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(71) Applicant (for all designated States except US): THE ROYAL ALEXANDRA HOSPITAL FOR CHILDREN [AU/AU]; Cnr Hawkesbury Road & Hainsworth Street, Westmead, NSW 2145 (AU).

(72) Inventor: and

(75) Inventor/Applicant (for US only): LITTLE, David, Graham [AU/AU]; Cnr Hawkesbury Road & Hainsworth Street, Westmead, NSW 2145 (AU).

(74) Agent: F B RICE & CO; 605 Darling Street, Balmain, NSW 2041 (AU).

(54) Title: A DEVICE FOR THE DELIVERY OF A DRUG TO A FRACTURED BONE

(57) Abstract: The present invention relates to a bone fixation device, such as a screw or fixation pin or wire, said device including a body containing or being coated with a drug selected from bisphosphonates. The device may be used in a number of procedures to promote healing of a bone, including promoting union of a fracture, to provide internal fixation due to osteoporosis, and in the treatment of osteonecrosis or osteochondritis dissecans.
A Device for the Delivery of a Drug to a Fractured Bone

Field of the Invention

The present invention relates to a drug for use in bone healing. More specifically, the present invention relates to the use of bisphosphonates to increase the amount of healing bone around a fracture when delivered to the region of a fracture by a fixation device.

Background of the Invention

While most commonly used for the prevention of bone resorption, bisphosphonates have recently been noted to promote bone growth and fracture repair. Research has shown that intravenous bisphosphonates increase the rate of bone formation in distraction osteogenesis. Administration of oral bisphosphonates may also be effective.

While intravenous and oral administration of the drug may be effective, it may also be useful in certain circumstances to provide a local or regional dose of bisphosphonate at and around the fracture site rather than administering the drug systemically.

The present invention aims to provide a delivery system for bisphosphonates which enables a therapeutically effective amount of a bisphosphonate to be delivered to a desired regional site of bone.

Summary of the Invention

In a first aspect, the present invention consists in a bone fixation device, the device including a body containing or coated with a drug selected from the group consisting of at least one bisphosphonate.

Typically, the drug is free to move from within or on the body of the device to a position outside the body of the device.

The bone fixation device may be a screw or a number of screws with or without a plate member.

Typically, the bone fixation device is made from a material including, but not limited to, stainless steel, a titanium alloy or any other biocompatible metal. Alternatively, the fixation device maybe made from a bioreabsorbable material.
such as polyglycolide (PGA), poly-(L-lactide), poly (D,L-lactide) (PLA) or another biocompatible polymer. In the latter embodiment, the bisphosphonate may be admixed with the material of the fixation device and delivered to the bone as the fixation device is resorbed thereby releasing the bisphosphonate.

In another embodiment, a metal device is coated with a layer of biodegradable material containing a bisphosphonate.

Whether used separately or together with a plate, the screw or screws may contain a therapeutically effective amount of a bisphosphonate. The external surface of the screws may be coated with the at least one bisphosphonate or, alternatively, each screw may include a receptacle member such as a hollow interior, compartment or groove or grooves to house the at least one bisphosphonate.

Where the screw(s) include a receptacle member, the screw(s) may further include an opening or other such structure such as a series of apertures which extend from the receptacle member to an outer surface of the screw(s) such that the at least one bisphosphonate may be released from within the screw to the surrounding bone. In this regard, the opening may be sealed with a resorbable material that breaks down upon placement of the device in a patient's body.

In another embodiment where the device is coated with the at least one bisphosphonate, the entire body of the device may be sealed with a resorbable material that breaks down upon placement of the device in a patient's body.

In another embodiment, the plate of the fixation device contains the at least one bisphosphonate. The plate may also be coated with the bisphosphonate or include some other structure to hold the bisphosphonate prior to its release into the fractured bone.

In a further embodiment, the bone fixation device of the invention comprises an intramedullary device. An example of such a device includes, but is not limited to, an intramedullary nail.

