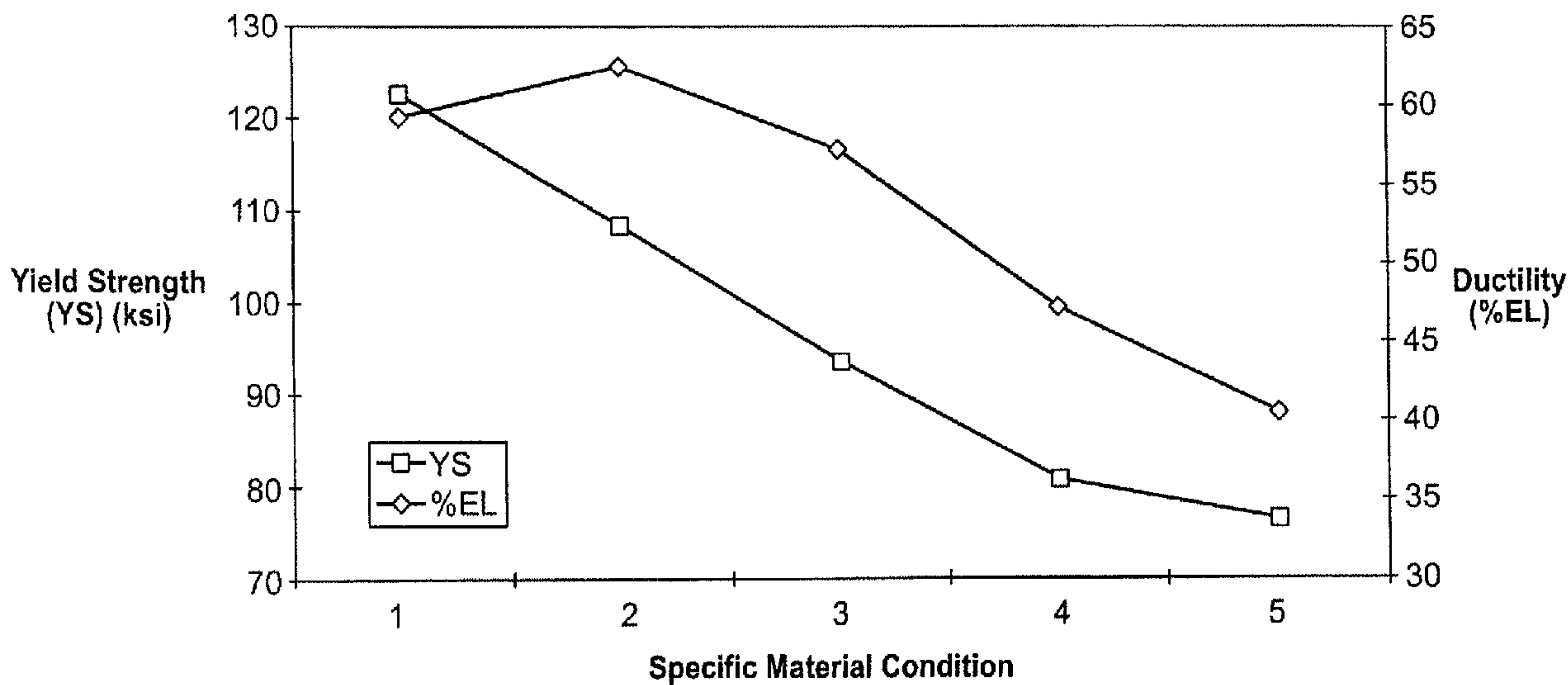




(22) Date de dépôt/Filing Date: 2005/06/08
 (41) Mise à la disp. pub./Open to Public Insp.: 2005/12/09
 (30) Priorité/Priority: 2004/06/09 (10/863,914) US

(51) Cl.Int.⁷/Int.Cl.⁷ C22C 38/44
 (71) Demandeur/Applicant:
CORDIS CORPORATION, US
 (72) Inventeurs/Inventors:
BURGERMEISTER, ROBERT, US;
GRISHABER, RANDY-DAVID BURCE, US
 (74) Agent: SIM & MCBURNEY

(54) Titre : ALLIAGE BIOCOMPATIBLE AMELIORE A BASE DE COBALT, DE NICKEL ET DE CHROME, POUVANT SERVIR A FABRIQUER DES DISPOSITIFS MEDICAUX IMPLANTABLES
 (54) Title: AN IMPROVED COBALT-NICKEL-CHROMIUM BIOCOMPATIBLE ALLOY FOR IMPLANTABLE MEDICAL DEVICES



(57) **Abrégé/Abstract:**

A biocompatible solid-solution alloy may be formed into any number of implantable medical devices. The solid-solution alloy comprises a combination of elements in specific ratios that make it magnetic resonance imaging compatible while retaining the characteristics required for implantable medical devices. The biocompatible solid-solution alloy is a cobalt-chromium alloy having substantially reduced iron and/or Silicon content.

CRD-5095 USANP

ABSTRACT OF THE INVENTION

A biocompatible solid-solution alloy may be formed into any number of implantable medical devices. The solid-solution alloy comprises a combination
5 of elements in specific ratios that make it magnetic resonance imaging compatible while retaining the characteristics required for implantable medical devices. The biocompatible solid-solution alloy is a cobalt-chromium alloy having substantially reduced iron and/or Silicon content.

10

CRD-5095 USANP

AN IMPROVED COBALT-NICKEL-CHROMIUM BIOCOMPATIBLE ALLOY
FOR IMPLANTABLE MEDICAL DEVICES

BACKGROUND OF THE INVENTION

5

1. Field of the Invention

The present invention relates to alloys for use in manufacturing or fabricating implantable medical devices, and more particularly, to implantable
10 medical devices manufactured or fabricated from alloys that are magnetic resonance imaging compatible.

2. Discussion of the Related Art

15 Percutaneous transluminal angioplasty (PTA) is a therapeutic medical procedure used to increase blood flow through an artery. In this procedure, the angioplasty balloon is inflated within the stenosed vessel, or body passageway, in order to shear and disrupt the wall components of the vessel to obtain an enlarged lumen. With respect to arterial stenosed lesions, the relatively
20 incompressible plaque remains unaltered, while the more elastic medial and adventitial layers of the body passageway stretch around the plaque. This process produces dissection, or a splitting and tearing, of the body passageway wall layers, wherein the intima, or internal surface of the artery or body passageway, suffers fissuring. This dissection forms a "flap" of underlying tissue,
25 which may reduce the blood flow through the lumen, or completely block the lumen. Typically, the distending intraluminal pressure within the body passageway can hold the disrupted layer, or flap, in place. If the intimal flap created by the balloon dilation procedure is not maintained in place against the expanded intima, the intimal flap can fold down into the lumen and close off the
30 lumen, or may even become detached and enter the body passageway. When the intimal flap closes off the body passageway, immediate surgery is necessary to correct the problem.

