EXPANDABLE INTERVERTEBRAL SPACERS

Laterally expanding vertebral spacer devices are provided for repairing damaged vertebral discs. The vertebral spacer devices maintain the height of a distracted vertebral disc space while providing stability to the spine. In one form of the invention, a vertebral spacer device (50) is provided with a first arm (52) movably coupled to a second arm (54). The first and second arms (52, 54) are laterally expandable from a first width for insertion into the disc space to a second width after insertion into the disc space. The first and second arms also define a cavity (66) therebetween for placement of bone growth material.
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EXPANDABLE INTERVERTEBRAL SPACERS

BACKGROUND OF THE INVENTION

The present invention is directed to implantable devices for stabilizing the spine. Specifically, the invention concerns intervertebral spacers expandable from a reduced size insertion configuration to an expanded size spacing configuration.

Intervertebral discs, located between the end plates of adjacent vertebrae, stabilize the spine, distribute forces between vertebrae and cushion vertebral bodies. An intervertebral disc may deteriorate due to trauma, aging or disease resulting in pain or discomfort to a patient. One common procedure for relief of patient discomfort is a discectomy, or surgical removal of a portion or all of an intervertebral disc. Often, this is followed by implantation of a device between adjacent vertebrae to maintain or restore disc space height. Typically, implantation of such a device is also intended to promote bony fusion between the adjacent vertebral bodies.

One limitation on the size of a device inserted into the disc space is the size of the opening through surrounding tissue that is available to gain access to the disc space. From a posterior approach to the spine, the dura and nerve roots must be mobilized to gain access to the disc space. Similarly, from an anterior approach, the aorta and vena cava must be mobilized to gain access to the disc space. Such mobilization is often limited by the anatomical structures, thus resulting in a relatively small access site. Removal of additional bone to enlarge an entrance to the disc space may weaken the joint between two adjacent vertebra.

Moreover, excessive retraction of vessels and neural structures to create a large access opening may damage these tissues. Thus, prior procedures have been limited to placing a first device passable through the available opening on one side of the spine and mobilizing the tissue or vessels to place another similar implant on
the opposite side of the spine. Each implant being limited in size by the available access site.

Thus, there remains a need for implantable devices that have a reduced size insertion form and are expandable in the disc space to a larger size for enhancing spine stability and facilitating immobilization via bony fusion.
SUMMARY OF THE INVENTION

The present invention contemplates an intervertebral spacer device that has a reduced size configuration for insertion into a disc space and an expanded size configuration to maintain the spacing of the disc space. In one aspect of the present invention, the device includes a pair of arms each having a first end and a second end, the arms being movably coupled at their first ends. When the arms are positioned adjacent one another, the device is in a reduced size configuration for insertion into the disc annulus. The device is laterally expandable in the disc space to an expanded configuration by moving the pair of arms about the first ends in order to increase the dimension of the device perpendicular to the longitudinal axis of the spine while maintaining the inter-space distraction. Preferably, the expanded device creates a cavity that may be filled with bone or bone substitute material for purposes of promoting fusion between the adjacent vertebrae. Preferably, the height of the device in the reduced size configuration is substantially the same as the height in the expanded configuration, with the expanded configuration providing an increased base of support.

In another embodiment of the present invention, the first and second arms each have laterally extending portions extending therefrom that cooperate to engage the first and second arms to one another. Preferably, each of the laterally extending portions defines a plurality of serrations, wherein the serrations of one laterally extending portion of the first arm cooperate in interdigitating fashion with serrations of the corresponding laterally extending portion of the second arm. In one preferred embodiment, the laterally extending portions are provided at the first and second ends of each of the arms. In another preferred embodiment, the pair of arms are pivotably coupled at their first ends, and laterally extending portions are provided at the second ends.

In still a further embodiment, the pair of arms are flexibly attached such that they are compressible into a first smaller configuration and laterally self-expand to a second larger configuration. In one such embodiment, the arms are
interconnected by a flexible hinge portion at one end of each arm. In another embodiment, each arm is flexibly connected to a first end portion and an opposing second end portion to form a substantially rectangular shape having flexible side walls. Preferably, the side walls are biased to assume the second larger configuration.

One object of the present invention is to provide a vertebral spacer device that is capable of insertion in a smaller form and laterally expandable within the disc space to an enlarged configuration for supporting the spine.

Other objects and advantages of the present invention will be readily discerned upon consideration of the following written description and accompanying figures.
BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1. is a perspective view of one embodiment of a vertebral spacer device according to the present invention.

Fig. 2 is a top view of the vertebral spacer device of Fig. 1.

Fig. 3 is a left end view of the vertebral spacer device of Fig. 2.

Fig. 4 is a right end view of the vertebral spacer device of Fig. 2.

Fig. 5 is an elevational view of the vertebral spacer device of Fig. 2.

Fig. 6 is a top view of the vertebral spacer of Fig. 1 shown in an expanded position.

Fig. 7 is an anterior-posterior view of a pair of vertebrae having a collapsed disc space therebetween.

Fig. 8 is an anterior-posterior view of the vertebrae of Fig. 7 showing the vertebrae after distraction of the disc space.

Fig. 9a is a partial cross-sectional top view of the vertebrae of Fig. 8 with the vertebral spacer device of Fig. 1 in an expanded position between the vertebrae.

Fig. 9b is a partial cross-sectional top view of a vertebral body as shown in Fig. 8, with a pair of vertebral spacer devices according to Fig. 1 inserted from a bilateral posterior approach.

Fig. 9c shows the vertebral spacer devices of Fig. 9b in an expanded configuration.

Fig. 10 is a side view of an insertion tool useable with the vertebral spacer devices of the present invention.

Fig. 10a is an end view of the insertion tool of Fig. 10.

Fig. 11 is a perspective view of an expansion tool useable with the vertebral spacer devices of the present invention.

Fig. 12 is a perspective view of an element of Fig. 11.

Fig. 13 is a perspective view of an alternate embodiment vertebral spacer device according to the present invention.

