After some 40 years of research and development, artificial disc technology has finally reached the point at which several designs are either near the end stage of preclinical study or in the early phase of clinical trial, with promising results so far. Due to the multicomponent structure of the disc, surgeons performing disc arthroplasty have the option of replacing either the entire disc or a portion of it. The decision will be largely dependent on the pathological entity addressed, the condition of the patient's spinal disc and surrounding tissues, and the cost and potential risk of the procedure. Driven by demand, almost all the emphasis in artificial disc development has been placed on the lumbar disc, with a smaller effort directed toward the cervical disc. No attempt has been made to develop an artificial thoracic disc. However, by examining the differences and similarities in structure, anatomy, function, mechanism of degeneration, pathology, surgical technique, and complications between the lumbar and thoracic disc, the authors believe it is feasible to apply artificial disc technology in the treatment of thoracic disc disease. Nonetheless, due to the rarity of thoracic disc disease and the more stable structure of this spinal component, the demand for artificial disc or artificial nucleus technology for the thoracic disc probably will be smaller than that for lumbar disc technology.

KEY WORDS • thoracic disc • artificial disc • artificial nucleus • disc degenerative disease

Abbreviations used in this paper: VATS = video-assisted thoracic surgery.
directions in any adjacent layers at approximately 30° to the horizontal plane. This structure provides the annulus with a high tensile modulus and strength, as well as equal torsional modulus in either direction. The nucleus is enclosed within the annulus and the endplates and is composed mainly of a very hydrophilic polymer called glycosaminoglycan, which is capable of absorbing a large amount of water and which forms a gellike matrix. The amount of water absorbed by the nucleus depends not only on the composition of the polymer matrix but also on the external pressure exerted on the disc. At a young age and when the disc is healthy, 80% of the nucleus is constituted by water, even under normal loading conditions, and this keeps the annulus well inflated. Again, somewhat like an automobile tire, when the annulus is inflated the disc becomes very stiff and stable. Although the cartilage endplate contributes little to the overall mechanical properties of the disc, it plays an important role in allowing nutrients to pass into the disc. It should be noted that the mechanical properties of a disc are a function of the structure and integrity of both annulus and nucleus in combination. A change in either one of these parameters can affect the overall properties of the disc.

Functionally, the intervertebral disc performs two important, but somewhat conflicting, duties: it maintains spinal column stability while providing the column with necessary flexibility. Without intervertebral discs, the human spine is unable to bend and its function is greatly impaired. However, because numerous nerves course alongside the spinal column as well as directly down the spinal cord, the disc has to keep the vertebral bodies separated so as to hold the foramen space open and provide adequate stability.

Disc degeneration often occurs in patients at a much younger age than hip and knee degeneration. Disc degeneration, which is believed to be the major cause of low-back pain, often begins with a structural change in which the nucleus loses its water-binding capacity and the disc consequently deflates. After this happens, more compressive loading shifts to the annulus, rendering this structure more susceptible to delamination and damage. Damage to the annulus, in turn, accelerates the disc degeneration process. Disc degeneration can also accelerate the degeneration of other surrounding tissues, such as facet joints. This degeneration cascade in the disc joint has been well documented.

Currently the two most common surgical procedures, discectomy and fusion, at best only address the symptom of low-back pain. Biologically and biomechanically, both procedures actually worsen the condition of the affected disc, adjacent discs, and surrounding tissues, leading to further degeneration. Hence the long-term results of both these procedures are relatively poor. Logically, a better solution is implantation of an artificial disc, which is intended not only to treat low-back pain but also to restore or maintain the normal anatomy and function of the diseased disc.

Several review articles on various types of artificial disc have been published. Somewhat like hemiarthroplasty in other joints involved, in intervertebral disc replacement one can replace either the entire disc or only its nucleus. The former prosthesis is called an artificial total disc and the latter an artificial nucleus.

