Abstract: A method for treating spinal nerve compression includes sequential dilation to position an instrument cannula along a patient’s spine. Instruments can be delivered through the instrument cannula to remove targeted tissue for a decompression procedure. One of the instruments can be a reamer instrument configured to abrade, cut, or otherwise affect tissue along the patient's spine.
SPINAL NERVE DECOMPRESSION SYSTEMS, DILATION SYSTEMS, AND METHODS OF USING THE SAME

CROSS-REFERENCE TO RELATED APPLICATIONS


TECHNICAL FIELD

[0002] The present disclosure relates generally to medical systems and, more particularly, to decompression systems, delivery instruments, visualization systems, and methods for treating spinal compression. In particular, the decompression systems can include dilation systems for providing access to treatment sites to treat spinal nerve compression.

BACKGROUND

[0003] Spinal nerve compression can be caused by narrowing of the spinal canal associated with arthritis (e.g., osteoarthritis) of the spine, degeneration of spinal discs, and thickening of ligaments. Arthritis of the spine often leads to the formation of bone spurs which can narrow the spinal canal and press on the spinal cord. In spinal disk degeneration, inner tissue of the disk can protrude through a weakened fibrous outer covering of the disk and can press on the spinal cord and/or spinal nerve roots. Ligaments located along the spine can thicken over time and press on the spinal cord and/or or nerve roots. Unfortunately, spinal nerve compression can cause lower back pain, hip pain, and/or leg pain and may also result in numbness, depending on the location of the compressed nerve tissue. For example, spinal stenosis that causes spinal cord compression in the lower back can cause numbness of the legs.

BRIEF DESCRIPTION OF THE DRAWINGS

[0004] Figure 1 is an isometric view of a dilation system in accordance with an embodiment of the disclosure.
Figures 2-7 illustrate a method of performing a spinal decompression procedure using the dilation system of Figure 1 in accordance with an embodiment of the disclosure.

Figure 8 is an isometric view of an introducer dilation assembly in accordance with an embodiment of the disclosure.

Figure 9 is an exploded isometric view of the introducer dilation assembly of Figure 8.

Figure 10 is a front view of an introducer dilator in accordance with an embodiment of the disclosure.

Figure 11 is a cross-sectional view of the introducer dilator taken along line 11-11 of Figure 10.

Figure 12 is a cross-sectional view of the introducer dilator taken along line 12-12 of Figure 10.

Figure 13 is a front view of a needle device with a handle suitable for use with the introducer dilator of Figures 10-12 in accordance with an embodiment of the disclosure.

Figure 14 is a cross-sectional view of the needle device taken along line 14-14 of Figure 13.

Figure 15 is a cross-sectional view of the introducer dilation assembly taken along line 15-15 of Figure 2 with a locking mechanism in a locked configuration in accordance with an embodiment of the disclosure.

Figure 16 is a cross-sectional view of the introducer dilation assembly taken along line 16-16 of Figure 2 with the locking device in an unlocked configuration.

Figure 17 is an isometric view of a cannula dilation assembly in accordance with an embodiment of the disclosure.

Figure 18 is an exploded isometric view of the cannula dilation assembly of Figure 17.

Figure 19A is an isometric view of an instrument cannula in accordance with an embodiment of the disclosure.
Figure 19B is a cross-sectional view of the instrument cannula taken along line 19B-19B of Figure 19C.

Figure 19C is a bottom view of the instrument cannula of Figure 19A.

Figure 20A is an isometric view of another embodiment of an instrument cannula.

Figure 20B is a cross-sectional view of the instrument cannula taken along line 20B-20B of Figure 20C.

Figure 20C is a bottom view of the instrument cannula of Figure 20A.

Figure 21A is an isometric view of an instrument cannula suitable for use with an optical system in accordance with an embodiment of the disclosure.

Figure 21B is a cross-sectional view of the instrument cannula taken along line 21B-21B of Figure 21C.

Figure 21C is a bottom view of the instrument cannula of Figure 21A.

Figure 22 is a front view of a dilation device with a handle in accordance with an embodiment of disclosure.

Figure 23 is a bottom view of the dilation device of Figure 22.

Figure 24 is a longitudinal cross-sectional view of the cannula dilation assembly of Figures 17 and 18 with a locking mechanism in a locked configuration.

Figure 25 is a longitudinal cross-sectional view of the cannula dilation assembly with the locking mechanism in an unlocked configuration.

Figure 26 is an isometric view of a reamer instrument in accordance with an embodiment of the disclosure.

Figure 27 is an isometric view of the reamer instrument of Figure 26 with a retracted reaming tip and an extended depth stop member.

Figure 28 is a longitudinal cross-sectional view of the reamer instrument of Figure 26.

Figure 29 is a detailed cross-sectional view of a distal portion of the reamer instrument with a reaming tip in a deployed position.
Figure 30 is a detailed cross-sectional view of the distal portion of the reamer instrument with the reaming tip in a retracted atraumatic position.

Figures 31-35 illustrate a method of accessing a treatment site and positioning an instrument cannula along a patient's spine.

Figure 36 is an isometric view of an instrument cannula positioned along a spine in accordance with an embodiment of the disclosure.

Figures 37-39 illustrate a method of assembling an instrument positioner assembly in accordance with an embodiment of the disclosure.

Figures 40-42 illustrate an instrument positioner assembly holding an instrument cannula in a patient.

Figure 43 is a side view of a device implanted in a patient in accordance with an embodiment of the disclosure.

Figure 44 is a side view of a cannula and a visualization system in accordance with an embodiment of the disclosure.

Figure 45 is a side view of a cannula and a visualization system coupled to the cannula in accordance with an embodiment of the disclosure.

Figure 46 is a side view of a spinal decompression instrument and a visualization system in accordance with an embodiment of the disclosure.

Figure 47 is a side view of a spinal decompression instrument and a visualization system in accordance with another embodiment of the disclosure.

DETAILED DESCRIPTION

The following disclosure describes various embodiments of treatment systems, delivery systems, dilations systems, visualization systems, and associated methods of use. At least some embodiments of a treatment system include a dilation system for accessing a treatment site. The dilation system can include a series of instruments sequentially delivered into the patient to sequentially dilate tissue and/or distract structures (e.g., adjacent vertebrae). One of the instruments can be a working cannula through which instruments can be passed. In one decompression procedure, a series of instruments can be delivered through the working cannula to alter tissue...
Visualization systems can be used to view the treatment site before and/or during tissue removal. Certain details are set forth in the following description and in Figures 1-47 to provide a thorough understanding of such embodiments of the disclosure. Other details describing well-known structures and systems often associated with, for example, dilating tissue, treating the spine, decompressing spinal nerves (e.g., nerves in the spinal cord, nerves in nerve roots exiting the spinal cord, etc.), or removing tissue are not set forth in the following description to avoid unnecessarily obscuring the description of various embodiments of the disclosure.

A. Overview

At least some embodiments are methods for treating spinal nerve compression and include making an incision and sequentially dilating tissue to position a working cannula in a patient. Sequential dilation can be used to gradually enlarge openings while minimizing or limiting trauma to tissue, thereby reducing recovery times and reducing patient discomfort. For example, sequential dilation provides an advantage in that it allows a surgeon to make an initially small incision, then gradually increase the size of the opening to the minimum size required for performing the surgical procedure, thus reducing tissue damage.

Instruments can be delivered through the working cannula to access targeted tissue. The targeted tissue can be, for example, bone, ligament, facet capsule, cyst material, and/or other tissue that contributes or causes stenosis, such as central and lateral recess stenosis. The decompression procedures can cause minimal or substantially no collateral tissue disruption and can be performed under anesthesia, such as local anesthesia. The method can further include, in some embodiments, delivering a spinal device (e.g., a spinal implant, a spacer device, prosthetics disk, or other spinal device) before and/or during tissue removal.

At least some embodiments are directed to a dilation system that includes a multiple dilation assemblies. Each dilation assembly can have an outer instrument and an inner instrument with a handle. The handle can be used to insert the dilation assembly into the subject. After inserting a dilation assembly into the subject, the inner instrument of that dilation assembly can be pulled from the outer instrument. A subsequent dilation assembly can be delivered over the outer instrument. This
process can be repeated to deliver any number of dilation assemblies to perform a
desired dilation procedure.

[0048] The dilation system, in some embodiments, includes first and second
dilation assemblies. The first dilation assembly can include a first inner instrument
with a handle and a first outer instrument. The first inner instrument can be configured
to be separated from the first outer instrument when the first dilation assembly is in an
unlocked configuration. The second dilation assembly can be moved over the first
outer instrument when the second dilation assembly is in a locked configuration. The
second dilation assembly can include a second inner instrument with a handle and a
second outer instrument. The second inner instrument can be removed from the
second outer instrument when the second dilation assembly is in an unlocked
configuration.