The fixation device may further comprise an external fixation pin or wire, useful in fractures, osteotomies or arthrodeses managed with external fixation devices. The external fixation pin or wire may also promote bone formation in distraction osteogenesis by providing regional delivery of a bisphosphonate to the bone undergoing distraction.
In another embodiment, the fixation device includes threaded or smooth Kirshner wires wherein the bisphosphonate is deposited on the surface of the wire, or in grooves or hollows of the wires.

In a preferred embodiment of the invention, the at least one bisphosphonate is zoledronic acid {1-hydroxy-2-{[1H-imidazol-1-yl]ethylidene} bisphosphonic acid}.

Alternatively, the at least one bisphosphonate may include, but is not limited to, any one of the following:
pamidronate {3-amino-1-hydroxypropylidene bisphosphonic acid};
aleendronate {4-amino-1-hydroxybutylidene bisphosphonic acid};
etidronate {1-hydroxyethylidene bisphosphonic acid};
clodronate {dichloromethylene bisphosphonic acid};
risedronate [2-(3-pyridinyl)-1-hydroxyethylidene bisphosphonic acid];
tiludronate {chloro-4-phenylthiomethylidene bisphosphonic acid};
ibandronate {1-hydroxy-3(methylpentylamino)-propylidene bisphosphonic acid};
incadronate {cycloheptyl-amino-methylene bisphosphonic acid};
minodronate {[1-hydroxy-2-(imidazo[1,2-a]pyridin-3-yl)ethylidene]bisphosphonic acid};
olpadronate {(3-dimethylamino-1-hydroxypropylidene) bisphosphonic acid};
neridronate (6-amino-1-hydroxyhexylidene-1,1-bisphosphonic acid);
EB-1053 1-hydroxy-3-(1-pyrrolidinyl)-propylidene-1,1-bisphosphonic acid;
or any other therapeutically effective bisphosphonate or pharmaceutically acceptable salts or esters thereof.

Preferably, the at least one bisphosphonate is administered in the following doses:
zoledronate at 0.0001 to 0.5 mg/kg body weight per dose;
pamidronate at 0.0001 to 5.0 mg/kg body weight per dose;
ibandronate (ibandronic acid) at 0.0001 to 0.5 mg/kg body weight per dose;
risedronate at 0.0001 to 0.5 mg/kg body weight per dose;
alendronate at 0.0001 to 5.0 mg/kg body weight per dose;
clodronate at 0.0001 to 20 mg/kg body weight per dose;
etidronate at 0.0001 to 20 mg/kg body weight per dose;
tiludronate at 0.0001 to 5.0 mg/kg body weight per dose;
incadronate at 0.0001 to 5.0 mg/kg body weight per dose;
minodronate at 0.0001 to 0.5 mg/kg body weight per dose;
olpadronate at 0.0001 to 0.5 mg/kg body weight per dose;
neridronate at 0.01 to 5.0 mg/kg body weight per dose;
EB-153 at from 0.0001 to 5.0 mg/kg body weight per dose.

The at least one bisphosphonate may be admixed with other substances
or carriers such as gelatin, glycerol, collagen, hyaluronan-based sponges, pads,
pastes and gels, chitosan, fibrin, synthetic polymers, calcium sulphate,
tricalcium phosphate, hydroxyapatite or other ceramics, and or their
combinations to alter the rate at which the bisphosphonate dose is
administered to the fractured bone.

The bone fixation device of the present invention may be used to
promote union of a fractured bone.

The bone fixation device may also be used to promote the fusion of a
joint during an arthrodesis procedure, including spinal arthrodesis (spinal
fusion).

In another embodiment, the bone fixation device may be used to
promote the union of a bone following an osteotomy procedure.

The device may also be used to promote union in a fracture healing
slowly (delayed union) or where healing has appreciably ceased (non-union).

The bone fixation device may be used to treat a patient suffering from
osteonecrosis or osteochondritis dissecans. Furthermore, the device may be
used to treat a patient in need of internal fixation because of osteoporosis and
the resultant risk of fracture. Alternatively, the patient may be at risk of
pathological fracture.

In another embodiment, the bone fixation device may be used to treat a
patient having a pathological fracture secondary to malignant disease. In this
embodiment, the at least one bisphosphonate contained within or coated on the
device would have the advantage of stimulating fracture repair in addition to
preventing further resorption due to the malignant process.