CRD-5095 USANP

Recently, transluminal prostheses have been widely used in the medical arts for implantation in blood vessels, biliary ducts, ureters, or other similar organs of the living body. These prostheses are commonly referred to as stents and are used to maintain, open, or dilate tubular structures. An example of a commonly used stent is given in U.S. Patent No. 4,733,665 to Palmaz. Such stents are often referred to as balloon expandable stents. Typically the stent is made from a solid tube of stainless steel. Thereafter, a series of cuts are made in the wall of the stent. The stent has a first smaller diameter, which permits the stent to be delivered through the human vasculature by being crimped onto a balloon catheter. The stent also has a second, expanded diameter, upon application of a radially, outwardly directed force, by the balloon catheter, from the interior of the tubular shaped member.

However, one concern with such stents is that they are often impractical for use in some vessels such as the carotid artery. The carotid artery is easily accessible from the exterior of the human body, and is close to the surface of the skin. A patient having a balloon expandable stent made from stainless steel or the like, placed in their carotid artery, might be susceptible to severe injury through day-to-day activity. A sufficient force placed on the patient's neck could cause the stent to collapse, resulting in injury to the patient. In order to prevent this, self-expanding stents have been proposed for use in such vessels. Self-expanding stents act like springs and will recover to their expanded or implanted configuration after being crushed.

The prior art makes reference to the use of alloys such as Nitinol (Ni-Ti alloy), which have shape memory and/or superelastic characteristics, in medical devices, which are designed to be inserted into a patient's body, for example, self-expanding stents. The shape memory characteristics allow the devices to be deformed to facilitate their insertion into a body lumen or cavity and then be heated within the body so that the device returns to its original shape. Superelastic characteristics, on the other hand, generally allow the metal to be deformed and restrained in the deformed condition to facilitate the insertion of the medical device containing the metal into a patient's body, with such

CRD-5095 USANP

deformation causing the phase transformation. Once within the body lumen, the restraint on the superelastic member can be removed, thereby reducing the stress therein so that the superelastic member can return to its original undeformed shape by the transformation back to the original phase.

5

One concern with self-expanding stents and with other medical devices formed from superelastic materials, is that they may exhibit reduced radiopacity under X-ray fluoroscopy. To overcome this problem, it is common practice to attach markers, made from highly radiopaque materials, to the stent, or to use radiopaque materials in plating or coating processes. Those materials typically include gold, platinum, or tantalum. The prior art makes reference to these markers or processes in U.S. Patent No. 5,632,771 to Boatman et al., U.S. Patent No. 6,022,374 to Imran, U.S. Patent No. 5,741,327 to Frantzen, U.S. Patent No. 5,725,572 to Lam et al., and U.S. Patent No. 5,800,526 to Anderson et al. However, due to the size of the markers and the relative position of the materials forming the markers in the galvanic series versus the position of the base metal of the stent in the galvanic series, there is a certain challenge to overcome; namely, that of galvanic corrosion. Also, the size of the markers increases the overall profile of the stent. In addition, typical markers are not integral to the stent and thus may interfere with the overall performance of the stent as well as become dislodged from the stent.

A concern with both balloon expandable and self-expandable stents is magnetic resonance imaging compatibility. Currently available metallic stents are known to cause artifacts in magnetic resonance generated images. In general, metals having a high magnetic permeability cause artifacts, while metals having a low magnetic permeability cause less or substantially no artifacts. In other words, if the stent or other medical device is fabricated from a metal or metals having a low magnetic permeability, then less artifacts are created during magnetic resonance imaging which in turn allows more tissue in proximity to the stent or other medical device to be imaged.

CRD-5095 USANP

Artifacts created under magnetic resonance imaging are promoted by local magnetic field inhomogeneities and eddy currents induced by the magnetic field generated by the magnetic resonance imaging machine. The strength of the magnetic field disruption is proportional to the magnetic permeability of the metallic stent or other medical device. In addition, signal attenuation within the stent is caused by radio frequency shielding of the metallic stent or other medical device material. Essentially, the radio frequency signals generated by the magnetic resonance imaging machine may become trapped within the cage like structure of the stent or other medical device. Induced eddy currents in the stent may also lead to a lower nominal radio frequency excitation angle inside the stent. This has been shown to attenuate the signal acquired by the receiver coil of the magnetic resonance imaging device. Artifact related signal changes may include signal voids or local signal enhancements which in turn degrades the diagnostic value of the tool.

15

Accordingly, there is a need to develop materials for implantable medical devices, such as stents, that are magnetic resonance imaging compatible while retaining the toughness, durability and ductility properties required of implantable medical devices such as stents.

20

SUMMARY OF THE INVENTION

The present invention overcomes the diagnostic tool limitations associated with currently available implantable medical devices as briefly described above.

25

In accordance with one aspect, the present invention is directed to a biocompatible, load-carrying metallic structure. The metallic structure being formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, manganese in the range from about 0

30

CRD-5095 USANP

weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, iron in an amount not to exceed 0.12 weight percent, Silicon in an amount not to exceed 0.12 weight percent and the remainder cobalt.

In accordance with another aspect, the present invention is directed to a biocompatible, load-carrying metallic structure. The metallic structure being formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, Silicon in the range from about 0.2 weight percent to about 0.5 weight percent, iron in an amount not to exceed 0.12 weight percent and the remainder cobalt.

20

In accordance with another aspect, the present invention is directed to a biocompatible, load-carrying metallic structure. The metallic structure being formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, iron in the range from about 0 weight percent to about 3 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, Silicon in an amount not to exceed 0.12 weight percent and the remainder cobalt.

30

CRD-5095 USANP

The biocompatible alloy for implantable medical devices of the present invention offers a number of advantages over currently utilized alloys. The biocompatible alloy of the present invention is magnetic resonance imaging compatible, is less brittle than other alloys, has enhanced ductility and toughness, and has increased durability. The biocompatible alloy also maintains the desired or beneficial characteristics of currently available alloys including strength and flexibility.

The biocompatible alloy for implantable medical devices of the present invention may be utilized for any number of medical applications, including vessel patency devices such as vascular stents, biliary stents, ureter stents, vessel occlusion devices such as atrial septal and ventricular septal occluders, patent foramen ovale occluders and orthopedic devices such as fixation devices.

The biocompatible alloy of the present invention is simple and inexpensive to manufacture. The biocompatible alloy may be formed into any number of structures or devices. The biocompatible alloy may be thermomechanically processed, including cold-working and heat treating, to achieve varying degrees of strength and ductility. The biocompatible alloy of the present invention may be age hardened to precipitate one or more secondary phases.

BRIEF DESCRIPTION OF THE DRAWINGS

The foregoing and other features and advantages of the invention will be apparent from the following, more particular description of preferred embodiments of the invention, as illustrated in the accompanying drawings.

Figure 1 is a graphical representation of the transition of critical mechanical properties as a function of thermomechanical processing for cobalt-chromium alloys in accordance with the present invention.

CRD-5095 USANP

Figure 2 is a graphical representation of the endurance limit chart as a function of thermomechanical processing for a cobalt-chromium alloy in accordance with the present invention.