Fig. 14 is a top plan view of the vertebral spacer device of Fig. 13 in an unexpanded position.
Fig. 15 is a top plan view of the vertebral spacer device of Fig. 13 in an expanded position.

Fig. 16 is a cross-sectional view of the vertebral spacer device of Fig. 14 taken along line 16-16.

Fig. 17 is a partial cross-sectional side view of an insertion tool device usable with the vertebral spacer device of Fig. 13.

Fig. 18a is a perspective view of another embodiment of a vertebral spacer device according to the present invention.

Fig. 18b is a perspective view of the vertebral spacer device of Fig. 18a constrained within a delivery system.

Fig. 19a is a top view of a laterally expandable implant according to another embodiment of the present invention.

Fig. 19b is a top view of the implant of Fig. 19a in a compressed configuration.

Fig. 19c is a side view of the implant of Fig. 19a.

Fig. 20a is a perspective view of another embodiment of a vertebral spacer device according to the present invention.

Fig. 20b is a perspective view of the space device of Fig. 20a without the ratchet mechanism.

Fig. 21 is a perspective view of yet another embodiment of a vertebral spacer device according to the present invention.

Fig. 22 is a perspective view of yet another embodiment of a vertebral spacer device according to the present invention.

Fig. 23 is a plan view of an expansion tool usable with the vertebral spacers of Figs. 20-23.

Fig. 23a is a fragmentary perspective view of a portion of the insertion tool device of Fig. 23.

Fig. 24 is a perspective view of another embodiment of a vertebral spacer device according to the present invention shown in a collapsed position.

Fig. 24a is a perspective view of the vertebral spacer device of Fig. 22 shown in an expanded position.
Fig. 25 is a perspective view of yet another embodiment of a vertebral spacer device according to the present invention.
DESCRIPTION OF THE PREFERRED EMBODIMENTS

For the purposes of promoting an understanding of the principles of the invention, reference will now be made to the embodiments illustrated in the drawings and specific language will be used to describe the same. It will nevertheless be understood that no limitation of the scope of the invention is thereby intended, any alterations and further modifications in the illustrated devices, and any further applications of the principles of the invention as illustrated therein being contemplated as would normally occur to one skilled in the art to which the invention relates.

In accordance with one embodiment of the invention, a vertebral spacer device 50 is depicted in Figs. 1-6. Device 50 includes a first lateral arm 52 and a second lateral arm 54. First arm 52 includes a first end 60 and an opposite connection end 61. Second arm 54 includes a first end 62 and an opposite connection end 63. First arm connection end 61 is fixedly coupled to second arm connection end 63 via connection pin 58 extending through a bore 59 defined through connection ends 61 and 63. Bore 59 extends transverse to the longitudinal axis 53 of spacer 50.

First arm 52 and second arm 54 each define a portion of a top bone engaging surface 56 adapted to engage a vertebral body and a portion of bottom bone engaging surface 57 substantially identical to top bone engaging surface 56. When first arm 52 and second arm 54 are in an opened position, as shown in Fig. 6, a central cavity 66 is defined therebetween. Cavity 66 is adapted to receive a graft or bone-growth inducing material therein.

Referring now to Figs. 3-6, the vertebral spacer device 50 is illustrated and described below in further detail. Connection end 63 of second arm 54 is fixedly coupled to connection end 61 of first arm 52 via connection pin 58 extending through bore 59. However, it should be understood that any type of connection mechanism contemplated herein, provide the principles of the current invention are adhered to. As an example, but without limitation, an alternative connection mechanism may be a hinge between and fixedly engaging first arm 52 and second
arm 54 to allow pivotal movement therebetween. Alternatively, first and second arms may be integrally formed of a flexible material, thereby permitting movement at the connection point.

First end 60 of first arm 52 and first end 62 of second arm 54 each define a corresponding socket portion 64 and 65, respectively. When device 50 is in a first closed position, as shown in Fig. 4, socket portions 64 and 65 define a socket for 67 for receiving a driving tool, which will be described more fully below. End 60 also includes an internally threaded bore 68 defined by device 50. Threaded bore 68 is provided to receive an attachment portion of an insertion tool configured for manipulation of device 50 into and out of a disc space.

It should be noted that in the illustrated embodiment first arm 52 and second arm 54 are configured such that the top bone engaging surface 56 defined on each of the arms 52 and 54 extends in a substantially uniform horizontal plane to make the bone engaging surface 56 substantially planar in a first plane. The bottom bone engaging surface 57 defined by arms 52 and 54 also extends in a substantially uniform horizontal plane making the bottom bone engaging surface 57 substantially planar in a second plane. In a preferred embodiment, the first and second planes are generally parallel and separated by a height. Preferably, the height between the first and second planes is substantially constant between the closed position of Fig. 2 and the open positions of Figs. 6 and 9. Thus, the disc space height during insertion may be substantially maintained in the expanded position.

Device 50 may be positioned in a closed position forming a reduced size configuration shown in Figs. 2-4. Preferably, arms 52 and 54 are closely adjacent in this position, although the exact arm positioning may vary depending on the application. In the closed position, device 50 has a lateral width $W_1$ extending transverse to longitudinal axis 53 of the device.

Device 50 may be positioned in an open position forming an expanded size configuration as shown in Figs. 1, 6 and 9. The extent of distance between first arm 52 and second arm 54 may be varied depending on the expanded size desired. In the open position, device 50 may have at least a lateral width $W_2$ extending
transverse to longitudinal axis 53 of the device. Lateral width $W_2$ being greater than lateral width $W_1$. As shown in Fig. 9a, the lateral width in the expanded configuration may be substantially greater than width $W_1$. This expanded width provides a much wider base of support than the device does in the closed position. The wider base of support provides greater stability of the device.