The bone fixation device may further be used to treat a patient suffering
from Paget’s disease, fibrous dysplasia, osteofibrous dysplasia, congenital
pseudarthrosis, osteogenesis imperfecta or other conditions requiring fixation of
a fracture.
In another embodiment, the bone fixation device may be used prophylactically in a patient who is receiving or has received systemic corticosteroid or other therapy, is an active smoker, intakes large quantities of alcohol or any other substance known to cause osteoporosis and resultant fracture and to interfere with fracture union.

In the embodiment wherein the bone fixation device comprises a plate member and a plurality of screws, said device may be particularly useful in the promotion of union of fractures of the femur, tibia, fibula, calcaneus, metatarsals, humerus, radius, ulna, metacarpals, maxilla, mandible and cranium, but could be used in any bone where plate fixation is feasible.

When used alone, the screws may be used to treat a patient suffering from a fracture of a small bone such as the scaphoid, talus and femoral neck. This embodiment may also be useful for the treatment of fractures of the spine, fractures involving articular surfaces and fractures around the foot and ankle or hand and wrist. The bone fixation device of this embodiment may also be used in the treatment of osteochondritis dissecans.

In one embodiment, the bone fixation device may be used in the treatment of a patient requiring a spinal or other joint arthrodesis. In this embodiment, the bone fixation device may be used to secure the arthrodesis in such a manner as to potentiate the bony healing necessary to accomplish an effective arthrodesis.

In a further embodiment, the bone fixation device may be used in the treatment of a patient requiring an osteotomy. In this embodiment, the fixation device may be used to secure the osteotomy in such a manner as to potentiate bony healing of the osteotomy.

The bone fixation device of the present invention may be used in humans or alternately in the field of veterinary medicine.

In a second aspect, the invention consists in a method of promoting union of a fracture in a patient in need of such treatment using the device according to the first aspect of the invention, the method including the steps of:

(a) carrying out reduction of the fractured bone;
(b) positioning the fixation device such that it fixes the fractured bone in a desired position;
(c) causing or allowing the delivery of the at least one bisphosphonate from the fixation device to a region of the fractured bone.
The entire dose of bisphosphonate is preferably delivered to a region of bone within the proximity of a fracture site such that the bisphosphonate has the desired effect of promoting bone growth between, and ultimately union of, the fractured ends of the bone. Accordingly, the bone fixation device may be positioned substantially adjacent or across the fracture site or substantially spaced from the fracture site. Furthermore, the bone fixation device may be positioned external the fractured bone or internal the bone.

Preferably, the at least one bisphosphonate is delivered to a sufficient region of bone around the fracture site by diffusion of the drug such that total osteoblastic production of bone is increased in addition to preventing against osteoporosis in the region around the fracture which may result from disuse or from stress shielding by the bone fixation device.

The entire dose of bisphosphonate may be delivered early in the course of healing of the fracture, that is, in approximately the first two weeks following the surgical procedure. The entire dose may, however, be delivered in a more prolonged manner in the course of healing of the fracture for example over approximately the first two months. Alternatively, the entire dose may be delivered over approximately one or more years following the fracture.

In a third aspect, the invention consists in a method of promoting union of an arthrodesis, including a spinal arthrodesis in a patient in need of such treatment using the device according to the first aspect of the invention, the method including the steps of:

(a) preparing the bone site for arthrodesis by exposing bone surfaces;
(b) positioning the fixation device such that it fixes the bone or bones in a desired position;
(c) causing or allowing the delivery of the at least one bisphosphonate from the fixation device to a region of the arthrodesis.

