Lumbar degenerative disc disease is extremely common. Current evidence supports surgery in carefully selected patients who have failed non-operative treatment and do not exhibit any substantial psychosocial overlay. Fusion surgery employing the correct grafting and stabilization techniques has long-term results demonstrating successful clinical outcomes. However, the best approach for fusion remains debatable. There is some evidence supporting the more complex, technically demanding and higher risk interbody fusion techniques for the younger, active patients or patients with a higher risk of non-union. Lumbar disc arthroplasty and hybrid techniques are still relatively novel procedures despite promising short-term and mid-term outcomes. Long-term studies demonstrating superiority over fusion are required before these techniques may be recommended to replace fusion as the gold standard. Novel stem cell approaches combined with tissue engineering therapies continue to be developed in expectation of improving clinical outcomes. Results with appropriate follow-up are not yet available to indicate if such techniques are safe, cost-effective and reliable in the long-term.

**Keywords:** Low back pain; Disc degeneration; Lumbar fusion; Disc replacement; outcome

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**Introduction**

Lumbar back pain affects 70%-85% of individuals during their lifetime [1]. The prevalence of back pain increases with age and lifetime recurrence is as high as 85% [2] with chronic low back pain (LBP) being the most common cause of disability in patients between 45 and 65 years of age [3]. Multiple factors may be relevant for the onset of LBP including degenerative disc disease (DDD), facet arthropathy, disc herniation, spondylolysis and spondylolisthesis. Isolating a single cause is difficult and often multiple morbidities co-exist contributing to the patient’s symptoms, as after all, most of these conditions share a common underlying process-degeneration.

Establishing a diagnosis of primary discogenic spinal pain is critical in successfully targeting surgical intervention. LBP, in fact, may be accompanied by radicular symptoms due to central or foraminal stenosis. This paper focuses on the surgical treatment of symptomatic lumbar degenerative disc disease (LDDD), a controversial entity which may not be, in philosophical terms, distinguishable from age-related and genetically predetermined changes that affect the intervertebral disc of the human race.

LDDD is a multifactorial process with changes in disc architecture and integrity, which may lead to pain through abnormal motion of the involved segment. Specifically, abnormal stresses, either through disturbances in oxygenation, nutrition, microvasculature and/or inflammation and collagenase release are thought to contribute to disc dehydration, annular tears and loss of disc height [4].
These processes are not only thought to incite a neuroinflammatory response that stimulates peptide containing nociceptive fibres [5] but also cause increased innervation to the inner annulus and nucleus pulposus with increased stimulation of segmental dorsal root ganglions (via sinuvertebral nerves) and paravertebral sympathetic fibres (Fig. 1) [6-8]. The loss of disc integrity and shock absorption capability from a dysfunctional disc is also thought to concurrently increase the biomechanical stress upon the facet joints [9]. LDDD is often accompanied by facet-related pain due to facet hypertrophy, osteophyte formation, distortion of innervating elements and pathological motion within the facet capsule. Contributing to the pathogenesis and potentially altering the natural history of disc degeneration are several patient factors which may act as confounders in studies comparing different treatments. These include age, gender, genetics, smoking, cardiovascular disease, obesity, physical inactivity, occupational factors (repetitive heavy lifting and vibration), constitutional annulus weakness, low-grade discitis, spinal instability and malalignment [4,9]. Age related degeneration, in particular, might be impossible to discriminate from adjacent segment degeneration (ASD) postoperatively [10-12].

**Clinical Presentation**

Low back pain (lumbago) generally manifests itself as prevalent axial lumbar back pain with radiation to the buttock region. Groin pain [13] and thigh pain [14] have also been described as potential areas of referred pain from the lower lumbar discs. Radicular symptoms suggest concurrent central or foraminal stenosis. A thorough history should be obtained to exclude ‘red flags’ such as a history of recent trauma, weight loss, night sweats, night pain and fever. On physical examination, lumbar range of motion may also be reduced. The pain is typically worsened with activities that increase intradiscal pressure such as forward flexion, coughing, sneezing and mechanical loading (e.g., lifting from a position of lumbar flexion with weight in front of the body) and is reduced with extension and lying down. Pain and stiffness related to sitting or standing for prolonged periods is common. Pain becoming worse with extension, however, is more likely to be due to facet arthropathy, which may be primary or secondary to disc degeneration.

The natural history of discogenic back pain is that up to 90% of patients will experience improvement of their symptoms within 6 weeks and resolution by 3 months with or without treatment [15] and approximately 20% experience recurrence of pain within 6 months [16]. Hence, the initial treatment for discogenic back pain should strictly involve non-operative care prior to referral to a surgeon. Exercise currently represents the main component of a non-operative care program, focusing on core muscles strengthening together with mobilization techniques under supervision, including extension exercises as well as hamstring and pelvic motion orientated programs [17].

