Abstract:
The invention relates to medicaments and novel methods of treating metabolic bone diseases, and more particularly for treating or preventing Critical Illness Related Metabolic Bone Disease or of critical illness induced Osteopenia [ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D00185] secondary to ICU Admission by sufficient autophagy inducing compound to inhibit or suppressing critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis or increased osteoclast differentiation. The disclosed therapeutic methods include administering of a autophagy activating compound to a mammal to: (1) treat or prevent a bone degenerative disorder; (2) slow bone deterioration; (3) restore lost bone; (4) maintain bone mass and/or bone quality or (5) inhibit bone resorption in particular by inhibiting or reducing a process by which osteoclasts break down bone and release the minerals resulting in a transfer of calcium from bone fluid to the blood. The invention also provides methods for administering the autophagy activating compound to treat a bone disorder of hyperresorption of bone and/or enhanced activation of osteoclasts.
OSTEOCLAST ACTIVITY

Background and Summary

BACKGROUND OF THE INVENTION

The invention relates to medicaments and novel methods of treating metabolic bone diseases, and more particularly for treating or preventing Critical Illness Related Metabolic Bone Disease or of critical illness induced Osteopenia [ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851] secondary to ICU Admission by sufficient autophagy inducing compound to inhibit or suppressing critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis or increased osteoclast differentiation. The disclosed therapeutic methods include administering of a autophagy activating compound to a mammal to: (1) treat or prevent a bone degenerative disorder; (2) slow bone deterioration; (3) restore lost bone; (4) maintain bone mass and/or bone quality or (5) inhibit bone resorption in particularly by inhibiting or reducing a process by which osteoclasts break down bone and release the minerals resulting in a transfer of calcium from bone fluid to the blood. The invention also provides methods for administering the autophagy activating compound to treat a bone disorder of hyperresorption of bone and/or enhanced activation of osteoclasts.

Several documents are cited throughout the text of this specification. Each of the documents herein (including any manufacturer's specifications, instructions etc.) are hereby incorporated by reference; however, there is no admission that any document cited is indeed prior art of the present invention.

B. Description of the Related Art

A number of conditions are associated with a loss of bone, particularly in the elderly and/or postmenopausal women. For example, osteoporosis is a debilitating disease characterized by a decrease in skeletal bone mass and mineral density, structural deterioration of the bone, and corresponding increases in bone fragility and susceptibility to fracture. Osteoporosis in humans is preceded by clinical osteopenia, a condition found in approximately 25 million people in the United States only.
Throughout adult life, bone continually undergoes a turnover through the coupled processes of bone formation and resorption. Bone resorption is mediated by bone resorbing cells, osteoclasts, which are formed by mononuclear phagocytic cells. New bone replacing the lost bone is deposited by bone-forming cells, osteoblasts, which are formed by mesenchymal stromal cells. Various other cell types that participate in the remodeling process are tightly controlled by systemic factors (e.g., hormones, lymphokines, growth factors, and vitamins) and local factors (e.g., cytokines, adhesion molecules, lymphokines, and growth factors). The proper spatiotemporal coordination of the bone remodeling process is essential to the maintenance of bone mass and integrity. A number of bone degenerative disorders are linked to an imbalance in the bone remodeling cycle which results in abnormal loss of bone mass (osteopenia) including metabolic bone diseases, such as osteoporosis, osteoplasia (osteomalacia) and osteodystrophy.

Critically ill patients have an increased osteoclast formation from circulating precursors in the blood, osteoclast maturation and osteoclast bone resorption activity in comparison to healthy patients.

There are currently two main types of pharmaceutical therapy available for the treatment of osteoporosis. The first, and most common, approach is the use of hormone therapy to reduce the resorption of bone tissue. Estrogen replacement therapy ("ERT") is known to prevent further deterioration and thus reduce the likelihood of fractures. However, the use of estrogen as a treatment is limited, as it is believed that long-term estrogen therapy may be associated with risk of uterine cancer, endometrial cancer, breast cancer, frequent vaginal bleeding, and thrombosis. Because of these serious side effects, many women choose to avoid this treatment. Further, few men agree to this type of therapy. The second major therapeutic approach to osteoporosis is the use of bisphosphonates, particularly alendronate, risedronate, and ibandronate. Although tests have shown that these compounds consistently increase the bone mineral density in osteoporosis patients, there are also significant problems with the treatment of osteoporosis by bisphosphonates, including irritation of the oesophagus and upper gastrointestinal tract.

Therefore, there exists a need to develop new therapeutic methods for treating and preventing bone disorders.
SUMMARY OF THE INVENTION

The present invention solves the problems of the related art by metabolic bone diseases.

The invention relates to medicaments and novel methods for treating or preventing bone degenerative disorders. The disorders treated or prevented include, for example, osteopenia, osteomalacia, osteoporosis, osteomyeloma, osteodystrophy, osteogenesis imperfecta, and bone degenerative disorders associated with chronic renal disease, hyperparathyroidism, long-term use of corticosteroids and critical illness.

In accordance with the purpose of the invention, as embodied and broadly described herein, the invention is broadly drawn to reverse decrease in bone formation markers (serum osteocalcin), and an increase in bone resorption markers (urinarypyridinoline) in a subject for instance a critically ill subject by spermidine or analogues thereof.

In one aspect of the invention, to treat by spermidine or analogues thereof a disorder of bone metabolism in a subject for instance to reverse a disorder of decrease in bone formation and an increase in bone resorption. Such disorder can be caused by critical illness.

Another aspect of present invention concerns to treat by an autophagy activator or enhancer a disorder of bone metabolism in a subject for instance to reverse a disorder of decrease in bone formation and an increase in bone resorption. Such disorder can be caused by critical illness. Particular aspects and embodiments of present invention are the following. An autophagy activator or autophagy inducing compound for use as a medicine to inhibit increased bone resorption. An autophagy activator or autophagy inducing compound for use as a medicine to inhibit osteoclast activity. An autophagy activator or autophagy inducing compound for use as a medicine to inhibit the formation of osteoclast (TRAP positive multinuclear cells) from blood progenitor cells. An autophagy activator or autophagy inducing compound for use as a medicine to activate bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is a bone metabolism disorder. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is caused by acute critical illness or prolonged critical illness. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is caused by (parenteral) nutrition
induced suppression of autophagy. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is caused by mTOR activation. And/or an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is hyperresorption of bone.

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An embodiment of present invention concerns an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is pathologically increased bone resorption and pathologically decreased bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is a disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness. Another embodiment of present invention concerns an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is pathologically increased bone resorption and pathologically decreased bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder caused by enhanced osteoclast formation from circular precursors in the blood during critical illness. Yet an embodiment of present invention concerns an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is pathologically increased bone resorption and pathologically decreased bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is increased osteoclast formation from circulating precursor leading to osteoporosis.