In a fourth aspect, the invention consists in a method of promoting union of an osteotomy in a patient in need of such treatment using the device according to the first aspect of the invention, the method including the steps of:

(a) carrying out reduction of the osteotomised bone;
(b) positioning the fixation device such that it fixes the osteotomised bone in a desired position;
(c) causing or allowing the delivery of the at least one bisphosphonate from the fixation device to a region of the osteotomy.
In a fifth aspect, the invention consists in a method of promoting healing of osteochondritis dissecans in a patient in need of such treatment using the device according to the first aspect of the invention, the method including the steps of:

(a) carrying out reduction of the osteochondritic fragment;
(b) positioning the fixation device such that it fixes the osteochondritic fragment in a desired position;
(c) causing or allowing the delivery of the at least one bisphosphonate from the fixation device to a region of the osteochondritic fragment.

In a sixth aspect, the invention consists in a method of promoting healing of osteonecrosis in a patient in need of such treatment using the device according to the first aspect of the invention, the method including the steps of:

(a) drilling a hole in the region of the osteonecrosis;
(b) positioning the fixation device such that it supports the bone in the region;
(c) causing or allowing the delivery of the at least one bisphosphonate from the fixation device to a region of the osteonecrosis.

As bisphosphonates have a strong affinity for bone mineral, when the at least one bisphosphonate is delivered to a region of bone surrounding a fracture, arthrodesis site, osteotomy site, site of osteochondritis dissecans or site of osteonecrosis, the bisphosphonate should be taken up locally in that region of bone thus minimising the distribution of the bisphosphonate to other regions of the body (i.e. systemic distribution).

The fixation device of the present invention does not include an orthopaedic prosthesis such as a prosthetic joint or dental replacement device wherein the provision of bisphosphonates would enhance securement of the prosthetic device in the surrounding bone.

**Brief Description of the Drawings**

By way of example only, a preferred embodiment of the invention is described with reference to the accompanying figures:

Fig. 1 is a generic formula for one class of bisphosphonates.
Fig. 2 is a schematic view of one embodiment of the present invention.
Fig. 3 is a schematic view of a further embodiment of the present invention.
Fig. 4 is a schematic view of another embodiment of the present invention.

Preferred Embodiment of the Invention

An orthopaedic fixation device according to the present invention is generally depicted in the accompanying drawings as 10. The fixation device 10 includes a central canal 11 which receives a therapeutic dose of bisphosphonate. The side vent holes 12 allow the bisphosphonate to diffuse out of the fixation device 10 and be taken up by the bone in the region of the fixation device 10. The fixation device is preferably positioned sufficiently close to a fracture site to allow for diffusion of the bisphosphonate to said fracture site. Such local or regional administration and binding of the bisphosphonate leads to the prevention of the unwanted bisphosphonate dosing of other areas of the body.

The device depicted in Figure 3 is positioned within a fractured scaphoid bone 13. The fixation device 10 comprises a screw 20 which may be countersunk under the articular surface 14. The screw 20 provides compression of the fractured bone. The screw 20 further includes a groove 15 which is adapted to receive and hold a therapeutically effective dose of bisphosphonate. Alternatively, an external surface of screw 20 may be coated with the therapeutically effective amount of bisphosphonate. Scaphoid fractures are known to have a high incidence of delayed union and osteonecrosis. Both these conditions may be ameliorated by the local administration of a bisphosphonate.

As depicted in Figure 4, the fixation device 10 may include a plate 30 having a series of screws 20 dependent therefrom. The screws 20 are drilled into the bone such that both the plate 30 and the screws 20 fix the bone in place. As previously discussed, either the plate 30 or the screws 20 may be contain a therapeutic amount of bisphosphonate. Further, both the plate 30 and the screws 20 may contain a therapeutic amount of bisphosphonate.
Example 1

Animal Model

The present experiment uses 20 to 24-week-old NZW rabbit model. Animals were approximately 3 kg at the time of surgery. A 10mm segmental fracture is surgically created in the right tibiae of the rabbits. The bones are held with a 7 hole plate and screws.