**Imaging**

Plain radiographs in patients with LDDD may demonstrate reduced disc height and in later stages, end plate sclerosis, osteophyte formation and foraminal stenosis. Magnetic resonance imaging (MRI) is the investigation of choice and, when symptoms persists over a number of months, MRI may show reduced disc height, and disc signal changes consistent with degeneration (reduced signal on a T2-weighted image). MRI may also assist in assessing other causes of lumbar back pain such as facet degeneration, an infection or a tumour. While MRI has high sensitivity in diagnosing LDDD, degenerative disc changes are commonly seen with MRI in asymptomatic patients [12,18]. However, a disc of normal dimensions, position and signal appearance on MRI is unlikely to be a substantial pain generator and should prompt a search for
other causes of back pain including extraspinal factors.

Preoperative MRI assessment can be suggestive but not diagnostic of LDDD (Fig. 2). Disc pathology itself is most commonly classified on T2-weighted MRI according to the modified Pfirrmann et al. [19] classification, which is based upon appearance of structure, distinction of nucleus and annulus, signal intensity and height. Modic et al. [20] proposed a classification system to describe degenerative vertebral endplate and subchondral marrow changes on MRI and in keeping with stages of LDDD. Three types of changes were described with this system: (1) Type I changes with hypointense T1-weighted imaging (T1) and hyperintense T2-weighted imaging (T2) representing marrow oedema and inflammation; (2) Type II changes with hyperintense T1 and isointense T2 as normal red marrow is replaced by yellow fatty marrow as a result of marrow ischemia; and (3) Type III changes with hypointense T1 and T2 representing subchondral sclerosis. Modic changes are frequently seen in patients with LDDD, and their presence has been reported as more frequently associated with clinical symptoms [21]. Among the these three types, Type I has been thought to be more strongly associated with LBP [21]. Further MRI parameters have been reported to be correlated with in vivo biochemical changes, consistent with painful LDDD, including the high intensity zone in the posterior annulus on T2-weighted MRI [22]. While MRI can image the internal morphology of the intervertebral disc, Zhou and Abdi [23] reported that relative low sensitivity (26.7% to 59%) as well as high false-positive (24%) and false-negative (38%) rates decrease the value of MRI in screening for the existence of internal disc disruption as a cause for discogenic LBP.

Discography has been utilised to confirm the clinical diagnosis of symptomatic LDDD via assessment for the presence of disruption of internal structure and reproduction of the patient’s typical pain. To be diagnostic for LDDD, injection of contrast into an abnormal disc must reproduce the patient’s typical symptoms at low pressure/volume, and symptoms should not be reproduced in adjacent normal discs at low pressure. Abnormal disc morphology such as dye extravasation from the injected annulus site indicates annular disruption, which may be interpreted as a positive finding (Fig. 3). However, typical pain reproduction is essential in classifying the discogram as positive. While the literature has mixed reports, some authors report that positive relief at provocative discography, with or without supporting MRI indicative of LDDD, is predictive of superior results for discogenic pain targeted lumbar fusion compared to patients with negative discography [24-26], and conversely, discography has the potential to be suggestive of those that may not benefit from surgical treatment targeted at LDDD [27-29].

Ideally, discography should be performed by an experienced “third party” and not upon patients with significant premorbid psychological conditions or secondary gain issues; this is to increase the independence and interpretive power of the test. In addition, discography should be carried out only for individuals deemed to be candidates for surgical intervention. Although many significant limitations of this technique have been proposed, fusion without preoperative discography has been reported as leading to poorer results compared to patients selected with preoperative discography [25,27]. With preoperative discography, it has shown to profoundly alter decision making (in as much as 71% of cases) with regard to the segments fused and the decision for surgery [30]. Nevertheless, the popularity of the technique has decreased over the years. Carragee et al. [31] highlighted the issue of iatrogenic degenerative changes from testing “healthy” control discs required for concordant testing, and drew attention to the significant influence of psychosocial overlay and second-
ary gains in false positive results. However, confusion still exists, with high sensitivity and specificity [32], high likelihood ratios [33,34] and low false positive rates found in selected studies [35] which may be related to the technique using a pain-free control disc and volume related pressure [36].

Diskoblocks using plain local anaesthetic have recently been reported to be a valuable adjunct in the preoperative assessment of patients with prevalent back pain and are thought to be predictive of satisfactory results, possibly representing an alternative diagnostic strategy to discography [37]. Some authors have found a positive correlation between pain relief from a local anaesthetic injection into the discs and contrast leakage from discography [38]. The issue of a possible cause and effect by diskoblocks for the disc has been dismissed as no radiologic indications were seen from a study seen at 5 years for any accelerated degenerative changes from injection of bupivacaine into the control discs of subjects [39]. This led to some groups previously reliant upon discography for preoperative assessment to transition to a preoperative diskoblock strategy [40].