[0049] In some embodiments, a dilation system for sequentially dilating
anatomical features to provide access to a treatment site along a subject's spine
includes first and second dilation assemblies. The first dilation assembly can include
a first dilator and a needle device. The first dilator includes a distal end, a proximal
end, and a lumen extending between the distal and proximal ends. The needle device
includes a handle and a needle. The needle can have an elongate body coupled to
the handle and a distal portion that protrudes from the distal end of the first dilator
when the elongate body extends through the lumen of the first dilator. The second
dilation assembly is configured to be moved over the first dilator after the needle
device has been removed from the first dilator. In one embodiment, the second
dilation assembly includes an instrument cannula and a second dilator. The
instrument cannula includes a distal cannula end, a proximal cannula end, and an
instrument passageway extending between the distal and proximal cannula ends.
The second dilator includes a handle and a passageway through which the first dilator
is capable of passing after the needle device has been removed from the first dilator.

[0050] A method for accessing a treatment site along a human subject's spine
comprises inserting an introducer dilation assembly into a human subject such that the
introducer dilation assembly is positioned between adjacent spinous processes of the
subject. The introducer dilation assembly can include an introducer dilator and a
needle assembly positioned in the introducer dilator. The needle assembly can be
removed from the introducer dilator after the introducer dilation assembly has been inserted into the subject. After removing the needle assembly from the introducer dilator, a cannula dilation assembly can be moved over the introducer dilator to position the cannula dilation assembly between the spinous processes. The cannula dilation assembly can include an instrument cannula and a cannula dilator positioned in the introducer dilator. The cannula dilator can be removed from the instrument cannula after the cannula dilation assembly has been inserted into the subject.

[0051] In further embodiments, a dilation system includes at least one dilator that includes a proximal portion and a tapered distal portion interconnected by an elongated body portion. The tapered distal portion can be configured for separating or splitting tissue (e.g., ligamentous tissue) for creating a pathway (e.g., a posterior midline pathway through the supraspinous ligament), as well as for distracting spinous processes. Two oppositely located and longitudinally extending channels or grooves are formed in the outer surface of the dilator for stabilizing the dilator with respect to the spinous processes. An accompanying cannula together with the dilator form an assembly for the distraction of the adjacent spinous processes, stabilization of the spinous processes with respect to the system, and/or creation of a suitable delivery path for the implantation of an interspinous spacer. In one embodiment, multiple dilators can be used to provided sequentially dilation. The dilators can be delivered over one another to gradually dilate tissue.

[0052] At least some embodiments are directed to a reamer instrument including a reaming assembly and a positioner element. The reaming assembly includes an outer reamer member having a lateral reaming element, an elongate body, and a lumen extending between first and second ends of the outer reamer member. The reaming assembly can also include an inner reamer member including a reaming tip and a rod. The rod is positioned in the lumen. The positioner element is connected to the inner reamer member. The positioner element can be moved to position at least a portion of the reaming tip outside of the outer reamer member and an atraumatic position for positioning the reaming tip within the outer reamer member.

[0053] At least one embodiment is directed to surgical instruments that can be delivered through a cannula. The surgical instruments can include a handheld reaming instrument that includes a reaming assembly and a handle assembly. The
reaming assembly can comprise an outer reamer member and an inner reamer member. The handle assembly can include a handle and a depth stop mechanism. The depth stop mechanism can be manually moved to adjust the maximum depth of penetration of the reaming assembly to avoid trauma to non-targeted tissue. The depth stop mechanism, in some embodiments, includes a locking assembly and a depth stop member. The locking assembly can have a locked configuration for holding the depth stop member and an unlocked configuration for moving the depth stop member. Other surgical instruments can be, without limitation, tissue removal instruments, debulker instruments, reamer instruments, or other types of instruments.

[0054] In some embodiments, a method for performing a procedure on a subject comprises positioning a visualization instrument relative to a cannula to view a vertebral column of the subject. The spinal decompression procedure can include, without limitation, crushing, separating, cutting, debulking, breaking, fracturing, removing, or otherwise altering tissue using decompression instruments sequentially positioned via the cannula. In non-fluoroscopic procedures, a physician can look through the lumen of the cannula to directly view the treatment site. The visualization instrument can illuminate and view the treatment site to help identify tissue (e.g., targeted tissue, non-targeted tissue, etc.), features of interest, or the like. In non-fluoroscopic procedures, the physician can use both direct viewing and viewing via fluoroscopy.

[0055] The visualization instrument, in some embodiments, can be mechanically coupled to the cannula such that the cannula and visualization instrument are moved together. For example, a coupler can fixedly couple the visualization instrument to the cannula. In one embodiment, the coupler can include a clamp having an open configuration for repositioning the visualization instrument and a closed configuration for holding the visualization instrument. In some embodiments, the visualization instrument can be positioned in an access feature in the form of a through-hole in a sidewall of the cannula and can include one or more light sources capable of outputting light for illuminating a treatment site distal to the cannula. The illustrated target tissue can be viewed with the naked eye. Additionally, the visualization instrument can include one or more imaging devices, such as cameras, for viewing on an electronic display (e.g., a color monitor).
In some embodiments, a visualization system can be used to view tissue to, among other things, prevent damaging non-targeted tissue. The visualization system can provide viewing of decompression instruments and/or treatment sites to help position decompression instruments. In one embodiment, the visualization system can be used for directly viewing of the treatment site and/or distal end of the decompression instrument. In other embodiments, visualization systems can provide viewing via a display, such as a color monitor.

Visualization systems can be used in decompression procedures for treating spinal nerve compression (e.g., spinal cord compression, spinal nerve root compression, or the like), spinal disk herniation, osteoporosis, stenosis, or other diseases or conditions. In one embodiment, a tissue removal instrument is used to perform a spinal cord decompression procedure, which can include removing bone from one or more vertebrae, separating the ligamentum flavum from one or more vertebrae, cutting or debulking the ligamentum flavum, and/or removing loose tissue while a physician views the treatment site using the visualization system.

The terms "distal" and "proximal" within this description, unless otherwise specified, reference a relative position of the portions of an systems, instruments, and/or associated access devices with reference to an operator and/or a location in the patient. For example, in referring to visualization systems described herein, the term "proximal" can refer to a position closer to the operator, and the term "distal" can refer to a position that is more distant from the operator.

B. Decompression Systems

Figure 1 is an isometric view of a dilation system 90 for sequentially dilating anatomical features of a human subject in accordance with an embodiment of the disclosure. The dilation system 90 can include an introducer or inner dilation assembly 100 ("introducer dilation assembly 100") for initially dilating anatomical features and an outer dilation assembly 170 for further dilating the anatomical features. The introducer dilation assembly 100 can include a hollow introducer dilator 166 and a needle device 167 extending through the introducer dilator 166. The outer dilation assembly 170 can include a dilator device 176 and a working or instrument cannula 172 ("instrument cannula 172") and can be delivered over the introducer.
dilator 166 after the needle device 167 has been removed from the introducer dilator 166. The introducer dilation assembly 100 and outer dilation assembly 170 can sequentially distract adjacent vertebrae to achieve a large amount of distraction while managing the pressure applied to the vertebrae.

[0060] Figures 2-5 illustrate a dilation procedure performed using the dilation system 90 of Figure 1. Figure 2 shows the introducer dilation assembly 100 after it has been driven into the subject. The introducer dilation assembly 100 can extend through a subject's skin 140, subcutaneous tissue 142, and supraspinous ligament 150 and, in midline procedures, can be positioned generally between adjacent spinous processes 160, 164. Figure 3 shows the introducer dilator 166 after the needle device 167 (Figure 2) has been removed therefrom. Figure 4 shows the dilation assembly 170 after it has been delivered over the introducer dilator 166 to position the instrument cannula 172 between the spinous processes 160, 164. Figure 5 shows the instrument cannula 172 after the introducer dilator 166 has been removed from the dilation assembly 170 and after the dilator device 176 has been pulled out of the instrument cannula 172. The cannula 172 can hold apart the spinous processes 160, 164 to maintain a desired amount of distraction for enlarging an interspinous space.