Experimental Groups

16 animals were allocated into one of four groups (4 per group)
   i) Saline (Control)
   ii) 1mg pamidronate, delivered via plate (1mg plate)
   iii) 8mg pamidronate, delivered via plate (8mg plate)
   iv) 5mg pamidronate, delivered via cannulated screw (5mg screw)

Device Preparation

All plates were identical in design, ¼ tubular plates with 7 holes and 55 mm in length, made to fit 2.7mm screws, but also compatible with 3.5 mm screws. Plates were prepared by autoclave sterilisation. For 1mg plates, a solution of pamidronate 10 mg/ml was prepared and 0.1 ml (1mg) was distributed on the concave side of the plate. This is the side in direct contact with bone. For 8 mg plate, a solution of pamidronate 25 mg/ml was prepared and 0.3 ml (approx 8mg) was applied to the concave side of the plate. Plates were placed in sterile Petrie dishes and incubated for 12 hrs at 37° C until dry, prior to implantation.

Delivery screws were of a special design. A 3.5 mm cortical bone screw had been cannulated with a 1.1 mm hole and a 0.7 mm side vent hole (Fig 1). 5mg of pamidronate was placed in the sterile screw immediately prior to surgery.

Operative Technique

Surgery was performed in an animal theatre. After premedication with IM ketamine 15 mg/kg, xylazine 4mg/kg given in combination 10 minutes before
surgery, anaesthesia was maintained with inhaled halothane 2%, nitrous oxide 2 litres/min and oxygen 1L/min.

The operative field was prepared by shaving with clippers, disinfected with povidone iodine 4% w/v in 70% alcohol. The tibia was exposed sub-periosteally along its length and a spare plate used as a template to allow pre-drilling and tapping of fixation holes. Drill sites were placed in holes 1, 2, 4, 6 and 7 (proximal to distal) – holes 3 and 5 were not used as they were close to the planned osteotomies. Two osteotomies were created with a saw and a 10 mm piece of mid-diaphyseal tibia removed and replaced. The designated 7-hole ¾ tubular plate was then attached to the tibia, 2 screws in each large segment and one screw in the de-vascularised central segment. All screws were 2.7mm cortical screws apart from the screw in the second hole, which was 3.5mm. In the 5mg screw group, the cannulated screw contained disodium pamidronate powder.

Wounds were closed with dissolvable suture and Buprenorphine 0.05 mg/kg was given at the end of surgery and 12 hours post-operatively. Animals were allowed to freely weight bear.

**Exclusions**

One rabbit in the screw group was excluded as the central segment had been split at surgery and fixation did not hold.

**Analysis**

Animals were culled at six weeks post operation. Both tibiae were harvested and the plates and screws removed. The whole bones were fixed in 4% paraformaldehyde. The tibiae were analysed with a Stratec XCT Research SA pQCT scanner and analysis software (Stratec Medizintechnik GmbH, Pforzheim, Germany). Scout scans were centred on the central screw hole and scans taken proximally 20mm, 10mm 5 mm and 2.5 mm from the centre, through the centre and 2.5 mm, 5 mm, 10mm and 20mm distally on the right (operated) leg. On the left (non-operated) leg, scans were taken in corresponding areas at the centre, and 10mm proximal and distal.

The scanner generated data for BMD, BMC, cross sectional area and polar moment of inertia. The callus volume was integrated from mean cross
sectional area and scan length. With n=3-4 data points, quantification of parametric statistical differences is limited, and was not used. Chi square analysis was applied to compare numbers of scan regions showing differences of greater than 10%.

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Results

Total bone volume is shown in Graph 1. Bone volume on the operated side was increased by 26% in the 5mg screw group. There were only slight differences in the plate groups.

Cross-sectional area for all regions in all groups is presented in Graph 2. There were increases in cross sectional area in the screw group over controls on the operated side ranging from 13% to 38% (median 27%), but no increase on the non-operated leg. The median increases in the 1mg and 8mg plate groups were 5% and 5%, respectively. The differences were greater than 10% in 9 of 9 scan regions for the screw group, in contrast in only 2 of 9 regions was there an increase of >10% in area in the 1mg plate group, and in only 4 of 9 regions was this the case in the 8mg plate group. This was significant by chi square p<0.01.