Operative Treatment

Operative treatment for lumbar back pain has long been a topic of debate with regards to its merits over non-surgical treatments. The Swedish Lumbar Spine Study Group (SLSSG) provided the first systematic evidence that fusion for DDD resulted in superior outcomes when compared to non-surgical treatments [1]. The operative group, in fact, had a 33% reduction in back pain score and a 25% decrease in disability measured using the Oswestry disability index (ODI) whilst the non-surgical group had 7% and 3% reductions in pain, respectively. The SLSSG also reported a lower total economic cost in the operative group when compared to long-term non-operative care [2]. Almost twice as many patients returned to work when fusion was carried out, a fact that has ramifications beyond the health economics alone [2]. There has subsequently been more evidence from randomized controlled
trials to support fusion surgery for pain relief in prevalent back pain with improvement in function in patients not responding to non-surgical treatments [41,42].

Important to both the arthroplasty and fusion techniques is the concept of ASD, thought to be due to the spinal motion segment behaviour changes leading to altered physiological loads. ASD is a radiographic observation of long-term degenerative changes at mobile segments above or below a fused spine. Although such changes are multifactorial, the altered biomechanical stresses are thought to play a key role in development of ASD [43,44]. Therefore, there may be an advantage in motion-sparing arthroplasty techniques, as the use of instrumented fusion is thought to increase the risk of ASD compared to non-instrumented fusion; this is presumably due to the rigidity caused by instrumentation as with literature demonstrating a trend towards lower prevalence of ASD with motion-sparing techniques [45].

It has, however, been shown that ASD is not the result of segmental stiffness alone; there are other contributing factors such as the type of approach (anterior-based approaches showing lower rates of ASD compared to the posterior-based ones [46]), sagittal alignment, and in addition surgical and patient factors. As an example, placement of superior pedicle screws can potentially damage the inferior facet of the adjacent segment above, accelerating the onset of ASD. Other factors influencing onset of ASD are age preference (where younger patients were often selected for motion preservation thus introducing a bias), damage to the posterior ligamentous complex, the level fused (more proximal levels have higher rates of ASD compared to distal), facet joint orientation and tropism and, importantly, sagittal imbalance [43,47-50]. Harrop et al. [51] contrasted ASD, which they described as the radiographic finding of degeneration at the level adjacent to the fusion, to adjacent segment disease, which is a clinical symptomatic degeneration, often resulting in additional surgery. Whilst radiographic findings of ASD are very common and approaching 100%, symptomatic adjacent segment disease, a combined clinical and radiographic diagnosis, is said to be less frequent with an incidence ranging between 5% and 18% [52].

1. Fusion for LDDD

The rationale behind arthrodesis of a selected lumbar spinal segment is to reduce the nociceptive load by wide removal of the disrupted disc material and the stabilization of the affected motion segment(s) that are thought to be causing pain due to continued motion of the involved sensitized levels. Broadly, as a technique, it has long been considered the “gold standard” for LDDD regardless of the approach employed due to its relatively reliable results in properly selected patients. There are three main fusion techniques currently employed, namely posterolateral fusion, interbody fusion and combined interbody fusion with posterolateral fusion, also known as a 360 degree or circumferential fusion.

2. Posterolateral fusion

Posterolateral fusion (PLF) targets only the posterior elements. Through a midline skin incision with either a posterior approach or bilateral muscle splitting fascial incisions, facets joints and transverse processes bilaterally are decorticated. Local or autologous bone graft from the iliac crest can be packed into the posterolateral gutters to promote fusion. Historically, posterolateral fusion was performed without instrumentation, but due to the relatively high non-fusion rate [48,49], pedicle screws instrumentation has become standard. Fusion rates have in fact been reported as ranging from 46 to 100% and the pseudoarthrosis rate have been reported as high as 32% for non-instrumented fusion [26]. Current evidence supports the use of instrumentation to ultimately increase fusion rates [48,49]. However, the evidence of an improved functional outcome for an instrumented fusion instead of a non-instrumented one remains very limited [48,49]. A few small clinical trials have shown improved clinical functional outcomes and better fusion rate amongst instrumented patients compared to those with non-instrumented fusion [47,53]. However, other studies have suggested no clinical benefit [48,54]. It should be noted that not all pseudoarthrosis results may be painful or unstable [49] and therefore, they may not necessarily warrant revision surgery.