5 Figure 3 is a flat layout diagrammatic representation of an exemplary stent fabricated from the biocompatible alloy in accordance with the present invention.

Figure 4 is an enlarged view of the "M" links of the exemplary stent of Figure 3 in accordance with the present invention.

10

Figure 5 is an enlarged view of a portion of the exemplary stent of Figure 3 in accordance with the present invention.

15 **DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS**

Biocompatible, solid-solution alloys may be utilized in the manufacture of any number of implantable medical devices. The biocompatible alloy for implantable medical devices in accordance with the present invention offers a number of advantages over currently utilized medical grade alloys. In particular, the biocompatible alloy of the present invention is magnetic resonance imaging compatible. Magnetic resonance imaging is a valuable diagnostic tool and thus any implantable medical device should preferably be magnetic resonance imaging compatible so that it and surrounding tissue may be accurately imaged. One such medical device where this is particularly relevant is stents.

Coronary stenting is currently the most widely utilized percutaneous coronary intervention. The primary benefit of stenting when compared with balloon-angioplasty alone is a reduction of the restenosis-rate. Nevertheless, in-stent restenosis remains a relatively common clinical scenario. If clinical symptoms suggest in-stent restenosis, x-ray coronary angiography is currently considered the standard for the evaluation of stent integrity. Conventional x-ray angiography has a number of disadvantages, including a small risk of potentially

CRD-5095 USANP

serious complications, the need for a contrast agent containing a form of iodine, and radiation exposure. Accordingly, a noninvasive imaging method for direct assessment of stent lumen integrity would be preferable. Magnetic resonance imaging provides such a method.

5

Currently, the majority of coronary artery stents are ferromagnetic but are considered to be magnetic resonance imaging safe. Although these devices are considered magnetic resonance imaging safe, they traditionally induce image artifacts that may pose inaccurate, clinically relevant inferences when inspected by a clinician. For example, traditional biocompatible cobalt-alloys such as Alloy 10 188 (common tradenames: HA-188 and Haynes 188 from the Haynes International Corporation) can have as much as 3 wt.% iron. Biocompatible metallic alloys that contain strongly ferromagnetic materials such as iron, but not limited thereto, generally exhibit a high magnetic permeability which tends to 15 induce unintended image artifacts. Moreover, traditional biocompatible ferrous-based alloys such as stainless steel may contain significantly greater concentrations of strongly ferromagnetic materials such as iron.

For reference, a traditional stainless steel alloy such as 316L (i.e. UNS 20 S31603) which is broadly utilized as an implantable, biocompatible device material may comprise chromium (Cr) in the range from about 16 to 18 wt.%, nickel (Ni) in the range from about 10 to 14 wt.%, molybdenum (Mo) in the range from about 2 to 3 wt.%, manganese (Mn) in the range up to 2 wt.%, silicon (Si) in the range up to 1 wt.%, with iron (Fe) comprising the balance (approximately 65 25 wt.%) of the composition.

Additionally, a traditional cobalt-based alloy such as Haynes 188 (i.e. UNS R30188) which is also broadly utilized as an implantable, biocompatible device material may comprise nickel (Ni) in the range from about 20 to 24 wt.%, 30 chromium (Cr) in the range from about 21 to 23 wt.%, tungsten (W) in the range from about 13 to 15 wt.%, iron (Fe) in the range up to 3 wt.%, manganese (Mn) in the range up to 1.25 wt.%, silicon (Si) in the range from about 0.2 to 0.5 wt.%, lanthanum (La) in the range from about 0.02 to 0.12 wt.%, boron (B) in the range

CRD-5095 USANP

up to 0.015 wt.% with cobalt (Co) comprising the balance (approximately 38 wt.%) of the composition.

5 In general, elemental additions such as chromium (Cr), nickel (Ni), tungsten (W), manganese (Mn), silicon (Si) and molybdenum (Mo) where added to iron- and/or cobalt-based alloys, where appropriate, to increase or enable desirable performance attributes, including strength, machinability and corrosion resistance within clinically relevant usage conditions.

10 The composition of the material of the present invention does not eliminate ferromagnetic components but rather shifts the 'susceptibility' (i.e. the magnetic permeability) such that the magnetic resonance imaging compatibility may be improved. In addition, the material of the present invention is intended to improve the measurable ductility by minimizing the deleterious effects induced by
15 traditional machining aides such as silicon (Si).

In accordance with an exemplary embodiment, an implantable medical device may be formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range
20 from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbonl in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in
25 the range from about 0 weight percent to about 0.015 weight percent, iron in an amount not to exceed 0.12 weight percent, Silicon in an amount not to exceed 0.12 weight percent and the remainder cobalt.

In accordance with another exemplary embodiment, an implantable
30 medical device may be formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, manganese in

CRD-5095 USANP

the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, Silicon in the range from about 0.2 weight percent to about 0.5 weight percent, iron in an amount not to exceed 0.12 weight percent and the remainder cobalt

In accordance with yet another exemplary embodiment, an implantable medical device may be formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, iron in the range from about 0 weight percent to about 3 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, Silicon in an amount not to exceed 0.12 weight percent and the remainder cobalt.

In contrast to the traditional formulation of this alloy (i.e. Alloy 188 / Haynes 188), the intended composition does not include any elemental iron (Fe) or silicon (Si) above conventional accepted trace impurity levels. Accordingly, this exemplary embodiment will exhibit a marked reduction in 'susceptibility' (i.e. the magnetic permeability) thereby leading to improved magnetic resonance imaging compatibility. Additionally, the exemplary embodiment will exhibit a marked improvement in material ductility and fatigue strength (i.e. cyclic endurance limit strength) due to the elimination of silicon (Si), above trace impurity levels.

The preferred embodiment may be processed from the requisite elementary raw materials, as set-forth above, by first mechanical homogenization (i.e. mixing) and then compaction into a green state (i.e. precursory) form. If

CRD-5095 USANP

necessary, appropriate manufacturing aids such as hydrocarbon based lubricants and/or solvents (e.g. mineral oil, machine oils, kerosene, isopropanol and related alcohols) be used to ensure complete mechanical homogenization. Additionally, other processing steps such as ultrasonic agitation of the mixture
5 followed by cold compaction to remove any unnecessary manufacturing aides and to reduce void space within the green state may be utilized. It is preferable to ensure that any impurities within or upon the processing equipment from prior processing and/or system construction (e.g. mixing vessel material, transfer containers, etc.) be sufficiently reduced in order to ensure that the green state
10 form is not unnecessarily contaminated. This may be accomplished by adequate cleaning of the mixing vessel before adding the constituent elements by use of surfactant based cleaners to remove any loosely adherent contaminants.