[0061] Figures 6 and 7 illustrate a method of performing at least a portion of the decompression procedure using the instrument cannula 172 held by a cannula holder 173. A surgical instrument in the form of a reamer instrument 190 has a distal end 192 that can scrape, abrade, or otherwise alter tissue within, adjacent to, or along the subject's spine. Other instruments can be delivered through the cannula 172 to perform a wide range of decompression procedures or other type of procedure. Details of the instruments and features shown in Figures 1-7 are discussed below.

[0062] Figure 8 is an isometric view of the introducer dilation assembly 100 with a relatively sharp tip or distal portion 200. Figure 9 is an exploded isometric view of the introducer dilation assembly 100. The needle device 167 can include a handle 210, a locking mechanism 212, and a needle 214. The handle 210 can be conveniently gripped by a user to push the introducer dilation assembly 100 into the subject. The locking mechanism 212 can have a locked configuration for holding the introducer dilator 166 and an unlocked configuration for releasing the introducer dilator 166.
Referring now to Figure 9, the needle 214 can be directly or indirectly coupled to the handle 210 and can include an elongate body 220 and a distal portion 222 with a sharp needle tip 202. To assemble the introducer dilation assembly 100, the needle tip 202 can be inserted into the introducer dilator 166. The needle 214 can be advanced along the introducer dilator 166 until a proximal end 240 of the introducer dilator 166 is received by the locking mechanism 212. The locking mechanism 212 can be moved from an unlocked configuration to a locked configuration to securely hold the introducer dilator 166. Details of the introducer dilator 166 are discussed in connection with Figures 10-12, details of the needle device 167 are discussed in connection with Figures 13 and 14, and details of the locking mechanism 212 are discussed in connection with Figures 15 and 16.

[0063] Figure 10 is a front view of the introducer dilator 166 in accordance with an embodiment of the present disclosure. Figure 11 is a longitudinal cross-sectional view of the introducer dilator 166 taken along line 11-11 of Figure 10. Figure 12 is a cross-sectional view of the introducer dilator 166 taken along line 12-12 of Figure 10. Referring now to Figure 10, the introducer dilator 166 can include a tapered distal end 204, proximal end 240, and main body 242. The distal end 204 can include an opening 274 (Figure 11) and a smooth outer surface 276 and can have a generally frusto-conical shape, truncated pyramidal shape, or other shape suitable for passing through an incision, spreading or stretching tissue, dilating openings or gaps, or the like. The proximal end 240 can include flanges 250a, 250a that define receiving windows 260a, 260b, respectively. Referring now to Figures 11 and 12, an inner surface 262 defines a passageway 272 extending between the openings 270, 274. The passageway 272 is configured to slidably receive the needle 214 (Figure 9).

[0064] Figure 12 shows two oppositely positioned outer alignment features in the form outer channels 280, 282 that extend longitudinally along the main body 242. The channels 280, 282 can have U-shaped profiles, V-shaped profiles, arcuate profiles (e.g., concave configurations), or other profiles suitable for engaging vertebrae, spinous processes, or other tissue. As shown in Figure 10, the channel 280 can extend from the distal end 204 toward the proximal end 240 to allow tissue to slide along the entire length of the introducer dilator 166 or portion thereof. Inner alignment features 290, 292 can be in the form of longitudinally-extending convex portions located on opposites sides of the passageway 272. The number, location, and
orientation of alignment features can be selected based on the instruments used with the introducer dilator 166.

[0065] Figure 13 is a front view of the needle device 167 in accordance with an embodiment of the disclosure. The needle 214 can include the needle tip 202, a proximal end 300, and a main body 302. The proximal end 300 can be fixedly or detachably coupled to the handle 210, illustrated as a T-shaped handle that a user can comfortably grip by wrapping his or her fingers about handle end portions 303, 304. Other types of handles can also be used. The needle tip 202 may be relatively sharp and may have a knife-like edge that can pierce tissue (e.g., ligaments) without first using a sharp edge and can therefore be used for percutaneous procedures. In other embodiments, the needle tip 202 can have a conical shape, a pyramidal shape, or other suitable shape for piercing tissue.

[0066] Figure 14 is a cross-sectional view of the needle 214 taken along line 14-14 of Figure 13. Two oppositely positioned alignment features in the form of channels 310, 312 extend longitudinally along the main body 302. Referring to Figure 13, the channels 310, 312 can have arcuate profiles, U-shaped profiles, V-shaped profiles, or other suitable convex or concave profiles for engaging the alignment features 290, 292 (Figure 12) of the introducer dilator 166. In some embodiments, the channels 310, 312 can slidably engage respective alignment features 290, 292 of the introducer dilator 166 to rotationally lock together the needle 214 and introducer dilator 166. Alignment features in the form of longitudinally-extending convex portions 320, 322 located on opposites sides of the needle 214 can slidably engage alignment features in the form of convex portions 330, 332 (Figure 12). In various embodiments, the needle 214 can have a polygonal cross-sectional profile (e.g. a square profile, a rectangular profile, etc.), an elliptical profile, or other profile suitable for maintaining desired alignment with introducer dilators or other components.

[0067] Figures 15 and 16 are longitudinal cross-sectional views of the introducer dilation assembly 100 with the locking mechanism 212 in locked and unlocked configurations, respectively. Referring to Figure 15, flanges 250a, 250b of the introducer dilator 166 can be held between upper surfaces of the retaining elements in the form of flanges 350a, 350b and an abutment 351 of the handle 210. To move the locking mechanism 212 to the unlocked configuration, a cylindrical body 352 of the
locking mechanism 212 can be rotated about an axis of rotation 354 (indicated by arrows 360) to move the flanges 350a, 350b. Figure 16 shows the locking mechanism 212 in the unlocked configuration after the flanges 350a, 350b (Figure 15) have been moved out of the windows 260a, 260b. To separate the introducer dilator 166 and the needle device 167, the user can push the introducer dilator 166 distally (indicated by arrow 370) away from the handle 210 and/or pull the needle 214 proximally (indicated by arrow 371) relative to the introducer dilator 166. Other types of locking mechanisms can be used and may include, without limitation, one or more pins, threaded members, or other features suitable for coupling together and releasing components.

[0068] Figure 17 is an isometric view of the outer dilation assembly 170 in accordance with an embodiment of the disclosure. Figure 18 is an exploded isometric view of the outer dilation assembly 170. Referring to Figures 17 and 18 together, the dilator device 176 can include a dilator handle 400, a locking mechanism 412, and an elongate dilator 402 with a distal end 410. The locking mechanism 412 has a locked configuration for coupling together the instrument cannula 172 and the dilator device 176 and an unlocked configuration for separating the instrument cannula 172 and the dilator device 176.

[0069] To assemble the outer dilation assembly 170 of Figure 18, the distal end 410 of the elongate dilator 402 can be inserted into an entrance opening 420 of the instrument cannula 172. The elongate dilator 402 can be moved along the cannula 172 until a head 430 of the cannula 172 is received by the locking mechanism 412. The locking mechanism 412 can be moved from the unlocked configuration to a locked configuration to hold together the dilation assembly 170 and instrument cannula 172 such that the distal end 410 protrudes from the instrument cannula 172 to expose sloped channels 442.

[0070] Figure 19A is an isometric view of the instrument cannula 172 in accordance with an embodiment of the disclosure. Figure 19B is a cross-sectional view of the instrument cannula 172 taken along line 19B-19B of Figure 19C. Figure 19C is a bottom view of the instrument cannula 172. The instrument cannula 172 can include the head 430, distal end 410, and main body 440 therebetween. The head 430 defines the opening 420 (Figures 19A and 19B) and keying features 450a, 450b.
(collectively "keying features 450"). An instrument passageway 470 (Figure 19B) extends between the opening 420 and the opening 472 and is configured to receive instruments.

[0071] The instrument cannula 172 can include positioning features 460 located along the bottom of guide channels 480. The positioning features 460 can be recesses (e.g., spherical recesses, elongated recesses, etc.), protrusions, grooves, notches, or other features suitable for engaging tissue or bone. The illustrated embodiment includes eleven positioning features 460, but a greater or lesser number of positioning features can be selected based on a desired number of available preferential positions. In some embodiments, the instrument cannula 172 can include an array of locators 461 for positioning relative to a holder, such as the clamp assembly discussed in connection with Figures 37-42. The guide channels 480 can have U-shaped cross-sectional profiles, V-shaped cross-sectional profiles, or other suitable profiles for interacting with anatomical features. The guide channels 480 can be sloped or angled to provide for distraction.