The present invention is predicated on the discovery by the inventors that the disorder of critical illness [MeSH Descriptor C23.550.291.625] correlates with increased osteoclast formation from circulating precursors in the blood, osteoclast maturation and osteoclast bone resorption activity in comparison to healthy patients, resulting in complications such as "Osteopenia" (ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851) or leading to the indication Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease, Osteoporosis Secondary to ICU Admission with more extreme bone loss which increases the risk of critically ill patients for fractures such as stress fractures (ICD-10-CM 733.93 Stress fracture of tibia or fibula, 733.94 Stress fracture of the metatarsals, 733.95 Stress fracture of other bone, 733.96 Stress fracture of femoral neck, 733.97 Stress fracture of shaft of femur and/or 733.98 Stress fracture of pelvis) and that usually needs that usually needs post-ICU ambulatory care. The applicants
found that trabecular bone mineral content and density were reduced in critically ill rabbits, coinciding with low serum levels of ionized calcium and osteocalcin. Moreover it was found that increased number of PBMC osteoclast precursors in the blood of critically ill humans resulted in increased in vitro osteoclast formation, which was further potentiated by the addition of serum from critically ill patients. Unexpectedly, neutralizing the inflammatory cytokines TNF-a and IL-6 in patient serum further increased osteoclast formation in patient PBMC cultures.

Further scope of applicability of the present invention will become apparent from the detailed description given hereinafter. However, it should be understood that the detailed description and specific examples, while indicating preferred embodiments of the invention, are given by way of illustration only, since various changes and modifications within the spirit and scope of the invention will become apparent to those skilled in the art from this detailed description. It is to be understood that both the foregoing general description and the following detailed description are exemplary and explanatory only and are not restrictive of the invention, as claimed. Some embodiments of the invention are set forth in claim format directly below:

**Detailed Description**

**DETAILED DESCRIPTION OF EMBODIMENTS OF THE INVENTION**

The following detailed description of the invention refers to the accompanying drawings. The same reference numbers in different drawings identify the same or similar elements. Also, the following detailed description does not limit the invention. Instead, the scope of the invention is defined by the appended claims and equivalents thereof.

The following detailed description of the invention refers to the accompanying drawings. The same reference numbers in different drawings identify the same or similar elements. Also, the following detailed description does not limit the invention. Instead, the scope of the invention is defined by the appended claims and equivalents thereof.

Several documents are cited throughout the text of this specification. Each of the documents herein (including any manufacturer's specifications, instructions etc.) are hereby incorporated by reference; however, there is no admission that any document cited is indeed prior art of the present invention.

The present invention will be described with respect to particular embodiments and with reference to certain drawings but the invention is not limited thereto but only by the claims. The drawings described are only schematic and are non-limiting. In the drawings, the size of
some of the elements may be exaggerated and not drawn to scale for illustrative purposes. The dimensions and the relative dimensions do not correspond to actual reductions to practice of the invention.

Furthermore, the terms first, second, third and the like in the description and in the claims, are used for distinguishing between similar elements and not necessarily for describing a sequential or chronological order. It is to be understood that the terms so used are interchangeable under appropriate circumstances and that the embodiments of the invention described herein are capable of operation in other sequences than described or illustrated herein.

Moreover, the terms top, bottom, over, under and the like in the description and the claims are used for descriptive purposes and not necessarily for describing relative positions. It is to be understood that the terms so used are interchangeable under appropriate circumstances and that the embodiments of the invention described herein are capable of operation in other orientations than described or illustrated herein.

It is to be noticed that the term "comprising", used in the claims, should not be interpreted as being restricted to the means listed thereafter; it does not exclude other elements or steps. It is thus to be interpreted as specifying the presence of the stated features, integers, steps or components as referred to, but does not preclude the presence or addition of one or more other features, integers, steps or components, or groups thereof. Thus, the scope of the expression "a composition comprising means A and B" should not be limited to the compositions consisting only of components A and B. It means that with respect to the present invention, the only relevant components of the composition are A and B.

Reference throughout this specification to "one embodiment" or "an embodiment" means that a particular feature, structure or characteristic described in connection with the embodiment is included in at least one embodiment of the present invention. Thus, appearances of the phrases "in one embodiment" or "in an embodiment" in various places throughout this specification are not necessarily all referring to the same embodiment, but may. Furthermore, the particular features, structures or characteristics may be combined in any suitable manner, as would be apparent to one of ordinary skill in the art from this disclosure, in one or more embodiments.

Similarly it should be appreciated that in the description of exemplary embodiments of the invention, various features of the invention are sometimes grouped together in a single embodiment, figure, or description thereof for the purpose of streamlining the disclosure and aiding the understanding of one or more of the various inventive aspects. This method of
disclosure, however, is not to be interpreted as reflecting an intention that the claimed invention requires more features than are expressly recited in each claim. Rather, as the following claims reflect, inventive aspects lie in less than all features of a single foregoing disclosed embodiment. Thus, the claims following the detailed description are hereby expressly incorporated into this detailed description, with each claim standing on its own as a separate embodiment of this invention.

Furthermore, while some embodiments described herein include some but not other features included in other embodiments, combinations of features of different embodiments are meant to be within the scope of the invention, and form different embodiments, as would be understood by those in the art. For example, in the following claims, any of the claimed embodiments can be used in any combination.

In the description provided herein, numerous specific details are set forth. However, it is understood that embodiments of the invention may be practiced without these specific details. In other instances, well-known methods, structures and techniques have not been shown in detail in order not to obscure an understanding of this description.

Other embodiments of the invention will be apparent to those skilled in the art from consideration of the specification and practice of the invention disclosed herein.

It is intended that the specification and examples be considered as exemplary only.

Each and every claim is incorporated into the specification as an embodiment of the present invention. Thus, the claims are part of the description and are a further description and are in addition to the preferred embodiments of the present invention.

Each of the claims set out a particular embodiment of the invention.

The following terms are provided solely to aid in the understanding of the invention.

An osteoclast is a type of bone cell that removes bone tissue by removing its mineralized matrix and breaking up the organic bone. For instance organic dry weight is about 90% collagen. This process is known as bone resorption. Osteoclasts and osteoblasts are instrumental in controlling the amount of bone tissue: osteoblasts form bone, osteoclasts resorb bone. Osteoclasts are formed by the fusion of cells of the monocyte-macrophage cell line and osteoclasts can be characterized by high expression of tartrate resistant acid phosphatase (TRAP) and cathepsin K.
"Microtubule associated protein light chain 3" (LC3) is an ubiquitin-like protein that binds to autophagosomes (AVs). Cellular biologists transfec mammalian cells with GFP tagged LC3 to track and follow the fate of AVs in the cell and to measure autophagic flux.

"Bone Remodeling" (or bone metabolism) is a life-long process where mature bone tissue is removed from the skeleton (a process called bone resorption) and new bone tissue is formed (a process called ossification or new bone formation). These processes also control the reshaping or replacement of bone during growth and following injuries like fractures but also micro-damage, which occurs during normal activity. Remodeling responds also to functional demands of the mechanical loading. As a result, bone is added where needed and removed where it is not required. It is a tightly regulated process involving the interaction of osteoblasts and osteoclasts.

