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(71) Applicant (for all designated States except US): THE REGENTS OF THE UNIVERSITY OF CALIFORNIA [US/US]; Office of the President, 1111 Franklin Street, 12th Floor, Oakland, CA 94607-5200 (US).

(72) Inventors; and

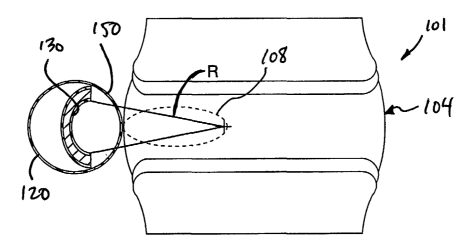
(75) Inventors/Applicants (for US only): DIEDERICH,
 Chris, J. [US/US]; 341 San Luis Way, Novato, CA 94945
 (US). LOTZ, Jeffrey, C. [US/US]; 129 Castillian Way,

San Mateo, CA 94402 (US). NAU, Will [US/US]; 8 Locksley Avenue, Apt. 9B, San Francisco, CA 94122 (US). BRADFORD, David, S. [US/US]; 9 Cloudview Road, Sausalito, CA 94965 (US). WISTROM, Elizabeth [US/US]; 1601 Dry Creek Road, Healdsburg, CA 95448 (US). BASS, Elisa [US/US]; 1727 9th Avenue, San Francisco, CA 94122 (US).

- (74) Agent: O'BANION, John, P.; O'Banion & Ritchey LLP, 400 Capitol Mall, Suite 1550, Sacramento, CA 95814 (US).
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(54) Title: SYSTEM AND METHOD PROVIDING DIRECTIONAL ULTRASOUND THERAPY TO SKELETAL JOINTS



(57) Abstract: An ultrasound therapy system and method is provided that provides directional, focused ultrasound to localized regions of tissue within body joints, such as spinal joints. An ultrasound emitter or transducer is delivered to a location within the body associated with the joint and heats the target region of tissue associated with the joint from the location. Such locations for ultrasound transducer placement may include for example in or around the intervertebral discs, or the bony structures such as vertebral bodies or posterior vertebral elements such as facet joints. Various modes of operation provide for selective, controlled heating at different temperature ranges to provide different intended results in the target tissue, which ranges are significantly effected by pre-stressed tissues such as in-vivo intervertebral discs. In particular, treatments above 70 degrees C, and in particular 75 degrees C, are used for structural remodeling, whereas lower temperatures achieves other responses without appreciable remodeling.



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SYSTEM AND METHOD PROVIDING DIRECTIONAL ULTRASOUND THERAPY TO SKELETAL JOINTS

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority from U.S. provisional application serial number 60/349,207 filed on January 15, 2002, and also from U.S. provisional application serial number 60/351,827 filed on January 23, 2002; and also U.S. provisional application serial number 60/410,603 filed on September 12, 2002; and also 60/411,401 filed on September 16, 2002. The disclosures of these U.S. provisional patent applications are herein incorporated in their entirety by reference thereto.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

Not Applicable

REFERENCE TO A COMPUTER PROGRAM APPENDIX Not Applicable

BACKGROUND OF THE INVENTION

1. Field of the Invention

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The present invention is a system and method for delivering therapeutic levels of energy to tissue in a living body. More specifically, it is a system and method for delivering therapeutic levels of ultrasound energy invasively within the body in order to treat disorders associated with the spine and other joints. Still more specifically, it is a system and method for delivering ultrasound energy to intervertebral discs in order to treat disorders associated therewith such as chronic lower back pain.

2. Description of the Background Art

For many years, much research and commercial development has been directed toward delivering energy to tissue in order to achieve various desired therapeutic results. Examples of energy modalities previously described for tissue treatment include: electrical current (both DC and AC, e.g. radiofrequency or "RF"

current), plasma ion energy, sonic energy (in particular ultrasound), light energy (e.g. laser, infrared or "IR", or ultraviolet or "UV"), microwave induction, and thermal energy (e.g. convection or conduction). Other modalities for treating tissue include without limitation: cryotherapy (cooling tissue to desired levels to affect structure of function), and chemical therapy (delivering chemicals to the tissue to affect the tissue structure of function). Each of these energy delivery and other treatment modalities has been extensively studied and characterized as providing unique benefits, as well as unique issues and concerns, with respect to tissue therapy. Accordingly, many specific energy delivery methods and systems have been disclosed to provide unique benefits for particular intended therapeutic applications.

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Various specific tissue responses to energy delivery have also been observed and reported during the course of significant study and characterization. In one regard, tissues or their function may be damaged by energy delivery such as thermal therapy. Examples of previously disclosed, differentiated effects of thermal tissue therapy generally characterized to damage tissue include, without limitation: ablation (which has been defined as either molecular dissociation or by achieving cellular necrosis), coagulation, degranulation, and desiccation. Alternatively, energy delivery in certain particular forms has also been characterized as promoting reproductive stimulation in certain tissues. Certain desired results have been disclosed with respect to intending controlled tissue damage with tissue thermal therapy; other desired results have been disclosed with respect to promoting tissue reproduction with tissue thermal therapy. In any event, because of the pronounced effects observed from tissue energy delivery, it is often desired to control and accurately select the localization of tissue/energy interaction in order to treat only the intended tissue, else normal surrounding tissue is effected with harmful results.

Accordingly, the different energy delivery modalities have been specifically characterized as providing particular benefits and problems versus other modalities with respect to various specific tissues and related medical conditions. Examples of specific medical conditions and related tissue that have been studied and characterized for tissue energy delivery include: tumors such as cancer (e.g. liver, prostate, etc.); vascular aneurysms, malformations, occlusions, and shunts; cardiac arrhythmias; eye disorders; epidermal scarring; wrinkling; and musculo-skeletal injury

repair. The nature of the condition to be treated, as well as the anatomy of the area, can have significant impact on the desired result of energy delivery, which directly differentiates between the appropriateness or inappropriateness of each of the different energy delivery modalities for such application (as well as the corresponding particular operating parameters, systems, and methods for delivering such energy).

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Depending upon the particular energy modality, various different parameters may be altered to affect the thermal effect in particular tissues, including which type of effect is achieved (e.g. ablation, coagulation, desiccation, etc.), as well as depth or degree of the effect in surrounding tissues. For the purpose of a general understanding, however, known tissue responses to thermal therapies, e.g. effect of changing temperatures to particular levels, have been previously characterized for certain tissues in prior disclosures which are summarized as follows.

As described above, temperature elevation of biological tissues is currently used for outright tissue destruction or to modify tissues to enhance other therapies. Low temperature elevations (41-45°C) of relatively short duration (<30-60 min) may damage cells but generally only to such extent to be repairable and considered nonlethal. In this range, it is believed that heat mediated physiological effects include heat induced acceleration of metabolism or cellular activity, thermal inactivation of enzymes, rupture of cell membranes, and delayed onset of increasing blood flow and vessel permeability. For temperature exposures in excess of 45°C and/or longer durations, it is believed that cellular repair mechanisms no longer function due to denaturation of key proteins or can't keep up with the accumulating damage. Complete cell death and tissue necrosis have been observed to be fully expressed in approximately 3-5 days. Temperature exposures in the 42-45°C regimen are commonly used for example as an adjuvant to radiation cancer therapy and chemotherapy, and have been considered for enhancing gene therapy and immunotherapy as well. Higher temperature elevations (50+°C) have been investigated for inducing desirable physical changes in tissue, ranging from applications such as controlled thermal coagulation for "tightening" ligaments and joint capsules, tissue reshaping, and selective tissue thermal coagulation for destroying cancerous and benign tumors. High temperature exposures (50+°C) are generally believed to produce rather lethal and immediate irreversible denaturation

and conformational changes in cellular and tissue structural proteins, thereby thermally coagulating the tissue.

In general, heat-induced cell damage or tissue structural changes described above are believed to be attributed to thermal denaturation and aggregation of key protein structures. The accumulation of this thermal damage can be modeled using the Arrhenius rate process equation, which establishes a relationship between rate of thermal damage and the duration and temperature of exposure:

$$\frac{1}{\tau} = A \bullet e^{-\Delta E/RT},$$

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where ΔE is activation energy (J mol⁻¹), R is the universal gas constant (8.32 J mol⁻¹ K⁻¹), A is the rate constant (s⁻¹), T is temperature in Kelvin, and $1/\tau$ is rate of thermal damage (s⁻¹). Using this expression (Eqn. 1), a relationship can be derived to determine an exposure time(τ_2) and/or temperature elevation (T₂) required to produce an equivalent observed biological effect associated with a specified temperature (T₁) and time exposure (τ_1). This is the basis of the thermal iso-effect equation as shown below, which is non-linear with respect to temperature exposure and linear with respect to exposure time:

(2)
$$\tau_2 = \tau_1 e^{(\Delta E/RT_1 T_2)(T_1 - T_2)} = \tau_1 K^{(T_1 - T_2)},$$

where the parameter K is approximated as constant for typical therapeutic temperature elevations (10-30°C). Furthermore, extensive *in vitro* and *in vivo* studies have demonstrated that ΔE for thermal damage is approximately constant at 140 J mol⁻¹ for temperatures greater than 43°C. Thus, the relationship between time and temperature for a given biological effect depends upon activation energy only. Thus, as determined from the hyperthermia biology literature, K≅2 for T ≥43°C and K≅4-6 for T<43°C. The different values split at approximately 43°C in order to model the biphasic behavior in the rate response, with faster damage accumulation after a break around 43°C. These values hold for lethal cellular damage, but not coagulation of structural proteins (collagen). Traditionally this iso-effect dose has been used to characterize hyperthermia cancer treatments with a target temperature elevation of 42.5-45°C, and has led to 43 °C becoming the historical reference dose temperature. This forms the basis of the thermal iso-effect dose (TID) equation,

which as shown below can be used to calculate thermal dose of a varying temperature exposure over time as an equivalent exposure duration at 43°C (or other reference temperature). Temperature time history is equated to a thermal dose at a known temperature reference.

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$$EM_{43} = \int_{0}^{t_f} K^{(T-43)} dt = \sum_{t=0}^{t_{final}} \Delta t K^{(T-43)},$$

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where dt is a time step (min) and EM ₄₃ is thermal dose expressed in equivalent minutes at 43 °C.

Various previously published disclosures have verified the Arrhenius model and the iso-effect relationship of different temperature-time exposures for generating trans-epidermal thermal necrosis in skin. Applying the TID analysis, a threshold of approximately 320 EM₄₃ (wherein "EM" represents "equivalent minutes" at the given temperature shown in subscript) as found for temperatures between 44-60 °C. Thermal dosages between 10-100 EM₄₃ have been shown to correlate with improved response to hyperthermia and radiation therapy. For a conservative approach 250 EM _{43°C} is a threshold dose for complete thermal necrosis, where reported values range from 25-240 EM43°C for brain and muscle tissue, respectively.

In addition, thermal coagulation or coagulation necrosis will occur in tissues exposed to temperatures greater than approximately 55°C for a duration of minutes in collagen in particular. Thermal coagulation of soft tissues requires temperatures in excess of 50 °C. Numerous investigators have validated the "TID" (or "temperature iso-dose") concept for predicting lesions using 240-340 EM 43°C as a threshold dose and critical temperatures of 53-54 °C for coagulating muscle.

Therapeutic Energy Delivery for Spinal Disorders

Spinal disorders have been the topic of significant study and commercial development for therapeutic energy delivery. In particular, various specific conditions that have been studied with respect to particular modes of therapeutic energy delivery.

Of particular interest has been chronic lower back pain. Chronic low back pain is a significant health and economic problem in the United States, being the most costly form of disability in the industrial setting. For a substantial number of

these patients the intervertebral disc is considered the principal pain generator. Traditionally, patients who fail conservative therapy have few treatment options beside discectomy or fusion, either of which can result in significant morbidity and variable outcomes. Recent efforts have been directed toward investigating thermal therapy for providing a healing effect on collagenous tissues, and therefore this modality has been incorporated into several minimally invasive back pain treatments.

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Early orthopedic use of high temperature heat therapy was to manage shoulder instability. In this application, the shoulder capsule is treated with laser or radio-frequency (RF) thermal energy to temperatures typically in the range of 70 to 80°C. This treatment has been disclosed to stabilize the joint by inducing tissue contraction. Such treatment also has been disclosed to result in an acute decrease in stiffness (e.g. as much as 50%) that may be recovered due to biologic remodeling. However, the long-term benefits of this treatment have been questioned since the collagenous tissue may re-lengthen over time.

The contraction associated with thermal therapy, which can reach as high as 50% along the fiber direction in the shoulder capsule, has been correlated with thermal denaturation. Thermal denaturation is an endothermic process in which the collagen triple helix unwinds after a critical activation energy is reached. Differential scanning calorimetry (DSC) is a technique to measure both the denaturation temperature (T_m - the peak temperature corresponding to this critical activation energy) and the total enthalpy of denaturation (ΔH - the total energy required to fully denature the collagen). This technique can be used to correlate thermal exposure with the resulting degree of denaturation for a specific collagenous tissue, and thus to guide the development of an optimal thermal dose.

Intradiscal electrothermal therapy (IDET) has been recently introduced as a minimally invasive, non-operative therapy in which a temperature elevation is applied in order to treat discogenic low back pain. In this procedure, a catheter containing a 5 cm long resistive-wire heating coil is introduced percutaneously into the disc under fluoroscopic guidance. The internal temperature of the device is then raised from 65°C to 90°C over a course of 16 minutes. This procedure is thought to produce temperatures sufficient to contract annular collagen and ablate annular nociceptors. A controlled, 12 month trial of IDET on a relatively small patient population (36

individuals) demonstrated some relief of back pain in 60% of patients and total relief in 23%. A two-year follow-up study of 58 patients found clinically significant improvement in pain, physical function, and quality of life. While these results have been considered by some to be promising, prospective placebo-controlled trials are lacking, and the therapeutic mechanisms of thermal therapy are unknown. Proposed therapeutic mechanisms of such technique have included: 1) collagen denaturation, causing annular stiffening, and tissue remodeling; 2) annular de-innervation; and 3) ablation of cytokine-producing cells. Due to mechanistic uncertainty, treatment optimization and patient selection are generally empirically based.

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The effect of heat on collagen denaturation and biomechanical properties has been investigated in various tissues: knee and shoulder capsule, tendon, and chordae tendineae. In general, at least one prior disclosure reports that significant denaturation and shrinkage occurred in tissue treated at 65°C and above for 1-5 minutes. However, given that the annular architecture of intervertebral discs is quite different from these other tissues it is has not been previously made clear that prior results can be directly extrapolated to the intervertebral disc.

Further more detailed background information related to various aspects of thermal tissue therapy and/or chronic back pain is variously disclosed in the following publications: Amonoo-Kuofi, H. S., 1991, "Morphometric changes in the heights and anteroposterior diameters of the lumbar intervertebral discs with age." J Anat 159-68; Arnoczky, S. P. and Aksan, A., 2001, "Thermal modification of connective tissues: basic science considerations and clinical implications." Instr Course Lect 3-11; Chen, S. S., Wright, N. T. and Humphrey, J. D., 1997, "Heat-induced changes in the mechanics of a collagenous tissue: isothermal free shrinkage." J Biomech Eng 4, 372-8; Chen, S. S., Wright, N. T. and Humphrey, J. D., 1998, "Heat-induced changes in the mechanics of a collagenous tissue: isothermal, isotonic shrinkage." J Biomech Eng 3, 382-8; Dewey, W. C., 1994, "Arrhenius relationships from the molecule and cell to the clinic." Int J Hyperthermia 4, 457-83; Flandin, F., Buffevant, C. and Herbage, D., 1984, "A differential scanning calorimetry analysis of the age-related changes in the thermal stability of rat skin collagen." Biochim Biophys Acta 2, 205-11; Gerber, A. and Warner, J. J., 2002, "Thermal capsulorrhaphy to treat shoulder instability." Clin Orthop 400, 105-16; Hall, B. K., 1986, "The role of movement and

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tissue interactions in the development and growth of bone and secondary cartilage in the clavicle of the embryonic chick." J Embryol Exp Morphol 133-52; Hayashi, K. and Markel, M. D., 2001, "Thermal capsulorrhaphy treatment of shoulder instability: basic science." Clin Orthop 390, 59-72; Hayashi, K., et al., 2000, "The mechanism of joint capsule thermal modification in an in-vitro sheep model." Clin Orthop 370, 236-49; Hayashi, K., et al., 1997, "The effect of thermal heating on the length and histologic properties of the glenohumeral joint capsule." Am J Sports Med 1, 107-12; Heary, R. F., 2001, "Intradiscal electrothermal annuloplasty: the IDET procedure." J Spinal Disord 4, 353-60; Hecht, P., et al., 1999, "Monopolar radiofrequency energy effects on joint capsular tissue: potential treatment for joint instability. An in vivo mechanical, morphological, and biochemical study using an ovine model." Am J Sports Med 6, 761-71; Karasek, M. and Bogduk, N., 2000, "Twelve-month follow-up of a controlled trial of intradiscal thermal anuloplasty for back pain due to internal disc disruption." Spine 20, 2601-7; Kronick, P., et al., 1988, "The locations of collagens with different thermal stabilities in fibrils of bovine reticular dermis." Connect Tissue Res 2, 123-34; Le Lous, M., et al. 1982. "Influence of collagen denaturation on the chemorheological properties of skin, assessed by differential scanning calorimetry and hydrothermal isometric tension measurement." Biochim Biophys Acta 2, 295-300; Lopez, M. J., et al., 2000, "Effects of monopolar radiofrequency energy on ovine joint capsular mechanical properties." Clin Orthop 374, 286-97; Miles, C. A. and Ghelashvili, M. 1999, "Polymer-in-a-box mechanism for the thermal stabilization of collagen molecules in fibers." Biophys J 6, 3243-52; Naseef, G. S., et al., 1997, "The thermal properties of bovine joint capsule. The basic science of laser- and radiofrequencyinduced capsular shrinkage." Am J Sports Med 5, 670-4; Nieminen, M. T., et al., 2000, "Quantitative MR microscopy of enzymatically degraded articular cartilage." Magn Reson Med 5, 676-81; (Nrc/lm), N. R. C. A. I. O. M. (2001). "Musculoskeletal Disorders and the workplace: low back and upper extremities." Washington D.C., National Academy Press; Saal, J. A. and Saal, J. S., 2002, "Intradiscal electrothermal treatment for chronic discogenic low back pain: prospective outcome study with a minimum 2-year follow-up." Spine 9, 966-73; discussion 973-4; Saal, J. S. and Saal, J. A., 2000, "Management of chronic discogenic low back pain with a

thermal intradiscal catheter. A preliminary report." Spine 3, 382-8; Schachar, R. A., 1991, "Radial thermokeratoplasty. Int Ophthalmol" Clin 1, 47-57; Schaefer, S. L., et al., 1997, "Tissue shrinkage with the holmium:yttrium aluminum garnet laser. A postoperative assessment of tissue length, stiffness, and structure." Am J Sports Med 6, 841-8; Schwarzer, A. C., et al., 1995, "The prevalence and clinical features of internal disc disruption in patients with chronic low back pain." Spine 17, 1878-83; Urban, J. P. and Mcmullin, J. F., 1985, "Swelling pressure of the intervertebral disc: influence of proteoglycan and collagen contents." Biorheology 2, 145-57; Vangsness, C. T., Jr., et al., 1997, "Collagen shortening. An experimental approach with heat." Clin Orthop 337, 267-71; Vujaskovic, Z., et al., 1994, "Effects of intraoperative hyperthermia on peripheral nerves: neurological and electrophysiological studies." Int J Hyperthermia 1, 41-9; Wallace, A. L., et al., 2000, "The scientific basis of thermal capsular shrinkage." J Shoulder Elbow Surg 4, 354-60; and Wallace, A. L., et al., 2002, "Creep behavior of a rabbit model of ligament laxity after electrothermal shrinkage in vivo. Am J Sports Med 1, 98-102. The disclosures of these references are herein incorporated in their entirety by reference thereto.

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Chronic lower back pain (e.g. discogenic lumbar pain) and related motor nerve deficit is typically due to damaged or herniated vertebral discs which either directly impinge on surrounding nerves or cause irritating inflammation. Traditional treatment options include surgery, anti-inflammatory drugs, physical therapy, etc., with surgery typically the last option. Due to difficulty of surgical procedure, complications and time of recovery, alternative procedures have been investigated. Recently, intradiscal thermal therapy has been introduced as a minimally-invasive alternative in the treatment of various spinal disorders such as chronic low back pain and otherwise disorders related to intervertebral disc abnormalities.

In particular, several different systems and methods have been disclosed for treating various abnormal conditions associated with intervertebral discs specifically by delivering electrical current in the RF range during invasive treatment procedures in and around the disc within the body. Other previously disclosed examples intended to invasively deliver therapeutic levels of energy in order to treat various spinal disorders include delivery of plasma ion energy (e.g. CoblationR from

Arthrocare, Inc.), laser light energy, or thermal energy from conductive heating elements (e.g. SpineCATH IDEC procedure, commercially available from Oratec Interventions). At least one other prior disclosure is intended to deliver heated thermoplastic material to allow it to flow into and then set upon cooling within the nucleus of an intervertebral disc in order to replace the nucleus pulposus.

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Further more detailed examples of energy delivery systems and methods such as of the types just described, that are intended to provide invasive therapy to treat various conditions associated with intervertebral disc disorders are variously disclosed in the following issued U.S. Patents: 4,959,063 to Kojima; 6,264,650 to Hovda *et al.*; 6,264,659 to Ross *et al.*. Examples are also disclosed in the following published U.S. Patent Application: US 2001/0029370 to Hodva *et al.*. Other examples are disclosed in the following published international patent applications: WO 00/49978 to Guagliano *et al.*; WO 00/71043 to Hovda *et al.*; WO 01/26570 to Alleyne *et al.*;. Additional disclosure is provided in the following published reference: Diederich CJ, Nau WH, Kleinstueck F, Lotz J, Bradford D (2001) "IDTT Therapy in Cadaveric Lumbar Spine: Temperature and thermal dose distributions, Thermal Treatment of Tissue: Energy Delivery and Assessment," Thomas P. Ryan, Editor, Proceedings of SPIE Vol. 4247:104-108. The disclosures of all these references provided in this paragraph are herein incorporated in their entirety by reference thereto.

<u>Ultrasound Energy Delivery Systems and Methods</u>

Ultrasound energy delivery and the effects of such energy on various different tissue structures has been the topic of significant recent study. The particular benefits of ultrasound delivery have been substantially well characterized, in particular with respect to different types of tissues as well as different ultrasound energy deposition modes. Many different medical device systems and methods have been disclosed for delivering therapeutic levels of ultrasound to tissues to treat wide varieties of disorders, including for example arterial blockages, cardiac arrhythmias, and cancerous tumors. Such disclosures generally intend to "ablate" targeted tissues in order to achieve a desired result associated with such particular conditions, wherein the desired response in the particular tissues, and the ultrasound delivery systems and methods of operation necessary for the corresponding energy

deposition modalities, may vary substantially between specific such "ultrasound ablation" systems and methods.

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Further more detailed examples of ultrasound delivery systems and methods such as of the type just described are disclosed in the following issued U.S. Patents which are incorporated herein by reference: 5,295,484 to Marcus *et al.*; 5,620,479 to Diederich; 5,630,837 to Crowley; 5,733,280 to Sherman *et al.*; 6,012,457 to Lesh; 6,024,740 to Lesh *et al.*; 6,117,101 to Diederich *et al.*; 6,164,283 to Lesh; 6,245,064 to Lesh *et al.*; 6,254,599 to Lesh *et al.*; and 6,305,378 to Lesh *et al.*. Other examples are disclosed in the following published foreign patent applications which are incorporated herein by reference: WO 00/56237 to Maguire *et al.*; WO 00/67648 to Maguire *et al.*; WO 00/67656 to Maguire *et al.*; WO 99/44519 to Langberg *et al.*

In addition, ultrasound enhanced drug delivery into tissues, e.g. to increase dispersion, permeability, or cellular uptake of therapeutic compounds such as drugs, has been well characterized and disclosed in many different specific forms.

Further more detailed examples of ultrasound energy delivery systems and methods such as those just described are disclosed in the following U.S. Patent References: 5,725,494 to Brisken; 5,728,062 to Brisken; 5,735,811 to Brisken; 5,846,218 to Brisken *et al.*; 5,931,805 to Brisken; 5,997,497 to Nita *et al.*; 6,210,393 to Brisken; 6,221,038 to Brisken; 6,228,046 to Brisken; 6,287,272 to Brisken *et al.*; and 6,296,619 to Brisken *et al.* The disclosures of these references are herein incorporated in their entirety by reference thereto.

Additional previously disclosed examples for ultrasound energy delivery systems and methods are intended to treat disorders associated with the spine in general, and in some regards of the intervertebral disc in particular. However, these disclosed systems are generally adapted to treat such disorders chronically from outside of the body, such as for example via transducers coupled to a brace worn externally by a patient. Therefore locally densified US energy is not achieved selectively within the tissues associated with such disorders invasively within the body. At least one further disclosure, however, proposes delivering focused ultrasound energy from outside the body for the intended purpose of treating intervertebral disc disorders, in particular with respect to degenerating the nucleus

pulposus to reduce the pressure within the disc and thus onto the adjacent spinal cord. However, the ability to actually achieve such targeted energy delivery at highly localized tissue regions associated with such discs, and to accurately control tissue temperature to achieve desired results, without substantially affecting surrounding tissues has not been yet confirmed or taught.

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Further more detailed examples of such systems and methods intended to treat spinal disorders with ultrasound energy from outside of the body are variously disclosed in the following issued U.S. Patents: 5,762,616 to Talish; 6,254,553 to Lidgren *et al.*. Other examples are disclosed in the following published international patent applications: WO 97/33649 to Talish; WO 99/19025 to Urgovich *et al.*; and WO 99/48621 to Cornejo *et al.* The disclosures of these references are herein incorporated in their entirety by reference thereto.

There is still a need for improved systems and methods for locally delivering therapeutic amounts of ultrasound energy invasively within the body in order to treat disorders associated with the spine and other joints.

There is in particular still a need for a system and method adapted to locally deliver ultrasound to a highly localized region of tissue, such as only a portion of a disc associated with

BRIEF SUMMARY OF THE INVENTION

An object of the invention is to deliver therapeutic levels of ultrasound energy to intervertebral discs in order to treat disorders associated therewith.

Another object of the invention is to provide a kit of energy delivery devices with varied shapes along the energy delivery portion thereof in order to specifically treat different regions of intervertebral discs having varied geometries from within the nucleus.

Another object of the invention is to deliver therapeutic levels of energy sufficient to cause necrosis of particular cellular structures associated with an intervertebral disc without substantially remodeling or affecting the structure integrity of the annulus fibrosus of the disc.

Another object of the invention is to provide thermal therapy to a region of tissue associated with a joint in the body without substantially remodeling structural support tissues associated with the joint.

Another object of the invention is to treat inflammation and pain associated with disorders of the spine in general and intervertebral discs in particular.

Another object of the invention is to denervate or necrose nociceptive fibers or cells in certain regions of tissue associated with an intervertebral disc.

Another object of the invention is to reduce inflammation associated with damaged intervertebral discs.

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Another object of the invention is to repair damaged regions of intervertebral discs.

A nother object of the invention is to achieve cellular necrosis of certain particular tissues associated with an intervertebral disc disorder without substantially altering the structure of the annulus fibrosus of the respective disc.

Another object of the invention is to remodel cartilaginous tissue associated with spinal joints, and in particular intervertebral discs.

Another object of the invention is to provide sufficient thermal therapy to a region of stressed tissue to cause a remodeling of the tissue.

Another object of the invention is to provide sufficient thermal therapy to a region of an intact mammalian intervertebral disc to cause a remodeling of at least one support structure.

Another object of the invention is to provide a thermal therapy device that can substantially heat deep regions of tissue from the site of therapy, such as with respect to spinal joints at least about 4 mm, 7 mm, or even 10 mm from the site of therapy.

Another object of the invention is to provide therapeutic levels of thermal therapy to tissue within relatively short periods of time.

Another object of the invention is to direct therapeutic energy into targeted tissues from remote locations within the body, such as in or around joints, without substantially harming closely adjacent tissues, such as nerves, vessels, or other tissues not intended to be treated.

Another object of the invention is to focus energy into targeted regions of tissue within the body.

Another object of the invention is to enhance cellular functions in certain tissue structures so as to provide a therapeutic effect.

Another object of the invention is to enhance drug delivery into remotely located body tissues, such as within or around joints such as spinal joints.

Another object of the invention is to treat back pain.

Another object of the invention is to treat arthritis.

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Another object of the invention is to locally enhance the delivery, permeability, or cellular uptake related to certain therapeutic compounds delivered within the spine and other joints.

Accordingly, one aspect of the invention is a spinal thermal therapy system with a spinal delivery system and a spinal thermal therapy device. The spinal delivery system is adapted to deliver a thermal therapy assembly of the spinal thermal therapy device to a location within the body such that the energy may be coupled to a region of tissue associated with a vertebral joint.

One mode of this aspect, the thermal therapy assembly is adapted to deliver sufficient energy to the region of tissue to heat the region to a temperature less than about 65 degrees C and to deliver a thermal dose of at least about 240 EM43.

According to one embodiment of this mode, the assembly is adapted to heat the region sufficiently to form cellular necrosis, but not to cause substantial denaturation of collagen, and in particular with respect to regions of tissue under mechanical stress.

Another mode of this aspect, the thermal therapy assembly is adapted to heat the region of tissue to a temperature of greater than about 65 degrees C.

According to one further embodiment of this mode, the thermal therapy assembly is adapted to heat the region of tissue to a temperature of greater than about 70 degrees C.

According to a further feature of this embodiment, the thermal therapy assembly is adapted to heat the region of tissue to a temperature of greater than a bout 75 degrees C.

According to another embodiment, the thermal therapy assembly that is adapted to heat the region of tissue to a temperature that is greater than about 65 degrees C but is less than about 100 degrees C.

In a further variation of this embodiment, the thermal therapy assembly is adapted to heat the region to a temperature between about 65 degrees C and about

85 degrees C.

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Another aspect of the invention is an ultrasound energy delivery system for treating a region of tissue associated with a skeletal joint, and includes an ultrasound treatment assembly with an ultrasound transducer; and a skeletal joint delivery assembly. The skeletal joint delivery assembly is adapted to deliver the ultrasound treatment assembly into the body with the ultrasound transducer positioned at a location within the body associated with the skeletal joint. The ultrasound treatment assembly is adapted to deliver a therapeutic level of ultrasound energy from the location and into the region of tissue.

According to one mode of this aspect, the ultrasound transducer is constructed to provide directional thermal therapy to the region of tissue. According to another mode, the ultrasound transducer is constructed to deliver a converging field of energy to the region of tissue. In another mode, the ultrasound transducer is constructed to deliver a diverging field of ultrasound energy to the region of tissue. In another mode, the ultrasound transducer is constructed to deliver a substantially collimated field of ultrasound energy to the region of tissue. In still another mode, the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue. Further embodiments for the coupling member it is a balloon that may be either elastomeric or pre-formed and noncompliant.