[0072] Figures 20A is an isometric view of an instrument cannula in accordance with an embodiment of the disclosure. Figure 20B is a cross-sectional view of the instrument cannula taken along line 20B-20B of Figure 20C. Figure 20C is a bottom view of the instrument cannula of Figure 20A. The description of the cannula 172 of Figures 19A-19C applies equally to the cannula 172 of Figures 20A-20C, except as detailed below. The cannula 172 has twelve (illustrated), thirteen, fourteen, or more positioning features 460 located along a central region of the guide channels 480. Guide rails 481 of Figures 20A and 20B are higher and longer than guide rails 481 of Figure 19C. In some embodiments, most or all of the positioning features 460 of Figures 20A-20C are positioned between the guide rails 481. The number, spacing, dimensions of the positioning features 460 can be selected based on the positions and configurations of the anatomical structures to be received, and the spacing and dimensions (e.g., lengths, heights, etc.) of the guide rails 481 can be selected based on the anatomical features to be moved along the guide channels 480.

[0073] Figures 21A is an isometric view of an instrument cannula 172 in accordance with an embodiment of the disclosure. Figure 21B is a cross-sectional view of the instrument cannula taken along line 21B-21 B of Figure 21C. Figure 21C is
a bottom view of the instrument cannula of Figure 21A. The description of the cannula 172 of Figures 19A-20C applies equally to the cannula 172 of Figures 21A-21C, except as detailed below. Referring to Figure 21A, the cannula 172 has positioning features (illustrated as notches) located along the guide channels. Instruments can be passed through access features 487 circumferentially spaced about the cannula 172. The access features 487 can be ports, through-holes, or other features through which visualization instruments, surgical instruments, or other instruments can be passed. For example, illumination instruments can be inserted through the ports 487 to illuminate tissue distal to the cannula 172.

[0074] Figures 22 and 23 are side and bottom views, respectively, of the dilator device 176. Referring to Figure 22, the handle 400 can include handle portions 500a, 500b, a guide 530, and an access opening or window 502 between the handle portions 500a, 500b. The guide 530 can include an opening 532 (Figures 17 and 24) positioned generally along the longitudinal axis 541 of the dilator device 176. Other types of handle assemblies can also be used. For example, T-shaped handles or spherical shaped handles can be used, if needed or desired.

[0075] Figures 24 and 25 are cross-sectional views of the dilator device 176 holding the introducer dilator 166 (shown in phantom line). The elongate dilator 402 can include the distal end 410, a proximal end 540, and an elongate body 544 therebetween. A passageway 545 extends between openings 546, 547. Figure 24 shows the locking mechanism 412 in a locked configuration for coupling together the instrument cannula 172 and outer dilation assembly 170. The head 430 of the instrument cannula 172 can be held between retaining features 550a, 550b of the locking mechanism 412 and an abutment 553 of the handle 400. A main body 572 of the locking mechanism 412 can be rotated about an axis of rotation 574 (indicated by arrows 552) until the retaining features 550a, 550b are aligned with the keying features (e.g., keying features 450a, 450b of Figures 19A-19B of the head 430). Figure 25 shows the locking mechanism 412 in the unlocked configuration to align the retaining features 550a, 550b with the keying features 450a, 450b, respectively. As such, the instrument cannula 172 is free to slide distally along the elongate dilator 402 away from the locking mechanism 412.
Figure 26 is an isometric view of a reamer instrument 190 in accordance with an embodiment of the disclosure. Figure 27 is an isometric view of the reamer instrument 190 with a retracted reaming tip and an extended depth stop member 650. Referring now to Figure 26, the reamer instrument 190 can include a reaming assembly 600 for abrading, scraping, or otherwise mechanically altering bone or tissue and a handle assembly 602 for operating the reaming assembly 600. The reaming assembly 600 can include a reaming tip 611 for contacting distal tissue and a lateral reaming element 612 for contacting lateral tissue. The reaming tips 611, 612 can include a roughened surface, array of sharp protrusions, texturing, or other features capable of loosening, separating, cutting, scraping, or otherwise affecting tissue. For example, the reaming tip 611 can be used to bore through tissue, and the lateral reaming element 612 can be used to ream laterally adjacent tissue. When the tip reaming tip 611 is retracted to an atraumatic position (Figure 27), it can be positioned inside of the lateral reaming element 612. As shown in Figure 27, an atraumatic edge 613 can be configured to inhibit or prevent injury to distal tissue. Accordingly, the reaming assembly 600 can be moved between different configurations to target specific tissue during a procedure.

The handle assembly 602 can include, without limitation, a depth stop mechanism 630, a reaming control element 632 ("control element 632"), and a handle housing 640 for protecting internal components. The depth stop mechanism 630 can include the stop member 650 and a positioning assembly 652. The stop member 650 can include a head 653 oriented generally perpendicular to a longitudinal axis 655 of the reaming assembly 600. The positioning assembly 652 can be used to move the stop member 650 distally (indicated by arrow 660 in Figure 26) or proximally (indicated by arrow 661 in Figure 27) to adjust, for example, a maximum depth of penetration of the reaming assembly 600. Once the stop member 650 is at the desired location, the positioning assembly 652 can be locked to hold the head 653 stationary relative to the reaming assembly 600. The reaming control element 632 can be moved proximally (indicated by arrow 670 in Figure 26) to move the reaming tip 611 (Figure 26) into the lateral reaming element 612. Referring now to Figure 27, the reaming control element 632 can be moved distally (indicated by arrow 672) to move the reaming tip 611 out of the lateral reaming element 612.
Figure 28 is a longitudinal cross-sectional view of the reamer instrument 190. In one embodiment, the reaming assembly 600 can include an outer reamer member 610 with the lateral reaming element 612 and an inner reamer member 750 with the reaming tip 611. The outer reamer member 610 can have a proximal end 720, a distal end 722, and a hollow elongate main body 724. The proximal end 720 can be fixedly coupled to the handle housing 640. The main body 724 can be a shaft (e.g., a tubular shaft made of metal, plastic, etc.) with an inner surface 740 (Figure 29) that closely surrounds the inner reamer member 750. The inner reamer member 750 can include a proximal end 752, a distal end 754, and an elongate body 756 (e.g., a solid or hollow rod or shaft made of metal, plastic, etc.). The proximal end 752 can be connected to the control element 632 by, for example, one or more fasteners, pins, welds, or other connection elements. In other embodiments, the proximal end 752 and a control element 632 can have a one-piece construction.

Figures 29 and 30 are detailed cross-sectional views of the distal end of the reaming assembly 600 in two different configurations. Referring now to Figure 29, the distal end 722 of the outer reamer member 610 can include an opening 732 and a stop in the form of a shoulder 734. A widened passageway 730 extends from the opening 732 to the shoulder 734. The distal end 754 of the inner reamer member 750 can include a shoulder 763 and a head 762. Figure 29 shows the reaming tip 611 in a distal reaming position such that the reaming tip 611 protrudes outwardly (distally) from the edge 613 of the distal end 722. The inner reamer member 750 can be moved proximally (indicated by arrow 770 in Figure 29) until the shoulder 763 contacts the shoulder 734 (Figure 30).

Referring again to Figure 7, the deployed reaming tip 611 can be used to abrade tissue (not shown) located posterior to the illustrated ligamentum flavum 615. The reaming assembly can be advanced distally to abrade tissue adjacent to the ligamentum flavum 615 or the tissue of the ligamentum flavum 615. The reaming tip 611 can be retracted to the atraumatic position (Figures 27 and 30) to avoid damaging non-targeted tissue, such as the spinal cord 617, which is located between the ligamentum flavum 615 and a ligament 184. The spinal cord 617 extends from the brain to the bottom of the spine and extends through vertebral foramina. Spinal nerves branch from the spinal cord 617 and exit the spine and extend to other parts of the body. The reaming tip 611 can be retracted to avoid traumatizing or damaging
nerve tissue, or other non-targeted tissue, such as the epidural sac. With the reaming tip 611, the reamer instrument 190 can be inserted deeper into the subject without risk of tearing or damaging the epidural sac or injuring the spinal cord 617. For example, the atraumatic edge 782 (Figure 30) can be blunted, rounded, and/or smooth to inhibit, limit, or substantially prevent damage and/or injury to the epidural sac.

[0081] Referring again to Figure 28, the depth stop mechanism 630 can include a control element 700, a slider locking element 702, and a rack 704. The control element 700 can include a cantilevered lever 709 movable between a first position (e.g. an undepressed position, an extended position, etc.) to a second position (e.g., a depressed position, an unextended position, etc.). When the control element 700 is depressed, an engagement feature (e.g., a U-shaped member) can be moved away from teeth of the rack 704. To move the positioning assembly 630 from a locked configuration (Figure 28) to an unlocked configuration, a user can press down on the control element 700 to overcome a biasing force provided by the element 709 and thereby move the control element 700 downwardly. After the control element 700 is depressed, it can be moved proximally or distally. After the stop member 650 is moved to the desired position, the control element 700 can be released to allow the control element 700 to move back to the undepressed position. Other types of positioning assemblies can be used and can include, without limitation, one or more biasing devices (e.g. springs, actuators, etc.), control elements, gears, or the like. The configuration and functionality of the positioning assemblies can be selected based on the desired operation of the reamer instrument 190.