The term "pharmaceutically acceptable" is used adjectivally herein to mean that the compounds are appropriate for use in a pharmaceutical product. The term "physiologically acceptable" also means that the compounds are appropriate for use in a pharmaceutical product.

As used herein, an "autophagy activator" is any compound that increases autophagy within a cell. An increase in autophagy may be determined as known in the art and described herein. Exemplary, non-limiting autophagy activators are known in the art and include, for example, proteasome inhibitor, tamoxifen, trehalose, vinblastine, rapamycin, Azithromycin macrolide or its analogues, that inhibit the mammalian target of rapamycin (mTOR) (a negative regulator of autophagy), ganima-benzene hexachloride, or of a derivative thereof which is obtainable by chemical substitution. Yet, azithromycin has retained said capacity of acting as an inducer or stimulator of autophagy maturation.

As used herein, the phrase "physiologically acceptable salts" or "pharmaceutically acceptable salts" or "nutraceutically acceptable salts" refers to salts prepared from pharmaceutically acceptable, preferably nontoxic, acids and bases, including inorganic and organic acids and bases, including but not limited to, sulfuric, citric, maleic, acetic, oxalic, hydrochloride, hydro bromide, hydro iodide, nitrate, sulfate, bisulfite, phosphate, acid phosphate, isonicitinate, acetate, lactate, salicylate, citrate, acid citrate, tartrate, oleate, tannate, pantothenate, bitartrate, ascorbate, succinate, maleate, gentisinate, fumarate, gluconate, glucaronate, saccharate,
fornate, benzoate, glutamate, methanesulfonate, ethanesulfonate, benzenesulfonate, p-toluensulfonate and pamoate (i.e. 1, r-methylene-bis-(2-hydroxy-3-naphthoate)) salts. Pharmaceutically acceptable salts include those formed with free amino groups such as, but not limited to, those derived from hydrochloric, phosphoric, acetic, oxalic, and tartaric acids. Pharmaceutically acceptable salts also include those formed with free carboxyl groups such as, but not limited to, those derived from sodium, potassium, ammonium, sodium lithium, calcium, magnesium, ferric hydroxides, isopropylamine, triethylamine, 2-ethylamino ethanol, histidine, and procaine.

As used herein, the term "carrier" refers to a diluent, adjuvant, excipient, or vehicle. Such carriers can be sterile liquids, such as saline solutions in water, or oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like. A saline solution is a preferred carrier when the pharmaceutical composition is administered intravenously. Saline solutions and aqueous dextrose and glycerol solutions can also be employed as liquid carriers, particularly for injectable solutions.

As used herein, the term "mineral" refers to a substance, preferably a natural substance that contains calcium, magnesium or phosphorus. Illustrative nutrients and minerals include beef bone, fish bone, calcium phosphate, egg shells, sea shells, oyster shells, calcium carbonate, calcium chloride, calcium lactate, calcium gluconate and calcium citrate.

The term "treatment" refers to any process, action, application, therapy, or the like, wherein a mammal, including a human being, is subject to medical aid with the object of improving the mammal's condition, directly or indirectly.

In its broadest sense, the term a "critically ill patient" (herein designated CIP) refers to a patient who is experiencing an acute life-threatening episode or who is diagnosed to be in imminent danger of such an episode. A critically ill patient is medically unstable, and when not treated, likely to die.

The term critically ill patient refers to a patient who has sustained or is at risk of sustaining acutely life-threatening single or multiple organ system failure due to disease or injury, a patient who is being operated and where complications supervene, and a patient who has been operated in a vital organ within the last week or has been subject to major surgery within the last week.

In a more restricted sense, the term a "critically ill patient", as used herein refers to a patient who has sustained or are at risk of sustaining acutely life-threatening single or multiple organ system failure due to disease or injury, or a patient who is being operated and where complications supervene.
In an even more restricted sense, the term a "critically ill patient", as used herein refers to a patient who has sustained or is at risk of sustaining acutely life-threatening single or multiple organ system failure due to disease or injury. Similarly, these definitions apply to similar expressions such as "critical illness in a patient" and a "patient is critically ill". A critically ill patient is also a patient in need of cardiac surgery, cerebral surgery, thoracic surgery, abdominal surgery, vascular surgery, or transplantation, or a patient suffering from neurological diseases, cerebral trauma, respiratory insufficiency, abdominal peritonitis, multiple trauma, severe burns, or critical illness polyneuropathy.

The term "critical illness" as used herein refers to the condition of a "critically ill patient". ICU scoring systems such as APACHE II & III, -Glasgow Coma Scale -PIM2 -SAPS II -SAPS III -SOFA define the measure the severity of critical illness disorder for adult patients admitted to intensive care units and predict hospital mortality risk for critically ill hospitalized adults.

The term "Intensive Care Unit" (herein designated ICU), as used herein refers to the part of a hospital where critically ill patients are treated. Of course, this might vary from country to country and even from hospital to hospital and the part of the hospital may not necessary, officially, bear the name "Intensive Care Unit" or a translation or derivation thereof. Of course, the term "Intensive Care Unit" also covers a nursing home, a clinic, for example, a private clinic, or the like if the same or similar activities are performed there. The term "ICU patient" refers to a "critically ill patient".

The term "parenteral administering" refers to delivery of substances given by routes other than the digestive tract, and covers administration routes such as intravenous, intra-arterial, intramuscular, intracerebroventricular, intraosseous intradermal, intrathecal, and intraperitoneal administration and intravesical infusion and intracavernosal injection.

Typically "parenteral administration" refers to intravenous administration. A particular form of parenteral administration refers to the delivery by intravenous administration of nutrition ("parenteral nutrition"). Parenteral nutrition is called "total parenteral nutrition" when no food is given by other routes.

"Parenteral nutrition" is a isotonic or hypertonic aqueous solution (or solid compositions to be dissolved, or liquid concentrates to be diluted to obtain an isotonic or hypertonic solution) comprising a saccharide such as glucose and further comprising one or more of lipids, amino acids.
"Peripheral blood mononuclear cells" (PBMCs) constitute a very important part of our peripheral immune system. The PBMCs consist mainly of monocytes, T-cells and B-cells, and smaller amounts of NK cells and dendritic cells of both myeloid and plasmacytoid origin. Bone-resorbing osteoclasts are formed from hemopoietic cells of the monocyte-macrophage lineage and the development of osteoclasts is called osteoclastogenesis.