Referring to Fig. 9b, there is shown a vertebral body with two laterally expandable implants according to the present invention. Implants 70 and 72, slightly smaller versions of device 50, have been inserted through posterior openings 74 and 76, respectively, into the disc space in their reduced size insertion form. It will be understood that this placement is approximately in the same position in the disc space into which known devices may be placed. In much the same manner that chairs, such as tall stools, are subject to tipping if the legs are too close, implants may also be subject to tipping if they lack a sufficiently wide base support area. However, referring to Fig. 9c, the present invention permits each of devices 70 and 72 to be expanded in the disc space to a greater width, thereby increasing the total width of the base of support. Moreover, material G promoting bone growth may be placed in the cavity between the arms and around the exterior of the implants.

The bone engaging surfaces 56 and 57 of device 50 are configured to provide an even distribution and transfer of the load from the upper vertebral body through the integral side walls of device 50 to the lower vertebral body. In a preferred embodiment, the endface plates 56 and 57 are knurled to provide frictional engagement between the vertebrae and the device 50. While knurling is shown as one configuration for the bone engaging surface, other configurations may be utilized. For example, but without limitation, grooves may be formed on the upper and lower bone engaging surfaces extending transverse to longitudinal axis 53 to resist expulsion. More specifically, arcuate grooves may be formed having a radius of curvature originating at pin 58 to follow the arc of the arms as they are expanded in the disc space to form the expanded open position shown in Fig. 6.
Referring to Figs. 7-9, a spinal segment with vertebrae V₁ and V₂ is illustrated to briefly describe a surgical procedure in which device 50 may be employed. More specifically, in Fig. 7 a damaged or diseased spinal segment is shown without the device 50. D₁ represents a degenerated or damaged disc between vertebrae V₁ and vertebrae V₂ that has resulted in the collapse of the disc space between the vertebrae. Vertebrae V₁ and V₂ form part of a spinal column having a longitudinal axis L extending therethrough.

In Fig. 8, the vertebrae V₁ and V₂ are shown distracted such that the disc space is restored to approximately its normal height, represented by distracted disc space D₂. Tensioning of annular structures that extend between D₁ and D₂ promotes disc stability. Also shown is an opening A made in the annulus fibrosus that may be created by the surgeon by an annulotomy or disectomy surgical procedure to gain access to the disc space from an anterior approach. As known in the art and not further described herein, the adjacent end plates of V₁ and V₂ may be prepared to promote bone fusion therebetween and accept device 50. Device 50 is inserted through opening A while in the reduced size configuration (as shown in Figs. 2 through 4). Once inserted into the disc space, the device 50 is laterally expanded to expanded size configuration (as shown in Figs. 1 and 6) by moving first arm 52 in relation to second arm 54 in the disc space. The lateral expansion of device 50 increases the lateral dimension of device 50 in a direction transverse to longitudinal axis L, while maintaining the height of distracted disc space D₂. In Fig. 9 the device 50 is shown in plan view inserted into D₂ between vertebrae V₁ and V₂ through opening A. It will be understood that use of the laterally expandable implant according to the present invention limits the amount of mobilization of overlapping vessels and permits insertion of an implant having a much wider spacing configuration than would otherwise be implantable with a non-expanding implant.

The expanded configuration of device 50 creates cavity 66 that may then be filled with a bone graft material or bone-growth inducing material G for the purposes of promoting fusion between vertebrae V₁ and V₂. The graft material G also helps to maintain the device 50 in the laterally expanded configuration. As
can be seen in Fig. 9, the expanded device 50 is larger than the opening A made through the annulus fibrosus. Thus, in addition to the knurled endface plates 56 and 57, the remaining annulus fibrosis may also act to limit displacement of device 50 from the disc space. While the device has been inserted with the wider end adjacent opening A, it is contemplated that the connection end may be disposed adjacent the opening. For this use, a biasing element, such as a spring, may be disposed between the arms to urge them to the expanded condition.

Figs. 7-9c illustrate two methods for inserting laterally expandable devices into the disc space D2. The present invention also contemplates the use of additional methods as known in the art for inserting interbody fusion implants. For example, more than one vertebral spacer device may be inserted through the same opening A. For example, a first device 50 could be inserted and laterally expanded, and packed with bone graft material. Then a second device may be inserted in the disc space and between the arms of the first device. The second device may be laterally expanded and packed with bone graft material G.

Referring to Figs. 9a and 9b, there is shown a vertebral body with two implants positioned in the disc space. In this procedure, bilateral access to the disc space is achieved by posterior openings 74 and 76. It will be understood that the size of openings may be limited by the amount of dural compression that may be safely achieved, nerve root location and the amount of bone removed adjacent the disc space. Devices 70 and 72 are inserted via opening 74 and 76, respectively. The devices are inserted into the disc space in the reduced size configuration. Once disposed in the disc space, devices 70 and 72 are expanded and graft material is positioned in the cavity formed between the arms. Preferably, as shown in Fig. 9c, material may be positioned between the implants before one or both are expanded to provide a further area for bone growth. While a device according to Fig. 1 has been shown for the purposes of illustrating the methods of insertion, it is contemplated that the other embodiments disclosed herein may be inserted in a like manner.

Referring now to Figs. 10-12, various instruments useful for insertion and lateral expansion of device 50 are shown therein. The insertion tool 260 of Fig. 10
is useable for insertion of device 50 into the disc space. Insertion tool 260 includes a handle portion 262, a threaded stem portion 266, and rod 264 extending between handle 262 and threaded portion 266. A sleeve 268 is slidably disposed about the stem 264. Sleeve 268 includes protrusion 270 extending therefrom and adapted to engage cavity 67 in device 50. While not illustrated, device 260 may include a stop mechanism operable to prevent sliding of sleeve 268 about rod 264 after device 50 is engaged thereto.

