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(54) **SYSTEMS AND COMPOSITIONS FOR LOCAL IMAGING AND TREATMENT OF PAIN**

SYSTEME UND ZUSAMMENSETZUNGEN FÜR DIE LOKALE DARSTELLUNG UND BEHANDLUNG VON SCHMERZEN

SYSTEMES ET COMPOSITIONS POUR REPRESENTER LOCALEMENT ET TRAITER LA DOULEUR

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Description

[0001] This invention pertains generally to imaging of tissues associated with skeletal joints. More particularly, it relates to identification and/or characterization of localized factors associated with musculoskeletal pain using labeled markers and related imaging tools.

[0002] Chronic back pain (i.e. generally persisting longer than 12 weeks) is among the most prevalent and expensive non-lethal conditions in the United States, and is believed to be the most common cause of disability in persons under 45 years old. The number of people suffering from chronic back pain is estimated to exceed 25% of the overall population. Every year, about 3-4% of the U.S. population is estimated to be disabled temporarily, and about 1% of the working age population is estimated to be disabled totally and permanently, due to intractable back pain. An estimated 11.7 Million patients present medically with chronic back pain. National disability expenses for this prevalent condition range from \$30-\$70 billion per year. Effectively treating this prevalent condition remains among the largest unmet clinical needs in medicine. Properly diagnosing and localizing the source of pain also remains a significant shortcoming on the critical path toward providing such therapy in a targeted manner with predictably successful outcomes.

[0003] Diagnosis of the location, mode, and extent of disc degeneration is often used as a precursor tool to drive therapy for treating back pain. However, such measures are often not specific enough to localize the exact site in or around a degenerating disc where pain is being experienced. Also, a direct correspondence is not always found between disc degeneration and back pain. Consequently, existing imaging modalities that identify (and even quantify) disc anatomy, such as CT or MRI, are not always helpful at localizing sources of back pain in many cases.

[0004] Accordingly, there is still a substantial need for new imaging modalities to objectively, accurately, and specifically identify and localize source(s) of pain, and in particular back pain, and still more particularly lower lumbar back pain. There is in particular such a need with respect to identifying painful discs in an improved way, and to localize within or around those discs the specific site of injury or source of pain in an improved, predictable, dependable manner.

[0005] WO 00/41514 and US 6,491,893 B1 each describe agents for specifically targeting and detecting or treating focal sites of infection or inflammation in a subject.

[0006] US 2003/0139652 A1 describes a system for pain diagnosis, comprising: an input module for receiving inputs of pain areas of a patient onto a three-dimensional human body model which comprises uniformly divided multiple cells; a diagnosis module for deriving diagnosis results, comprising a database for storing data on various pain patterns, a submodule A for checking over a surface of the human body model according to divided multiple

blocks, comparing the inputted pain areas with respect to the blocks, and deriving referred pain patterns with respect to the blocks corresponding to the patient's pain areas from the database, a submodule B for allowing the patient to confirm symptoms of the derived pain patterns and assigning weighted values to confirmed pain patterns, according to degrees of matching upon confirmation, a submodule C for comparing images of the confirmed pain patterns with the inputted pain areas, and a submodule D for calculating degrees of matching between the confirmed pain patterns and the inputted pain areas; and an output module for outputting the diagnosis results derived from the diagnosis module.

15 SUMMARY OF THE INVENTION

[0007] The invention provides a method for preparing a system comprising a targeted agent for performing a medical procedure on a patient having been diagnosed with back pain, characterized by: based upon the back pain diagnosis, preparing a volume of a targeted agent in combination with a delivery assembly in a configuration that is operable to allow delivery of the targeted agent to a region of the patient that comprises at least a portion of a spine, wherein the targeted agent is characterized by differentially binding to a pain factor at a location of a spinal joint associated with a source of the back pain when delivered into the patient; configuring at least one of an energy delivery system to deliver energy to the patient and an imaging system to image the pain factor at a location of a spinal joint; and wherein the targeted agent when delivered into a region of the patient that comprises at least a portion of a spine is further characterized to enhance at least one of (i) diagnostic localization of the source of the back pain at the location in the patient's body when the imaging system is operated in the configuration to image the location of a spinal joint and (ii) selective tissue therapy of the source of the back pain at the location when the energy delivery system is operated in a configuration to deliver energy to the location of a spinal joint; and wherein the targeted agent comprises at least one of: a nerve factor, a blood vessel factor, an inflammatory cytokine, or a combination thereof; or a factor associated with pH or pO₂ in tissue, optionally associated with a relatively low pH or pO₂, respectively, or a binding agent or antibody respectively thereof.

[0008] The invention also provides a system, which comprises a targeted agent for performing a medical procedure on a patient, said targeted agent prepared for dosed delivery into a patient diagnosed with back pain, characterized by:

a delivery assembly for the targeted agent that is configured and operable to allow delivery of the targeted agent into a location of a spinal joint in the patient; and
said targeted agent when delivered to a region of the

patient that comprises at least a portion of a spine via the delivery assembly is characterized to differentially bind to a pain factor at a location of a joint associated with the back pain in a manner further characterized to enhance at least one of (i) diagnostic localization of a source of the back pain at the location of a spinal joint and (ii) selective tissue therapy to a source of the back pain at the location containing the bound pain factor in response to a delivered energy to the region of the patient that comprises at least a portion of a spine; and wherein the pain factor comprises at least one of: a nerve factor, a blood vessel factor, an inflammatory cytokine, or a combination thereof; or a factor associated with pH or pO₂ in tissue, optionally associated with a relatively low pH or pO₂, respectively, or a binding agent or antibody respectively thereof.

[0009] Accordingly, certain aspects of the present invention provide a system, composition of matter, and method that better describe, diagnose, and localize of the sources of pain in and around musculoskeletal joints, and in particular beneficial modes in and around spinal discs in relation to back pain.

[0010] Among the various modes employed according to this aspect, one particular beneficial mode involves artificially labeling substances locally in the area of back pain, such as in a particular beneficial example the spinal motion segment, that are known suspects to pain generation and transmission, such as for example disc, facet joints, and vertebral bodies.

[0011] Two particularly beneficial embodiments according to this mode, useful either alone or in combination, include: (a) labeling nerves, and in particular beneficial embodiments nociceptors, and (b) labeling chemical factors that irritate nerves, (c) labeling cells that produce chemical factors that irritate nerves; and (d) labeling blood vessels that are typically in close approximation to nerves.

[0012] In addition to the significant benefit provided by these approaches for clinical diagnosis, they are also considered highly beneficial in providing new avenues to drive choices for therapeutic approaches.

[0013] One aspect of the invention is a method for conducting a medical procedure related to a localized, active source of pain at a location within a patient. This method includes artificially labeling a pain factor at the location in a manner substantially increasing the ability to image the pain factor with an imaging tool. The labeled pain factor is then labeled in a manner sufficient to selectively differentiate a first concentration of the labeled pain factor at the location versus a second concentration of the labeled pain factor in tissue adjacent to the location.

[0014] According to one highly beneficial mode, the location is associated with a skeletal joint.

[0015] Another mode of this aspect further includes delivering a substantially targeted label into the patient that is adapted to differentially bind to and label a pain factor

associated with the source of pain at the location. The pain factor at the location is artificially labeled by binding the pain factor with the targeted label.

[0016] According to one embodiment, the differential binding comprises specific binding to the pain factor.

[0017] According to another embodiment, the differential binding comprises non-specific binding to the pain factor.

[0018] According to another mode, the pain factor comprises at least one of a nerve factor, an inflammatory factor, a cellular factor, or a blood vessel factor, or a combination thereof.

[0019] In one more particular mode, the pain factor comprises a nerve factor.

[0020] According to one embodiment of this mode, the nerve factor comprises at least one substance associated with at least one of a nerve fiber or a cellular structure associated with the nerve fiber.

[0021] In another embodiment, the nerve factor comprises a substance associated with a nerve fiber. According to one particularly beneficial embodiment, the substance is in particular associated with nociceptors.

[0022] In another more particular mode, the pain factor comprises a blood vessel factor.

[0023] According to one embodiment of this mode, the blood vessel factor comprises at least one of a blood vessel or a substance or structure associated with the blood vessel.

[0024] In another embodiment of this mode, the blood vessel factor comprises a substance or structure associated with microvessels.

[0025] According to another more particular mode, the pain factor comprises a cellular factor.

[0026] According to one embodiment of this particular mode, the cellular factor is associated with a cell that produces at least one inflammatory factor.

[0027] In another embodiment, the cellular factor is associated with at least one inflammatory factor.

[0028] In another embodiment, the cellular factor is associated with cells actively producing inflammatory factors.

[0029] In another embodiment, the cellular factor is associated with an inflammatory cell of a type that is attracted to a second pain factor at the location. According to one particular variation of this embodiment, the inflammatory cell comprises a leukocyte or macrophage.

[0030] According to another more particular mode, the pain factor comprises an inflammatory factor.

[0031] According to another mode, the pain factor comprises a cytokine.

[0032] According to another mode of the present aspect, the pain factor comprises substance P or an analog or derivative or binding agent or antibody thereof.

[0033] According to another mode, the pain factor comprises CGRP or an analog or derivative or binding agent or antibody thereof.

[0034] According to another mode, the pain factor comprises receptor tyrosine kinase A (TrkA) or an analog or

derivative thereof.

[0035] According to another mode, the pain factor comprises a TrkA binding agent or antibody.

[0036] According to another mode, the pain factor comprises a TrkA receptor or a binding agent or antibody thereof.

[0037] According to another mode, the pain factor comprises nerve growth factor (NGF) or an analog or derivative thereof.

[0038] According to another mode, the pain factor comprises an NGF binding agent or antibody.

[0039] According to another mode, the pain factor comprises an NGF antagonist or an analog or derivative thereof.

[0040] According to another mode, the pain factor comprises an NGF-antagonist binding agent or anti-NGF antagonist antibody.

[0041] According to another mode, the pain factor comprises a nerve binding agent or antibody or an analog or derivative thereof.

[0042] According to another mode, the pain factor comprises protein gene product 9.5 (PGP 9.5) or an analog or derivative or binding agent or antibody thereof.

[0043] According to another mode, the pain factor comprises SYN or an analog or derivative or binding agent or antibody thereof.

[0044] According to another mode, the pain factor comprises peripherin or an analog or derivative or binding agent or antibody thereof.

[0045] According to another mode, the pain factor comprises Neurofilament 200kD (NF200) or an analog or derivative or binding agent or antibody thereof.

[0046] According to another mode, the pain factor comprises tissue necrosis factor alpha (TNF- α) or an analog or derivative or binding agent or antibody thereof.

[0047] According to another mode, the pain factor comprises a TNF- α blocker or binding agent or antibody thereof.

[0048] According to another mode, the pain factor comprises macrophage migration inhibitory factor (MIF) or an analog or derivative or binding agent or antibody thereof.

[0049] According to another mode, the pain factor comprises infliximab, or an analog or derivative thereof, or a binding agent or an antibody thereof.

[0050] According to another mode, the pain factor comprises PECAM or an analog or derivative or binding agent or antibody thereof.

[0051] According to another mode, the pain factor comprises CD34 or an analog or derivative or binding agent or antibody thereof.

[0052] According to another mode, the pain factor comprises vascular cell adhesion molecule-1 (VCAM-1) or an analog or derivative or binding agent or antibody thereof.

[0053] According to another mode, the pain factor comprises an interleukin or an analog or derivative or binding agent or antibody thereof.

[0054] According to one embodiment of this mode, the interleukin comprises IL-1 or an analog or derivative or binding agent or antibody thereof.

[0055] According to another embodiment, the interleukin comprises IL-6 or an analog or derivative or binding agent or antibody thereof.

[0056] According to another embodiment, the interleukin comprises IL-8 or an analog or derivative or binding agent or antibody thereof.

[0057] According to another mode of the present aspect, the pain factor comprises prostaglandin E2 (PGE₂) or an analog or derivative or binding agent or antibody thereof.

[0058] According to another mode, the pain factor comprises a factor associated with pH in tissue or a binding agent or an antibody thereof.

[0059] According to one embodiment of this mode, the labeled pain factor is indicative of a relatively low pH below a predetermined threshold at the location.

[0060] According to another mode, the pain factor comprises a factor associated with pO₂ in tissue or a binding agent or an antibody thereof.

[0061] In one embodiment according to this mode, the labeled pain factor is indicative of a relatively low pO₂ at the location.

[0062] According to another mode, the pain factor comprises glial fibrillary acidic protein (GFAP) or an analog or derivative or binding agent or antibody thereof.

[0063] According to another mode, the pain factor comprises synuclein (SYN) or an analog or derivative or binding agent or antibody thereof.

[0064] According to another mode of the present aspect, the targeted label comprises at least one of a nerve factor, a blood vessel factor, a cellular factor, an inflammatory factor, or an antibody thereof.

[0065] According to one embodiment of this mode, the targeted label comprises a nerve factor or a binding agent or an antibody thereof.

[0066] In one variation according to this embodiment, the nerve factor comprises at least one substance associated with at least one of a nerve fiber or a cellular structure associated with the nerve fiber or an antibody thereof.

[0067] In another variation, the nerve factor comprises a substance associated with a nerve fiber or a binding agent or an antibody thereof.

[0068] In another embodiment, the targeted label comprises a blood vessel factor or a binding agent or an antibody thereof.

[0069] In one variation of this embodiment, the blood vessel factor comprises a substance associated with a structure of a blood vessel or a binding agent or an antibody thereof.

[0070] In another variation, the blood vessel factor comprises a substance associated with a structure of a microvessel or a binding agent or an antibody thereof.

[0071] According to another embodiment, the targeted label comprises a cellular factor or a binding agent or an

antibody thereof.

[0072] In one variation, the cellular factor is associated with a cell that produces at least one inflammatory factor, or a binding agent or an antibody thereof.

[0073] In another variation, the cellular factor is associated with at least one inflammatory factor or a binding agent or an antibody thereof.

[0074] In another variation, the cellular factor is associated with an intervertebral disc cell that is actively producing inflammatory factors, or a binding agent or an antibody thereof.

[0075] In another variation, the cellular factor is associated with an inflammatory cell of a type that is attracted to the pain factor at the location, or a binding agent or an antibody thereof.

[0076] According to one feature of this variation, the inflammatory cell comprises a leukocyte, or a binding agent or an antibody thereof.

[0077] According to another embodiment, the targeted label comprises an inflammatory factor, or a binding agent or an antibody thereof.

[0078] In one variation of this embodiment, the inflammatory factor comprises a cytokine, or an analog or derivative thereof, or a binding agent or an antibody thereof.

[0079] According to another mode of the present aspect, the targeted label comprises a binding agent or antibody to substance P.

[0080] According to another mode, the targeted label comprises a binding agent or antibody to calcitonin gene-related peptide (CGRP).

[0081] According to another mode, the targeted label comprises a TrkA antibody or binding agent.

[0082] According to another mode, the targeted label comprises nerve growth factor (NGF), or an analog or derivative thereof.

[0083] According to another mode, the targeted label comprises a NGF binding agent or an anti-NGF antibody.

[0084] According to another mode, the targeted label comprises a NGF antagonist or a binding agent or an antibody thereof.

[0085] According to another mode, the targeted label comprises an anti-NGF antagonist antibody or binding agent.

[0086] According to another mode, the targeted label comprises a nerve antibody or binding agent.

[0087] According to another mode, the targeted label comprises PGP 9.5, or an analog or derivative thereof, or a binding agent or an antibody thereof.

[0088] According to another mode, the targeted label comprises a binding agent or antibody to peripherin.

[0089] According to another mode, the targeted label comprises Neurofilament 200kD (NF200), or an analog or derivative thereof, or a binding agent or an antibody thereof.

[0090] According to another mode, the targeted label comprises TNF- α , or an analog or derivative thereof, or a binding agent or an antibody thereof.

[0091] According to another mode, the targeted label

comprises a TNF- α blocker.

[0092] According to another mode, the targeted label comprises infliximab, or an analog or derivative thereof, or a binding agent or an antibody thereof.

[0093] According to another mode, the targeted label comprises a PECAM binding agent or antibody.

[0094] According to another mode, the targeted label comprises a binding agent or antibody to CD34.

[0095] According to another mode, the targeted label comprises an interleukin binding agent or antibody.

[0096] In one embodiment of this mode, the interleukin binding agent or antibody comprises an IL-1 binding agent or antibody.

[0097] In another embodiment of this mode, the interleukin binding agent or antibody comprises an IL-6 binding agent or antibody.

[0098] In another embodiment of this mode, the interleukin binding agent or antibody comprises an IL-8 binding agent or antibody.

[0099] According to another mode of the present aspect, the targeted label comprises a binding agent or antibody to PGE₂.

[0100] According to another mode, the targeted label comprises a binding agent or antibody to MIF.

[0101] According to another mode, the targeted label comprises an antibody or binding agent to a factor associated with pH in tissue.

[0102] According to one embodiment of this mode, the labeled pain factor is indicative of a relatively low pH below a predetermined threshold at the location.

[0103] According to another mode, the targeted label comprises an antibody or binding agent to a factor associated with pO₂ in tissue.

[0104] According to one embodiment of this mode, the labeled pain factor is indicative of a relatively low pO₂ at the location.

[0105] According to another mode, the targeted label comprises a radioactive material.

[0106] According to one embodiment of this mode, the targeted label comprises a radio-labeled TNF- α antibody, or an analog or derivative thereof.

[0107] According to another embodiment, the targeted label comprises radiolabeled iodine. In one variation of this embodiment, the radiolabeled iodine comprises I-125.

[0108] According to another mode, the targeted label comprises a nanoparticle.

[0109] According to another mode, the targeted label comprises gold.

[0110] According to another mode, the targeted label comprises iron oxide.

[0111] According to another mode, the targeted label comprises gadolinium.

[0112] According to another mode of the present aspect, the method further includes imaging the labeled pain factor using an imaging tool that comprises a phosphor imaging plate.

[0113] According to another mode, the method in-

cludes imaging the labeled pain factor using MRI.

[0114] According to another mode, a first binding agent is delivered into the body that is adapted to bind to a first pain factor. The targeted label is delivered into the patient's body after the first binding agent is bound to the first pain factor. The targeted label is adapted to bind to a site located on the bound combination of the first binding agent and the first pain factor.

[0115] According to one embodiment, the first binding agent comprises a bi-specific antibody with a first binding site adapted to bind to the first pain factor and a second binding site adapted to bind to the targeted label.

[0116] According to another mode, the targeted label comprises a cell bound to an antibody having an exposed binding site that is adapted to bind to the pain factor.

[0117] According to another mode, the method further includes conducting a therapeutic procedure in a substantially localized manner to the location where the targeted labeled pain factor is locally imaged.

[0118] In one embodiment of this mode, the therapeutic procedure is adapted to substantially alleviate generation or transmission of pain at the location.

[0119] According to another embodiment, the therapeutic procedure is adapted to substantially ablate at least one nerve at the location.

[0120] In another embodiment, the therapeutic procedure comprises delivering at least one therapeutic chemical in a substantially localized manner to the location.

[0121] In another embodiment, the therapeutic procedure comprises delivering a therapeutic dose of energy in a substantially localized manner to the location.

[0122] In one variation of this embodiment, the therapeutic procedure further comprises ablating at least one nerve at the location with the therapeutic dose of energy.

[0123] In another variation, the therapeutic procedure further comprises delivering ultrasound energy to the location. In a further variation, the method further includes delivering the ultrasound energy in a directed manner locally into the location from a second location. In still a further variation, the second location is outside of the patient, and the ultrasound energy is delivered via high intensity focused ultrasound (HIFU) that is adapted to focus the ultrasound energy to the location. In yet another variation, the second location is adjacent to the location within the patient, and the ultrasound energy is delivered via a directional ultrasound probe. In still a further feature of this variation, the second location is adjacent to an intervertebral disc and the location receiving the directional ultrasound therapy is within the intervertebral disc.