Bone mineral content was likewise increased in the screw group (Graph 3). For clarity, remaining graphs show only the control and screw groups. The increases in the different scan regions ranged from 9% to 22% (median 16%). There was no increase in the non-operated leg. The median increases in the 1mg and 8mg plate groups were 6% and 6%, respectively. The differences were greater than 10% in 8 of 9 scan regions in the screw group, in contrast, in only 2 of 9 regions was there an increase of >10% in BMC in the 1mg plate group, and in only 3 of 9 regions was this the case in the 8mg plate group. This was significant by chi square p<0.025.

Polar moment of inertia was increased in the 5mg screw group by 49% to 87% (median 71%) over controls. There was no increase in the non-operated leg. In contrast the 1mg plate and 5 mg plate showed little or no improvement in polar moment (median 0% and –8%, respectively). All 9 areas showed increases of over 40% in the 5mg screw group, whereas only 1 of 9 and 0 of 9 showed this level of increase in the 1mg and 8mg plate groups (P<0.01 chi square).
There was no notable difference in volumetric BMD between any of the operated groups.

**Discussion**

This experiment looks at the effect of regional bisphosphonate dosing via a delivery device for fracture care. The inventor has previously documented that systemic bisphosphonate delivery can increase the amount, mineral content and strength of callus during bone healing. In a systemic pamidronate study, increases in BMC were noted on the operative side, but also increases in BMC of 10%-12% were seen on the non-operative side.

The advantages of regional delivery are numerous, including: increased amount, mineral content and strength of callus; reduced bone exposure away from the target region; and reduced renal exposure and therefore reduced risk of nephrotoxicity.

The results of the experiment show that bisphosphonates may be successfully delivered to the region of a fracture. Dosing on the plate itself did not lead to a notable improvement in the parameters measured. In this regard, it is possible that the pamidronate was “trapped” between the plate and the bone and was only delivered to the area immediately beneath the plate. It is notable that one of the plate groups received a higher dose than the screw group, with lesser effect on outcome.

In contrast the dosing screw allowed the pamidronate to diffuse into the region of the fracture haematoma, and its effects can be seen to have occurred at both the proximal and distal osteotomy sites (Graphs 3-5). This indicates that the dosing was regional and not directly local. There was no change in BMC on the non-operated side in the 5mg screw group. This indicates that only very small amounts of bisphosphonate are likely to have travelled systemically to the other limb. We have noted increases of around 10% in BMC on the non-operated side in both a previous systemic pamidronate and zoledronic acid experiment.

The increases in moment of inertia in the screw group were large (49% to 87%) which is likely to correlate with increases in strength.

This technique allows the regional treatment of fractures where there is concern about union or stress-shielding osteopenia while minimising systemic exposure.
Conclusions

This experiment indicates that regional delivery of pamidronate via an internal fixation device, a cannulated screw in this case, can result in increases in area, mineralisation and moment of inertia in a double osteotomy model in NZW rabbits. No effect was seen in the contralateral limb, in contrast to previous systemic dosing experiments. Delivery of pamidronate via the plates was ineffective; it is possible the drug was sequestered under the plate.

It will be appreciated by persons skilled in the art that numerous variations and/or modifications may be made to the invention as shown in the specific embodiments without departing from the spirit or scope of the invention as broadly described. The present embodiments are, therefore, to be considered in all respects as illustrative and not restrictive.
Graph 1 Bone Volume