According to further modes, a temperature control system is provided that is adapted to control the temperature of at least one of the ultrasound transducer or a tissue interface between a region of tissue and the ultrasound treatment assembly during ultrasound energy delivery to the region of tissue.

Another mode further includes an ultrasound control system with a temperature monitoring system, a controller, and an ultrasound drive system. The ultrasound treatment assembly is adapted to couple to the ultrasound control system. The ultrasound control system is adapted to control the ultrasound energy being emitted by the ultrasound transducer as a function of at least one parameter related to thermal therapy in the region of tissue.

In a further embodiment, the parameter comprises a parameter related to the effects of ultrasound delivery from the ultrasound transducer in tissue, such as

thermal dose, tissue temperature, depth of thermal penetration, or rates of change thereof. Such parameter may be empirically based, or a value that is monitored during ultrasound therapy, and/or calculated based upon another measured parameter.

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In other embodiment the controller is adapted to control the operation of the ultrasound transducer such that the temperature in at least a portion of the tissue within at least a 4 mm depth from the ultrasound transducer is elevated to at least about 70 degrees C. In another embodiment, the controller is adapted to control the operation of the ultrasound transducer such that the temperature of the tissue within the 4mm depth is elevated to at least about 75 degrees C. In another embodiment, the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up to at least about a 7 mm depth from the ultrasound transducer is elevated to at least about 70 degrees C. In still a further embodiment, the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up to at least a 10mm depth from the ultrasound transducer is elevated to greater than about 45 degrees C.

In still a further embodiment, the controller is adapted to control the operation of the ultrasound transducer such that the temperature within the tissue are limited to not exceed certain values. In one such embodiment the control is set for tissue at up to at least a 4 mm depth from the ultrasound transducer not to exceed an elevated temperature of about 75 degrees C or less. In another embodiment, the controller is adapted to control the operation of the ultrasound transducer such that the temperature of the tissue within the 4mm depth is elevated to an elevated temperature of no more than about 70 degrees C or less. In another embodiment the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up to at least a 7 mm depth from the ultrasound transducer is elevated to at least 70 degrees C.

Further modes provide the treatment assembly in a manner adapted to provide certain ideal temperature delivery objectives. In one such mode, the device is adapted to heat tissue within up to at least a 10mm depth from the ultrasound transducer is elevated to greater than 45 degrees C. In another embodiment, the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least

about 4 mm from the ultrasound transducer to a temperature of at least about 70 degrees C.

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In another embodiment, the ultrasound heating assembly is adapted to heat the tissue within the 4mm depth to a temperature of at least about 75 degrees C. In still another embodiment, the ultrasound treatment assembly is adapted to heat the tissue at least about 4 mm away from the transducer to the temperature of at least about 75 degrees C in less than about 5 minutes of ultrasound energy delivery into the tissue. In yet a further embodiment, the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least about 7 mm from the ultrasound transducer to a temperature of at least about 70 degrees C. In further embodiments, the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least about 10mm from the ultrasound transducer to a temperature of at least about 45 degrees C. In another embodiment, the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least 4 mm from the ultrasound transducer to a temperature of no more than about 75 degrees C or less.

According to another mode, the ultrasound treatment assembly is adapted to be positioned within an intervertebral disc and to heat at least portion of the disc.

In another mode, the ultrasound treatment assembly is adapted to be positioned at a location adjacent to an intervertebral disc and to provide thermal therapy to the intervertebral disc from the location.

In another mode, the ultrasound treatment assembly is adapted to positioned within at least a portion of a vertebral body, or posterior vertebral element such as facet joints, and to provide thermal therapy at least in part to such structure.

In another mode, it is adapted to be positioned adjacent to such spinal joint bony structures and to heat at least a portion of the vertebral body.

Another aspect of the invention is a skeletal joint thermal therapy device with a thermal treatment assembly on the distal end of a delivery member. The thermal treatment assembly includes an energy emitter. The distal end portion is adapted at least in part to deliver the thermal treatment assembly into a body of an animal with the energy emitter positioned at a location such that the energy emitter is adapted to deliver energy into a region of tissue associated with a skeletal joint. The thermal treatment assembly is adapted to heat tissue up to a distance of at least 4mm from

the energy emitter to a temperature of at least 75 degrees C, and is adapted to heat tissue up to a distance of at least about 7 mm from the energy emitter to a temperature of at least about 55 degrees C, and is adapted to heat tissue up to a distance of at least about 10 mm from the energy emitter to a temperature of at least about 45 degrees C.

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In one further mode of this aspect, the thermal treatment assembly is adapted to heat tissue up to a distance of at least 4 mm from the energy emitter to a temperature of at least 80 degrees C. In another mode, the thermal treatment assembly is adapted to heat tissue up to a distance of at least 4 mm from the energy emitter to a temperature of at least 85 degrees C. In still another mode, the thermal treatment assembly is adapted to heat tissue up to a distance of at least 7 mm from the energy emitter to a temperature of at least 60 degrees C. In yet another mode, the thermal treatment assembly is adapted to heat tissue up to a distance of at least 7 mm from the energy emitter to a temperature of at least 65 degrees C. A further modes provides the thermal treatment assembly adapted to heat tissue up to a distance of at least 7 mm from the energy emitter to a temperature of at least 70 degrees C. Still further, the thermal treatment assembly may be further adapted to heat tissue up to a distance of at least 10 mm from the energy emitter to a temperature of at least 50 degrees C. It may also be adapted to heat tissue up to a distance of at least 10 mm from the energy emitter to a temperature of at least 55 degrees C. Even still further, it may be adapted to heat tissue up to a distance of at least 10 mm from the energy emitter to a temperature of at least 60 degrees C.

In yet another mode, the thermal treatment assembly is constructed to provide directional thermal therapy to the region of tissue.

In another highly beneficial mode, the energy emitter comprises an ultrasound transducer.

In another mode, the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.

Another aspect of the invention is an ultrasound thermal treatment system with an ultrasound treatment assembly with an ultrasound transducer on a distal end portion of a rigid delivery probe. The probe's distal end portion has a proximal

section with a longitudinal axis, a distal section with a distal tip, and a bend between the proximal and distal section. The ultrasound treatment assembly is located along the distal section of the distal end portion and extending at an angle from the proximal section. The distal end portion is adapted to be delivered into the body of an animal by manipulating the proximal end portion externally of the body and such that the ultrasound treatment assembly is positioned at a location associated with a region of tissue to be treated. Moreover, the ultrasound treatment assembly is adapted to deliver a therapeutic level of ultrasound energy into the region of tissue from the location within the body.

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In one further mode of this aspect, the proximal end portion of the rigid delivery probe comprises a metal tube.

In another mode, the distal end portion of the rigid delivery probe comprises a metal tube.

In another mode, the ultrasound transducer is constructed to provide directional ultrasound delivery to the region of tissue.

In another mode, the ultrasound transducer is constructed to deliver a converging field of ultrasound energy to the region of tissue.

Another aspect of the invention is an ultrasound thermal therapy system with an ultrasound heating assembly with an ultrasound transducer and a therapy control system coupled to the ultrasound heating assembly. The ultrasound heating assembly is adapted to be delivered into a body of an animal with the ultrasound transducer positioned at a location such that the ultrasound transducer is adapted to deliver a therapeutic amount of ultrasound energy into a targeted region of tissue in the body from the location. The therapy control system is adapted to control operation of the ultrasound heating assembly such that a substantial portion of the region of tissue being heated by the ultrasound heating assembly does not exceed a maximum temperature of at least about 70 degrees C.

According to one further mode of this aspect, the therapy control system is adapted to control operation of the ultrasound heating assembly such that a substantial portion of the region of tissue being heated by the ultrasound heating assembly does not exceed a maximum temperature of at least about 75 degrees C.

Another aspect of the invention is a directional ultrasound spinal thermal

therapy system with an ultrasound delivery assembly. The ultrasound delivery assembly has a directional ultrasound transducer that is adapted to be positioned at a location associated with a spinal joint and to deliver a directed, therapeutic amount of ultrasound energy from the location and to a region of tissue associated with the spinal joint.

In one mode of this aspect, the ultrasound delivery assembly is adapted to direct the ultrasound delivery into substantially only a particular radial zone around the circumference of the distal end portion of the support member.

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In another mode, the ultrasound delivery assembly is constructed to provide directional thermal therapy to the region of tissue.

In another mode, the ultrasound delivery assembly is constructed to deliver a converging field of energy to the region of tissue.

In another mode, the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.

In another mode, the ultrasound delivery assembly is adapted to be positioned at the location within at least a portion of an intervertebral disc associated with the spinal joint and to deliver the therapeutic ultrasound energy into the region of tissue from that location.

In another mode, the ultrasound delivery assembly is adapted to be positioned at the location outside of an intervertebral disc associated with the spinal joint and to deliver the therapeutic level of ultrasound energy into the intervertebral disc from that location.

In another mode, the ultrasound delivery assembly is adapted to be positioned at the location within at least a portion of a vertebral body and/or posterior vertebral elements such as facet joints associated with the spinal joint and to be ultrasonically coupled with the region of tissue from that location.

In another mode, the ultrasound delivery assembly is adapted to be positioned at the location outside of a vertebral body and/or posterior vertebral elements such as facet joints associated with the spinal joint and to deliver the therapeutic level of ultrasound energy into the intervertebral disc from that location.

Another aspect of the invention is a skeletal joint ultrasound delivery system

with an ultrasound treatment assembly with an ultrasound transducer and a coupling member. The ultrasound delivery assembly is adapted to be delivered into a body of a mammal with the ultrasound transducer positioned at a location within the body associated with a skeletal joint. The ultrasound transducer is adapted to deliver a therapeutic amount of ultrasound energy to a region of tissue associated with the skeletal joint via the coupling member.

In one further mode, the ultrasound delivery assembly is constructed to provide directional ultrasound delivery to the region of tissue.

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In another further mode, the ultrasound delivery assembly is constructed to deliver a converging field of energy to the region of tissue.

In another mode, the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.

Another aspect of the invention is an ultrasound thermal therapy system with an ultrasound heating assembly and a control system. The ultrasound heating assembly is adapted to be positioned at a location within a body of a mammal so as to deliver ultrasound energy into a region of tissue in the body from the location. The control system is adapted to couple to the ultrasound heating assembly and to control operation of the ultrasound heating assembly such that a region of tissue being heated by the ultrasound heating assembly exceeds a temperature of at least about 70 degrees C.

Another aspect of the invention is an ultrasound thermal therapy system with a an ultrasound heating assembly located along the distal end portion of a delivery member. The ultrasound heating assembly has a curvilinear ultrasound transducer having a concave surface with a radius of curvature around a reference axis that is transverse to the longitudinal axis of the distal end portion.

The invention according to another mode is a method for invasively treating a medical condition associated with a skeletal joint within a body of an animal. This method includes delivering a therapeutic level of ultrasound energy to a region of tissue associated with the joint from a location within the body of the patient.

Another aspect of the invention is a method for treating a medical condition associated with a skeletal joint within a body by delivering sufficient energy to a

region of tissue associated with the skeletal joint that is sufficient to necrose nociceptive nerve fibers or inflammatory cells in such tissue region without substantially affecting collagenous structures associated with the skeletal joint.

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One mode of this aspect further includes delivering the energy into the region of tissue while the region of tissue is under mechanical stress; and heating the tissue to a temperature of up to no more than 75 degrees C.

Another mode includes heating the tissue to a temperature of up to no more than 70 degrees C.

Another aspect of the invention is a method for treating a region of tissue associated with an intervertebral disc in a body of an animal by delivering an ultrasound transducer to a location within the body such that a therapeutic level of ultrasound may be coupled from the transducer to the tissue.

One mode of this aspect includes delivering energy to a region of tissue associated with the spine, wherein such tissue does not experience a rise in temperature of more than about 55 degrees C.

Another aspect of the invention is a method for treating an animal by delivering energy to a region of tissue associated with the spine, wherein such energy delivery is between about 10 and about 300 equivalent minutes at 43 degrees C.

Another aspect is a method for invasively treating a medical condition associated with an intervertebral disc within a body of animal by delivering a therapeutic level of ultrasound energy to a region of tissue associated with an intervertebral disc from a location within the body of the patient.

Another aspect of the invention is a method for treating medical condition associated with a joint between two bony structures in a body of an animal by delivering an ultrasound transducer to a location within the patient's body associated with the joint; and emitting ultrasound energy from the transducer at the location so as to provide a therapeutic effect to at least a portion of the joint.

Another aspect of the invention is a method for treating an animal by introducing an ultrasound transducer into a body of the animal; positioning the ultrasound transducer at a location within the animal that is adjacent to at least one of an annulus fibrosus of an intervertebral disc, a nucleus pulposus of the

intervertebral disc, or a vertebral body associated with a spinal joint in the body; and emitting ultrasound energy from the ultrasound transducer at the location.

Another aspect of the invention is a method for providing ultrasound energy delivery within a body of an animal by introducing an ultrasound transducer into a body of a patient; positioning the ultrasound transducer at a location within the patient that is within at least one of an annulus fibrosus of an intervertebral disc, a nucleus pulposus of the intervertebral disc, or a vertebral body associated with a spinal joint in the body; and

emitting ultrasound energy from the ultrasound transducer at the location.

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Another aspect of the invention is a method for treating a patient by ultrasonically heating a region of tissue associated a spinal joint to a temperature between about 45 to about 90 degrees Fahrenheit for sufficient time to cause a therapeutic result in the tissue.

The method according to one further mode includes providing sufficient thermal dose so as to cause necrosis effect in nociceptive nerve fibers or inflammatory cells in the region of tissue.

Another mode includes providing sufficient thermal dose at appropriate temperature so as to stimulate cellular metabolism without substantially killing cells in the region of tissue.

Another mode includes delivering sufficient thermal dose and temperature in the region of tissue to cause a substantially non-necrotic cellular effect.

Another mode includes delivering sufficient thermal dosing to the region of tissue to cause preferential regeneration.

Another mode includes delivering sufficient thermal dose to the region of tissue to cause inhibition of inflammatory factors or cytokines.

Another mode includes delivering sufficient thermal dose to the region of tissue to cause modification of a healing response to injury in the region of tissue.

Further objects and advantages of the invention will be brought out in the following portions of the specification, wherein the detailed description is for the purpose of fully disclosing preferred embodiments of the invention without placing limitations thereon.

BRIEF DESCRIPTION OF THE DRAWINGS

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

FIG. 1A shows a side perspective view of an illustration of a typical human spine for treatment according to the systems and methods of the invention.

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- FIG. 1B shows an exploded, cross-sectioned side view of the region depicted as 1B in FIG. 1A, and shows an intervertebral disc between two adjacent vertebral bodies.
- FIG. 2 shows an angular perspective view of a transversely cross-sectioned intervertebral disc in relation to adjacent spinal structures.
 - FIG. 3A shows an angular perspective view of an ultrasonic intervertebral disc therapy system according to the invention, and includes a partially segmented view of an ultrasound treatment device, a schematic view of an ultrasound drive system, and an angular perspective view of an introduction device, respectively, of the system.
 - FIG. 3B shows a longitudinal side view taken along lines 3B-3B in FIG. 3A.
 - FIG. 3C shows a transverse cross-sectioned view taken along lines 3C-3C in FIG. 3A.
 - FIG. 4 shows a transverse cross-sectioned view of the distal end portion of another ultrasound treatment device of the invention with a different support structure under the ultrasound transducer of the device than the support structure shown in FIG. 3C.
 - FIG. 5A shows a slightly angled side view of the distal end portion of another ultrasound treatment device according to the invention, and also shows a schematic view of a guide wire included in the system.
 - FIG. 5B shows a cross-sectioned side view of the device taken along lines 5B-5B in Figure 5A.
 - FIG. 6A shows a slightly angled side view of an ultrasound transducer component assembly for use in the distal end portion of another ultrasound treatment device according to the invention.
 - FIG. 6B shows a cross-sectioned transverse view taken along lines 6B-6B in FIG. 6A.

FIG. 6C shows a schematic cross-sectioned side view of an alternative ultrasound transducer component assembly within an ultrasound treatment device according to the invention versus that shown in FIG. 6B.

FIG. 6D shows an angular perspective view of a semi-spherical disc-shaped transducer for use according to a further embodiment of the invention.

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- FIGS. 7A-F show a top perspective view of a laterally cross-sectioned intervertebral disc during respectively sequential modes of operating the ultrasound treatment system for intervertebral disc therapy according to the invention.
- FIGS. 8-9 show alternative modes of operating an ultrasound treatment device according to the invention for treating different respective regions of an annulus fibrosus of a disc from within the nucleus and according to a posterior-lateral approach into the disc.
- FIG. 10 shows an alternative mode of operating an ultrasound treatment device according to the invention using an anterior approach to treat a posterior wall region of the annulus fibrosus of a disc from within the nucleus of the disc.
- FIGS. 11-13 show alternative modes of operating an ultrasound treatment device for treating different respective regions of an annulus fibrosus of a disc from externally of the annulus fibrosus and without entering the nucleus, wherein FIGS. 11 and 13 show a posterior-lateral approach to right lateral and posterior wall regions of the annulus, respectively, and FIG. 12 shows an anterior approach to a left lateral wall region of the annulus.
- FIGS. 14A-B show plan perspective views of the distal end portion of another ultrasound treatment device of the invention during various modes of operation, wherein FIG. 14A shows a straight configuration for the device, and FIG. 14B shows two modes of angular deflection for the distal end portion of the device.
- FIG. 14C shows a top perspective view of another ultrasound treatment device of the invention with a distal ultrasound treatment section that is adapted to be rotated about a hinge point for minimally invasive treatment of intervertebral discs and other spine or joint disorders.
- FIG. 15A shows a plan perspective view of the distal end portion of another ultrasound treatment device of the invention having a predetermined shape and operative region for ultrasound delivery that corresponds to particular desired

approaches and ultrasound therapy to certain specified regions of tissue associated with an intervertebral disc.

FIG. 15B shows a perspective view of another ultrasound treatment device having a similar shape to that shown in FIG. 15A, but having a different operative region for ultrasound delivery corresponding to delivering invasive therapy to a different desired region of a respective intervertebral disc.

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FIG. 16 shows a perspective view of another ultrasound treatment device having a different unique shape and operative region for ultrasound delivery corresponding to delivering invasive therapy to a different desired region of a respective intervertebral disc.

FIG. 17A shows a top view of an ultrasound treatment device assembly with transducers inside of an outer cooling jacket that is interfaced with a fluid circulation pump to actively cool the transducers.

FIG. 17B shows a top view of another partially cross-sectioned ultrasound treatment device assembly similar to that shown in FIG. 17A, except showing the cooling fluid to circulate from within the ultrasound transducer device and into the surrounding sheath.

FIG. 18A shows an x-ray picture of an explanted disc being treated according to one aspect of the invention, and shows various data points along temperature monitoring probes inserted along certain desired locations for monitoring across the disc.

FIG. 18B shows a graph of temperature vs. time for an ultrasound heating study in an explanted cadaver spine disc, and shows curves for tissue depths of 1mm, 4mm, 7mm, and 10mm away from the directional heating transducer.

FIG. 19 shows a typical modulus versus applied stress plot according to a study performed in Example 2, and shows results before (solid line) and after (dashed line) heat treatment at 85°C, and indicates the following biomechanical parameters: change in modulus at the inflection point (MI), change in modulus at 150kPa (M150), and change in residual stress at the inflection point (RSI).

FIG. 20 shows a typical stress-strain plot before (solid line) and after (dashed line) treatment at 85°C, and shows for each cycle an upper line indicating the loading phase, and a lower curve indicating the unloading phase.

FIG. 21 shows graphs (a)-(e) that variously represent respective changes in certain tissue parameters that were observed after varied heat treatments.

FIG. 22 shows various light microscopy photographs of various tissue samples, and includes bright light (left panel) and polarized light (right panel) microscopy of specimens treated, and shows pictures (a, b) for samples intact at 37°C; pictures (c, d) for intact samples at 85°C; and pictures (d, e) representing excised at 85°C.

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- FIG. 23 shows a schematic side view of a distal end portion of an external directional ultrasound thermal treatment ("ExDUSTT™") device of the invention incorporating a directional, focused ultrasound emitter assembly that is adapted for external use adjacent to an intervertebral disc.
- FIG. 24 shows an illustrated side view of a partially cross-sectioned ultrasound spinal treatment assembly similar to that shown in the ExDUSTT device shown in FIG. 23, and shows the assembly during one mode of use in treating a region of an intervertebral disc associated with a spinal joint.
- FIG. 25 shows a plan view of a distal end portion of a further ExDUSTT embodiment that incorporates a substantially rigid, pre-shaped probe device platform.
- FIG. 26 shows a plan view of a distal end portion of another ExDUSTT device embodiment that incorporates a substantially flexible, catheter device platform according to another embodiment of the invention.
- FIG. 27A shows a longitudinally cross-sectioned view of a distal end portion of an ExDUSTT device on a rigid probe platform similar to that shown in FIG. 25, and shows a substantially compliant elastomeric balloon over a curvilinear ultrasound transducer.
- FIG. 27B shows a transverse cross-sectioned view through an ultrasound transducer mounting region of the ExDUSTT device shown in FIG. 27A.
- FIG. 28A shows a longitudinally cross-sectioned view of a distal end portion of another ExDUSTT device on a rigid probe platform that is also similar to that shown in FIG. 25, except with a substantially non-compliant pre-formed balloon over the curvilinear ultrasound transducer.
 - FIG. 28 B shows a transverse cross-sectioned view through an ultrasound

transducer mounting region of the ExDUSTT device shown in FIG. 28A.

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FIG. 29A shows a transverse cross-sectioned view of a distal end portion of another ExDUSTT device on a polymeric catheter delivery chassis similar to that shown in FIG. 26, and shows a substantially compliant elastomeric balloon over a transversely aligned, curvilinear ultrasound transducer.

FIG. 29 B shows a transverse cross-sectioned view through an ultrasound transducer mounting region of the ExDUSTT device shown in FIG. 29A.

FIG. 30A shows a transverse cross-sectioned view of a distal end portion of another ExDUSTT device on a catheter delivery platform similar to that shown in FIG. 29A, except with an axially aligned, curvilinear ultrasound transducer within a substantially compliant elastomeric balloon.

FIG. 30B shows a transverse cross-sectioned view through an ultrasound transducer mounting region of the ExDUSTT device shown in FIG. 30A.

FIGS. 31A-B show two respective graphs for acoustic efficiency and acoustic output power, respectively, for one exemplary working embodiment of a rigid probe ExDUSTT device similar to that shown in FIG. 25.

FIGS. 32A-B show two respective graphs for acoustic efficiency and acoustic output power, respectively, for one exemplary working embodiment of a catheter-based ExDUSTT device similar to that shown in FIG. 26.

FIG. 33 shows a schematic drawing of an ultrasound heating assembly portion of a catheter-based ExDUSTT device similar to that shown in FIG. 26 superimposed over an X-ray picture of the intervertebral disc to illustrate one experimental set-up to evaluate the device, and also shows superimposed reference numbers designating certain monitored temperatures at various locations within the disc during one mode of treatment.

FIG. 34A-B show two respective graphs of temperature monitored over time at various thermocouples locations 105(C)-109(C) during two respective in-vivo thermal therapy treatments in an intervertebral pig disc using a catheter-based ExDUSTT device similar to that shown in FIG. 26 and according to an experimental set-up similar to that shown in FIG. 33.

FIG. 35 shows two, respective ExDUSTT devices of pre-shaped, rigid probe construction similar to that shown in FIG. 25, except the devices shown are

constructed according to different size embodiments incorporating ultrasound transducers having varied respective widths of 2.5mm and 3.5mm, respectively.

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FIGS. 36A-B show top and angular perspective views, respectively, of certain power output profile across the face of the 2.5mm wide transducer shown in FIG. 35A.

FIGS. 37A-B show top and angular perspective views, respectively, of certain power output profiles across the face of the 3.5mm wide transducer shown in FIG. 31B.

FIG. 38 shows an X-ray picture of a top view of an ex-vivo experimental arrangement similar to that shown schematically in FIG. 33, except showing a directional ultrasound heating assembly of a working embodiment for a probe-based ExDUSTT device such as shown in FIG. 25 positioned for desired heating of an intervertebral disc according to one modes of use, and shows various thermocouple probes within the disc to monitor experimental temperatures.

FIG. 39 shows an exploded view of the same Ex-DUSTT intervertebral disc treatment arrangement shown for the 3.5mm probe-like ExDUSTT device in FIG. 38, and shows monitored temperature values at various respective locations along the axial and radial temperature probes during one relatively high temperature mode of use with active cooling at 0 degrees C.

FIG. 40 shows a graph of temperature vs. time for the various temperature probes according to the thermal therapy arrangement shown in FIG. 39.

FIG. 41 shows a graph of temperature monitored along the 5mm and 10mm deep axial temperature sensor probes and the radial temperature sensor probes shown within the intervertebral disc and during treatment with the probe-based ExDUSTT with the 3.5mm wide transducer shown in FIG. 38, and according to an ex-vivo study performed with active transducer cooling at 0 degrees C.

FIG. 42 shows the same exploded view of the experimental arrangement shown in FIG. 39, except shows thermocouple values corresponding to a relatively low temperature mode of operation with room temperature cooling.

FIG. 43 shows a similar graph of temperature vs. time as that shown in FIG. 40, except with respect to data measured according to the arrangement illustrated for FIG. 42.

FIG. 44 shows another graph of certain temperature vs. transducer position results similar to the graph shown in FIG. 41, except showing results according to the mode of operation also variously illustrated in FIGS. 42-43.

FIG. 45 shows a similar exploded X-ray picture to that shown in FIGS. 39 and 42, except showing thermocouple values according to a relatively low temperature mode of operation using a 3.5mm wide transducer and cooling at 0 degrees C.

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- FIG. 46 shows another temperature vs. time graph, except with respect to the arrangement also illustrated in FIG. 45.
- FIG. 47 shows a graph of temperature vs. thermocouple position results according to the thermal therapy arrangement illustrated in FIGS. 45 and 46.
- FIG. 48 shows another exploded X-ray picture of the same ExDUSTT arrangement, except according to use of a 2.5mm wide transducer with relatively low temperature heating mode and cooling at 0 degrees C.
- FIG. 49 shows another temperature vs. time graph for the ExDUSTT heating arrangement illustrated in FIG. 48.
- FIG. 50 shows another temperature vs. thermocouple position graph for a 2.5mm curvilinear ExDUSTT ex-vivo disc treatment using 0 degree C cooling and relatively low temperature mode of operation.
- FIG. 51 shows another exploded X-ray picture, except with thermocouple values corresponding to use of a 2.5mm transducer ExDUSTT device at a relatively low temperature mode of use with room temperature cooling.
- FIG. 52 shows a temperature vs. time graph according to the ExDUSTT mode of therapy also illustrated in FIG. 51.
- FIG. 53 shows a temperature vs. thermocouple position graph for the modes of ExDUSTT therapy illustrated in FIGS. 51 and 52.
 - FIG. 54 shows a perspective view of an internal directional ultrasound thermal therapy system ("InDUSTT™") that includes a spinal disc delivery probe and an InDUSTT device that fits within the spinal disc delivery probe.
 - FIG. 55A shows a plan view of a schematic representation of an internal ultrasound thermal spine therapy device according to another embodiment of the invention.
 - FIG. 55B shows an exploded view taken at region B shown in FIG. 55A, and

shows enhanced detail of various aspects of the ultrasound heating assembly along the distal end portion of the InDUSTT system according to a further feature of that embodiment.

FIG. 55C shows a transverse cross-sectioned view taken along line C-C in FIG. 55A.

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FIGS. 56A-B show respective X-ray pictures of the distal end portion of an InDUSTT system similar to that shown in FIGS. 54-55B positioned within an intervertebral disc during in-vivo thermal spinal treatments according to certain modes of the invention.

FIG. 57 shows a table providing thermal dosimetry data collected during certain modes of in-vivo operation for various working embodiments of an InDUSTT system similar to that shown in FIGS. 54-55B and providing therapeutic ultrasonic heating at various temperature modes of powered operation corresponding to C2/3, C3/4, C4/5, and C5/6 intervertebral sheep discs, respectively.

FIG. 58 shows various X-ray pictures of the placement of the InDUSTT transducer within the C2/3, C3/4, C4/5, and C5/6 intervertebral discs corresponding to the results provided in the table in FIG. 57.

FIG. 59A shows a graph of temperature vs. time during InDUSTT heating of a C3/4 intervertebral disc according to a relatively high temperature mode of use, and shows curves for various respective thermocouple probe positions.

FIG. 59B shows a graph of temperature vs. time using the same InDUSTT device as that used for creating the data shown in FIG. 59A, except shows results according to a relatively low temperature mode of use in a C4/5 intervertebral disc location.

FIG. 59C shows a graph of temperature monitored at various temperature sensor positions during 10 minute InDUSTT heating, and shows curves for results in two separate intervertebral discs each heated with a different one of two separate InDUSTT systems of the invention.

FIG. 60 shows another table providing thermal dosimetry data collected during modes of in-vivo operation for various working embodiments of a directly coupled InDUSTT system providing therapeutic ultrasonic heating from within C2/3, C3/4, and C4/5 intervertebral discs of a sheep.

FIG. 61 shows various respective X-ray pictures of certain transducer placements for the directly coupled InDUSTT during in-vivo spinal disc thermal therapy at the C2/3, C3/4, and C4/5 intervertebral sheep discs corresponding to the similarly designated rows of data illustrated in the table of FIG. 60.

FIG. 62A shows a graph of temperature vs. time corresponding to the C2/3 disc treatment shown in FIG. 61 and according to a relatively high temperature mode of use.

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FIG. 62B shows a graph of temperature vs. time corresponding to the C3/4 disc treatment shown in FIG. 61, and according to a relatively low temperature mode of use.

FIG. 62C shows a graph of temperature monitored at various temperature sensor positions during 10 minute directly coupled InDUSTT heating, and shows curves for thermal treatment results at both dead sectors and active sectors of the transducer in the C2/3 disc at relatively high temperature power level, and at similar locations in the C3/4 disc at the corresponding, relatively low temperature power level.

FIG. 62D shows a graph of accumulated thermal dose versus temperature sensor position for the 10 minute treatments at the C2/3 and C3/4 discs at the relatively high and low temperature power levels, respectively.

FIG. 63 shows another table providing thermal dosimetry data collected during modes of in-vivo operation for various working embodiments of a catheter cooled InDUSTT system providing ultrasonic heating from within C2-3, C3-4, C4-5, and C5-6 intervertebral discs of a sheep.

FIG. 64 shows various respective X-ray pictures of certain transducer placements for the catheter cooled InDUSTT during in-vivo thermal spinal disc therapy at the C2-3, C3-4, C4-5 and C5-6 intervertebral discs corresponding to the similarly designated rows of data illustrated in the table of FIG. 63.

FIG. 65A shows a graph of the relatively high temperature, catheter cooled InDUSTT therapy of the C2/3 disc as monitored over multiple temperature sensors along first and second temperature probes positioned within the disc.

