI and partial signal suppression on FSI, indicating a combination of fatty degeneration and inflammatory edema (Type 3, Fig. 2E.F) was observed in 25 patients (35.71%, 43.08 ± 5.79 years).

Histological Evaluation Of Multifidus

The demographic information of the 25 surgical patients is listed in Table 2. From the histological analyses, adipocytes infiltration was observed in 14 patients (56.00%, 60.14 ± 6.27 years). Adipocytes could be seen in muscle tissue by HE staining and the muscle cell nuclei were squeezed into periphery without inflammatory cells infiltration (Fig. 3A). A total of 3 patients (12.00%, 27.00 ± 4.36 years) showed inflammatory cells infiltration. HE staining showed inflammatory cells (mainly neutrophils) in muscle tissue without adipocytes (Fig. 3B). Finally, 8 patients (32.00%, 45.88 ± 6.15 years) showed both
adipocytes and inflammatory cells infiltration. The HE staining showed both adipocytes and inflammatory cells (mainly lymphocytes) in muscle tissue, light staining of inflammation edema area, and tissue gap narrowed or disappeared (Fig. 3C). Degenerative lumbar disease of 25 surgical patients are shown in Fig. 4.

Histological results of these 25 patients were consistent with their imaging. The 14 patients with adipocytes infiltration showed type 1 on imaging; the 3 patients with inflammatory cells infiltration showed type 2 and the 8 patients with both adipocytes and inflammatory cells infiltration showed type 3 on imaging.

**Demographic Data Evaluation**

For patients of the different types of MRI evaluation, age (p < 0.001) and BMI (p < 0.001) indicated significant difference, while gender (p = 0.424) indicated no statistical difference (Table. 1). For patients of the different types of histological evaluation, age (p < 0.001) and BMI (p = 0.025) indicated significant difference (Table. 2). In addition, there was a statistical difference in the disease duration (p < 0.001, Table. 3). The duration of type 1 patients (58.33%) mainly were longer than 12 weeks, all type 2 patients (100%) lasted for less than 4 weeks and type 3 patients (52.00%) were mostly between 4 to 12 weeks.

**Discussion**

The PVM mainly comprises psoas, erector spinae, and multifidus muscle and is considered to have two functions: to stabilize and move the lumbar vertebral column. Kalichman et al. found that fatty degeneration in the PVM was common in adults and was strongly associated with LBP [5]. Therefore, studying PVM can help to discover the potential mechanism of lumbar instability and LBP. Moreover, Guo et al. found no significant difference in psoas or erector spinae between the patients with LBP and normal people [12]. Barker et al. reported multifidus as the largest and most medial PVM that is significantly stronger in normal persons than LBP patients and is sensitive to pathological changes [13]. This indicates that multifidus might be more involved in maintaining spinal stability and preventing chronic LBP.

MRI, computerized tomography (CT) and ultrasound are ideal methods for assessing the morphology of PVM. Many researchers have inclined to use MRI to analyze PVM in patients with lumbar diseases due to its clear muscular contour and high-fat resolution [14]. Hu et al. advised MRI rather than CT for the measurement of CSA and FI by comparing the intra- and inter-reliability [8]. Fortin et al. found that an increase in fat was caused by age and BMI in a 15-years longitudinal MRI study [15]. Previous studies mainly evaluated the degree of fat degeneration through signal intensity on T2WI. But since both fat and liquid show high signal on T2WI, measurement on T2WI cannot clearly distinguish between fatty degeneration and inflammatory edema of PVM. Thus, combining T2WI and FSI could provide a reasonable contrast between fat and liquid.
CSA and FI are major indicators to evaluate PVM degeneration. Chon et al. found that CSA of multifidus was smaller in patients with chronic lumbar radiculopathy than in normal people [16]. Resorlu et al. observed that patients with ankylosing spondylitis had decreased CSA of multifidus, and it was related to the disease duration [17]. In a cross-sectional observation study, Barker et al. revealed that CSA of multifidus was smaller in patients with chronic LBP than in normal [13]. Moreover, Fortin et al. systematically reviewed the studies on the morphology of PVM in patients with LBP and the control group. They concluded that CSA of PVM in patients with LBP was significantly smaller than that in the control group [18]. Nonetheless, studies on the pathological mechanism of PVM degeneration are still rare. Therefore, in the present study, we mainly examined the pathological process for assessing PVM change.