Bone Volume

- Control
- 1mg Plate
- 8mg Plate
- 5mg Screw

mm$^3$

- 0
- 500
- 1000
- 1500
- 2000
- 2500
- 3000
- 3500
- 4000
- 4500

3% 8% 26%
Graph 2 Cross-Sectional Area (Screw denoted position of dosing screw, arrows denote sites of osteotomies)
Graph 3 BMC in Control and 5mg screw dosing groups (Screw denoted position of dosing screw, arrows denote sites of osteotomies)
Graph 4 Polar moment of inertia in Control and 5mg screw dosing groups (Screw denoted position of dosing screw, arrows denote sites of osteotomies)
CLAIMS:
1. A bone fixation device, the device including a body containing or coated with a drug selected from the group consisting of at least one bisphosphonate.
2. The bone fixation device of claim 1 wherein said device comprises at least one screw.
3. The bone fixation device of claim 2 wherein the at least one screw includes a receptacle member to receive the at least one bisphosphonate.
4. The bone fixation device of claim 3 wherein the receptacle member comprises a hollow interior, compartment or groove and wherein the device further includes an aperture or a series of apertures extending from the hollow interior, compartment or groove to an external surface of the screw.
5. The bone fixation device of claim 4 wherein the aperture or series of apertures are sealed with a material that is resorbable within a body of a subject.
6. The bone fixation device of claim 2 wherein the external surface of the at least one screw is coated with the at least one bisphosphonate.
7. The bone fixation device of any one of claims 2 to 6 further including a plate member having an aperture or a series of apertures therein to receive the at least one screw.
8. The bone fixation device of claim 7 wherein the plate member is coated with the at least one bisphosphonate.
9. The bone fixation device of any one of the preceding claims wherein said device is made from a material selected from the group comprising stainless steel, a titanium alloy or any other biocompatible metal.
10. The bone fixation device of any one of claims 1 to 8 wherein said device is made from a biodegradable material including polyglycolide (PGA), poly-(L-lactide), poly-(D,L-lactide) (PLA).
11. The bone fixation device of claim 10 wherein the at least one bisphosphonate is admixed with the material of the bone fixation device.
12. The bone fixation device of claim 1 wherein said device comprises an intramedullary device including an intramedullary nail.
13. The bone fixation device of claim 1 wherein said device comprises an external fixation pin or wire.
14. The bone fixation device of claim 1, said device comprising threaded or smooth Kirshner wires wherein the at least bisphosphonate is deposited on the surface of the wire, or in grooves or hollows of the wires.
15. The bone fixation device of any one of the preceding claims wherein the at least one bisphosphonate is zoledronic acid {1-hydroxy-2-[(1H-imidazol-1-yl)ethylidene] bisphosphonic acid}.

16. The bone fixation device of any one of claims 1 to 14 wherein the at least one bisphosphonate is selected from the group consisting of pamidronate {3-amino-1-hydroxypropylylidene bisphosphonic acid}; alendronate {4-amino-1-hydroxybutylylidene bisphosphonic acid}; etidronate {1-hydroxyethylylidene bisphosphonic acid}; clodronate {dichloromethylene bisphosphonic acid}; risedronate {2-(3-pyridinyl)-1-hydroxyethylylidene bisphosphonic acid}; tiludronate {chloro-4-phenylthiomylylidene bisphosphonic acid}; ibandronate {1-hydroxy-3(methylpentylamino)-propylylidene bisphosphonic acid}; incadronate {cycloheptyl-amino-methylene bisphosphonic acid}; minodronate {[1-hydroxy-2-(imidazo[1,2-a]pyridin-3-yl)ethylidene]bisphosphonic acid}; olpadronate {(3-dimethylamino-1-hydroxypropylylidene) bisphosphonic acid}; neridronate (6-amino-1-hydroxyethylidene-1,1-bisphosphonic acid); EB-1053 1-hydroxy-3-(1-pyrrolidinyl)-propylylidene-1,1-bisphosphonic acid; or any other therapeutically effective bisphosphonate or pharmaceutically acceptable salts or esters thereof.

17. The bone fixation device of any one of the preceding claims wherein the at least one bisphosphonate is admixed with a carrier medium including gelatin, glycerol, collagen, hyaluronan-based sponges, pads, pastes and gels, chitosan, fibrin, bioresorbable polymers, silicon, calcium sulphate, tricalcium phosphate, hydroxyapatite or other ceramics, or any drug delivery system.

18. The bone fixation device of any one of the preceding claims when used to promote union of a fractured bone including the promotion of union in delayed or non-union fractures.

19. The bone fixation device of any one of claims 1 to 17 when used to promote the fusion of a joint during an arthrodesis procedure, including spinal arthrodesis (spinal fusion).

20. The bone fixation device of any one of claims 1 to 17 when used to promote the union of a bone following an osteotomy procedure.

21. The bone fixation device of any one of claims 1 to 17 when used in the treatment of osteonecrosis or osteochondritis dissecans.