FIG. 65B shows a temperature vs. time graph of the relatively low temperature mode of operation for the catheter cooled InDUSTT therapy in the C3/4 disc and as

monitored over multiple temperature sensors along first and second temperature monitoring probes positioned within the disc.

FIG. 65C shows a temperature vs. time graph of the relatively high temperature, catheter cooled InDUSTT therapy of the C5/6 disc as monitored over the multiple temperature sensors along first and second temperature monitoring probes positioned within the disc.

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FIG. 65D shows a graph of temperature monitored at various axial positions relative to the transducer during 10 minute catheter cooled InDUSTT heating, and shows curves for thermal treatment results at various locations in the C2/3 disc at relatively high temperature power level, in the C3/4 disc at relatively low temperature power level, and C5/6 disc at relatively high temperature power levels.

DETAILED DESCRIPTION OF THE INVENTION

Referring more specifically to the drawings, for illustrative purposes the present invention is intended to provide thermal treatment to spinal joints, and in particular intervertebral discs as illustrated in FIGS.1A-2, as embodied in the apparatus shown and characterized by way of the various modes of operation with respect to certain intended anatomical environments of use variously throughout FIGS. 3-65D. 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.

As an initial introduction, various respective aspects, modes, embodiments, variations, and features of the invention are herein shown and described, both broadly and in variously increasing levels of detail. Each provides individual benefit, either in its own regard, or in the ability to provide enhanced modes of operation and therapy by way of combinations with other aspects or features. Moreover, their various combinations, either as specifically shown or apparent to one of ordinary skill, provide further benefits in providing useful healthcare to patients.

In one regard, two illustrative ultrasound spinal thermal therapy probe configurations are described for applying thermal (heat) therapy or ultrasound (US) exposure to tissues within the spine or other joints. Heat at high temperatures and thermal doses can shrink tissues, change the structural matrix, generate physiological changes and/or kill cells. Heat at relatively lower temperatures and US

exposure can generate permeability changes or changes in the cellular transport/metabolism that increase effectiveness or deposition of certain pharmaceutical agents. The heat or US can be delivered with the present invention in a highly controlled fashion to selected tissue regions in order to exploit these physiological effects for therapeutic purposes. Ultrasound applicators may achieve more precise targeting or heating control not possible with current RF and Hot Source techniques. For soft tissue or bone surfaces within the spine or other joints, the high temperature exposure can be used to shrink tissue impinging on nerves, restructure and possibly strengthen mechanical properties of the disc or joint material, destroy abnormal or undesirable cells or tissue, destroy nerves responsible for pain, seal leaks from the disc annulus/nucleus, joint capsules, etc. Novel ultrasound applicators and treatment methodologies are thus herein shown and described which allow for the interstitial insertion or laparoscopic or arthroscopic placement of these applicators within or upon targeted tissue to receive such treatments or prophylaxis.

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As will be further developed by reference to the Figures below, one exemplary type of such an applicator and treatment methodology provides a segmented array of tubular, sectored tubular, plate, hemispherical, or portions of cylinders (e.g. convex) with linear control of US exposure or heating via power level adjustments and angular control of US exposure or heating via directional characteristics of the applicators. (e.g. angularly directive with an inactive zone). These transducers are mounted over a guidewire lumen or tube or structure to facilitate placement, wires, and/or cooling structures. Thermometry sensors can be placed directly on the transducer/tissue or applicator/tissue interface. Internal cooling via gas or liquid or external cooling via an outer plastic sheath or catheter can be accomplished, though may not be necessary in many instances. These can be inserted within the disc or laparoscopically placed against the target tissue or directed toward the target tissue. Acoustic gain and temperature regulation of applicator surface(s) can help control distance of heated regions and effects from the applicator surface. Frequency and depth of focus can be selected to control heating pattern, and time can be varied to control heating effects and distribution. Some of the device and method embodiments provided herein may incorporate various features similar to those previously disclosed such as in US Patent No. 5,620,479 to Diederich, though in

many instances will be modified specifically for heating within the special environment of use within or around intervertebral discs or other joints.

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Another illustrative type of applicator according to the present invention incorporates a segmented array similar to that described above, but using concave sections of cylindrical or tubular transducers or spherical or semi-spherically focused transducers. The outer diameter (OD) of the tubes used to form such transducers are much larger than the applicator diameter for the tubes – the sectors activated are a small arc of the tube they otherwise would be a part of. Thus, a line of convergence, e.g. focus, is produced at depth over a small arc angle, producing an intense US exposure or heating pattern which is approximately the same length as the tubular segment transducer emitting the US, though very narrow (e.g. 1-5 degrees) in the angular dimension. The length and number of segments can be varied in either applicator type described here for introduction purposes (or elsewhere herein), and may be a single transducer versus an array. These applicators can also have internal cooling or external cooling as described above and further detail with respect to the particular embodiments below. The applicators can be inserted within the disc or laparoscopically placed against the target tissue or directed toward the target tissue. Acoustic gain and temperature regulation of applicator surface can help control distance of the heated region and effects from the applicator surface. Further applicable features may by incorporated from other prior disclosures, such as U.S. Patent No. 5,391,197 to Burdette et al. disclosing prostate therapy devices and methods, and may be modified to suit the particular needs for the present invention. Frequency and depth of focus can be selected to control heating pattern, and time can be varied to control heating effects and distribution.

The various embodiments herein described have applications in other soft and/or hard tissue sites and body parts where ultrasound exposure, high temperature, low temperature, or combination effects are desired.

Each type of applicator can be designed with or without cooling balloons, distendable (e.g. compliant and/or elastomeric) or pre-shaped (e.g. substantially non-compliant with relatively fixed inflation size and shape), and symmetric or asymmetric shapes are considered. The devices' respective chasses may be substantially stiff, e.g. rigid probes, or flexible. They may further be either implantable within the target

tissue, or be used on surface contact. They may be delivered on a guidewire rail platform, through pre-shaped insertion or placement guides, or have their own steerability or deflectability. For spinal treatments, they may be placed surgically following for example a posterior approach, or laparoscopic/arthroscopic lateral/anterior directed to the spinal joint for treatment.

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Treatment methodologies contemplated include implanting the devices within or positioning them next to the target tissue for heating, such as for example inserted into a disc or joint capsule, or placed outside of the disc or joint.

Directivity and cooling aspects, when incorporated, protect sensitive non-targeted tissue, which is highly beneficial for example in spinal applications protecting spinal nerves. Applicators herein described are repositionable according to various modes to control angular thermal profile according to their directed energy delivery. In one example for further illustration, a specially adapted spinal disc insertion apparatus is adapted to deflect an applicator being delivered therethrough into the spinal disc from an angle. Other special procedures and tools are also herein described to align the applicators with target areas of tissue such as with respect to spinal joints and intervertebral discs in particular.

Though many different configurations, sizes, shapes, and dimensions are contemplated consistent with the overall intent to meet the various objects of the invention, exemplary devices may be provided with outer diameters between about 1.2 to about 3mm, though may be up to 5mm in some instances, deliverable as desired to spinal joint areas from 18 gauge.

Insertion techniques into tissue to be treated may progress according to several example. In one mode, a relatively stiff (e.g. sufficient to support the intended use), pre-shaped guidewire is used which may be with or without memory metal alloy such as nickel titanium for example. The guidewire is inserted under fluoroscopy and positioned in an annulus fibrosus or posterior annulus, avoiding the nucleus of the disc. An applicator of the relatively more flexible variety is then inserted over the guidewire and into position. In another regard, a relatively stiff (e.g. sufficient support) pre-shaped insertion tool guides the applicator with a sharp tip into the annulus from outside without requiring the guidewire (though they may be used in conjunction). Similar insertion techniques may be used for thermometry placement,

if desired. Such delivery tool may thus be multi-lumened to integrate both placements (e.g. applicator and temperature probes) simultaneously for better positioning, etc.

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Contact therapy techniques of operation may also proceed according to a variety of modes. An arthroscopic approach is suitable for many applications, such as for example as follows. Internal tip deflection may be used to align (e.g. steer) the applicator with or along the outside of an annulus –e.g. similar to certain intracardiac catheters (such as mapping or ablation devices). Such may be integrated to a steerable catheter. The device according to these modes may be placed lateral or posterior behind the disc and nerves, or ventral. The device is aligned with the disc, the region is targeted and then treated with directional thermal therapy.

Various of the components herein described for the various embodiments may be provided together, or may be provided separately. For example, implements for providing streaming liquid or balloon to protect tissue from transducer conductive heating may be an integral part of the respective applicator, or may be separate as an accessory.

The applicators and respective insertion and/or guidance tools herein described may be further adapted to be compatible with magnetic resonance imaging for real time monitoring of the procedure. Other imaging modalities may also be used for positioning, monitoring, thermal monitoring, lesion assessment, real-time monitoring of coagulation, etc. This includes ultrasound monitoring.

Further to the ultrasound aspects of the various embodiments, use of such energy modality provides temperature elevation as one mode of creating an intended effect, but also provides other non-thermal effects on tissues, such as for example drug activation, etc., such as for example to treat arthritis at joints where the applicator is being used.

It is to be appreciated that the invention is in particular well adapted for use in treating intervertebral disc disorders of the spine, such as at spinal joints, and in particular at an intervertebral disc 1 shown in various relation to surrounding spinal structures of a spinal joint in FIGS. 1A-2. In particular, as will be further developed below, disc disorders associated with chronic lower lumbar back pain are to be beneficially treated according to the invention. However, it is to be appreciated that

other disorders of the discs in particular, and of other joints (e.g. hips, knees, shoulders, etc.) may also be treated according to the device systems and methods herein shown and described. For example, other regions of the vertebrae will be beneficially treated with invasive ultrasound delivery according to the invention in order to promote bone growth, such as for example to assist in the healing of injuries or bone-grafts. Areas between the vertebral bodies, or the spinal processes, for example, may be treated with US application from the present devices and according to the methods as herein shown and described. In particular, regions such as graft between inner bodies through nucleus; anterior inner body fusion; posterior lateral fusion are contemplated. Ultrasound energy may be delivered with collagen matrixes or autograft/allograft materials to bone.

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A typical intervertebral disc 1 such as shown in FIGS. 1A-2 generally includes an annulus fibrosus 2 that surrounds a nucleus pulposus 3 along a plane that lies between two adjacent vertebrae 8,9, respectively, that are located above and below, also respectively, disc 1 along the spine. More specifically, disc 1 lies between two the two cartilaginous endplates 8a,9a that border two adjacent vertebral bodies 8b,9b of vertebrae 8,9, respectively.

As will be further developed below, an ultrasound treatment device according to the invention may be located in various places in and around a disc 1. A variety of such locations is shown for the purpose of illustration at locations a-d in FIG. 1B, wherein device 11 is shown: within the middle of the nucleus at location a; along the border between the nucleus 3 and the annulus 2 such as shown at proximal wall at location b; in the wall of the annulus itself, as shown for example at location c; or outside of the disc 1 around the outer periphery of annulus 3, as shown at location d. Moreover, the device may also be delivered into and around bony structures associated with the spinal joint, such as for example shown at locations e, f, g, or h in FIG. 1B. Such positioning may be accomplished for example by drilling a bore into the vertebral body from a posterolateral approach through an associated pedicle, as shown in shadow at location E in FIG. 2, or via a more lateral approach as shown directly into the body at location F in FIG. 2. Such positioning and heating within bone structures associated with the joint may be in particular useful in one regard for treating bone cancer, destroying nociceptive nerves, stimulating growth or drug

uptake (e.g. low thermal dose applications). Either the vertebral body itself may the target for heating, or the end plate, or the disc from such location. A further particular useful application of this is treatment of osteoporotic back pain.

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As shown in particular in FIG. 2, disc 1 also has a shape similar to a "kidney"-shape with a concave curvature along a proximal wall 4 that borders the spinal cord (not shown), as well as along opposite anterior wall 5. Right and left anterior walls 6,7 are generally characterized by a more acute radius of curvature than posterior and anterior walls 4,5. As will be further developed below, each of these uniquely located and anatomical wall regions may be selectively treated with localized therapeutic ultrasound energy according to the system and method of the present invention. In general, intervertebral discs (with respect to the lumbar region associated with lower back pain) are typically 30mm wide (e.g. laterally from right wall 6 to left lateral wall 7, about 20mm front-back, e.g. anterior wall 5 to posterior wall 4); and approximately 10mm tall, e.g. from end plate 8a to endplate 9a. Accordingly, the devices herein shown and described are to be particularly adapted to operate within this general description of the intended environment of use within intervertebral discs.

As will be appreciated by the description below of the various modes of operating the ultrasound treatment system of the invention, treatment of the annulus fibrosus 2 from within the nucleus may be achieved via various approaches. In particular, regions A and B shown in FIG. 2 correspond to right and left anterior approaches, whereas regions C and D correspond to right and left posterior-lateral approaches around right and left vertebral prostheses 8c,8d, respectively

Multiple ultrasound probe configurations are herein described for applying thermal (heat) therapy or ultrasound (US) exposure to tissues within the spine in particular, though other joints such as knee, hip, etc. are contemplated. It is to be appreciated that the two specific probe configurations shown and described provide highly beneficial embodiments, though they are exemplary and other configurations, improvements, or modifications according to one of ordinary skill based upon this disclosure in view of the known art are contemplated.

In any event, heat produced according to the present invention at high temperatures and thermal doses can shrink tissues, change the structural matrix,

generate physiological changes, and/or kill cells within the targeted region of tissue associated with a disc. Heat at low temperatures and US exposure can generate permeability changes or changes in the cellular transport/metabolism that increase effectiveness or deposition of certain pharmaceutical agents. The heat or US can be delivered with this technology in a highly controlled fashion to selected tissue regions in order to exploit these physiological effects for therapeutic purposes.

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Ultrasound applicators may achieve a degree of precise targeting or heating control generally not possible with previously disclosed RF, plasma ion, or heat source techniques. In addition, ultrasound energy actually penetrates surrounding tissues, rather than according to other modes (e.g. RF and laser) that heat the closest tissues the hottest and allowing conduction therefrom in a diminishing temperature profile curve with distance away. For soft tissue or bone surfaces within the spine or other joints, high temperature exposure by use of the invention is used to shrink tissue impinging on nerves, re-structure and possibly strengthen mechanical properties of the disc or joint material, destroy abnormal or undesirable cells or tissue, destroy nerves responsible for pain, and seal leaks from the disc annulus/nucleus, joint capsules, etc.

In particular to disc applications, three general goals are intended to be achieved according to use of the present invention: (1) collagen associated with the annulus fibrosus may be reorganized to reshape the annulus; (2) nerve ingrowth in and around the annulus or nucleus may be killed; or (3) inflammatory cells around areas of injury or otherwise penetrating areas in or around a disc may be killed or ablated. In particular with respect to causing nerve damage, this may include regions of the annulus itself, at the endplates, usually is located posteriorly, and rarely but at times may be within the region of the nucleus itself. In any event, such nervous ingrowth is typically related to structural disc damage that is identified e.g. in a discogram and therefore predicted to be where pain/nerve treatment should be directed.

In one particular non-limiting application, either or both of nerve and inflammatory cells are necrosed by US delivery without achieving sufficient heating to denature or weaken, or to denature but not weaken, or to reshape the disc annulus. This is possible using the devices and methods of the invention herein described at

levels of energy delivery between about 10 to about 300 EM43degC (e.g. may be from 1 to 60 min at between about 42degC and about 45degC). Where collagen denaturation, modification, or reshaping is desired, energy delivery from the ultrasound devices herein described may be from between about 55degC to about 85degC for between about 10sec to about 30min.

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The novel ultrasound applicators and treatment methodologies herein disclosed allow for the interstitial insertion or laparoscopic or arthroscopic placement of these applicators within or upon targeted tissue, in particular with respect to intervertebral discs.

One particularly beneficial embodiment of the invention is shown at ultrasound treatment system 10 in FIGS. 3A-C. This system 10 includes an ultrasound device 11, ultrasound drive system 40, and intervertebral disc delivery assembly 50.

Device 11 is shown to couple proximally to an a proximal end portion (not shown) that generally includes a handle (not shown) that is adapted to couple to ultrasound drive system 40, which includes an ultrasound actuator 41. Drive system 40 may be operated empirically, such that a predetermined delivery of energy is achieved at a desired level known to produce a desired result. Or, external therapy monitoring may be employed during treatment, e.g. MRI, CT, fluoroscopy, X-ray, discogram, or PET in order to control energy delivery and determine appropriate levels and time duration for a particular case. These monitoring modalities may be effective prior to treatment in order to identify the area of concern to be treated, which may impact the choice of particular device to be used as provided according to the embodiments herein. Still in a further alternative embodiment, a treatment feedback device 42, such as a temperature monitoring system, may be incorporated in a feedback control system, as shown in FIG. 3A.

Device 11 is also adapted to be delivered to the desired location for treatment through delivery assembly 50 and therefore has a length corresponding to length L of delivery assembly 50 that is adapted for use in standard access procedures for intervertebral disc repair. For posterior-lateral approaches, such as for example in order to invade the nucleus 3 through posterior-lateral sites B or C shown in FIG. 2, delivery assembly 50 is typically a spinal needle of about 18 Gauge. Accordingly, the length for device 11 may be about 30 cm long, with a corresponding outer diameter

for device 11 adapted to fit within such a needle, generally less than about 3 mm, generally between about 1 and about 3 mm, typically between about 1.2 and 3 mm. However, other sizes may be realized for applications not requiring delivery through size-limiting delivery assemblies such as spinal needles, and up to or greater than 5mm OD is realizable (e.g. in particular for applications outside of the disc nucleus or within the annulus). For anterior delivery such as at sites A or B shown in FIG. 2, delivery assembly 50 may be minimally invasive delivery device such as an arthroscope or laparoscopic assembly.

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Device 11 is of a type that contains a linear array of segmented transducers 16 that are adapted to provide selective, localized ultrasonic heating via radial, collimated energy delivery in tissue adjacent to the array. The particular device 11 of the present invention, including corresponding elements such as transducers 16 located thereon, are generally smaller and more flexible than elsewhere previously described for other linear array transducer devices. In addition, fewer transducers 16 are typically required for treating the generally smaller regions of the intervertebral discs as contemplated herein. These substantial modifications are believed to significantly enhance the controllability and performance of ultrasound therapy within the unique (and often dangerous) anatomy of an intervertebral disc. Otherwise, the basic components for segmented, linear array transducer device 11 may be similar to those previously described in U.S. Patent No. 5,620,479, which has been previously incorporated by reference above.

Referring more specifically to FIGS. 3A through 4, an ultrasound applicator 11 of the invention preferably includes a cylindrical support member such as a tube, conduit or catheter 12 which may be compatible for adjunctive radiation therapy of the spine such as according to remote afterloaders and standard brachytherapy technology. Since catheter 12 includes a coaxial longitudinal inner lumen 14, a source of radiation, a drug or a coolant can be inserted therein, as well as guidewires, deflection members, stylets, etc., according to modified embodiments elsewhere herein described. One highly beneficial embodiment for example uses a polyurethane or other similar polymer that is of a soft, low modulus type according to uses as contemplated herein within the sensitive intervertebral discs and elsewhere along the spine and related, highly sensitive nervous tissues. Where direct access to

the desired treatment location is possible without risk of damaging soft, sensitive tissues around the spine, a more rigid support may be used, and may even include a thin-walled stainless steel hypodermic tubing or stiff conduits (though shapes may be important as further developed below).

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Catheter 12 is coaxially disposed through a plurality of tubular piezoceramic transducers 16 which are spaced apart and electrically isolated as shown, thus forming a segmented array of tubular ultrasound transducers which radiate acoustical energy in the radial dimension. Transducers 16 may be formed from a variety of materials as has been previously disclosed. Transducers 16 may have an outer diameter between about .5 to about 6 mm, though with respect to energy therapy from within the nucleus 3, is more typically between about 0.5 mm and about 1.5 mm.

It is preferred that the wall thickness for transducers 16 be substantially uniform in order to generate uniform acoustic patterns where such energy delivery is desired. Further to conjunctive radiation therapy (e.g. spinal tumor therapy), the transducer material is preferably stable when exposed to typical radiation sources.

The frequency range of the transducers will vary between approximately 5-12MHz depending upon the specific clinical needs, such as tolerable outer diameter, depth of heating, and inner catheter outer diameter. Inter-transducer spacing is preferably approximately 1 mm or less. Those skilled in the art will appreciate that, while three transducers 16 are shown in FIG. 3A-B, the number and length of transducers can be varied according to the overall desired heating length and spatial resolution of the applicator and depth of penetration desired. This may vary for example for devices 11 intended to treat along the entire length of posterior wall 4 of disc annulus 2, versus a lateral wall 6,7 thereof, versus the anterior wall 5. Even the desired length along a given one of these regions may vary depending upon the particular patient, or even within a given patient depending upon a particular region of the spine being treated (e.g. lower discs along the spine increase in size). Therefore, a kit of devices 11 having varied lengths and sizes for the array of treatment transducers 16 is contemplated according to such variances. Transducers 16 are also shown to be substantially cylindrical for the purpose of illustration, and which design may be desired where uniform heating around the circumference of the

device 11 is desired. However, as developed below, the highly selective tissue therapy typically desired within and around intervertebral discs may require in many cases more radial selectivity around the device 11, as further developed below.

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Each transducer 16 is electrically connected to actuator 41 of drive assembly 40, which is typically an RF current supply. This electrically coupling is achieved via separate pairs of signal carrying wires 18, such as 4-8 mil silver wire or the like, soldered directly to the edges of the transducer surface to form connections 20a,20b. One wire in the pair is connected to the edge of transducer 16 at its outer surface, while the other is connected to the edge of transducer 16 at its inner surface, although other connection points and modalities are also contemplated. Each wire 18 is routed through the center of the transducers between the outer wall of catheter 12 and the inner wall of the transducer 16, as can be seen in FIG. 3B, and from there to the connection point through the spaces between the individual transducer elements.

In order to ensure that each transducer 16 in the array is kept centered over catheter 12 while still maintaining flexibility and not impending transducer vibration, a plurality of spacers 22 are disposed between the transducers and catheter 12. These spacers 22 may take various forms as previously described. For the particularly smaller designs herein contemplated for spinal applications, a "springground lead" comprising 3-4 mil stainless steel wire or the like wound to form a coaxial spring may be placed between a transducer 16 and an electroded outer surface of inner catheter 12 where such coil is soldered directly thereto. Such electroded outer surface may be a common ground for all transducers 16.

As previously disclosed, transducers 16 are preferably "air-backed" to thereby produce more energy and more even energy distribution radially outwardly from device 11. To ensure such air-backing and that the transducers 16 are electrically and mechanically isolated, a conventional sealant 24 as previously described is injected around exposed portions of catheter 12, wires 18, and spacers 22 between transducers 16. Sealant 24 serves multiple functions in this application, as has been previously described.

As a means for monitoring temperature of tissue surrounding the transducers 16, and to provide for temperature control and feedback where desired, a plurality of

small (e.g. 25.4um) thermocouple sensors 26, such as copper-constantan or constantan-maganin fine wire junctions, are placed along the outer surface of each transducer at points which are approximately one-half of the length of the transducer, and connected to individual temperature sensing wires 28 which run along one-half of the length of the transducer 16 and then through the space between catheter 12 and the transducers 16. A conventional acoustically compatible flexible epoxy 30 such as has been described is then spread over the transducers, thereby embedding the temperature sensors. The epoxy coated transducers are then sealed with an ultra thin walled (e.g. about 0.5 to about 3 mil) tubing 32 that may for example be a heat shrink tubing such as polyester or the like. Or, the epoxy coated transducers may otherwise covered, as is known. Heat shrink tubing 32 extends beyond the area over transducers 16 and covers substantially the length of device 11. To support tubing 32 in such extended area, a filler 33 of chosen composition (preferably flexible) is placed around catheter 12 and between it and tubing 32.

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According to the present embodiments and those elsewhere herein shown and described, a cooling system may be included, which has been characterized to increase heating efficiency by about 20-30% versus non-cooled embodiments. In addition, as shown in FIG. 4, standoffs 34 may be used to support transducers 16, which are preferably flexible and may be an integral part of catheter 12.

For spinal disc therapies herein contemplated, device 11 is generally designed to be sufficiently flexible to be delivered in a substantially straight configuration through delivery device 50, and thereafter be adapted to assume a configuration appropriate for delivering energy along a length corresponding to an interface between the linear array of transducers 16 and the desired region of tissue to treat. This flexibility may be modified according to various different modes elsewhere herein described in order to achieve appropriate positioning and shape conformability used in a particular case.

In one particularly beneficial further embodiment shown variously in FIGS. 5A-B, device 11 is modified from the previous embodiment of FIGS. 3A-4 such that inner support catheter, similar to catheter 12 in FIGS. 3A-4, provides a through lumen 13 that is adapted to slideably receive a guidewire 60 therethrough. Guidewire 60 includes a stiff proximal end portion and either a shaped or shapeable, more flexible

distal end portion 62. According to this guide wire-based embodiment for system 10, guidewire 60 is adapted to be placed within the desired region of treatment by steering and advancing the shaped distal end portion 62 with manipulation of proximal end portion 61 externally of a patient's body. Device 11 along the array of treatment transducers 16 is sufficiently flexible to track over guidewire 60 in order to be positioned for treatment along the desired treatment region. Guidewire 60 may be of a shape memory type, or may be designed according to many other previous guidewire disclosures. Moreover, device 11 may be adapted to receive and track over guidewire 60 over substantially the length of device 11, also known as "overthe-wire", or may be of a "rapid exchange" or "monorail"-type wherein guidewire 60 exits proximally from device 11 at a port along device 11 that is distal to the most proximal end of device 11.

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FIG. 5A also shows the linear array of transducers 16 to be of the segmented type, wherein each linear location for a transducer 16 corresponds to two opposite transducer regions 16a,16b that may be independently or alternatively actuated for energy delivery. This allows localization of US energy along only one radial aspect surrounding device 11. Particular designs and methods incorporating linear array of segmented or partially activated US transducer regions is disclosed in US Patent No. 5,733,315 to Burdette et al., which has been previously incorporated herein above. It is to be appreciated herein that each transducer segment 16a or 16b is in effect an independently actuatable transducer, though shown and described as sub-parts of an overall transducer 16 for illustrative simplicity. Corresponding transducer regions 16a may be all actuated simultaneously along the array, without actuating the opposite regions 16b, in order to ablate along a length only along one radial aspect of device 11 corresponding to actuated regions 16a. Or, the other regions 16b may be actuated without regions 16a emitting US energy. The device of FIG. 5A provides this selectivity, which may be useful for treating different regions of a disc annulus as further discussed below.

Various different shaft structures may be appropriate for housing the corresponding functional components of a device 11 according to the embodiment of FIG. 5A, though one particular cross-section is shown in FIG. 5B for illustration. Due to the radially segmented aspect of the transducer array of this embodiment, the

number of actuatable transducer elements corresponding to a given length of the array is doubled compared to the earlier embodiment (e.g., each linear region is split into two radial regions). Therefore, twice the number of electrode leads 18 and thermocouple leads 28 must be housed. The cross-sectioned, multi-lumen tubing 12A shown in FIG. 5B therefore has a plurality of surrounding lumens 14 surrounding a central guidewire lumen 13. This is adapted to give an organized shaft structure for adapting a proximal pin connector to interface with drive and control system 40, as well as uniform flexibility in multiple planes to enhance delivery and position control in use.

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Device 11 as illustrated by the FIG. 5A embodiment may include transducers 16 having many different geometries that may be customized for a particular energy delivery profile along the transducer array. Examples include tubular (e.g. FIGS. 3A-4), sectored tubular (e.g. FIG. 5A), and also as further examples planar or plate, hemispherical, or portions of cylinders (convex), depending upon the desired energy delivery for a particular application. Further to the FIG. 5A embodiment, a distinct radial region of a transducer location along the array may be rendered completely inoperative for US delivery, such as for example in order to protect against delivery into sensitive tissues such as the spinal chord. In any event, according to the various embodiments, device 11 may be adapted for linear control of US exposure or heating via power level adjustments and angular control of US exposure or heating via directional characteristics of transducer emitters 16 (e.g., angularly directive with an inactive zone).

A further embodiment shown in FIGS. 6A-C further provides semi-hemispherically shaped transducers 16 that are essentially flipped to have an opposite radial orientation relative to the radius of device 11 as compared to otherwise similar transducer sectors 16a,b in FIG. 5A. More specifically, transducer segments 16a,b each have their concave surfaces 17 facing outwardly from device 11. This is believed to allow for a focused US signal to be emitted therefrom to a focal point or depth in relation to device 11 that is controlled by the radius of curvature R for the corresponding transducer. Such an arrangement may include two radial sides of linearly spaced transducers, e.g. 16a,b as shown in FIGS. 6A-B. Or, device 11 may be adapted to house a single such transducer segment 16, as

shown in FIG. 6C, which limits the radial emission choices to one region around the device periphery, but increases the design real estate and options for device 11 to house this unique transducer configuration. In any event, the applicator device 11 may be repositioned to control angular thermal profile. Where angular control is of important necessity (such as treatment immediately adjacent spinal column), device 11 may be designed to be in particular torqueable from outside the body in order to rotate the radially focused transmission with the shaft, such as via use of metal reinforcements or tubular members, or in general composite shaft designs such as wire or braid reinforcements.

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As elsewhere herein shown and described, device 11 along the US path from transducer 16 generally includes an ultrasonically translucent medium, such as a fluid. Acoustic gain and temperature regulation of applicator surface can help control distance of heated region and effects from applicator surface. Frequency and depth of focus can be selected to control heating pattern, and time can be varied to control heating effects and distribution.

The segmented array of FIG. 6C can use many different types of transducers 16, such as for example concave sections of cylindrical or tubular transducers or spherical or semi-spherically focused transducers. The OD of the tubes used to form the transducer shown in FIG. 6C is much larger than the diameter D for device 11 that supports the transducer 16. The transducer sectors are a small arc; thus, a line focus is produced at depth over a small arc angle, producing an intense US exposure or heating pattern which is approximately the same length as the tubular segment, but very narrow in the angular dimension, which can beneficially be as narrow as from about 1 to about 10 deg., preferably as narrow as from about 1 deg. to about 5 deg. The length and number of segments can be varied.

Referring to the particular embodiment shown in FIG. 6D, a conical or semi-spherical disc-shaped transducer 16 is shown which focuses energy not only radially along a length of the corresponding transducer, but instead focuses along the entire surface of the disc-shape. Therefore such shaped transducer more precisely and densely focuses and localizes the energy being delivered into a very small region of tissue. While an array of such shaped disc transducers 16 is contemplated, the focused pattern may create energy gaps between adjacent elements – therefore this

design may be more applicable to precise treatment in one area by one transducer, which may be followed by moving the transducer, either together with or along the supporting device 11 to another location to be treated.