In search of better outcomes, spinal surgeons have switched their attention back to the disc, postulating it as the main pain generator. Weatherley et al. [55] investigated a subgroup of patients who had solid posterior fusion with persistent LBP reproduced by provocative discography. These patients experienced relief of the symptoms following subtotal disc clearance and anterior interbody fusion demonstrating the importance of disc clearance and interbody fusion.
3. Interbody fusion

Interbody fusion can be performed with either a posterior or anterior approach. Interbody fusion techniques offer several biologic and biomechanical advantages over posterolateral fusion: (1) They address the disc as the pain generator in patients with discogenic pain. (2) The anterior column of the spine supports 80% of the body load and consequently interbody devices are subjected to compressive load. Together with a large surface area, this biomechanical advantage facilitates fusion. (3) Insertion of interbody devices allows restoration of intervertebral height, correction of lumbar lordosis and restoration of sagittal balance depending on design and final positioning on the endplates, with potential for indirect decompression of the neural elements.

Various approaches have been described to achieve this purpose: (1) Posterior lumbar interbody fusion (PLIF) accesses both columns (anterior and posterior) through a single posterior approach and involves a wide laminectomy, with partial or complete facetectomy, subtotal disectomy, endplate preparation and insertion of interbody device with or without supplemental pedicle screw instrumentation. This approach allows excellent spinal canal decompression, which can address concurrent central stenosis. The wider the resection, the lesser the neural retraction to access the diseased intervertebral disc, although this approach may destabilize the segment and it is recommended that in order to achieve stability of the construct, pedicle screws be employed especially above L5–S1 and when multiple levels are involved [56]. The disc is approached through the axilla formed by the dural sac and exiting upper nerve root. The interbody device consists of either a cage (typically manufactured from Peek and carbon fibre) or a bone graft spacer, which is then inserted after end plate preparation with curettes and rongeurs. PLIF can also be supplemented with posterolateral fusion to increase fusion rates. (2) Transforaminal lumbar interbody fusion (TLIF) is a modification of the PLIF technique where the intervertebral disc is exposed unilaterally through a transforaminal approach with subtotal facetectomy in conjunction with pedicle screw instrumentation (Fig. 4). Nerve root retraction is required to access the disc, exposing the patient to a risk of nerve injury and radiculitis. Bilateral pedicle screw instrumentation is recommended to restore stability. Similar to PLIF, contralateral posterolateral fusion can be added to increase fusion rates. While some advocates of this procedure maintain that they can treat a wide spectrum of pathologies through this approach, including degenerative spondylolisthesis, a potential disadvantage is that it is a more demanding technique for implantation of the interbody device, which can affect the ability to adequately achieve sufficient lordosis and subtotal disc clearance. (3) The anterior approach for lumbar interbody fusion (ALIF) is performed through either a transperitoneal or retroperitoneal approach. This approach allows access to the lumbar discs from L2 to the sacrum. Renal vessels and the diaphragm limit the more proximal extension of the

Fig. 4. Lateral and anteroposterior lumbar plain radiographs demonstrating transforaminal lumbar interbody fusion with titanium cage and posterior instrumentation for lumbar degenerative disc disease.
exposure. The pubic symphysis occasionally also precludes appropriate visualisation and instrumentation of the L5–S1 disc space in patients with a deep-seated L5–S1 level relative to the pelvis. Lateral radiographs including the pubis are obtained during the preoperative planning stages to assess suitability of this approach at that level. Preparation of end plates and insertion of graft or implant are similar to PLIF and TLIF but the direct front-on approach allows better visualization and larger size of implants with greater capacity to increase disc height and lordosis.

4. Minimally invasive surgery fusion

The minimally invasive surgery (MIS) fusion techniques have been developed that allow access while limiting soft tissue dissection and muscle damage. These are often modifications of pre-existing open techniques. This is in response to higher patient demand and expectation for faster recovery times in line with minimally invasive technique advances in other areas of surgery in the last three decades. Proponents of MIS interbody fusion techniques have also highlighted reduced intraoperative blood loss and muscle damage [57,58] as well as faster return to work and decreased opioid dependence for the patient [59].

MIS adaptation of TLIF utilises the Wiltse paraspinal approach [60] through multifidus and longissimus intermuscular plane for unilateral facetectomy and pedicle screw instrumentation. This technique reduces muscle stripping, preserves blood supply and muscular attachment and thereby reduces erector muscle dysfunction and fibrosis [61].

More recently lateral lumbar interbody fusion and anterior to psoas approaches have been developed, which are MIS or mini-open in nature. The former technique gains access to the lateral lumbar spine through a transpsoas approach while the latter occurs anterior to psoas [62]. The patient is placed in a lateral decubitus and a small lateral incision is made to gain access through the retroperitoneal space. Neural monitoring is then placed in the posts to identify the lumbar sacral plexus. Safe access to the lumbar spine is usually obtained within the anterior one third to one half of the psoas; however, neurologic adverse events overall are reported for 18%–40% of patients [63–65].