A "critically ill" patient is a patient receiving intensive-care where under mechanical ventilation and intensive-care nutrition support under the form of parenteral and/or enteral nutrition and is being treated in an Intensive Care Unit ("ICU"), e.g., of a hospital. Often the intensive-care nutrition support concerns parenteral nutrition (PN) which is feeding an individual intravenously, bypassing the usual process of eating and digestion. The person thereby receives nutritional formulas that contain nutrients such as glucose, amino acids, lipids and added vitamins and dietary minerals. It is called total parenteral nutrition (TPN) or total nutrient admixture (TNA) when no food is given by other routes. This unnatural way of feeding the body is far from perfect and comes with several significant complications. Administration of large volumes of blood products, especially packed red cells, undergoing dialysis, especially continuous veno-venous hemofiltration, receiving multiple antibiotics, having a pulmonary artery catheter or an arterial blood pressure catheter inserted can be further part of the intensive care. These further criteria for critically ill patients are exemplary only, and one skilled in the art will understand that other indicia of a patient in a critically ill state are possible and are considered to be encompassed by the term "critically ill" as it is used herein. The typical condition of treatment in an ICU environment and patient condition at entry leads to disease attributes [MeSH Descriptor C23.550.291] and pathological conditions [MeSH Descriptor C23] recognized by the medical community as a typical disorder of Critical Illness [MeSH Descriptor C23.550.291.625] a disease or state in which death is possible or imminent and with typical complication. Important complications specifically induced by the critical illness condition are critical illness myopathy (2012 ICD-9-CM Diagnosis Code 359.81) and critical illness polyneuropathy (2011 ICD-10-CM G62.81) and "Osteopenia" (ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851) which is a condition where bone mineral density is lower than normal more specifically (osteopenia is defined as a bone mineral density T-score between -1.0 and -2.5). Another important indication related to critical illness condition of critically ill patients is critical illness bone Atrophy. Critically ill patients often present with extreme bone loss, are at increased risk of
fractures such as stress fractures (ICD-10-CM 733.93 Stress fracture of tibia or fibula, 733.94 Stress fracture of the metatarsals, 733.95 Stress fracture of other bone, 733.96 Stress fracture of femoral neck, 733.97 Stress fracture of shaft of femur and/or 733.98 Stress fracture of pelvis) during rehabilitation, and can experience impaired healing of traumatic and surgical bone fractures. Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease, Osteoporosis Secondary to ICU Admission is a serious disorder that usually needs post-ICU ambulatory care. The bone resorption related to this disorder occurs in nearly all prolonged critically ill patients (B.Nierman (1998) Chest 114(4): 1122-8). It very often leads to complications such as hip fracture and vertebral fracture (A.Volk (2009) Am fam Physician 79(6): 459-64).

Critical illness uniformly predisposes to prolonged dependency on vital organ support. Lean tissue wasting of skeletal muscle and bone characterize the hypercatabolic state of prolonged critical illness (Hermans G, Vanhorebeek I, Derde S, Van den Berghe G. Metabolic aspects of critical illness polyneuromyopathy. Crit Care Med. 2009;37(10 Suppl):S391-7.). Specifically, circulating biomarkers of bone breakdown are extremely elevated whereas markers of bone formation are low (Van den Berghe G, Van Roosbroeck D, Vanhove P, Wouters PJ, De Pourcq L, Bouillon R. Bone turnover in prolonged critical illness: effect of vitamin D. J Clin Endocrinol Metab. 2003;88(10):4623-32.), an imbalance which may predispose critically ill patients to skeletal morbidity such as impaired fracture healing, osteoporosis, and increased risk of new fractures during rehabilitation. Examples of the latter are patients who underwent sternotomy for cardiac surgery and who develop slow or poor healing of the sternal fracture. This is referred to as sternal dehiscence, a serious complication that carries high morbidity and even mortality (Hashemzadeh K, Hashemzadeh S. In-hospital outcomes of delayed sternal closure after open cardiac surgery. J Card Surg. 2009;24(1):30-3.). Normal sternal bone healing may be further threatened by the necessity, in complicated cases, to leave the sternotomy wound open for a number of days in order to reduce intrathoracic pressure and allow the damaged heart to recover (Abid Q, Podila SR, Kendall S. Sternal dehiscence after cardiac surgery and ACE inhibitors [correction of ACE type 1 inhibition]. Eur J Cardiothorac Surg. 2001;20(1):203-4.). In addition, a recent retrospective case-cohort study has revealed a significant increase in fracture risk in patients who were in intensive care for a variety of reasons (Orford NR, Saunders K, Merriman E, Henry M, Pasco J, Stow P, et al. Skeletal morbidity among survivors of critical illness. Crit Care Med. 2011; 39(6): 1295-300.).
Together these data suggest a strong link between prolonged critical illness and skeletal morbidity.

Normal bone turnover depends on a tight coupling between function of mature osteoclasts, osteoblasts, and vascularization. This coupling requires a complex equilibrium of mechanical, endocrine, and nutritional factors. Prolonged critically ill patients are immobilized and suffer from a wide variety of endocrine and inflammatory disturbances, including hypercortisolism, hyposomatotropism, secondary hypothyroidism, hypogonadism, vitamin D deficiency and elevated cytokine levels such as tumor necrosis factor a (TNF-a), interleukin (IL-) 1 (IL-1) and IL-6 (2), all of which may contribute to the increase in bone resorption markers and reduced bone formation markers observed.

In other disease states characterized by excessive bone loss, such as postmenopausal osteoporosis or cystic fibrosis, increased osteoclast formation and activity of peripheral blood mononuclear cells (PBMC) has been reported (D'Amelio P, Grimaldi A, Pescarmona GP, Tamone C, Roato I, Isaia G. Spontaneous osteoclast formation from peripheral blood mononuclear cells in postmenopausal osteoporosis. FASEB J. 2005;19(3):410-2), both in the presence and absence of the canonical osteoclast activation factors, receptor activator of NF-κB ligand (RANKL) and macrophage colony stimulating factor (M-CSF) (Neale SD, Schulze E, Smith R, Athanasou NA. The influence of serum cytokines and growth factors on osteoclast formation in Paget’s disease. QJM. 2002;95(4):233-40). Increased osteoclast formation in a number of disease states characterized by excessive bone loss, such as postmenopausal osteoporosis, rheumatoid arthritis and cystic fibrosis appear to be related to elevated serum TNF-a, IL-1 and IL-6 levels (Shead EF, Haworth CS, Barker H, Bilton D, Compston JE. Osteoclast function, bone turnover and inflammatory cytokines during infective exacerbations of cystic fibrosis. J Cyst Fibros. 2009; Kaur K, Hardy R, Ahasan MM, Eijken M, van Leeuwen JP, Filer A, Thomas AM, Raza K, Buckley CD, Stewart PM, Rabbitt EH, Hewison M, Cooper MS. Synergistic induction of local glucocorticoid generation by inflammatory cytokines and glucocorticoids: implications for inflammation associated bone loss. Ann Rheum Dis. 2010). This is in contrast with our findings, where inflammatory cytokines do not appear to contribute to increased osteoclast formation. Hence these data show that increased osteoclast formation during critical illness differs from classical bone resorption diseases, which is a very novel insight. New bone formation and fracture healing requires precursor cells, located in the periosteum, a specialized connective tissue forming a thin but tough fibrous membrane firmly anchored to bone. Human periosteal-derived cells
(hPDCs) have been utilized as a clinically relevant model to examine the effects of illnesses on bone formation.