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As previously mentioned above, the embodiments of FIG. 6A-D, in addition to the other embodiments contemplated herein, can cooled either externally or internally, an can be inserted within the disc or laparoscopically placed against the target tissue or directed toward the target tissue. Acoustic gain and temperature regulation of applicator surface can help control distance of heated region and effects from applicator surface. Frequency and depth of focus can be selected to control heating pattern, and time can be varied to control heating effects and distribution. Other examples of ultrasound array techniques for adjusting the selectivity of ultrasound transmission from a device are disclosed in US 5,391,197, which has been previously incorporated by reference above; the various embodiments of that disclosure are contemplated in combination with this disclosure with respect to segmented array of US transducers, where appropriate and as modified according to this disclosure according to one of ordinary skill.

As elsewhere provided herein, the illustrative embodiments and procedures have applications in many different soft/hard tissue sites and body parts where ultrasound exposure, high temperature, low temperature or combination of effects are desired. Joints in particular are locations where the present invention is well suited for providing therapy. However, as stated above of particular benefit is use of the present invention for treating intervertebral discs.

Therefore, one example of a method for treating an intervertebral disc according to the present invention is provided according to various sequential modes of use shown in FIGS. 7A-F. This procedure for the purpose of illustration more specifically shows posterior-lateral approach to treating a posterior wall of an intervertebral disc from a location within the nucleus. However, because selectivity in disc treatment is so very important, clearly other procedures are contemplated as will be further developed through other examples below.

More specifically, according to FIG. 7A a posterior wall 4 of annulus 2 is observed to require thermal treatment, either due to physical damage to the annulus 2 structure (e.g. herniation), or otherwise, such as for example innervation with

unwanted nervous tissue causing pain or other inflammatory cells (which may be directly or indirectly related to disc damage such as herniation). As shown in FIG. 7A, a sharp, pointed tip 51 of a spinal needle 50 is used in a posterior-lateral approach to puncture through the posterior-lateral region of the wall of annulus 2. This gives lumenal access through needle bore 53 into the nucleus 3 for ultrasound probe delivery.

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As shown in FIG. 7B, a guidewire 60 having a steerable distal tip 62 is then advanced through needle bore 53 and into nucleus 3. It is contemplated that guidewire 60 may be used for many purposes within the nucleus 3, or otherwise in or around disc 1. However, one particularly beneficial use is shown, wherein guidewire is tracked along proximal wall 4, and then further around the more severe radius along lateral wall 7. This gives a rail along proximal wall 4 along which a highly flexible device 11 may track for ablation there. This is shown in FIG. 7C, wherein ultrasound transducers 16 are of length, size, and location along device 11 such that they are positioned over guidewire 60 to coincide with the area along posterior wall 4 to be treated. According to the embodiment shown in FIG. 7D, only one radial aspect of device 11 is actuated for US emission and treatment, which is shown to be transducer segments 16b in an array on one radial aspect of device 11 interfaced with or facing proximal wall 4. Thus US energy is transmitted into wall 4 to treat that region without substantial treatment elsewhere.

After treatment, other regions may be treated by further manipulating guidewire 60 and/or device 11 within the nucleus 3 (or outside of annulus if desired). Once treatment within the annulus is completed, device 11 may be withdrawn. In the embodiments shown in FIGS. 7E-F, the ultrasound transducers 16 may be used to assist in closing the wound formed at entry site C through annulus, such as by elevating the temperature in that area sufficient to cause collagen shrinkage to aid in closing that aperture, as shown in FIG. 7F for example. In this application, the entire circumference of device 11 may be activated for US emission, or if only a radial region is adapted for such emission, it may be rotated to heat all aspects of the wound aperture to seal it. In addition, a sealant may be administered through the distal end of device 11 to close the wound, such as at introduction region C, either instead of or in conjunction with ultrasonic emission from device 11.

Other regions of disc 1 may also require localized, selective therapy with US, and the present invention allows for highly specialized treatments in the various regions. FIGS. 8 and 9 for example show use of device 11 from the right posterior-lateral approach through site C shown in FIGS. 7A-F, but for treating anterior wall 5 and left lateral wall 7 regions, respectively. For the embodiments shown wherein device 11 is highly flexible for sufficient trackability over guidewire 60, the same device may be used for treating the very different areas shown in FIGS. 7A-F, FIG.8 (anterior wall), and FIG. 9 (left lateral wall). However, as shown in comparing FIGS. 7A-F, 8, and 9, different radial regions of device 11 may be activated to treat the respective interfacing wall (where angular specificity is provided, which may not be required though often preferred). In particular, the guidewire trackability of the present embodiments provides for highly beneficial flexibility between the ability to treat these very different segments, in particular in view of the generally more significant curvature associated with the anatomy around the lateral wall regions as shown at radius R for lateral wall region 7 shown in FIG. 9.

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For the purpose of further illustration, FIG. 10 shows a device 11 according to the invention used to treat a posterior wall 4 of disc 1, but according to an anterior approach through a region of anterior wall 5 adjacent to left lateral wall 7. Again, this may for example be a similar device as that shown and described with respect to FIGS. 7A-9.

The devices and methods of the invention are also adapted for use in treating spinal disorders from outside of the annulus 2, though preferably still from an invasive location within the patient's body in order to provide the necessary and desired amounts of energy at only the highly localized, target locations. For example, FIG. 11 shows a right posterior-lateral approach to treat a region of annulus 2 from outside of disc 1. Because of surrounding tissues, it is highly desired (though not always required), to deliver only highly selective, directed US energy into only the region of annulus 2 being treated. In particular, truncal nerves often extend along such areas, as well as the spinal cord being located not to far away from such region. Therefore, a controllable, selective array of transducers as shown at transducers 16a radiates only toward the disc annulus 2, whereas the opposite side of device 11 is non-emissive. In fact, this opposite side may be selectively cooled to

prevent from thermal heating of the area, and may include sensors to monitor such temperature around that area opposite the active US treatment zone, such as for example at thermocouples 27 shown in FIG. 11.

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For the purpose of further illustration, FIG. 12 also shows US treatment from outside of a disc 1 according to the invention, but according to an anterior approach. FIG. 13 shows still another exterior treatment modality, however this particular location along the posterior wall 4 is particularly sensitive as the spinal cord is located immediately adjacent device 11 opposite transducers 16a and must not be harmed. Therefore, not only the radial emission of energy (either US or thermal heat) from device 11 must be insulated from that radial region corresponding to the spinal cord.

Though guidewire tracking mechanisms provide the illustrative embodiments for positioning in FIGS. 7A-13, other embodiments are contemplated. Moreover, positioning of a device 11 may include simultaneous or sequential positioning of thermometry probes for monitoring of sensitive tissue areas, etc.

A pre-shaped or otherwise directional introduction/delivery device may assist to point a device 11 to a localized area for treatment, such as shown for example in shadow in FIG. 3A for shaped tip 51 for delivery device 50. Such directionality from the delivery device 50 may be provided in addition to, or in the alternative to, providing guidewire tracking of device 11 or other additional positioning modes herein discussed. In addition, other positioning control mechanisms may be incorporated into device 11 itself as follows.

One particular deflectable tip design is shown for device 11 in FIG. 14A-B. According to this embodiment, the region of device 11 that includes the array of transducers 16 is deflectable to take a variety of shapes, such as shown in right and left deflection modes around a radius r in FIG. 14B. Such deflection may be achieved using conventional deflection mechanisms, such as for example using an arrangement of one or more pull-wires integrated into the tip region so that tension causes catheter deflection. In addition, such deflection may be achieved instead of along a length of an arc, around a pivot point, as shown in FIG. 14C. Such may be achieved in order to achieve different angles for surgical approaches into the desired treatment areas, such as in or around intervertebral discs. The transducer elements

may be along a deflectable segment that is either round, or may be more planar as desired.

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Pre-shaped distal regions for device 11 may also provide for desired treatment of highly unique anatomies. A kit of devices, each having a particular shape is contemplated. Such shapes may be integrated in procedures with or without conjunctively using guidewire tracking. For example, FIG. 15A shows a pre-shaped distal end for device 11 having a simple distal curve around radius r. The transducers 16 are around the outer radius of such shaped end, and therefore this shape and orientation is suitable for example for treating an anterior wall 5 from within a disc 1 via a posterior-lateral approach, such as for example according to FIG. 8. A similar shaped end is shown in FIG. 15B, but the transducers 16 are instead on the inside radius r. This is more suitable for posterior-lateral approach with posterior wall treatment, such as for example in FIGS. 7A-F.

A further beneficial shape and orientation is shown in FIG. 16. Here, an acute bend shown around radius r2 is adapted to correspond to the more drastically rounded lateral wall regions 6,7 of a disc annulus 2 with an energy emission region on the outside of that bend. An additional bend region may be highly beneficial, though not always required, shown proximally of the distal bend around radius r2 and having a less drastic bend, in the opposite direction of r2, shown around radius r2. This configuration is highly beneficial for treating lateral wall regions 6,7 from a posterior-lateral approach (e.g. FIG. 9), though may be used in the same configuration or slightly modified for anterior approach.

Though ultrasound transducers and their many benefits for invasive energy delivery into tissues has been extensively herein described, various of the embodiments further contemplate use with other energy sources or treatment modalities, either instead of or in conjunction with ultrasound. Thus, treatment region 16 in FIG. 16 does not specifically show individual ultrasound segments as in the other figures, for the purpose of illustrating other energy sources or treatment modalities that my be incorporated thereon and still gain the benefit of the unique shape provided for inner disc ablation according to that figure. Other sources such as electrical (e.g. RF), light (e.g. laser), microwave, or plasma ion may be used. In addition, cryotherapy or chemical delivery may be achieved along the regions

variously designated as "transducer 16", which may be accompanied by other modifications corresponding therewith, without departing from the scope contemplated by the embodiments.

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According to the various deflectable or pre-shaped modes, or modes where energy delivery is limited to only one side of the device, the device 11 is preferably torqueable, such as by integrating into the shaft design a composite of braided fibers or other stiff members. This allows for more precise control of the distal tip regions as it deflects or takes its shape along a plane within the desired area of the body to treat.

The various embodiments for device 11 above may be adapted to incorporate active cooling, such as circulating cooling fluids within or around active energy emitting elements such as transducers 16 variously shown or described. Such cooling may be integrated into the particular device 11, or may be achieved by interfacing the particular device 11 inside of or otherwise with another device.

FIG. 17A shows for example device 11 within an outer jacket 15 that may or may not be distendable, as shown in shadow at 15'. Outer jacket 15 is adapted to circulate fluids around transducers 16 and therefore is interfaced with a circulation pump 70 in an overall system. In an alternative embodiment sharing many common features as FIG. 17A, the device 11 shown in FIG. 17A provides for the cooling fluid to be delivered through an interior passageway of the interior ultrasound device, out the distal tip thereof within the outer jacket 15, and back over the outer surface of the interior device including the transducers 16.

In either the FIG. 17A or 17B embodiments, device 11 may be fixed within outer jacket 15, or may be moveable relative to outer jacket 15, or visa versa. Fluids provided in an outer jacket surrounding transducers 16 according to the invention may also be used beneficially for ultrasound coupling to intended tissues to be treated. This may be in addition to or instead of being used for cooling. In particular, such coupling fluids may be provided in a jacket 15 that is conformable, such that irregular surfaces to be treated receive uniform energy coupling from the assembly. Or, pre-shaped, and symmetric or asymmetric shapes, may be provided as appropriate to provide such coupling. Ultrasound coupling may be further achieved by providing a non-liquid coupling member as a stand-off over a transducer in order

to couple that transducer to the tissue – such as for example a sonolucent coupling gel pad, etc.

In addition to the various designs for device 11 described above for achieving positioning, e.g. guidewire tracking, pre-shaped, or deflectable, other mechanisms may also be incorporated for accurate positioning. For example, stiff or flexible distendable member(s) may be incorporated on device 11, e.g. a balloon or expandable cage, that distends to a predetermined-shaped (or just generally distends). This may help positioning, such as for example where the nucleus 3 is void of pulposus in order to position the transducers 16 within a balloon at a desired location within the annulus 2. In addition to positioning, such a member may also be used to aid coupling, tissue deforming, and tissue repositioning during a treatment procedure.

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As previously discussed, the intervertebral disc applications of ultrasound herein contemplated require high selectivity for US or otherwise thermal therapy due to the presence of highly sensitive, non-targeted tissues in close proximity (e.g. spinal cord and other nerves). Therefore, though heat conduction may not be the intended mode of therapy with transducers 16, their concomitant heating during US sonic wave delivery may cause unwanted damage in either the targeted or nontargeted tissues. Accordingly, cooled lumens or balloons over the transducers may be employed to protect such tissues from such heat, or directivity of the ultrasound per the embodiments herein described my adequately protect sensitive non-targeted tissue. In the case of an active cooling mechanism, it is to be appreciated that such mechanism may be integrated directly onto device 11 that carries the transducers 16. Or, a separate co-operating device such as an outer sleeve carrying cooling fluids may be used. Such cooling chamber may be on the side of the transducer delivering the targeted US wave, in which case fluid in the chamber must be substantially sonolucent for efficient energy delivery. In the event the cooling is intended to protect a "back side" of the device only, other fluids may be used.

Applicators, such as the various embodiments shown for device 11 among the FIGS., and insertion tools, e.g. delivery device 50, may be adapted to be MR compatible for real time monitoring of a particular procedure. Also other imaging modalities may be used instead, or in conjunction with one another, in order to

control and optimize the US treatment procedure, including for example for monitoring positioning, temperature, lesion assessment, coagulation, or otherwise changes in tissue structures related to the treatment (e.g. targeted tissue to be heated or adjacent tissues to monitor safety, such as regions of concern to preserve nerves associated with the spinal chord). In fact, US itself is an energy source that has been widely used for acoustic imaging in and around internal body structures. It is contemplated that imaging US devices may be incorporated into a device 11 directly, or indirectly incorporated as a separate cooperating device in system 10, and further that the US treatment transducers 16 herein shown and described may be operated in imaging modes before, during, or after thermal US therapy is performed with those same transducers 16.

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In addition to the spine, the device systems and methods according to the embodiments may be used in other regions of the body, in particular other joints. Examples of such regions include knee, ankle, hip, shoulder, elbow, wrist, knuckles, spinal processes, etc. In such case, further modifications from the illustrative embodiments herein provided may be made in order to accommodate the unique anatomy and target tissue regions, without departing from the spirit and scope of the present invention.

While the device systems and methods have been herein described with respect to treating tissue via US exposure in order to provide hyperthermia effects, other non-thermal results may also be intended, either in conjunction with hyperthermia or in the alternative to. For example, drug activation and or enhanced drug delivery, such as for example via enhanced dispersion or cellular permeability or uptake, may be achieved by delivering certain specific therapeutic dosing of US energy, as has been well studied and characterized in the art. Such methods may for example aid in the treatment for example of arthritis in joints, etc.

The invention as described herein according to the particular embodiments is highly beneficial for treatment of the body, in particular joints, and in particular the spine. In general, these devices and methods are adapted for such treatment invasively from within the body. However, external applications are contemplated as well. In addition, treating living bodies according to the invention is believed to provide a highly therapeutic result for improved living. Nevertheless, use of the

devices and methods as described herein are also contemplated for conducting scientific studies, in particular with respect to characterizing tissues in their relation to applied energy. Therefore, "therapeutic" applications may include those sufficient to induce a measurable change in tissue structure or function, whether living or postmortem, prophylactic or ameliorative, research or clinical applications.

Example:

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External Directional Ultrasound Thermal Therapy of Cadaver Spinal Discs

FIG. 18A shows an X-ray photograph of a cadaver intervertebral disc during invasive ultrasound treatment with a device and method according to the invention as follows. Temperature monitoring measurements are shown as overlay dotted lines and numbers over the X-ray.

An ultrasound probe was provided as follows. Two PZT ultrasound transducers were provided on a hypotube, each being 1.5mm OD x 10mm long (0.012" wall thickness), and being spaced by about 1mm. The ultrasound probe was inserted within a 13-g Brachytherapy Implant Catheter having a 2.4mm O.D., which is commercially available from Best Industries. Water at room temperature was circulated through the outer catheter and over the transducers at about 40ml/min during ultrasound transmission. The assembly of the outer catheter with inner transducers and probe was inserted laterally into a cadaver disc along the border of the nucleus pulposus and posterior wall of the annulus fibrosus. The approximate location of the transducers is shown in two rectangles in FIG. 18A. Thermocouple probes were inserted into the disc as shown in the X-ray, and with measurement locations reflected by the sample measurements in the overlay. The above test sample and instrumentation was placed within a 37degC water bath during testing. Each transducer was run at approximately 10W power, wherein temperature numbers shown in FIG. 18 generally represent temperatures at substantially steady state after actuating the transducers. Heat generated by the ultrasound probe was sufficient to cause therapeutic effects in surrounding tissues of the annulus and nucleus.

Another similar study was performed using ultrasound to heat a post-mortem intervertebral cadaver disc using a curvilinear ultrasound applicator directly coupled to tissue at 5.4 MHz and 10W power. A temperature vs. time graph of the results at

varied depths from the transducer surface are shown in FIG. 18B, which shows among other information that temperatures reached 70-85 degrees within 7mm from the disc treatment surface and within 5 minutes of treatment..

As shown in the graph of FIG. 18B, the elevated temperature achieved at 1 mm from the ultrasound transducer reached: over 45 degrees C within 90 seconds (1 ½ minutes); over 70 degrees C within 120 seconds (2 minutes); over 75 degrees C in nearly 120 seconds; over 80 degrees C within 180 seconds (3 minutes); and over 85 degrees C by about 300 seconds (5 minutes).

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Temperatures at 4 mm depth from the transducer reached: 45 degrees C in less than 120 seconds (2 minutes); 55 degrees in close to about 120 seconds; 65 degrees C within 150 seconds (2 ½ minutes); over 70 degrees C within less than 210 seconds (3 ½ minutes); and over 75 degrees C and still rising by about 240 seconds (4 minutes).

Temperatures at 7 mm depth from the transducer reached: 45 degrees C by about 120 seconds (2 minutes); 55 degrees C by about 150 seconds (2 ½ minutes); 60 degrees C in nearly 180 seconds (3 minutes); 65 degrees C within 240 seconds (4 minutes); and 70 degrees C within 300 seconds (5 minutes).

Temperatures at 10 mm depth from the transducer reached: 45 degrees C in less than 150 seconds (2 ½ minutes); 55 degrees C in less than about 210 seconds (3 ½ minutes); over 60 degrees C in less than 270 seconds (4 ½ minutes); and slightly less than about 65 degrees C by 300 seconds (5 minutes).

In another regard, the graph in FIG. 18B also shows that temperatures above 45 degrees C were reached within 90 seconds at 1mm, 120 seconds at 4mm and 7mm, and 150 seconds at 10mm depths from the transducer. Similarly, temperatures of at least 55 degrees C were reached within about 120 seconds at 1mm and 4mm, 150 seconds at 7mm, and 210 seconds at 10mm depths.

Temperatures of at least 65 degrees C were reached within less than 120 seconds at 1 mm, 150 seconds at 4mm, 240 seconds at 7mm depths. Still further, temperatures above 70 degrees C were reached within 120 seconds at 1mm, 210 seconds at 4 mm, and 300 seconds at 7mm depths. Even further heating to above 75 degrees C were reached within close to 120 seconds at 1mm, and 240 seconds at 4 mm depths.

Further observation of FIG. 18B in the time domain, in less than 150 seconds temperatures at up to 7mm depth from the transducer reached at least 55 degrees C. Within about 210 seconds temperatures in tissue as deep as 4 mm deep reached over 70 degrees C, up to 7mm deep reached at least 60 degrees C, and up to 10 mm deep reached over 55 degrees C. Within 270 seconds, temperatures 4 mm deep reached over 75 degrees C, up to 7mm deep reached over 65 degrees C, and up to 10 mm deep reached over 60 degrees C. In still a further regard, by 300 seconds temperatures up to 7mm deep reached at least 70 degrees C, and up to 10mm deep reached almost 65 degrees C.

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Upon further comparison of temperatures vs. depth according to the FIG. 18B graph: temperatures over 60 degrees C (and for the most part up to 65 degrees or more) were achievable up to 10 mm deep; temperatures up to at least 70 degrees C were achieved up to 7mm deep; temperatures over 75 degrees were achieved up to at least 4mm deep; and at 1 mm depth temperatures of over 80 degrees and even 85 degrees were observed.

As will be further developed below and elsewhere herein, such elevated heating, including at tissues as deep as 4mm, 7mm, and in some regards even 10mm, is a highly beneficial aspect of the present invention. For example, other more conventional intervertebral disc heating devices, in particular the "IDTT" device elsewhere herein described, have been observed to be limited as to the extent and depth of heating possible.

For example, according to at least one study observing the heating effects of the "IDTT" radiofrequency electrical heating device (elsewhere herein described) also on cadaveric lumbar spine disc samples, the following observations were made. During intended modes of use for internal disc heating, and over treatment times of 17 minutes (1020 seconds), the IDTT devices tested were able to heat only the closest 1-2mm of intervertebral disc tissue to temperatures just barely exceeding 60 degrees, with no tissue of 1mm depth or greater exceeding 65 degrees C despite reaching 90 degrees C on the probe itself. Moreover, only tissues within a 7mm radius of the heating probe exceeded 48 degrees C during the 17 minute treatment time. Still further thermal dosing was limited such that the maximum predicted depth for damaging nociceptive fibers infiltrating the discs was believed to be only within a

6-7mm radius.

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Accordingly, substantial benefit is gained by using the ultrasound treatment device of the present invention to the extent depth of heating and heating to substantial temperatures and within reasonable times is desired.

Example: Thermal Therapy of Pre-Stressed Spinal Joints

This Example provides an abstract summary, introduction, methods, results, and conclusions with respect to a certain group of studies performed to evaluate heat-induced changes observed in intervertebral discs, related structures such as in particular annulus fibrosus, and the related biomechanics, in particular with respect to "intact" discs, as follows.

1. Abstract.

The intervertebral disc is considered a principal pain generator for a substantial number of patients with low back pain. Thermal therapy has been disclosed to have a healing effect on other collagenous tissues, and has been incorporated into various minimally invasive treatments intended to treat back pain. Since the therapeutic mechanisms of thermal therapy have generally been previously unknown, proper dosage and patient selection has been difficult. Thermal therapy in one regard has been disclosed to acutely kill cells and denature and deinnervate tissue, leading to a healing response.

The purpose of this study was to quantify the acute biomechanical changes to the intact annulus fibrosus after treatment at a range of thermal exposures and to correlate these results with the denaturation of annular tissue. Intact annulus fibrosus from porcine lumbar spines was tested *ex vivo*. Changes in biomechanical properties, microstructure, denaturation temperature, and enthalpy of denaturation before and after hydrothermal heat treatment (at 37, 50, 60, 65, 70, 75, 80, and 85°C) were determined. Shrinkage of excised annular tissue was also measured after treatment at 85°C. Significant biomechanical changes in the intact annulus were observed after treatment at 70°C and above, but the effects were much smaller in magnitude than those observed in excised tissues. Histological and mDSC data indicated that denaturation had occurred in intact annular tissue treated to 85°C for 15 minutes, though such effect was observed to be slight. It is believed based on observations made that constraints imposed on the tissue by the joint structure

retard changes in properties. These findings have implications for dosing regimens when thermally treating disc tissue.

2. Introduction.

The goals of this study were to: 1) quantify acute biomechanical changes to the intact annulus fibrosus induced by a broad range of *ex vivo* thermal exposures; and 2) to correlate these results with denaturation of annular tissue using modulated differential scanning calorimetry (mDSC) and histological data.

3. Methods.

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a. Mechanical Testing.

Forty-one spinal motion segments (18 L₁₂, 19 L₃₄, 19 L₅₆) consisting of the intervertebral disc (IVD) and each adjacent vertebral body were cut from 22 fresh frozen porcine lumbar spines (domestic farm pig weight range: 115-135 lbs). Muscular and ligamentous structures, facet joints, transverse processes, and posterior elements were dissected from the vertebral bodies to isolate the disc. Saline-soaked gauze was wrapped around the discs during preparation to minimize dehydration. Next, the nucleus was depressurized by drilling holes first through the vertebral bodies to the center of the nucleus in the superior-inferior direction, and then from the anterior faces of the vertebral bodies to the central hole. Plastic tubing was inserted into the anterior openings and affixed with cyanoacrylate. The vertebral bodies, anchored with 2.5mm threaded rod, were embedded into fixation cups using polymethylmethacrylate (PMMA). An alignment bar mated with grooves in the fixation cups to ensure that the plane of the disc remained normal to the vertical loading axis. X-rays (Faxitron Cabinet X-Ray System, Hewlett-Packard, McMinnville, OR) were taken of the specimens in the dorsal-ventral plane after equilibration in a 37°C saline bath. Disc heights were determined by averaging three caliper readings from the dorsal-ventral x-rays.

Specimens were secured in fixation cups, mounted into a hydraulic materials testing machine (MTS Bionix 858, Eden Prairie, MN), and placed into a temperature controlled 0.15M saline bath at 37°C to equilibrate. Saline at bath temperature was also circulated through the center of the discs via the tubing attached to the vertebral bodies; this allowed for a more rapid and uniform heat distribution within the annulus.

Temperatures were measured using two stainless steel thermocouple needle probes, one placed in the bath, and one inserted approximately halfway into the anterior annular wall. These fine-needle temperature probes were fabricated inhouse using 25 micron constantan-manganin thermocouple junctions embedded within a 30 gauge (0.30 mm OD) needle. Superior-inferior x-rays were used to verify proper placement of the annular temperature probe.

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The testing protocol consisted of a 20-minute thermal equilibration at 37°C, a 15-minute heat treatment, and another 20-minute equilibration at 37°C. Fast temperature changes were facilitated by exchanging the saline in the bath with that in a reservoir heated to the desired temperature and then maintained with temperature-controlled circulation. The target temperature (to within 7%) was reached within 5 minutes of exchanging the saline. During the equilibrations, the disc stress was maintained at 0 kPa.

Mechanical testing was performed at 37°C just prior to heat treatment and again subsequent to heat treatment and re-equilibration at 37°C. Testing consisted of nine preconditioning cycles in axial tension-compression (-25 to +150N at 0.25 Hz), followed by one testing cycle to the same limits. The applied load was measured using a precision force transducer (Load Cell 662, MTS, Eden Prairie, MN), and the deformation of the disc was assumed to be the change in distance between fixtures, measured using the test system LVDT. Data was collected every 0.01 seconds during mechanical testing and every 15 seconds during heat treatment and equilibration. Heat treatment was to one of the following temperatures: 37 (Controls), 50, 60, 65, 70, 75, 80, or 85°C. Five specimens were tested at each treatment temperature except for the 60°C group that had six specimens.

After testing, specimens were removed and the discs were cut in the transverse plane and scanned at a resolution of 600 dpi (CanoScan N656U, Canon, Inc., Costa Mesa, CA). Annulus areas were measured using imaging software (Scion Image, v. 4.0.2B, Frederick, MD).

Two additional experiments were conducted to allow us to explore the limits of annular thermal response. In the first study, a specimen was prepared as described above and treated at 85°C until the thermal contraction stabilized (within 0.01mm). For the second study, sections of anterolateral annulus were excised from five

lumbar discs (2 L₂₃, 3 L₄₅) from four different spines and treated at 85°C using the same heating protocol as above. X-rays were taken before and after treatment, and changes in circumferential and radial dimensions after heat treatment were measured using digital calipers.

b. Microstructure

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Tissue samples were excised from 37°C and 85°C mechanical test specimens, and from an excised specimen treated at 85°C. Samples were embedded in paraffin, sectioned in the circumferential plane at 6 microns, and stained with HBQ (Hall, 1986). The sections were imaged on a Nikon Eclipse E800 microscope (Nikon, Melville, NY) under bright field to examine tissue structure, and under polarized light to assess collagen birefringence.

c. Modulated Differential Scanning Calorimetry

Traditional DSC measures the combined effects of reversible and nonreversible heat flow, but the two components can be measured separately if the modulated DSC (mDSC) technique is used. mDSC was performed on samples of anterolateral annulus fibrosus removed from fifteen previously treated specimens (Cambridge Polymer Group, Somerville, MA). Punches (approximately 10mg) were removed from the control (37°C) mechanical test specimens (n=5), mechanical test specimens treated at 85°C (n=5), and from the excised annular specimens treated at 85°C (n=5). Each sample was placed in 0.1% NaCl solution for 20 minutes, blotted, weighed, and crimped into an aluminum anodized hermetic DSC pan. Samples were placed into a Q1000 differential scanning calorimeter (TA Instruments, New Castle, DE), equilibrated at 55°C, and then ramped from 55°C to 95°C at 0.5°C/min. Using an empty pan as a reference, total enthalpy of denaturation (ΔH) and the temperature corresponding to the nonreversible endothermic peak (T_m) were recorded. Following the mDSC procedure, samples were vacuum dehydrated, and the fractional dry mass (ratio of dry weight to wet weight) was recorded.

4. Data Analysis

The force and displacement data from the mechanical tests were converted to stress and strain. The stress and strain data for each mechanical test were then fit to a high-order polynomial, and an equation for the specimen tangent modulus was calculated as the derivative of this polynomial. A plot of modulus vs. applied stress

was constructed. The stress at the inflection point - the transition between tension and compression - was the stress at which the second derivative of the polynomial was zero. The reference configuration was defined as the stress and strain at the pre-treatment inflection point. Three biomechanical parameters were calculated from the modulus vs. applied stress curves to quantify heat-induced changes in the mechanical response (Fig. 19): the change in modulus at the inflection point (MI), the change in modulus at 150kPa (M150), and the change in residual stress at inflection point (RSI). The percent change in hysteresis (HYST%) was calculated from the preand post-treatment load-displacement curves. The change in the modulus at the inflection point is a measure of the increase or decrease in the stability of the joint, while the change in the modulus at 150 kPa is an indication of how well the joint will withstand physiologic loading. The percent change in strain at 0 stress (E0%) was used to quantify axial shrinkage of the tissue.

Differences in each parameter with treatment temperature were compared using a one-way analysis of variance (ANOVA). *Post-hoc* multiple pairwise comparison tests (Fisher's Least Significant Difference) were performed to determine differences between treatment groups with a significance of p<0.05.

5. Results.

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a. Mechanical Testing

FIG. 21 includes various graphs representing observed biomechanical parameters after varied heat treatments according to the present study as follows: graph (a) represents change in modulus at the inflection point (MI); graph (b) represents change in modulus at 150 kPa (M150), graph (c) represents percent change in strain at 0 stress (E0%), graph (d) represents change in residual stress at the inflection point (RSI), and graph (e) shows percent change in hysteresis (HYST%). Further to the graphs in FIG. 21, the reference letter "X" is used to designate where data is significantly different from 37°C group, p<0.05; whereas the symbol "+" is used to designate where data is observed to be significantly different from the 85°C group, p<0.05.