**Artificial Total Disc**

The artificial total disc is designed to replace the entire disc: annulus, nucleus, and (very often) endplates. Because the function of the endplate is more biological than biomechanical, it often does not need to be preserved after the disc is replaced with nonbiological material(s), unless the design entails articulating the artificial disc surface with the endplate for reduced friction and wear. Most artificial disc designs require removal of the endplates and fixation of the superior and inferior surfaces of the implant to the vertebral bodies. The main benefit of replacing the entire disc is that the disc is consequently less dependent on the integrity of the annulus and the stage of degeneration. Conceptually, artificial discs can be used in patients with disc degeneration at any stage of progression. Because of the added cost and risk involved in implanting such a device, however, in practice its use can often be justified only in patients with more severe disc degeneration.

Due to the complexity of the structure and function of the disc, it has proven difficult to design an artificial total disc that mimics all the mechanical properties of a natural disc while retaining the required durability. A multicomponent design incorporating several materials with very different mechanical properties or a composite structure is often required. In a multicomponent design, researchers are also often faced with problems such as interfacial bonding and wear.

For flexibility, either the material must be elastic itself or the design must have elastic characteristics at least in one direction (preferably in multiple directions). On the other hand, because the implant must maintain a firm fixation to the vertebrae, a hard material such as metal must often be used for the superior and inferior surfaces of the device. Fixation is often achieved by one or a combination of the following mechanisms: 1) anchoring through one or several pegs or posts inserted into the vertebrae; 2) physical interfacing via a threaded surface; 3) promotion of bone ingrowth by means of a porous surface; or 4) fixation with screws through a side wing extending from the plate.

To minimize contact stress, researchers tend to design the device to cover the entire cross-sectional area of the vertebra so that the load can be spread over a large surface area. Although this makes mechanical sense, it renders surgical insertion more difficult and requires almost every artificial disc to be implanted via an open anterior approach with a large incision, and also entails prolonged operative time. Use of a hard or rigid material for superior or inferior surfaces further increases the bulkiness of the implant. Given the current trend of using less or minimal invasive surgery (even for some fusion surgeries), this type of device may have difficulty gaining wide acceptance among surgeons.

In cases of fixation failure, the bulkiness of the artificial total disc could prove catastrophic. Although artificial disc implantation offers some benefits over fusion, it would be difficult to convince surgeons to use the device if the potential risk to patients is higher than that associated with fusion. In view of the fact that the current fusion rate is higher than 90% for the fusion cage (possibly even higher when using some forms of bone growth factor),
Artificial disc technology

and given the relatively low incidence of cage migration, the fusion procedure has set a high standard for the artificial disc to meet.

Although the concept of a disc prosthesis was first set forth in 1956 in a French patent by van Steenbrugghe, it was not until 17 years later that Urbaniak, et al., reported the first disc prosthesis, which was prototyped and implanted into a chimpanzee. Since then, many other disc prosthesis concepts have been proposed, mostly in the patent literature.

Basic design concepts and component material(s) can be divided into three groups: metal, nonmetal, and metal in combination with nonmetal. The main advantage of a disc prosthesis design using metal alone is its inherently high fatigue strength compared with a nonmetal design. Because patients with back pain are on average approximately 40 years old—that is, much younger than the population of patients requiring total hip or total knee patients—it is proposed that the device should have at least 40 years of fatigue life. Researchers who use exclusively metal materials designs believe that they are thereby more effectively addressing the fatigue issue. However, because most metals are much stiffer than the natural disc they are used to replace, they must be designed in special forms to reduce this stiffness and provide needed flexibility.

The most widely tested all-metal disc prosthesis, developed by Kostuik and associates, features two Ti-6Al-4V springs poking between two forged or hot isostatically pressed cobalt-chromium-molybdenum alloy plates with a posterior hinge allowing flexion and extension. In designing the implant an effort was made to meet various design criteria such as biocompatibility, endurance, kinematics, dynamics, and bone fixation. The hinge provides the implant with a full range (15–20°) of axial rotation in the sagittal plane as well as a small amount of rolling lateral rotation (3–6°) to match the physiological range. The springs were chosen to duplicate the stiffness of a natural disc. Vertically projecting lugs are positioned at the front and side of the plate members, through which screws can be placed for fixation. Fatigue tests were conducted on the individual components (100 million cycles), and no failure was found in a total of 17 springs. In a sheep model, it was shown that fibrous tissue does not grow between the hinges or around the coils, at least in the short term. Such ingrowth would be expected to interfere significantly with implant mechanics. However, due to the bulkiness of the device and the potential risk associated with its use, it has yet to be applied in humans.