5. Comparison of fusion techniques

Over the past few decades, there has been an explosion of different instrumentation types, fusion methods and bone graft sources developed to improve fusion rates and improve functional outcomes. Many studies have since been published to establish their utility, but results are often contradictory, with inclusion of a wide range of clinical diagnoses such as isthmic and degenerative spondylolysis and differing methods of patient selection that are often uncontrolled for comorbidities such as smoking, diabetes mellitus and steroid use. This makes comparisons difficult when attempting to advocate for one technique over another.

With regards to ALIF, the advantages of include preservation of posterior tension-band functioning musculature, improved access to the intervertebral space for disc removal and precise placement of interbody graft or implant for ease of height and lordosis restoration. The approach has several other advantages such as obviating the need for prone positioning, reduced blood loss (assuming no great vessel injury occurring intraoperatively), reduced neurological injury and adjacent segment disease [66,67]. ALIF also allows insertion of a larger structural graft without endangering neural structures. However, it can be technically demanding, particularly in the case of an obese patient and those with widespread atherosclerosis, and requires confidence with dealing with potential vascular complications due to retraction of vessels with a potential for tears to the common iliac and iliolumbar vessels [68]. Furthermore, injury to the parasympathetic plexus may result in retrograde ejaculation and autonomic sensory disturbances in the lower limbs. These disadvantages have led to a movement of surgeons employing the expertise of “access” vascular surgeons and intraoperative heparin administration to facilitate access to the intervertebral disc in the hope of reducing complications for the patient [68–70]. Bateman et al. [71] undertook a systematic review of the complication of anterior approach surgery finding intraoperative and postoperative complication rates of 9.1% and 5.2%, respectively. The most common complications reported were venous injury (3.2%), retrograde ejaculation (2.7%), neurologic injury (2%), prosthesis related (2%), postoperative ileus (1.4%), superficial infection (1%), and others (1.3%). In addition, should an ALIF technique be considered for a candidate with poorer bone stock or previous wide laminectomies, this would be a relative indication for supplemental posterior instrumentation or posterolateral fusion compared to a standalone procedure.
Early experience with ALIF as a standalone procedure was associated with high rates of non-union [72], but reports on newer techniques and devices designed for ALIF have shown good clinical outcomes and comparable fusion rates with other techniques [73] as well as comparable biomechanical properties to the addition of posterior instrumentation (Fig. 5) [74]. ALIF combined with posterolateral-instrumented fusion or also known as circumferential fusion provides the best fusion rates; however, this involves intraoperative repositioning of the patient and the added morbidity of increased operative time.

The main theoretical benefit of PLIF and TLIF over ALIF is the significant lower risk of injury to anterior vascular elements. PLIF also allows for simultaneous decompression of central canal and bilateral foraminae. While TLIF allows for only unilateral decompression of foramina compared to PLIF, it may lead to a lower incidence of neural complications (dural tears and epidural fibrosis because of the minimal and unilateral exposure of the cauda equina). PLIF is not recommended for cases involving the upper lumbar level (L1–2) due to the need to manipulate the conus medullaris. While TLIF is considered by some authors as safer to PLIF at this level due to lesser neural retraction, great caution is still required. In a meta-analysis comparing PLIF to TLIF, PLIF was associated with a significantly higher complication rates [75], in particular dural tears and root injury as well as graft

Fig. 5. Preoperative magnetic resonance imaging of a 40-year-old male who had recurrent disc prolapse with radiculopathy following an initial discectomy (A, B). Postoperative radiographs of anterior lumbar interbody fusion performed for the same patient demonstrating progression of fusion at 6-month follow-up (C, D).
malposition. However, there were no differences with regards to clinical satisfaction, blood loss and radiographic evidence of fusion between the two groups, even if PLIF required longer operative times.

Unilateral instrumentation has also been advocated as an alternative to bilateral instrumentation for spinal fusion with interbody device. The theoretical advantage is shorter operative times and lower operative costs for implants [76]. In a systematic review of ten randomized trials, Molinari et al. [76] reported no difference in clinical outcomes, union rates nor complications when single-level unilateral instrumentation was used for degenerative conditions in the lumbar spine compared to bilateral instrumented fusion with PLIF and TLIF.

When comparing the three main types of fusion techniques—posterolateral, posterolateral with instrumentation, and circumferential—SLSSG found that circumferential fusion had the highest fusion rate, followed by instrumented posterolateral fusion, for all groups having significantly reduced pain and disability and no overall significant differences in clinical outcomes between the groups. The circumferential group, however, had significantly more complications, prolonged operating times and prolonged hospital stays.