Significant differences between the control group and the heat-treated specimens were observed at temperatures of 70°C and above (Figs. 19, 20, 21). The variation increased with increasing treatment temperature. No significant

changes were observed between the control group and the 50 and 60°C groups, and the 65°C treatment group showed a change only in the hysteresis parameter. The modulus at the inflection point (MI) increased by 152 kPa after treatment at 70°C (p<0.05), and continued to increase with increasing heat treatment: the 85°C group, with an average increase of 343 kPa after treatment as shown in Fig. 19), and was significantly different from both the control group (p<0.001) and the 70°C group (p<0.05) according to the graph in Fig. 21a. The modulus at 150 kPa (M150) significantly decreased for groups treated at 70°C and up, but did not continue to decrease with increasing treatment temperature; the decrease was 17% at 70°C and 18% at 85°C as shown in the graphs of Figs. 19, 21b.

Relative to the control group, significant axial shrinkage (E0%) was first observed at the 70°C treatment temperature. There was no significant difference observed in this particular experiment between the axial shrinkage after treatment at 70 and 85°C, although there was a trend towards continued increase (p<0.10) according to the graphical results in Fig 21c. The change in the residual stress at the inflection point (RSI) after heat treatment was significantly larger for groups treated at temperatures of 75°C and up with a trend towards increasing stress with increasing treatment temperature (75 vs. 85°C, p=0.084), as shown in the graph of Fig. 21d. A 36% percent increase in hysteresis (HYST%) was observed for the 65°C group; this was significantly larger than that of the control group (p<0.05) per the Fig. 21e graph.

The disc heights of the specimen exposed to long heat treatment time at 85°C stabilized after approximately 2.5 hours. As a result of treatment, M150 decreased 47%, MI increased 625 kPa, RSI was 47.3 kPa, and hysteresis increased 98%. The percent change in strain at 0 stress (E0%) was 22.5%.

Heat treatment of the excised annulus at 85° C resulted in shrinkage of 45.1% \pm 5.5% in the circumferential direction and expansion of $56.9\% \pm 25.4\%$ in the radial direction. The shrinkage was accompanied by a color change from white to translucent, a finding that which was not present in our whole-disc samples.

b. Microstructure

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The structure of the annular collagen, as indicated by its birefringence under polarized light microscopy, varied with heat treatment (Figs 22a-f). The structure of the excised sample treated at 85°C changed dramatically relative to that of the

control specimen: the 37°C specimen was strongly birefringent under polarized light as shown in Fig 22b, while the 85 °C excised specimen showed no birefringence as shown in Fig 22f. The 85°C mechanical test specimen appeared less birefringent than the control, as shown in Fig. 22d. Bright light microscopy revealed a structure consistent with that observed under polarized light. The heated excised specimen exhibited a homogenous morphology, as shown in Fig. 22e, with a complete loss of the original structure relative to the control shown in Fig. 22a. Tissue organization decreased, but was not absent, in the 85°C mechanical test specimen shown in Fig. 22c.

c. Modulated Differential Scanning Calorimetry

The excised specimens did not exhibit an endothermic peak, and thus, values for T_m and ΔH were not calculable. Both intact groups exhibited a full and clear endothermic denaturation event. There were no significant differences in T_m and ΔH between the intact (37°C & 85°C) specimens. T_m for the control group and the 85°C intact group were 65.4 ± 1.5°C and 65.3 ± 0.9°C respectively, while ΔH was 11.5 ± 2.4 W/g and 12.2 ± 4.6 W/g. The fractional dry mass of the 85°C intact group (0.33 ± 0.03) and the 85°C excised group (0.37 ± 0.043) were both significantly higher than the control group (0.26 ± .04; p<0.05 and p<0.01, respectively).

6. Discussion

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In this study we examined the acute biomechanical effects of thermal treatment on the annulus fibrosus. The data demonstrate that treatment for 15 minutes at 70°C or above is required to produce statistically significant biomechanical modification of the intact motion segment *ex vivo*. Heat treatments of 70°C and higher resulted in stiffening of the annulus at low loads (i.e. in the 'toe' region, parameter MI) and a decrease in stiffness at higher applied loads (M150).

These results suggest that thermal therapy at temperatures 70°C and greater leads to a more stable transition from flexion to extension. The depressurization we performed during specimen preparation created a neutral zone at the transition between tension and compression, within which small changes in force resulted in relatively large changes in displacement. After treatment at higher temperatures, this neutral zone was reduced or eliminated, as reflected in the graph shown in Fig. 20. It is further believed that the experimental model was illustrative of, and that this

response would similarly affect, the neutral zone of intact discs. By contrast, our observation that heat treatment at temperatures greater than 70°C softens the annulus at higher forces suggests that the acutely treated intact disc may be at increased risk of injury when brought to its range-of-motion limits. This confirms, in the unique setting of vertebral discs, similar behavior that has been noted in the shoulder capsule, where heat has been observed to both acutely shrink and decrease the linear-region stiffness of the joint. As a result of such prior observations in that setting, clinical practitioners have recommended joint protection for six to twelve weeks after treatment.

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While the trends in our data are comparable to those reported for other tissues such as the shoulder capsule, the magnitude of the annular treatment effect in intact tissue is smaller. For instance, while shoulder capsule contraction has been reported in at least one study to be 60% after 80°C treatment, we observed annular contraction of only 7.8% (E0%) after heating the intact disc to 85°C. Similarly, shoulder capsule stiffness reductions at high loads were much greater than those observed in the intact vertebral discs: we observed stiffness decreases of 20% (M150), while shoulder capsule stiffness decreases were on the order of 50%. It is believed that these differences are likely due to either the unique joint structure or the fiber orientation of the intact annulus, or both. The shoulder data was derived from experiments in which the capsule, a linearly oriented collagenous tissue, was cut into strips along the collagen fiber direction before testing. In contrast, intact annular collagen is oriented in two directions at ± 65° to the spinal axis, and it is highly constrained both axially, by the adjacent vertebrae, and circumferentially, by its annular structure. When the in situ constraints on the annulus were removed by excising the tissue before heat treatment, we observed a 45% circumferential shrinkage, which is similar in magnitude to that reported for linear collagenous tissues. It is believed, therefore, based upon our observations, that in situ tissue constraint, rather than fiber orientation, may be the dominant mechanism responsible for the observed differences.

Our conclusion that *in situ* tissue constraint reduces the effects of thermal therapy on the annulus fibrosus, though not previously known or confirmed prior to this study, is further supported by results observed in several other previously

reported studies. In one previous report, for example, only 6.6% shrinkage was observed in the patellar tendon, a linearly oriented collagenous tissue, after *in situ* treatment with laser energy. This difference was attributed to constraints imposed by the intact joint. Similarly, a number of other studies have been reported examining heat-induced changes in the mechanics of chordae tendineae. Tissue stress was observed to have a retarding effect: when tissue was stressed during heating, increases in the temperature, the heating time, or both, were required to achieve effects noted for unstressed tissue in these studies.

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The mechanism by which tissue stress retards thermal denaturation has a thermodynamic basis. Tensile stress straightens tissue collagen and decreases configurational entropy, which in turn, increases the activation energy required for thermal denaturation. This retarding effect was clearly evident in intact annulus, where we observed that several hours of thermal treatment at 85°C were required to achieve maximum contraction. In contrast, at least two groups of prior researchers examining excised collagenous tissues achieved maximum contraction within 5 minutes. Also, while M150 for an intact specimen treated at 85°C for 15 minutes was only 18%, the decrease in stiffness (47%) after several hours of treatment at 85°C was comparable to that elsewhere reported for excised shoulder capsule tissue.

Our polarized light microscopy data provides further evidence that tissue constraint effects both the temperature and time required to achieve a given amount of thermal damage. Collagen birefringence disappeared completely after heat treatment for 15 minutes at 85°C in the unconstrained specimen, but it remained in the intact treated annulus. Clearly 15 minutes of treatment was not sufficient to fully denature the intact annular tissue. While it was not possible to quantify the degree of birefringence in the intact tissue after treatment at 85°C relative to that at 37°C with only one specimen, it appears that that the treated specimen was less birefringent than the control. These observations are consistent with the results of our mechanical tests.

Differences in the mechanical behavior of the intact annulus after treatment at temperatures greater than 70°C indicate that the tissue underwent a thermally mediated change. However the results of the mDSC experiments indicate that tissue constraint prevented significant collagen denaturation: the main denaturation peak

and enthalpy of denaturation of the intact annulus were unaffected by 15 minutes of treatment at 85°C. Although the increase in hysteresis after treatment implies an energetic change, the mechanisms by which the tissue was thermally modified are unclear. One possible explanation is provided by studies examining both the structure of collagenous tissue using scanning electron microscopy (SEM), and endothermic events, using DSC. Using these techniques, several investigators identified discrete stages of the denaturation process. They attributed the earliest denaturation (< 56°C) to the destruction of heat-labile cross-links (which are more pronounced in young animals), and showed that the structure of the fibrils remain intact during this process.

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A second contributing factor for the biomechanical changes is suggested by the observed increase in fractional dry mass in both our constrained and unconstrained treated tissue relative to the control tissue. The increase in fractional dry mass indicates that the tissues heated at 85°C swell less when equilibrated in saline. Since annular tissue hydration has been disclosed to be related to proteoglycan content, our finding indicates that the proteoglycans of the annulus have been affected by the heat treatment. Similar to collagen, proteoglycans are susceptible to denaturation through destruction of heat-labile hydrogen bonds. Alteration of annular proteoglycan can affect tissue properties since they have been previously disclosed to play a role in stabilizing the collagen matrix, as had been observed according to at least one prior disclosure in articular cartilage where the modulus decreases significantly when the proteoglycans are removed. It is thus believed that a portion of the observed biomechanical changes is due to changes in proteoglycan, the thermal properties of which are not extensively understood according to prior publications. Confirmation of such belief as to the specific mechanism with respect to proteoglycans may be achieved according to further study and observation by one of ordinary skill based upon review of this disclosure.

The retarding effect of stress on annular denaturation has a number of clinically relevant implications. First, to achieve a significant degree of collagen denaturation *in vivo*, the annulus should be heated either for long times or at high temperatures, or both. Second, thermal treatment according to the devices and methods of the present invention may be applied in a selective fashion. Since

unstressed annular fibers are more susceptible to thermal treatment than stressed fibers, areas of slack tissue (e.g. the inner annulus in degenerating discs) are preferentially heated, while preserving structurally competent areas that are carrying stress (e.g. the outer annulus that retains stress into later stages of degeneration). Further, patient pre-positioning is desired for certain circumstances, allowing the practitioner to selectively stress particular annular regions, thereby further controlling the zone of biomechanical alterations.

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In another regard, the present invention provides a useful tool when applied to selectively shrink proliferative fibrocartilage responsible for annular protrusion and prolapse. This is accomplished for example by providing the thermal therapy to degrade proteoglycans and decrease swelling.

In still a further regard, and as further supported by the results of this study, the present invention is used to provide thermal therapy in a manner specifically adapted to ablate annular nociceptors and cytokine producing cells while sparing tissue material properties. Thermal therapy in the range of 48-60 °C is sufficiently low to avoid collagen denaturation and biomechanical changes, yet this temperature region is desired for modes of thermal spine treatment intended to induce nerve injury and cellular death without significant biomechanical change from the heating (or with biomechanical change if desired and brought about by other means).

It is to be further appreciated that the results of this study, as to specific ranges and/or numbers, are potentially limited by the use of non-degenerate porcine intervertebral discs. While porcine discs are similar to human discs in many ways, there may be differences in denaturation temperature, which is dependent on a number of factors such as collagen cross-link type and density. However, the consistent tissue quality and size afforded by the porcine model minimizes interspecimen variability and therefore provides a good system by which to investigate mechanisms of thermal/biomechanical interactions. The disc height also differs between human and porcine lumbar discs. Since the lumbar human disc is generally taller (averaging approximately 11mm) than the porcine disc (averaging 3mm in this study), it may be less influenced by vertebral constraint and therefore more able to thermally contract. In this regard, as with many previously disclosed devices and treatment methods, the exact extent of effect may vary even between species

according to varied anatomy.

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Notwithstanding the foregoing, future studies may be performed on human discs according to one of ordinary skill based upon this disclosure to confirm effects of specific treatment regimens. Moreover, it is further believed that the relationship between varied temperatures (and/or ranges) and predictably varied results are well correlated across species, though specific temperatures, temperature-time dosing, or magnitudes of observed results may differ. Accordingly, it is believed that the studies disclosed herein and aspects of the invention related thereto provide beneficial treatment regimens, though such may clearly require further tuning in order to be particularly adapted for specified use in treating a particular patient, patient group, or even animal type.

Further to the experimental model of the present Example, nuclear depressurization allowed for the biomechanical response of the intact annulus to be isolated. However, for intact discs, nuclear pressure will increase annular stress and therefore is believed to further retard thermal effects beyond that observed here. Finally, the *ex vivo* study summarized herein does not characterize any subsequent biologic remodeling that would occur after heat treatment *in vivo*. Remodeling likely further modifies annular tissue properties, and the magnitude and temporal sequence of this response may be further characterized in a suitable *in vivo* model. However, the acute effects provided hereunder provide significant benefit notwithstanding such potential for remodeling.

Despite these limitations, the foregoing observations and related description demonstrates a number of mechanisms by which thermal therapy influences the biomechanical response of the annulus fibrosus. Unique features of the disc – specifically tissue structure and stress-strain constraints due to attachment to adjacent vertebrae – have significant impact on the thermal treatment effect size. Future *in vivo* animal studies and controlled human trials may be further performed by one of ordinary skill in the art based at least in part on this disclosure in order to further link biomechanical and biological consequences of tissue heating to the various beneficial patient outcomes.

External Directional Ultrasound Thermal Treatment ("ExDUSTT") System and Method

The following description relates generally to FIGS. 23-53 and provide further illustrative embodiments of the invention according to modes previously described above for providing an external directional ultrasound thermal treatment (or "ExDUSTT") device, and method for treating spinal disorders therewith.

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As illustrated in FIG. 23, the illustrative ExDUSTT applicator 110 of the present invention preferably has a support member 120 with an ultrasound transducer 130 mounted thereon within an outer covering 150 that is typically an inflatable coupling balloon such as is shown.

According to the further view shown in FIG. 24 during one mode of use in treating a region 108 of an intervertebral disc 104 associated with a spinal joint 101, the transducer is generally chosen to be a curvilinear panel that is both directional and focusing (e.g. converging signals) to help highly localized deep heating, in particular useful for applications from outside the disc as shown. It is to be appreciated that "spinal joint" where used throughout this disclosure generally includes intervertebral discs, adjacent vertebral bodies, and associated structures such as posterior vertebral elements such as facet joints.

As shown in FIGS. 25 and 26, respectively, ExDUSTT devices of the present invention can be on many different delivery platforms, such as a rigid pre-shaped platform shown in FIG. 25 with the transducer 130 on the distal bend section 116 and canted at an angle A for angular directional ultrasound relative to the proximal shaft 112 axis, or on a catheter-based platform shown in FIG. 26. Both chassis are beneficial for respective purposes, and each for common purposes. However, the pre-shaped probe is in particular useful for angular directional heating at hard to reach places in the body, such as certain spinal locations, and the rigidity helps position control.

FIG. 27A shows an illustrative rigid probe device 200 in finer detail for further understanding. Distal shaft 202 includes a 4mm outer diameter brass tube 204 with .008" silver lead wires 206 and water flow lines (e.g. 0.0226" polyimide tubing) contained therein and coupled at the proximal end portion (not shown) to a proximal adapter. A distal 0.1135" stainless steel tubing 208 is shown secured with epoxy 210

within the brass tube 204, and has a window 212 cut out leaving support ridges upon which the transducer 230 is mounted with Nusil 1137 Silicone, as shown in finer detail in FIG. 27B. Further included beneath the transducer 230 are rubber threads 236 strung across the window 212 to help provide a good non-dampening support system. An outer inflation balloon is shown at 240 and in shadow in the expanded condition for tissue coupling. As can be appreciated from the figure, an air backing is thus provided at 238 to provide highly directional ultrasound delivery away from the central shaft of the device and out through the eccentric inflation balloon 240 and into tissue there. Moreover, as illustrated in FIG. 27B, the transducer 230 is curvilinear around an axis that is aligned with the long axis of the support shaft and thus is focused into tissue along the transducer and balloon length as such. For the purpose of a complete description, one exemplary transducer that has been observed to be useful in this and other ExDUSTT designs herein shown and described has for example the following specifications: 0.394" long x 0.98" wide x 0.013" thick PZT4, 0.59" radius of curvature.

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Various modifications may be made to the device just shown and described. For example, the balloon according to that Figure was elastomeric type, such as 0.005" wall silicone balloon. However, better repeatability of size and shape may be required than what such elastomers can offer, and thus a less compliant balloon of the preformed type may be used. This is shown for example at balloon 248 in FIGS. 28A-B that is for example a pre-shaped PET balloon having a wall thickness for example of 0.001". Further considerations for materials may be considered, such as for example thermal properties, ultrasound transmissivity, profile, etc.

Moreover, similar features as just described for the ExDUSTT device may be incorporated onto a different catheter chassis without much required modification, as referenced in FIGS. 29A-B. Here a proximal catheter shaft 250 is shown coupled to a distal 4mm OD brass tube 254. Everything else may be the same as described above for the rigid probe designs. The catheter shaft 250 may be multi lumen, or may be a bundle of lumens, etc.

The transducers shown in the previous FIGS. are not the only configurations contemplated, either. For example, FIGS. 30A-B show a curvilinear transducer 260 with its radius of curvature R around an axis that is transverse (e.g. orthogonal) to

the long axis L of the support shaft 280.

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Further understanding of various modes of operating devices of the rigid probe type just shown and described are provided in FIGS. 31A-32B, which reflect operation at 5.6 MHz optimal frequency, with peak efficiency at 40%, and linear output and efficiency out to 12W applied with 5.5 W emitted from the transducer.

FIG. 33 shows a test set-up for ex-vivo pig spine treatment using a catheter-based ExDUSTT device over 5 minute heating period, and shows certain measured temperatures during a relatively low temperature mode of operation. T1 is a temperature probe 5mm deep into tissue from the transducer coupling interface, whereas T2 temperature probe shows the temperature profile over varied depths from the transducer.

In contrast to the ex-vivo data shown in FIG. 33, FIGS. 34A-B show results for in-vivo treatments in pig discs, and show temperatures all exceeding 55 degrees, though temperatures close to the transducer exceeded well over 65 degrees and even up to 80 degrees.

As illustrated in FIG. 35, various different sizes may be used depending upon the particular need, and a kit of different sizes, lengths, angles, etc. may be provided. The devices 270, 280 shown in FIG. 35 generally differ in that the larger device supports a 3.5mm wide ultrasound transducer, whereas the smaller device supports a 2.5mm wide transducer. In general, other features may be similar unless desired to change them, whereas for the embodiment kit shown, each device has other components scaled to meet the 2.5:3.5 comparison for the transducer widths. Other variations may be made, however.

For the purpose of further characterization, and understanding of directivity and focus of energy delivery as relates to the present invention, FIGS. 36A-B and 37A-B show certain output power profiles across the transducer faces for both 2.5mm and 3.5mm curvilinear transducers, respectively. The radius of curvature for these transducers is around an axis that is aligned with the long axis of these plots.

Various thermal treatment studies have been performed with working prototypes of the present invention and will be explained hereafter in part by reference to the test set-up for the rigid, pre-shaped bent ExDUSTT device shown in FIG. 38.

For example, as shown in FIG. 41, all temperatures monitored using 0 degree C cooling during relatively high temperature mode of operation were above 60 degrees C, even out to 10mm deep, and in particular were above 70 degrees C for most all data taken at 5mm depth, whereas temperatures predicted at 7mm deep were also in excess of 70 degrees. Other illustrative and informative results are reported up through FIG. 53 with respect to additional modes of ExDUSTT operation, which are best understood by further reference to the Brief Description of the Drawings above.

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Various different modes and embodiments for curvilinear transducers may be suitable for use according to the various embodiments herein described, such as for example the various ExDUSTT device embodiments just described. However, the following provides some further detail for particular modes and variations contemplated for the purpose of providing a more complete understanding

In one regard, these transducer segments such as used in the ExDUSTT devices (and per for example the earlier embodiment in FIG. 6C) are sectors of larger diameter cylinders or plated tubes, with ultrasound energy emitted from the concave surface. Examples include 0.394" (+/-.005")x 0.098" (+/- 0.002") linear wide x 0.013 thick which form an 14.2 deg. Arc segment of 0.4 " inner radius tube. One can realize different diameters or arc segments, such as 9.5 deg. Arc segment of 0.591 " inner radius tube. These can be purchased for example from vendors such as Boston Piezo-optics using materials such as PZT-4, 5, or 8.

The radius of curvature can be selected to sharpen or decrease the amount of energy concentration or apparent focusing (i.e., radius of curvature of 0.5, 0.75, 1.0, 1.5, 2.0 cm can be appreciated) with the higher radius of curvature and wider transducers giving more penetrating distributions. The width can vary to suit particular needs for operation or device compatibility, but may be for example between about 1.5 mm to about 6 mm; whereas the length can also vary to meet particular needs, such as for example from between about 2mm to about 10 mm or greater. Transducers meeting these specifications are in particular useful for various of the embodiments herein described, provided however that such embodiments nor other aspects of the invention should not be considered to be so limited to only these dimensions.

These transducers can be mounted in transducer assemblies using a variety of suitable means. Flexible adhesives (e.g. silicone adhesive, Nusil), rigid epoxies or conformal coatings (Dow Corning) may be used. Rigid metal (brass or stainless steel) or plastic assemblies can be machined to hold the transducers and maintain air-backing. One more detailed example incorporated into many of the ExDUSTT devices shown and described includes a filed down, 15 mm long portion (or specified length) across a 180 degree plane transversely through a distal stainless-steel support hypotube. :This forms a shelf for either side of the transducers to be mounted. Lead wires (such as for example either silver lead wire or miniature coaxial cable (Temflex, Inc)) are soldered to the transducer surfaces for power application and can be run within the central lumen of the SS.

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A thin layer of silicone adhesive can be placed upon the edges of the tube structure, and the transducer segment placed. The transducer can then be sealed using silicone adhesive and/or conformal coating. The conformal coating can be accelerated using elevated heat for about 60 min. Alternatively, rubber thread can be used for a spacer with silicone adhesive, to keep transducers from contacting the metal surface. Other holding devices can be implemented, including pieces machined from brass bar or rod with gaps for the air space and offsets to support. In some implementations, it may be desired to circulate water or fluid behind the surface of the transducers.

These transducer assemblies can be either modular catheter form insertion into target tissue (intra-discal) or rigid external applicator. It is not necessary, but these transducers can be sealed using epoxy and polyester layers a previously described, or using mineral oil or other type of oil instead of Epotek, or the transducer can be left bare, though in many applications would be sealed on its edges and possibly top surface with conformal coating for watertight integrity and durability. Custom multilumen extrusions in materials such as pebax can form the flexible catheter member of which the transducer assembly is attached. The transducers are rigid, but if multiple segments are used, they may be coupled in a manner providing flexible hinges for better bendability in use.

Pre-shaped high-pressure balloons such as those herein shown for ultrasound tissue coupling can be provided in various shapes. Suitable sources include for

example custom fabrication, such as for example by Advanced Polymers, or may be made in house by heat-stabilizing the PET heat-shrink in pre-determined shape using molds and Teflon-coated mandrels. These balloons can have a neck that is 3 mm OD and a one sided inflation with a 2 mm radius. Compliant balloons using silicone, c-flex, polyurethane, or other material can also be used for various applications indicating such compliance or elastomeric properties.

These devices can have temperature regulated flow, flow in general, or no flow at all. In addition, devices without encapsulating balloons can be realized with sterile saline or fluid flow used to cool and couple US to the interface.

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Internal Directed Ultrasound Thermal Therapy ("InDUSTT™") System and Method

The following description relates to device and method embodiments in particular adapted for use internally within intervertebral discs or other joints, e.g. "InDUSTT" devices and methods.

The following FIGS. and accompanying description is to be read in conjunction with prior description herein made above, and in any event relate to InDUSTT devices such as that shown at catheter 290 in FIG. 54. More specifically, FIG. 54 illustrates arrangements for two alternative modes of InDUSTT assemblies and treatments as follows. Catheter 290 is shown at the top of the Figure in a "direct coupling mode" and is adapted to be delivered directly into a region of a spinal joint, e.g. within the disc or bony structure, with the transducer coupled directly to tissue. Thus, there is only shown a coupling 294 such as for electrical leads 296, and fluid cooling coupling 298. As shown in the assembly of FIG. 54 on the bottom, however, a catheter cooled or "cc" arrangement differs from the direction coupling or "cc" arrangement in that a catheter 290 is buried within a closed housing of an outer delivery device 300 that has a sharp pointed tip 302 for puncturing into an intervertebral disc. Water circulation ports 298,310 according to this arrangement cycle cooling fluids between a lumen within the internal catheter 290 and over the internal catheter 290 but within the outer sheath 300.

Further details of the cc arrangement are variously shown in FIGS. 55A-C, wherein a transducer 320 is of a cylindrical tubular type that has sectored grooves

322 (FIG. 55C) on either side of an electroded and active portion 326 for directional ultrasound delivery along about a 90 degree span of space radially outward from that section, and thus both directionality is achieved, as well as diverging US signal. Couplers shown include power lead coupler 294, water inflow coupler 298, water outflow coupler 310. As shown in FIG. 55B, the water cooling is aided by a distal port 299 in the internal catheter device 290. A thermocouple 330 is shown along the active sector, as well as others may be provided elsewhere (not shown) such as along the other dead sector as temperature monitoring may still be important there to protect certain tissues from conductive heating during US therapy on the opposite side of the catheter. The transducer shown may be for example 1.5mm outer diameter, 0.9 mm inner diameter, by 10 mm long and mounted over a plastic support ring. The outer catheter 300 may be for example constructed from a simple polymeric tubing such as made from CELCON™ from Best Industries typically used conventionally for implanting radiation seeds in tumor therapy and of 13 gauge construction.

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For the purpose of providing a thorough understanding of the many different aspects and considerations of using the InDUSTT device just described, a significant amount of summary results from multiple studies of working embodiments is herein provided by reference to FIGS. 56A and beyond. A thorough understanding of various aspects of these results and experimental arrangements shown is gleaned by review of the Brief Description of the Drawings above. In general, where discs are indicated by an arrangement such as "C3-4" or "C4/5" such designates the vertebrae in the animal between which the disc was treated. In addition, curve legends designating numbers for graphs (such as for example "Probe 1-1", "Probe 1-2", etc. in FIG. 65A) designate thermocouples along temperature monitoring probes in the disc tissue, typically spaced by 5 mm apart.

Accordingly, as is reflected in the graphs and other pictures and Figures, many different discs were treated. Still further, the test results shown also reflect an understanding of the effects of cooling at different temperatures, as well as direct coupling versus catheter cooled coupling, as well as relatively high versus relatively low temperature modes of use.

While the results shown in these latter FIGS are in non-human animal models,

the results, and in particular the relationships between results between different treatment groups, correlate to the human condition and are confirmed by earlier human cadaver studies performed. Actual values may of course differ, however, but it is believed that the extreme ends of the results would apply across vertebrate animal species. Moreover, the date suggests that directivity is confirmed, as is the ability to achieve high temperatures over 70 degrees or even75 degrees, as well as control heating to lower temperatures for other intended treatments.

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In one example, FIGS 59A and B show two sets of lines that cross at time equal to about 900 seconds. This indicates a period of time when the InDUSTT device, with sectored, directional ultrasound emission, was rotated. Accordingly, the uniform change in temperature reflects the preferential heating that can be achieved with such device and not heretofore possible. For further illustration, FIG. 65A shows similar results according also to turning a directional device to realign the active sector to different transducers.

Various embodiments have been herein described, including ExDUSTT, InDUSTT, rigid-probe based, catheter based, directly coupled, actively cooled, sectored transducers, curvilinear transducers, axially aligned transducers, transversely aligned transducers, relatively large transducers, relatively small transducers, compliant elastomeric coupling balloons, relatively non-compliant preformed coupling balloons, relatively high temperature modes of operation, relatively low temperature modes of operation, low temperature cooling, room temperature cooling, preshaped, flexible, guidewire delivery, and deflectable/steerable delivery platforms. It is to be appreciated that the more detailed description for such embodiments provided herein is for the purpose of illustration, and other modes of achieving such may be suitable for inclusion according to the invention without departing from the present scope. Moreover, the combinations of such features herein shown are highly beneficial, but not intended to be limiting. Other combinations may be made without departing from the intended scope hereof. For example, the particular embodiments shown and described for "ExDUSTT" applications are described as such merely according to their highly beneficial ability to perform in that arrangement, but they may be used as InDUSTT devices as well, despite their particular external use benefits. The opposite is true, as well, with

respect to InDUSTT devices which may also be used in other external locations such as for disc heating. The devices shown and described may be used within or around the bony structures of spinal joints, too. Moreover, where various of the features may be highly beneficial for particular applications, they may not be necessary for other applications. For example, directional energy delivery is a highly beneficial aspect of the various ultrasound embodiments herein shown and described, in particular where highly localized heating is desired while other surrounding tissues need to be protected such as nerves. However, in other applications, such as some complete disc remodeling applications for example, non-directional emission may be suitable to heat all the surrounding tissue equally.

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It is to be appreciated that the various modes of devices and operation herein described, together with tissue characterization studies performed and herein presented, provide a significant understanding with respect to adapting and controlling thermal therapy, or other modes of ultrasound delivery for therapy, in special areas in the body such as joints, and in particular spinal joints and their discs and bony structures. Back pain and other issues in these joints are significant medical issues that may be addressed with the present invention according to its many different modes and aspects.

Although the description above contains many specificities, 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. Thus the scope of this invention should be determined by the appended claims and their legal equivalents. Therefore, it will be appreciated that the scope of the present invention fully encompasses other embodiments which may become obvious to those skilled in the art, and that the scope of the present invention is accordingly to be limited by nothing other than the appended claims, in which reference to an element in the singular is not intended to mean "one and only one" unless explicitly so stated, but rather "one or more." All structural, chemical, and functional equivalents to the elements of the above-described preferred embodiment that are known to those of ordinary skill in the art are expressly incorporated herein by reference and are intended to be encompassed by the present claims. 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. Furthermore, no element, component, or method step in the present disclosure is intended to be dedicated to the public regardless of whether the element, component, or method step is explicitly recited in the claims. No claim element herein is to be construed under the provisions of 35 U.S.C. 112, sixth paragraph, unless the element is expressly recited using the phrase "means for."

CLAIMS

What is claimed is:

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1. An ultrasound energy delivery system for treating a region of tissue associated with a skeletal joint, comprising:

an ultrasound treatment assembly with an ultrasound transducer;

a skeletal joint delivery assembly that is adapted to deliver the ultrasound treatment assembly into the body with the ultrasound transducer positioned at a location within the body associated with the skeletal joint; and

wherein the ultrasound treatment assembly is adapted to deliver a therapeutic level of ultrasound energy from the location and into the region of tissue.