To use insertion tool 260 to insert the implant device 50, threaded portion 266 threadedly engages device 50 via threaded bore 68. Once the device 50 is threadedly engaged to insertion tool 260, sleeve 268 may be slid down rod 264 toward the device 50 until protrusion 270 resides within cavity 67. Rod 264 and protrusion 270 prevent rotation between device 50 and insertion tool 260 during insertion. The vertebral spacer device 50 may then be inserted into a prepared disc space using the insertion tool 260. Once device 50 is placed in the disc space, sleeve 268 may be retracted towards handle 262 to disengage protrusion 270 from cavity 67. Threaded stem portion 266 may then be removed from threaded bore 68. Alternatively, if it is desired to remove the device 50 from the disc space after initial insertion or to reposition the device 50 within the disc space, the threaded stem portion 266 allows the device 50 to be withdrawn or repositioned. It is contemplated herein that insertion of device 50 into the disc space via insertion tool 260 is accomplished with device 50 in a closed position, as shown in Fig. 2.

Once the device 50 is inserted into the desired position in the disc space, first arm 52 and second arm 54 may be laterally expanded to increase the lateral dimension of device 50 with respect to spinal longitudinal axis L in order to stabilize the spinal column and fill a larger portion of the disc space. In a preferred embodiment, each bone engaging surface 56 and 57 includes a beveled edge, around the perimeter of device 50. The beveled edge facilitates insertion between adjacent vertebrae and eases expansion in the disc space.

Fig. 11 illustrates one type of driving tool 250 operable to at least initially laterally expand device 50 to a laterally expanded configuration. Driving tool 250 includes T-handle portion 254, a square driving end 258 adapted to engage cavity
67, and a hollow tube 256 extending between handle portion 254 and driving end 258. In order to laterally expand device 50, driving tool 250 is rotated via the T-handle 254 with driving end 258 disposed within cavity 67. Rotation of driving end 258 causes first arm 52 and second arm 54 to move laterally with respect to one another in a manner that laterally expands the arms 52 and 54 of device 50.

In order to further laterally expand first arm 52 and second arm 54, a spreader 280 as shown in Fig. 12 may be used in conjunction with tool 250. Spreader 280 includes a first end 282, a wedge portion 286, and stem 284 extending therebetween. As shown in Fig. 11, spreader 280 may be disposed within hollow tube 256 and advanced beyond its distal end to more fully expand the device. Wedge portion 286 may be placed between first arm 52 and second arm 54. A force applied to first end 282 drives wedge portion 286 between arms 52, 54 in order to further laterally expand the device 50.

While the above-described spreader is disclosed as a preferred embodiment, it is contemplated that other instruments may be used to expand the device without deviating from the scope of the invention. Specifically, spreader 280 may be used may be used alone to laterally spread the expandable device.

As shown in Figs. 6 and 9, when device 50 is in a laterally expanded position, a cavity 66 is formed between first arm 52 and second arm 54. A graft material G may then be placed or packed into cavity 66. The graft material G could be cancellous bone or bone chips, or a suitable bone graft substitute material known to those skilled in the art. One advantage of the device 50 is that it allows bone graft material G to be placed at or near the central portion of the vertebrae while the expandable spacer engages more lateral portions of the vertebra. This central portion is known to be highly vascular and biologically active, so that it is an excellent location for bone graft incorporation and fusion. In addition, bone-growth enhancing materials may be introduced with the graft material to enhance initial and ultimate fusion of the vertebrae \( V_1 \) and \( V_2 \).

It should be appreciated that device 50 may be delivered to the disc space for insertion through a cannula employed in a minimally-invasive surgical technique. Device 50 is sized for placement through the cannula in its unexpanded
configuration. Once positioned in the disc space, the lateral dimension of the device is increased by expanding the first and second arms 52, 54 as described above. Other surgical techniques for insertion are contemplated, for example, open surgical procedures with direct access to the spine. Device 50 thus allows minimization of the size of the entry into the disc space and the resulting damage to tissue surrounding the surgical site. Further, the reduced size configuration of the implant permits insertion of a relatively large spacer where anatomical features, such as the dura, nerve roots or blood vessels, would have prevented placement of a larger, non-expanding sized spacer.

Referring now to Figs. 13-16, another embodiment of the present invention is illustrated. The expandable vertebral spacer 80 includes a first arm 82 having a distal end 90, and a second arm 84. Second arm 84 is movable coupled to main body portion 82 via hinge portion 98. First arm 82 is provided with a tapering guide 88 protruding therefrom as it extends from hinge portion 98 towards distal end 90. Guide 88 is received within a recess 86 defined in second arm 84. Vertebral spacer 80 also defines tool receiving opening 99 defined in hinge 98.

Tool receiving opening 99 is configured to have an internal thread to accommodate an insertion tool, such as tool 300 illustrated in Fig. 17.

Second arm 84 includes a locking arm 94 adjacent its distal end that is integrally formed with laterally expandable portion 84 via locking arm hinge portion 95. Locking arm 94 is configured to be positioned adjacent distal end portion 90 in the closed position shown in Fig. 14. In the closed position the device is in a reduced size configuration suitable for insertion. In this configuration, device 80 has a lateral width W₁ extending transverse to the longitudinally axis of the device. Preferably, spacer 80 is formed of an at least partially resilient material and distal end portion 90 may be biased toward cavity 85. In this configuration the arms tend to move to the locked position once the spacer is sufficiently expanded. Distal end portion 90 includes a catch 92 formed thereon, and locking arm 94 includes a catch-receiving portion 96. When the device 80 is laterally expanded to a second lateral position, as shown in Fig. 15, locking arm hinge 95 urges locking arm 94 towards distal end portion 90 until
catch-receiving portion 94 engages catch 92. Catch 92 prevents displacement of expandable portion 84 towards main body portion 82 after the device 80 is inserted in the disc space. The device 80 is then held in the expanded position, and cavity 85 may be packed with bone growth material through opening 99. Further openings for bone ingrowth or bone growth material packing may be provided. In the laterally expanded configuration of Fig. 15, device 80 has a maximum lateral width $W_4$, width $W_4$ being greater than $W_3$.

It should be noted that the device 80 defines a top vertebral bearing surface 97 and a bottom vertebral bearing surface 93. The bearing surfaces 93 and 97 are composed of the surfaces provided on first arm 82, second arm 84, and hinge 98. In a preferred embodiment, bearing surfaces 93 and 97 are spaced apart a height that remains relatively constant from the closed to expanded positions. The bearing surfaces contact the adjacent vertebrae endplates to provide an even distribution of loads through the endplates and balanced loading conditions. While not shown, it will be understood that these surfaces may include roughening to inhibit expulsion.