Until now, no studies have directly investigated the effect of critical illness on bone metabolism and repair at the tissue and cellular level. We here performed a series of studies to address this issue. Although biomarkers suggested excessive bone loss during critical illness, this had not been confirmed at the tissue level. Hence, we first quantified bone loss in an in vivo rabbit model of prolonged critical illness. Second, in an in vitro model, we investigated whether PBMCs isolated from critically ill patients are predisposed, more than those from healthy subjects, to differentiate into osteoclasts and assessed their osteoclastic activity. In addition, we studied the role of humoral factors in the patient's serum on these processes. Third, we assessed osteogenesis during critical illness, in an in vitro model using human periosteal cells, and in an in vivo murine model.

**Experimental**

By present invention it was demonstrated that osteoclasts isolated from ICU patient PBMCs form sooner than osteoclasts from healthy controls, whereby that osteoclasts isolated from patient PBMCs form in the absence and presence of RANKL and M-CSF.

Furthermore Autophagy marker p62 protein expression in healthy and sick (from ICU patient) osteoclasts (RANKL/MCSF stimulated) studies revealed that p62 protein is increased in osteoclasts from critically ill patients, suggesting a deficiency in autophagy, demonstrating role of autophagy increased osteoclast formation and activity in critical illness.

A dose finding of present demonstrated that spermidine inhibits osteoclast resorption. While osteoclast formation is significantly increased in Stimulated (+RANKL/MCSF) PBMCs from sick (ICU) patients vs. healthy controls, this increase is completely blocked by spermidine treatment.

It has been also demonstrated that osteoclasts isolated from patient PBMCs form sooner than osteoclasts from controls. Osteoclasts isolated from patient PBMCs form in the absence and presence of RANKL and M-CSF. p62 expression is increased in patient osteoclasts suggesting deficient autophagy. Increased osteoclast formation and activity in patient PBMCs is completely blocked with spermidine.
The present invention relates to novel therapeutic methods useful for the treatment or prevention of such bone deterioration or bone loss in the group of severe, life threatening diseases. Moreover the present invention provides compositions and treatments of Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease or Osteoporosis Secondary to ICU Admission.

The present invention relates to novel therapeutic methods useful for the treatment or prevention of stress fractures (ICD-10-CM 733.93 Stress fracture of tibia or fibula, 733.94 Stress fracture of the metatarsals, 733.95 Stress fracture of other bone, 733.96 Stress fracture of femoral neck, 733.97 Stress fracture of shaft of femur and/or 733.98 Stress fracture of pelvis) of patients that experienced extreme bone loss due to Metabolic Bone Disease Following ICU or due to bone resorption during ICU treatment that occurs in nearly all prolonged critically ill patients.

The methods, treatments and compositions of present invention are also for in use to treat or prevent disuse osteoporosis also called bone disuse atrophy, disuse atrophy of bone (ICD-9-CM Diagnosis Code 733.03), a form of osteoporosis due to immobilization or inactivity.

In general the invention can be considered as relating to the field of intensive care medicine. More specifically, the invention is based on the surprising finding that during critical illness there occurs an uncoupling between bone formation and degradation occurs. In a rabbit model of critical illness, pQCT analysis of proximal tibiae revealed a decrease in bone mineral content.

This was investigated further in a clinically relevant human in vitro model of critical illness, where peripheral blood mononuclear cells (PBMCs) from critically ill patients formed mature, multi-nuclear actively resorbing osteoclasts both in the presence and absence of osteoclastogenic factors RANKL and M-CSF, potentially due to an increase in circulating osteoclast precursors detected by flow cytometry. Treatment with 10% critically ill patient serum further increased osteoclast formation and activity.

No apparent differences in osteogenesis were observed in human perisoteal-derived cells (hPDCs) treated with patient serum in vitro, however, a decrease in the expression of VEGF-R1 suggested impaired vascularization. This was confirmed using serum-treated hPDCs
implanted onto calcium phosphate scaffolds in a murine in vivo model of bone formation, where decreased vascularization and increased osteoclast activity led to a decrease in bone formation in scaffolds with patient serum-treated hPDCs. In summary, we have shown that the disruption in bone metabolism observed in critically ill patients is predominantly explained by an increase in osteoclastogenesis coupled with a moderate decrease in bone formation which may be due to deficient vascularization.

Present invention provides composition and pharmaceutical compositions for suppressing osteoclastogenesis. The invention is based on the surprising finding on a novel mechanism that is at the basis of enhanced osteoclastogenesis in critically ill patients which is a cause of Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease or Osteoporosis Secondary to ICU Admission and is the reason why critically ill patients often present with extreme bone loss and are at increased risk of fractures such as stress fractures (ICD-10-CM 733.93 Stress fracture of tibia or fibula, 733.94 Stress fracture of the metatarsals, 733.95 Stress fracture of other bone, 733.96 Stress fracture of femoral neck, 733.97 Stress fracture of shaft of femur and/or 733.98 Stress fracture of pelvis) during rehabilitation and can experience impaired healing of traumatic and surgical bone fractures, can be determined by measuring peptides indicating a high physiological release of the potent vasoconstrictors as endothelin-1, in critically ill patients with high levels of adrenomedullin can exert a beneficial, potentially life-saving effect. Based on the novel mechanism of suppressing osteoclastogenesis compounds have been demonstrated to suppress such enhanced osteoclastogenesis and compositions and pharmaceutical compositions have been presented to treat or prevent Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease or Osteoporosis Secondary to ICU Admission.

The applicants have determined that the treatment of patient's peripheral blood mononuclear cell (PBMCs), in particular PBMCs of critically ill patients with autophagy inducers can be used to suppress enhanced osteoclastogenesis. It turned out that a particular advantage is that such autophagy inducers can be targeted to particularly treat the PBMCs. The applicants also determined that for such treatment surprisingly low doses of autophagy inducers can be used, possibly because patient's PBMCs are the treatment target cells. For instance by present invention the applicants used PBMCs from critically ill patients cultured for 14 days in critically ill patient serum and compounds added from dl-dl4 and cells staining with TRAP
for osteoclast formation at day 14 and demonstrated the inhibiting effect on critical illness enhanced osteoclast formation of low doses autophagy inducers such as Spermidine, Rapamycin, Everolimus, Promethazine and of methylation inhibitor such as the global methylation inhibitor-5-azacytidine.