- 2. The system of claim 1, wherein the skeletal joint delivery assembly comprises a needle.
- 3. The system of claim 1, wherein the skeletal joint delivery assembly comprises a cannula with a sharpened tip that is adapted to puncture a tissue barrier associated with the joint.
 - 4. The system of claim 1, wherein: the skeletal joint delivery assembly comprises a guidewire; and the ultrasound treatment assembly is adapted to be delivered over the guidewire to the location.
- 5. The system of claim 1, wherein:
 the skeletal joint delivery assembly comprises a cannula; and the ultrasound treatment assembly is adapted to be delivered through the cannula to the location.
- 6. The system of claim 1, wherein the ultrasound treatment assembly is adapted to be steerable to the location.
 - 7. The system of claim 6, wherein the ultrasound treatment assembly is

located on a deflectable member.

8. The system of claim 1, wherein the ultrasound treatment assembly is located on a pre-shaped support member.

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- 9. The system of claim 1, wherein the ultrasound transducer is constructed to provide directional thermal therapy to the region of tissue.
- 10. The system of claim 9, wherein the ultrasound transducer is constructed to deliver a converging field of energy to the region of tissue.
 - 11. The system of claim 9, wherein the ultrasound transducer is constructed to deliver a diverging field of ultrasound energy to the region of tissue.
- 15 12. The system of claim 9, wherein the ultrasound transducer is constructed to deliver a substantially collimated field of ultrasound energy to the region of tissue.
- 13. The system of claim 1, wherein the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.
 - 14. The system of claim 13, wherein:

the coupling member comprises a radially extendable member that is adapted to radially extend from a first contracted position relative to the ultrasound transducer to a second extended position relative to the ultrasound transducer that is further away from the ultrasound transducer than the first position; and

the ultrasound transducer is adapted to delivery the ultrasound energy to the region of tissue via the coupling member with the radially extendable member in the second extended position.

15. The system of claim 14, wherein:

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the extendable member comprises an inflatable balloon that is adapted to be coupled to a source of inflation medium and to inflate with a pressurized volume of fluid from the source; and

the ultrasound transducer is adapted to deliver the ultrasound energy to the tissue via the fluid and inflatable balloon.

16. The system of claim 1, further comprising:

a temperature control system that is adapted to control the temperature of at
least one of the ultrasound transducer or a tissue interface between a region of
tissue and the ultrasound treatment assembly during ultrasound energy delivery to
the region of tissue.

- 17. The system of claim 16, wherein the ultrasound treatment assembly is adapted to actively cool the ultrasound transducer or tissue interface.
 - 18. The system of claim 1, further comprising:

a cooling system with a controller;

wherein the ultrasound treatment device is adapted to couple to the cooling system; and

wherein the cooling system via the controller is adapted to cool at least one of the ultrasound transducer and a region of tissue being affected by the ultrasound delivery.

- 25 19. The system of claim 18, wherein the cooling system is adapted to cool the ultrasound transducer with a fluid having a temperature less than room temperature.
- 20. The system of claim 18, wherein the fluid temperature is about zero degrees C.
 - 21. The system of claim 18, wherein the fluid temperature is equal to or

greater than about room temperature.

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22. The system of claim 18, wherein the cooling system comprises: a source of fluid medium; and

wherein the cooling system is adapted to circulate fluid from the source through the ultrasound treatment device so as to cool the ultrasound transducer.

23. The system of claim 1, further comprising:

an ultrasound control system with a temperature monitoring system, a controller, and an ultrasound drive system;

wherein the ultrasound treatment assembly is adapted to couple to the ultrasound control system; and

wherein the ultrasound control system is adapted to control the ultrasound energy being emitted by the ultrasound transducer as a function of at least one parameter related to thermal therapy in the region of tissue.

- 24. The system of claim 23, wherein the at least one parameter comprises a parameter related to the operation of the ultrasound transducer.
- 25. The system of claim 24, wherein the at least one parameter comprises impedance, voltage, current, power, temperature, reflected power, or rates of change thereof.
- 26. The system of claim 23, wherein the at least one parameter comprises
 a parameter related to the effects of ultrasound delivery from the ultrasound transducer in tissue.
 - 27. The system of claim 26, wherein the ultrasound control system is adapted to control operation of the ultrasound delivery assembly based upon a temperature monitored at a coupling tissue surface between the tissue and the ultrasound delivery assembly.

28. The system of claim 26, wherein the at least one parameter comprises thermal dose, tissue temperature, depth of thermal penetration, or rates of change thereof.

- 29. The system of claim 23, wherein the at least parameter is empirically based.
 - 30. The system of claim 23, wherein the at least one parameter comprises a value that is monitored during ultrasound therapy.

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- 31. The system of claim 23, wherein the at least one parameter comprises a value that is calculated based upon another measured parameter.
- 15 32. The system of claim 23, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature in at least a portion of the tissue within at least a 4 mm depth from the ultrasound transducer is elevated to at least about 70 degrees C in response to ultrasound energy being delivered thereto from the ultrasound transducer.

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33. The system of claim 32, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature of the tissue within the 4mm depth is elevated to at least about 75 degrees C in response to the ultrasound energy delivery.

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34. The system of claim 23, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up to at least about a 7 mm depth from the ultrasound transducer is elevated to at least about 70 degrees C.

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35. The system of claim 23, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up

to at least a 10mm depth from the ultrasound transducer is elevated to greater than about 45 degrees C.

36. The system of claim 23, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature within the tissue at up to at least a 4 mm depth from the ultrasound transducer is elevated to an elevated temperature of no more than about 75 degrees C or less.

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- 37. The system of claim 36, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature of the tissue within the 4mm depth is elevated to an elevated temperature of no more than about 70 degrees C or less.
- 38. The system of claim 23, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up to at least a 7 mm depth from the ultrasound transducer is elevated to at least 70 degrees C.
- 39. The system of claim 23, wherein the controller is adapted to control the operation of the ultrasound transducer such that the temperature in tissue within up to at least a 10 mm depth from the ultrasound transducer is elevated to greater than 45 degrees C.
- 40. The system of claim 1, wherein the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least about 4 mm from the ultrasound transducer to a temperature of at least about 70 degrees C.
 - 41. The system of claim 40, wherein the ultrasound heating assembly is adapted to heat the tissue within the 4 mm depth to a temperature of at least about 75 degrees C.
 - 42. The system of claim 41, wherein the ultrasound treatment assembly is

adapted to heat the tissue at least about 4 mm away from the transducer to the temperature of at least about 75 degrees C in less than about 5 minutes of ultrasound energy delivery into the tissue.

- 43. The system of claim 1, wherein the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least about 7 mm from the ultrasound transducer to a temperature of at least about 70 degrees C.
- 44. The system of claim 1, wherein the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least about 10mm from the ultrasound transducer to a temperature of at least about 45 degrees C.
 - 45. The system of claim 1, wherein the ultrasound treatment assembly is adapted to heat tissue up to a distance of at least 4 mm from the ultrasound transducer to a temperature of no more than about 75 degrees C or less.
 - 46. The system of claim 1, wherein the ultrasound treatment assembly is adapted to be positioned within an intervertebral disc and to heat at least portion of the disc.

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- 47. The system of claim 1, wherein the ultrasound treatment assembly is adapted to be positioned at a location adjacent to an intervertebral disc and to provide thermal therapy to the intervertebral disc from that location.
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- 48. The system of claim 1, wherein the ultrasound treatment assembly is adapted to be positioned within at least a portion of a vertebral body and to provide thermal therapy at least in part to the vertebral body.
- 49. The system of claim 1, wherein the ultrasound treatment assembly is adapted to be positioned at a location adjacent to a vertebral body and to heat at least a portion of the vertebral body.

50. The system of claim 1, wherein the ultrasound treatment assembly and the skeletal joint delivery assembly are integrated.

51. The system of claim 50, wherein:

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- the skeletal joint delivery assembly comprises a substantially rigid probe device with a proximal end portion and a distal end portion; and the ultrasound heating assembly is located along the distal end portion.
- 52. The system of claim 1, wherein the ultrasound treatment assembly and skeletal joint delivery assembly are separate devices that cooperate together.
 - 53. The system of claim 1, wherein the skeletal joint delivery assembly comprises a spinal joint delivery assembly.
- 15 54. The system of claim 1, wherein the ultrasound treatment assembly and spinal joint delivery assembly together comprise an external directional ultrasound thermal therapy system.
- 55. The system of claim 1, wherein the ultrasound treatment assembly and spinal joint delivery assembly together comprise an internal directional ultrasound thermal therapy system.
 - 56. The system of claim 1, wherein the ultrasound treatment assembly comprises a single curvilinear ultrasound transducer.
 - 57. The system of claim 1, wherein the ultrasound treatment assembly comprises multiple curvilinear ultrasound transducers.
- 58. The system of claim 1, wherein the ultrasound transducer is adapted to emit directed, therapeutic ultrasound energy within a treatment zone of less than or equal to about one hundred eighty degrees relative to the ultrasound transducer.

59. The system of claim 58, wherein the treatment zone is less than or equal to about ninety degrees.

60. The system of claim 1, wherein the ultrasound transducer is comprises a curved shape with a radius of curvature that is around a reference axis that is transverse to the longitudinal axis of a support member upon which the ultrasound transducer is mounted.

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- 61. The system of claim 1, wherein the ultrasound transducer is comprises
 a curved shape with a radius of curvature that is around a reference axis that is
 aligned with the longitudinal axis of a support member upon which the ultrasound
 transducer is mounted.
 - 62. A skeletal joint thermal therapy device, comprising:
 a delivery member with a proximal end and a distal end portion;
 a thermal treatment assembly with an energy emitter on the distal end portion;
 wherein the distal end portion is adapted at least in part to deliver the thermal
 treatment assembly into a body of a mammal with the energy emitter positioned at a
 location such that the energy emitter is adapted to deliver energy into a region of
 tissue associated with a skeletal joint; and

wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 4mm from the energy emitter to a temperature of at least 75 degrees C, and is adapted to heat tissue up to a distance of at least about 7 mm from the energy emitter to a temperature of at least about 55 degrees C, and is adapted to heat tissue up to a distance of at least about 10 mm from the energy emitter to a temperature of at least about 45 degrees C.

- 63. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 4 mm from the energy emitter to a temperature of at least 80 degrees C.
 - 64. The device of claim 62, wherein the thermal treatment assembly is

adapted to heat tissue up to a distance of at least 4 mm from the energy emitter to a temperature of at least 85 degrees C.

- 65. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 7 mm from the energy emitter to a temperature of at least 60 degrees C.
- 66. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 7 mm from the energy emitter to a temperature of at least 65 degrees C.
- 67. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 7 mm from the energy emitter to a temperature of at least 70 degrees C.

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- 68. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 10 mm from the energy emitter to a temperature of at least 50 degrees C.
- 20 69. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 10 mm from the energy emitter to a temperature of at least 55 degrees C.
- 70. The device of claim 62, wherein the thermal treatment assembly is adapted to heat tissue up to a distance of at least 10 mm from the energy emitter to a temperature of at least 60 degrees C.
 - 71. The system of claim 62, wherein the thermal treatment assembly is constructed to provide directional thermal therapy to the region of tissue.

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72. The system of claim 62, wherein the thermal treatment assembly is constructed to deliver a converging field of energy to the region of tissue.

73. The system of claim 62, wherein the thermal treatment assembly is constructed to deliver a diverging field of energy to the region of tissue.

- 74. The system of claim 62, wherein the thermal treatment assembly is constructed to deliver a substantially collimated field of energy to the region of tissue.
 - 75. The system of claim 62, wherein the energy emitter comprises an ultrasound transducer.
- 76. The system of claim 62, wherein the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.
 - 77. An ultrasound thermal treatment system, comprising:
 an ultrasound treatment assembly with an ultrasound transducer;
 a rigid delivery probe having a elongate body with a proximal end portion and
 a distal end portion:

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the distal end portion has a proximal section with a longitudinal axis, a distal section with a distal tip, and a bend between the proximal and distal sections;

wherein the ultrasound treatment assembly is located along the distal section of the distal end portion and extending at an angle from the proximal section;

wherein the distal end portion is adapted to be delivered into the body of a mammal by manipulating the proximal end portion externally of the body and such that the ultrasound treatment assembly is positioned at a location associated with a region of tissue to be treated; and

wherein the ultrasound treatment assembly is adapted to deliver a therapeutic level of ultrasound energy into the region of tissue from the location within the body.

- 78. The system of claim 77, wherein the proximal end portion of the rigid delivery probe comprises a metal tube.
 - 79. The system of claim 77, wherein the distal end portion of the rigid

delivery probe comprises a metal tube.

80. The system of claim 77, wherein the ultrasound transducer is constructed to provide directional ultrasound delivery to the region of tissue.

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- 81. The system of claim 77, wherein the ultrasound transducer is constructed to deliver a converging field of ultrasound energy to the region of tissue.
- 82. The system of claim 77, wherein the ultrasound transducer is
 constructed to deliver a diverging field of ultrasound energy to the region of tissue.
 - 83. The system of claim 77, wherein the ultrasound transducer is constructed to deliver a substantially collimated field of ultrasound energy to the region of tissue.

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- 84. The system of claim 77, wherein the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.
- 20 85. The system of claim 77, further comprising a temperature control system coupled to the ultrasound treatment assembly.
 - 86. The system of claim 77, further comprising a drug delivery system that is adapted to deliver a drug into the region of tissue when the ultrasound treatment assembly is delivering the therapeutic ultrasound to the region of tissue.
 - 87. An ultrasound thermal therapy system, comprising:
 an ultrasound heating assembly with an ultrasound transducer;
 a therapy control system coupled to the ultrasound heating assembly;
 wherein the ultrasound heating assembly is adapted to be delivered into a
 body of a mammal with the ultrasound transducer positioned at a location such that
 the ultrasound transducer is adapted to deliver a therapeutic amount of ultrasound

energy into a targeted region of tissue in the body from the location; and
wherein the therapy control system is adapted to control operation of the
ultrasound heating assembly such that a substantial portion of the region of tissue
being heated by the ultrasound heating assembly does not exceed a maximum

- 88. The system of claim 87, wherein the therapy control system is adapted to control operation of the ultrasound heating assembly such that a substantial portion of the region of tissue being heated by the ultrasound heating assembly does not exceed a maximum temperature of at least about 75 degrees C.
- 89. A directional ultrasound spinal thermal therapy system, comprising: an ultrasound delivery assembly with a directional ultrasound transducer that is adapted to be positioned at a location associated with a spinal joint and to deliver a directed, therapeutic amount of ultrasound energy from the location and to a region of tissue associated with the spinal joint.
 - 90. The system of claim 89, further comprising:

temperature of at least about 70 degrees C.

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a support member comprising an elongate body with a proximal end portion and a distal end portion;

wherein the ultrasound delivery assembly is located along the distal end portion.

- 91. The system of claim 90, wherein the ultrasound delivery assembly is adapted to direct the ultrasound delivery into substantially only a particular radial zone around the circumference of the distal end portion of the support member.
 - 92. The system of claim 89, wherein the ultrasound delivery assembly is constructed to provide directional thermal therapy to the region of tissue.
 - 93. The system of claim 89, wherein the ultrasound delivery assembly is constructed to deliver a converging field of energy to the region of tissue.

94. The system of claim 89, wherein the ultrasound delivery assembly is constructed to deliver a diverging field of energy to the region of tissue.

- 95. The system of claim 89, wherein the ultrasound delivery assembly is constructed to deliver a substantially collimated field of energy to the region of tissue.
 - 96. The system of claim 89, wherein the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.

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97. The system of claim 89, wherein the ultrasound delivery assembly is adapted to be positioned at the location within at least a portion of an intervertebral disc associated with the spinal joint and to deliver the therapeutic ultrasound energy into the region of tissue from that location.

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98. The system of claim 89, wherein the ultrasound delivery assembly is adapted to be positioned at the location outside of an intervertebral disc associated with the spinal joint and to deliver the therapeutic level of ultrasound energy into the intervertebral disc from that location.

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99. The system of claim 89, wherein the ultrasound delivery assembly is adapted to be positioned at the location within at least a portion of a vertebral body associated with the spinal joint and to be ultrasonically coupled with the region of tissue from that location.

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100. The system of claim 89, wherein the ultrasound delivery assembly is adapted to be positioned at the location outside of a vertebral body associated with the spinal joint and to deliver the therapeutic level of ultrasound energy into the intervertebral disc from that location.

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101. A skeletal joint ultrasound delivery system, comprising: an ultrasound treatment assembly with an ultrasound transducer and a

coupling member;

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wherein the ultrasound delivery assembly is adapted to be delivered into a body of a mammal with the ultrasound transducer positioned at a location within the body associated with a skeletal joint; and

wherein the ultrasound transducer is adapted to deliver a therapeutic amount of ultrasound energy to a region of tissue associated with the skeletal joint via the coupling member.

- 102. The system of claim 101, wherein the ultrasound delivery assembly is constructed to provide directional ultrasound delivery to the region of tissue.
 - 103. The system of claim 101, wherein the ultrasound delivery assembly is constructed to deliver a converging field of energy to the region of tissue.
- 15 104. The system of claim 101, wherein the ultrasound delivery assembly is constructed to deliver a diverging field of energy to the region of tissue.
 - 105. The system of claim 101, wherein the ultrasound delivery assembly is constructed to deliver a substantially collimated field of energy to the region of tissue.
 - 106. The system of claim 101, wherein the ultrasound treatment assembly comprises a coupling member positioned so as to couple ultrasound energy from the ultrasound transducer to the region of tissue.
 - 107. The system of claim 106, wherein:

the coupling member comprises a radially extendable member that is adapted to radially extend from a first contracted position relative to the ultrasound transducer to a second extended position relative to the ultrasound transducer that is further away from the ultrasound transducer than the first position; and

the ultrasound transducer is adapted to delivery the ultrasound energy to the region of tissue via the coupling member with the radially extendable member in the second extended position.

108. The system of claim 107, wherein:

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the extendable member comprises an inflatable balloon that is adapted to be coupled to a source of inflation medium and to inflate with a pressurized volume of fluid from the source; and

the ultrasound transducer is adapted to deliver the ultrasound energy to the tissue via the fluid and inflatable balloon.

- 109. The system of claim 108, wherein the inflatable balloon is substantially non-compliant and preformed.
- 110. The system of claim 108, wherein the inflatable balloon comprises a substantially compliant, elastomeric balloon material.
- 111. The system of claim 101, wherein the ultrasound delivery assembly is
 adapted to be positioned at the location within at least a portion of an intervertebral
 disc associated with the spinal joint and to deliver the therapeutic ultrasound energy
 into the region of tissue from that location.
- 112. The system of claim 101, wherein the ultrasound delivery assembly is
 20 adapted to be positioned at the location outside of an intervertebral disc associated
 with the spinal joint and to deliver the therapeutic level of ultrasound energy into the
 intervertebral disc from that location.
- 113. The system of claim 101, wherein the ultrasound delivery assembly is
 adapted to be positioned at the location within at least a portion of a vertebral body
 associated with the spinal joint and to be ultrasonically with the region of tissue from
 that location.
- 114. The system of claim 101, wherein the ultrasound delivery assembly is
 adapted to be positioned at the location outside of a vertebral body associated with
 the spinal joint and to deliver the therapeutic level of ultrasound energy into the
 intervertebral disc from that location.

115. An ultrasound thermal therapy system, comprising:

an ultrasound heating assembly that is adapted to be positioned at a location within a body of a mammal so as to deliver ultrasound energy into a region of tissue in the body from the location; and

a control system coupled to the ultrasound heating assembly and adapted to control operation of the ultrasound heating assembly such that a region of tissue being heated by the ultrasound heating assembly exceeds a temperature of at least about 70 degrees C.

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116. An ultrasound thermal therapy system, comprising:

a delivery member with a proximal end portion and a distal end portion with a longitudinal axis;

an ultrasound heating assembly located along the distal end portion of the delivery member and with a curvilinear ultrasound transducer having a concave surface with a radius of curvature around a reference axis that is transverse to the longitudinal axis of the distal end portion.

117. A method for invasively treating a medical condition associated with a skeletal joint within a body of a patient, comprising:

delivering a therapeutic level of ultrasound energy to a region of tissue associated with the joint from a location within the body of the patient.

118. A method for treating a medical condition associated with a skeletal joint within a body of a patient, comprising:

delivering sufficient energy to a region of tissue associated with the skeletal joint that is sufficient to necrose nociceptive nerve fibers or inflammatory cells in such tissue region without substantially affecting collagenous structures associated with the skeletal joint.

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119. The method of claim 118, further comprising: delivering the energy into the region of tissue while the region of tissue is

under mechanical stress; and

heating the tissue to a temperature of up to no more than 75 degrees C.

120. The method of claim 119, further comprising: heating the tissue to a temperature of up to no more than 70 degrees C.

121. A method for treating a region of tissue associated with an intervertebral disc in a body of a patient, comprising:

delivering an ultrasound transducer to a location within the body such that a therapeutic level of ultrasound may be coupled from the transducer to the tissue.

122. A method for treating a patient, comprising:

delivering energy to a region of tissue associated with the spine, wherein such tissue does not experience a rise in temperature of more than about 55 degrees C.

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123. A method for treating a patient, comprising:

delivering energy to a region of tissue associated with the spine, wherein such energy delivery is between about 10 and about 300 equivalent minutes at 43 degrees C.

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124. A method for invasively treating a medical condition associated with an intervertebral disc within a body of a patient, comprising:

delivering a therapeutic level of ultrasound energy to a region of tissue associated with an intervertebral disc from a location within the body of the patient.

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125. A method for treating medical condition associated with a joint between two bony structures in a body of a patient, comprising:

delivering an ultrasound transducer to a location within the patient's body associated with the joint;

emitting ultrasound energy from the transducer at the location so as to provide a therapeutic effect to at least a portion of the joint.

126. The method of claim 125, further comprising: focusing the ultrasound energy from the transducer toward the portion.

- 127. The method of claim 125, wherein the location is within the joint between the two bony structures.
 - 128. The method of claim 125, wherein the joint is between two vertebral bodies in a spine of the patient.
- 10 129. The method of claim 125, wherein the portion heated comprises at least a portion of an intervertebral disc.
 - 130. The method of claim 125, wherein the portion heated comprises at least a portion of an annulus fibrosus of the intervertebral disc.
 - 131. The method of claim 125, further wherein the location is at least in part within an annulus fibrosus of the intervertebral disc.
- 132. The method of claim 125, wherein the location is adjacent to and outside of an annulus fibrosus of the intervertebral disc.

- 133. The method of claim 125, wherein the location is adjacent to a vertebral body associated with the spinal joint.
- 25 134. The method of claim 125, wherein the location is within a vertebral body associated with the spinal joint.
- 135. A method for treating a patient, comprising:
 introducing an ultrasound transducer into a body of a patient;
 positioning the ultrasound transducer at a location within the patient that is
 adjacent to at least one of an annulus fibrosus of an intervertebral disc, a nucleus
 pulposus of the intervertebral disc, or a vertebral body associated with a spinal joint

in the body; and

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emitting ultrasound energy from the ultrasound transducer at the location.

136. A method for providing ultrasound energy delivery within a body of an animal, comprising:

introducing an ultrasound transducer into a body of a patient;

positioning the ultrasound transducer at a location within the patient that is within at least one of an annulus fibrosus of an intervertebral disc, a nucleus pulposus of the intervertebral disc, or a vertebral body associated with a spinal joint in the body; and

emitting ultrasound energy from the ultrasound transducer at the location.

137. A method for treating a patient, comprising:

ultrasonically heating a region of tissue associated a spinal joint to a temperature between about 45 to about 90 degrees C for sufficient time to cause a therapeutic result in the tissue.

- 138. The method of claim 137, further comprising heating the region of tissue to a temperature that is between about 45 to about 50 degrees.
- 139. The method of claim 137, further comprising providing a thermal dose to the tissue between about 10 and about 300 equivalent minutes at 43 degrees C.
- 140. The method of claim 137, further comprising providing sufficient thermal dose so as to cause necrosis effect in nociceptive nerve fibers or inflammatory cells in the region of tissue.
- 141. The method of claim 137, further comprising ultrasonically heating the region to a temperature between about 50 and about 60 degrees C.
 - 142. The method of claim 137, further comprising providing sufficient

thermal dose at appropriate temperature so as to stimulate cellular metabolism without substantially killing cells in the region of tissue.

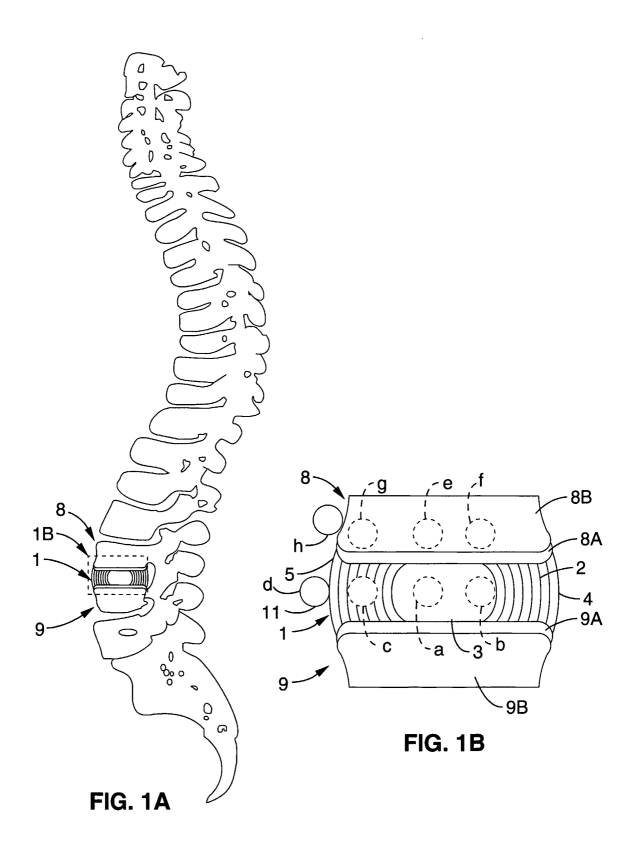
- 143. The method of claim 137, further comprising heating the regionsufficient to cause cellular necrosis in the tissue.
 - 144. The method of claim 137, further comprising delivering sufficient thermal dosing to the region of tissue to cause a substantially non-necrotic cellular effect.

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- 145. The method of claim 137, further comprising delivering sufficient thermal dosing to the region of tissue to cause preferential regeneration.
- The method of claim 137, further comprising delivering sufficient
 thermal dose to the region of tissue to cause inhibition of inflammatory factors or cytokines.
 - 147. The method of claim 137, further comprising delivering sufficient thermal dose to the region of tissue to cause modification of a healing response to injury in the region of tissue.
 - 148. The method of claim 137, further comprising heating the region from within an annulus fibrosus of an intervertebral disc.

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- 149. The method of claim 137, wherein the region comprises a portion of an intervertebral disc.
- 150. The method of claim 137, wherein the region comprises a portion of an annulus fibrosus.