Whereas the principle advantage of a metal disc prosthesis is its fatigue strength, the primary benefit of a nonmetal design is its mechanical similarity to the natural disc. With its lower elasticity modulus, it more closely replicates disc kinematics. The challenge, however, arises when attempting to develop a long-lasting device (longevity of ≥ 40 years) and an interface that promotes ingrowth of the bone implant.

The concept of a disc prosthesis was introduced with a nonmetal design by van Steenbrugghe. The implant comprises a multicomponent disc encompassing intermediate cushion inlayers with a plastic body of varying shapes. However, the inventor did not mention how the implant was to stay in place. No laboratory test has been reported on this device. The most widely tested device in the nonmetal category is that designed by Lee and colleagues. This design features a soft elastomer central core to mimic the function of the natural disc nucleus and reinforcing fiber sheets with specific alternating fiber orientation in six to 15 laminae embedded in a second elastomer to mimic the function of the annulus, as well as two stiff plates. By selecting appropriate materials, the device is able to mimic both the compressive modulus and the compressive–torsional stiffness of the natural disc. Data on compression–flexion stiffness and compression–extension have not been reported. Lack of adequate implant and vertebra fixation is believed to be the major obstacle to the clinical use of this device.

To take advantage of the benefits of both metal and nonmetal materials and overcome the drawbacks involved in using either of them alone, many researchers have combined both types of materials in their designs. Most commonly this has taken the form of a metal-polymer-metal sandwich design. A metal plate is used to improve fixation by means of spikes, tabs with screws, or porous coating for ingrowth. With the component thus stable and fixed, the polymer may provide the needed flexibility.

Although many artificial disc designs have been proposed over the years, only three have been tested extensively and used clinically. The LINK SB Charité device has undergone the largest and longest clinical trial of any existing artificial disc. It was developed by Buttner-Janz and colleagues in the mid-to-late 1980s. The prosthesis has undergone several major design, structural, and manufacturing modifications from the first to the third and current generation. The design features two concave endplates composed of a cobalt-chromium-molybdenum alloy. The plates have spikes or teeth that enable them to be fixed without cement to the vertebral body. A biconvex oval polyethylene spacing piece with contours to match the endplates is placed between them. Similar to the ball-and-socket design of hip joint prosthesis, this device allows rotation in all three directions (unpublished data). However, due to the lack of elasticity of the material, this device cannot replicate the normal compressive stiffness of the natural disc.

Since 1984, well over 2000 prostheses of all three generations have been implanted in Europe. The most detailed clinical data for this device were reported by Griffith, et al., after a multicenter retrospective study conducted on 93 patients in whom a total of 139 third-generation prostheses were implanted. The predominant surgical indication was for degenerative disc disease, usually occurring at L4–5 or L5–S1. The average length of follow-up review was 11.9 months (range 1–37 months). Using a 10-point analog pain scale, the data indicated a statistically significant alleviation of pain in both leg and back after disc implantation, with greater gains in relieving leg pain than back pain. The authors stated that the improvements in leg pain might be a result of the degree of decompression achieved by the surgeon and not necessarily a direct effect of the use of prosthesis placement.

Parallel to European developments, Steffee conducted his pilot clinical study in six patients in the United States between 1988 and 1989 after some design work and laboratory testing. This disc consists of a hexene-based, carbon black–filled polyolefin rubber core vulcanized to two
titanium plates. Fixation is accomplished with a porous coating to promote ingrowth and four 7-mm conical posts that are socketed into the vertebral body. In the six patients in the preliminary clinical series (as reported in Fernstrom and McMilland and Steffee) there were two prostheses failures caused by fracture of the rubber core. Several reasons could be evinced to explain this high rate of device failure. First, implants were used at disc levels adjacent to the fusion site in four of the six patients; this may have led to the exertion of severe stress at the implant level. Second, one of the patients had degenerative scoliosis; without secure plate fixation and resistance to intense shear force, deformity should be a contraindication for implanting an artificial total disc. Third, the shear-fatigue strength of this rubber core might not have been sufficient.