It is contemplated that devices according to the present invention may be manufactured from bio-compatible materials having at least some flexibility without fracture. Further, it is anticipated that portions of bone may be used provided the hinge points have been at least partially demineralized to provide flexibility. Demineralization of bone is known in the art and will not be described further herein. More preferably, device 80 is formed from material having a degree of resiliency tending to urge locking arm 94 into the locking position with the catch 92 engaged with catch-receiving portion 94. Such materials may include, but are not limited to, stainless steel, shape memory alloys, composites and plastics. Moreover, while flexible hinge portions have been disclosed, it will be understood that hinge pin and channel connections may replace the flexible hinges without deviation from the spirit of the invention. Optionally, a biasing mechanism, such as a spring, may be placed between the arms to urge the device to the expanded configuration.
The present invention also contemplates an instrument for inserting and expanding an implant according to the present invention. Referring now to Fig. 17, an insertion tool 300 is illustrated. Tool 300 includes a hollow outer sleeve 302 that receives a portion of an inner sleeve 304. Inner sleeve 304 defines connecting portion 322 that engages mating portion 320 of outer sleeve 302. In the illustrated embodiment, inner sleeve 304 is threadedly received within the outer sleeve 302. Inner sleeve 304 further defines an opening 324 therethrough for receiving rod 310. Inner sleeve 304 also includes a pair of movable arms 306 and 308 having gripping portions 309 and 311, respectively, configured for holding device 80 during insertion. In order for arms 306 and 308 to grip the device 80, outer 302 is moved with respect to inner handle 304 such that inclined portion 318 of outer sleeve 302 urges gripping portions 309 and 311 of arms 306 and 308 against device 80. In the illustrated device 300, this accomplished by rotating outer handle 302 about a thread on connecting portion 322 towards the device 80.

Once device is engaged by gripping portions 309 and 311, it may be inserted into the disc space. After insertion of device 80 to the desired location, rod 310 is operable to laterally expand device 80. Rod 310 has a handle portion 312, and opposite a threaded portion 314, and a shaft 313 extending therebetween. In a preferred embodiment, shaft 313 has a distal end 316 that is beveled to engage the inclined surfaces 87 and 89 of first arm 82 and second arm 84, respectively. Handle 310 may be engaged with device 80 during insertion into the disc space via threaded engage with tool receiving opening 99. The threaded engagement between threaded portion 314 and the device 80 allows the device 80 to be positioned within the disc space. In order to position the device 80 to its expanded configuration, mechanism 310 is threaded within receiving portion 99 in order to urge distal end 316 against surfaces 87 and 89 to laterally expand device 80 to the expanded or second lateral configuration as shown in Fig. 15.

While the above-described spacer embodiments of Figs. 1 and 13 have been described as having a first arm and a second arm movable coupled, it will be understood that the invention contemplates a main body portion and laterally expandable portion movably coupled thereto. Specifically, while first arm and
second arm may simultaneously move laterally to form the expanded configuration, it is contemplated that one arm may remain stationary while the other arm moves. Moreover, the device may be formed such that the device includes a stationary main body with one or more movable laterally expandable portions movable to give the device both a reduced size configuration and a laterally expanded size configuration.

Referring now to Figs. 18a and 18b, another embodiment of the present invention is illustrated. Vertebral spacer device 100 includes a pair of lateral arms 102 and 103 extending between a distal end 106 and a proximal end 108. The device 100 includes a top vertebral bearing surface 112 and an identical bottom vertebral bearing surface (not shown). A central cavity 114 is formed between the lateral arms 102. The device 100 also includes openings 104 and 105 defined by lateral arms 102 and 103, respectively. Openings 104 and 105 permit communication between the interior and exterior of the device and reduce the material in walls 102 and 103, thereby increasing the flexibility of device 100. Device 100 also includes at least one insertion tool opening 110 formed in proximal end 108. Preferably, opening 110 is threaded to receive a correspondingly threaded insertion tool (not shown).

The embodiment of Fig. 18 is preferably formed of a resiliently flexible material. Such materials may include, without limitation, bio-compatible metals (including shape memory alloys), composites, and plastics. In a preferred embodiment, the device 100 is expanded and contracted by making the device 100 from a shape memory material, such as nitinol, exhibiting super elasticity and/or temperature induced shape memory. The device 100 is initially formed in a laterally expanded or second position. In order to insert the device 100 through a small opening and into the disc space, it is contracted to a first lateral position by applying a force to lateral arms 102 and 103 in the direction indicated by the arrows "R". Thus, the device is laterally compressed into a smaller sized configuration. Often, the device will experience some elongation, as shown by dimension "l". When the device is contracted, as shown in Fig. 19, it may be inserted through a tubular delivery system, such as the cannula 120. Once the
device is inserted in the disc space, it is no longer confined by the cannula 120, and it self-expands laterally to a second position within the disc space approximating its pre-insertion condition. Cavity 114 may be filled with bone growth material delivered through opening 110. Cavity 114 may also be partially loaded with bone growth material prior to insertion. It is also contemplated herein that device 100 may be inserted into the disc space without use of cannula 120, such as by an open surgical procedure. Temporary compression may be achieved by an external device such as, but without limitation, pliers adapted to compress the implant.

Figs. 19a through 19c illustrate a further embodiment of a laterally expandable spacer according to the present invention. Spacer 121 includes arms 122 and 123 connected by a flexible portion. Arm 122 terminates in an end wall 125 and arm 123 terminates in an end wall 126. As shown in Fig. 19b, the respective lengths of arms 122 and 123 allow end wall 126 to nest within end wall 125.