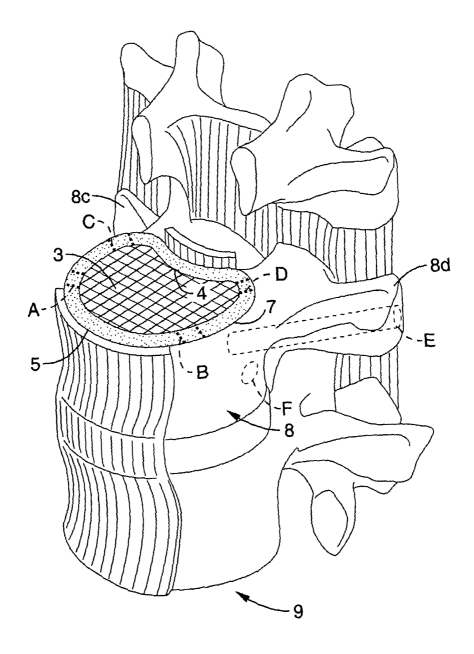
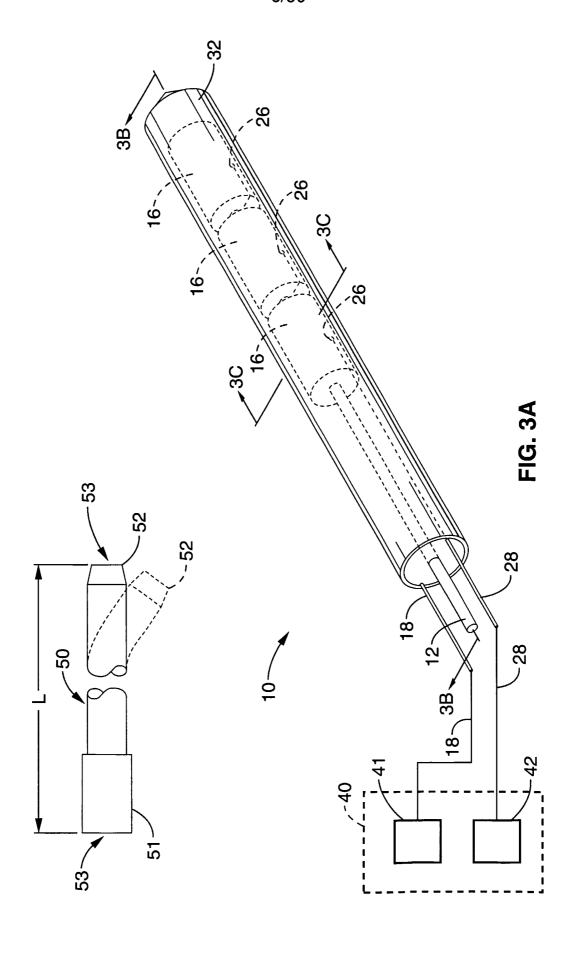
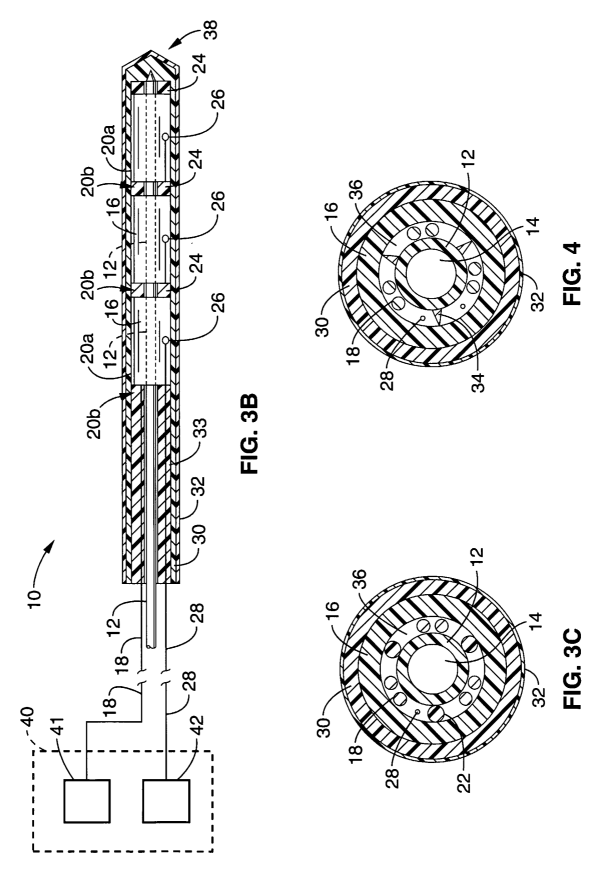
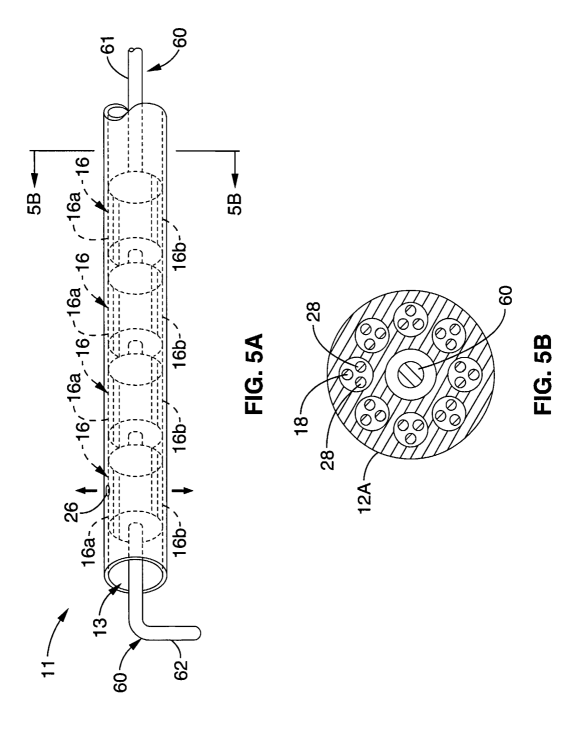


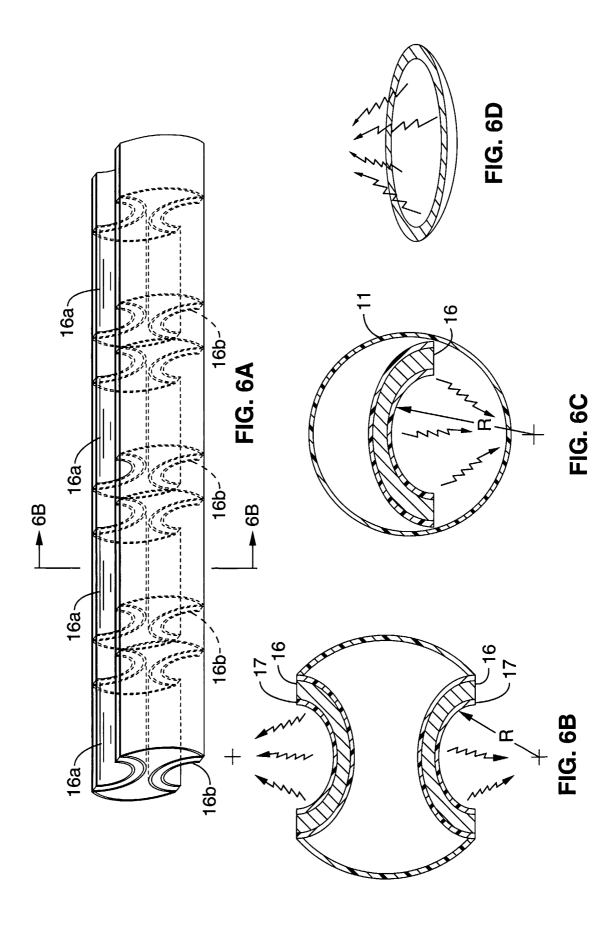
FIG. 2

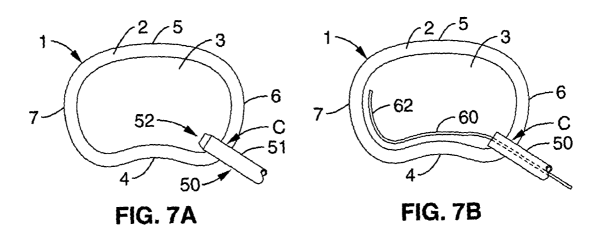


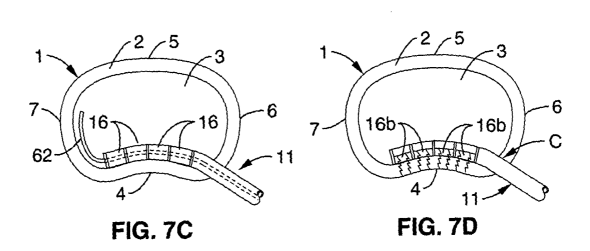
4/56

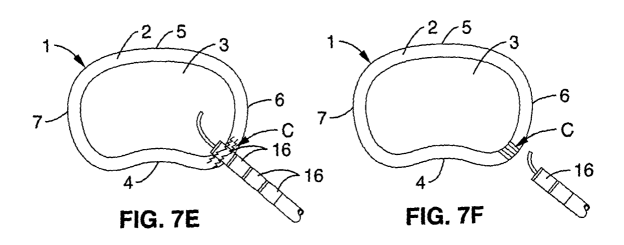


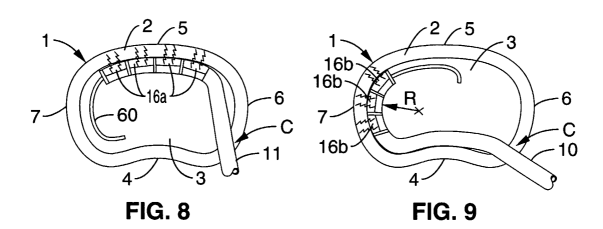


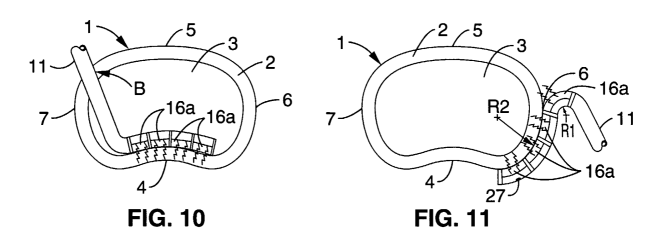


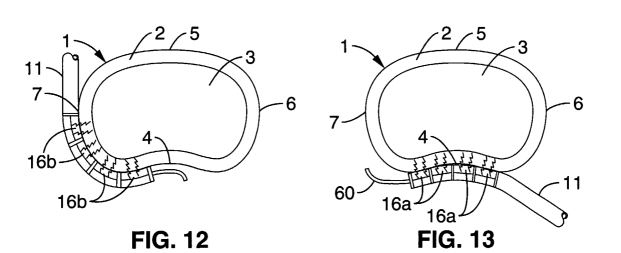




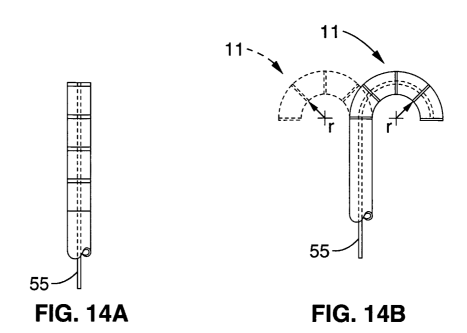


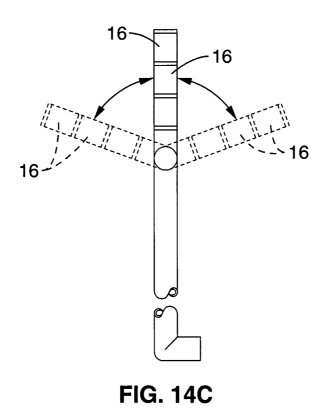


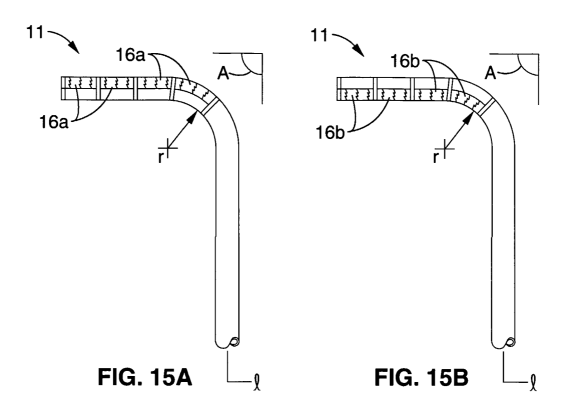












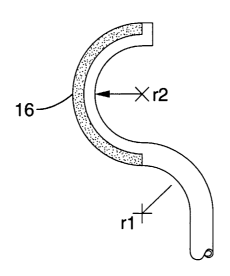
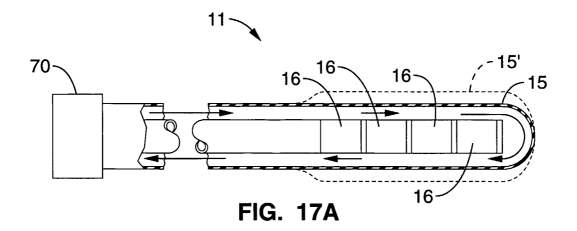
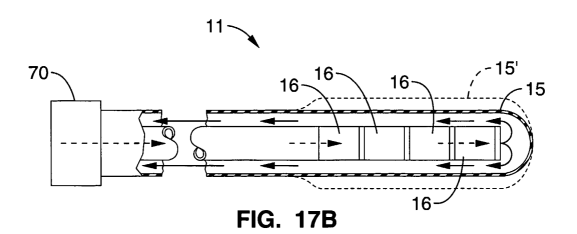


FIG. 16





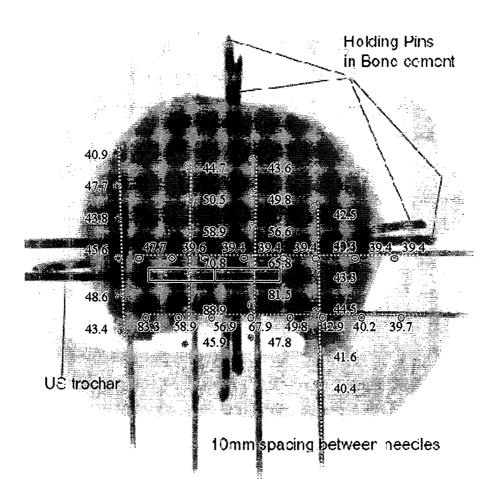
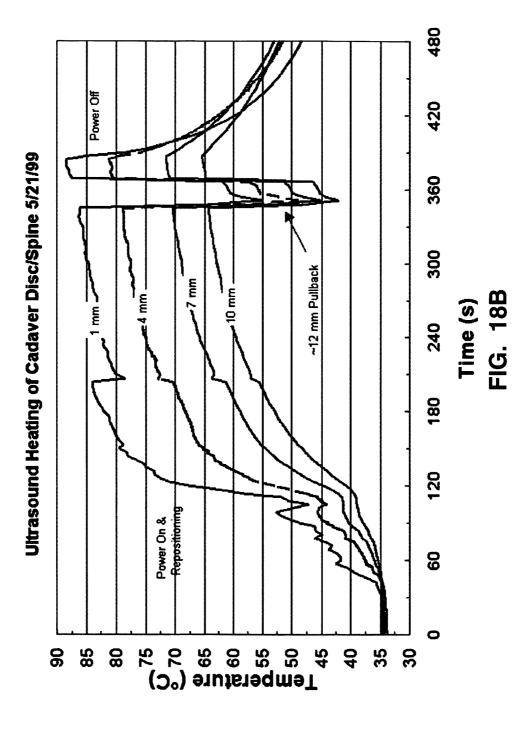
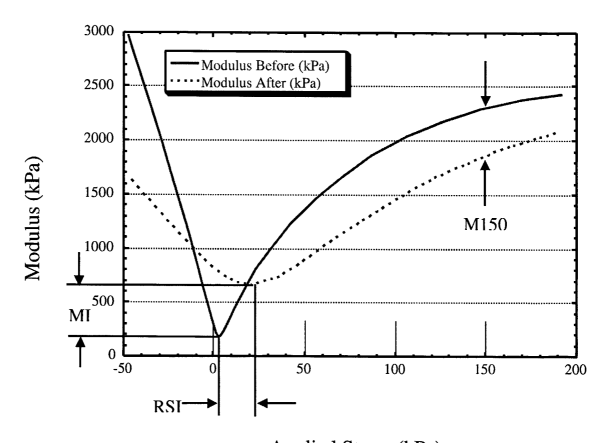


FIG. 18A





Applied Stress (kPa)

FIG. 19

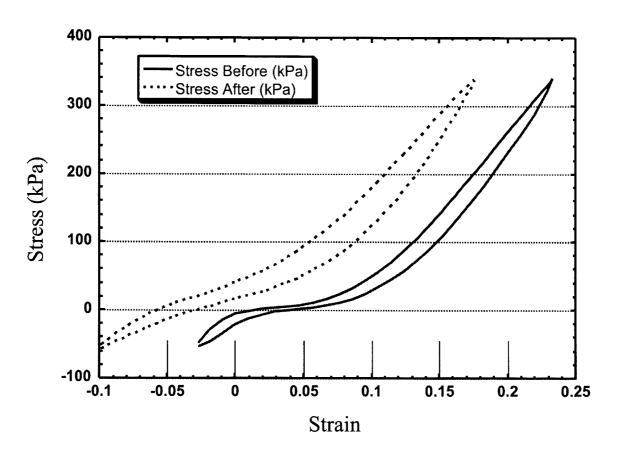
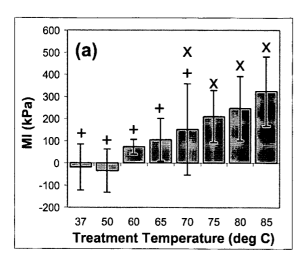
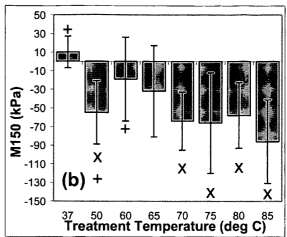
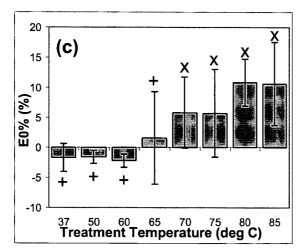
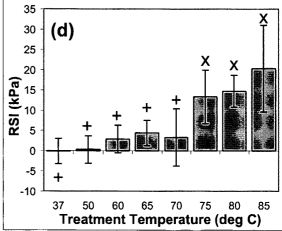


FIG. 20









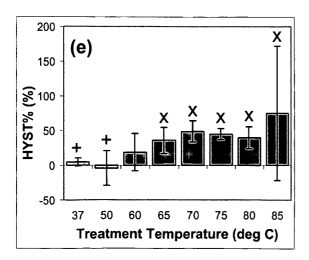


FIG. 21

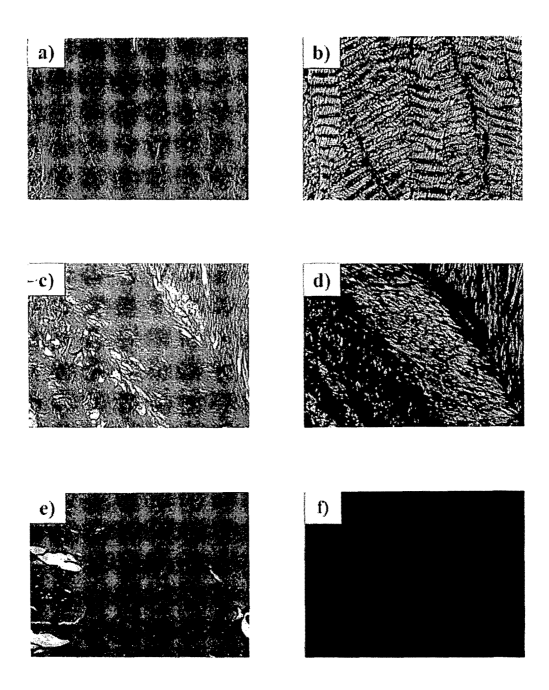


FIG. 22

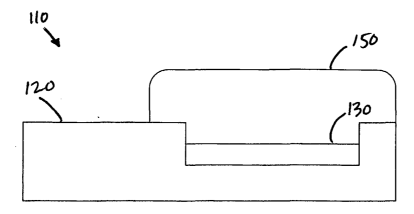


FIG. 23

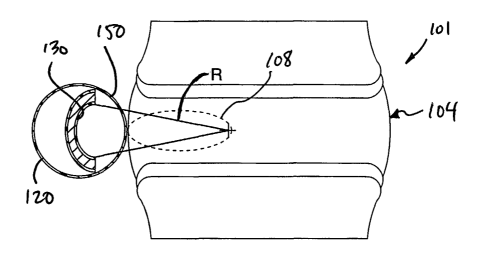
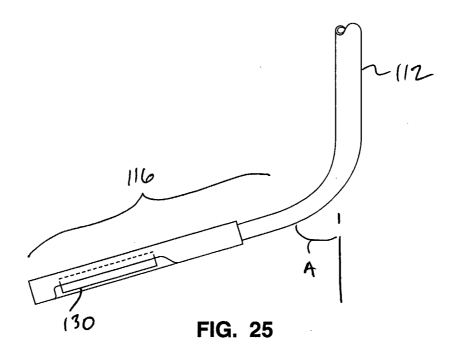


FIG. 24



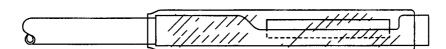
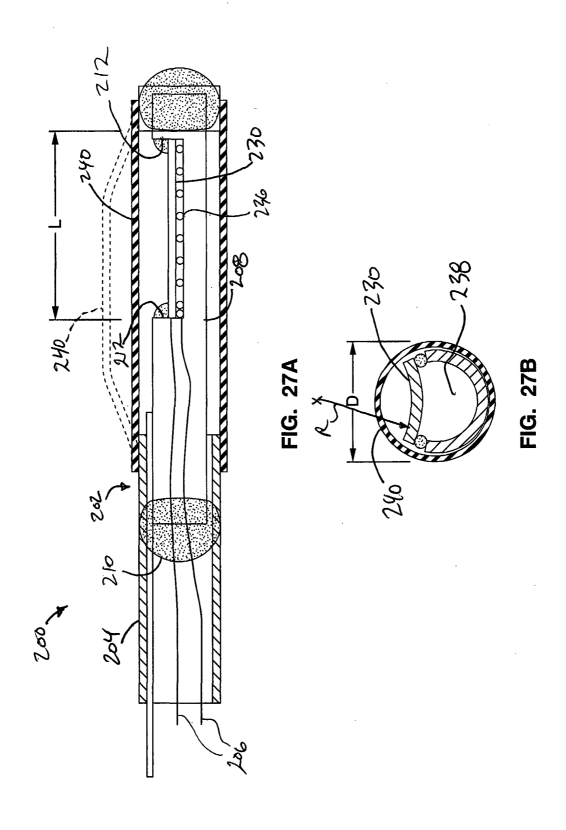
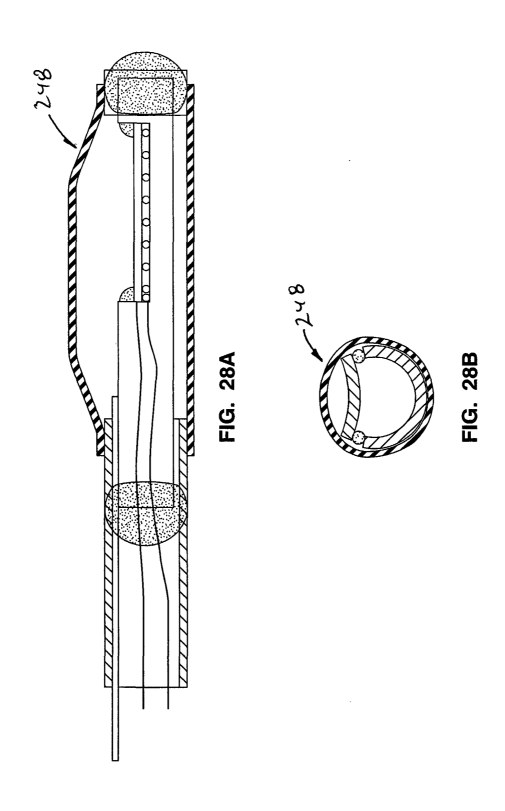
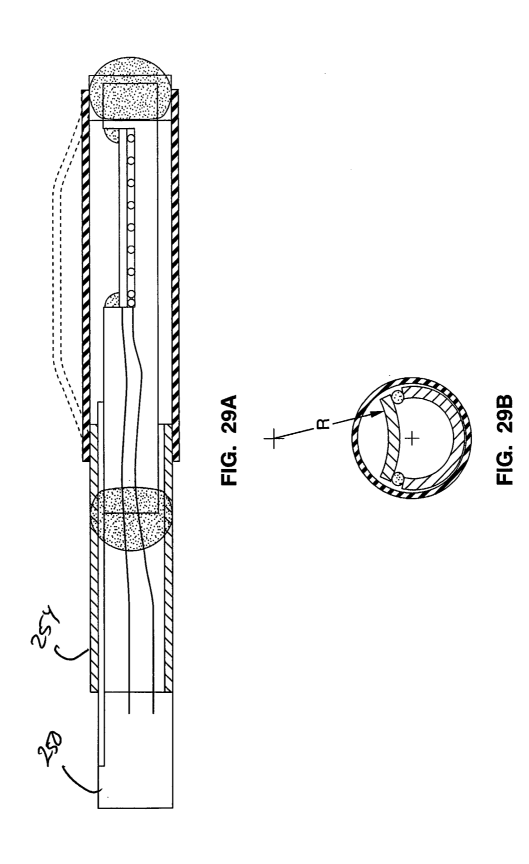
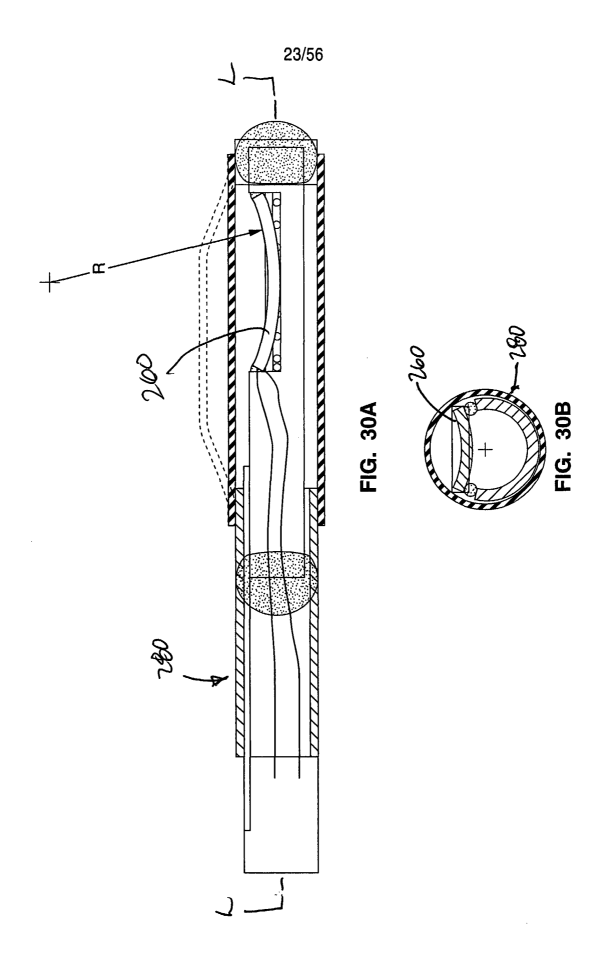


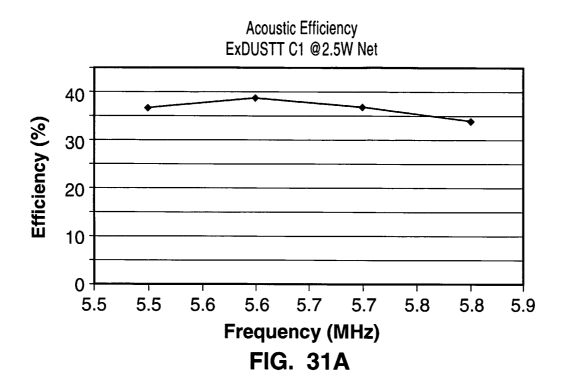
FIG. 26

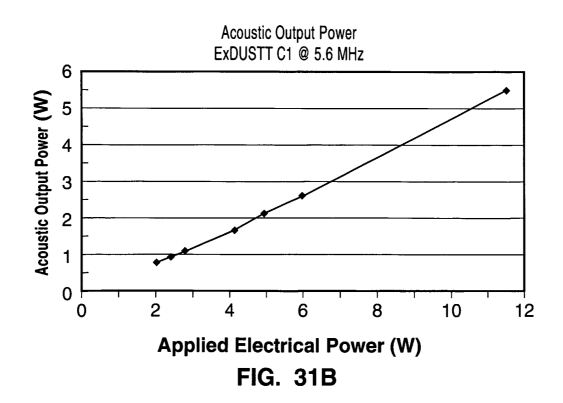


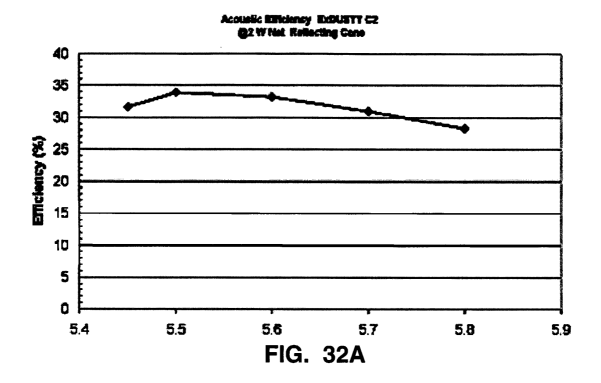












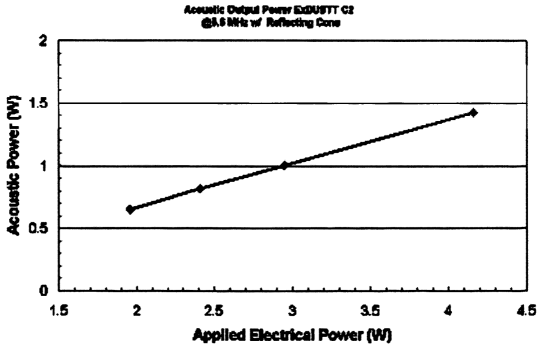


FIG. 32B

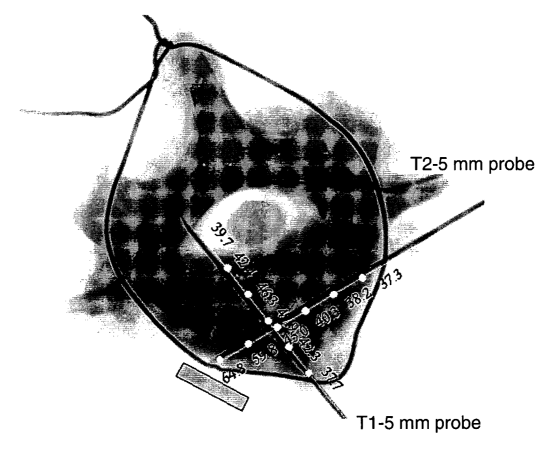
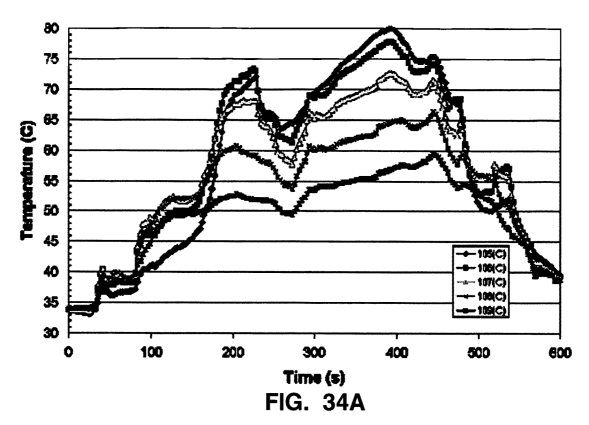
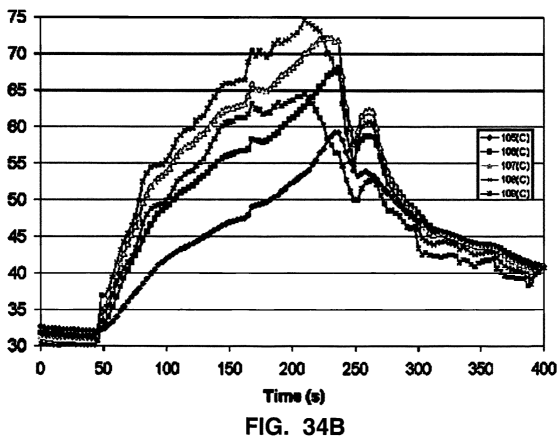
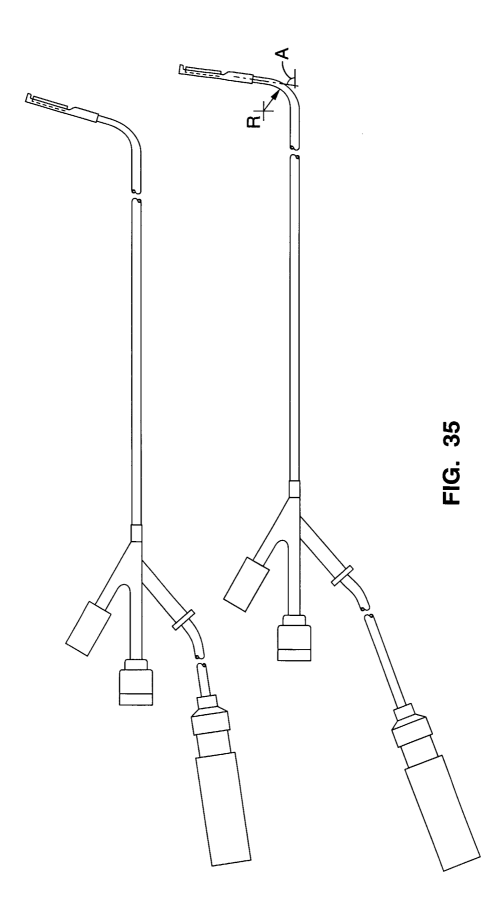
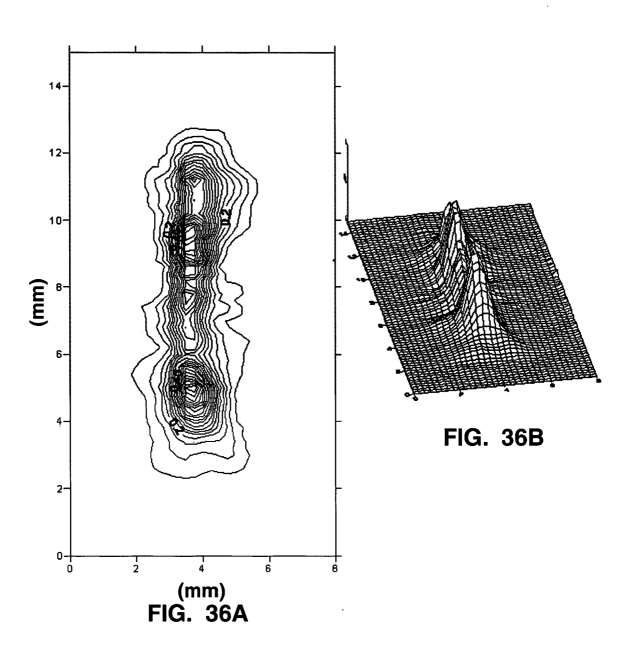