More recently, a cap–cup matching articulation device was designed by Marnay. The implant (ProDisc) is composed of titanium endplates with titanium plasma spray coating on the outer surfaces and a polyethylene core. Marnay (unpublished data) recently reported his clinical results in 50 patients after a minimum 7 years’ follow up, with excellent or good results in 78% of these patients.

Artificial Nucleus

The nucleus prosthesis approach has several obvious advantages over the total disc prosthesis. By replacing only the nucleus, it preserves the remaining disc tissues—annulus and endplates—and therefore preserves their functions. Because the nucleus has a much simpler structure and function than the annulus and endplates, the design of the nucleus prosthesis allows surgeons to leave the annulus and endplates intact, making the surgical procedure much easier than that required to replace the entire natural disc.

The primary objectives of the nucleus prosthesis are to reinflate the annulus and to relieve the compressive load on this disc component by sharing a significant portion of that load. Another major advantage of the nucleus prosthesis is that it can be implanted by means of a minimally invasive surgical procedure. Depending on the design and choice of material, it is even feasible to implant the device using an endoscopic technique with only a small incision in the annulus. Because the implant is not designed to be affixed to the vertebrae, no fixation component is required. The surgical time required should be comparable to that required for a discectomy.

The risk associated with placement of a nucleus prosthesis should be lower than that associated with a total disc prosthesis. Although implant extrusion remains a primary concern in the design and use of a nucleus implant, it is less likely than a total disc prosthesis to cause permanent injury to the nerve because of its relatively small size and soft consistency. In case of prosthesis failure, it is relatively easy to remove the implant and convert to a primary fusion or perform a disc replacement procedure.

The major limitation of the nucleus prosthesis is that it can be used only in patients in whom disc degeneration is at an early or intermediate stage because it requires the presence of a competent natural annulus. In a disc at a late stage of degeneration, with severe annular tears and delamination, the annulus may not be strong enough to provide the needed constraint. Most nucleus prostheses would also not be suitable for discs with significant height loss, which also often occurs during the later stages of disc degeneration. When a disc loses height over time, the collagen fibers may adapt into their new structural characteristics and would hence be difficult to stretch significantly at surgery.

Without much prior laboratory testing, Fernstrom initiated the clinical use of a spherical endoprosthesis consisting of a stainless-steel ball. The diameter of the ball ranged from 10 to 16 mm for the lumbar disc and from 6 to 10 mm for the cervical disc. This solid-ball device was meant to serve as a spacer allowing movement of adjacent vertebrae. It did not, for obvious reasons, restore the normal stiffness of the disc. Although the only published long-term clinical study of this device showed reasonably good clinical results in a fairly large group of patients, its use has been discontinued due to concerns about implant migration and subsidence.

Aware that metals are too stiff for a nucleus device, many researchers have experimented with various elastomers (either preformed or formed in situ) for nucleus replacement. Other than the device invented by Hou, which was used in a limited number of patients, many of these devices did not meet the basic requirements, such as biocompatibility and mechanical strength, for a clinical trial.

In 1990, Bao and Higham pioneered research into using hydrogel material for intervertebral nucleus replacement. After having evaluated many different hydrogel materials, they developed a hydrogel nucleus implant designed to contain approximately 70% water content under physiological loading conditions while also meeting the mechanical requirements. As reported at the 11th annual meeting of the North American Spine Society, these researchers, in collaboration with others were also the first to demonstrate that the hydrogel nucleus implant can mimic the capacity of the nucleus tissue to absorb and release water under cyclic loading conditions. Biomechanical studies (reported at the aforementioned meeting) have confirmed restoration of disc anatomy and function after implantation of a hydrogel nucleus. A study using a baboon model (unpublished data) has also shown that there is no adverse local or systemic tissue reaction. This nucleus device may well reach the clinical trial stage in the near future.