[0082] Figures 31-35 illustrate a method of performing at least a portion of a decompression procedure on a patient in accordance with an embodiment of disclosure. Generally, an incision can be made along the patient's back using, for example, a scalpel 780. The introducer dilation assembly 100 can be moved through the incision and inserted between the spinous processes 160, 164. The needle device 167 can be removed from the introducer dilator 166. The dilation assembly 170 can be advanced over the introducer dilator 166 and into the subject. The introducer dilator 166 can then be removed from the dilation assembly 170, and the dilator device 176 can then be removed from the instrument cannula 172. Details of the procedure are discussed below.
Figure 31 shows the scalpel 780 ready to make an incision along the midline of the subject. An entry point can be selected on the patient's skin to obtain access to the targeted surgical site, and an incision of appropriate length is made through the dermal layers of a patient's body at the entry point. The length and depth of the incision may be larger depending on whether the clinician is using an open, mini-open, or minimally invasive, percutaneous approach. In some procedures, a targeted surgical level can be identified and a midline incision (e.g., 5 mm to 15 mm length incision) can be made under direct visualization, fluoroscopic guidance, or other suitable visualization technique. In some procedures, the supraspinous ligament 150 can be dissected (e.g., longitudinally dissected) to provide access to an interspinous space 174. The scalpel 780, or other cutting instruments, can form incisions at other locations to access the spine using non-midline approaches, such as lateral approaches.

Figure 32 shows the introducer dilation assembly 100 after it has been passed through the incision and moved through the supraspinous ligament 150. As the introducer dilation assembly 100 is advanced distally, the sharp tip 202 and distal end 204 dilates tissue (e.g., spreads or separates tissue) and/or otherwise affect tissue to facilitate penetration into the patient. As the introducer dilation assembly 100 is initially inserted between the spinous processes 160, 164, it can drive apart the spinous processes. The distracted spinous processes 160, 164 can be positioned in and slide along the channels 280, 282, respectively, (Figure 12) until the distal end 204 is at the desired depth. The spinous processes 160, 164 in the channels 280, 282, respectively, can inhibit or limit rotation of the introducer dilator 166. Such placement of the introducer dilation assembly 100 with respect to the spinous processes 160, 164 therefore stabilizes the introducer dilation assembly 100.

The introducer dilation assembly 100 can be monitored using fluoroscopy, direct visualization, or other visualization technique. After the introducer dilation assembly 100 is at the desired location, the locking mechanism 212 can be moved from the locked configuration to the unlocked configuration and the needle device 167 can then be pulled out of the introducer dilator 166.

Figure 33 shows the needle device 167 separated from the introducer dilator 166. A longitudinal axis 781 of the introducer dilator 166 can be generally
perpendicular to the patient's spine or at another suitable orientation. For example, the longitudinal axis 781 can be generally parallel to the anterior-to-posterior direction.

[0087] Referring to Figure 34, the dilation assembly 170 has been inserted over the introducer dilator 166. The dilation assembly 170 can be aligned (e.g., rotationally aligned) with the proximal end 240 of the introducer dilator 166 and then slid over the proximal end 240. As shown in Figure 23, the alignment features 560, 562 (e.g., convex features) of the elongate dilator 402 can be received by the channels 280, 282 of the introducer dilator 166 to rotationally lock the dilator 402 and the introducer dilator 166. As the dilator 402 moves along the introducer dilator 166, the tapered distal end (Figure 17) can dilate the incision, spread or separate tissue, and/or otherwise affect tissue to facilitate penetration into the patient. If the spinous processes 160, 164 are sufficiently close together, the distal end 410 can contact and push apart the spinous processes 160, 164.

[0088] As the dilation assembly 170 is advanced over the introducer dilator 166, the channels 280, 282 (Figure 12) of the introducer dilator 166 can be aligned with the respective alignment features 560, 562 (Figure 23). The spinous processes and/or tissue can move from the channels 280, 282 of the dilator 166 to the respective alignment features 560, 562 of the dilator 402 as the dilation assembly 170 is advanced along the stationary dilator 166. Referring to Figures 21 and 34, the channels 480 of the cannula 172 (Figure 21) can be generally aligned with the channels 442 (Figure 17) of the elongate dilator 402 such that the spinous processes and/or tissue move from the channels 442 of the dilator 402 to the respective channels 480 of the cannula 172.

[0089] Figure 35 shows the dilator device 176 spaced apart from the instrument cannula 172. Figure 36 shows an instrument cannula 172 positioned between the spinous processes 160, 164. Tips of the spinous processes 160, 164 can be received by the receiving features 460 (see, e.g., Figure 19A) to set the cannula 172. Any number of different instruments can be delivered through the instrument cannula 172 to treat a wide range of symptoms, conditions, and/or diseases, including, without limitation, spinal nerve compression (e.g., spinal cord compression, spinal nerve root compression, or the like), spinal disk herniation, osteoporosis, stenosis, or other diseases or conditions. In some procedures, the cannula 172 provides access for
surgical instruments for performing a spinal cord decompression procedure that includes, without limitation, delivering visualization media, removing bone from one or both vertebrae 804, 806, separating the ligamentum flavum from one or both vertebrae 804, 806, cutting or debulking the ligamentum flavum, and removing loose tissue. A wide range of decompression procedures can be performed and can include, without limitation, a discectomy, osteophyte removal, laminotomy, or other type of decompression procedures for removing bone and/or soft tissue. Each stage of the decompression procedure can be performed with a different instrument or series of instruments.

[0090] Instruments can be advanced through the cannula 172 to remove tissue (e.g., bone, connective tissue, etc.) to, for example, reduce spinal compression, increase access to the treatment site, and can be viewed under fluoroscopy or other suitable visualization technique. The cannula 172 can be sufficiently large to allow repositioning of the instruments to access different treatment sites, such as the lateral recesses, facets, ligamentum flavum, or the like. In some simultaneous bilateral access procedures, the cannula 172 can be repositioned while remaining in the patient to remove tissue from opposing lateral recesses or other lateral treatment sites. Additionally, the cannula 172 can provide direct visualization. For example, a user can view the treatment site and/or instrument by looking through the passageway of the cannula 172. Additionally or alternatively, visualization devices (e.g., fiber optics, cameras, light sources, or the like) can be coupled to or incorporated into the cannula 172. After removing the desired amount of bone (or other tissue), the instrument can be withdrawn from the subject.

[0091] Fluoroscopy (e.g., anterior-posterior imaging, lateral imaging, contralateral-oblique imaging, etc.) can be used to view the treatment site, tools, and delivery path. In certain procedures, visualization techniques can be used to identify margins of the epidural space, dura, ligamentum flavum, and/or nerve roots relative to the lamina and interlaminar space, as well as the features of instruments. Contrast media can be refreshed to maintain desired imaging. When reaming instruments (e.g., reaming instrument 190) are near nerve tissue, the reaming instruments can be in an atraumatic configuration.
[0092] Figures 37-39 illustrate a method of assembling a holder in the form of an instrument positioner assembly for holding an instrument cannula in accordance with an embodiment of the disclosure. Generally, the cannula 172 can be installed in a collar 800 positionable in an open clamp assembly 802. The cannula 172 can be rotated relative to the clamp assembly 802. After the cannula 172 is at the desired orientation, the clamp assembly 802 can be closed to securely hold the collar 800. Instruments can be delivered through the cannula 172 while the clamp assembly 802 holds the cannula 172 at the desired orientation. The clamp assembly 802 can be opened to reorient the cannula 172. Non-limiting exemplary methods of using the clamp assembly 802 are discussed below.

[0093] A user can select a desired axial position along the cannula 172 for the collar 800 based on, for example, the distance from the patient's skin to the treatment site. Figure 37 shows the collar 800 coupled to the cannula 172 and ready for installation in the clamp assembly 802. The collar 800 can include protrusions or other features matable with one or more locators 461 of the cannula 172. The illustrated cannula 172 includes an array of spaced apart locators 461 that can be, for example, recess, holes, or the like. The cannula 172 can be inserted into the patient before or after installing the collar 800.