Initial melting of the green state form into a ingot of desired composition,
15 is achieved by vacuum induction melting (VIM) where the initial form is inductively heated to above the melting point of the primary constituent elements within a refractory crucible and then poured into a secondary mold within a vacuum environment (e.g. typically less than or equal to 10^{-4} mmHg). The vacuum process ensures that atmospheric contamination is significantly
20 minimized. Upon solidification of the molten pool, the ingot bar is substantially single phase (i.e. compositionally homogenous) with a definable threshold of secondary phase impurities that are typically ceramic (e.g. carbide, oxide or nitride) in nature. These impurities are typically inherited from the precursor elemental raw materials.

25

A secondary melting process termed vacuum arc reduction (VAR) is utilized to further reduce the concentration of the secondary phase impurities to a conventionally accepted trace impurity level (i.e. < 1,500 ppm). Other methods maybe enabled by those skilled in the art of ingot formulation that
30 substantially embodies this practice of ensuring that atmospheric contamination is minimized. In addition, the initial VAR step may be following followed by repetitive VAR processing to further homogenize the solid-solution alloy in the ingot form. From the initial ingot configuration, the homogenized

CRD-5095 USANP

alloy will be further reduced in product size and form by various industrially accepted methods such as, but not limited too, ingot peeling, grinding, cutting, forging, forming, hot rolling and/or cold finishing processing steps so as to produce bar stock that may be further reduced into a desired raw material
5 form.

In this exemplary embodiment, the initial raw material product form that is required to initiate the thermomechanical processing that will ultimately yield a desired small diameter, thin-walled tube, appropriate for interventional
10 devices, is a modestly sized round bar (e.g. one inch in diameter round bar stock) of predetermined length. In order to facilitate the reduction of the initial bar stock into a much smaller tubing configuration, an initial clearance hole must be placed into the bar stock that runs the length of the product. These tube hollows (i.e. heavy walled tubes) may be created by 'gun-drilling' (i.e. high
15 depth to diameter ratio drilling) the bar stock. Other industrially relevant methods of creating the tube hollows from round bar stock may be utilized by those skilled-in-the-art of tube making.

Consecutive mechanical cold-finishing operations such as drawing
20 through a compressive outer-diameter (OD), precision shaped (i.e. cut), circumferentially complete, diamond die using any of the following internally supported (i.e. inner diameter, ID) methods, but not necessarily limited to these conventional forming methods, such as hard mandrel (i.e. relatively long traveling ID mandrel also referred to as rod draw), floating-plug (i.e. relatively
25 short ID mandrel that 'floats' within the region of the OD compressive die and fixed-plug (i.e. the ID mandrel is 'fixed' to the drawing apparatus where relatively short workpieces are processed) drawing. These process steps are intended to reduce the outer-diameter (OD) and the corresponding wall thickness of the initial tube hollow to the desired dimensions of the finished
30 product.

When necessary, tube sinking (i.e. OD reduction of the workpiece without inducing substantial tube wall reduction) is accomplished by drawing

CRD-5095 USANP

the workpiece through a compressive die without internal support (i.e. no ID mandrel). Conventionally, tube sinking is typically utilized as a final or near-final mechanical processing step to achieve the desired dimensional attributed of the finished product.

5

Although not practically significant, if the particular compositional formulation will support a single reduction from the initial raw material configuration to the desired dimensions of the finished product, in process heat-treatments will not be necessary. Where necessary to achieve intended mechanical properties of the finished product, a final heat-treating step is utilized.

Conventionally, all metallic alloys in accordance with the present invention will require incremental dimensional reductions from the initial raw material configuration to reach the desired dimensions of the finished product. This processing constraint is due to the material's ability to support a finite degree of induced mechanical damage per processing step without structural failure (e.g. strain-induced fracture, fissures, extensive void formation, etc.).

In order to compensate for induced mechanical damage (i.e. cold-working) during any of the aforementioned cold-finishing steps, periodic thermal heat-treatments are utilized to stress-relieve (i.e. minimization of deleterious internal residual stresses that are the result of processes such as cold-working) thereby increasing the workability (i.e. ability to support additional mechanical damage without measurable failure) the workpiece prior to subsequent reductions. These thermal treatments are typically, but not necessarily limited to, conducted within a relatively inert environment such as an inert gas furnace (e.g. nitrogen, argon, etc.), a oxygen rarified hydrogen furnace, a conventional vacuum furnace and under less common process conditions, atmospheric air. When vacuum furnaces are utilized, the level of vacuum (i.e. subatmospheric pressure), typically measured in units of mmHg or torr (where 1 mmHg is equal to 1 unit torr), shall be sufficient to ensure that excessive and deteriorative high temperature oxidative processes are not

CRD-5095 USANP

functionally operative during heat treatment. This process may usually be achieved under vacuum conditions of 10^{-4} mmHg (0.0001 torr) or less (i.e. lower magnitude).

5 The stress relieving heat treatment temperature is typically held constant between 82 to 86% of the conventional melting point (i.e. industrially accepted liquidus temperature, 0.82 to 0.86 homologous temperatures) within an adequately sized isothermal region of the heat-treating apparatus. The workpiece undergoing thermal treatment is held within the isothermal
10 processing region for a finite period of time that is adequate to ensure that the workpiece has reached a state of thermal equilibrium and for that sufficient time is elapsed to ensure that the reaction kinetics (i.e. time dependent material processes) of stress-relieving and/or process annealing, as appropriate, is adequately completed. The finite amount of time that the
15 workpiece is held within the processing is dependent upon the method of bringing the workpiece into the process chamber and then removing the workpiece upon completion of heat treatment. Typically, this process is accomplished by, but not limited to, use of a conventional conveyor-belt apparatus or other relevant mechanical assist devices. In the case of the
20 former, the conveyor belt speed and appropriate finite dwell-time, as necessary, within the isothermal region is controlled to ensure that sufficient time at temperature is utilized so as to ensure that the process is completed as intended.

25 When necessary to achieve desired mechanical attributes of the finished product, heat-treatment temperatures and corresponding finite processing times may be intentionally utilized that are not within the typical range of 0.82 to 0.86 homologous temperatures. Various age hardening (i.e. a process that induces a change in properties at moderately elevated
30 temperatures, relative to the conventional melting point, that does not induce a change in overall chemical composition change in the metallic alloy being processed) processing steps may be carried out, as necessary, in a manner consistent with those previously described at temperatures substantially below

CRD-5095 USANP

0.82 to 0.86 homologous temperature. For Co-based alloys in accordance with the present invention, these processing temperatures may be varied between and inclusive of approximately 0.29 homologous temperature and the
5 aforementioned stress relieving temperature range. The workpiece undergoing thermal treatment is held within the isothermal processing region for a finite period of time that is adequate to ensure that the workpiece has reached a state of thermal equilibrium and for that sufficient time is elapsed to ensure that the reaction kinetics (i.e. time dependent material processes) of age hardening, as appropriate, is adequately completed prior to removal from the processing
10 equipment.