The nucleus implant in most widespread clinical use is the one developed by RayMedica (Bloomington, MN). In 1988 Ray and Corbin introduced the use of dual disc cylinders for disc nucleus replacement. The threaded outer layer corresponding to the light threads cut into the endplates was a double-woven spiral of flexible, high-tensile-strength polymeric fibers and tissue ingrowth–promoting polyglycolic acid filaments. The flexible and semipermeable cylinders contained a viscous hygroscopic semifluid. Perhaps because of the technical difficulty involved in sealing the aforementioned semifluid into the capsule, this concept was abandoned. In the mid-1990s, Ray, et al., working for RayMedica, decided to change their approach, using for the core a hydrogel constrained by a flexible but inelastic woven polyethylene-fiber jacket. The shape of the implant was also changed to resemble that of a pillow. The implant position was changed from side-by-side in an anterior–posterior direction to a medial–lateral
Artificial disc technology

direction, mainly to reduce implant extrusion. Due to the size of implant, an incision larger than the normal size for a discectomy procedure has to be made using a posterior approach. To minimize annulus dissection, a flap is made and sutured back after implantation.

RayMedica initiated the clinical trial of this implants in humans in Germany in 1996.36 A pilot United States Investigational Device Exemption study was begun in 1998. At the 1st Annual North American Artificial Disc Summit in October 1999, RayMedica reported that 101 patients had received its nucleus implant (unpublished data). Among patients in whom there was no implant extrusion, pain relief was achieved in the majority. However, the company reported that 17 of the first 101 patients did suffer implant extrusion.

More recently, in an effort to minimize implant extrusion and improve artificial disc technology with minimally invasive surgical methods, Advanced BioSurfaces (Minnetonka, MN) has developed a system that features a polymer formulation, a delivery balloon and balloon catheter, and a polymer-injection gun.13 The polymer is a polyurethane that can be cured in situ; it has strong mechanical properties comparable to those of other industrially cured medical polyurethanes. The company has conducted extensive in vivo and in vitro studies (unpublished data) to demonstrate the biocompatibility of this polymer during the course of developing this technology for application in knee surgery. In these studies it was shown that monomer-leachable elements are minimal. The balloon is made of a compliant material yet is very robust, which allows significant expansion during polymer injection to fill and conform with the cavity. Due to these characteristics, the device allows the surgeon to distract the disc space under controlled pressure. A biomechanical study in which a human cadaveric disc model was used demonstrated that this device is able to restore both disc height and disc modulus.

Disc Prosthesis for Thoracic Disc Replacement

Thus far, the majority of research-and-development efforts in artificial disc technology have been focused on the lumbar disc, with only a small percentage devoted to the cervical disc. There has been almost no significant research into prostheses for the thoracic disc. One of the undisputed reasons for this disparity is the relative rarity of thoracic disc disease and, therefore, thoracic disc surgery. This type of procedure accounts for less than 1% of all disc corrective surgeries.43 However, because thoracic disc diseases do nevertheless exist and many of them require the same type of surgical treatment as that used for lumbar disc diseases, it may be worthwhile to speculate on the applicability of artificial disc technology in treating thoracic disc disease.

Many different factors, such as structure, anatomy, function, degeneration mechanism, pathology, surgical approach, and complications of thoracic disc surgery, can affect the applicability of artificial disc technology in the treatment of thoracic disc disease.