Spacer 121 is preferably formed of a flexible and resilient material. The spacer is in a relaxed form in the expanded configuration of Fig. 19a having a lateral width W_6. Width W_6 is decreased to lateral width W_5 by the application of compressive force on arms 122 and 123 urging end walls 125 and 126 towards one another. Preferably, spacer 121 self-expands from the reduced size configuration of Fig. 19b to the expanded configuration of 19a. Preferably W_6 is approximately twice W_5, although a greater or lesser amount of lateral expansion may be provided. Preferably, spacer 121 is formed of a fiber reinforced polymer composite. The fibers, shown by the parallel shading marks in Figs. 19a through 19c, extend generally parallel to the length of side walls 122 and 123. It will be understood that this arrangement of fibers provides a degree of flexibility between the arms but resists compression from the upper to lower surfaces engaging the vertebral bodies.

Referring to Fig. 20a, another embodiment of a vertebral spacer device is illustrated. Vertebral spacer device 130 includes a first arm 132 and a second arm 134 fixedly connected via hinge portion 136. In this embodiment, hinge portion 136 is integrally formed with first arm 132 and second arm 134. First arm 132
includes first lateral extending portion 138, and second arm 134 includes a second laterally extending portion 140. First laterally extending portion 138 includes first serrations 139 and second laterally extending portion 140 includes corresponding second serrations 141 disposed adjacent first serrations 139. Serrations 139 and 141 cooperate in interdigitizing fashion to restrain lateral contracting of the first arm 132 with respect to the second arm 134. The device 130 also includes tool opening 142, which allows engagement of device 130 to insertion and/or expansion tools. As previously disclosed, opening 142 may be threaded to receive a corresponding threaded tool. As with earlier disclosed embodiments, device 130 also defines a cavity 146, and includes substantially planar vertebral bearing surfaces 148 and 149 for engaging respective end plates of adjacent vertebrae.

The device 130 is shown in a contracted position, and once inserted the device may be expanded by applying a force in the direction of the arrows “R”. The interdigiting serrations 139 and 141 must yield sufficiently to allow movement of first arm 132 with respect to second arm 134, while maintaining the separation of arm 132 and second arm 134 when the force is removed.

Fig. 20b represents a modified version of Fig. 20a lacking serrations 139 and 141. Preferably, spacer 130 is formed of a flexible material that may be plastically deformed. Thus, force applied to arms 132 and 134 to expand the device plastically deforms hinge portion 136. Plastic deformation of hinge portion 136 maintains the device in the expanded condition.

Fig. 21 illustrates another embodiment of the vertebral spacer device of the present invention. Device 150 includes a first arm 152 and a second arm 154. The term arm as used throughout the disclosure is used broadly to define sections and portions of devices. Arms may not necessarily move within a device configuration. First arm 152 includes a first extension 158 and a second extension 155. Second arm 154 includes third extension 156 and fourth extension 166. First arm 152 is sized to receive extensions 156 and 166 within extensions 158 and 155. First extension 158 defines first serrations 159 and second extension defines second serrations 161. Third extension 156 defines third serrations 157 and fourth extension 166 defines fourth serrations 168. First serrations 159 and third
serrations 157 cooperate in interdigitating fashion in cooperation with interdigitating engagement of second serrations 161 and fourth serrations 168 to maintain lateral spacing between first arm 152 and second arm 154. Device 150 also defines an upper vertebral engaging surface 164, and an identical lower vertebral engaging surface, and tool openings 160. Device 150 also defines a cavity 162, which may be filled with bone growth material. Once the device 150 is inserted into the disc space, it may be expanded by applying force in the direction indicated by arrow “R” to move first arm 152 with respect to second arm 154.

Referring now to Fig. 22, yet another embodiment of a vertebral spacer device in accordance with the present invention is illustrated. Device 170 includes a first arm 172 and a second arm 174. First arm 172 includes first a pair of extensions 176 and second arm 174 includes a pair of extensions 178. Extensions 176 include projections 175, and extensions 178 define receptacles 177. Projections 175 are configured to be placed within a respective one of receptacles 177. Projections 178 define first serrations 179 thereon, and receptacle 177 define second serrations 181 thereon. First serrations 179 and second serrations 181 cooperate in interdigitating fashion to resist displacement at first arm 172 with respect to second arm 174. However, first serrations 179 and second serrations 181 yield sufficiently to allow lateral expansion of the device 170. Device 170 includes an upper vertebral engaging surface 186 and an identical lower vertebral engaging surface. Arms 172 and 174 define a central cavity 182 for receiving bone growth material.

A tool 340 for expanding the devices illustrated in Figs. 20-22 is illustrated in Figs. 23 and 23a. Tool 340 includes a first lever 350 pivotably coupled to a second lever 360 by pin 346. First lever 350 includes a first handle portion 351 pivotably coupled to a first extension 353 via pin 352. Second lever 360 has a second handle portion 361 pivotably coupled to a second extension 363 via pin 362. Extensions 353 and 363 are pivotable engaged via pin 348. Handle 340 also includes ratchet mechanism 342 coupled to one of the handle portions 351, 361. In the illustrated embodiment, ratchet mechanism 342 is coupled to second handle portion 361 via pin 344. Ratchet mechanism 342 has teeth 346 for engaging first
handle portion 351. Ratchet mechanism 342 is operable to maintain the relative spacing between handle portions 351 and 361 when engaged thereto.

First extension 353 has a first engagement portion 354 and second extension 363 has a cooperable second engagement portion 364 located at respective distal ends of each extension 353 and 363. First engagement portion 354 includes a first coupling 356, and second engagement portion 364 includes a second coupling 366, each for coupling respective lever arms 350 and 360 to a vertebral spacer device, such as device 150 illustrated in Fig. 21. Couplings 356 and 366 extend through a corresponding one of tool openings 160 to engage the device 150. As shown in detail in Fig. 23a with respect to first engagement portion 354, first and second couplings 356 and 366 each include a first and second head 358 and 368 and a first and second recess 359 and 369, positioned between first and second extensions 353 and 363, respectively. The first and second recesses 359 and 369 are configured to receive a portion of the arms 152 and 154 therein to allow head 358 and 368 to engage the device 150. The device 150 may then be laterally expanded or contracted as needed by manipulation of first and second lever arms 351 and 361. Tool 340 may then be uncoupled from device 150 by withdrawing the first and second coupling 356 and 366 from device 150.