Example 1. In vivo model of critical illness

Animals

All animals were treated according to the Principles of Laboratory Animal Care formulated by the U.S. National Society for Medical Research and the Guide for the Care and Use of Laboratory Animals prepared by the National Institutes of Health. The study protocol was approved by the Leuven University ethical review board for animal research (PI 08/2009). The model has been described in detail previously (Weekers F, Giuliani AP, Michalaki M, Coopmans W, Van Herck E, Mathieu C, et al. Metabolic, endocrine, and immune effects of stress hyperglycemia in a rabbit model of prolonged critical illness. Endocrinology. 2003;144(12):5329-38.). At day -1, adult, 3- to 4-month-old male New Zealand white rabbits weighing 3 kg were anesthetized and catheters were inserted in the right jugular vein and right carotid artery, allowing intravenous infusion of insulin and fluids and repetitive blood sampling. Fluid resuscitation consisting of hartmann (Baxter, Lessines, Belgium), enriched with glucose to prevent hypoglycemia, was started after the operation. At day 0, animals were anesthetized again, and a full thickness 15-20% body surface area third-degree burn injury was inflicted on the flanks, after having performed a paravertebral block with lidocaine (Xylocaine, AstraZeneca, Brussels, Belgium). Fluid resuscitation consisted of hartmann-glucose, and was changed to total parenteral nutrition at day 1. Animals were targeted to hyperglycemia, with concomitant hyperinsulinemia. This model has been validated as representative of the human critically ill condition (9, 36). Daily arterial blood was sampled and ionized calcium levels were measured immediately by a blood gas analyser (ABL725, Radiometer, Copenhagen, Denmark). A daily 4cc blood sample was collected and plasma stored at -80°C until further analysis. On day 7 animals were sacrificed and the left and right tibiae were dissected, dipped in polyvinylalcohol (PVA) and snap-frozen in liquid nitrogen, after which they were stored at -80°C until further analysis. For comparison, healthy rabbits were sacrificed and tissue samples were collected as described above. Plasma osteocalcin was measured by an in-house rabbit osteocalcin RIA as previously described (Van den Berghe G, Van Roosbroeck D, Vanhove P, Wouters PJ, De Pourcq L, Bouillon R. Bone turnover in prolonged critical illness: effect of vitamin D. J Clin Endocrinol Metab. 2003;88(10):4623-32., Bouillon R,

Peripheral Quantitative Computed Tomography

Trabecular and cortical BMC and BMD and the geometry of the right tibia were assessed ex vivo by peripheral quantitative computed tomography (pQCT) using the Stratec XCT Research densitometer (Norland Medical Systems, Fort Atkinson, WI, USA). Slices of 0.2 mm thickness were obtained using a voxel size of 0.070 mm. One scan was taken 2.4 mm from the proximal end of the tibia to measure trabecular volumetric density. The trabecular bone region was defined by setting an inner threshold to 30% of the total cross-sectional area. A second scan was taken 7 mm from the proximal end of the tibia (an area containing mostly cortical bone). These mid-diaphyseal scans were performed to determine cortical volumetric density, cortical thickness, and periosteal and endocortical perimeter. Cross-sectional moment of inertia, based on cortical bone measurements, was calculated using the circular ring model algorithm of the software program.

Example 2. In-vitro model of bone resorption during critical illness

Experimental subjects

Human peripheral blood was collected from prolonged critically ill patients (n = 12, 26-80 years of age, mean age 57 ± 16.39 years of age) and healthy control volunteers, matched for age, sex and body mass index (BMI) (n = 12, 23-81 years of age, mean age 57 ± 17.44 years of age). All protocols were approved by the Institutional Review Board of the Leuven University. Written informed consent was obtained from all healthy volunteers and from the patients or, when the patient was unable to give consent, from the closest family member. Prior to sample collection, it was ensured that no steroidal drugs or bisphosphonates had been taken by patients or healthy volunteers in the past 12 months.

Flow cytometry

Osteoclast precursors were detected in healthy volunteers or critically ill patients by staining fresh blood samples with allophycocyanin (APC)-conjugated anti-VNR, phycoerythrin (PE)-conjugated anti-CD14 and fluorescein isothiocyanate (FITC)-conjugated anti-CD14-labeled cells, or with the corresponding isotype control followed by incubation at 4°C for 30 min. A human Fc-gamma receptor (FcyR)-binding inhibitor was used to inhibit non-specific FcyR mediated binding. We treated double positive CD14+/CD11b+ cells as early osteoclast precursors, and triple positive CD14+/CD11b+/VNR+ cells as osteoclast precursors, according to previous literature (22, 38). Flow cytometry was performed on a FACSCalibur® flow cytometer (BD
The expression of membrane antigens was analyzed using BD FACSDiva® software (BD Biosciences).

**Peripheral blood mononuclear cell isolation and culture**

PBMCs were isolated from the whole blood of patients or healthy volunteers by means of Ficoll-Paque Plus (GE Healthcare, Brussels, Belgium) density gradient centrifugation according to the manufacturer's instructions. Heparinized whole blood (20 ml) was drawn from each individual and diluted 1:1 with alpha minimal essential medium (α-MEM; Lonza, Braine-l'Alleud, Belgium), before being layered over Ficoll-Paque and centrifuged at 450g for 40 min. The mononuclear cell interface was then removed and washed 3 times with α-MEM. Cells were stored in liquid nitrogen until use and seeded at a density of 5x10^5 cells per well. All cultures were performed in quadruplicate in 16-well culture slides (VWR, Leuven, Belgium) or 16-well BD Biocoat Osteologic Slides (BD Biosciences), and cells were cultured in α-MEM supplemented with 10% FBS, 100 IU/ml penicillin and 100 µg/ml streptomycin (‘complete medium’), or complete medium plus M-CSF (25 ng/ml) and RANKL (30 ng/ml).

In order to study the effect of circulating factors on osteoclast formation, PBMCs were also grown in α-MEM containing 10% human healthy volunteer serum (HS) or 10% patient serum (PS), in the presence or absence of M-CSF (25 ng/ml) and RANKL (30 ng/ml). For cytokine neutralization experiments, PBMCs were grown in α-MEM/10% PS for 14 days, in the presence of varying concentrations of anti-TNF-a and/or anti-IL-6 (R&D Systems, Abingdon, UK). All cultures were maintained at 37°C in a humidified atmosphere with 5% CO2.

**Osteoclast formation and activity**

After 14 days, PBMCs were fixed and stained for the osteoclast-specific marker tartrate-resistant acid phosphatase (TRAP), and the nuclear marker DAPI, as previously described. The formation of TRAP positive multi-nucleated (more than three nuclei) cells was quantified by counting the multi-nuclear stained cells in each well. (D'Amelio P, Grimaldi A, Pescarmona GP, Tamone C, Roato I, Isaia G. Spontaneous osteoclast formation from peripheral blood mononuclear cells in postmenopausal osteoporosis. FASEB J. 2005;19(3):410-2.). Filamentous (F)-actin rings representing active osteoclasts were visualized after staining of the actin cytoskeleton with the toxin phalloidin conjugated to FITC, and quantified by counting as described above. To evaluate osteoclast activity, cells were removed from the hydroxyapatite-coated wells with 14% sodium hypochlorite, and the mineral layer was stained with Von-Kossa as previously described. (Ariyoshi W, Takahashi T, Kanno T, Ichimiya H, Shinmyouzu K, Takano H, et al. Heparin inhibits osteoclastic differentiation and function. J Cell Biochem. 2008;103(6):1707-17). Lacunar resorption was determined by measuring the
total area unstained by Von-Kossa (total area resorbed) using ImageJ software (National Institute of Health, MD, USA), and was expressed as the total percentage of the surface reabsorbed.

