DISENTANGLING THE LUMBAR PIVD PUZZLE:
A NARRATIVE REVIEW

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ABSTRACT
Lumbar prolapse intervertebral disc (PIVD) is a common back related disability throughout the world. It is one of common cause for work absenteeism and pose high economic burden on society. Till date, the etiology of lumbar PIVD has not been clearly established. Mechanical compression and chemical irritation are the major pathophysiological changes in lumbar PIVD. The accurate diagnosis of lumbar PIVD is a prerequisite for appropriate therapy. Lumbar PIVD results in significant disability and loss of productivity. Therefore, it is pertinent to summarize the intricacies and nuances of Lumbar PIVD as well as its evidence based management. Various cost effective and time saving protocols are available in surgical and non-surgical management of lumbar PIVD. In the present review, an attempt has been made to highlight etiology, underlying pathophysiological mechanisms of lumbar PIVD, as well as evidence based management of lumbar PIVD.

Keywords: lumbar prolapsed intervertebral disc, diagnosis, pathophysiology, management

INTRODUCTION
Lumbar prolapsed intervertebral disc is one of most common spinal problem in both men and women across the world. It causes significant disability as well as economic burden [1,2]. Lumbar disc undergoes axial loading, bending and shear forces. These forces may result in disc prolapse. Avascular nature of disc further adds to this vulnerability [3]. Pain from a prolapsed lumbar disc can vary, depending on level and severity of disc prolapse. Mechanical compression or chemical irritation of lumbar nerve root typically produces pain on one side of body [3]. After lumbar disc prolapse, if pain and numbness persists, it may require surgical intervention. Lumbar disc prolapse can be managed by various means ranging from simple bed rest to endoscopic micro-discectomy. Early diagnosis along with proper management leads to better outcome [4]. Most of the cases recover in 12 weeks. Conservative treatment is recommended to decrease pain and disability. Patient may need to undergo surgery. Failure of conservative treatment and progressive neurological deficit or cauda equina syndrome are indications of Surgery [5].

ANATOMY OF THE INTERVERTEBRAL DISC
Disc is made of three components: cartilaginous endplates, annulus fibrosus (AF), and nucleus pulposus (NP). Peripheral part of disc is annulus fibrosus while nucleus pulposus is the central one. The cartilaginous end plates are located on bodies of adjacent vertebra. The annulus fibrosus has a unique multilayer structure made up of collagen type I, type II fibers and proteoglycans, each layer oriented at 30° to horizontal and successive layer in opposite direction in such a way that leads to a criss-cross pattern. Such unique structure gives the annulus more tensile modulus against torsional, axial, and tensile loads [3]. Nucleus pulposus contains approximately 70% water, notochordal cells along with fibroblast and chondrocyte like cells. Noto-
chordal cells stimulate collagen and proteoglycans production and control apoptosis of chondrocyte like cells [6]. Hydrostatic pressure is generated in disc due to imbibed water. Hydrophilic proteoglycan macro-molecules of nucleus pulposus make a unique composition within collagen matrix. It is encircled by annulus peripherally and end plates above and below. The nucleus pulposus provides resistance to axial compression. The endplates are composed of mainly of water followed by chondrocytes, proteoglycans (PGs), and type II collagen. Capillary network of cartilaginous layer may extend into the outer portions of annular fibrosus upto a short distance that provide nourishment to otherwise avascular disc [7]. Nutrition of disc cells occurs via diffusion through the vertebral end plates. Disc is poorly innervated. Normally, nucleus pulposus and inner annulus fibrosus has no innervations. However, outer annulus fibrosis is innervated [3].

**DISC PROLAPSE OR HERNIATION**

As per combined statement of the North American Spine Society, the American Society of Spine Radiology, and the American Society of Neuroradiology, disc herniation can be defined as “localized or focal displacement of disc material beyond the limits of the intervertebral disc space”. Almost three fourth cases of disc pathology are of degeneration and one fourth are actual disc herniation [8]. Herniated discs can be classified as “protrusion”, “extrusion”, or “sequestration”. Protrusion is focal bulging of disc than diameter in have wider based herniations canal. In Extrusion, nucleus pulposus ruptures through annulus with a smaller base and larger herniation in the canal. In sequestration, disc has migrated away and no contact between herniated part and the remaining disc [8].