Referring now to Figs. 24-24a, another embodiment of a vertebral spacer device according to the present invention is illustrated. Device 190 includes first arm 192 and second arm 194. First arm 192 is pivotally coupled to second arm 194 via sidewalls 196 extending therebetween. In the illustrated embodiment, two sidewalls 196 are shown with one at the proximal end of the device 190 and the other sidewall 196 at the distal end of device 190. The first and second arms 192 and 194 are engaged to sidewalls 196 via hinge pins 197. The device 190 also defines an upper vertebral engaging surface 200 and a lower vertebral engaging surface, and tool insertion openings 198. In one embodiment, the device 190 is provided with ridges 204 extending from vertebral engaging surfaces for engaging the adjacent vertebral endface plates. A central cavity 202 for receipt of bone growth material is defined by the sidewalls 96, first arm 192, and second arm 194.
As shown in Fig. 24, the device 190 is collapsible to a first reduced size configuration for insertion into the disc space having a lateral width \( W_7 \). Once the device is inserted, it may be pivoted about hinge portions 197 to an expanded position having greater width \( W_8 \) as shown in Fig. 24a. The device may be expanded by a tool inserted through one or more of the openings 198. Bone growth material may be placed in cavity 202 through openings 198.

Another embodiment of the vertebral spacer device of the present invention is illustrated in Fig. 25. The vertebral spacer device 210 includes a first arm 212 and a second arm 214. First arm 212 includes a first laterally extending portion 215, and second arm 214 includes a second laterally extending portion 216. Second arm 214 also includes offset portion 217 extending to engage first arm 212 at connection 226. Preferably, connection 226 is a hinge-type connection. The device 210 also includes vertebral engagement surfaces 220 and 221 and, in a preferred embodiment, ridges 224 for engaging vertebral endface plates after insertion. A central cavity 222 is formed between first arm 212 and second arm 214. Bone growth material may be placed in central cavity 222. Openings 218 may also be provided in the device for receiving various tools for inserting and expanding the device. A force applied in the direction indicated by the arrows “R” will act to expand the device 210 from the first reduced size lateral position of Fig. 25 to a second expanded lateral position (not shown) after insertion of the device 210 into the disc space.

The vertebral spacers of the present invention may be placed and maintained in position within the disc space by additional fixation. The vertebral spacer devices are generally retained in position by the compressive forces of the vertebral bodies acting on the bone engaging surfaces of the implant. The spacer devices are preferably configured to transmit the compressive forces from the upper vertebral body directly through a one-piece side wall to the lower vertebral body and to limit concentration of compressive loads at the movable couplings of the arms. Moreover, it is contemplated herein that fixation devices may be used in conjunction with the vertebral spacer device of the present invention.

Alternatively, the vertebral spacer devices may be provided with an opening for
receiving a fixation device, such as a bone screw, allowing the vertebral spacer to be attached to adjacent vertebrae. Moreover, it is contemplated that the bone engaging surfaces may be configured, without limitation, to be tapered, concave or convex in order to approximate the disc space. More specifically, upper and lower bone engaging surfaces may define an angle therebetween for enhancing lordosis of the spine.

Preferably, implants according to the present invention may have lengths varying from 20mm to 26mm. Further, implants may have reduced size insertion configurations with widths varying preferably between 16mm and 20mm. Although these dimensions may be used, larger or smaller dimensions may be used without deviating from the scope of the invention.

While the invention has been illustrated and described in detail in the drawings and foregoing description, the same is to be considered as illustrative and not restrictive in character, it being understood that only the preferred embodiment has been shown and described and that all changes and modifications the come within the spirit of the invention are desired to be protected.
What is claimed is:

1. A vertebral spacer device configured for introduction into a space between two adjacent vertebrae, the device comprising:
   a spacer having a longitudinal axis and opposite bone engaging surfaces adapted to contact each of the adjacent vertebrae, said spacer further including a first arm and a second arm movable coupled to said first arm to permit lateral movement transverse to said longitudinal axis,
   wherein said first arm and second arm are coupled in a first lateral position for insertion through a portal in a disc annulus into the intradiscal space, said first arm being laterally movable with respect to said second arm to a second expanded lateral position in which said first arm is spaced a distance from said second arm.

2. The device of claim 1, wherein said first arm and said second arm each include a coupling end, said arms are pivotally coupled at said coupling ends.

3. The device of claim 2, wherein said first arm and said second arm each include a second end opposite said coupling end, said second ends include a catch mechanism disposed therebetween for maintaining said arms in said second lateral position.

4. The device of claim 3, wherein said catch mechanism includes a notch on said first arm engageable with a locking arm attached to said second arm.

5. The device of claim 3, wherein said catch mechanism includes a first plurality of serrations formed on said first arm and a corresponding second plurality of serrations formed on said second arm and interdigitably with said first plurality of serrations.

6. The device of claim 1, wherein each of said first and second arms include opposing first and second ends, first said arm is movably coupled to said
second arm by a pair of laterally extending portions at each of said first and second ends.

7. The device of claim 6, wherein said laterally extending portions define a catch mechanism for maintaining said pair of arms in spaced relation.

8. The device of claim 7, wherein said laterally extending portion of said first arm includes a projection and said laterally extending portion of said second arm defines a receptacle, said projection being movably received within said receptacle.

9. The device of claim 8, wherein a plurality of interdigitizing serrations are defined by said projections and said receptacles.

10. The device of claim 7, wherein said first arm is pivotably coupled to said second arm at said first ends, said laterally extending portions being included only at said second ends.

11. The device of claim 10, wherein said pivotable connection is a resilient hinge.

12. The device of claim 6, wherein said catch mechanism includes a plurality of interdigitizing serrations formed on each of said laterally extending portions.