Structurally, the composition of the discs in all three spinal regions is fairly similar. However, the thoracic spine has several unique characteristics. In this region the spinal cord is relatively small, yet the ratio of spinal cord to spinal canal is larger than that in other regions. Also, the blood supply is more variable in the thoracic spine, with a particularly vulnerable region between T-4 and T-9. The cross-sectional area of thoracic discs is larger than that of cervical discs, but smaller than that of lumbar discs. The variation in disc height does not follow the same trend along the spine, with midthoracic discs being the highest. The shape of thoracic discs is more circular, whereas that of lumbar and cervical discs is more elliptical.34 The thoracic spine has a kyphotic curvature, whereas both cervical and lumbar discs are more wedge shaped.34

Structurally, it should not be too difficult to transfer artificial disc technology (either total disc prosthesis or disc nucleus prosthesis) from the lumbar region to the thoracic region, apart from the problem of adjusting for differences in size and geometry. The similarity of shape across the entire length of the thoracic region should entail less emphasis on maintaining normal lordosis than when replacing lower lumbar discs. The greater disc height in the midthoracic region should, in general, make disc replacement easier than in the case of narrower discs. The structural similarity in disc composition should also allow nucleus replacement if annular integrity is not grossly compromised.

The major functions of the thoracic discs are similar to those of cervical and lumbar discs. The compressive load borne by the thoracic discs is lower than that borne by lumbar discs but higher than that borne by cervical discs. Without the support of the rib cage, one would probably expect that thoracic discs would be more susceptible to motion, given their position in the middle of the spinal column. Actually, thoracic disc joints are stiffer, with less range of motion in flexion–extension, than lumbar and cervical joints; this is due to the additional constraining force exerted by costovertebral articulations and the rib cage. The variation in range of motion in lateral bending throughout the thoracic region is not large. The range of motion of axial rotation of lower thoracic discs is similar to that of lumbar discs, whereas that of mid- to upper-thoracic discs is similar to that of cervical discs.37

The greater stability of the thoracic disc joint has implications for the applicability of artificial disc technology in treating thoracic disc disease. First, even in degenerative thoracic discs there is less joint instability than in degenerative lumbar and cervical discs. Therefore, in treating thoracic disc pathology, it is not as necessary to reestablish joint stability. Second, the superior stability contributed by nondisc elements such as the rib cage and costovertebral articulation can often better maintain disc joint stability even after disc surgery. For these reasons it is less common to fuse the thoracic disc after discectomy. Because one of the main objectives of artificial disc implantation is to maintain or restore disc function and stability, there will be less need for surgeons to place an artificial disc if they can treat the disease by using current techniques without significantly altering normal mechanical function.

Due to the rarity of thoracic disc disease, the exact mechanism of its degeneration has not been studied as thoroughly as that of lumbar disc degeneration. However, based on structural and functional similarity among discs in all three regions, it is believed that the mechanism of thoracic disc degeneration should be similar to that of lumbar and cervical disc degeneration. The most common result of thoracic disc disease is disc herniation, in which
the herniated disc material compresses spinal nerves and causes pain. As is the trend in the lumbar region, the majority of disc herniations in the thoracic region occur at the lower levels (T8–T12), probably due to the fact that the lower thoracic discs bear greater stress than the upper thoracic discs. Most herniations occur at either central or costotransversectomy; the total overall morbidity rate associated with symptomatic herniated tissue. Although symptomatic herniation at multiple levels has been reported in thoracic discs, it is not as common as it is in lumbar discs.

Discectomy is the surgical procedure most commonly used by surgeons to treat thoracic disc disease. Although autograft bone block placement is sometimes used with discectomy, interbody fusion is not a common procedure for this disease because of the thoracic spine’s relative stability. For this reason it would be more difficult to use an artificial total disc because it entails more radical surgery. It may be more effective to replace only the disc nucleus with a prosthesis, if it can be easily implanted using current discectomy methods.

Laminectomy, one of the early surgical treatments for thoracic disc disease, was found to produce disappointing results, due largely to inadequate decompressive effect of the posterior approach as opposed to direct removal of herniated tissue. Since then, many other surgical techniques have been developed to treat thoracic disc disease, including the following: lateral rachotomy, first described by Capener; modified costotransversectomy; transthoracic surgery; transthoracic surgery; lateral extracavitary surgery; and transthoracic and costotransversectomy procedures. These methods involve the same three approaches used for lumber and cervical discs surgery: posterior, anterior, and lateral. Therefore, it would be feasible to apply any artificial disc or disc nucleus developed for these three approaches to the thoracic disc.