In some cases for Co-based alloys in accordance with the present invention, the formation of secondary-phase ceramic compounds such as carbide, nitride and/or oxides will be induced or promoted by age hardening heat
15 treating. These secondary-phase compounds are typically, but not limited to, for Co-based alloys in accordance with the present invention, carbides which precipitate along thermodynamically favorable regions of the structural crystallographic planes that comprise each grain (i.e. crystallographic entity) that make-up the entire polycrystalline alloy. These secondary-phase carbides can
20 exist along the intergranular boundaries as well as within each granular structure (i.e. intragranular). Under most circumstances for Co-based alloys in accordance with the present invention, the principal secondary phase carbides that are stoichiometrically expected to be present are M_6C where M typically is cobalt (Co). When present, the intermetallic M_6C phase is typically expected to
25 reside intragranularly along thermodynamically favorable regions of the structural crystallographic planes that comprise each grain within the polycrystalline alloy in accordance with the present invention. Although not practically common, the equivalent material phenomena can exist for a single crystal (i.e. monogranular) alloy.

30

Additionally, another prominent secondary phase carbide can also be induced or promoted as a result of age hardening heat treatments. This phase, when present, is stoichiometrically expected to be $M_{23}C_6$ where M typically is

CRD-5095 USANP

chromium (Cr) but is also commonly observed to be cobalt (Co) especially in Co-based alloys. When present, the intermetallic $M_{23}C_6$ phase is typically expected to reside along the intergranular boundaries (i.e. grain boundaries) within a polycrystalline alloy in accordance with the present invention. As previously
5 discussed for the intermetallic M_6C phase, the equivalent presence of the intermetallic $M_{23}C_6$ phase can exist for a single crystal (i.e. monogranular) alloy, albeit not practically common.

In the case of the intergranular $M_{23}C_6$ phase, this secondary phase is
10 conventionally considered most important, when formed in a manner that is structurally and compositionally compatible with the alloy matrix, to strengthening the grain boundaries to such a degree that intrinsic strength of the grain boundaries and the matrix are adequately balanced. By inducing this equilibrium level of material strength at the microstructural level, the overall
15 mechanical properties of the finished tubular product can be further optimized to desirable levels.

In addition to stress relieving and age hardening related heat-treating steps, solutionizing (i.e. sufficiently high temperature and longer processing
20 time to thermodynamically force one of more alloy constituents to enter into solid solution – ‘singular phase’, also referred to as full annealing) of the workpiece may be utilized. For Co-based alloys in accordance with the present invention, the typical solutionizing temperature can be varied between and inclusive of approximately 0.88 to 0.90 homologous temperatures. The
25 workpiece undergoing thermal treatment is held within the isothermal processing region for a finite period of time that is adequate to ensure that the workpiece has reached a state of thermal equilibrium and for that sufficient time is elapsed to ensure that the reaction kinetics (i.e. time dependent material processes) of solutionizing, as appropriate, is adequately completed
30 prior to removal from the processing equipment.

The sequential and selectively ordered combination of thermomechanical processing steps that may comprise but not necessarily

CRD-5095 USANP

include mechanical cold-finishing operations, stress relieving, age hardening and solutionizing can induce and enable a broad range of measurable mechanical properties as a result of distinct and determinable microstructural attributes. This material phenomena can be observed in Figure 1. which
5 shows a chart that exhibits the affect of thermomechanical processing (TMP) such as cold working and in-process heat-treatments on measurable mechanical properties such as yield strength and ductility (presented in units of percent elongation) in accordance with the present invention. In this example, thermomechanical (TMP) groups one (1) through five (5) were subjected to
10 varying combinations of cold-finishing, stress relieving and age hardening and not necessarily in the presented sequential order. In general, the principal isothermal age hardening heat treatment applied to each TMP group varied between about 0.74 to 0.78 homologous temperatures for group (1), about 0.76 to 0.80 homologous temperatures for group (2), about 0.78 to 0.82 homologous
15 temperatures for group (3), about 0.80 to 0.84 homologous temperatures for group (4) and about 0.82 to 0.84 homologous temperatures for group (5). The each workpiece undergoing thermal treatment was held within the isothermal processing region for a finite period of time that was adequate to ensure that the workpiece reached a state of thermal equilibrium and to ensure that
20 sufficient time was elapsed to ensure that the reaction kinetics of age hardening was adequately completed.

More so, the effect of thermomechanical (TMP) on cyclic fatigue properties is on Co-based alloys, in accordance with the present invention, is
25 reflected in Figure 2. Examination of Figure 2. shows the affect on fatigue strength (i.e. endurance limit) as a function of thermomechanical processing for the previously discussed TMP groups (2) and (4). TMP group (2) from this figure as utilized in this specific example shows a marked increase in the fatigue strength (i.e. endurance limit, the maximum stress below which a material can
30 presumably endure an infinite number of stress cycles) over and against the TMP group (4) process.

CRD-5095 USANP

The above-described alloy may be utilized in any number of implantable medical devices. The alloy is particularly advantageous in situations where magnetic resonance imaging is a useful diagnostic tool such as determining in-stent restenosis. Accordingly, although the alloy may be utilized for any implantable medical device, an exemplary stent constructed from the alloy is described below.

Figure 3 is a flat layout of an exemplary embodiment of a stent that may be constructed utilizing the alloy of the present invention. The stent 10 comprises end sets of strut members 12 located at each end of the stent 10 and central sets of strut members 14 connected each to the other by sets of flexible "M" links 16. Each end set of strut members 12 comprises alternating curved sections 18 and diagonal sections 20 connected together to form a closed circumferential structure. The central sets of strut members 14 located longitudinally between the end sets of strut members 14 comprise curved sections 22 and diagonal sections 24 connected together to form a closed circumferential ring-like structure.

Referring to Figure 4 there is illustrated an enlargement of the flexible "M" links 16 of the stent 10. Each "M" link 16 has a circumferential extent, i.e. length, L' above and L'' below line 11. The line 11 is drawn between the attachment points 13 where the "M" link 16 attaches to adjacent curved sections 18 or 22. Such a balanced design preferably diminishes any likelihood of the flexible connecting link 16 from expanding into the lumen of artery or other vessel.

As illustrated in Figure 3, the diagonal sections 20 of the end sets of strut members 12 are shorter in length than the diagonal sections 24 of the central sets of strut members 14. The shorter diagonal sections 20 will preferably reduce the longitudinal length of metal at the end of the stent 10 to improve deliverability into a vessel of the human body. In the stent 10, the widths of the diagonal sections 20 and 24 are different from one another.

CRD-5095 USANP

Referring to Figure 5, there is illustrated an expanded view of a stent section comprising an end set of strut members 12 and a central set of strut members 14. As illustrated, the diagonal sections 24 of the central sets of strut members 14 have a width at the center thereof, T_c , and a width at the end thereof, T_e , wherein T_c is greater than T_e . This configuration allows for increased radiopacity without affecting the design of curved sections 22 that are the primary stent elements involved for stent expansion. In an exemplary embodiment, the curved sections 22 and 18 may be tapered and may have uniform widths with respect to one another as is explained in detail subsequently. The diagonal sections 20 of the end sets of strut members 12 also have a tapered shape. The diagonal sections 20 have a width in the center, T_{c-end} , and a width at the end, T_{e-end} , wherein T_{c-end} is greater than T_{e-end} . Because it is preferable for the end sets of strut members 12 to be the most radiopaque part of the stent 10, the diagonal section 20 center width T_{c-end} of the end sets of strut members 12 is wider than the width T_c of the diagonal section 24. Generally, a wider piece of metal will be more radiopaque. Thus, the stent 10 has curved sections with a single bend connecting the diagonal sections of its sets of strut members, and flexible connecting links connecting the curved sections of its circumferential sets of strut members.