[0124] According to another variation of the present embodiment, the therapeutic dose of energy comprises thermal energy.

[0125] According to another variation, the therapeutic dose of energy comprises electrical energy. In one further variation, the method involves delivering the electrical energy via a radiofrequency (RF) probe.

[0126] According to another variation, the therapeutic dose of energy comprises microwave energy.

[0127] According to another variation, the therapeutic dose of energy comprises light energy.

[0128] According to another mode of the present aspect, the location comprises at least a portion of an intervertebral disc.

[0129] According to another mode, the location comprises a region of tissue located within only a portion that is equal to less than an entire circumference of an intervertebral disc.

[0130] In one embodiment of this mode, the portion comprises a region of tissue located within less than or equal to one-half of the circumference of the intervertebral disc.

[0131] In one variation of this embodiment, the portion comprises a region of tissue located within less than or equal to one-quarter of a circumference of the intervertebral disc.

[0132] According to another mode of the present aspect of the invention, the location comprises an end-plate associated with a vertebral body.

[0133] According to another mode, the location comprises a facet joint.

[0134] The method of the present aspect according to another mode includes delivering the targeted label in a localized manner to the location.

[0135] One embodiment of this mode further includes injecting the targeted label into a region of tissue associated with the location using a local injection assembly.

[0136] Another embodiment includes delivering the targeted label systemically to the patient.

[0137] One further embodiment includes injecting the targeted label into the patient's systemic blood circulation.

[0138] Another further embodiment includes delivering the targeted label into the patient's gastrointestinal system.

[0139] Another mode includes artificially labeling the pain factor at multiple said locations by binding the pain factor with the targeted label delivered into the patient. The labeled pain factor is then imaged with an imaging tool adapted to image at least one of the targeted label or the labeled pain factor and in a manner sufficient to differentiate a first concentration of the labeled pain factor at the multiple said locations versus a second concentration of the labeled pain factor in tissue adjacent to the multiple said locations.

[0140] According to one embodiment of this mode, the method further includes conducting at least one therapeutic procedure in a substantially localized manner to each of the locations where the targeted labeled pain factor is locally and selectively imaged.

[0141] Another aspect of the invention involves a system for treating pain at a location within a body of a patient. This aspect includes a targeted label that is adapted to bind to and label a pain factor associated with a source of pain at the location. Also included is a delivery assembly that is adapted to deliver the targeted label into the patient. An imaging system also included in the system

is adapted to image at least one of the targeted label or the labeled pain factor and in a manner sufficient to selectively differentiate a first concentration of the labeled pain factor at the location versus a second concentration of the labeled pain factor in tissue adjacent to the location. A therapeutic device assembly is also included, and is adapted to provide therapy in a substantially localized manner that is substantially isolated to the location.

[0142] According to one mode of this aspect, the targeted label is adapted to bind and label a pain factor associated musculoskeletal joint pain, and the location is associated with at least one musculoskeletal joint.

[0143] According to one embodiment, the therapeutic device assembly comprises an energy delivery assembly that is adapted to deliver a therapeutic dose of energy in a substantially localized manner that is substantially isolated to the location associated with the musculoskeletal joint.

[0144] According to one further embodiment, the energy delivery assembly is adapted to be delivered into the patient to a position at or adjacent to the location.

[0145] According to another further embodiment, an introducer is provided in the system and is adapted to deliver the energy delivery assembly to the location.

[0146] In one variation of this embodiment, the introducer comprises a needle assembly. This may provide the additional feature in that the needle assembly is adapted to be advanced through bone and to deliver the therapeutic device assembly to a position within the bone. According to another further feature, the therapeutic device assembly may be adapted to ablate an intraosseous nerve within the bone and that is associated with pain related to the labeled pain factor visualized at the location. In another further beneficial feature, the needle assembly is adapted to be advanced through bone of a vertebral body and to deliver the therapeutic device assembly to a position within the vertebral body associated with a basivertebral nerve, and the therapeutic device assembly is adapted to ablate the basivertebral nerve from the position.

[0147] According to another mode of the present aspect, the therapeutic device assembly comprises a radiofrequency (RF) current ablation assembly.

[0148] In one embodiment, the RF current ablation assembly comprises a first electrode and a second electrode adapted to be positioned at first and second positions adapted to straddle at least a portion of the basivertebral nerve. The RF current ablation assembly is adapted to deliver the RF current between the first and second electrodes sufficient to ablate nerve tissue between the first and second positions.

[0149] According to one variation of this embodiment, the RF current ablation assembly comprises a delivery probe with an elongated body that carries the first and second electrodes in a bipolar lead assembly arrangement.

[0150] According to another mode of the present embodiment, the targeted label is adapted to bind and label

a pain factor comprising at least one of a nerve factor, a blood vessel factor, a cellular factor, an inflammatory factor, or an antibody thereof.

[0151] It is to be appreciated that further more detailed particularly beneficial modes provided hereunder are contemplated with respect to the present aspect described. In particular, further modes of the present aspect include the various beneficial examples for pain factors and targeted labels described for use under the method aspect of the invention described above

[0152] According to another mode, the system further includes an imaging tool that is adapted to image the labeled pain factor in a manner sufficient to differentiate a first concentration at the location associated with pain versus a second concentration at a second location adjacent to the location and associated with less pain that at the location.

[0153] Another aspect of the invention is a method for imaging and identifying a localized, active source of pain at a location associated with a region of tissue in a patient, such as in particular beneficial further modes a skeletal joint in a patient, and in still further beneficial more detailed modes spinal joints in a patient. This method includes delivering a substantially targeted label into the patient that is adapted to differentially bind to and label a pain factor. A pain factor that is resident at the location is artificially labeled by binding the pain factor with the targeted label delivered into the patient. The labeled pain factor is imaged with an imaging tool adapted to image at least one of the label or the labeled pain factor and in a manner sufficient to differentiate a first concentration of the labeled pain factor at the location versus a second concentration of the labeled pain factor in tissue adjacent to the location.

[0154] According to various modes of this aspect, the pain factor may be related to at least one of a nerve fiber, a substance associated with a nerve fiber, a blood vessel, a substance associated with a blood vessel, a cell actively producing at least one inflammatory factor, a cell attracted to inflammation or other pain factors, or a chemoinflammatory factor, or a combination thereof.

[0155] Another aspect of the invention is a system for identifying or characterizing a property of tissue associated with a skeletal joint. Such aspect may further include any one or more of the various aspects, modes, embodiments, variations, or features herein shown or described, or combinations thereof.

[0156] According to one mode of this aspect, the system is adapted to provide information indicative of a degree of a property of at least a portion of an intervertebral disc.

[0157] Another aspect is a system for identifying or characterizing a property of tissue associated with a skeletal joint in a patient. This includes labeling at least one of: pain factors, nerve factors, blood vessel factors, cellular factors, or inflammation factors. Or, the system may include a combination of one or more of the foregoing.

[0158] According to one mode of this aspect, the infor-

mation is related to a degree of a property of at least a portion of an intervertebral disc.

[0159] Another aspect of the invention is a system for characterizing at least a portion of an intervertebral disc with respect to a degree of a property of that disc, such as in particular related to pain or degeneration. This system includes a labeled marker delivery system and a labeled marker imaging system. The labeled marker imaging system provides information that is useful to indicate at least in part the degree of the property.

[0160] According to one further embodiment of the foregoing aspects and modes, the respective system is adapted to produce the information based on either or both of an annular portion or a nucleus portion of the intervertebral disc.

[0161] According to another embodiment, the system is adapted to display a geographical representation related to the spatial concentration of the labeled factor, and a portion of the geographical representation provides the information.

[0162] According to another embodiment, the information is adapted to distinguish a degree of degradation of the disc. According to one highly beneficial further embodiment, the information is adapted to distinguish as to the degree of degradation by reference to a Thompson scale.

[0163] According to another embodiment, the property comprises at least one of pain, or at least one factor that correlates with pain.

[0164] According to another embodiment, the information is related to ratios of concentration of one or more pain factors.

[0165] According to another embodiment, the information is related to presence of secondary or other indirect materials that generally, though indirectly, correlate well with presence of other more direct pain factors.

[0166] According to another embodiment, the information relates to at least one chemical constituent of an intervertebral disc.

[0167] According to another embodiment, the property comprises at least one of a degree of dehydration of the disc, a degree of breakdown of a proteoglycan matrix of the disc, and a degree in a breakdown of a collagen matrix.

[0168] According to another embodiment, the system further includes a radiolabel imaging system that is adapted to produce the information.

[0169] Another aspect of the invention is a method for identifying or characterizing a property of tissue associated with a skeletal joint. One or more of the foregoing method aspects, modes, embodiments, variations, or features herein described, or combinations thereof, may be employed to advance this method.

[0170] One further mode of this aspect further includes providing information indicative of a degree of a property of at least a portion of an intervertebral disc.

[0171] Another aspect is a method for identifying or characterizing a property of tissue associated with a skeletal joint in a patient, and includes at least one of the following steps: labeling a pain factor in the tissue; imaging the labeled pain factor in the tissue; comparing different imaged regions having different concentrations of the labeled pain factor; identifying a location of increased presence of pain factors based upon the comparison; and treating the location with local treatment modality based upon the identification. Or a combination of one or more of the foregoing may be used.

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10 **[0172]** One mode of this aspect includes determining a degree of a property of at least a portion of an intervertebral disc based upon the information.

15 **[0173]** Another aspect of the invention is a method for characterizing at least a portion of an intervertebral disc with respect to a degree of a property thereof, and includes capturing a signal related to the portion using a signal imaging system. The signal imaging system provides information that indicates at least in part the degree of the property.

20 **[0174]** According to one embodiment of the various method aspects and modes just described, the information produced is based on either or both of an annular portion or a nucleus portion of the intervertebral disc.

25 **[0175]** In another embodiment, a curve is displayed that is related to the presence of the labeled pain factor, and wherein a portion of the curve provides the information.

30 **[0176]** Another embodiment includes distinguishing a degree of degradation of the disc based upon the information. A still further embodiment includes distinguishing the degree of degradation of the disc in relation to a Thompson grade based upon the information.

35 **[0177]** Another embodiment includes correlating the disc with degree of pain, or at least one factor that correlates with pain, based upon the information.

[0178] According to another embodiment, the information is related to a ratio of magnitude of a signal imaged that corresponds with the amount of labeled pain factor in a given area or volume of tissue.

40 **[0179]** According to another embodiment, the information is related to a cytokine, a precursor material thereof, an analog or derivative thereof, or a metabolite or degradation product thereof.

45 **[0180]** According to another embodiment, the information relates to at least one chemical constituent of an intervertebral disc.

[0181] According to another embodiment, the property relates to at least one of a degree of dehydration of the disc, a degree of breakdown of a proteoglycan matrix of the disc, and a degree in a breakdown of a collagen matrix.

[0182] Another embodiment includes producing the information at least in part using a radiation imaging system.

55 **[0183]** Another aspect is a method for preparing a system for performing a medical procedure on a patient, comprising: diagnosing the patient with pain; and based upon the diagnosis, preparing a volume of a targeted

agent for delivery into the patient. The prepared volume of targeted agent is configured to differentially bind to a pain factor associated with the pain in a manner adapted to enhance at least one of (i) diagnostic localization of the pain and (ii) selective tissue therapy in an area associated with the bound pain factor in response to a delivered energy to the area.

[0184] Another aspect is a system for performing a medical procedure on a patient, comprising: a therapeutic volume of a targeted agent prepared for delivery into a patient diagnosed with pain and that is configured to differentially bind to a pain factor associated with the pain in a manner adapted to enhance at least one of (i) diagnostic localization of the pain and (ii) selective tissue therapy to a location containing the bound pain factor in response to a delivered energy to an area containing the location.

[0185] Another aspect is a method for selectively treating one or more tissue regions associated with pain in a patient, comprising delivering a targeted agent into the patient configured to differentially bind to a pain factor associated with the pain; and allowing the delivered targeted agent to differentially bind to the pain factor so as to form a differentially bound pain factor; and delivering energy into the patient in a manner that differentially treats the one or more regions associated with the differentially bound pain factor.

[0186] Another aspect is a system for selectively treating one or more tissue regions associated with pain in a patient, comprising: a volume of targeted agent; and an energy delivery system that is configured to deliver energy into the patient. The volume of targeted agent is configured for delivery into a patient and to differentially bind to a pain factor associated with the pain in a manner such that tissue regions containing a first concentration of the differentially bound pain factor exhibit a differential and selective therapeutic response to the delivered energy versus other regions with lower concentrations of the differentially bound pain factor.

[0187] Each aspect, mode, embodiment, variation, or feature herein described is considered independently beneficial without requiring combination with the others. However, such further combinations and sub-combinations thereof are also considered yet further beneficial independent aspects invention. For example, where particular modes, embodiments, variations, or features are herein described with respect to one aspect hereunder, it is to be appreciated by one of ordinary skill that such description is further applicable to other aspects also described though such particular combination may not be specifically mentioned. In further example, a more detailed description provided with respect to a method aspect may provide information that is to be clearly combined as further development of a similar system-related aspect or description, or visa versa.

[0188] Further aspects of the invention will be brought out in the following portions of the specification, wherein the detailed description is for the purpose of fully disclos-

ing preferred embodiments of the invention without placing limitations thereon.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWING(S)

[0189] The invention will be more fully understood by reference to the following drawings which are for illustrative purposes only:

FIG. 1 shows a schematic of certain cascades associated with inflammation and pain.

FIGS. 2A-D shows stained cross-sectioned histology slides indicating presence of certain factors associated with pain as follows, wherein "N" is nucleus pulposus, "A" is annulus fibrosus, and "G" designates growth plate.

FIG. 2A shows a mid-sagittal section of normal mouse-tail disc demonstrating TNF-alpha localization in periphery of nucleus pulposus (brown stain). FIG. 2B shows a normal mouse disc wherein localization of TNF-alpha is present in the hypertrophic zone of the growth plate as generally expected.

FIG. 2C shows in the compressed disc wherein increased amounts of TNF-alpha are apparent within the nucleus and inner annulus.

FIG. 2D shows increased TNF-alpha in the nucleus, inner annulus and irregularities in growth plate observed in compressed disc.

FIG. 3 shows a schematic view of a mouse 30 according to an experimental model wherein the mouse tail 36 is injured by a fixture 40 for evaluating pain factors.

FIG. 4 shows an experimental set-up related to the mouse injury model illustrated in FIG. 3, wherein a series of mice 30 are positioned for viewing their respective tails via a phosphor imaging plate 50.

FIG. 5 shows an image 60 taken from a phosphor imaging plate according to the set-up shown in FIG. 4 for four treatment mice and one control mouse tail (located centrally in the figure).

FIG. 6 shows a schematic view of MAPK signaling pathways associated with certain pain factors.

FIG. 7 shows a schematic view of a NF- κ B pathway associated with certain pain factors.

FIG. 8 shows a schematic view of a prostaglandin pathway associated with certain pain factors.

DETAILED DESCRIPTION OF THE INVENTION

[0190] Referring more specifically to the drawings, for illustrative purposes the present invention is embodied in the systems and methods generally shown in or illustrated by reference to FIG. 1 through FIG. 8. It will be appreciated that the apparatus may vary as to configuration and as to details of the parts, and that the method may vary as to the specific steps and sequence, without

departing from the basic concepts as disclosed herein.

Label Disc Features Associated With Pain

[0191] Discogenic pain is generally believed to be a multifactoral phenomenon in many cases. In particular, three illustrative factors are summarized in varying levels of detail here as examples that are considered contributors in various ways to (or otherwise indicative of) the generation or transmission of discogenic pain. It is believed that these illustrative factors frequently act as a co-existent combination, often acting simultaneously. These types of factors are summarized as follows.

[0192] One such factor type relates to the presence of nociceptors. Normally, intervertebral discs are substantially avascular and only sparsely innervated at the outer margins of the disc annulus. These unmyelinated, substance P (SP) or calcitonin gene-related peptide (CGRP) containing fibers are typically unresponsive and termed silent nociceptors [Cavanaugh, 1996]. SP and CGRP are believed to be the sensory transmitters of nociceptive information. As degeneration proceeds, nerves can follow microvessels and grow deeper into discs, which may occur for example either peripherally or via the endplate. This nerve and vessel in-growth is facilitated by degeneration-related decreases in disc pressure and proteoglycan content.

[0193] A second such factor type is generally embodied by the need for the intradiscal nociceptors to be sensitized, and thus generally involves agents providing such sensitization. This can occur for example via cytokines, which are typically small, secreted proteins that mediate and regulate inflammation. Elevated levels of certain cytokines have been measured in human discs, and are associated with degeneration and pain. Such major cytokines have been observed to include interleukin-1, -6, and -8, tissue necrosis factor-alpha (TNF- α), macrophage migration inhibitory factor (MIF), and prostaglandin E₂ (PGE₂). The source of cytokines can be circulating inflammatory cells, such as for example in the case of herniated discs, or disc cells, such as for example in the case of contained disc degeneration. These pro-inflammatory stimuli can trigger cells to initiate a number of catabolic programs meant to stimulate tissue repair and remodeling that includes production of matrix metalloproteinases 1, 9 and 13. During this wound healing process, cytokines are also often involved in stimulating angiogenesis and granulation tissue formation.

[0194] In one particular beneficial embodiment of the present invention, cytokines and/or their cell-surface receptors are imaged at sites of inflammation *in vivo* using labeled markers, such as radiolabels. In particular beneficial examples, cytokines are tagged with one or more of the following, without limitation: iodine-123, iodine-125, iodine-131, technetium-99m, fluorine-18, or indium-111. In addition, positron-emitting radioisotopes (for example and without limitation fluorine-18) can be imaged using positron emission tomography (PET) or positron

emission tomography-computed tomography (PET-CT). Other radiolabeled compounds can be imaged for example using single photon emission computerized tomography (SPECT).

[0195] It is also to be appreciated that MRI may be employed according to further embodiments for visualizing or observing accumulation or binding of various labeled markers variously herein described, such as for example in applying gadolinium as a marker tagged to or conjugated with certain labels to be bound to pain factors. Moreover, nanoparticles such as gold or iron oxide may be used as labels or markers to bind and thereafter be viewed or selectively targeted for therapy using appropriate visualization or treatment modalities, respectively.

[0196] A third such factor related to discogenic back pain involves disc depressurization that leads to mechanical instability while a pre-stress in the annulus and inter-spinal ligaments is diminished. Depressurization and instability, in turn, lead to abnormal internal disc stress that may stimulate nerves, leading to discogenic pain. Abnormal disc stress may also cause disc cells to be pro-inflammatory, compounding the adverse effects of an abnormal mechanical environment.

Labeling and Imaging Nerve Factors

[0197] According to certain particular embodiments, one or more materials associated with nerves in or around intervertebral discs are labeled with markers that are imaged for localization of pain. This is premised in part on the presence of certain such factors as indicators that pain may originate or transmit in the area. These embodiments include, without limitation, labeling structures or substances associated with nerves themselves. Further detailed modes of this include labeling substances within nerves, such as in particular but without limitation substance P or "CGRP". Other nerve fiber factors, substances or components that may be labeled according to such further embodiment(s) include, without limitation: TRK- α ; anti-TRK- α antibody; nerve growth factor (NGF); anti-NGF antibody; NGF antagonist; anti-NGF antagonist antibody; PGP 9.5; SYN; peripherin; or other form of nerve antibodies or related materials in general. Other materials such as neurofilament 200kD (NF200) [Johnson, 2001; Ashton, 1994] may also be the target of such labeling and subsequent imaging.