Recently, minimally invasive spinal surgery techniques have also been applied to thoracic discectomy. Video-assisted thoracic surgery, which involves the same anterolateral approach entailed by open thoracotomy, allows surgeons access to the disc spaces, vertebral bodies, paravertebral soft tissues, spinal cord, spinal nerves, and sympathetic chain at all thoracic spine levels. Although the indications for VATS are the same as those for open thoracotomy, VATS offers the advantages of reduced postoperative pain, lower complication rates, and shorter recovery times. If this procedure continues to gain in popularity, artificial disc or nucleus devices that can be implanted by means of endoscopic procedures should have an advantage over those that must be implanted via open surgery.

The last but not the least important factor that can affect the applicability of artificial disc technology in treating thoracic disc disease is the potential risk it may add to currently used surgical procedures. Because artificial disc or disc nucleus devices are designed mainly to improve the long-term results of existing procedures that have known (and relatively low) risk factors and complication rates, it is vital that the use of these prosthetic devices not be allowed to increase significantly these potential liabilities, especially when the clinical benefits have not yet been proven.

Early laminectomy procedures were associated with high rates of paresis, paralysis, or even death. Coupled with poor clinical outcomes, laminectomy was essentially abandoned as a treatment for thoracic disc disease after 1960. The paresis and paralysis rate for anterolateral thoracotomy, one of the most popular procedures for thoracic disc disease, has been reported to be approximately 1%, with no deaths. The total overall morbidity rate associated with the transthoracic procedure—including pneumonia, atelectasis, pulmonary embolus, postoperative occult compression fracture, deep or superficial wound infections, bowel obstruction, and postoperative seizure—is only approximately 11%. Another common method of thoracic disc surgery is the lateral extracavitary approach, which is associated with approximately the same overall morbidity rate as transthoracic surgery, with no reported deaths. The morbidity rate for other less commonly used approaches, such as transpedicular and costotransversectomy, is also reported to be in the same range. Due to its brief history, the morbidity rate for VATS has been less well documented. Based on the small number of publications, the overall morbidity rate associated with VATS appears to be reasonably low, but only after a learning curve that is steeper than that required for open surgery.

Overall, the morbidity rate associated with current surgical techniques for thoracic disc disease appears to be low, with almost no deaths. Again, when applying artificial disc technology to the treatment of thoracic disc disease, it is important that the morbidity rate not be increased.

CONCLUSIONS

Although it is growing more and more clear that artificial disc or disc nucleus implantation will become the treatment of choice for lumbar and cervical disc disease, its application in treating thoracic disc disease may be further away. The rarity of thoracic disc surgery is the major reason for the lack of research and development activities related to artificial thoracic disc or disc nucleus technology. However, after the safety and efficacy of the technology have been demonstrated in clinical use for the treatment of lumbar and cervical discs, there should be few obstacles to prevent its transfer to the treatment of dis eased thoracic discs. Nevertheless, because the thoracic disc joint is more stable, the potential benefits of using an artificial nucleus may not be as great for thoracic as they would be for lumbar or cervical discs. To make artificial disc or disc nucleus implantation more acceptable as a treatment for thoracic disc disease, any added risk and cost associated with this new procedure must be reduced so that the benefit-to-cost and benefit-to-risk ratios are high enough to justify its clinical use.

Disclosure

Dr. Bao is vice president of research and development at Disc Dynamics, Inc., which is a spin-off of Advanced Biosurfaces, Inc. He was employed by Howmedica, Inc., when he pioneered research into the use of hydrogel material for nucleus prostheses.

References


Q. B. Bao and H. A. Yuan

6 Neurosurg. Focus / Volume 9 / October, 2000
Artificial disc technology


Manuscript received August 14, 2000. Accepted in final form September 13, 2000.

Address reprint requests to: Qi-Bin Bao, Ph.D., Disc Dynamics, Inc., 5909 Baker Road, Suite 550, Minnetonka, Minnesota 55345. email: cbao@advbiosurf.com.