20

The width of the curved sections 22 and 18 taper down as one moves away from the center of the curve until a predetermined minimum width substantially equal to that of their respective diagonal sections 24 and 20. To achieve this taper, the inside arc of the curved sections 22 and 18 have a center that is longitudinally displaced from the center of the outside arc. This tapered shape for the curved sections 22 and 18 provides a significant reduction in metal strain with little effect on the radial strength of the expanded stent as compared to a stent having sets of strut members with a uniform strut width.

This reduced strain design has several advantages. First, it can allow the exemplary design to have a much greater usable range of radial expansion as compared to a stent with a uniform strut width. Second, it can allow the width at the center of the curve to be increased which increases radial strength without

CRD-5095 USANP

greatly increasing the metal strain (i.e. one can make a stronger stent). Finally, the taper reduces the amount of metal in the stent and that should improve the stent thrombogenicity.

5 The curved sections 18 of the end sets of strut members 12 and the curved sections 22 of the central sets of strut members 14 have the same widths. As a result of this design, the end sets of strut members 12, which have shorter diagonal sections 20, will reach the maximum allowable diameter at a level of strain that is greater than the level of strain experienced by the central
10 sets of strut members 14.

It is important to note that although a stent is described, the alloy may be utilized for any number of implantable medical devices.

15 Although shown and described is what is believed to be the most practical and preferred embodiments, it is apparent that departures from specific designs and methods described and shown will suggest themselves to those skilled in the art and may be used without departing from the spirit and scope of the invention. The present invention is not restricted to the particular constructions
20 described and illustrated, but should be constructed to cohere with all modifications that may fall within the scope for the appended claims.

CRD-5095 USANP

WHAT IS CLAIMED IS:

1. A biocompatible, load-carrying metallic structure being formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, iron in an amount not to exceed 0.12 weight percent, Silicon in an amount not to exceed 0.12 weight percent and the remainder cobalt.
2. The biocompatible, load-carrying metallic structure according to claim 1, wherein the solid-solution alloy is constructed through thermomechanical processing to exhibit relatively high strength and low ductility characteristics in the fully cold-worked state.
3. The biocompatible, load-carrying metallic structure according to claim 1, wherein the solid-solution alloy is constructed through thermomechanical processing to exhibit relatively moderate strength and moderate ductility characteristics in the partially cold-worked state.
4. The biocompatible, load-carrying metallic structure according to claim 1, wherein the solid-solution alloy is further constructed through age hardening for a predetermined time within a gaseous environment at a temperature less than the annealing temperature to precipitate one or more secondary phases, including at least one of intragranular and intergranular phases, from a substantially single phase structure.

CRD-5095 USANP

5. The biocompatible, load-carrying metallic structure according to claim 4, wherein the age hardening temperature is in the range from about 750 degrees Fahrenheit to about 2,150 degrees Fahrenheit.

5 6. The biocompatible, load-carrying metallic structure according to claim 4, wherein the age hardening gaseous environment comprises hydrogen, nitrogen, argon and air.

10 7. The biocompatible, load-carrying metallic structure according to claim 3, wherein the solid-solution alloy is further constructed through stress relieving for a predetermined time within a gaseous environment at a temperature less than the annealing temperature while maintaining a substantially single phase to increase toughness and ductility.

15 8. The biocompatible, load-carrying metallic structure according to claim 7, wherein the stress relieving temperature is about or less than 100 degrees Fahrenheit below the annealing temperature.

20 9. The biocompatible, load-carrying metallic structure according to claim 7, wherein the stress relieving gaseous environment comprises hydrogen, nitrogen, argon and air.

25 10. The biocompatible, load-carrying metallic structure according to claim 1, wherein the solid-solution alloy is constructed through thermomechanical processing to exhibit relatively low strength and high ductility characteristics in the fully annealed state.

30 11. The biocompatible, load-carrying metallic structure according to claim 1, wherein the medical device comprises a fixation device.

 12. The biocompatible, load-carrying metallic structure according to claim 1, wherein the medical device comprises an artificial joint implant.

CRD-5095 USANP

13. The biocompatible, load-carrying metallic structure according to claim 3, wherein the solid-solution alloy is further constructed through stress relieving for a predetermined time with a vacuum environment at a temperature less than the annealing temperature while maintaining a substantially single phase to increase toughness and ductility.

14. The biocompatible, load-carrying metallic structure according to claim 13, wherein the stress relieving temperature is about or less than one hundred degrees Fahrenheit below the annealing temperature.

10

15. A biocompatible, load-carrying metallic structure being formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, Silicon in the range from about 0.2 weight percent to about 0.5 weight percent, iron in an amount not to exceed 0.12 weight percent and the remainder cobalt.

20

16. A biocompatible, load-carrying metallic structure being formed from a solid-solution alloy comprising nickel in the range from about 20 weight percent to about 24 weight percent, chromium in the range from about 21 weight percent to about 23 weight percent, tungsten in the range from about 13 weight percent to about 15 weight percent, iron in the range from about 0 weight percent to about 3 weight percent, manganese in the range from about 0 weight percent to about 1.25 weight percent, carbon in the range from about 0.05 weight percent to about 0.15 weight percent, lanthanum in the range from about 0.02 weight percent to about 0.12 weight percent, boron in the range from about 0 weight percent to about 0.015 weight percent, Silicon in an amount not to exceed 0.12 weight percent and the remainder cobalt.