[0094] The clamp assembly 802 can be placed over the collar 800. A base 810 (Figure 38) can rest against the patient's skin and can extend in the superior direction (or other direction). The cannula 172 can be rotated in the lateral direction (indicated by arrows 812, 814) or other desired direction. The base 810 can inhibit or limit rocking movement of the clamp assembly 802 (e.g., rocking in the superior direction), thereby stabilizing the cannula 172.

[0095] Referring to Figure 38, a lever mechanism 816 can be used to close and open the clamp assembly 802. Figure 39 shows the closed clamp assembly 802 with the rotationally fixed collar 800. The cannula 172 can be keyed to the collar 800 to prevent axial movement of the cannula 172, and the clamp assembly 802 can be opened to adjust the orientation of the cannula 172. When an instrument is positioned in the cannula 172, a depth stop mechanism of the instrument can contact the cannula 172 to limit movement of the instrument in the distal direction. By way of example, the stop member 650 discussed in connection with Figures 26-27 can contact the
proximal end (e.g., head 430) of the cannula 172 and thereby limit the penetration depth of the reamer instrument 190. By adjusting the position of the stop member 650 (Figures 26-27), the penetration depth of the reamer instrument 190 can be adjusted to safely access targeted tissue.

[0096] Figures 40-42 illustrate the clamp assembly 802 positioned on a patient. The cannula 172 of Figure 40 is positioned to access left regions of the subject's left lateral vertebral body. The cannula 172 of Figure 41 is positioned to access the right regions of the subject's right lateral recess of the vertebral body. Figure 42 shows the reamer instrument 190 ready to be delivered through the cannula 172. Visualization techniques can be used to confirm the position, trajectory, and depth of the end of instrument cannula 172, instrument(s), etc. The dimensions (e.g., diameter) of the passageway 470 (Figure 19B) of the cannula 172 can be sufficiently large to allow repositioning of the instrument to access different treatment sites, such as the lateral recesses, facets, ligamentum flavum, or the like. In some simultaneous bilateral access procedures, the cannula 172 can be repositioned while remaining in the patient to remove tissue from opposing lateral recesses or other lateral treatment sites. Additionally, the cannula 172 can provide direct visualization. For example, a user can view the treatment site and/or instrument by looking through the passageway 470 (Figure 19B). Additionally or alternatively, visualization devices (e.g., fiber optics, cameras, or the like) can be coupled to or incorporated into the cannula 172 and/or instruments for viewing.

[0097] In some procedures, the reamer instrument 190 can extend a distance (e.g., 10 mm, 15 mm, 20 mm, etc.) past the distal end of the cannula 172 when the stop member 650 contacts the head 430. The reamer instrument 190 can be rotated to abrade, loosen, tear, or otherwise alter tissue and can be removed any number of times to remove residual tissue (e.g., ligament tissue, bone tissue, etc.) attached to the reamer instrument. Different types of instruments can be used to cut bone, create one or more defects (e.g., a generally hemispherical defect) in the inferior medial aspect of the superior lamina, or otherwise prepare the treatment site.

[0098] To remove midline tissue, the cannula 172 can be oriented towards the midline interlaminar region. A reamer instrument can be inserted through the cannula 172 and positioned towards the midline position of the superior lamina. In one
exemplary embodiment, the depth stop mechanism 630 of the reamer instrument 190 can be used to, for example, prevent injury to the dural or other non-targeted tissue. Visualization techniques can be used to monitor the position on the reamer head. In some procedures, the reamer head can be moved from midline to left lateral or the right lateral. Any number of reamer instruments can be used to remove the desired amount of midline lamina bone. The depth stop mechanism can be used to allow access to the targeted region while maintaining a desired distance from the epidural space and other vital structures. After performing the reaming procedure, the reamer can be removed from the patient and a preparation procedure can be performed. The preparation procedure can include, without limitation, irrigating the treatment site, removing residual tissue (e.g., via suction), applying one or more agents (e.g., hemostatic agents), or other procedures.

[0099] A debulker instrument can be used to provide a complete blunt dissection of the ligamentum flavum from the lamina and disrupt ligamentous tissue. In some procedures, the debulker instrument is inserted through the cannulas and positioned at a midline position of the superior lamina. The depth stop mechanisms can facilitate positioning of the distal tip (e.g., debulking head) at the most dorsal margin of the superior lamina. Intraoperative fluoroscopy and/or tactile feedback can be used to confirm positioning. While maintaining a midline trajectory, the distal tip of the debulker instrument can be gently moved around the inferior lamina lip and repositioned against the bony underside. The adjustable depth stop can be reset, if desired, to allow access to the targeted region while maintaining a desired distance from the epidural space and other vital structures. The properly positioned distal tip can engage the underside of the lamina and resist attempts to gently withdraw the instrument.

[00100] The debulking tip can dissect and separate the ligamentum flavum from the lamina when it is move from midline toward the lateral recess. A subtle left-right sweeping motion can be used disrupt ligamentous tissue and help extend the desired tissue plane. The distal tip can be moved until it reaches the most lateral desired position. The depth stop mechanism can be adjusted to allow access to the lateral recesses. The debulker tip can be moved slightly inferior and out from the lamina underside. The debulker tip can be used to continually debulk the ligamentum flavum. The depth stop mechanism can be adjusted to allow access to the targeted region,
while intraoperative fluoroscopy is used to verify the distal tip position and maintain a safe working distance from the epidural space and/or other vital structures. After performing the debulking procedure, the debulker instrument can be removed from the patient and a preparation procedure can be performed.

[00101] The lamina can be removed using a tissue removal instrument. The cannula 172 can be oriented towards the desired interlaminar region (e.g., left or right interlaminar region). A closed jaw assembly of a tissue removal instrument can be moved through the cannula 172 towards a generally midline position. A depth stop mechanism can be used to adjust the depth of penetration until the jaw assembly is positioned proximate the most dorsal margin of the superior lamina. The jaw assembly can be closed to remove tissue. While maintaining midline trajectory, the jaw assembly can be moved around the inferior lamina lip and positioned against the bony underside. The depth stop can be adjusted to allow access to the targeted region while maintaining a desired distance from the epidural space and other vital structures. The distal or lower jaw of the jaw assembly can engage the underside of the lamina and the proximal or upper jaw can be positioned just dorsal to the lamina. The jaw assembly can be held against the targeted lamina bone while the jaw assembly is closed. The tissue removal instrument can be withdrawn from the patient. The jaw assembly can be opened to release the captured material. This process can be repeated to remove bone and other tissue in the lateral direction until the desired decompression is achieved.

[00102] Systems, components, and instruments disclosed herein can be disposable or reusable. For example, the reamer instrument 190 can be disposable to prevent cross-contamination. As used herein, the term "disposable" when applied to a system or component (or combination of components), such as an instrument, a tool, or a distal tip or a head (e.g., a reamer head, a jaw assembly, etc.), is a broad term and generally means, without limitation, that the system or component in question is used a finite number of times and is then discarded. Some disposable components are used only once and are then discarded. In other embodiments, the components and instruments are non-disposable and can be used any number of times.

[00103] The cannula 172 can be used deliver one or more spinal implants before, after, or during tissue removal. The methods of delivery, delivery instruments,
dilators, spinal implants, and other features of U.S. Pat. No. 8,012,207; U.S. Pat. No. 8,123,807; U.S. Pat. No. 8,152,837; U.S. App. No. 12/217,662 (corresponding U.S. Pub. No. 20080287997); U.S. App. No. 13/844,173; U.S. App. No. 12/358,010, and U.S. App. No. 13/844,324. U.S. Pat. No. 8,012,207; U.S. Pat. No. 8,123,807; U.S. Pat. No. 8,152,837; U.S. App. No. 12/217,662 (corresponding U.S. Pub. No. 20080287997); U.S. App. No. 13/844,173; U.S. App. No. 12/358,010, and U.S. App. No. 13/844,324 are hereby incorporated by reference in their entireties. Figure 43 shows an implanted device 830 positioned between the spinous processes 160, 164. The device 830 can be delivered via the cannula or other access device. In one embodiment, the device 830 is a SUPERION® Interspinous Spacer from VertiFlex, Inc. (San Clemente, CA) or a similar device. The device 830 can be implanted while imaging using visualization media and/or direct visualization.

C. Visualization Systems and Procedures

[00104] Visualization can be used throughout an entire decompression procedures or stage(s) of decompression procedures. Visualization systems and components disclosed herein can be incorporated into or used with dilation systems, introducer dilation assemblies, cannula dilation assemblies, instrument cannulas, dilation devices, instrument positioner assemblies, reamer instruments, and other systems and components disclosed herein.