**EPIDEMIOLOGY**

There are a number of risk factors that may contribute to lumbar PIVD. Lumbar disc herniation is more common in males and an average age of 41 years [9]. Bostman found that in Finland, 27% of patients who underwent a lumbar disc surgery were obese, whereas at same time obesity prevalence was only 16% among population of Finland. Overweight and obese patients are more prone to lumbar radiculopathy as compared to Non-obese people [9]. Obese persons are thirty times more prone to re-operation after microdiscectomy [10]. Mobbs et al. stated that reoperation for lumbar PIVD in diabetics is more frequent as compared to non-diabetics [10]. Due to some comorbidities, impaired microcirculation of disc increases the chances of lumbar PIVD. Certain occupations may be a cause for lumbar PIVD. Most commonly affected level is L4-L5 [4]. Twisting mechanism in combination with axial load or flexion with axial load can be a biomechanics of lumbar PIVD [11]. In a retrospective study Alpesh et al. [12] support an inheritable predisposition of lumbar disc herniation [12]. Lumbar disc herniation is multifactorial, including contributions from mechanical stresses to the spine, age-dependent disc degeneration, biochemical factors and genetics.

**PATHOPHYSIOLOGY**

Natural degenerative changes in intervertebral disc include decrease in number of capillaries, altered cell morphology and density of nucleus pulposus. Annular clefts and apoptosis of fibroblast-like cells are increased in intervertebral disc [13]. Disc herniation is categorized in three steps: protrusion, extrusion and sequestration (Figure 1). Mechanical compression is commonly considered as cause of radiculopathy. Nerve roots are composed of endoneurium similar to peripheral nerve while cerebrospinal fluid and dural lining are present instead of perineurium and epineurium respectively. Such type of structure make nerve root soft and prone to get compressed mechanically. Nerve roots are in close approximation to vertebral bodies. Disc herniation exerts tensile force on nerve root similar to bowstring effect. Nutrition is impaired in mechanically compressed nerve root as both blood flow and nutrient diffusion are compromised. Intraneural edema, nerve fibrosis and injury may lead to compartment syndrome in mechanically compressed nerve. A large number of studies conclude that mechanical effect is main contributing factor in radiculopathy, while some studies claims that chemical irritation along with mechanical compression are responsible for radiculopathy in lumbar PIVD [14]. Disc herniation can be seen in magnetic resonance imaging (Figure 2). On the basis of magnetic resonance imaging, it can be in-
ferred that compression of nerve root is mostly asymptomatic. According to a study, 20% of asymptomatic participants of age below 60 years and 36% of participants of age above 60 years have radiological evidences of disc herniation. It is

![FIGURE 1. Classification of lumbar disc herniation. A. Protrusion; B. Extrusion; C. Sequestration](image)

![FIGURE 2. MRI of lumbar disc herniation](image)

![FIGURE 3. Pathophysiology of lumbar disc herniation](image)
claimed that irritated nerve root are more prone to mechanical compression as compared to normal nerve roots [14]. Some bioactive molecules found in nucleus pulposus like interleukins, tumor necrosis factor-(TNF) alpha and some other factors are responsible for nerve root sensitization making it prone to mechanical compression [15]. TNF-alpha can be the factor responsible for decrease in nerve conduction velocity. The pathophysiology of lumbar disc herniation has been summarized in Figure 3.

CLINICAL PRESENTATION

Radiating pain, abnormal sensation and weakness in area of lumbosacral nerve roots in lower limb are primary clinical presentations of lumbar disc herniation [16]. There may be localized paresis, coldness in leg, limited trunk flexion, exacerbation in pain with sneezing, straining and coughing [16]. Patients complain difficulty in sitting, as it increases disc pressure upto 40% as compared to upright standing [17]. Forward flexion also contributes to increase in pain as the pressure on lumbar disc is increased by 100-400% [17]. So lumbar disc prolapse induces radiating pain, abnormal sensation, weakness, paresis, cold leg, limited range of motion and difficulty in sitting.

DIAGNOSTIC GUIDELINES

Initial screening for diagnosis can be done by straight leg raise (SLR) test along with Hancock rule [18,19]. Three symptoms should be positive out of following four: pain along dermatome, abnormal reflex, motor weakness and sensory deficits [18]. Importance and validity of negative SLR test in clinical diagnosis is supported by a number of studies, irrespective of level of involvement [20-22]. Importance of crossed SLR tests for clinical diagnosis is also supported by a number of studies [20-22]. Various imaging tools are used to confirm the diagnosis.