**Example 3. In-vitro model of osteoblast differentiation during critical illness**

**Cell Culture**

Human Periosteal Derived Cells (hPDCs) were obtained from the Laboratory for Skeletal Development and Joint Disorders, Katholieke Universiteit Leuven, Leuven, Belgium. Cells were expanded in monolayer at 37°C in a humidified atmosphere of 5% CO2 in growth medium, which consisted of high-glucose Dulbecco's modified Eagle medium (DMEM; Invitrogen, Merelbeke, Belgium) containing 10% γ-irradiated and filtered FBS (Gibco), 1% sodium pyruvate (100 mM; Invitrogen) and 1% antibiotic-antimycotic solution (100 units/ml penicillin, 100 µg/ml streptomycin and 0.25 µg/ml amphotericin B; Invitrogen). The medium was replaced every 3 days. All experiments were carried out with expanded cell populations between passage 5 and 7, with a seeding density of 4500 cells/cm². After 48 hours in culture, the growth medium of in vitro osteogenic assays was replaced using osteogenic medium, which consisted of FBS-free growth medium supplemented with 100 nM dexamethasone (Sigma-Aldrich, Bornem, Belgium), 10 mM glycerol-2-phosphate disodium salt hydrate (Sigma-Aldrich), 50 µM L-ascorbic acid 2-phosphate sesquimagnesium salt hydrate (Sigma-Aldrich) and 10% serum (10% FBS, 10% PS, or 10% HS). The medium was replaced every 3 days.

**In vitro Mineralization Assay**

hPDCs were analyzed for in vitro mineralization using Alizarin Red to detect mineral deposits, as previously described. Briefly, hPDCs were seeded at 4500 cells/cm² and treated for 21 days with osteogenic medium and FBS, HS or PS. At day 21, the cultures were rinsed with PBS, fixed with 4% formaldehyde, and then rinsed with distilled water before staining with an Alizarin Red solution (Sigma; pH 4.2). Nonspecific staining was rinsed off carefully with distilled water. Quantification of calcium mineral deposits was performed by dissolving the dye with 10% cetylpyridinium chloride (in demineralized water) for 10 min at RT. Absorbance was measured spectrophotometrically at 570 nm.

**Gene Expression Analysis**

hPDCs were seeded at 4500 cells/cm² and treated for 7 days with osteogenic medium and FBS, HS or PS. Total RNA was isolated using the RNeasy kit (Qiagen Benelux, Venlo, Netherlands) and cDNA was synthesized with the Superscript III First Strand synthesis system for real-time PCR (Invitrogen). Quantitative real-time SYBR Green (Invitrogen) PCR
was performed according to the manufacturer's protocol, with mRNA levels normalized to \( \beta \)-actin expression. SYBR Green qPCR primers were designed to span an intron so that only RNA-specific amplification was possible (RUNX2-F, 5'-CGCATCTCATCCCAGTAT-3'; RUNX2-R, 5'-GCCTGGGTCTGTGAATCTGA-3'; COL1A1-F, 5'-GACGAAGACATCCACCAAT-3'; COL1A1-R, 5'-AGATCACGTCATCGACAAC-3'; ALP-F, 5'-GGACATGCAGTACGTGAA-3'; ALP-R, 5'-GTCAATTCTGCTCTTCCA-3'; VEGFA-F, 5'-CCCACCTGAGGAGTCCAACAT-3'; VEGFA-R, 5'-GCATCACATTGGTCTGCTG-3'; VEGF-R1-F, 5'-AAGCAAAAACTGCGCTTC-3'; VEGF-R1-R, 5'-CGGAGTTACGTGACATCTCT-3'). Total RNA samples subjected to cDNA synthesis reactions in the absence of reverse transcriptase were included as negative controls and relative differences in expression were calculated using the 2\( \Delta \)ACT method (Livak KJ, Schmittgen TD, Analysis of relative gene expression data using real-time quantitative PCR and the 2(\(-\Delta\Delta C(T)\)) Method. Methods. 2001;25(4):402-8).

**Western Immunoblotting**

For protein expression analysis, hPDCs were seeded at 4500 cells/cm\(^2\) and treated for 7 days with osteogenic medium and FBS, HS or PS. After 7 days the cell monolayer was washed with ice-cold 1xPBS, and lysates were prepared by adding lysis buffer (RIPA buffer; 50 mM Tris, 150 mM NaCl, 0.1% SDS, 0.5% Na-deoxycholate, 1% TritonX-100, 1 mM phenylmethanesulfonylfluoride (PMSF)) containing protease and phosphatase inhibitors. Lysates were scraped into a 1.5 ml microcentrifuge tube, and centrifuged for 15 min at 4°C. Supernatants were transferred to fresh microcentrifuge tubes and stored at -80°C. The protein concentration was determined by the Pierce BCA Protein Assay (VWR). Equal amounts of protein were loaded onto each lane of a 4-12% Bis-Tris gel and subjected to electrophoresis under reducing conditions. After blotting, polyvinylidene difluoride membranes were blocked for one hour (5% milk powder in 0.1% PBS/Tween) and incubated with Rabbit anti-human VEGFRI primary antibody (Abeam, Cambridge, UK; 1:500) overnight at 4°C. Binding of Goat anti-rabbit IgG HRP secondary antibody (DakoCytomation, Glostrup, Denmark; 1:2000) was visualized by enhanced chemiluminescence (ECL). Normalization for total protein was performed by re-probing the membrane with Mouse anti-mouse Beta-actin (Abeam, Cambridge, UK; 1:1000) for 1 hour at RT, followed by Goat anti-mouse IgG HRP (DakoCytomation; 1:2000) for 1 hour at RT.

**Example 4. In-vivo model of bone formation during critical illness**

*In vivo Bone Formation*
In vivo bone formation was analyzed as described recently (Roberts SJ, Geris L, Kerckhofs G, Desmet E, Schrooten J, Luyten FP. The combined boneforming capacity of human periosteal derived cells and calcium phosphates. Biomaterials.32(19):4393-405). Briefly, hPDCs (passage 5) were seeded at 4500 cells/cm² and treated for 7 days with osteogenic medium and FBS, HS or PS. After 7 days, the cells were trypsin released, centrifuged and resuspended at a concentration of 50 million cells/ml. Subsequently, 20 μl of the cell suspension was applied to the upper surface of each scaffold. To allow cell attachment, the seeded scaffolds were incubated overnight at 37°C. After incubation, the constructs were either directly implanted subcutaneously in the back at the cervical region of NMRI-nu/nu mice. The remaining cells in the supernatant were counted to estimate the seeding efficiency which was calculated as follows: [(number of seeded cells - number of cells in the supernatant)/number of seeded cells] × 100. The implants were collected after 56 days of implantation. Each explant was fixed in 4% formaldehyde, scanned by μCT, decalcified in EDTA/PBS (pH 7.5) for 2 weeks, paraffin embedded and processed for histology. All procedures on animal experiments were approved by the local ethical committee for Animal Research (Katholieke Universiteit Leuven). The animals were housed according to the guidelines of the Animalium Leuven (Katholieke Universiteit Leuven).