The results of our study showed different types of signal changes in PVM. Type 1 included the patients with high signal on T2WI and low signal on FSI on MRI imaging and with adipocytes infiltration on histological analyses. Type 2 comprised of the patients with high signal on both T2WI and FSI and showed inflammatory cells (mainly neutrophils) infiltration. Type 3 included the patients with high signal on T2WI and partial signal suppression on FSI showed both adipocytes and inflammatory cells (mainly lymphocytes) infiltration. We further analyzed the age, BMI and disease duration and proportion of each type of patient. Among the 70 patients studied, type 1 had the largest proportion and mainly composed of middle-aged and elderly, the largest BMI, and chronic LBP. Type 2 had the least proportion and was mainly made up of the young, with the lowest BMI, and mainly acute LBP. The age, BMI, duration and proportion of type 3 cut across both type 1 and type 2. Inflammatory cells in type 2 were mainly neutrophils, which indicated acute inflammation whereas in type 3 the inflammation cells were mainly lymphocytes indicating chronic inflammation. Bahar Shahidi et al. found that muscle cells actively degenerated as opposed to simple atrophy, a process that may be facilitated by inflammation [9]. Hatice et al. found that chronic inflammation and cytokine-mediated fibrosis in patients with ankylosing spondylitis contributed to fatty degeneration and atrophy in the PVM [19]. In addition, Paul et al. reported a parallel increase in the expression of pro-inflammatory cytokines during the degeneration of PVM [20]. Therefore, we reasonably speculated that PVM degeneration in patients with LBP is a gradual process, and inflammation is an early pathological manifestation that might gradually transform into fatty degeneration.

Currently, many treatment modalities are used for LBP such as bed-rest, functional training of back muscle, physiotherapy, and medication, but their effectiveness are still inconclusive. Previous treatment strategies did not take into account the different pathological types of PVM degeneration. According to the imaging and histological results of this study, we speculate that personalized treatments are more likely to be effective for patients with different pathological changes. For patients with type 1 PVM degeneration, functional training for back muscle should be encouraged, because fatty degeneration is difficult to reverse with medication treatments or physiotherapy only. Bed-rest seems to not be of much benefit and may actually be detrimental because of fat deposit and muscle atrophy in these patients [21]. Storheim et al. reported that comprehensive training had a significant tendency of reversing the degeneration of PVM in patients with subacute LBP [22]. Hides et al. found that stability training could
improve the CSA of multifidus in young athletes with LBP and reduce the pain [23]. However, according to the results of this study, we do not recommend high-intensity back muscle training for type 2 patients due to its acute inflammation. Bed-rest and non-steroidal anti-inflammatory drugs could be the best option, which can inhibit the progress of inflammation. Although most acute LBP is considered self-limiting, previous evidence found that a high percentage of individuals will experience recurrent symptoms leading to poor functional outcomes over time [24]. Bed-rest and drug treatment can effectively prevent type 2 patients from turning acute to chronic LBP. Because type 3 patients have a combined fatty degeneration and inflammation, we recommend that they should be treated in stages: firstly bed-rest and use of non-steroidal anti-inflammatory drugs to inhibit the progress of inflammation, then performing functional training. For many patients with acute LBP, clinicians may recommend two days of bed-rest rather than longer periods [25]. The results of this study can provide certain scientific guidance for the specific time of bed-rest. In individuals with persistent recurrence or chronic symptoms, muscle tissue is relatively slow to respond to traditional rehabilitation measures [26]. Therefore, understanding different types of pathological changes in PVM is crucial to determining appropriate treatment strategies.

There were some limitations in this study that need further discussion and investigations. The sample size in the study was small hence a need for further clinical research to support our personalized treatment methods for LBP and demographic data evaluation. Secondly, the study did not establish the exact interaction between PVM degeneration and LBP, hence should be further studied. Based on this study, the specific transformation mechanism that may exist between fatty degeneration and inflammation in PVM needs further clarification.

**Conclusion**

Based on the findings of this study, there are three types of pathology in the process of PVM degeneration, which may provide a basis for the personalized treatments of PVM-induced LBP. Besides, inflammation presents as an early pathological manifestation of PVM degeneration that gradually transforms into fatty degeneration.

**Declarations**

**Acknowledgments:** Science and Technology Department of Zhejiang Province (LGF20H060013) and Health Bureau of Zhejiang Province (2020358020) funds were received in support of this work.