[0198] As apparent from these highly beneficial illustrative embodiments just noted immediately above (and elsewhere herein), endogenous substances such as TrkA or NGF may be targeted as the pain factor for labeling, or related antibodies or other substances having particular binding affinity or specificity to such resident materials may be bound to them in the area of pain and then thereafter provide the binding site for targeted labels to be subsequently delivered. In this regard, it is to be appreciated that various forms binding agents are broadly contemplated hereunder this description, though they

may not be particularly antibodies affecting function of the target for binding. For example but without limitation, an antibody mimetic may be employed according to the present embodiments. Furthermore, various such substances described hereunder as targeted pain factors may be themselves labeled as markers and delivered to other targets. For example, NGF may be labeled and artificially delivered as the agent to mark TrkA as the targeted pain factor for imaging. In each of these different types of exemplary cases, the ultimate target for labeling via a separately delivered agent (e.g. whether the target is an endogenous resident substance or an artificially delivered substance) is considered a "nerve factor" as a pain factor according to the present embodiments.

[0199] The following description provides further understanding of the role of these types of chemicals and other materials with respect to these present embodiments. Further description of the benefits of various particular illustrative examples are also provided elsewhere herein for a further understanding.

[0200] The intervertebral disc is normally avascular and only sparsely innervated at the outer layers of the annulus fibrosus and the vertebral endplate [Fagan, 2003]. The outer 1/3 of the posterior annulus is believed to be most typically innervated by the afferent fibers from the sinovertebral nerve, which is considered a 'recurrent branch' of the ventral ramus of the spinal nerve at the same level [Nakamura, 1996]. The ventral and lateral aspects of the annulus are believed to be most typically innervated by the dorsal root ganglion (DRG) [Aoki, 2004]. Also, it has been reported that sensory fibers from upper level DRGs are believed to most typically innervate the dorsal portion of discs via the paravertebral sympathetic trunk [Ohtori, 2001].

[0201] The endplate is also suggested to be innervated by the basivertebral nerve, which as further suggested may be a branch of the sinovertebral nerve entering the vertebral body through the posterior neurovascular foramen [Antonacci, 1998].

[0202] Nerves usually accompany blood vessels, but can be found as isolated nerves in disc matrix. These non-vessel-associated fibers found in back pain patients have been observed to express growth-associated protein 43 (GAP43) as well as SP [Freemont, 1997]. Small disc neurons contain CGRP and also express the high-affinity nerve growth factor (NGF) receptor, tyrosine kinase A (trkA)[Aoki, 2004]. Disc inflammation has been observed to cause an increase in CGRP positive neurons [Aoki, 2004]. A recent study showed that NGF is expressed in microvascular blood vessels in a painful lumbar disc, and that there are trkA (TRK- α) expressing nerve fibers adjacent to the vessels that enter painful discs primarily through the endplate [Freemont, 2002; Brown, 1997]. Along with nerves growing into degenerated discs are specialized nerve support cells termed 'glia' or Schwann cells localized using glial fibrillary acidic protein (GFAP) [Johnson, 2001].

[0203] Accordingly, various such materials may pro-

vide the requisite binding affinity or specificity to painful regions (or highly innervated regions) to play the role as the labeled marker agent for delivery to pain factor targets. Or, these materials may provide the particular target as the pain factor to be labeled with selectively bound markers according to various embodiments of the present invention. In one particular beneficial example, TrkA antibody (or other binding agent) is labeled and delivered as a marker for binding and visualization at a location associated with pain. In another beneficial example, NGF itself is labeled and delivered as a marker to itself bind to TrkA. In further embodiments, the resident quantities of these materials are treated as the pain factors themselves for targeted labeling, e.g. using anti-bodies or other agents with beneficial binding affinity and/or specificity to these types of resident compounds in painful regions.

[0204] The following Published PCT Patent Applications are herein incorporated in their entirety by reference thereto: WO 2004/032870 to Shelton et al. as "Inventors" and Rinat Neuroscience Corp. as "Applicant" for "Methods for Treating Post-Surgical Pain By Administering a Nerve Growth Factor Antagonist and Compositions Containing the Same"; WO 2004/058184 to Shelton et al. as "Inventors" and Rinat Neuroscience Corp. as "Applicant" for "Anti-NGF Antibodies and Methods Using Same"; WO 2004/073653 to Shelton et al. as "Inventors" and Rinat Neuroscience Corp. as "Applicant" for "Methods for Treating Pain by Administering A Nerve Growth Factor Antagonist and an NSAID And Compositions Containing The Same"; WO 2004/096122 to Shelton et al. as "Inventors" and Rinat Neuroscience Corp. as "Applicant" for "Methods for Treating Pain By Administering A Nerve Growth Factor Antagonist And An Opioid Analgesic and Compositions Containing The Same"; and WO 2005/000194 to Shelton et al. as "Inventors" and Rinat Neuroscience Corp. as "Applicant" for "Methods for Treating Post-Surgical Pain By Administering An Anti-Nerve Growth Factor Antagonist Antibody and Compositions Containing The Same."

[0205] The various compositions and methods described in these incorporated references may be adopted where appropriate to one of ordinary skill as label/marker vehicles and/or pain factor targets according to further embodiments of the various aspects and modes of the present invention herein described. For example without limitation, NGF antagonists, anti-NGF antibodies, anti-NGF antagonist antibodies, and various combinations or blends of these, or analog or derivatives thereof, may be so incorporated as further embodiments of the aspects herein described. Moreover, additional compounds may also be included in the agent delivery scheme, or as additional targets for labeled markers, such as for example opioids, NSAID, or other molecules or drug agents related to pain therapy.

Labeling and Imaging Blood Vessel Factors

[0206] Since blood vessels typically run along side and co-existent with nerves, factors related to blood vessels may also be labeled and imaged as indicia regarding vascularity itself, or as a measure of concomitant innervation in an area. Such constitutes a further embodiment contemplated hereunder, and described in some further detail as follows. In one regard, PECAM and/or CD34 [Freemont, 2002; Brown, 1997] may be appropriate targets as factors related to blood vessels and thus indicating their presence in a particular location or region. Another example of an appropriate target includes GFAP for endothelial cells [Johnson, 2001]. Other microvessel-related factors are considered as included, though not specifically listed here, as would be apparent to one of ordinary skill based upon review of this disclosure and other available information.

Labeling and Imaging Inflammatory Factors

[0207] According to still further embodiments contemplated hereunder, inflammatory factors themselves may be labeled with targeted markers and imaged as indicators of pain in a location or area. One exemplary type of such factor includes cytokines, such as for example but without limitation (though considered of particular benefit): *tnf- α* , or certain interleukins such as IL-1, 6, or 8 (or other interleukins). Another exemplary pro-inflammatory factor includes MIF and PGE₂.

[0208] Other factors considered indicative of certain activities or environmental considerations believed linked to pain, and thus appropriate targets for labeling and imaging using targeted markers, include: pH (e.g. in particular marking low pH as indicator of pain; or O₂ levels, e.g. in particular marking low O₂ as indicator of pain).

[0209] Cytokines, in the present context, are generally described as small, secreted proteins that mediate and regulate inflammation. They generally act over short distances, short times, and at very low concentrations. They typically function by binding to specific membrane receptors, which often then signal the cell via second messengers (discussed below) to alter gene expression. Responses to cytokines include increasing or decreasing expression of membrane proteins (including cytokine receptors), cell proliferation, and secretion of effector molecules. Cytokines may act on the cells that secrete them (autocrine action), on nearby cells (paracrine action), or in some instances on distance cells (endocrine action). It is common for different cells types to secrete the same cytokine or for a single cytokine to act on several different cell types (pleiotropy). Cytokines are redundant in their activity, and are often produced in a cascade, as one cytokine stimulates its target cells to make additional cytokines. Cytokines can also act synergistically or antagonistically.

[0210] Elevated levels of certain cytokines have been measured in human discs, and have been associated

with degeneration and pain. Among the major cytokines found are, for example and without limitation: interleukin-1, -6, and -8, tissue necrosis factor- α (TNF- α), and prostaglandin E₂ (PGE₂) [Miyamoto, 2000; Ahn, 2002; Olmarker, 1998; Weiler, 2005]. The source of cytokines can be circulating inflammatory cells in the case of herniated discs [Kawaguchi, 2002; Woertgen, 2000], or disc cells in the case of contained disc degeneration [Burke, 2002].

[0211] For disc cells, inflammatory factor production may be stimulated for example as part of several signaling cascades (described below), by fragments of degraded extracellular matrix, or matrix deformation (FIG. 1). These exemplary pro-inflammatory stimuli can trigger cells to initiate a number of catabolic programs meant to stimulate tissue repair and remodeling that includes production of matrix metalloproteinases 1, 9 and 13 [Anderson, 2002]. During this wound healing process, cytokines are also involved in stimulating angiogenesis and granulation tissue formation [Gillitzer, 2001].

IL-1 and TNF- α

[0212] IL-1b and TNF- α have been observed to demonstrate overlapping pro-inflammatory effects, activate common signaling cascades, and induce similar target genes (see ref in Faur). Effector cascades mediating inflammatory responses to IL-1 and TNF- α include the mitogen-activated protein kinases (MAPK), NF- κ B, and prostaglandin signal transduction pathways (shalombarak). The signaling molecule nitric oxide may also form important component of the inflammatory cascade.

[0213] Imaging via labeling tissue necrosis factor- α (TNF- α) provides one particular beneficial example of marking for imaging a pro-inflammatory cytokine that can chemically hypersensitize the intervertebral disc and spinal nerve roots, thereby contributing to low back pain. Studies have been conducted that utilize immunohistochemistry to localize TNF- α in histologic sections of normal and degenerated mouse-tail discs. These studies suggest that the levels of TNF- α are increased after compression-induced degeneration of the intervertebral disc (FIGS. 2A-D).

[0214] To demonstrate a TNF- α based localization modality of the present invention, compositions and methods have been developed that label TNF- α antibodies with I-125 so that variations in TNF- α content can be imaged *in vivo*. An experiment was conducted to observe and confirm the beneficial use of this approach as follows. Mice such as mouse 30 shown in FIG. 3 were subjected to conditions that initiate tail-disc degeneration (FIG. 3), and were then injected intravenously with I-125 labeled TNF- α antibody. These animals were then imaged with a phosphor imaging plate, such as plate 50 shown in FIG. 4. Use of this composition and imaging methods demonstrated readily observed increased uptake in the regions of the injured discs, such as seen in image 60 in FIG. 5 wherein four injured tails are shown in 2-group sets on either side of a centrally located control tail in the image

that was not injured though received similar labeled marker injection.

[0215] This particular experiment was performed using a particular radio-labeled TNF- α blocker, more specifically infliximab (Trade name "Remicade™" commercially available from Johnson & Johnson), and demonstrates one exemplary embodiment adapted for beneficial use according to the present invention. While this particular modality is considered highly beneficial in the specific mode described, it is also exemplary of a number of broad aspects of the present invention that may be illustrated by many alternative or combinatorial approaches that are herein contemplated.

[0216] In one regard, the present illustrative embodiment provides an example of using a therapeutic compound that actually provides some pain-related therapy (e.g. TNF- α antibody or other form of blocker) that is also used to image the location of the pain being treated (as the labeled marker, as conducted in the illustrative experiment, or targeted factor itself). This step may be followed by additionally treating the imaged region thereafter with additional spacially localized or directed therapies. Examples include, without limitation, directed energy therapies such as those elsewhere herein described, or further localized injection of similar or other therapeutic compound(s).

[0217] In another more specific regard, TNF- α blockers or antibodies are contemplated as a class of therapeutic compounds beneficially adapted for use according to the invention, within which infliximab or Remicade™ (or analogs or derivatives thereof) is used in a particular beneficial embodiment as just described. These provide the benefit of selective uptake at nerve endings where pain may be occurring, and thus a particular beneficial target agent for labeling to image pain. They also provide the benefit of some therapeutic value to the pain itself.

[0218] Furthermore, it is to be appreciated that targeted agents, such as antibodies as herein described by way of example, may provide the label for imaging, or may take the form of the targeted factor (either by itself or by virtue of its conjugation or binding with a first resident factor). In the later case, delivery of the first factor is then subjected to subsequent labeling by delivery of a second agent as the labeled marker (again either by its imagability itself or as bound, associated, or conjugated with the first delivered agent to the region imaged).

MAPK Pathway

[0219] MAPKs form an intracellular signaling pathway built upon a self-propagating phosphorylation system (FIG. 6). Activation of MAPKs are one of the pivotal intracellular pathways triggered by cytokine receptors (Shalom-berak). Three MAPK subgroups have been identified: extracellular signal regulated kinase (ERK); the Jun NH₂-terminal kinases (JNK); and p38 (geng, others). In chondrocytes, ERK activation occurs in response to diverse stimuli, while JNK and p38 is only seen in re-

sponse to IL-1 and TNF- α (Firestein, liancini): this signaling pathway is thought responsible for cartilage degradation (geng). JNK and p38 are collectively termed stress activated protein kinases (SAPKs). The signal is initiated by membrane-proximal small GTPases of the Rho family, activation of MLK, and phosphorylation and activation of MKK3/6 that in turn phosphorylates and activates p38. Faur).

[0220] One important endpoint of MAPK activation is the production of the phosphorylated active activator protein 1 (AP-1) transcription factor (heterodimer of c-Jun and c-Fos), which in turn, can influence chondrocyte collagenase activity (mengshol, Ferreria refs). AP-1 plays a central role in the transcriptional regulation of many MMP genes including collagenase and stromelysin (mengshol refs, Firestein). Similarly, MIF activates the MAPK pathway and AP-1 leading to cell proliferation, and PGE₂ production, which eventually promotes monocyte/macrophage activation. Certain published data suggests that MIF is in particular upregulated under conditions of chronic emotional stress and can potentiate elevated levels of other inflammatory factors such as for example those examples herein described. Accordingly, labeling MIF provides yet a further embodiment of the various present aspects.

[0221] JNK and p38 are essential for IL-1 induction of mmp-13, while ERK pathway is not. p38 is essential for multiple inflammatory genes, including IL-1, TNF- α , IL-6, stromelysin-1 (mmp-3) and mmp-1 (mengeshol).

[0222] It is to be appreciated that various such materials associated with pathways or molecular cascades associated with pain may provide the target for labeled markers and subsequent imaging as herein described, and various such materials are provided here as beneficial examples which, though of particular value, are also not intended to limit broad aspects contemplated hereunder. In addition, such otherwise indigenous materials may also demonstrate selective uptake in tissues associated with pain. In such case, these otherwise indigenous materials (or synthetic or other biologic constructs similar to them, such as analogs or derivatives thereof) may also be harnessed and labeled for delivery as the labeled marker. Moreover, due to their selective uptake, particular accumulated concentrations of certain molecules in areas of pain also render them viable targets as the pain factors themselves for labeling with labeled markers that bind to them.

NF- κ β Pathway

[0223] In addition to the MAPK induction, IL-1 and TNF- α activate NF- κ β . NF- κ β is a transcription factor that exists in a latent form in the cytoplasm of unstimulated cells and is composed of a transcriptionally active dimer (p65 and p50) bound to an inhibitor protein (I κ β) (Bowie, Magnani). NF- κ β is activated by a large number of different signals that include similar cell stress signals that activate SAPKs. IL-1 and TNF- α trigger the phosphor-

ylation and degradation of I κ B, resulting in the release of NF- κ B to enter the nucleus (refs in Shalom; Baeuerle). NF- κ B activation occurs through a cascade starting with NF- κ B-inducing kinase (NIK), which then phosphorylates and activates the inhibitor of NF- κ B (I κ B) kinases. Phosphorylation of I κ B results in ubiquitination and degradation of I κ B inhibitory subunit, allowing NF- κ B to translocate to the nucleus where it acts as a transcription factor and regulates its target genes, which include collagenase (MMP-1; Barchowsky) (Mengshol, magnani) and COX-2 (Mifflin). FIG. 7 shows certain further details of this cascade and relationship between components.

Prostaglandin Pathway

[0224] Eicosanoids are signaling molecules that act in an autocrine fashion. Pro-inflammatory stimuli can lead to increased phospholipid-derived eicosanoid synthesis that involves a cascade of three enzyme reactions (FIG. 8). First, arachidonic acid (AA) is liberated from its phospholipid storage sites by phospholipase A2 (PLA2). The next rate-limiting step is conversion of AA to prostaglandin H2 by cyclooxygenase (COX).

[0225] The prostaglandin pathway is stimulated by IL-1b. This cytokine increases the activity of PLA2 and induces COX-2 gene expression by binding to a specific cell-surface receptor (IL-1RI) that ultimately leads to increases in COX-2 promoter activity via the NF- κ B pathway (Faur refs, geng). In chondrocytes, COX activity is not increased by TNF- α . Rather, TNF- α can amplify COX activity in IL-1 stimulated cells. (Berenbaum).

[0226] Prostaglandin E₂ (PGE₂) stimulates the catabolism of chondrocytes, having both anti-proliferative and pro-apoptotic effects (berenbaum ref, also goldring ref in liancici). An increase in PGE₂ may therefore tip the balance toward catabolism.

Nitric Oxide

[0227] Nitric oxide (NO) is a small signaling molecule that is part of the catabolic program in chondrocytes induced by IL-1 and TNF- α (Lotz; Goldring). It is produced within the cell by the inducible isoform of NO synthase (iNOS), and then passes readily through the cell membrane to affect neighboring cells. Because it has a short half-life (5 to 10 seconds) it acts only locally, yet it plays an important role in the pathophysiology of arthritic disease (Ferreira Mendes). It has been shown to: induce apoptosis (by stimulating release of cytochrome c from mitochondria) and inflammation (by activating COX and PLA2 (Vassalle, clancey)); suppress collagen and proteoglycan synthesis; and upregulate MMP synthesis (Scheurwegh).

[0228] IL-1 and TNF- α increase the gene expression and synthesis of iNOS, through the transcription factors NF- κ B and AP-1. Activation of NF- κ B is an essential step for iNOS induction (see Mendes refs). Also, there is some evidence that the MAPK p38 may be involved in the ac-

tivation of NF- κ B and subsequent iNOS expression, since p38 is reported to be required for IL-1-induced iNOS expression in chondrocytes (Mendes).

5 Labeling/Imaging Cellular Factors Associated with Inflammation

[0229] Cells that produce or are associated with inflammatory factors can also be labeled with targeted markers and thereafter imaged as an indicator that pain exists in the area. For example, disc cells that are actively synthesizing inflammatory factors may be labeled as such (or components thereof may be labeled). Inflammatory cells that are attracted to painful discs, such as for example leukocytes, may be labeled and imaged for this purpose.

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[0231] It is to be appreciated based upon the foregoing disclosure that pain factors are labeled and imaged in order to identify, with a useful degree of geographic specificity, active pain sites in and around skeletal joints. Such is considered highly beneficial in particular for use in diagnosing the cause of pain, and understanding where and how to treat for pain relief, such as for example with local ablation or energy delivery systems, and/or local drug delivery.

[0232] Various terms have been used herein of a certain technical nature, and should be given their standard technical meaning in the context of the particular art to which this disclosure pertains, and in the context of their use in this description together with other accompanying disclosure, unless otherwise given a specific meaning hereunder. Notwithstanding the foregoing, it is understood that certain specific materials or types of materials are identified, whereas other similar materials or types of materials are also intended to be implicated within the broad scope intended for the current invention. For example, "pain factors" are herein identified as playing a role in various of the present embodiments. Such terms are intended to mean any and all materials, whether structural, chemical, or otherwise, that have an association, either directly or indirectly, with pain such that binding them provides a vehicle to enhance diagnosis or therapy in relation to the associated pain. In one particular example, factors related to transmitting pain signals along or between nerves are to be included. Or, factors that stimulate pain, such as "inflammatory" materials, are indicated. Materials related to other points in a chemical or biological cascade related to pain are also implicated, such as factors that relate to secondary or tertiary products or components of such pain generation or transmission process. If a factor is distinctly present (or absent) in a somewhat recognizable manner when and where pain is present, and in a different level or manner than when and where pain is not present, then it is considered a "pain factor" as herein described. This use of the term "factor" similarly applies in other contexts herein provid-

ed, such as for example "inflammatory factors", "cellular factor(s)", "nerve factors", etc.