[00105] Figure 44 is a side view of a visualization system 1100 in accordance with one embodiment of the disclosure. The visualization system 1100 can include an illumination instrument 1110 and an access device in the form of a cannula 1120. The cannula 1120 can be similar or identical to the cannula 172 discussed in connection with Figures 21A-21C. The cannula 1120 of Figure 44 can extend through a subject's skin, subcutaneous tissue, and/or a supraspinal ligament. The illumination instrument 1110 can extend through a sidewall 1122 of the cannula 1120 and can direct light toward a working space 1130. In some embodiments, the instrument 1110 passes through an access features 1487 and can include a light source 1140 and a waveguide. The light source 1140 can output light suitable for viewing tissue with the naked eye or with an optical aid, such as loupes. The waveguide can include a flexible fiber optic cable (illustrated) configured to deliver the light from the light source 1140 towards the space 1130.
An inner surface 1144 of the cannula 1120 can reflect the light to enhance light delivery to the working space 1130. In some embodiments, the inner surface 1144 can include one or more optically reflective coatings. In other embodiments, the cannula 1120 can include one or more reflective elements (e.g., mirrors) for directing light out an open distal end 1127 of the cannula 1120. The cannula 1120 can have one or more imaging devices 1146 positioned to image the working space 1130, and the imaging devices 1146 can include one or more light sources oriented to illuminate tissue within its field of view.

Figure 45 is a side view of a visualization system 1200 in accordance with one embodiment of the disclosure. The visualization system 1200 can include an imaging instrument 1210 and a cannula 1220. The imaging instrument 1210 is positioned within a lumen of the cannula 1220 and can provide a field of view for viewing the working space 1130 (including regions of the working space 1130 not viewable by direct viewing). The imaging instrument 1210 can be an endoscope or other imaging device for providing a desired field of view. A coupler 1212 can help keep the imaging instrument 1210 positioned to view the working space 1130 while allowing rotation of the imaging instrument 1210 relative to the cannula 1220. In one embodiment, the imaging instrument 1210 can be rotated 360 degrees to provide complete peripheral viewing of tissue not viewable with the naked eye. The orientation of the field of view can be selected based on the desired peripheral viewing. The imaging instrument 1210 can be a visualization instrument with an endoscope 1216 and a viewing device, such as monitor 1218.

Figure 46 is a side view of a visualization system 1300 in accordance with one embodiment of the disclosure. The visualization system 1300 can include a visualization instrument 1310 extending through a lumen 1340 of a decompression instrument 1350. The decompression instrument 1350 can be identical or similar to the decompression instrument 190 of Figures 26 and 27. A distal end 1342 shown in Figure 46 can be positioned in the field of view 1344 of the instrument 1310. In some embodiments, the visualization instrument 1310 can include a monitor and an endoscope connected to the monitor. The configuration of the visualization instrument 1310 can be selected based on the configuration of the decompression instrument 1350.
[00109] Figure 47 is a side view of the visualization system 1370 including a monitor 1372 and a visualization instrument 1380 extending through a decompression instrument in the form of a rongeur 1400. The field of view of the visualization instrument 1380 can include at least a portion of the rongeur 1400 to accurately positioning jaws 1402a, 1402b. The configuration and position of the instrument 1380 can be selected to provide viewing of the tissue grabbed by the jaws 1402a, 1402b.

[00110] The visualization systems disclosed herein can be utilized with a wide range of different types of decompression instruments. A spinal procedure can be performed while viewing the treatment to help, for example, remove tissue to perform a decompression procedure and avoid damaging non-targeted tissue. For example, viewing can help perform one or more of the steps discussed in connection with Figures 2-7 and 31-43. In some procedures, a spinal procedure can be performed without utilizing additional view, such as fluoroscopic viewing. In some embodiments, visualization systems disclosed herein can be used with fluoroscopic viewing or other imaging techniques. The various embodiments described herein may also be combined to provide further embodiments. For example, features from various instruments can be combined with features and methods disclosed in U.S. Pat. No. 8,012,207; U.S. Pat. No. 8,123,807; U.S. Pat. No. 8,152,837, U.S. App. No. 12/217,662 (U.S. Publication No. 2008/0287997), and U.S. App. No. 13/844,324, which are incorporated by reference in their entitities and a part of the present specification. A wide range of visualization instruments and treatment instruments can be used to address a wide range of symptoms, conditions, and/or diseases, including, without limitation, spinal nerve compression (e.g., spinal cord compression, spinal nerve root compression, or the like), spinal disk herniation, osteoporosis, stenosis, or other diseases or conditions.

[00111] The above detailed descriptions of embodiments of the technology are not intended to be exhaustive or to limit the technology to the precise form disclosed above. Although specific embodiments of, and examples for, the technology are described above for illustrative purposes, various equivalent modifications are possible within the scope of the technology, as those skilled in the relevant art will recognize. For example, while steps are presented in a given order, alternative embodiments may perform steps in a different order. For example, visualization media can be delivered before, during, or after positioning a cannula (e.g., instrument
cannula 172 of Figures 19-21). Additionally, the instruments (e.g., tissue removal instruments, reamer instruments, debulker instruments, dilators, syringes, etc.) can have one or more stops (e.g., depth stops) to inhibit or prevent injury or damage to tissue. Additionally or alternatively, the stops can be incorporated into the cannulas (e.g., cannulas or instruments disclosed herein). The various embodiments described herein may also be combined to provide further embodiments. For example, features from various instruments can be combined with features disclosed in U.S. Pat. No. 8,012,207; U.S. Pat. No. 8,123,807; U.S. Pat. No. 8,152,837, U.S. App. No. 12/217,662 (U.S. Publication No. 2008/0287997), and U.S. App. No. 12/358,010, which are hereby incorporated by reference herein and made a part of this application.

[0012] Where the context permits, singular or plural terms may also include the plural or singular term, respectively. Moreover, unless the word "or" is expressly limited to mean only a single item exclusive from the other items in reference to a list of two or more items, then the use of "or" in such a list is to be interpreted as including (a) any single item in the list, (b) all of the items in the list, or (c) any combination of the items in the list. Additionally, the term "comprising" is used throughout to mean including at least the recited feature(s) such that any greater number of the same feature and/or additional types of other features are not precluded. It will also be appreciated that specific embodiments have been described herein for purposes of illustration, but that various modifications may be made without deviating from the technology. Further, while advantages associated with certain embodiments of the technology have been described in the context of those embodiments, other embodiments may also exhibit such advantages, and not all embodiments need necessarily exhibit such advantages to fall within the scope of the technology. Accordingly, the disclosure and associated technology can encompass other embodiments not expressly shown or described herein.
What is claimed is:

1. A dilation system for sequentially dilating anatomical features to provide access to a treatment site along a subject's spine, the dilation system comprising:
   a first dilation assembly configured to be inserted between adjacent spinous processes of the subject, the first dilation assembly including—
   a first dilator having a distal end, a proximal end, and a lumen extending between the distal and proximal ends, and
   a needle device having a handle and a needle, wherein the needle has an elongate body coupled to the handle and a distal portion configured to protrude from the distal end of the first dilator when the elongate body extends through the lumen of the first dilator; and
   a second dilation assembly movable over the first dilator after the needle device has been removed from the first dilator, the second dilation assembly including—
   an instrument cannula having a distal cannula end, a proximal cannula end, and an instrument passageway extending between the distal and proximal cannula ends, and
   a second dilator including a dilation handle and an elongate dilator configured to move over the first dilator after the needle device has been removed from the first dilator, and wherein the elongate dilator of the second dilator is configured to be removed from the instrument passageway of the instrument cannula after the second dilation assembly has been advanced over the first dilator.

2. The dilation system of claim 1 wherein the first dilation assembly has a locking mechanism with a locked configuration for coupling together the first dilator
and the needle device and an unlocked configuration for allowing the needle device to be removed from the first dilator.

3. The dilation system of claim 1 wherein the second dilation assembly has a locking mechanism with a locked configuration for mechanically coupling together the instrument cannula and the second dilator and an unlocked configuration for allowing the elongate dilator to be removed from the instrument cannula.

4. The dilation system of claim 1 wherein the distal portion of the needle extends distally past the distal end of the first dilator when the needle device and the first dilator are coupled together for insertion between the spinous processes.

5. The dilation system of claim 1 wherein the elongate body of the needle has a longitudinally-extending outer channel, wherein at least a portion of the first dilator is positioned in the outer channel when the first dilator and the needle device are coupled together.