30

FIG. 1

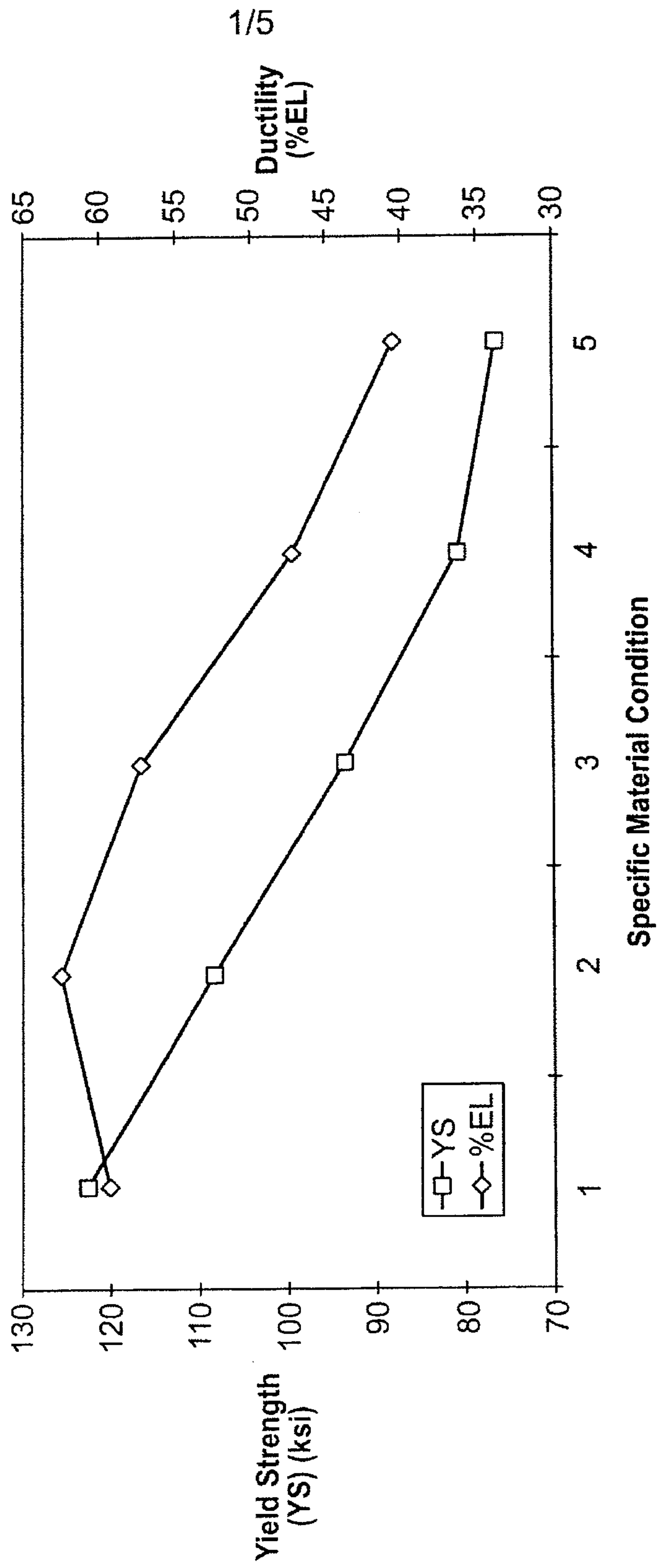


FIG. 2

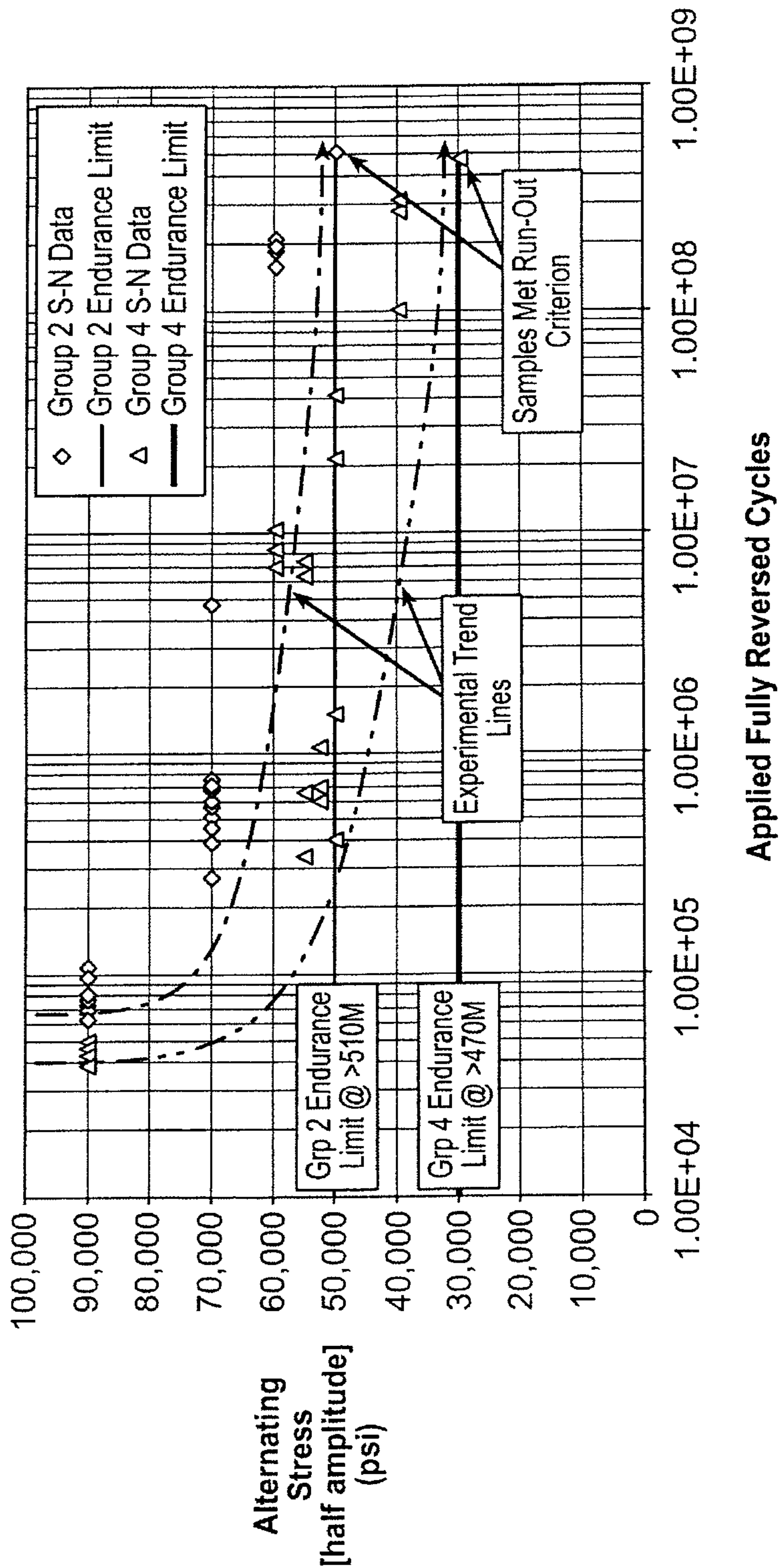