[0233] In another regard, it is also contemplated that, where certain specific examples of chemicals or materials are herein provided, other related compounds may be interposed in addition or in the alternative to such specified compound. For example, agents related to a certain material may be suitable substitutes and may include for example precursor materials, such as a material that may be metabolized or otherwise altered to produce the specified "factor" or "label" or other compound or material referenced. Analogs or derivatives of the specified material may also be suitable in similar uses or preparations or systems. This includes for example modified molecular forms of a specified material that retain the related binding or other activity of the specified material so as to perform as herein described as a labeled pain factor or targeted label.

[0234] Moreover, use of a "marker" or "label" to tag or label a "factor" is generally herein described in fairly simple terms for the purpose of providing a general overall understanding of the broad aspects contemplated hereunder. However, the actual steps and/or materials used in order to achieve such "labeled pain factor" result may be more extensive than herein described, though may be carried out by one of ordinary skill in the art based upon review of this disclosure in its entirety in combination with other available related information and thus further contemplated hereunder. For example, intermediary tagging, labeling, or binding may be beneficially used in order to achieve the labeled marking necessary to provide differential imaging of the labeled result in a useful manner.

[0235] In one further exemplary embodiment, bi-specific antibodies may be used in such a manner as follows. One binding site of the bi-specific antibody provides a particular binding affinity for the pain factor being targeted, and thus differentially binds to that factor. However, this is done in a manner leaving a second binding site exposed, and which second binding site has binding specificity to a second material as a label agent. This second material thereafter binds to the second binding site of the bi-specific antibody bound at the first site to the pain marker. The result provides a labeled marker on the pain factor via the second material, which is tagged to the pain factor via use of this intermediary bi-specific antibody.

[0236] It is also to be understood that the labeled marking of pain factors herein described is of particular benefit with respect to thereafter image the result. While imaging the "labeled pain factor" may be generally described, it is to be understood that what is imaged by the particular imaging modality may include without limitation: the overall conjugate or combination of label-plus-factor; the label itself; the factor itself (e.g. to the extent modified in a recognizable way by the labeled marking); or combinations of the above, including in further modes use of intermediary binding materials such as for example bi-specific antibodies as herein described.

[0237] One particular example of a labeled marker and pain factor combination believed to be useful according to certain of the embodiments herein described is provided in finer detail to provide a further understanding. This relates to radiolabeled TNF- α antibodies and related imaging tools herein described. However, it is to be appreciated that this approach, though in particular highly beneficial, is exemplary of broader aspects of the present invention and other labeling and/or marker modalities, or targeted vehicles such as without limitation antibodies, and/or imaging tools are contemplated and may be used without departing from the intended broad scope according to various aspects of the invention.

[0238] The invention according to further aspects provides a unique ability to direct therapy to pain, including without limitation pain associated with musculoskeletal joints and in particular the spine. Accordingly, the systems and methods of the invention according to further embodiments also include therapeutic device assemblies for delivering such therapy. Such may include local drug or other chemical delivery modalities. Or, therapeutic dosing of energy may be delivered, such as for example radiofrequency (RF) energy delivery probes, ultrasound probes, high intensity focused ultrasound (HIFU), light energy (e.g. lasers for example), microwave energy, or cryovascular therapeutic tools may be used. By identifying where treatment is required due to the selectively visualized pain factors there, these tools may be used in a more efficient manner. Accordingly, the compositions of labeled markers, the visualization or imaging tools, and the therapeutic tools are thus used in an overall symphony that together provides beneficial healthcare results in treating pain.

[0239] This is in particular the case with respect to back pain. For example, a disc may be identified as a source of pain, whereas lack of further clarity may render it difficult to treat the pain in a selective way. Often, ablation of the entire disc is not desired. According to certain further embodiments, the labeled marking of pain factors and related imaging is used to identify more specifically where pain occurs. In one mode, at least one-half of the disc is identified as the target for therapy. In another mode, the labeled marker visualization localizes the target for therapy to one or more quarter quadrants of the disc. In still further embodiments, directionally localized energy delivery, e.g. laser, ultrasound, or microwave, may be particularly beneficial for isolating the therapy to the isolated region of visualized, labeled pain factors. Furthermore, local injections of pain medication may be directed via such targeted labeling and related imaging of pain localization.

[0240] In another highly beneficial aspect, pain factors that are visualized with targeted markers as described hereunder may relate to nerves that are located at least in part within bones. This may be the case for example with respect to bony end-plates that are innervated with nociceptive nerve fibers. In one particular beneficial embodiment, pain factor imaging as herein described is

used to locally identify one or more particular end-plates of vertebral bodies as the pain source. Accordingly in many such instances, a basivertebral ablation tool set and method may be used to ablate the basivertebral nerve that innervates that end-plate. This may be done for example using a mono- or bi-polar electrode assembly that is delivered via one or more needle or drill probes into the vertebral body that is used to RF ablate the nerve closer to a root trunk section within the bone. Despite this particular beneficial combination of tools and methods for treating pain in a uniquely localized manner, however, it is to be appreciated that other localized pain sources may be selectively visualized using a variety of useful targeted markers, and a variety of tools or methods may be used to direct therapy accordingly, without departing from the present intended scope of the present invention.

[0241] The following US Patents are referred to herein : 5,391,197 to Burdette et al.; 6,074,352 to Hynynen et al.; 6,126,682 to Sharkey et al.; 6,231,528 to Kaufman et al.; 6,368,292 to Ogden et al.; 6,470,220 to Kraus, Jr. et al.; 6,562,033 to Shah et al.; 6,575,969 to Rittman III et al.; 6,699,242 to Heggeness; 6,736,835 to Pellegrino et al.; 6,827,716 to Ryan et al.; 6,907,884 to Pellegrino et al. The following published PCT Patent Applications are referred to herein: WO 2003/059437 to Diederich et al.; and WO 03/061756 to Diederich et al. The following Published US Patent Applications are also referenced herein: US 2004/0064137 to Pellegrino et al.; and US 2004/0064136 to Papineau et al.

[0242] Various different modes of "imaging" and related tools are herein contemplated, as apparent to one of ordinary skill to match the targeted marker modalities employed to accomplish the general objectives hereunder. In one regard, a variety of diagnostic tools may be used to acquire information related to the targeted pain factor(s) and related spacial location relative to surrounding tissues. This information may be processed and converted into a representation that may be displayed or otherwise conveyed to a healthcare provider in a manner sufficient and useful to understand the spacial location of the associated pain. Accordingly, various different types of sensors, data acquisition systems, processors, and displays may be used in various combinations to convert the labeled marking to useful information to such healthcare providers. Many of these are commercially available in sufficient form to readily integrate with the targeted marker agents and delivery systems herein described (which may further include therapeutic aspects) in an overall system sufficient to provide useful information in medical patient management.

[0243] Although the description above contains many details, these should not be construed as limiting the scope of the invention but as merely providing illustrations of some of the presently preferred embodiments of this invention. Moreover, it is not necessary for a device or method to address each and every problem sought to be solved by the present invention, for it to be encompassed by the present claims.

Claims

- 1. A method for preparing a system comprising a targeted agent for performing a medical procedure on a patient having been diagnosed with back pain, **characterized by:**

based upon the back pain diagnosis, preparing a volume of a targeted agent in combination with a delivery assembly in a configuration that is operable to allow delivery of the targeted agent to a region of the patient that comprises at least a portion of a spine, wherein the targeted agent is **characterized by** differentially binding to a pain factor at a location of a spinal joint associated with a source of the back pain when delivered into the patient;

configuring at least one of an energy delivery system to deliver energy to the patient and an imaging system to image the pain factor at a location of a spinal joint; and

wherein the targeted agent when delivered into the region of the patient that comprises at least a portion of a spine is further characterized to enhance at least one of (i) diagnostic localization of the source of the back pain at the location in the patient's body when the imaging system is operated in the configuration to image the location of a spinal joint and (ii) selective tissue therapy of the source of the back pain at the location when the energy delivery system is operated in a configuration to deliver energy to the location of a spinal joint; and

wherein the targeted agent comprises at least one of: a nerve factor, a blood vessel factor, an inflammatory cytokine, or a combination thereof; or a factor associated with pH or pO₂ in tissue, optionally associated with a relatively low pH or pO₂, respectively, or a binding agent or antibody respectively thereof.

- 2. A system, which comprises a targeted agent for performing a medical procedure on a patient, said targeted agent prepared for dosed delivery into a patient diagnosed with back pain, **characterized by:**

a delivery assembly for the targeted agent that is configured and operable to allow delivery of the targeted agent into a location of a spinal joint in the patient; and

said targeted agent when delivered to a region of the patient that comprises at least a portion of a spine via the delivery assembly is characterized to differentially bind to a pain factor at a location of a joint associated with the back pain in a manner further characterized to enhance at least one of (i) diagnostic localization of a source of the back pain at the location of a spinal joint

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and (ii) selective tissue therapy to a source of the back pain at the location containing the bound pain factor in response to a delivered energy to the region of the patient that comprises at least a portion of a spine; and

wherein the pain factor comprises at least one of: a nerve factor, a blood vessel factor, an inflammatory cytokine, or a combination thereof; or a factor associated with pH or pO₂ in tissue, optionally associated with a relatively low pH or pO₂, respectively, or a binding agent or antibody respectively thereof.

- 3. The system of claim 2, configured for diagnosing or treating a source of back pain at a location of a spinal joint within a body of a patient, wherein:

the targeted agent is labeled so as to comprise a targeted label that is characterized to bind to and label a pain factor associated with a source of back pain at the location when delivered into the patient;

the delivery assembly for the targeted agent is in a configuration that is operable to allow delivery of the targeted label into the patient; and

the system comprises at least one of (a) an imaging system in a configuration that is operable to image at least one of the targeted label or the labeled pain factor in a manner sufficient to selectively differentiate a first concentration of the labeled pain factor at the location versus a second concentration of the labeled pain factor in a second tissue location adjacent to the location, and (b) a therapeutic device assembly in a configuration that is operable to provide therapy in a substantially localized manner that is substantially isolated to the location comprising the labeled pain factor.

- 4. The method of claim 1 or the system of claim 2 or 3, further **characterized by:**

a volume of the targeted agent that is labeled such that the targeted agent comprises a targeted label that is configured for delivery into the patient and that is characterized to differentially bind to and label the pain factor associated with the source of back pain at the location; and

the targeted label being configured for artificially labeling the pain factor at the location by binding the pain factor with the targeted label when delivered into the region of the patient that comprises at least a portion of a spine.

- 5. The method of claim 1 or the system of claim 2 or 3, wherein the pain factor comprises:

a blood vessel factor that comprises at least one of a blood vessel or microvessel, or a substance or

- structure associated with the blood vessel or microvessel; or substance P, CGRP, trkA, nerve growth factor (NGF), a NGF antagonist, PGP 9.5, SYN, peripherin, Neurofilament 200kD (NF200), TNF- α , infliximab, PECAM, CD34, GFAP, PGE-2, or an analog or derivative or binding agent or antibody thereof; or a nerve binding agent or antibody; or an interleukin optionally comprising IL-1, IL-6, or IL-8, or an analog or derivative thereof; or MIF or a binding agent or antibody thereof; or a CGRP receptor.
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6. The method of claim 1, the system of claim 2 or 3, or the method or system of claim 4, wherein the pain factor or targeted agent comprises:
a nerve factor that comprises at least one substance associated with at least one of a nerve fiber or a cellular structure associated with the nerve fiber.
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7. The method of claim 1 or the system of claim 2 or 3, wherein the pain factor or targeted agent comprises a TNF- α blocker.
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8. The method of claim 1 or the system of claim 2 or 3, wherein the pain factor comprises: an interleukin optionally comprising IL-1, IL-6, or IL-8, or an analog or derivative thereof; or a factor associated with pH or pO₂ in tissue, optionally associated with a relatively low pH or pO₂, respectively, or a binding agent or antibody respectively thereof.
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9. The method of claim 1 or the system of claim 2 or 3, wherein the targeted agent comprises:
at least one of a nerve factor, a blood vessel factor, an inflammatory cytokine, or a binding agent or an antibody thereof; or a binding agent or an antibody of the nerve factor defined in claim 5; or a blood vessel factor that comprises a substance associated with a structure of a blood vessel or a microvessel, or a binding agent or an antibody thereof; or an inflammatory cytokine binding agent or antibody; or a Substance P binding agent or antibody; or a CGRP or CGRP receptor binding agent or antibody; or a trkA binding agent or antibody; or an anti-trkA antibody binding agent or antibody thereof; or a nerve growth factor (NGF) or an analog or derivative thereof; or a NGF antagonist or an analog or derivative thereof; or a nerve binding agent or antibody thereof or an analog or derivative thereof; or a NGF binding agent; or an anti-NGF antibody; or a NGF antagonist binding agent; or an anti-NGF antagonist antibody; or a PECAM binding agent or antibody; or a CD34 binding agent or antibody; or a GFAP binding agent or antibody; or an interleukin binding agent or antibody; or an interleukin binding agent or antibody that comprises an IL-1, IL-6, or IL-8 binding agent or antibody; or a PGE-2 binding agent or antibody; or a factor associated with pH or pO₂ in tissue or a binding agent or antibody thereof; or PGP 9.5, or
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- SYN, or peripherin, or Neurofilament 200kD (NF200), or TNF- α , or infliximab, or an analog or derivative thereof, or a binding agent or an antibody thereof; or a radioactive material, optionally a radiolabeled TNF- α , antibody or an analog or derivative thereof, or radiolabeled iodine, optionally I-125; or an MIF binding agent or antibody; or a nanoparticles; or at least one of gold or iron oxide; or an MRI contrast agent, optionally gadolinium; or an ultrasound contrast agent; or a radiographic contrast agent.
10. The system of claim 3 or the method or system of claim 4, wherein the labeled pain factor when labeled with the targeted agent is indicative of a relatively low pH or pO₂ at the location.
11. The system of claim 3 or the method or system of claim 4, comprising:
an imaging system configured for imaging the labeled pain factor using an imaging tool that comprises a phosphor imaging plate; or wherein the targeted agent comprises an MRI contrast agent, and comprising an MRI imaging system configured for imaging an area of increased concentration of the MRI contrast agent bound to the pain factor; or wherein the targeted agent comprises an ultrasound contrast agent, and comprising an ultrasound imaging system configured for ultrasonically imaging an area of increased concentration of the ultrasound contrast agent bound to the pain factor; or wherein the targeted agent comprises a radiographic contrast agent, and comprising an X-ray system configured for imaging an area of increased concentration of the radiographic contrast agent bound to the pain factor using X-ray.
12. The method of claim 1, the system of claim 2 or 3, or the method or system of claim 4, further comprising:
a first binding agent prepared for delivery into the body, wherein the first binding agent is adapted to bind to a first pain factor;
wherein the targeted agent is prepared for delivery into the patient's body after the first binding agent is delivered into the body and bound to the first pain factor; and
wherein the targeted agent when delivered to the region of the patient that comprises at least a portion of a spine characterized to bind to a site located on the bound combination of the first binding agent and the first pain factor; and
wherein the first binding agent optionally comprises a bi-specific antibody with a first binding site is characterized to bind to the first pain factor and a second binding site characterized to bind to the targeted agent.

13. The method of claim 1, the system of claim 2 or 3, or the method or system of claim 4, comprising:

an imaging system configured in a configuration that is operable to image a region with diagnostic localization of the source of the back pain at the location via the targeted agent selectively bound to the pain factor at the location; and a therapeutic system configured in a configuration that is operable for conducting a therapeutic procedure in a substantially localized manner to the location where the targeted agent bound to the pain factor is locally imaged.

14. The system or method of claim 13, wherein the therapeutic system is configured in a configuration that is operable to substantially alleviate generation or transmission of pain at the location, such as by being configured in an operating mode to substantially ablate at least one nerve at the location and/or to deliver at least one therapeutic chemical to the location.

15. The system or method of claim 13, wherein the therapeutic system comprises: an energy delivery system configured in a configuration that is operable for delivering a therapeutic dose of energy in a substantially localized manner to the location, optionally configured for ablating at least one nerve at the location with the energy.

16. The method or system of claim 15, wherein: the therapeutic system is configured for delivering ultrasound energy to the location in a directed manner from a second location; the second location is outside of the patient; and the ultrasound energy is configured to be delivered via high intensity focused ultrasound (HIFU) that is adapted to focus the ultrasound energy to the location.

17. The method or system of claim 16, wherein: the second location is adjacent to the location within the patient; and the system is configured to deliver ultrasound energy via a directional ultrasound probe.

18. The method or system of claim 15, wherein therapeutic system is configured in a configuration to deliver the therapeutic dose of energy that comprises thermal, electrical, microwave, or light energy.

19. The method or system of claim 18, comprising a radiofrequency (RF) probe coupled to an electrical energy source to the RF probe, such that the system is configured in a configuration that is operable for delivering the therapeutic dose of electrical energy

via the radiofrequency (RF) probe.

20. The method of claim 1 or system of claim 2 or 3, wherein the location of the spinal joint comprises:

- (a) at least a portion of an intervertebral disc; or (b) a region of tissue located within only a portion that is equal to less than an entire circumference of an intervertebral disc, optionally a region of tissue located within less than or equal to one-half or one-quarter of the circumference of the intervertebral disc; or (c) an end-plate associated with a vertebral body.

21. The method of claim 1 or system of claim 2 or 3, wherein: the targeted agent is prepared for delivery in a localized manner to the location, optionally for injection into a region of tissue associated with the location using a local injection assembly.

22. The method of claim 1 or system of claim 2 or 3, wherein: the targeted agent is prepared for delivery systemically into the patient, optionally for delivery into the patient's systemic blood circulation, gastrointestinal system, or respiratory system.

23. The method of claim 1, the system of claim 2 or 3, or the method or system of claim 4, wherein:

the targeted agent is prepared for delivery into the patient that is characterized to allow for artificially labeling the pain factor at multiple said locations by binding the pain factor with the targeted agent when delivered into the patient; and an imaging tool is provided that is configured in a configuration that is operable to image at least one of the targeted agent or the labeled pain factor and in a manner sufficient to differentiate a first concentration of the labeled pain factor at one of the multiple said locations versus a second concentration of the labeled pain factor at another of said multiple locations.

24. The system or method of claim 23, comprising: at least one therapeutic system configured in a configuration that is operable to perform a therapeutic procedure in a substantially localized manner to each of the locations where the targeted labeled pain factor is locally and selectively imaged.

25. The method of claim 1, the system of claim 2 or 3, or the method or system of claim 4, comprising: an energy delivery system configured and operable for delivering energy into the patient in a manner that differentially treats one or more regions associated

with differentially bound pain factor when the targeted agent is delivered into the patient.