6. The dilation system of claim 1 wherein the elongate dilator of the second dilation assembly has oppositely located outer channels, and wherein the instrument cannula has oppositely located outer cannula channels alignable with the outer channels of the elongate dilator of the second dilator such that the spinous processes move from the outer channels of the elongate dilator to the respective outer cannula channels when the second dilation assembly is moved along the first dilator positioned between the spinous processes.

7. The dilation system of claim 1 wherein the first dilation assembly is configured to cause distraction of the spinous processes; and the second dilation assembly is configured to cause additional distraction of the spinous processes.

8. The dilation system of claim 1 wherein the second dilation assembly is movable over the proximal end of the first dilator and movable along the first dilator.
until the proximal end of the first dilator extends proximally from a proximal opening of the second dilator device and accessible such that a user is capable of pulling the first dilator from the second dilation assembly.

9. The dilation system of claim 1 wherein the dilation handle has an access window through which the first dilator moves when the first dilator is pulled proximally out of the lumen of the second dilator.

10. A dilation system, comprising:

    a first dilation assembly configured to be inserted between spinous processes when the first dilation assembly is in a locked configuration, the first dilation assembly including a first inner instrument with a handle and a first outer instrument, wherein the first inner instrument is configured to be separated from the first outer instrument when the first dilation assembly is in an unlocked configuration; and

    a second dilation assembly movable over the first outer instrument when the second dilation assembly is in a locked configuration, the second dilation assembly including a second inner instrument with a handle and a second outer instrument, wherein the second inner instrument is configured to be removed from the second outer instrument when the second dilation assembly is in an unlocked configuration.

11. The dilation system of claim 10 wherein the first outer instrument of the first dilation assembly is removable from the second dilation assembly when the second dilation assembly is in the locked configuration.

12. The dilation system of claim 10 wherein

    the first inner instrument of the first dilation assembly includes a needle; and

    the first outer instrument is a dilator with a passageway through which the needle extends when the first dilation assembly is in the locked configuration.
13. The dilation system of claim 10 wherein
the second inner instrument of the second dilation assembly includes a dilator
with a passageway through which the first outer instrument of the first
dilation assembly is capable of passing through; and
the second outer instrument of the second dilation assembly is a cannula with
an instrument passageway for receiving one or more instruments.

14. The dilation system of claim 10 wherein
the first dilation assembly includes a first locking mechanism with a locked
configuration for mechanically coupling together the first inner and outer
instruments while the first inner and outer instruments are inserted
together into the subject; and
the second dilation assembly includes a second locking mechanism with a
locked configuration for mechanically coupling together the second inner
and outer instruments while the second inner and outer instruments are
inserted together into the subject.

15. The dilation system of claim 10 wherein at least one of the first outer
instrument and the second inner instrument has a tapered distal end configured to
push apart the spinous processes.

16. The dilation system of claim 10 wherein a proximal end of the first outer
instrument is insertable into a passageway of the second dilation assembly to allow
the second dilation assembly to be advanced along the first outer instrument towards
a distal end of the first outer instrument such that the proximal end of the first outer
instrument extends proximally out of the passageway of the second dilation assembly.

17. The dilation system of claim 16 wherein the handle of the second inner
instrument has an access window for accessing the proximal end of the first outer
instrument when the second dilation assembly has been delivered over the first outer
instrument.
18. A method for accessing a treatment site along a human subject's spine, the method comprising:

inserting an introducer dilation assembly into a human subject such that the introducer dilation assembly is positioned between adjacent spinous processes of the subject, wherein the introducer dilation assembly includes an introducer dilator and a needle assembly positioned in the introducer dilator;

removing the needle assembly from the introducer dilator positioned in the subject;

after removing the needle assembly from the introducer dilator,

moving a cannula dilation assembly over the introducer dilator to position the cannula dilation assembly between the adjacent spinous processes, wherein the cannula dilation assembly includes an instrument cannula and a cannula dilator positioned in the introducer dilator; and

removing the cannula dilator from the instrument cannula.

19. The method of claim 18, further comprising removing the introducer dilator from the cannula dilation assembly before removing the cannula dilator from the instrument cannula.

20. The method of claim 18 wherein removing the needle assembly from the introducer dilator includes:

moving a locking device of the needle assembly from a locked configuration for fixedly coupling together the introducer dilator to the needle assembly to an unlocked configuration for separating the needle assembly and the introducer dilator; and

removing a needle of the needle assembly from the introducer dilator while the introducer dilator is positioned between the adjacent spinous processes.
21. The method of claim 18 wherein removing the cannula dilator from the instrument cannula includes-

moving a locking device of the cannula dilation assembly from a locked configuration for fixedly coupling together the instrument cannula and the cannula dilator to an unlocked configuration for separating the cannula dilator from the instrument cannula; and

removing the cannula dilator from an instrument passageway of the instrument cannula while the instrument cannula is positioned between the spinous processes.

22. The method of claim 18, further comprising delivering a surgical instrument through the instrument cannula while the instrument cannula is positioned between the spinous processes; and performing at least a portion of a spinal decompression procedure on the subject using the surgical instrument while the surgical instrument is positioned in the subject.

23. The method of claim 18, further comprising advancing the cannula dilation assembly into the subject such that the cannula dilation assembly wedges apart the spinous processes.

24. The method of claim 23 wherein the spinous processes include a first spinous process and a second spinous process, the method further comprising:

positioning the first spinous processes in a first channel of the cannula dilator;
positioning the second spinous processes in a second channel of the cannula dilator; and
advancing the cannula dilator into the subject to distract and/or maintain distraction of the first and second spinous processes positioned within the first and second channels.

25. The method of claim 18 wherein inserting the introducer dilation assembly into the human subject includes moving distal ends of the introducer cannula and the needle assembly through the supraspinous ligament of the subject.
26. The method of claim 18 wherein inserting the introducer dilation assembly into the human subject includes moving the introducer assembly using an midline path relative to the subject.

27. A reamer instrument, comprising:
   a reaming assembly including—
   an outer reamer member having a lateral reaming element, an elongate body, and a lumen extending between first and second ends of the outer reamer member, and
   an inner reamer member including a reaming tip and a rod, wherein the rod is positioned in the lumen; and
   a positioner element connected to the inner reamer member and movable between a distal ream position for positioning at least a portion of the reaming tip outside of the outer reamer member and an atraumatic position for positioning the reaming tip within the outer reamer member.

28. The reamer instrument of claim 27 wherein the reaming assembly includes a longitudinal axis, wherein the reaming tip extends in a direction substantially parallel to the longitudinal axis past the entire lateral reaming element when the positioner element is in the distal ream position.

29. The reamer instrument of claim 27 wherein the reaming tip moves into the lateral reaming element when the positioner element moves from the distal reaming position to the atraumatic position.

30. The reamer instrument of claim 27, further comprising:
   a depth stop member movable along the reaming assembly so as to adjust a depth of penetration of the reaming assembly when the reaming assembly extends through a cannula configured to physically contact the depth stop member.
INTERNATIONAL SEARCH REPORT

Box No. II  Observations where certain claims were found unsearchable (Continuation of item 2 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. ☒ Claims Nos.: 18-26 because they relate to subject matter not required to be searched by this Authority, namely:
   Claims 18-26 pertain to methods for treatment of the human body by surgery, and thus relates to a subject matter which the International Searching Authority is not required to search, under PCT Article 17(2)(a)(i) and PCT Rule 39.1(iv).

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 No. III  Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

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

1. Claims 1-17 directed to a dilation system.
2. Claims 27-30 directed to a reamer instrument.

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 fees, this Authority did not invite payment of any additional fees.

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 and, where applicable, the payment of a protest fee.
☐ The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
☐ No protest accompanied the payment of additional search fees.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

A61B 17/70(2006.01)i, A61B 17/3205(2006.01)i

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)

A61B 17/70; A61M 29/00; A61M 5/00; A61B 17/16; A61B 17/3211; A61B 17/00; A61M 5/178; A61B 18/18; A61B 17/88; A61B 17/3205

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

Korean utility models and applications for utility models

Japanese utility models and applications for utility models

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

eKOMPASS/KIPO internal) & Keywords: treat, spinal nerve compression, dilator, inner, outer, needle, cannula, movable, reamer, lateral, distal

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Citation of document, with indication, where appropriate, of the relevant passages

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Further documents are listed in the continuation of Box C.

See patent family annex.

* 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 application or patent but published on or after the international filing date
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  "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

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"&" document member of the same patent family

Date of the actual completion of the international search

24 July 2015 (24.07.2015)

Date of mailing of the international search report

03 August 2015 (03.08.2015)

Name and mailing address of the ISA/KR

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