Radiographic imaging

Radiograph is an important tool used to screen patients for diagnosis. Flexion and extension views may also be helpful along with antero-posterior (AP) and lateral views. Radiographs show compensatory scoliosis, osteophytes and reduced intervertebral space in lumbar disc prolapse.

Magnetic resonance imaging

Magnetic resonance imaging (MRI) is used as a gold standard tool for confirmation of lumbar disc prolapse due to its high inter-observer reliability and 97% accuracy [23].

Computed tomography scan

Computed tomography scan (CT-Scan) is also an important diagnostic tool. Clinical importance of CT-Scan is supported by a number of studies [24]. It can be used as alternative tool to MRI for suspected cases.

TREATMENT

Characteristics of major included studies are summarized in Table1. Treatment is broadly classified into operative and non-operative management. Non-operative management is further sub classified as pharmacological and non-pharmacological management.

Non-operative management

Non-operative management for lumbar disc prolapse is first choice of treatment in majority of cases. Conservative and surgical procedures both are equally effective outcomes at midterm and long term [25]. Conservative management for lumbar disc prolapse consists of non-steroidal anti-inflammatory drugs (NSAIDs), steroids, acupuncture, core exercise, taping, ergonomic advise and Physiotherapy [26-29]. Non-steroidal anti-inflammatory drugs and pregabalin are most commonly used agents and have different clinical impact [30]. A variety of lumbar epidural injections are used clinically [31-36]. However, effectiveness of such injections varies from 20-95% decrease in pain on long term follow up. Trans-foraminal injections give better result than caudal or interlaminar technique [36]. Position of patient during epidural injection can also affect the success of treatment. Decubitus position during epidural injection gives better outcome at 6 month and 12 month as compared to prone position [37]. Outcomes of epidural injections can be predicted by radiologic evaluation [38]. Epidural and subcutaneous injection of TNF-α inhibitors are effective in clinical improvement [39].
### TABLE 1. Lumbar PIVD management

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Author</th>
<th>No. of subjects</th>
<th>Intervention</th>
<th>Outcome measure</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Nakashima H et al., 2018</td>
<td>n = 60</td>
<td>Pregabalin, NSAID's</td>
<td>Visual analog scale, Patient global impressions of change</td>
<td>Pregabalin plus NSAID's group reported lesser sleep disturbance but comparatively non-significant improvement in pain then NSAID's group</td>
</tr>
<tr>
<td>2</td>
<td>Singh S et al., 2017</td>
<td>n = 80</td>
<td>Caudal epidural steroid, selective nerve root block</td>
<td>Oswestry disability index (ODI), Visual analog scale (VAS)</td>
<td>Caudal epidural steroid injection is safe and better to decrease pain and disability than selective nerve root block (SNRB) in prolapsed lumbar intervertebral disc.</td>
</tr>
<tr>
<td>3</td>
<td>Dagar A. et al. 2017</td>
<td>n = 33</td>
<td>Transforaminal epidural etanercept</td>
<td>VAS, Modified oswestry disability index (MODI)</td>
<td>Clinically and statically significant improvement in pain and disability without any adverse effect in prolapsed lumbar intervertebral disc</td>
</tr>
<tr>
<td>4</td>
<td>van Helvoirt H et al., 2017</td>
<td>n = 77</td>
<td>Transforaminal epidural injection</td>
<td>VAS, Roland-Morris Disability Questionnaire</td>
<td>Transforaminal epidural steroid injection (TESIs) centralizes pain in 62% patients and improves disability in lumbar disc herniation</td>
</tr>
<tr>
<td>5</td>
<td>Pandey RA, 2016</td>
<td>n = 140</td>
<td>Epidural steroid injection</td>
<td>Japanese Orthopaedic Association Score</td>
<td>Transforaminal route in epidural steroid injection is most effective to manage pain in patients of lumbar prolapsed intervertebral disc</td>
</tr>
<tr>
<td>6</td>
<td>Bhatia R et al., 2016</td>
<td>n = 10</td>
<td>Platelet rich plasma (PRP) via interlaminar epidural</td>
<td>VAS, Modified oswestry disability questionnaire</td>
<td>Pain and disability improves significantly without any complications. PRP can be an alternative to surgery and epidural steroid in lumbar prolapsed intervertebral disc</td>
</tr>
</tbody>
</table>