- 26. The method of claim 1, the system of claim 2 or 3, or the method or system of claim 4, comprising: 5

an energy delivery system;
 wherein the energy delivery system is configured to deliver energy into the patient; and
 wherein the volume of targeted agent is configured and operable for delivery into a patient and to differentially bind to a pain factor associated with the pain in a manner characterized to allow tissue regions containing a first concentration of the differentially bound pain factor to exhibit a differential and selective therapeutic response to the energy when delivered versus other regions with lower concentrations of the differentially bound pain factor. 10 15 20

- 27. The method of claim 1, the system of claim 2 or 3, or the system or method of claim 26 **characterized by:**

a therapeutic device assembly is provided that comprises an energy delivery assembly that is configured and operable to deliver a therapeutic dose of energy in a substantially localized manner that is substantially isolated to the location associated with the spinal joint. 25

- 28. The system or method of claim 27, wherein: the energy delivery assembly is configured and operable to be delivered into the patient to a position at or adjacent to the location. 30

- 29. The system or method of claim 27, further comprising: an introducer that is configured and operable to deliver the energy delivery assembly to the location. 35

- 30. The system or method of claim 29:

wherein the introducer comprises a needle assembly; or
 wherein the introducer comprises a needle assembly configured and operable to be advanced through bone and to deliver the therapeutic device assembly to a position within the bone; or
 wherein the introducer comprises a needle assembly configured and operable to be advanced through bone and to deliver the therapeutic device assembly to a position within the bone, and wherein the therapeutic device assembly is configured and operable to ablate an intraosseous nerve within the bone and that is associated with pain related to the labeled pain factor visualized at the location; or
 wherein the introducer comprises a needle as- 40 45 50 55

sembly configured and operable to be advanced through bone of a vertebral body and to deliver the therapeutic device assembly to a position within the vertebral body associated with a basivertebral nerve, and wherein the therapeutic device assembly is configured and operable to ablate the basivertebral nerve from the position; or

wherein the introducer comprises a needle assembly configured and operable to be advanced through bone and to deliver the therapeutic device assembly to a position within the bone, wherein the therapeutic device assembly comprises a radiofrequency (RF) current ablation assembly, wherein the RF current ablation assembly comprises a first electrode and a second electrode adapted to be positioned at first and second positions adapted to straddle at least a portion of the basivertebral nerve, and wherein the RF current ablation assembly is configured and operable to deliver the RF current between the first and second electrodes sufficient to ablate nerve tissue between the first and second positions, and further wherein the RF current ablation assembly optionally comprises a delivery probe with an elongated body that carries the first and second electrodes in a bipolar lead assembly arrangement. 30

- 31. The method of claim 1 or system of claim 2 or 3, wherein:

the targeted agent comprises a material that has a preferential binding affinity to the pain factor located within the region at the location sufficient to preferentially bind to the pain factor versus other structures within the region, and such that the material accumulates at a higher concentration within a first portion of the region having a higher amount of the pain factor than other portions within the region; and
 including an energy delivery system configured and operable for delivering energy to the region in a manner that selectively treats the location of the first portion versus the other portions, after delivering the material to the region, and such that pain is reducible in the region. 35 40

- 32. The method of claim 1 or system of claim 2 or 3, or method or system of claim 31, wherein the targeted agent comprises a material that comprises: a metal, optionally gold; or a nanoparticle; or a gold nanoparticle; or an antibody; or an antibody and a metal, optionally a metal nanoparticle, optionally a gold nanoparticle, associated with the antibody. 50 55

- 33. The method of claim 1 or system of claim 2 or 3, wherein:

the targeted agent is configured in a configuration that is operable in a method for delivery into at least a first region that comprises the location and at least one other region that is different than the first region;

the first region comprises at least one spinal joint level along the spine, optionally a single spinal joint, or an area of a spinal joint, optionally at least a part of an intervertebral disc, or a vertebral body, or a vertebral body end-plate, or a facet joint or transverse process; and said at least one other region comprises at least one other spinal joint level along the spine.

34. The method or system of claim 31, comprising: an imaging system configured and operable for imaging the region in a manner that sufficiently differentiates spatial relationships between different concentrations of the material so as to substantially identify the location of the first portion relative to the other portions within the region.

35. The method or system of claim 31, wherein: the energy delivery system is configured and operable to deliver the energy principally to the first portion in a substantially localized manner sufficient to differentially treat the first portion with the energy versus the other portions.

36. The method of claim 1, wherein: the targeted agent is prepared for delivery into the patient after the patient is diagnosed in a manner that identifies the region as a painful region of the patient's body.

37. The method of claim 1 or the method or system of claim 15, wherein: the location is not diagnosed to include cancer cells prior to conducting the medical procedure.

38. The method or system of claim 17, wherein:

the second location is adjacent to an intervertebral disc; and

the directional ultrasound probe is configured and operable to deliver the directional ultrasound therapy into the location within the intervertebral disc.

Patentansprüche

1. Verfahren zum Herstellen eines Systems, das ein abgezieltes Mittel umfasst, zur Verwendung bei einem medizinischen Eingriff an einem Patienten mit diagnostiziertem Rückenschmerz, **gekennzeichnet durch:**

Herstellen eines Volumens eines abgezielten Mittels in Kombination mit einer Abgabebaugruppe in einer Konfiguration, die so betreibbar ist, dass sie die Abgabe des abgezielten Mittels an eine Region des Patienten ermöglicht, die zumindest einen Abschnitt einer Wirbelsäule umfasst, auf Basis der Rückenschmerzdiagnose, wobei das abgezielte Mittel **dadurch** gekennzeichnet ist, dass es differentiell an einen Schmerzfaktor an einer Stelle eines Wirbelsäulengelenks bindet, die mit einer Quelle des Rückenschmerzes assoziiert ist, wenn es in den Patienten abgegeben wird;

Konfigurieren zumindest eines von einem Energieabgabesystem, um Energie an den Patienten abzugeben, und eines Bildgebungssystems, um den Schmerzfaktor an einer Stelle eines Wirbelsäulengelenks abzubilden; und

wobei das abgezielte Mittel, wenn es in die Region des Patienten abgegeben wird, die zumindest einen Abschnitt einer Wirbelsäule umfasst, ferner so gekennzeichnet ist, dass es zumindest eines von (i) einer diagnostischen Lokalisierung der Quelle des Rückenschmerzes an der Stelle in dem Körper des Patienten, wenn das Bildgebungssystem in der Konfiguration zum Abbilden der Stelle eines Wirbelsäulengelenks betrieben wird, und (ii) einer selektiven Gewebetherapie der Quelle des Rückenschmerzes an der Stelle, wenn das Energieabgabesystem in einer Konfiguration zum Abgeben von Energie an die Stelle eines Wirbelsäulengelenks betrieben wird, verbessert; und

wobei das abgezielte Mittel zumindest eines umfasst von: einem Nervenfaktor, einem Blutgefäßfaktor, einem entzündungsfördernden Zytokin oder einer Kombination davon; oder einem Faktor, der mit pH-Wert oder pO₂-Wert in Gewebe assoziiert ist, der optional mit einem relativ geringen pH-Wert bzw. pO₂-Wert assoziiert ist, oder einem Bindungsmittel oder jeweiligen Antikörper davon.

2. System, das ein abgezieltes Mittel umfasst, zur Durchführung eines medizinischen Eingriffs an einem Patienten, wobei das abgezielte Mittel für eine dosierte Abgabe in einen Patienten mit diagnostiziertem Rückenschmerz hergestellt ist, **gekennzeichnet durch:**

eine Abgabebaugruppe für das abgezielte Mittel, die so konfiguriert und betreibbar ist, dass sie die Abgabe des abgezielten Mittels in eine Stelle eines Wirbelsäulengelenks in dem Patienten ermöglicht; und

wobei das abgezielte Mittel, wenn es über die Abgabebaugruppe in eine Region des Patienten abgegeben wird, die zumindest einen Abschnitt

- einer Wirbelsäule umfasst, so gekennzeichnet ist, dass es differentiell an einen Schmerzfaktor an einer Stelle eines Gelenks bindet, die mit dem Rückenschmerz assoziiert ist, auf eine Weise, die ferner so gekennzeichnet ist, dass sie zumindest eines von (i) einer diagnostischen Lokalisierung der Quelle des Rückenschmerzes an der Stelle eines Wirbelsäulengelenks und (ii) einer selektiven Gewebetherapie an einer Quelle des Rückenschmerzes an der Stelle, die den gebundenen Gewebefaktor enthält, in Reaktion auf eine Energie, die an die Region des Patienten abgegeben wird, die zumindest eines Teils der Wirbelsäule umfasst, verbessert; und wobei der Schmerzfaktor zumindest eines umfasst von: einem Nervenfaktor, einem Blutgefäßfaktor, einem entzündungsfördernden Zytokin oder einer Kombination davon; oder einem Faktor, der mit pH-Wert oder pO₂-Wert in Gewebe assoziiert ist, der optional mit einem relativ geringen pH-Wert bzw. pO₂-Wert assoziiert ist, oder einem Bindungsmittel oder jeweiligen Antikörper davon.
3. System nach Anspruch 2, das zum Diagnostizieren oder Behandeln einer Rückenschmerzquelle an einer Stelle eines Wirbelsäulengelenks im Körper eines Patienten konfiguriert ist, wobei:
- das abgezielte Mittel so markiert wird, dass es eine abgezielte Markierung umfasst, die so gekennzeichnet ist, dass sie an einen Schmerzfaktor, der mit einer Rückenschmerzquelle an der Stelle assoziiert ist, bindet und diesen markiert, wenn sie in den Patienten abgegeben wird; die Abgabebaupruppe für das abgezielte Mittel in einer Konfiguration vorliegt, die so betreibbar ist, dass sie die Abgabe der abgezielten Markierung in den Patienten ermöglicht; und das System zumindest eines von (a) einem Bildgebungssystem in einer Konfiguration, die so betreibbar ist, dass sie zumindest eine abgezielte Markierung oder den markierten Schmerzfaktor in einer Weise abbildet, die ausreichend ist, um eine erste Konzentration des markierten Schmerz factors an einer Stelle von einer zweiten Konzentration des markierten Schmerz factors an einer zweiten Gewebestelle benachbart zu der Stelle zu unterscheiden, und (b) einer therapeutischen Vorrichtungsbaugruppe in einer Konfiguration, die so betreibbar ist, dass sie eine Therapie in einer im Wesentlichen lokalisierten Weise bereitstellt, die auf die Stelle, die den markierten Schmerzfaktor umfasst, im Wesentlichen isoliert ist, umfasst.
4. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, ferner **gekennzeichnet durch**:
- ein Volumen des abgezielten Mittels, das so markiert wird, dass das abgezielte Mittel eine abgezielte Markierung umfasst, die für eine Abgabe in den Patienten konfiguriert ist und die so gekennzeichnet ist, dass sie differentiell an den Schmerzfaktor, der mit der Rückenschmerzquelle an der Stelle assoziiert ist, bindet und diesen markiert; und wobei die abgezielte Markierung so konfiguriert ist, dass sie den Schmerzfaktor an der Stelle künstlich markiert, indem der Schmerzfaktor mit der abgezielten Markierung gebunden wird, wenn in die Region des Patienten abgegeben, die zumindest einen Abschnitt einer Wirbelsäule umfasst.
5. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei der Schmerzfaktor umfasst: einen Blutgefäßfaktor, der zumindest eines von einem Blutgefäß oder einem Mikrogefäß oder einer Substanz oder Struktur, die mit dem Blutgefäß oder Mikrogefäß assoziiert ist; oder einer Substanz P, CGRP, trkA, Nervenwachstumsfaktor (NGF), einem NGF-Antagonisten, PGP 9.5, SYN, Peripherin, Neurofilament 200 kD (NF200), TNF- α , Infliximab, PECAM, CD34, GFAP, PGE-2 oder einem Analogon oder Derivat oder Bindungsmittel oder Antikörper davon; oder einem Nervenbindungsmittel oder Antikörper; oder einem Interleukin, das optional IL-1, IL-6 oder IL-8 umfasst, oder einem Analogon oder Derivat davon; oder MIF oder einem Bindungsmittel oder Antikörper davon; oder einem CGRP-Rezeptor umfasst.
6. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 4, wobei der Schmerzfaktor oder das abgezielte Mittel umfasst: einen Nervenfaktor, der zumindest eine Substanz umfasst, die mit zumindest einem von einem Nervenfaktor oder einer mit Zellstruktur, die mit der Nervenfasern assoziiert ist, assoziiert ist.
7. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei der Schmerzfaktor oder das abgezielte Mittel einen TNF- α -Blocker umfasst.
8. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei der Schmerzfaktor umfasst: ein Interleukin, das optional IL-1, IL-6 oder IL-8 umfasst, oder ein Analogon oder Derivat davon; oder einen Faktor, der mit pH-Wert oder pO₂-Wert in Gewebe assoziiert ist, der optional mit einem relativ geringen pH-Wert bzw. pO₂-Wert assoziiert ist, oder ein Bindungsmittel oder einen jeweiligen Antikörper davon.
9. Verfahren nach Anspruch 1 oder System nach An-

spruch 2 oder 3, wobei das abgezielte Mittel umfasst: zumindest eines von einem Nervenfaktor, einem Blutgefäßfaktor, einem entzündungsfördernden Zytokin oder einem Bindungsmittel oder Antikörper davon; oder einem Bindungsmittel oder einem Antikörper des Nervenfaktors, wie in Anspruch 5 definiert; oder einem Blutgefäßfaktor, umfassend eine Substanz, die mit einer Struktur eines Blutgefäßes oder eines Mikrogefäßes assoziiert ist, oder einem Bindungsmittel oder Antikörper eines entzündungshemmenden Zytokins; oder einem Bindungsmittel oder Antikörper einer Substanz P; oder einem CGRP- oder CGRP-Rezeptor-Bindungsmittel oder -Antikörper; oder einem trkA-Bindungsmittel oder -Antikörper; oder einem Anti-trkA-Antikörper-Bindungsmittel oder -Antikörper davon; oder einem Nervenwachstumsfaktor (NGF) oder einem Analogon oder Derivat davon; oder einem NGF-Antagonisten oder einem Analogon oder Derivat davon; oder einem Nervenbindungsmittel oder Antikörper davon oder einem Analogon oder Derivat davon; oder einem NGF-Bindungsmittel; oder einem Anti-NGF-Antikörper; oder einem NGF-Antagonist-Bindungsmittel; oder einem Anti-NGF-Antagonist-Antikörper; oder einem PECAM-Bindungsmittel oder -Antikörper; oder einem CD34-Bindungsmittel oder -Antikörper; oder einem GFAP-Bindungsmittel oder -Antikörper; oder einem Interleukin-Bindungsmittel oder -Antikörper; oder einem Interleukin-Bindungsmittel oder -Antikörper, umfassend ein/en IL-1-, IL-6- oder IL-8-Bindungsmittel oder -Antikörper; oder einem PGE-2-Bindungsmittel oder -Antikörper; oder einem Faktor, der mit pH-Wert oder pO₂-Wert in Gewebe assoziiert ist, oder einem Bindungsmittel oder Antikörper davon; oder PGP 9.5 oder SYN oder Peripherin oder Neurofilament 200 kD (NF200) oder TNF- α oder Infliximab oder einem Analogon oder Derivat davon oder einem Bindungsmittel oder Antikörper davon; oder einem radioaktiven Material, optional einem radiomarkierten TNF- α -Antikörper oder einem Analogon oder Derivat davon, oder radiomarkiertem Iod, optional I-125; oder einem MIF-Bindungsmittel oder -Antikörper; oder einem Nanopartikeln; oder zumindest einem von Gold oder Eisenoxid; oder einem MRT-Kontrastmittel, optional Gadolinium; oder einem Ultraschallkontrastmittel; oder einem radiographischen Kontrastmittel.

10. System nach Anspruch 3 oder Verfahren oder System nach Anspruch 4, wobei der markierte Schmerzfaktor, wenn mit dem abgezielten Mittel markiert, für einen relativ geringen pH-Wert oder pO₂-Wert an der Stelle indikativ ist.

11. System nach Anspruch 3 oder Verfahren oder System nach Anspruch 4, umfassend: ein Bildgebungssystem, das für das Abbilden des markierten

Schmerz factors unter Verwendung eines Bildgebungswerkzeugs konfiguriert ist, das eine Phosphor-Bildgebungsplatte umfasst; oder wobei das abgezielte Mittel ein MRT-Kontrastmittel umfasst, und umfassend ein MRT-Bildgebungssystem, das für das Abbilden eines Bereichs mit erhöhter Konzentration des MRT-Kontrastmittels konfiguriert ist, das an den Schmerzfaktor gebunden ist; oder wobei das abgezielte Mittel ein Ultraschallkontrastmittel umfasst, und umfassend ein Ultraschallbildgebungssystem, das für das Ultraschallabbilden eines Bereichs mit erhöhter Konzentration des Ultraschallkontrastmittels konfiguriert ist, das an den Schmerzfaktor gebunden ist; oder wobei das abgezielte Mittel ein radiographisches Kontrastmittel umfasst, und umfassend ein Röntgensystem, das für das Abbilden eines Bereichs mit erhöhter Konzentration des radiographischen Kontrastmittels, das an den Schmerzfaktor gebunden ist, unter Verwendung von Röntgenstrahlen konfiguriert ist.

12. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 4, ferner umfassend:

ein erstes Bindungsmittel, das für eine Abgabe in den Körper hergestellt ist, wobei das erste Bindungsmittel so ausgelegt ist, dass es an einen ersten Schmerzfaktor bindet;

wobei das abgezielte Mittel für eine Abgabe in den Körper des Patienten hergestellt ist, nachdem das erste Bindungsmittel in den Körper abgegeben wurde und an den ersten Schmerzfaktor gebunden hat; und

wobei das abgezielte Mittel, wenn es an die Region des Patienten abgegeben wird, die zumindest einen Abschnitt einer Wirbelsäule umfasst, so gekennzeichnet ist, dass es an eine Stelle bindet, die sich auf der gebundenen Kombination aus dem ersten Bindungsmittel und dem ersten Schmerzfaktor befindet; und

wobei das erste Bindungsmittel optional einen bispezifischen Antikörper mit einer ersten Bindungsstelle, die so gekennzeichnet ist, dass sie an den ersten Schmerzfaktor bindet, und eine zweite Bindungsstelle, die so gekennzeichnet ist, dass sie an das abgezielte Mittel bindet umfasst.

13. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 4, umfassend:

ein Bildgebungssystem, das in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie eine Region mit diagnostischer Lokalisierung der Rückenschmerzquelle an der Stelle über das abgezielte Mittel abbildet, das selektiv

- an den Schmerzfaktor an der Stelle gebunden ist; und
ein therapeutisches System, das in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie eine therapeutische Verfahrensweise in einer im Wesentlichen lokalisierten Weise an der Stelle durchführt, an der das abgezielte Mittel, das an den Schmerzfaktor gebunden ist, lokal abgebildet ist.
- 5
14. System oder Verfahren nach Anspruch 13, wobei das therapeutische System in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie die Erzeugung oder Übertragung von Schmerz an der Stelle im Wesentlichen lindert, z. B. dadurch, dass es in einem Betriebsmodus konfiguriert ist, um zumindest einen Nerv an der Stelle zu abladien und/oder zumindest eine therapeutische Chemikalie an die Stelle abzugeben.
- 10
15. System oder Verfahren nach Anspruch 13, wobei das therapeutische System umfasst:
ein Energieabgabesystem, das in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie eine therapeutische Dosis an Energie in einer im Wesentlichen lokalisierten Weise an die Stelle abgibt, optional konfiguriert, um zumindest einen Nerv an der Stelle mit der Energie zu abladien.
- 15
16. Verfahren oder System nach Anspruch 15, wobei:
das therapeutische System zur Abgabe von Ultraschallenergie an die Stelle auf gerichtete Weise von einer zweiten Stelle konfiguriert ist; die zweite Stelle außerhalb des Patienten liegt; und
die Ultraschallenergie so konfiguriert ist, dass sie über fokussierten Hochintensitätultraschall (HIFU) abgegeben wird, der so ausgelegt ist, dass er die Ultraschallenergie an die Stelle fokussiert.
- 20
17. Verfahren oder System nach Anspruch 16, wobei:
die zweite Stelle zu der Stelle in dem Patienten benachbart liegt; und
das System so konfiguriert ist, dass es Ultraschallenergie über eine directionale Ultraschallsonde abgibt.
- 25
18. Verfahren oder System nach Anspruch 15, wobei ein therapeutisches System in einer Konfiguration zur Abgabe der therapeutischen Dosis an Energie, die Wärme-, elektrische, Mikrowellen- oder Lichtenergie umfasst, konfiguriert ist.
- 30
19. Verfahren oder System nach Anspruch 18, umfassend eine Funkfrequenz-(HF-)Sonde, die mit einer
- Quelle elektrischer Energie für die HF-Sonde verbunden ist, so dass das System in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie die therapeutische Dosis an elektrischer Energie über die Funkfrequenz-(HF-)Sonde abgibt.
20. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei die Stelle des Wirbelsäulengelenks umfasst:
(a) zumindest einen Abschnitt einer Bandscheibe; oder
(b) eine Region von Gewebe, das innerhalb nur eines Abschnitts angeordnet ist, der einem gesamten Umgang einer Bandscheibe gleicht bzw. kleiner als dieser ist, optional eine Region von Gewebe, das sich innerhalb weniger als oder gleich einer Hälfte oder einem Viertel des Umfangs der Bandscheibe befindet; oder
(c) eine Endplatte, die mit einem Wirbeltierkörper assoziiert ist.
21. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei:
das abgezielte Mittel für eine Abgabe in lokalisierter Weise an die Stelle hergestellt ist, optional für eine Injektion in eine Region von Gewebe, das mit der Stelle assoziiert ist, unter Verwendung einer Lokalinjektionsbaugruppe.
22. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei:
das abgezielte Mittel für eine systemische Abgabe in den Patienten hergestellt ist, optional für eine Abgabe in den systemischen Blutkreislauf, das Magen-Darm-System oder das Atmungssystem des Patienten hergestellt ist.
23. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 4, wobei:
das abgezielte Mittel für eine Abgabe in den Patienten hergestellt ist, die so gekennzeichnet ist, dass sie eine künstliche Markierung des Schmerzfaktors an den mehreren Stellen durch Bindung des Schmerzfaktors mit dem abgezielten Mittel, wenn es in den Patienten abgegeben wurde, ermöglicht; und
ein Bildgebungswerkzeug bereitgestellt ist, das in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie zumindest eines des abgezielten Mittels oder des markierten Schmerzfaktors abbildet, und in einer Weise, die ausreichend ist, um eine erste Konzentration des markierten Schmerzfaktors an einer der mehreren Stellen von einer zweiten Konzentration des markierten Schmerzfaktors an einer anderen

der mehreren Stellen zu unterscheiden.