The issue of fusion rate needs careful interpretation. Radiological determination of lumbar fusion is known to have poor intra- and inter-observer consistency and depends on the method utilised. Sensitivity and specificity of plain radiographs have been reported to be as low as 43% and 50%, respectively [48]. Additionally reported fusion rates in the literature can be flawed with bias, as in many cases the designation of fusion status is assigned by the operating surgeon. In addition, fusion rates are not the only measure of success after lumbar spine fusion and indeed some pseudoarthroses (such as the “locked” pseudoarthrosis [77]) are not incompatible with good outcomes.

A randomized controlled study by Videbaek et al. [78] was in contrast to the findings of the Swedish Lumbar Spine Study Group, and reported improved mid-term outcomes with circumferential fusion when compared to instrumented posterolateral fusion alone. In this trial, there was a statistically significant improvement in pain score and functional score with a 5 to 9 year follow up [78]. The study population was significantly younger when compared to the Swedish group with a majority of patients being at work. A further study by Soegaard et al. [79] supported these results, reporting superior cost saving in the circumferential group compared with posterolateral fusion group in the long term.

In line with current evidence, one may suggest choosing interbody fusion techniques in the younger, more active and working individuals and in patients with an increased risk of non-union i.e. smokers. Also, the debate between conventional or minimally invasive techniques will continue until long-term comparisons of these techniques in well-designed studies are available. In order to displace conventional techniques as the standard of care, MIS techniques would need to demonstrate at least equivalent outcomes without additional costs and increased complications, in addition to the currently reported beneficial short term outcomes of reduced hospital time and quicker return to employment [80].

6. Interbody and graft options

Previously, tricortical blocks of autologous iliac crest bone graft were used in surgery. This presented substantial technical challenges, including the sizing of the grafts, its stability once impacted into the intervertebral disc space and the issue of subsequent collapse and subsidence. The development of interbody devices with instruments designed for ease of introduction has increased the technical ease of interbody fusion. There is currently a wide range of materials used as interbody devices with which various graft options can be used to enhance fusion. These interbody devices also have design characteristics that may help in restoring lumbar lordosis, sagittal balance, indirect neuroforaminal decompression and physiological load through the anterior spinal column. Titanium cages have over the years been popular but can be associated with subsidence. Interbody cages made of carbon fibre and the non-absorbable, biocompatible material polyetheretherketone (PEEK) are radiolucent and are thought to have a modulus of elasticity closer to bone. These cages may have lower subsidence rates and, in addition, their usage allows easier assessment of fusion rates on radiographs. However, some debate exists on the enhancement by titanium implants for bone incorporation into the implant as opposed to PEEK, which is thought to stimulate fibrous, non-mineralised tissue.

Iliac crest graft has been long been considered the gold standard for spinal fusion, but it has several disadvantages compared to currently available alternatives. These include
associated donor site morbidity, increased blood loss, prolonged hospital stay and often limited graft volume in older patients. Bone graft substitutes can be broadly classified into two categories: (1) There are osteoinductive agents that induce formation of bone, and these include agents such as recombinant human bone morphogenetic protein, bone marrow aspirate etc. (2) There are also osteoconductive agents, including demineralized bone matrix, tricalcium phosphate etc., that provide a scaffold for new bone growth.

Allograft yields comparable fusion rates to autograft without donor site morbidity [81], but with an increased cost and a potential for infectious disease transmission. Demineralised bone matrix (DBM), while still on-going investigation regarding its efficacy, has had early positive reports with high fusion rates [82]. Currently DBM is used as mass extender, combined with local autologous bone graft, rather than used in isolation. Bone morphogenetic proteins (BMPs) are reported to be at least as effective as autologous bone graft substitutes or extenders [83]; however, there are concerns regarding their safety. With BMPs, extensive osteolysis resulting in graft subsidence and cage migration has been reported [84]. Higher rates of retrograde ejaculation have also been reported when used in ALIF, thought to be due to the inflammatory response associated with release of cytokines due to BMP in closed vicinity of the parasympathetic plexus [85]. However, Scott-Young [86] reported the incidence of the above to be only 0.7% in a large cohort of males who underwent ALIF and similarly another study did not find the high rates of complication previously reported [87]. Although not proven, there are theoretical concerns for a cancer when applying exogenous transforming growth factors such as BMPs [88]. A multitude of other graft substitutes and biological mediators are being introduced into clinical practice, but are beyond the scope of this review.