### Non-pharmacological management

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Author</th>
<th>No. of subjects</th>
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<th>Outcome measure</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Satpute K et al., 2019</td>
<td>n = 60</td>
<td>Spinal mobilization with leg movement (SMWLM)</td>
<td>VAS, ODI, Lumbar flexion ROM, Global rating of change (GROC)</td>
<td>Spinal mobilization with leg movement significantly improves pain, disability, straight leg raise range of motion (SLR ROM) and patient satisfaction in short term as well as in long term in lumbar radiculopathy</td>
</tr>
<tr>
<td>2</td>
<td>Kocak FA et al., 2018</td>
<td>n = 48</td>
<td>Motorized traction, Non-surgical spinal decompression</td>
<td>VAS, ODI, Short form (SF) 36, Beck depression inventory</td>
<td>Conventional motorized traction and non-surgical spinal decompression both are effective in pain, disability, depression and quality of life in lumbar disc herniation</td>
</tr>
<tr>
<td>3</td>
<td>Mobeen A et al., 2018</td>
<td>n = 44</td>
<td>Mobilization, Spinal decompression</td>
<td>MODI, ROM, Numeric pain rating scale (NPRS)</td>
<td>Spinal decompression with mobilization are effective to improve lumbar range of motion, pain, walking time and disability in posterolateral lumbar disc protrusion</td>
</tr>
<tr>
<td>4</td>
<td>Demirel A et al., 2017</td>
<td>n = 20</td>
<td>Non-invasive spinal decompression therapy (NSDT)</td>
<td>Numeric analog scale, Straight leg raise (SLR) test, ODI</td>
<td>NSDT can be used along with conventional physiotherapy to improve pain, functional restoration, and reduction in thickness of herniated disc</td>
</tr>
<tr>
<td>5</td>
<td>Keles BY et al., 2017</td>
<td>n = 60</td>
<td>Kinesio taping</td>
<td>Health assessment questionnaire, ODI, Numeric rating scale</td>
<td>Kinesio taping significantly improve quality of life, disability, pain and decreases the need of analgesic upto 12 weeks follow-up in lumbar disc herniation</td>
</tr>
<tr>
<td>6</td>
<td>Kiran R et al., 2017</td>
<td>n = 40</td>
<td>Stretching, exercises, Mobilization</td>
<td>Modified schober’s test, Visual analog scale, ODI</td>
<td>Thoracic mobilization and periscapular soft tissue manipulations are effective to improve pain, disability and lumbar ROM in chronic prolapsed intervertebral disc</td>
</tr>
<tr>
<td>7</td>
<td>de Carvalho MEIM et al., 2016</td>
<td>n = 54</td>
<td>LASER, LED 945</td>
<td>VAS, ODI, Flexion range of motion of hip</td>
<td>LED with lateral decubitus position and flexion exercise of lower limb shows better improvement in pain, gait claudication and disability in lumbar disc herniation</td>
</tr>
<tr>
<td>Study</td>
<td>Authors</td>
<td>n</td>
<td>Intervention</td>
<td>Outcome Measures</td>
<td>Summary</td>
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<tr>
<td>8</td>
<td>Lopez J, 2016</td>
<td>30</td>
<td>Maintained resonant oscillation</td>
<td>Visual analog scale, Range of motion</td>
<td>Maintained resonant oscillation improves pain, range of lumbar flexion and rapid centralization in lumbar disc herniation</td>
</tr>
<tr>
<td>9</td>
<td>Bayraktar D et al., 2016</td>
<td>31</td>
<td>Core stability exercise</td>
<td>VAS, ODI, RMDQ, Nottingham health profile (NHP)</td>
<td>Water based and land based core stability exercises both are equally effective to improve pain, disability and quality of life in lumbar disc herniation</td>
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<tr>
<td><strong>Surgical management</strong></td>
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<tr>
<td>1</td>
<td>Dutta A et al., 2017</td>
<td>14</td>
<td>Instrumented posterior lumbar interbody fusion</td>
<td>VAS, ODI, Modified Lee’s criteria of fusion</td>
<td>One level instrumented posterior lumbar interbody fusion gives significant improvement in pain as well as disability and restores disc height in prolapsed intervertebral disc</td>
</tr>
<tr>
<td>2</td>
<td>Wang SJ et al., 2017</td>
<td>92</td>
<td>Discectomy with different anesthesia</td>
<td>VAS, ODI</td>
<td>Discectomy with continuous epidural anesthesia has more advantages in improving pain and disability in prolapse of lumbar intervertebral disc</td>
</tr>
<tr>
<td>3</td>
<td>Wankhade UG et al., 2016</td>
<td>50</td>
<td>Discectomy by fenestration</td>
<td>Back pain functional score (BPFS), Prolo rating scale</td>
<td>Fenestration technique significantly improves in terms of complete relief of pain and return to work at six month follow-up in lumbar disc prolapse</td>
</tr>
<tr>
<td>4</td>
<td>Nikoobakht M et al., 2016</td>
<td>177</td>
<td>Plasma disc decompression</td>
<td>VAS, ODI, Short form (SF)-36</td>
<td>Plasma disc decompression is safe and effective to improve functional mobility, pain and disability in lumbar disc herniation in long term follow-up</td>
</tr>
<tr>
<td>5</td>
<td>Gugliotta M et al., 2016</td>
<td>370</td>
<td>Open discectomy</td>
<td>North american spine society questionnaire, SF-36</td>
<td>Surgical treatment provides faster relief in pain and physical function in lumbar disc herniation at short term, but no benefit in mid-term and long-term</td>
</tr>
</tbody>
</table>
Shin et al. concluded the combined effect of multiple therapies including acupuncture, bee-venom pharmacopuncture, herbal supplementation and spinal manipulation are effective in long term to improve VAS as well as ODI score [40]. Spontaneous resorption is also possible [41]. Traction and spinal decompression is also effective in decreasing disability and VAS score [42-45]. Joint mobilization, core training and active exercises are effective in improving recovery outcomes in lumbar disc prolapse [46-48]. Mesenchymal stem cell (MSC) therapy and platelet rich plasma (PRP) injection are also used clinically nowadays. Studies corroborate clinical improvement in ODI and VAS score without any complications [49-52].