24. System oder Verfahren nach Anspruch 23, das umfasst:
 zumindest ein therapeutisches System, das in einer Konfiguration konfiguriert ist, die so betreibbar ist, dass sie eine therapeutische Verfahrensweise in einer im Wesentlichen lokalisierten Weise an jeder der Stellen, an denen der abgezielte markierte Schmerzfaktor lokal und selektiv abgebildet ist, durchführt. 5
25. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 4, das umfasst:
 ein Energieabgabesystem, das so konfiguriert und betreibbar ist, dass es Energie in den Patienten auf eine Weise abgibt, die eine oder mehrere Regionen, die mit differentiell gebundenem Schmerzfaktor assoziiert sind, wenn das abgezielte Mittel in den Patienten abgegeben wurde, differentiell behandelt. 15
26. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 4, das umfasst:
 ein Energieabgabesystem;
 wobei das Energieabgabesystem so konfiguriert ist, dass es Energie in den Patienten abgibt; und
 wobei das Volumen von abgezieltem Mittel so konfiguriert und betreibbar ist, dass es in einen Patienten abgegeben wird und differentiell an einen Schmerzfaktor bindet, der mit dem Schmerz assoziiert ist, in einer Weise, die so gekennzeichnet ist, dass sie ermöglicht, dass Geweberegionen, die eine erste Konzentration des differentiell gebundenen Schmerzfaktors enthalten, ein differentiell und selektives therapeutisches Ansprechen auf die Energie, wenn abgegeben, in Bezug auf andere Regionen mit geringeren Konzentrationen des differentiell gebundenen Schmerzfaktors aufweist. 20
27. Verfahren nach Anspruch 1, System nach Anspruch 2 oder 3 oder System oder Verfahren nach Anspruch 26, **gekennzeichnet dadurch, dass:**
 eine therapeutische Vorrichtungsbaugruppe bereitgestellt ist, die eine Energieabgabebaugruppe umfasst, die so konfiguriert und betreibbar ist, dass sie eine therapeutische Dosis an Energie in einer im Wesentlichen lokalisierten Weise abgibt, die auf die Stelle, die mit dem Wirbelsäulengelenk assoziiert ist, im Wesentlichen isoliert ist. 25
28. System oder Verfahren nach Anspruch 27, wobei: die Energieabgabebaugruppe so konfiguriert und betreibbar ist, dass sie in den Patienten an eine Position an oder benachbart zu der Stelle abgegeben 30

wird.

29. System oder Verfahren nach Anspruch 27, das ferner umfasst:
 ein Einbringelement, das so konfiguriert und betreibbar ist, dass es die Energieabgabebaugruppe an die Stelle abgibt. 35
30. System oder Verfahren nach Anspruch 29:
 wobei das Einbringelement eine Nadelbaugruppe umfasst; oder
 wobei das Einbringelement eine Nadelbaugruppe umfasst, die so konfiguriert und betreibbar ist, dass sie durch Knochen fortbewegt wird und die therapeutische Vorrichtungsbaugruppe an eine Position innerhalb des Knochens abgibt; oder
 wobei das Einbringelement eine Nadelbaugruppe umfasst, die so konfiguriert und betreibbar ist, dass sie durch Knochen fortbewegt wird und die therapeutische Vorrichtungsbaugruppe an eine Position innerhalb des Knochens abgibt, und wobei die therapeutische Vorrichtungsbaugruppe so konfiguriert und betreibbar ist, dass sie einen intraossären Nerv abladiert, der innerhalb des Knochens liegt und der mit Schmerz in Zusammenhang mit dem markierten Schmerzfaktor assoziiert ist, der an der Stelle visuell dargestellt ist; oder
 wobei das Einbringelement eine Nadelbaugruppe umfasst, die so konfiguriert und betreibbar ist, dass sie durch Knochen eines Wirbeltierkörpers fortbewegt wird und die therapeutische Vorrichtungsbaugruppe an eine Position innerhalb des Wirbeltierkörpers abgibt, die mit einem basivertebralen Nerv assoziiert ist, und wobei die therapeutische Vorrichtungsbaugruppe so konfiguriert und betreibbar ist, dass sie den basivertebralen Nerv von der Position abladiert; oder
 wobei das Einbringelement eine Nadelbaugruppe umfasst, die so konfiguriert und betreibbar ist, dass sie durch Knochen fortbewegt wird und die therapeutische Vorrichtungsbaugruppe an eine Position innerhalb des Knochens abgibt, wobei die therapeutische Vorrichtungsbaugruppe eine Funkfrequenz-(HF-)Stromablationsbaugruppe umfasst, wobei die HF-Stromablationsbaugruppe eine erste Elektrode und eine zweite Elektrode umfasst, die so ausgelegt sind, dass sie an ersten und zweiten Positionen positioniert werden, die so ausgelegt sind, dass sie zumindest einen Abschnitt des basivertebralen Nervs überspannen, und wobei die HF-Stromablationsbaugruppe so konfiguriert und betreibbar ist, dass sie den HF-Strom zwischen der ersten und der zweiten Elektrode in einer 40

Weise abgibt, die ausreichend ist, um Nervengewebe zwischen den ersten und zweiten Positionen zu abladieren, und ferner wobei die HF-Stromablationsbaugruppe optional eine Abgabesonde mit einem länglichen Körper, der die erste und die zweite Elektrode trägt, in einer bipolaren Leitungsbaugruppenanordnung umfasst.

31. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei:

das abgezielte Mittel ein Material umfasst, das eine bevorzugte Bindungsaffinität für den Schmerzfaktor aufweist, der sich innerhalb der Region an der Stelle befindet, die ausreichend ist, um in Bezug auf andere Struktur innerhalb der Region bevorzugt an den Schmerzfaktor zu binden, und so dass sich das Material in einer höheren Konzentration innerhalb eines ersten Abschnitts der Region mit einer höheren Menge des Schmerzfaktors als andere Abschnitte innerhalb der Region ansammelt; und ein Energieabgabesystem beinhaltend, das für eine Abgabe von Energie an die Region auf eine Weise konfiguriert und betreibbar ist, die die Stelle des ersten Abschnitts im Vergleich zu den anderen Abschnitten selektiv behandelt, nachdem das Material an die Region abgegeben wurde, und so dass Schmerz in der Region verringert ist.

32. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3 oder Verfahren oder System nach Anspruch 31, wobei das abgezielte Mittel ein Material umfasst, das umfasst: ein Metall, optional Gold; oder einen Nanopartikel; oder einen Goldnanopartikel; oder einen Antikörper; oder einen Antikörper und ein Metall, optional einen Metallnanopartikel, optional einen Goldnanopartikel, assoziiert mit dem Antikörper.

33. Verfahren nach Anspruch 1 oder System nach Anspruch 2 oder 3, wobei:

das abgezielte Mittel in einer Konfiguration konfiguriert ist, die in einem Verfahren zur Abgabe in zumindest eine erste Region, die die Stelle umfasst, und zumindest eine andere Region, die sich von der ersten Region unterscheidet, betreibbar ist; die erste Region zumindest eine Wirbelsäulengelenkebene entlang der Wirbelsäule umfasst, optional ein einzelnes Wirbelsäulengelenk, oder einen Bereich eines Wirbelsäulengelenks, optional zumindest einen Teil einer Bandscheibe oder eines Wirbeltierkörpers oder einer Wirbeltierkörper-Endplatte, oder ein Facettengelenk

oder einen Querfortsatz; und wobei die zumindest eine andere Region zumindest eine andere Wirbelsäulengelenkebene entlang der Wirbelsäule umfasst.

34. Verfahren oder System nach Anspruch 31, das umfasst:

ein Bildgebungssystem, das zum Abbilden der Region auf eine Weise konfiguriert und betreibbar ist, die räumliche Beziehungen zwischen verschiedenen Konzentrationen des Materials ausreichend unterscheidet, um die Stelle des ersten Abschnitts relativ zu den anderen Abschnitten innerhalb der Region im Wesentlichen zu identifizieren.

35. Verfahren oder System nach Anspruch 31, wobei: das Energieabgabesystem zum Abgeben der Energie vorwiegend an den ersten Abschnitt auf eine im Wesentlichen lokalisierten Weise konfiguriert und betreibbar ist, die ausreichend ist, um den ersten Abschnitt gegenüber den anderen Abschnitten differenziell mit der Energie zu behandeln.

36. Verfahren nach Anspruch 1, wobei: das abgezielte Mittel für eine Abgabe in den Patienten hergestellt ist, nachdem der Patient diagnostiziert wurde, und zwar auf eine Weise, die die Region als schmerzhafte Region des Körpers des Patienten identifiziert.

37. Verfahren nach Anspruch 1 oder Verfahren oder System nach Anspruch 15, wobei: die Stelle nicht als Krebszellen enthaltend diagnostiziert wird, bevor der medizinische Eingriff durchgeführt wird.

38. Verfahren oder System nach Anspruch 17, wobei:

sich die zweite Stelle benachbart zu einer Bandscheibe befindet; und die directionale Ultraschallsonde zur Abgabe der directionalen Ultraschalltherapie in die Stelle innerhalb der Bandscheibe konfiguriert und betreibbar ist.

Revendications

1. Procédé pour la préparation d'un système comprenant un agent ciblé pour réaliser une procédure médicale sur un patient chez lequel une douleur dorsale a été diagnostiquée, **caractérisé par** :

sur la base du diagnostic de douleur dorsale, la préparation d'un volume d'un agent ciblé en combinaison avec un ensemble d'administration dans une configuration qui est exploitable pour permettre l'administration de l'agent ciblé

dans une région du patient qui comprend au moins une partie d'une colonne vertébrale, l'agent ciblé étant **caractérisé en ce qu'**il se lie différenciellement à un facteur de douleur à un emplacement d'une articulation vertébrale associée avec une source de la douleur dorsale lorsqu'il est administré au patient ;
 la configuration d'un système d'administration d'énergie pour administrer de l'énergie au patient et/ou d'un système d'imagerie pour obtenir une image du facteur de douleur à un emplacement d'une articulation vertébrale ;
 l'agent ciblé lorsqu'il est administré dans la région du patient qui comprend au moins une partie d'une colonne vertébrale étant en outre **caractérisé en ce qu'**il améliore (i) une localisation diagnostique de la source de la douleur dorsale à l'emplacement dans le corps du patient lorsque le système d'imagerie est exploité dans la configuration pour obtenir une image de l'emplacement d'une articulation vertébrale et/ou (ii) un traitement tissulaire sélectif de la source de la douleur dorsale à l'emplacement lorsque le système d'administration d'énergie est exploité dans une configuration pour administrer de l'énergie à l'emplacement d'une articulation vertébrale ; et
 l'agent ciblé comprenant au moins un parmi : un facteur nerveux, un facteur de vaisseau sanguin, une cytokine inflammatoire ou une combinaison de ceux-ci ; ou un facteur associé avec le pH ou le pO₂ dans le tissu, éventuellement associé avec un pH ou pO₂ faible, respectivement, ou un agent de liaison ou anticorps respectif de celui-ci.

2. Système, qui comprend un agent ciblé pour réaliser une procédure médicale sur un patient, ledit agent ciblé étant préparé pour une administration dosée dans un patient chez qui une douleur dorsale a été diagnostiquée, **caractérisé par** :

un ensemble d'administration pour l'agent ciblé qui est configuré et exploitable pour permettre une administration de l'agent ciblé dans un emplacement d'une articulation vertébrale dans le patient ; et
 ledit agent ciblé qui, lorsqu'il est administré dans une région du patient qui comprend au moins une partie d'une colonne vertébrale via l'ensemble d'administration, est **caractérisé en ce qu'**il se lie différenciellement à un facteur de douleur à un emplacement d'une articulation associée avec la douleur dorsale d'une manière en outre **caractérisée en ce qu'**il améliore (i) une localisation diagnostique d'une source de la douleur dorsale à l'emplacement d'une articulation vertébrale et/ou (ii) un traitement tissulaire sélectif

d'une source de la douleur dorsale à l'emplacement contenant le facteur de douleur lié en réponse à une énergie administrée dans la région du patient qui comprend au moins une partie d'une colonne vertébrale ; et
 le facteur de douleur comprenant au moins un parmi : un facteur nerveux, un facteur de vaisseau sanguin, une cytokine inflammatoire ou une combinaison de ceux-ci ; ou un facteur associé avec le pH ou le pO₂ dans le tissu, éventuellement associé avec un pH ou pO₂ relativement faible, respectivement, ou un agent de liaison ou anticorps respectif de celui-ci.

3. Système selon la revendication 2, configuré pour le diagnostic ou le traitement d'une source de douleur dorsale à un emplacement d'une articulation vertébrale dans un corps d'un patient, dans lequel :

l'agent ciblé est marqué de manière à comprendre un marqueur ciblé qui est **caractérisé en ce qu'**il se lie à et marque un facteur de douleur associé avec une source de douleur dorsale à l'emplacement lorsqu'il est administré dans le patient ;
 l'ensemble d'administration pour l'agent ciblé est dans une configuration qui est exploitable pour permettre l'administration du marqueur ciblé dans le patient ; et
 le système comprend (a) un système d'imagerie dans une configuration qui est exploitable pour obtenir une image du marqueur ciblé et/ou du facteur de douleur marqué d'une manière suffisante pour différencier sélectivement une première concentration du facteur de douleur marqué à l'emplacement d'une deuxième concentration du facteur de douleur marqué à un deuxième emplacement tissulaire adjacent à l'emplacement, et/ou (b) un ensemble dispositif thérapeutique dans une configuration qui est exploitable pour fournir un traitement d'une manière essentiellement localisée qui est essentiellement isolée à l'emplacement comprenant le facteur de douleur marqué.

4. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, en outre **caractérisé** :

par un volume de l'agent ciblé qui est marqué tel que l'agent marqué comprenne un marqueur ciblé qui est configuré pour l'administration dans le patient et qui est **caractérisé en ce qu'**il se lie différenciellement à et marque le facteur de douleur associé avec la source de la douleur dorsale à l'emplacement ; et
en ce que le marqueur ciblé est configuré pour marquer artificiellement le facteur de douleur à l'emplacement par liaison du facteur de douleur

- avec le marqueur ciblé lorsqu'il est administré dans la région du patient qui comprend au moins une partie d'une colonne vertébrale.
5. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel le facteur de douleur comprend :
 - un facteur de vaisseau sanguin qui comprend au moins un parmi un vaisseau ou microvaisseau sanguin, ou une substance ou structure associée avec le vaisseau ou microvaisseau sanguin ; ou la substance P, le PRGC, le trkA, le facteur de croissance nerveuse (NGF), un antagoniste du NGF, le PGP 9.5, le SYN, la périphérine, le neurofilament 200kD (NF200), le TNF- α , l'infliximab, le PECAM, le CD34, le GFAP, le PGE-2, ou un analogue ou dérivé ou agent de liaison ou anticorps de ceux-ci ; ou un agent de liaison nerveuse ou un anticorps ; ou une interleukine comprenant éventuellement IL-1, IL-6 ou IL-8, ou un analogue ou dérivé de celle-ci ; ou le MIF ou un agent de liaison ou anticorps de celui-ci ; ou un récepteur du PRGC.
 6. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 4, dans lequel le facteur de douleur ou l'agent ciblé comprend :
 - un facteur nerveux qui comprend au moins une substance associée avec une fibre nerveuse et/ou une structure cellulaire associée avec la fibre nerveuse.
 7. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel le facteur de douleur ou l'agent ciblé comprend un bloquant de TNF- α .
 8. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel le facteur de douleur comprend : une interleukine comprenant éventuellement IL-1, IL-6 ou IL-8 ou un analogue ou dérivé de celle-ci ; ou un facteur associé avec le pH ou le pO₂ dans le tissu, éventuellement associé avec un pH ou pO₂ relativement faible, respectivement, ou un agent de liaison ou anticorps respectif de celui-ci.
 9. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel l'agent ciblé comprend :
 - au moins un parmi un facteur nerveux, un facteur de vaisseau sanguin, une cytokine inflammatoire ou un agent de liaison ou anticorps de ceux-ci ; ou un agent de liaison ou anticorps du facteur nerveux défini dans la revendication 5 ; ou un facteur de vaisseau sanguin qui comprend une substance associée avec une structure d'un vaisseau ou microvaisseau sanguin, ou un agent de liaison ou anticorps de celui-ci ; ou un agent de liaison ou anticorps d'une cytokine inflammatoire ; ou un agent de liaison ou anticorps de la substance P ; ou un agent de liaison ou anticorps du PRGC ou récepteur du PRGC ; ou un agent de liaison ou anticorps de trkA ; ou un agent de liaison d'un anticorps anti-trkA ou un anticorps de celui-ci ; ou un facteur de croissance nerveuse (NGF) ou un analogue ou dérivé de celui-ci ; ou un antagoniste de NGF ou un analogue ou dérivé de celui-ci ; ou un agent de liaison nerveuse ou un anticorps de celui-ci ou un analogue ou dérivé de celui-ci ; ou un agent de liaison à NGF ; ou un anticorps anti-NGF ; ou un agent de liaison d'un antagoniste de NGF ; ou un anticorps d'un antagoniste anti-NGF ; ou un agent de liaison ou anticorps de PECAM ; ou un agent de liaison ou anticorps de CD34 ; ou un agent de liaison ou anticorps de GFAP ; ou un agent de liaison ou anticorps d'une interleukine ; ou un agent de liaison ou anticorps d'une interleukine qui comprend un agent de liaison ou anticorps d'IL-1, IL-6 ou IL-8 ; ou un agent de liaison ou anticorps de PGE-2 ; ou un facteur associé avec le pH ou le pO₂ dans le tissu ou un agent de liaison ou anticorps de celui-ci ; ou le PGP 9.5, ou le SYN, ou la périphérine, ou le neurofilament 200kD (NF200), ou le TNF- α , ou l'infliximab, ou un analogue ou dérivé de ceux-ci, ou un agent de liaison ou anticorps de ceux-ci ; ou un matériau radioactif, éventuellement un TNF- α radiomarqué, un anticorps ou un analogue ou dérivé de celui-ci, ou de l'iode radiomarqué, éventuellement I-125 ; ou un agent de liaison ou anticorps de MIF ; ou une nanoparticule ; ou l'or et/ou l'oxyde de fer ; ou un agent de contraste d'IRM, éventuellement le gadolinium ; ou un agent de contraste pour ultrasons ; ou un agent de contraste radiographique.
 10. Système selon la revendication 3 ou procédé ou système selon la revendication 4, dans lequel le facteur de douleur marqué lorsqu'il est marqué avec l'agent ciblé est indicateur d'un pH ou pO₂ relativement faible à l'emplacement.
 11. Système selon la revendication 3 ou procédé ou système selon la revendication 4, comprenant :
 - un système d'imagerie configuré pour obtenir une image du facteur de douleur marqué en utilisant un outil d'imagerie qui comprend une plaque d'imagerie au phosphore ; ou dans lequel l'agent ciblé comprend un agent de contraste d'IRM, et comprenant un système d'imagerie IRM configuré pour obtenir une image d'une zone de concentration augmentée de l'agent de contraste d'IRM lié au facteur de douleur ; ou dans lequel l'agent ciblé comprend un agent de contraste pour ultrasons, et comprenant un système d'imagerie par ultrasons configuré pour obtenir par ultrasons une image d'une zone de concentration augmentée de l'agent de contraste pour ultrasons lié au facteur de douleur ; ou dans lequel

l'agent ciblé comprend un agent de contraste radiographique, et comprenant un système à rayons X configuré pour obtenir une image d'une zone de concentration augmentée de l'agent de contraste radiographique lié au facteur de douleur en utilisant des rayons X.

12. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 4, comprenant en outre :

un premier agent de liaison préparé pour l'administration dans le corps, le premier agent de liaison étant conçu pour se lier à un premier facteur de douleur ;

l'agent ciblé étant préparé pour l'administration dans le corps du patient après que le premier agent de liaison ait été administré dans le corps et lié au premier facteur de douleur ; et

l'agent ciblé lorsqu'il est administré dans la région du patient qui comprend au moins une partie d'une colonne vertébrale étant **caractérisé en ce qu'il se lie à un site situé sur la combinaison liée du premier agent de liaison et du premier facteur de douleur** ; et

le premier agent de liaison comprenant éventuellement un anticorps bispécifique comprenant un premier site de liaison qui est **caractérisé en ce qu'il se lie au premier facteur de douleur** et un deuxième site de liaison qui est **caractérisé en ce qu'il se lie à l'agent ciblé**.

13. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 4, comprenant :

un système d'imagerie configuré dans une configuration qui est exploitable pour obtenir une image d'une région avec localisation diagnostique de la source de la douleur dorsale à l'emplacement via l'agent ciblé lié sélectivement au facteur de douleur à l'emplacement ; et

un système thérapeutique configuré dans une configuration qui est exploitable pour réaliser une procédure thérapeutique d'une manière essentiellement localisée à l'emplacement où une image de l'agent ciblé lié au facteur de douleur est obtenue localement.

14. Système ou procédé selon la revendication 13, dans lequel le système thérapeutique est configuré dans une configuration qui est exploitable pour atténuer sensiblement la génération ou la transmission de douleur à l'emplacement, tel qu'en étant configuré dans un mode d'exploitation pour essentiellement réaliser l'ablation d'au moins un nerf à l'emplacement et/ou pour administrer au moins un produit chimique thérapeutique à l'emplacement.

15. Système ou procédé selon la revendication 13, dans lequel le système thérapeutique comprend : un système d'administration d'énergie configuré dans une configuration qui est exploitable pour administrer une dose thérapeutique d'énergie d'une manière essentiellement localisée à l'emplacement, éventuellement configuré pour réaliser l'ablation d'au moins un nerf à l'emplacement avec l'énergie.

16. Procédé ou système selon la revendication 15, dans lequel :

le système thérapeutique est configuré pour administrer une énergie ultrasonore à l'emplacement d'une manière dirigée à partir d'un deuxième emplacement ;

le deuxième emplacement est à l'extérieur du patient ; et

l'énergie ultrasonore est configurée pour être administrée via des ultrasons focalisés de haute intensité (HIFU) qui sont conçus pour focaliser l'énergie ultrasonore à l'emplacement.

17. Procédé ou système selon la revendication 16, dans lequel :

le deuxième emplacement est adjacent à l'emplacement dans le patient ; et

le système est configuré pour délivrer une énergie ultrasonore via une sonde à ultrasons directionnels.

18. Procédé ou système selon la revendication 15, dans lequel le système thérapeutique est configuré dans une configuration qui administre la dose thérapeutique d'énergie qui comprend une énergie thermique, électrique, micro-onde ou lumineuse.

19. Procédé ou système selon la revendication 18, comprenant une sonde à radiofréquence (RF) couplée à une source d'énergie électrique pour la sonde RF, de telle sorte que le système soit configuré dans une configuration qui est exploitable pour administrer la dose thérapeutique d'énergie électrique via la sonde à radiofréquence (RF).

20. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel l'emplacement de l'articulation vertébrale comprend :

(a) au moins une partie d'un disque intervertébral ; ou

(b) une région de tissu située dans uniquement une partie qui est égale à moins d'une circonférence entière d'un disque intervertébral, éventuellement une région de tissu située dans une partie qui est inférieure ou égale à la moitié ou un quart de la circonférence du disque

- intervertébral ; ou
(c) un plateau associé avec un corps vertébral.
21. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel :
l'agent ciblé est préparé pour l'administration d'une manière localisée à l'emplacement, éventuellement pour l'injection dans une région de tissu associée avec l'emplacement en utilisant un ensemble d'injection locale.
22. Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel :
l'agent ciblé est préparé pour l'administration systémique dans le patient, éventuellement pour l'administration dans la circulation sanguine systémique, le système gastrointestinal ou le système respiratoire du patient.
23. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 4, dans lequel :
- l'agent ciblé est préparé pour l'administration dans le patient qui est **caractérisée en ce qu'**elle permet un marquage artificiel du facteur de douleur à plusieurs desdits emplacements par liaison du facteur de douleur avec l'agent ciblé lorsqu'il est administré dans le patient ; et un outil d'imagerie est prévu, qui est configuré dans une configuration qui est exploitable pour obtenir une image de l'agent ciblé et/ou du facteur de douleur marqué, et d'une manière suffisante pour différencier une première concentration du facteur de douleur marqué à un des multiples desdits emplacements d'une deuxième concentration du facteur de douleur marqué à un autre desdits multiples emplacements.
24. Système ou procédé selon la revendication 23, comprenant :
au moins un système thérapeutique configuré dans une configuration qui est exploitable pour réaliser une procédure thérapeutique d'une manière essentiellement localisée à chacun des emplacements où une image du facteur de douleur marqué ciblé est obtenue localement et sélectivement.
25. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 4, comprenant :
- un système d'administration d'énergie configuré et exploitable pour administrer de l'énergie dans le patient d'une manière qui traite différenciellement une ou plusieurs régions associées avec un facteur de douleur lié différenciellement lorsque l'agent ciblé est administré dans le patient.
26. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 4, comprenant :
- un système d'administration d'énergie ;
le système d'administration d'énergie étant configuré pour administrer de l'énergie dans le patient ; et
le volume d'agent ciblé étant configuré et exploitable pour l'administration dans un patient et pour se lier différenciellement à un facteur de douleur associé avec la douleur d'une manière **caractérisée en ce qu'**elle permet à des régions tissulaires contenant une première concentration du facteur de douleur lié différenciellement de présenter une réponse thérapeutique différencielle et sélective à l'énergie lorsqu'elle est administrée en comparaison d'autres régions présentant des concentrations plus faibles du facteur de douleur lié différenciellement.
27. Procédé selon la revendication 1, système selon la revendication 2 ou 3, ou système ou procédé selon la revendication 26, **caractérisé par** :
- un ensemble dispositif thérapeutique, qui comprend un ensemble d'administration d'énergie qui est configuré et exploitable pour administrer une dose thérapeutique d'énergie d'une manière essentiellement localisée qui est essentiellement isolée à l'emplacement associé avec l'articulation vertébrale.
28. Système ou procédé selon la revendication 27, dans lequel :
l'ensemble d'administration d'énergie est configuré et exploitable pour être administré dans le patient à une position à ou adjacente à l'emplacement.
29. Système ou procédé selon la revendication 27, comprenant en outre :
un introducteur qui est configuré et exploitable pour administrer l'ensemble d'administration d'énergie à l'emplacement.
30. Système ou procédé selon la revendication 29 :
- dans lequel l'introducteur comprend un ensemble aiguille ; ou
dans lequel l'introducteur comprend un ensemble aiguille configuré et exploitable pour être avancé au travers d'un os et pour administrer l'ensemble dispositif thérapeutique à une position dans l'os ; ou
dans lequel l'introducteur comprend un ensemble aiguille configuré et exploitable pour être avancé au travers d'un os et pour administrer l'ensemble dispositif thérapeutique à une position dans l'os, et dans lequel l'ensemble dispositif thérapeutique est configuré et exploitable

pour réaliser l'ablation d'un nerf intra-osseux dans l'os et qui est associé avec la douleur relative au facteur de douleur marqué visualisé à l'emplacement ; ou

dans lequel l'introducteur comprend un ensemble aiguille configuré et exploitable pour être avancé au travers d'un os d'un corps vertébral et pour administrer l'ensemble dispositif thérapeutique à une position dans le corps vertébral associée avec un nerf basivertébral, et dans lequel l'ensemble dispositif thérapeutique est configuré et exploitable pour réaliser l'ablation du nerf basivertébral à partir de la position ; ou dans lequel l'introducteur comprend un ensemble aiguille configuré et exploitable pour être avancé au travers d'un os et pour administrer l'ensemble dispositif thérapeutique à une position dans l'os, l'ensemble dispositif thérapeutique comprenant un ensemble d'ablation à courant de radiofréquence (RF), l'ensemble d'ablation à courant RF comprenant une première électrode et une deuxième électrode conçues pour être positionnées à une première et une deuxième position conçues pour chevaucher au moins une partie du nerf basivertébral, et l'ensemble d'ablation à courant RF étant configuré et exploitable pour administrer le courant RF entre la première et la deuxième électrode d'une manière suffisante pour réaliser l'ablation du tissu nerveux entre la première et la deuxième position et, par ailleurs, l'ensemble d'ablation à courant RF comprenant éventuellement une sonde d'administration à corps allongé qui porte la première et la deuxième électrode dans un agencement d'ensemble de conducteurs bipolaires.

- 31.** Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel :

l'agent ciblé comprend un matériau qui présente une affinité de liaison préférentielle avec le facteur de douleur situé dans la région à l'emplacement, suffisante pour se lier préférentiellement au facteur de douleur en comparaison d'autres structures dans la région, et de telle sorte que le matériau s'accumule à une concentration plus élevée dans une première partie de la région contenant une quantité plus importante du facteur de douleur que d'autres parties dans la région ; et

comprenant un système d'administration d'énergie configuré et exploitable pour administrer de l'énergie dans la région d'une manière qui traite sélectivement l'emplacement de la première partie en comparaison des autres parties, après l'administration du matériau dans la région, et de telle sorte que la douleur puisse être

réduite dans la région.

- 32.** Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, ou procédé ou système selon la revendication 31, dans lequel l'agent ciblé comprend un matériau qui comprend : un métal, éventuellement l'or ; ou une nanoparticule ; ou une nanoparticule d'or ; ou un anticorps ; ou un anticorps et un métal, éventuellement une nanoparticule de métal, éventuellement une nanoparticule d'or, associée avec l'anticorps.

- 33.** Procédé selon la revendication 1 ou système selon la revendication 2 ou 3, dans lequel :

l'agent ciblé est configuré dans une configuration qui est exploitable dans un procédé pour l'administration dans au moins une première région qui comprend l'emplacement et au moins une autre région qui est différente de la première région ;

la première région comprend au moins un niveau d'articulation vertébrale le long de la colonne vertébrale, éventuellement une articulation vertébrale individuelle, ou une zone d'une articulation vertébrale, éventuellement au moins une partie d'un disque intervertébral, ou un corps vertébral, ou un plateau de corps vertébral, ou une articulation facettaire ou une apophyse transverse ; et

ladite au moins une autre région comprend au moins un autre niveau d'articulation vertébrale le long de la colonne vertébrale.

- 34.** Procédé ou système selon la revendication 31, comprenant :

un système d'imagerie configuré et exploitable pour obtenir une image de la région d'une manière qui différencie suffisamment des relations spatiales entre différentes concentrations du matériau de manière à identifier essentiellement l'emplacement de la première partie par rapport aux autres parties dans la région.

- 35.** Procédé ou système selon la revendication 31, dans lequel :

le système d'administration d'énergie est configuré et exploitable pour administrer l'énergie principalement dans la première partie d'une manière essentiellement localisée suffisante pour traiter différenciellement la première partie avec l'énergie en comparaison des autres parties.

- 36.** Procédé selon la revendication 1, dans lequel : l'agent ciblé est préparé pour l'administration dans le patient après que le patient ait été diagnostiqué d'une manière qui identifie la région comme une région douloureuse du corps du patient.

37. Procédé selon la revendication 1 ou procédé ou système selon la revendication 15, dans lequel :
l'emplacement n'est pas diagnostiqué comme incluant des cellules cancéreuses avant la réalisation de la procédure médicale. 5

38. Procédé ou système selon la revendication 17, dans lequel :

le deuxième emplacement est adjacent à un disque intervertébral ; et 10

la sonde à ultrasons directionnels est configurée et exploitable pour administrer le traitement par ultrasons directionnels dans l'emplacement dans le disque intervertébral. 15

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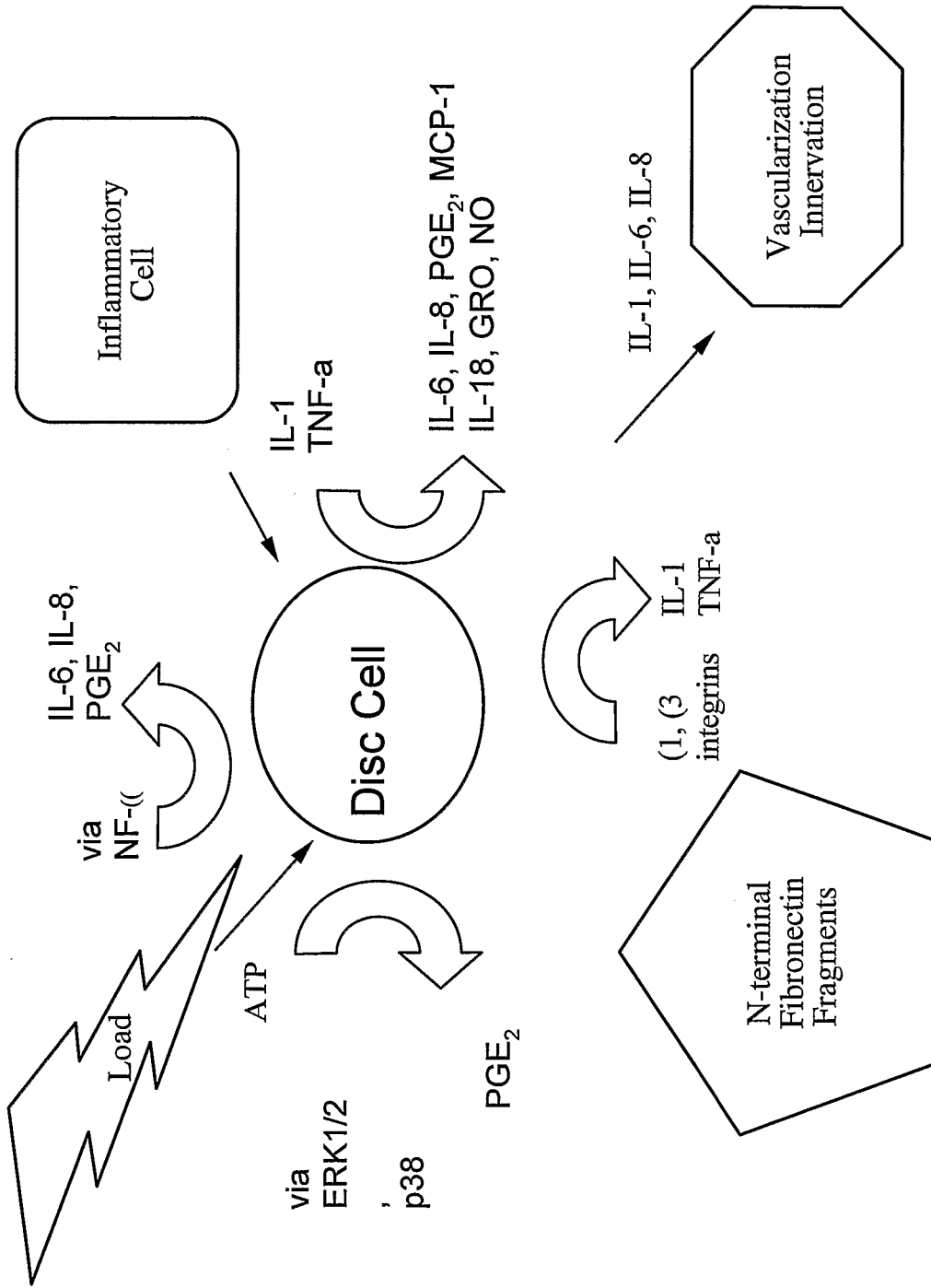