FIG. 3

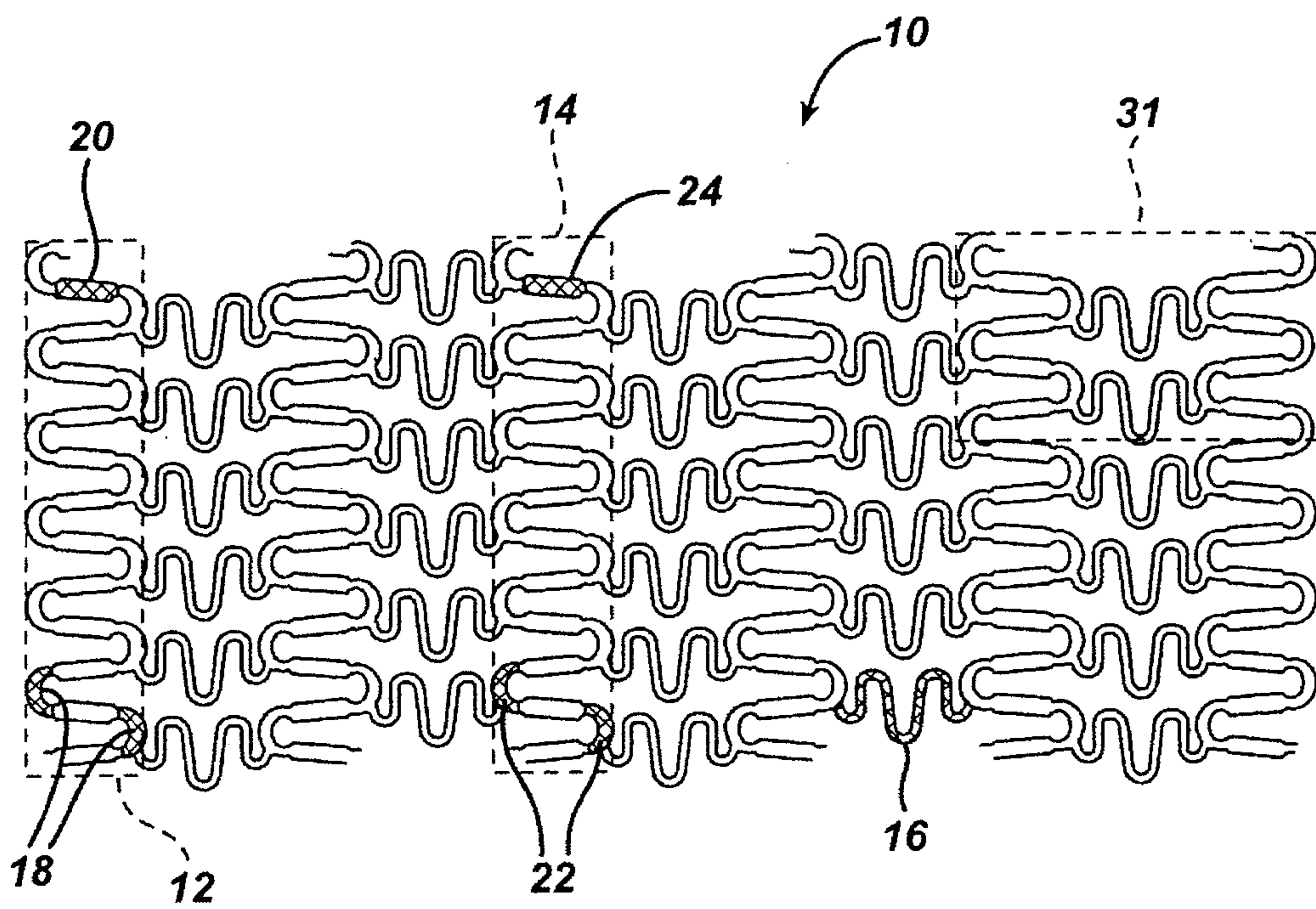


FIG. 4

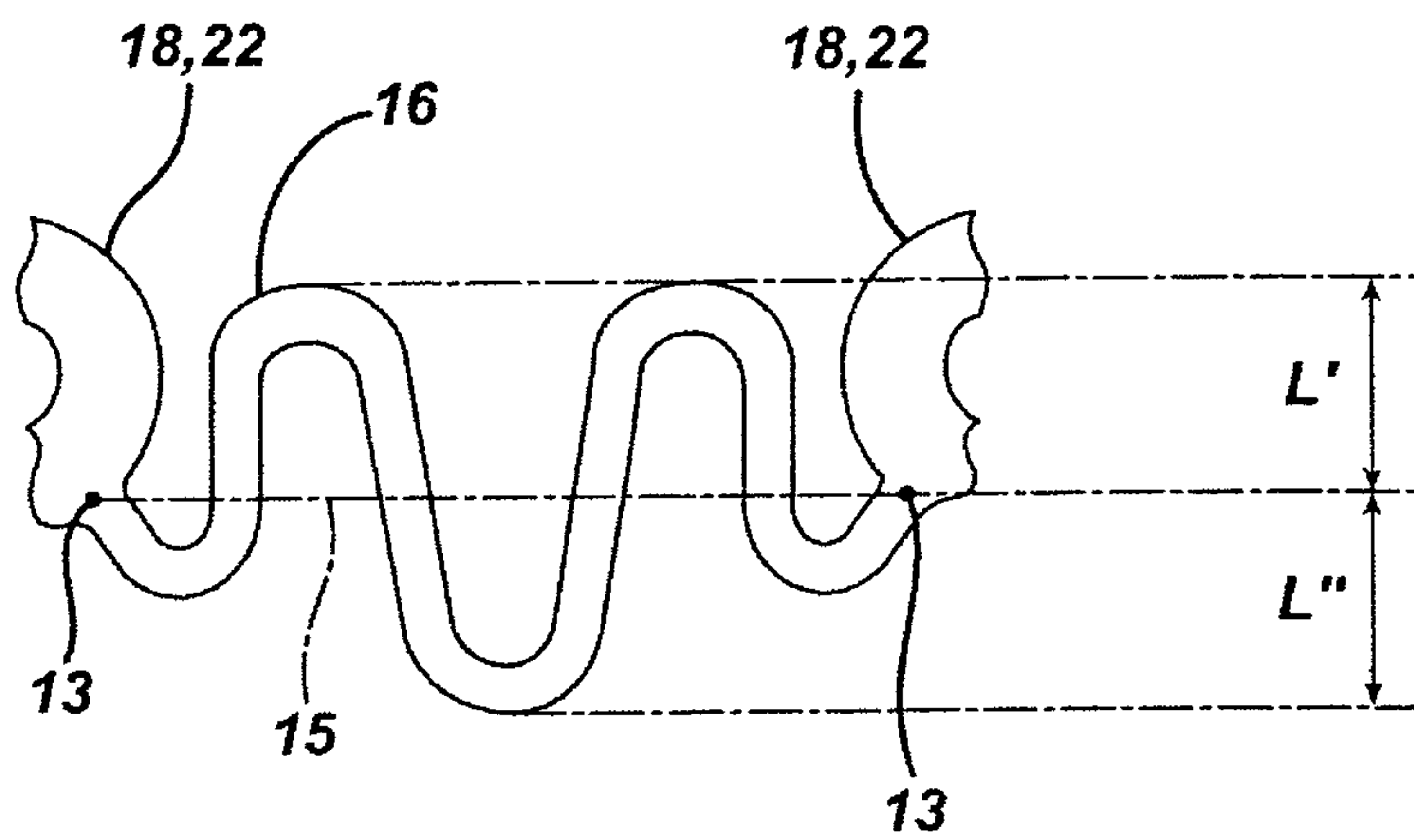


FIG. 5