Operative treatment

Surgical management is beneficial in short term as compared to non-operative management. However, in mid to long term both have no significant differences [25].

Minimal invasive surgery

Such type of intervention leads to less soft tissue lesion, reduced hospital stay and early joining back to work [53]. Trans-foraminal, trans-iliac, inter-laminar and posterolateral are few percutaneous endoscopic minimal invasive approach used commonly for spine surgery [54-56]. These interventions are associated with reduced blood loss, re-operation rate, operative time and other complications as compared to open procedure [57].

Other surgical procedure

Discectomy, instrumental posterior lumbar interbody fusion, dynamic stabilization with nucleotomy and plasma disc decompression are used having significant results [58,59]. Previously discectomy was used as a gold standard surgical procedure in lumbar disc prolapse. Discectomy procedure has a number of complications like dural tear, postoperative infection; nerve root injury and increased hospital stay [60-62]. Recent meta-analysis shows advantages of microdiscectomy over open discectomy, such as shorter hospital stay and reducing initial post-operative pain. Rongqing et al. [63] stated that percutaneous endoscopic lumbar discectomy showed shorter hospital stay and time of return to work [63].

CONCLUSIONS

Lumbar disc is almost avascular. It has small regenerative ability and bears significant axial load. Multidirectional movements along with rotational forces and axial loads commonly lead to lumbar disc herniations that may be associated with radiculopathy. Proper and accurate diagnosis for assessment of level involved, severity of compression and neurological involvement can be assessed by clinical examination, patient history assisted by radiological evidences of magnetic resonance imaging (MRI) and computed tomography scanning (CT-scan) etc. There are number of interventions including conservative and surgical approaches that are used for management of lumbar PIVD. Authors opine that for an effective and evidence based practice, attention of the therapist should be focused on the biomechanics of lumbar PIVD and specific biochemical factors that leads to pain, disability and other associated complications due to lumbar PIVD.

REFERENCES

26. de Carvalho ME, de Carvalho RM Jr, Marques AP et al. Low
25.
24.
23.
22. Knutsson B. Comparative Value of Electromyographic, Myelographic
20. de Boos N, Weissbach S, Rohrbach H et al. Classification of age-