22. The bone fixation device of any one of claims 1 to 17 when used to treat a patient in need of internal fixation due to osteoporosis and the associated risk or fracture.
23. The bone fixation device of any one of claims 1 to 17 when used to treat a patient having a pathological fracture secondary to malignant disease.
24. A method of promoting union of a fracture in a patient in need of such treatment using the bone device of claim 1, the method including the steps of:
   (a) carrying out reduction of the fractured bone;
   (b) positioning the bone fixation device such that it fixes the fractured bone in a desired position;
   (c) causing or allowing the delivery of a dose of the at least one bisphosphonate from the bone fixation device to a region of the fractured bone.
25. The method of claim 24 wherein the bone fixation device is positioned substantially adjacent or substantially spaced from the fracture site.
26. The method of claim 24 or claim 25 wherein the entire dose of the at least one bisphosphonate is delivered to a region of bone within the proximity of the fracture site within the first two weeks following the surgical procedure.
27. The method claim 24 or claim 25 wherein the entire dose is delivered to a region of bone within the proximity of the fracture site over a period of two months following the surgical procedure.
28. The method of claim 24 or claim 25 wherein the entire dose is delivered to a region of bone within the proximity of the fracture site over a period of one or more years following the surgical procedure.
29. A method of promoting union of an arthrodesis, including a spinal arthrodesis in a patient in need of such treatment using the bone fixation device of claim 1, the method including the steps of:
   (a) preparing the bone site for arthrodesis by exposing bone surfaces;
   (b) positioning the bone fixation device such that it fixes the bone or bones in a desired position;
   (c) causing or allowing the delivery of the at least one bisphosphonate from the bone fixation device to a region of the arthrodesis.
30. A method of promoting union of an osteotomy in a patient in need of such treatment using the bone fixation device of claim 1, the method including the steps of:
   (a) carrying out reduction of the osteotomised bone;
   (b) positioning the bone fixation device such that it fixes the osteotomised bone in a desired position;
   (c) causing or allowing the delivery of the at least one bisphosphonate from the bone fixation device to a region of the osteotomy.
31. A method of promoting healing of osteochondritis dissecans in a patient in need of such treatment using the bone fixation device of claim 1, the method including the steps of:
   (a) carrying out reduction of the osteochondritic fragment;
   (b) positioning the bone fixation device such that it fixes the osteochondritic fragment in a desired position;
   (c) causing or allowing the delivery of the at least one bisphosphonate from the bone fixation device to a region of the osteochondritic fragment.
32. A method of promoting healing of osteonecrosis in a patient in need of such treatment using the bone fixation device of claim 1, the method including the steps of:
   (a) drilling a hole in the region of the osteonecrosis;
   (b) positioning the fixation device such that it supports the bone in the region;
   (c) causing or allowing the delivery of the at least one bisphosphonate from the bone fixation device to a region of the osteonecrosis.
Fig. 1

[Chemical structure diagram]

O=\cancel{\text{P}}-\text{C}--\cancel{\text{P}}=\cancel{\text{O}}

\begin{align*}
\text{OH} & \quad \text{R}_1 & \quad \text{OH} \\
\text{OH} & \quad \text{R}_2 & \quad \text{OH}
\end{align*}
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

Int. Cl.?: A61B 17/86; A61K 031/663; A61P 19/00

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

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC: A61K, SEARCH TERMS AS BELOW

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

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

WPAT, CAPLUS, MEDLINE: (bone OR osteotomy OR arthrodex) AND (screw OR plate OR device OR prosthesis OR implant) AND (bisphosphonate OR diphosphonate OR biphosphonate OR zoledron: OR pamidron: OR alendron: OR cladron: OR risedron:)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Date of the actual completion of the international search 4 June 2002  
Date of mailing of the international search report 20 JUN 2002

Name and mailing address of the ISA/AU
AUSTRALIAN PATENT OFFICE
PO BOX 200, WODEN ACT 2600, AUSTRALIA
E-mail address: pct@ipaustalia.gov.au
Facsimile No. (02) 6285 3929

Authorized officer
MICHAEL GRIEVE
Telephone No: (02) 6283 2267

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