13. The device of claim 1, wherein said pair of arms define a cavity when in said second lateral position.

14. The device of claim 13, further including bone-growth material placed within said cavity.
15. The device of claim 1, wherein said device is made of a bio-compatible metal material.

16. The device of claim 1, wherein said device is made of a fiber reinforced polymer composite.

17. The device of claim 1, wherein said device is made of bone material.

18. The device of claim 1, wherein opposite end-face plates of said pair of arms define an angle for enhancing lordosis of the spine.

19. The device of claim 1, wherein said bone engaging surfaces of said pair of arms defines an area that is substantially smaller than the area of the intradiscal space.

20. A vertebral spacer device configured for introduction into an intradiscal space defined by two adjacent vertebrae, the device comprising:

   a first arm having a first coupling end and a first free end, said first arm further defining a first portion of an upper and a lower endface plate adapted to contact each of the adjacent vertebrae; and

   a second arm having a second coupling end and a second free end, said arm further defining a second portion of an upper and a lower endface plate adapted to contact each of the adjacent vertebrae;

   said first coupling end and said second coupling end being movably coupled, wherein said arms are movable from a first configuration for insertion in the intradiscal space and to a second expanded configuration in the intradiscal space.

21. The device of claim 20, wherein said first free end and said second free end each define a portion of a socket, said socket configured to receive an
expansion tool for moving said first arm and said second arm from said first configuration to said second expanded configuration.

22. The device of claim 21, wherein said first arm and said second arm are composed of a bio-compatible material.

23. The device of claim 22, wherein said bio-compatible material is metal.

24. The device of claim 22, wherein said bio-compatible material is ceramic.

25. The device of claim 22, wherein said bio-compatible material is fiber reinforced polymer composite.

26. A method for maintaining the longitudinal intradiscal space between two adjacent vertebrae, the method comprising:
   (a) providing a vertebral spacer having a first portion moveably coupled to a second portion, the first portion and second portion moveable between an insertion configuration and an expanded spacer configuration;
   (b) inserting the vertebral spacer into the disc space, the vertebral spacer in the insertion configuration; and
   (c) moving the vertebral spacer into the expanded spacer configuration.

27. The method of claim 26, further comprising packing bone growth material between the first portion and second portion when the vertebral spacer is in the expanded spacer configuration.

28. The method of claim 26, wherein said moving includes inserting a tool in a socket defined by the first portion and second portion; and rotating the tool to move the vertebral spacer to the expanded spacer configuration.
29. A vertebral spacer device configured for introduction into an intradiscal space between two adjacent vertebrae, the device comprising:
   a body portion including a first end and an opposite second end; and
   a laterally expandable portion attached to said body to form a spacer, said
   laterally expandable portion movably coupled to said body portion at said first end,
   said laterally expandable portion being moveable to a first position whereby said
   spacer defines a first lateral width,
   wherein said laterally expandable portion is movable to a second position
   whereby said spacer defines a second lateral width, said second width greater than
   said first lateral width.

30. The device of claim 29, wherein said second end includes a notch
   and said laterally expandable portion includes a catch proximate said second end,
   said catch engaging said notch when said laterally expandable portion is in said
   second position.

31. The device of claim 30, wherein said catch is positioned adjacent
   said notch when said laterally expandable portion is in said first position, said
   catch being pivotably biased away from said first end to engage said notch when
   said laterally expandable portion is in said second position.

32. The device of claim 29, wherein said laterally expandable portion is
   coupled to said spacer via a resilient hinge.

33. The device of claim 29, wherein said spacer has a maximum height,
   the maximum height substantially identical in said first position and said second
   position.

34. A spacer body, comprising:
a first portion and a second portion, said first portion moveably coupled to said second portion to form a spacer, said spacer having a reduced size configuration with a maximum height and a first maximum width and an expanded size configuration having said maximum height and a second maximum width, wherein said second width is greater than said first width,

said spacer movable between said reduced size configuration and said expanded size configuration by movement of said first and second portions.

35. The spacer body of claim 34, further including a third portion joined to said first portion opposite said second portion, and a fourth portion extending between said second and third portions opposite said first portion;

wherein said second and third portions define end walls and said first and fourth portions define flexible side walls, said spacer movable between said reduced size configuration and said expanded size configuration by movement of said flexible side walls.

36. The spacer body of claim 35, wherein said flexible side walls are biased to urge said spacer to said expanded configuration.
**INTERNATIONAL SEARCH REPORT**

**A. CLASSIFICATION OF SUBJECT MATTER**

IPC 7 A61F2/44 A61F2/46

According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 A61F

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<td>the whole document</td>
<td>5-8, 10, 11, 21-24</td>
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<td>Y</td>
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Further documents are listed in the continuation of box C.

Date of mailing of the international search report

07/04/2000

Name and mailing address of the ISA

European Patent Office, P.B. 5318 Patentlaan 2 NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epc nl, Fax (+31-70) 340-3018

Authorized officer

Klein, C

*Special categories of cited documents:
*A* document defining the general state of the art which is not considered to be of particular relevance
*E* earlier document but published on or after the international filing date
*L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
*O* document referring to an oral disclosure, use, exhibition or other means
*P* document published prior to the international filing date but later than the priority date claimed

*"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
*"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
*"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
*"A" document member of the same patent family

Date of the actual completion of the international search

30 March 2000
**INTERNATIONAL SEARCH REPORT**

**C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT**

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INTERNATIONAL SEARCH REPORT

Box i Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. [X] Claims Nos.: 26-28
   because they relate to subject matter not required to be searched by this Authority, namely:
   Rule 39.1(iv) PCT - Method for treatment of the human or animal body by surgery

2. [ ] Claims Nos.:  
   because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:

3. [ ] Claims Nos.:  
   because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box ii Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1. [ ] As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.

2. [ ] As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. [ ] As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:

4. [ ] No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims: it is covered by claims Nos.: 

Remark on Protest

[ ] The additional search fees were accompanied by the applicant's protest.

[ ] No protest accompanied the payment of additional search fees.
<table>
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