7. Lumbar disc arthroplasty

Lumbar disc arthroplasty (LDA) has been increasingly adopted into routine spinal practice since its introduction in 1966 [89]. After a lag period, the Charité Hospital in Berlin, East Germany led the trend for widespread implantation beginning in the 1980s. LDA typically is performed through an anterior approach at lower lumbar levels (L4/5 and L5/S1) in younger and active individuals with limited ASD, adequate bone stock, absence of spondylolisthesis and normal facet joints (Fig. 6). Non-inferiority studies involving both cohorts of patients implanted with the Charité prosthesis compared to ALIF [90], the ProDisc prosthesis compared against circumferential fusion [91] and the Maverick compared to ALIF [92], respectively, showed favourable results for LDA but each had methodological flaws. There is no current conclusive evidence of LDA superiority in the long-term in level I studies and the surgery is technically more challenging. However, mid-term studies of LDA have reported satisfactory clinical results and implant survival and comparable complication profiles to fusion [93,94] with proponents of LDA supporting its use for several theoretical and clinical advantages.

The major driver for development of this technique has been the motion preserving philosophy as an alternative to fusion with theoretically lower risk for ASD. European studies have cited rates of ASD at 10-year follow-up of LDA to be 2.0%–2.8% and up to 4.5% in a large North American study [95], which compares favourably to historical rates in lumbar fusion of between 14 and 29%. However, it may be unfair to compare ASD to fusion procedures indiscriminately as the anterior approach itself has been reported as being associated with lower rates of ASD [96,97].

Other advantages cited for LDA include having no need either for grafting or screw based fixation [98] and having a lower reoperation rate with at least equivalent rates of return to work and clinical outcomes to fusion at mid-term [99]. In one study, David [100] concluded that the “rate of reoperation secondary to adjacent segment disease is ten times lower than the rates reported in the literature for fusion.” Furthermore, the rates of secondary sacroiliac joint dysfunction following LDA were low in the mid-term [101]. Studies of modern LDAs have also reported faster patient recovery, higher satisfaction in the short-term compared to the fusion techniques [95]. Patient satisfaction rates have been in the order of 88% or more [90,102,103], at least equivalent to fusion satisfaction rates quoted as 85%–95% [104,105], although some series have reported a mild but statistically significant deterioration in outcomes of patients transitioning from early to mid-term follow-up [94]. High rates of return to work and lower long-term unemployment have also been reported as other benefits for the LDAs [90]. These results may also be further improved with focused physiotherapy [106].
Disadvantages cited in the literature are the disappointing outcomes from some of the earlier designs (perhaps confounded also by patient selection), the lack of long-term outcomes of modern designs, the lower versatility in indications compared to fusion and the difficulty in acquiring adequate surgical expertise in performing this procedure. Many of the cited complications such as implant extrusion and vascular injury can be linked to inadequate training, improper sizing and lack of confirmation of satisfactory placement on imaging [107] with implant impingement linked to the aforementioned factors [108]. Design variables relating to constraint and core mobility have also been shown to alter wear and force transmission in in vitro studies [109,110].

Earlier, some of the initial LDA designs led to inconsistent outcomes [111]. For example, the cumulative survival for the Acroflex LDA was only 60.7% at 10 years when first revision surgery was used as the end point. Etiology of the implant failure reported by authors [112] included failure of osteointegration, midsurface elastomeric tears and osteolysis. Complication rates, while reported to be lower than fusion in the Charité trials (29.1% for arthroplasty and 50.2% for fusion at 2-year follow-up), are still significant [113]. These can be broadly separated into those related to the anterior surgical approach (e.g., vascular injury, nerve root injury, retrograde ejaculation), prosthesis/fusion failure (e.g., subsidence, osteolysis, migration, implant fracture, endplate fracture, pseudarthrosis), heterotopic ossification (up to 76% at 3 years) and subsequent hypomobility of the implant and donor-site complications [93]. Device failures necessitating repeat operations have been reported at 5.4% to 6.3%. Salvage procedures involving conversion to spinal fusion are technically demanding; however, they appear to improve outcomes modestly in failed LDA [112]. Earlier designs were also associated with a high rate, in the order of 60%,
of spontaneous fusion in the long term [111]. Metal ion and polyethylene wear particle release from the articulation interface also add to the disadvantages [114]. Limited versatility exists compared to fusion techniques as there are relative contraindications for three or more lumbar levels, high lumbar disease (above L3), spondylolisthesis, severe ligamentous instability, facet degeneration, adjacent prior fusion and conditions affecting bony healing (e.g. rheumatoid arthritis, diabetes mellitus and steroid use) [115,116].