FIG. 1

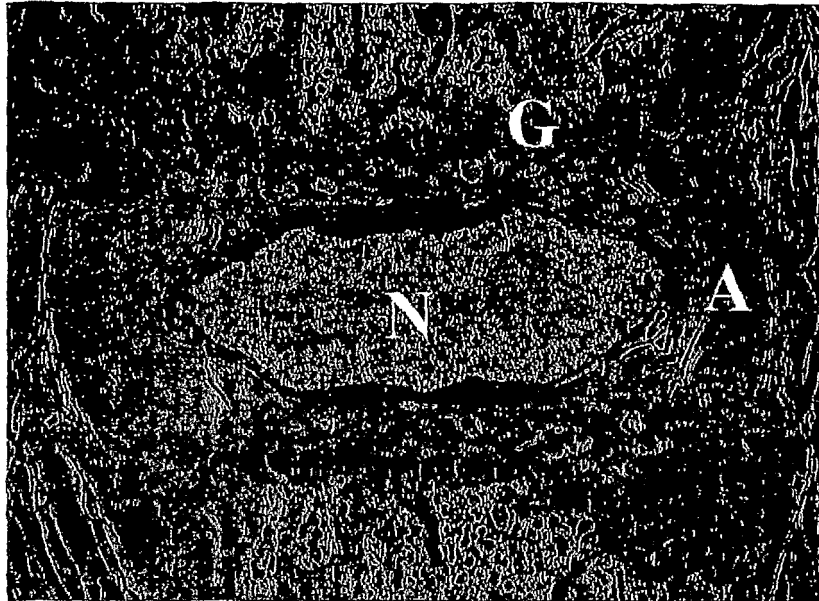


FIG. 2A

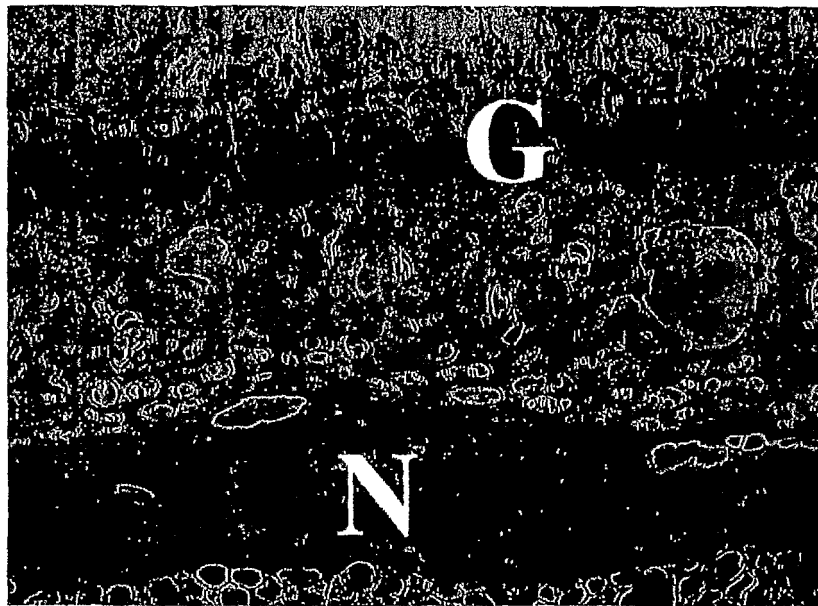


FIG. 2B

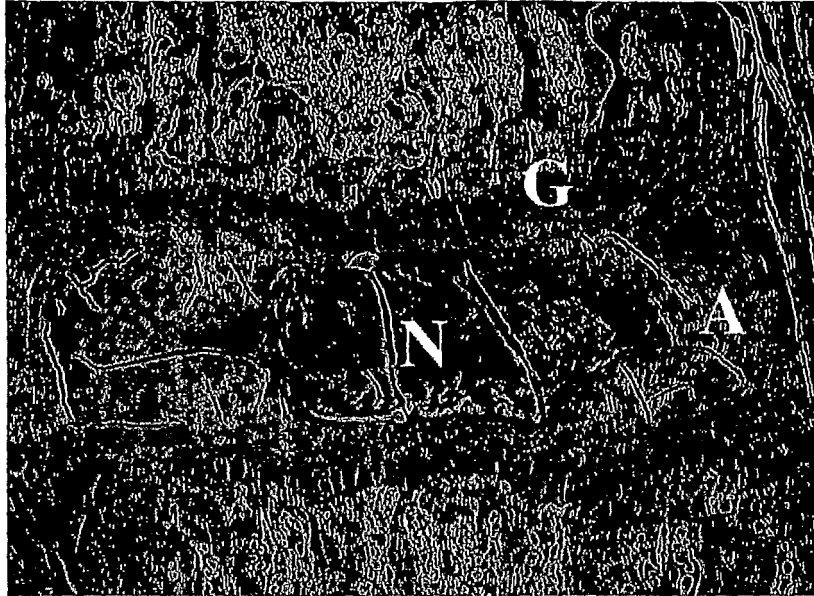


FIG. 2C



FIG. 2D

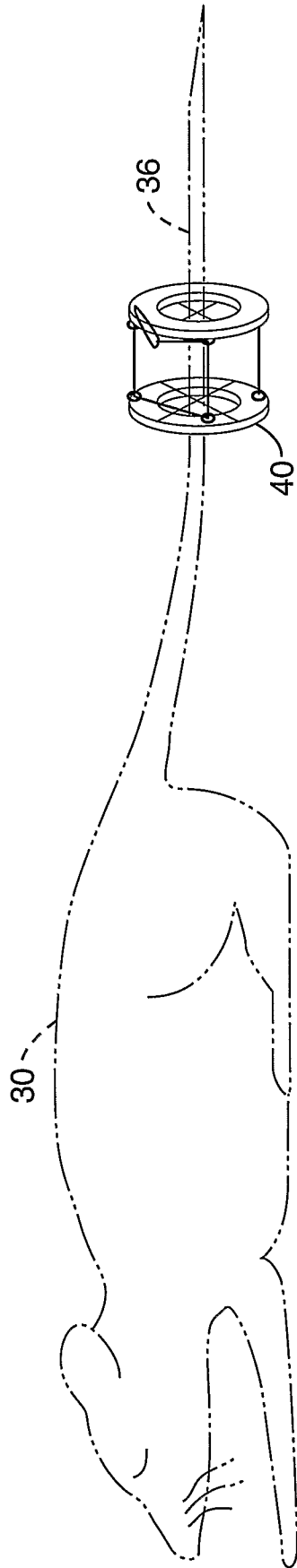


FIG. 3

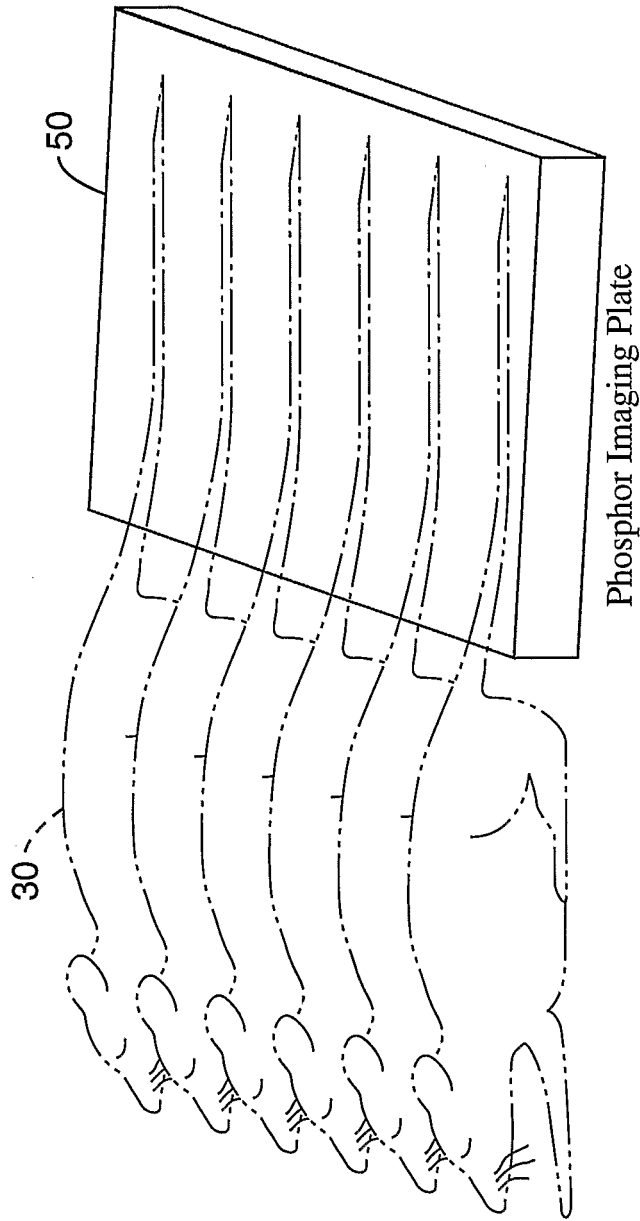
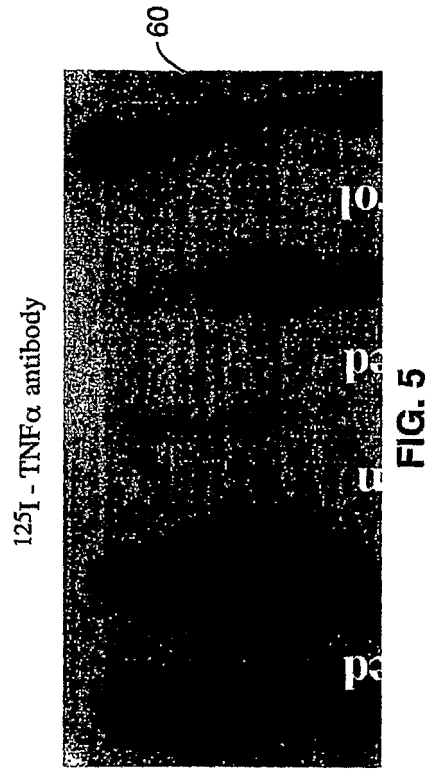


FIG. 4



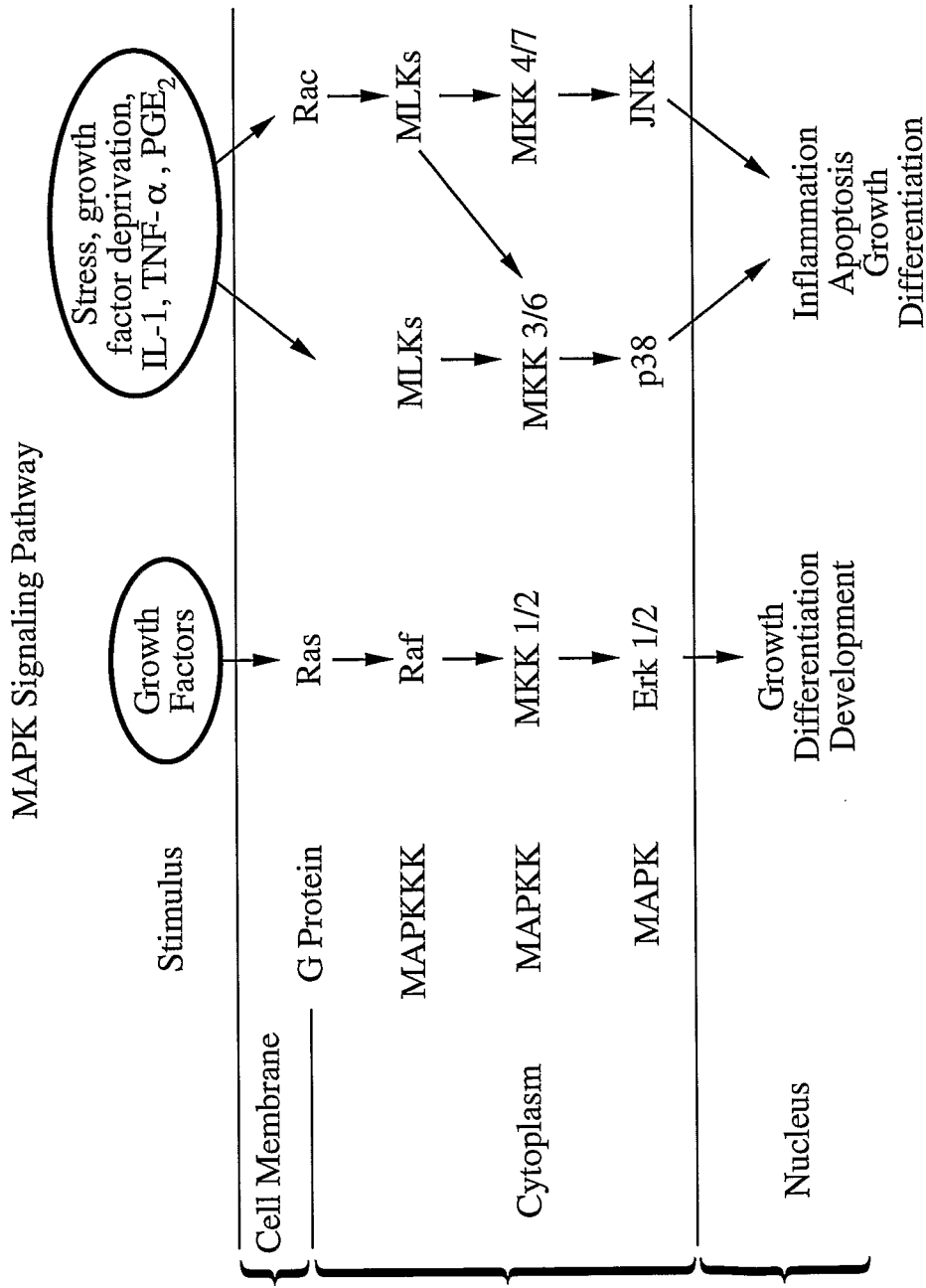


FIG. 6

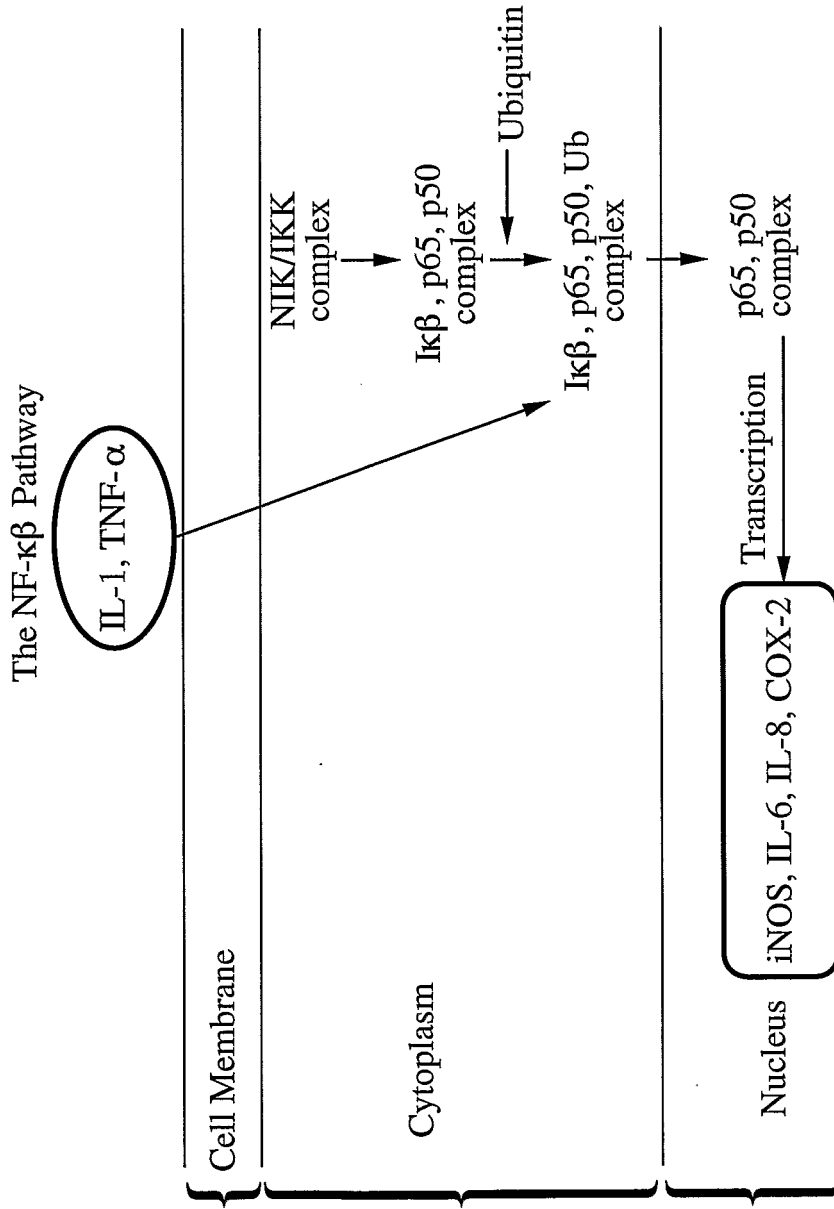


FIG. 7

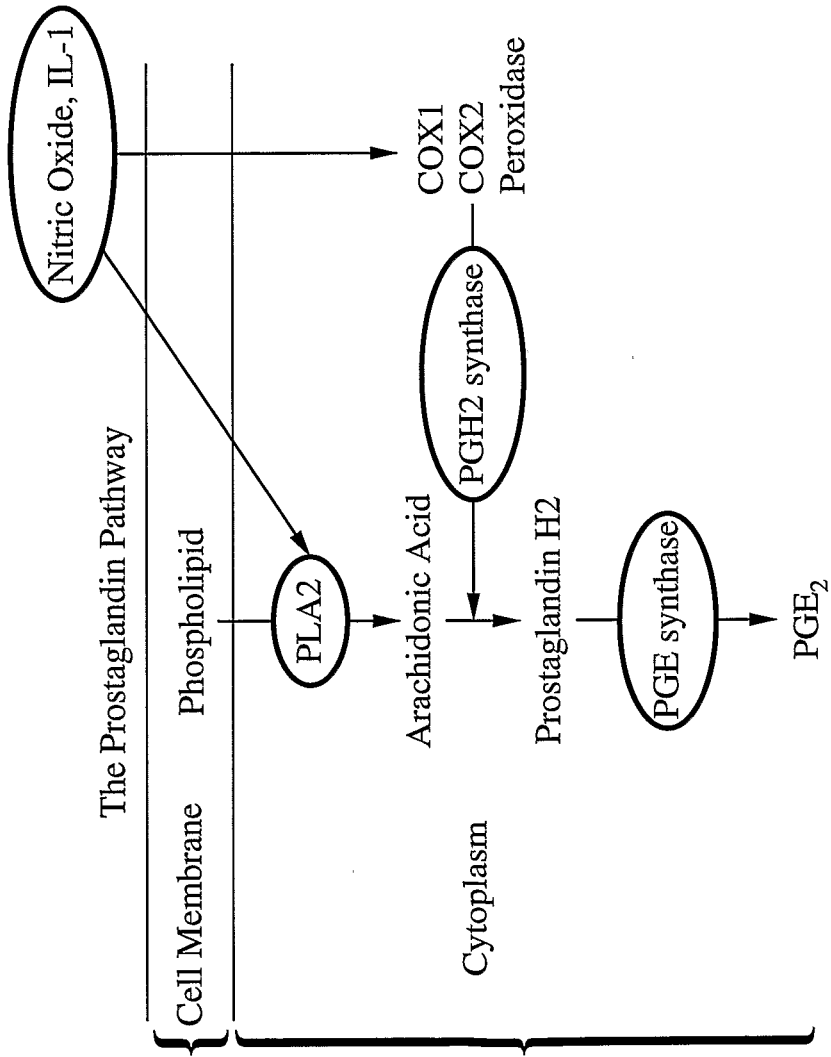


FIG. 8

REFERENCES CITED IN THE DESCRIPTION

This list of references cited by the applicant is for the reader's convenience only. It does not form part of the European patent document. Even though great care has been taken in compiling the references, errors or omissions cannot be excluded and the EPO disclaims all liability in this regard.

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专利名称(译)	局部显示和治疗疼痛的系统 and 组合物		
公开(公告)号	EP1933714B1	公开(公告)日	2020-03-18
申请号	EP2006815161	申请日	2006-09-21
[标]申请(专利权)人(译)	加利福尼亚大学董事会		
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当前申请(专利权)人(译)	加利福尼亚大学董事会		
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发明人	BRADFORD, DAVID, S. LOTZ, JEFFREY, C.		
IPC分类号	A61B5/00 A61K51/00		
CPC分类号	A61B5/4824 A61B6/506 A61K49/0002 A61N7/02 A61P19/00 A61P19/02 A61P21/00 A61P25/00 A61P29/00 A61B5/4041 A61B18/14 A61B2018/00339 A61B2018/00577 A61B2018/00642 A61K51 /1096 A61M5/007 A61M5/1723 A61M2230/005		
优先权	60/719670 2005-09-21 US 60/750990 2005-12-15 US		
其他公开文献	EP1933714A4 EP1933714A2		
外部链接	Espacenet		

摘要(译)

疼痛因子用传递到体内的靶向药物或标记物标记。用适当的成像工具以允许选择性识别和定位疼痛源或传播区域的方式对标记的疼痛因素进行成像。标记的疼痛因素允许成像中的空间差异足以指定疼痛的位置，从而驱动治疗决策和技术以治疗疼痛。以这种方式标记和成像的疼痛因子可以包括神经因子，血管因子，细胞因子和炎症因子中的一种或多种。标记的标记物可以包括例如放射性材料（例如三化或碘化的分子）或其他材料例如金属（例如金）纳米粒子。

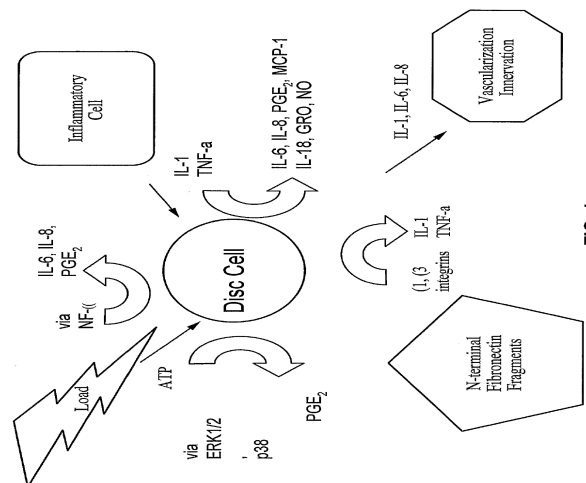


FIG. 1