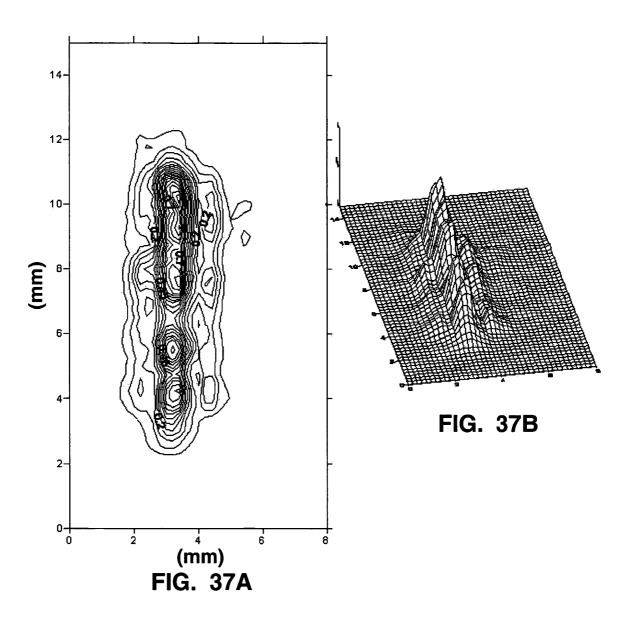
FIG. 33











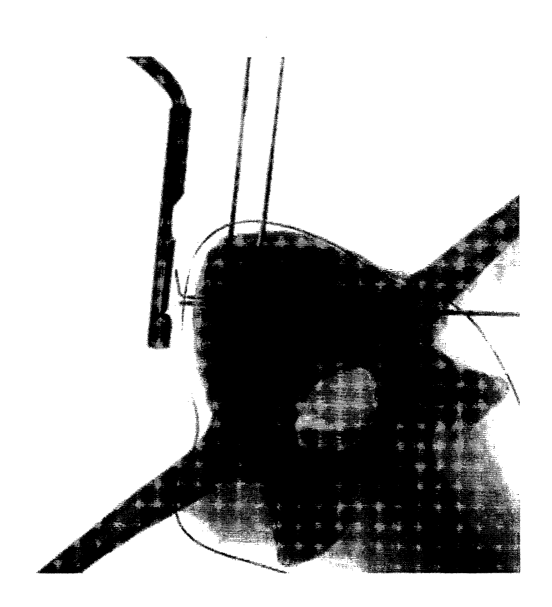


FIG. 38

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3.5mm curvilinear ExDUSTT ex vivo (0°C cooling)

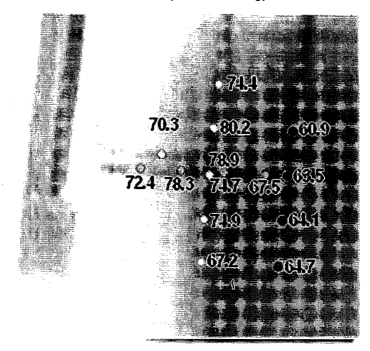


FIG. 39

3.5 curvilinear ExDUSTT ex vivo (0 C cooling)

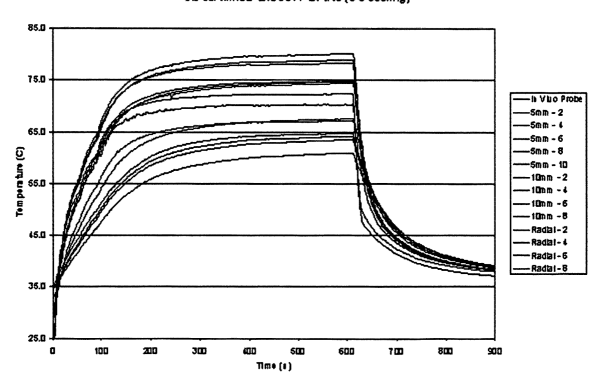


FIG. 40

3.5mm curvilinear ExDUSTT ex vivo (0 C cooling)

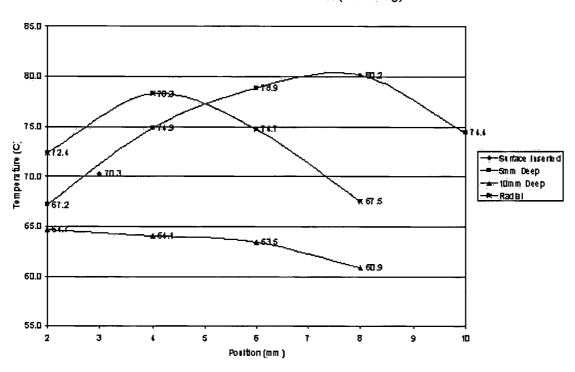


FIG. 41

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3.5mm curvilinear ExDUSTT ex vivo (RT cooling)

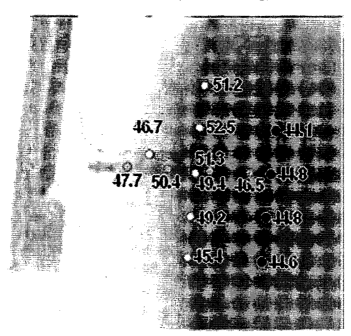


FIG. 42

3.5mm curvilinear Ex DUSTT ex vivo (RT cooling)

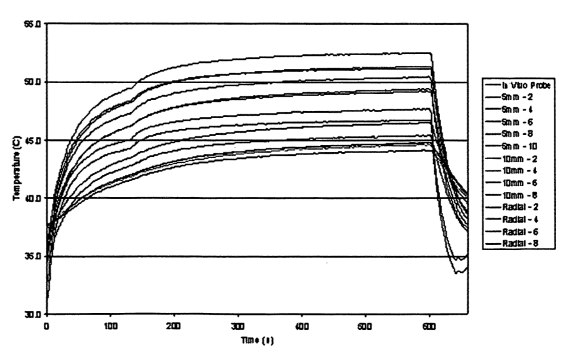


FIG. 43

3:5mm curvilinear ExDUSTT ex vivo (RT cooling)

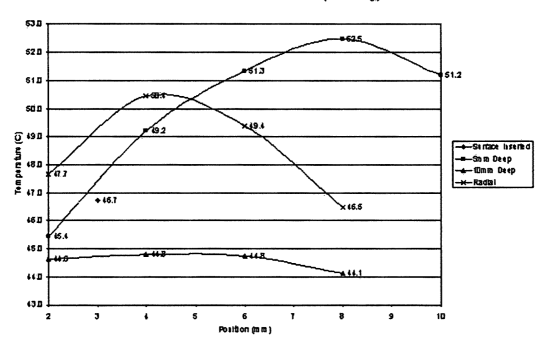


FIG. 44

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3.5mm curvilinear ExDUSTT ex vivo (0°C cooling)

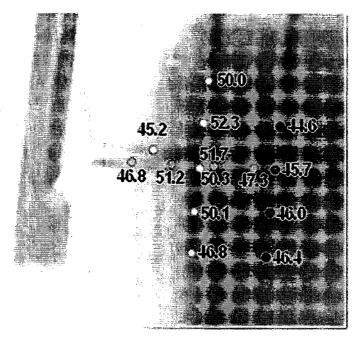


FIG. 45

3.5mm curvilinear ExDUSTT ex vivo (0 C cooling)

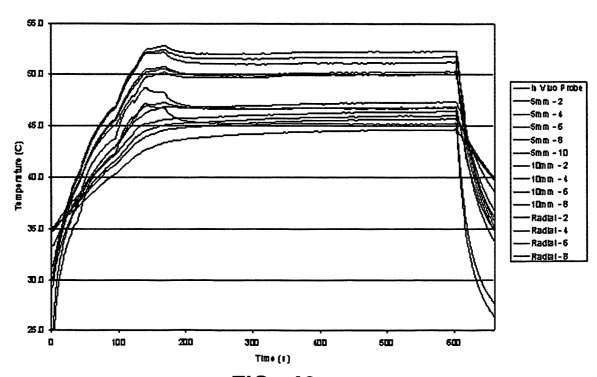


FIG. 46

3.5mm curvilinear ExDUSTT ex vivo (0 C cooling)

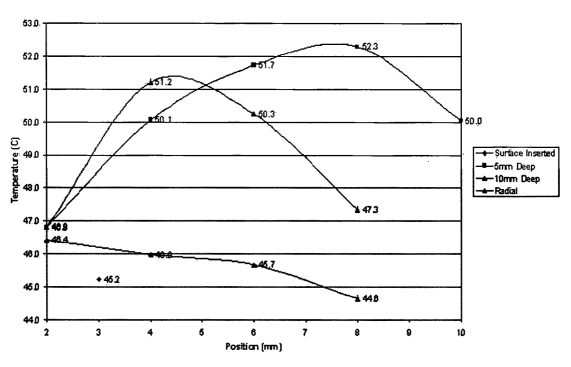


FIG. 47

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2.5mm curvilinear ExDUSTT ex vivo (0°C cooling)

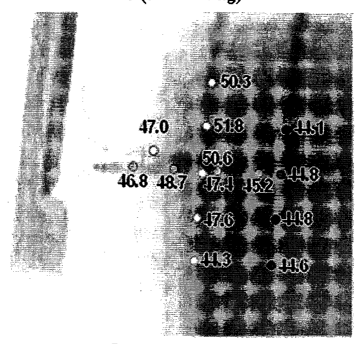


FIG. 48

2.5mm curvillear ExDUSTT ex vivo (0 C Cooling)

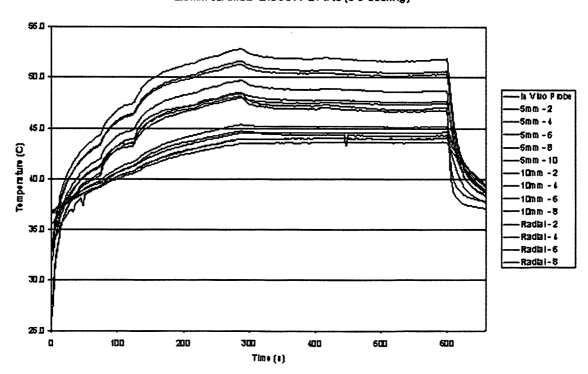


FIG. 49

2.5mm curvilinear ExDUSTT ex vivo (0 C cooling)

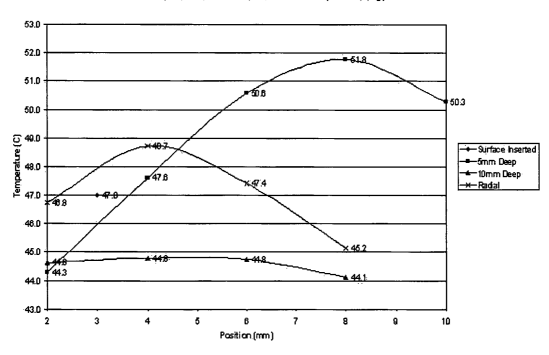


FIG. 50

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2.5mm curvilinear ExDUSTT ex vivo (RT cooling)

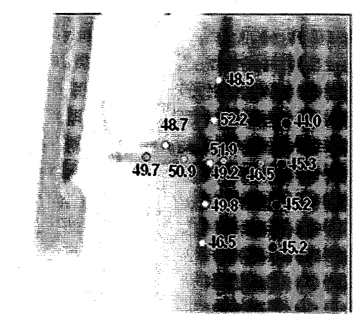


FIG. 51

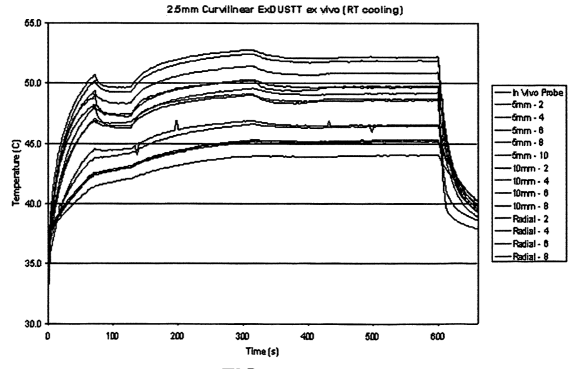


FIG. 52

2.5mm curvilinear ExDUSTT ex vivo (RT cooling)

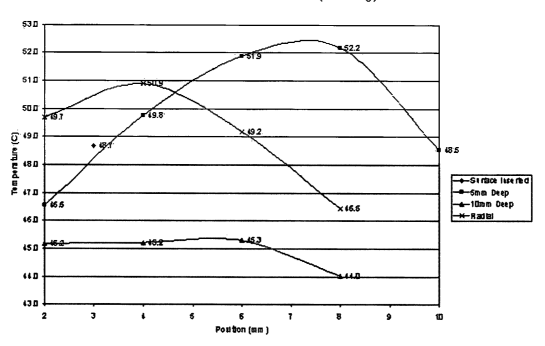
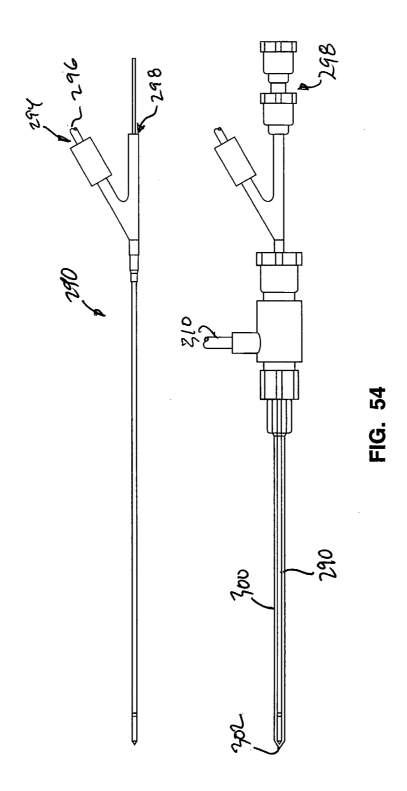
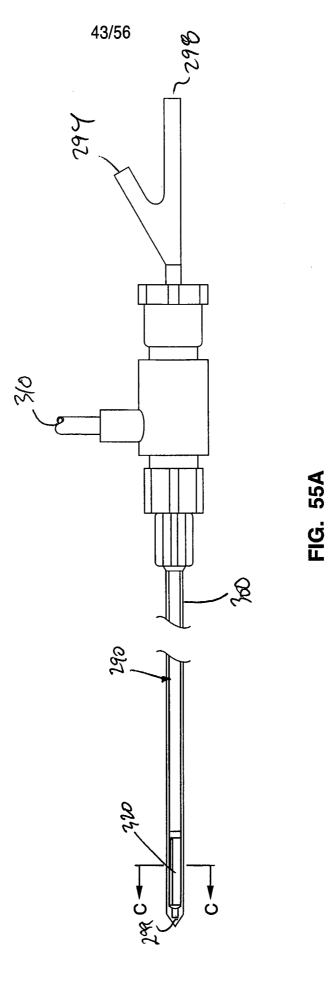
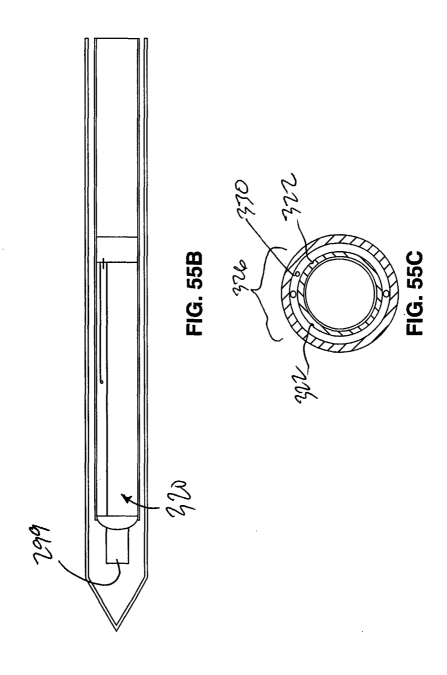


FIG. 53







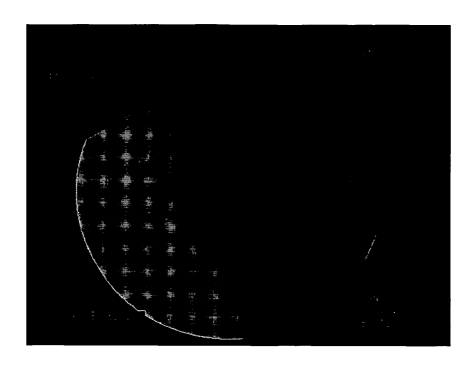


FIG. 56A

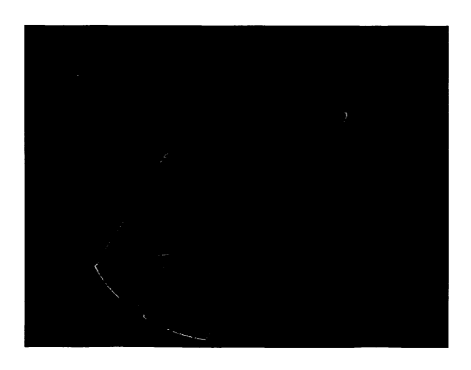
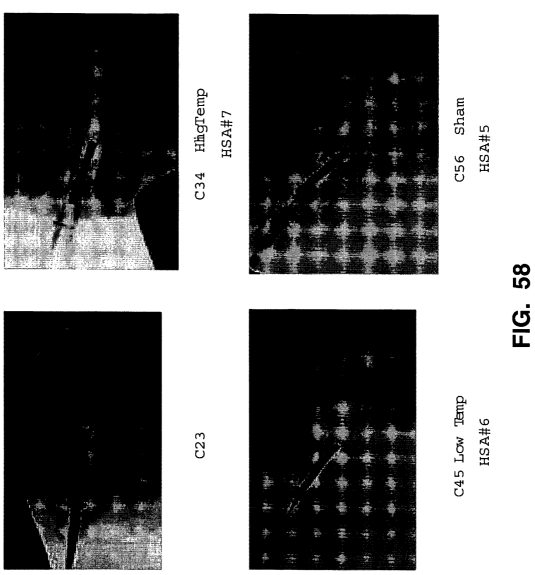
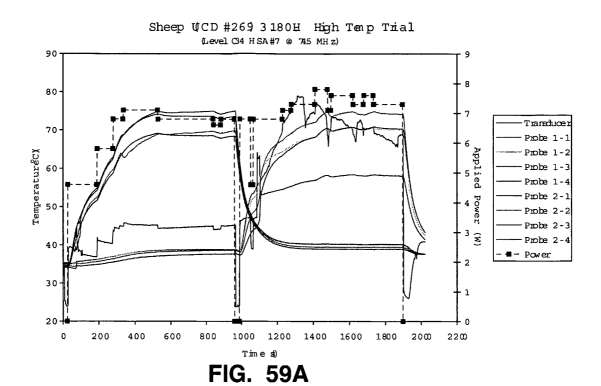


FIG. 56B

 2.99×10^{10} 3.34×10^{5} 1.07×10^4 7.35 x 10^1 Dose (EM_{43°C}) **StDev**_{Tave} () () 1.1 Table 3. Thermal Dosimetry Data (Hedgehog Sheep Spine Study) T_{Ave} 74.2 57.8 52.8 43.9 75.0 58.4 55.4 50.7 (C) Max
Min
Min
Max
Min
Min
Min
Max
Min
Min
Max Applica tor HSA#6 HSA#7 HSA#5 None C23 Sham C34 C56 Sham High Case C45 Low

FIG. 57





Sheep U(CD #29) 3180H bw Temp Trial (Level C45 H SA #6 @ 735 MHz) 80 Transducer 70 Applied TemperaturéCX 60 Power Probe 2-3 (W) Probe 2-4 40 ■ - Power 30 20 0 200 800 1000 1800 0 4 00 600 1200 1400 1600 Time s)

FIG. 59B

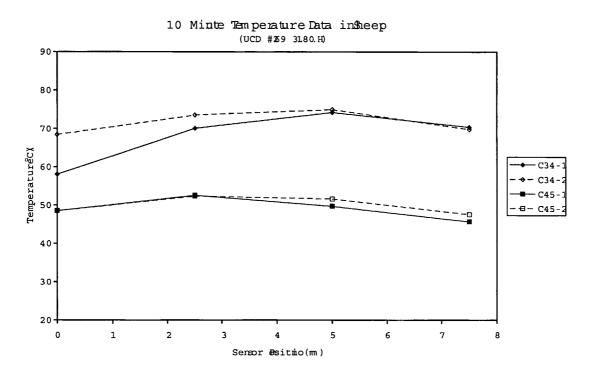


FIG. 59C

	Dose	$(EM_{43}\cdot c)$	1.7×10^9	72786	347	9		
	StDev _{Tave}	(C)	9.0	9.0	0.3	0.7		
metry Data	Tave	(C)	70.2	55.7	48.0	42.3		
Spine Study Thermal Dosimetry Data	T	(C)	71.3	56.3	49.2	43.0		
ine Study			Max	Min	Max	Min		
			ISA#D	ISA#11	ISA#D		ISA#D	
Table: Hedgehog Sheep	Case		C23	High	C34	Low	C45	i

Rise Time Period of time to reach target treatment temperature

StDev_{Tave} Standard deviation of temperature during 10 minute steady-state treatment period Tave Average temperature during the 10 minute steady- state treatment period

Dose Accumulated thermal dose expressed in equivalent minutes at 43°C (EM43°C)

FIG. 60



High Temp Trial C2/3

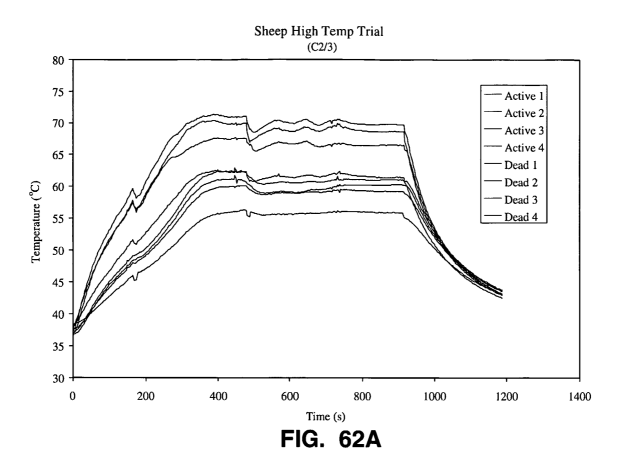


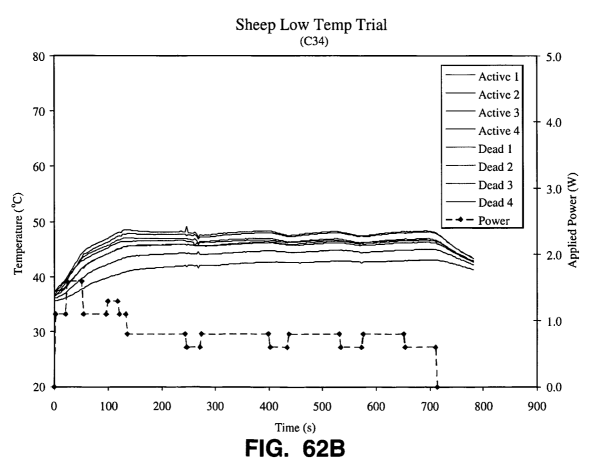
Low Temp Trial C3/4

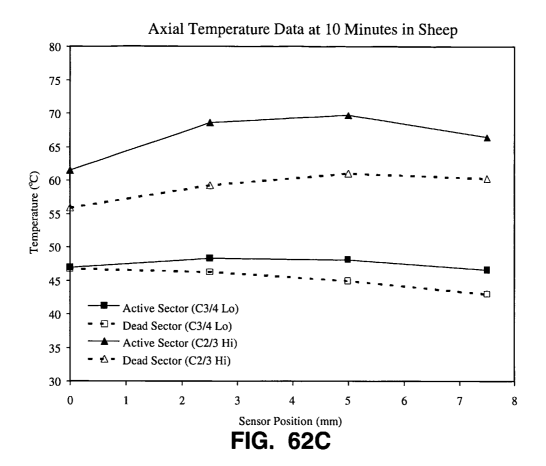


C4/5 Sham

FIG. 61







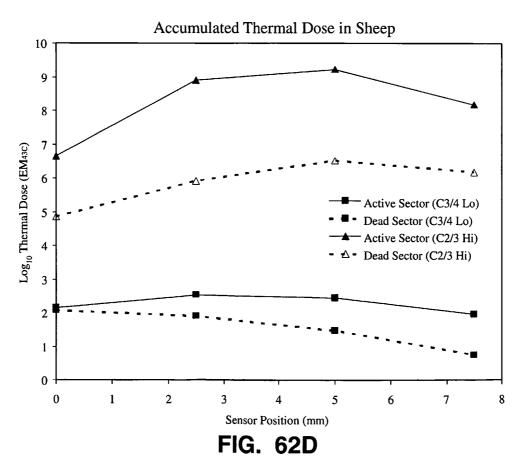
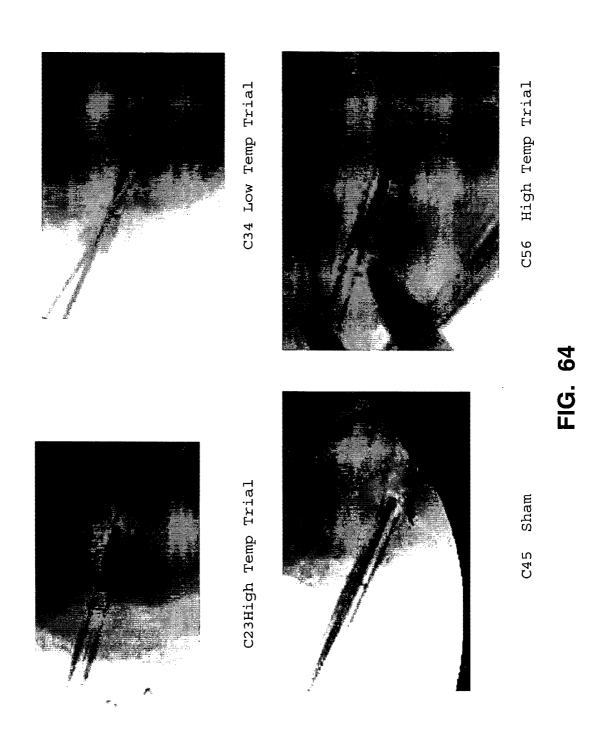


Table: Hedgehog Sh	dgehog Sheep Spine St	ndy	 Thermal Dosimetry Data 	metry Data		
Case	Applicator		L	TAve	StDev _{Tave}	Dose
			(C)	(C)	(°C)	$(EM_{43} \cdot c)$
C23	HSA#6	lax	80.0	77.4	6.0	2.54×10^{11}
High	M	lin .	63.6	61.6	0.0	5.05×10^6
C34	HSA#6 M	Max	53.7	53.0	0.4	1.12×10^4
Low	M	lin	48.2	47.8	0.1	3.12×10^2
C45	M	ax				
Sham	M	lin				
C26	HSA#6 M	lax	80.2	77.5	1.2	2.91×10^{11}
High	M	lin	63.6	69.3	1.1	2.72×10^6

 T_{ave} Average temperature during the 10 minute steady - state treatment period StDev_{Tave} Standard deviation of temperature during 10 minute steady - state treatment period Dose Accumulated thermal dose expressed in equivalent minutes at 43 $^{\circ}$ C (EM43 $^{\circ}$ C)

<u>:1G.</u> 63



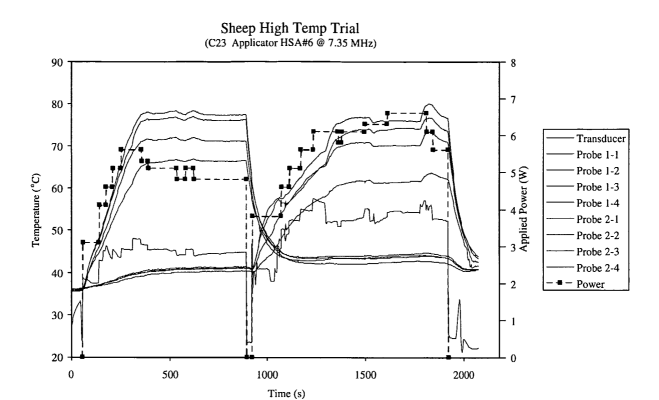


FIG. 65A

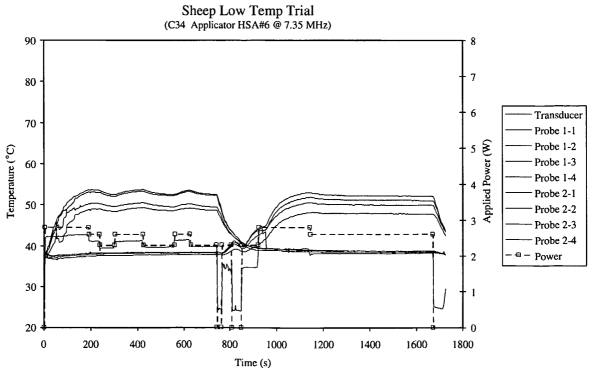


FIG. 65B

Sheep High Temp Trial (C56 Applicator HSA#6 @ 7.35 MHz)

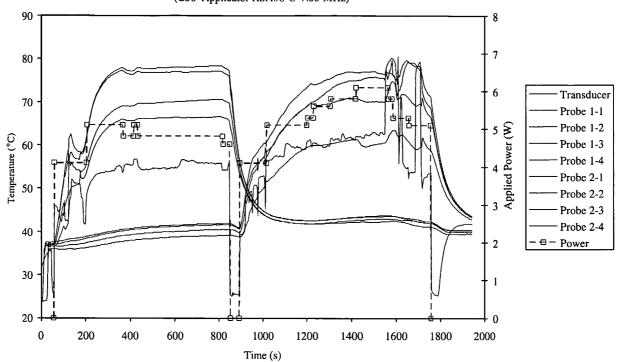


FIG. 65C

Axial Temperature Data at 10 Minutes in Sheep

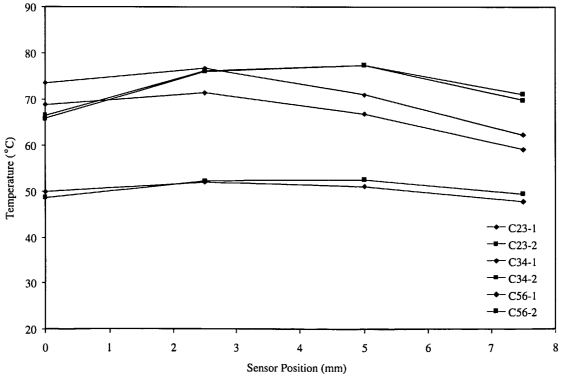


FIG. 65D