**Multiple-level LDA and Hybrid LDA/Fusion Techniques**

Despite good results for single-level LDA, there is evidence to suggest that the two-level arthroplasty constructs have comparatively poorer outcomes and have an increase in the rate of complications compared to the two-level fusions [94,117,118]. However, conflicting data exists with some authors reporting good outcomes with two-level LDA [119] and outcomes comparable to single-level LDA [117,120,121], and thus calling into question the role of design and constraint of the implant itself [122,123]. The poorer outcomes were in fact thought to be due to increased segmental instability and postoperative facet and sacroiliac pain from increased loading [94].

Some authors have proposed hybrid constructs of LDA/ALIF in addressing multilevel disease in the lumbar spine, and combining the perceived advantages of motion-sparing and fusion techniques (Fig. 7). However, there are currently limited published results in this area. This approach currently has only been reportedly used for two-level disease in the literature, with ALIF performed on the more severe (usually inferior) level and LDA on the less severely affected or hypermobile level. Hoff et al. [124] reported results of a randomized trial of ALIF and LDA compared with circumferential fusion using TLIF posteriorly with a mean of 37 months follow-up. The hybrid group was associated with a lower VAS with and superior lordosis at the lower lumbar levels as well as having comparatively increased range of motion at the L4/5 level with a similar low complication profile and overall. Similarly, Aunoble et al. [125] published their experience in 42 patients over 2 years follow-up and showed a 53% improvement in the functional score and a 65% improvement in the visual analogue score. For the same group of patients, however, symptoms derived from parasympathetic changes were noted in only 9.5%. Kasis et al. [126] additionally presented improvement in all outcome parameters at minimum 2 years follow-up with a low complication rate. Long-term studies are required in this area to assess viability of this technique in addressing multiple-level LDDD and, specifically, to assess whether these techniques are superior clinically to multiple-level fusion. This would need assessing not just the clinical outcomes but also the complication profiles, ASD, revision rates and economic costs.

1. Novel technologies

Stem cell therapy combined with tissue engineering approaches for effective delivery into the degenerative disc has been proposed as a minimally invasive alternative to manage LBP. This therapy focuses on the restoration of extracellular matrix of the “damaged” disc where autologous nucleus pulposus cells or bone marrow derived mesenchymal stem cells are harvested, expanded in vitro and delivered into the nucleus pulposus, with the hope of stabilizing or reversing LDDD pathology. In a recent systematic review evaluating the effectiveness of a tissue
engineered approach in the treatment of DDD in controlled animal studies, the technique has shown promising results in improvement in disc height and MRI signal intensity but none of the studies showed restoration to the properties of a healthy disc [127]. Human trials on the use of mesenchymal stem cells from autologous bone marrow in patients with LDDD are currently on-going in multiple centres. Finally, short term results of intradiscal pulsed radiofrequency have been promising and have shown not to be significantly different from intradiscal electrothermal therapy at 6 months [128]. Results with appropriate follow-up are not yet available to indicate if such techniques are safe, cost-effective and reliable in the long-term.

Conclusions

While LDDD is extremely common, the role of operative treatment and the choices of which operative treatments to select for which patients remain controversial. There is scientific evidence to support surgery in carefully selected patients who have failed to respond to appropriate non-operative treatments over a minimum of six months and do not exhibit any substantial psychosocial overlay. Confirming a clinical diagnosis of discogenic back pain with supportive imaging is, however, crucial for targeted treatment of the condition with either disc clearance and fusion or LDA.

LDA for LDDD is still a relatively novel procedure despite promising mid-term outcomes. As revision procedures for LDA can be complex with expenses considerable, long-term studies demonstrating superiority over fusion are required before this technique may be recommended to replace fusion as the gold standard. While early evidence suggests comparable results between single-level LDA and fusion, the role of multilevel LDA has not yet been fully scrutinized in comparison to multiple-level fusion. There is no current long-term evidence that motion-sparing LDA reduces symptomatic adjacent segment disease or robust long-term evidence for hybrid techniques combining arthroplasty and fusion. Should multiple-level LDA or hybrid constructs demonstrate long-term evidence in line with their promising short- to mid-term outcomes, they may surpass fusion surgery as the treatment of choice for patients with multilevel LDDD who meet indications for either fusion or LDA.

Novel and minimally invasive fusion techniques continue to be developed in expectation of improving clinical outcomes through minimizing soft-tissue damage. However, the only long-term evidence for predictably successful clinical outcomes currently exists in carefully selected patients undergoing open fusions and employing correct grafting and stabilization techniques. As such, the best approach for fusion remains debatable. In addition, the more complex, technically demanding and higher risk interbody fusion techniques, such as ALIF or TLIF, might be advocated for younger, active patients or patients with a higher risk of non-union.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

Acknowledgments

We would like to thank Ass. Prof. Matthew Scott-Young (Gold Coast Spine, Queensland, Australia) for enlightening perspectives on anterior approach surgery and lumbar disc arthroplasty.

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