**Funding:** No relevant financial activities outside the submitted work.

**Conflict of interest:** The authors declare that they have no financial or other conflict of interests.

**Ethics approval:** The manuscript submitted does not contain information about medical device(s)/drug(s).

**Consent to participate:** All participants signed written informed consent before the experiments.
Consent for publication: All authors involved in this study agree to publish this manuscript.

Code availability: Not applicable.

Availability of data and material: All data generated or analysed during this study are included in this published article.

References


Tables

Table 1

Demographic data of patients in the MRI evaluation

<table>
<thead>
<tr>
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<th>Type 1</th>
<th>Type 2</th>
<th>Type 3</th>
<th>p value</th>
</tr>
</thead>
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<tr>
<td>Cases</td>
<td>36 (51.42%)</td>
<td>9 (12.86%)</td>
<td>25 (35.71%)</td>
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</tr>
<tr>
<td>Gender (M:F)</td>
<td>15:21</td>
<td>6:3</td>
<td>11:14</td>
<td>0.424</td>
</tr>
<tr>
<td>Age (years)</td>
<td>57.33±7.24</td>
<td>26.00±5.12</td>
<td>43.08±5.79</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>28.71±3.40</td>
<td>23.46±2.72</td>
<td>26.60±2.78</td>
<td>&lt;0.001*</td>
</tr>
</tbody>
</table>

BMI: body mass index

*indicates significant difference

Table 2

Demographic data of patients in the histological evaluation
Adipocytes infiltration

Inflammatory cells infiltration

Adipocytes and inflammatory cells infiltration

<table>
<thead>
<tr>
<th>Cases</th>
<th>14 (56.00%)</th>
<th>3 (12.00%)</th>
<th>8 (32.00%)</th>
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</thead>
<tbody>
<tr>
<td>Gender (M:F)</td>
<td>6:8</td>
<td>2:1</td>
<td>5:3</td>
</tr>
<tr>
<td>Age (years)</td>
<td>60.14±6.27</td>
<td>27.00±4.36</td>
<td>45.88±6.15</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>30.82±3.12</td>
<td>25.18±2.12</td>
<td>29.44±3.03</td>
</tr>
</tbody>
</table>

p value

<0.001*

BMI: body mass index

*indicates significant difference

Table 3

Disease duration of patients with different types

<table>
<thead>
<tr>
<th>Duration</th>
<th>Type 1</th>
<th>Type 2</th>
<th>Type 3</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Less than 4 weeks</td>
<td>3 (8.33%)</td>
<td>9 (100%)</td>
<td>8 (32.00%)</td>
<td>&lt;0.001*</td>
</tr>
<tr>
<td>4 to 12 weeks</td>
<td>12 (33.33%)</td>
<td>0</td>
<td>13 (52.00%)</td>
<td></td>
</tr>
<tr>
<td>More than 12 weeks</td>
<td>21 (58.33%)</td>
<td>0</td>
<td>4 (16.00%)</td>
<td></td>
</tr>
</tbody>
</table>

The duration of less than 4 weeks manifests as acute LBP, 4 to 12 weeks manifests as subacute LBP and more than 12 weeks manifests as chronic LBP

*indicates significant difference

Figures
Figure 1

Demonstration of threshold technology of Image J. A female patient aged 66 years old: High signal of multifidus muscle surrounded by green line colored in white on T2WI (A); High signal of multifidus muscle surrounded by green line colored in red using the threshold technique of Image J on T2WI (B).

Figure 2

Representative T2WI magnetic resonance images in patients with PVM degeneration. A female patient aged 66 years old (A and B): High signal of multifidus muscle colored in red using the threshold technique of Image J on T2WI and FSI. A male patient aged 25 years old (C and D): High signal of multifidus muscle colored in red using the threshold technique of Image J on T2WI and FSI. A male patient aged 43 years old (E and F): High signal of multifidus muscle colored in red using the threshold technique of Image J on T2WI and FSI.
Degenerative lumbar pathology of 25 surgical patients. There were 2 patients with lumbar disc herniation, 4 patients with lumbar spinal stenosis, and 8 patients with lumbar disc herniation combined with lumbar spinal stenosis in type 1 (56.00%); All three patients in type 2 (12.00%) presented lumbar disc herniation; Type 3 patients (32.00%) included 4 of lumbar disc herniation, 1 of lumbar spinal stenosis, and 3 of lumbar disc herniation combined with lumbar spinal stenosis.

Figure 4