Bone quantification

To quantify ectopic bone formation, μCT was used to quantify the volume of new bone formed in three dimensions by segmenting the newly formed mineralized tissues from the calcium phosphate grains in each material. For segmentation, a manually selected, but consistent global threshold value was used for each scaffold. The choice of threshold value was confirmed by visual comparison to the corresponding histological sections.

Histological analysis

Histological staining was performed on paraffin embedded sections of cell/biomaterial constructs. Briefly, paraffin sections were deparaffinized in Histoclear™ (Laborimpex, Brussels, Belgium) and methanol, and rinsed with distilled water. For TRAP staining the tissue sections were incubated in TRAP buffer (50 mm Sodium Acetate, 30 mm Sodium Tartrate, 0.1% Triton X-100, pH 5) for 20 min. This was then replaced with TRAP stain (0.5 mg/ml napthol AS-MX phosphate and 1.1 mg/ml Fast red violet LB in TRAP buffer prewarmed to 37°C) for 1 h before washing twice in PBS, counterstaining with hematoxylin and mounting with aqueous mounting medium. TRAP-positive osteoclasts were visualized using fluorescent microscopy, and quantified using ImageJ® software (National Institutes of Health, USA). For CD31 immunohistochemistry, the tissue sections underwent antigen
retrieval in proteinase K for 15min at 95°C and were blocked in 1% normal rabbit serum, before being incubated overnight at 4°C with Rat Anti-mouse CD3 1 (BD Pharmingen; 1:20). Sections were then probed with the biotinylated rabbit anti-rat IgG antibody (Vector Laboratories; 1:100), and the signal amplified with the Vector Laboratories 'ABC system'. The signal was detected using 3,3’-Diaminobenzidine (DAB) the sections counterstained with hematoxylin, and the number of CD3 1 positive vessels per scaffold was visually quantified.

**Statistical analysis**

The data were processed using the Statistical software package StatView 5.0.1 (SAS Institute Inc.). Student's t test was used for the comparison of normally distributed data (presented as mean ± SD), and the Mann-Whitney U test for data that were not normally distributed (presented as median and IQR, unless otherwise indicated). P values less than or equal to 0.05 were considered statistically significant. Statistical significance is indicated on all graphs as follows: *: p < 0.05, **: p < 0.01, ***: p < 0.001.

**Example 5 Osteoclasts in critical illness**

Isolation of osteoclasts from human peripheral blood mononuclear cells

15-20ml peripheral blood collected in lithium-heparin tubes

Mixed with aMEM and add to Ficoll-Paque, centrifuged and mononuclear cells separated

Plate down cells in aMEM with 10% serum and +/-RANKL and MCSF

By week 3 extensive resorption can be observed and osteoclast formation can be measured by staining for TRAP. Also Markers of osteoclast activity significantly increased during critical illness

**Example 6 Spermidine induced reduction of osteoclast activity**

On peripheral blood mononuclear cells (PBMCs) isolated from ICU patients (4) and healthy controls (5) present invention demonstrates by a dose response experiment using healthy cells only (Figure 1) that osteoclast activity is reduced at 10⁻⁶, 10⁻⁷, and 10⁻⁸ M Spermidine (Sigma S4139, lot 000141 1140>.

**Example 7 Spermidine induced reduction of osteoclast activity in critically ill derived PBMCs (stimulated)**

In the next experiment, we measured osteoclast formation (Figure 2), in healthy and sick (from critically ill) stimulated (with RANKL and MCSF) PBMCs with and without
spermidine (10^{-8} \text{ M}). By day 7 there was already a noticeable difference between the treatment groups (Figure 2A top panel), and when the cells were stopped at day 14, there was a significant increase in TRAP positive multinuclear cells (indicative of increased osteoclast formation) in sick vs. healthy cells, and a significant reduction in TRAP positive cells (indicative of reduced osteoclast activity) in spermidine treated sick PBMCs compared to 'control' sick PBMCs (Figure 2A, bottom 2 panels, and Figure 2B). Spermidine treatment did not, however, have a significant effect on the healthy PBMCs.

**Example 8** Spermidine induced reduction of osteoclast activity in critically ill derived PBMCs (unstimulated)

We also looked at spontaneous osteoclast formation in healthy and sick unstimulated (no RANKL or MCSF) PBMCs, with and without spermidine. As previously, we found a significant increase in osteoclast formation in sick PBMCs vs. healthy, and this increase was completely knocked down with spermidine (Figure 3A and B). Again we found there was no effect on the healthy cells with spermidine.

**Example 9** Osteoclast resorption measurement in stimulated PBMCs, unstimulated PBMCs of healthy and sick (critically ill) subjects and the effect of spermidine thereon.

In addition to osteoclast formation, we also looked at osteoclast activity, by measuring osteoclast resorption of a hydroxyapatite layer. The area of hydroxyapatite 'eaten into' can be measured using ImageJ software. In stimulated PBMCs, there was a significant increase in resorption in sick cells compared to healthy, and again this increased was reduced with spermidine treatment (Figure 4A and B). In unstimulated PBMCs, only a very small amount of resorption was seen in any of the treatment groups (Figure 5A and B), however, there was a small increase in resorption in the sick cells compared to healthy.

**Example 10** Effects of spermidine on autophagy markers.

We also measured markers of autophagy in the cell lysates from these experiments. The concentration of proteins we obtained were quite low, and there was insufficient protein to measure anything by western blotting in the 'sick spermidine' group. However, we were able to measure p62 in the healthy, sick, and healthy spermidine cells, and found that p62 protein was significantly increased in sick cells compared to healthy, suggesting an accumulation of p62 and therefore defective autophagy in sick osteoclasts. There was no effect of spermidine on the healthy PBMCs (Figure 6).
Example 11. Trabecular, but not cortical, bone is lost in a rabbit model of critical illness
A 7-day rabbit model of critical illness has previously been validated to examine localized
effects of critical illness (Weekers F, Giulietti AP, Michalaki M, Coopmans W, Van Herck E,
Mathieu C, et al. Metabolic, endocrine, and immune effects of stress hyperglycemia in a
rabbit model of prolonged critical illness. Endocrinology. 2003;144(12):5329-38). Like in
human critically ill patients, serum ionized calcium levels were significantly reduced in
critically ill rabbits (Figure 19A). Infliction of the critical illness evoked an immediate
decrease in ionized calcium on day 1 (3.4%), which remained lower than in healthy rabbits
for the entire 7-day period of illness. Also similar to what is described in human patients,
serum osteocalcin levels were lowered in critically ill rabbits by day 4 of illness (Figure 19B).
In order to examine the direct effect of critical illness on skeletal integrity, bone mineral
content (BMC) and bone mineral density (BMD) were measured in the proximal tibia after 7
days of illness. Importantly, trabecular BMC and BMD were significantly lower in critically
ill rabbits than in healthy controls (Figures 19C and D), but no differences were observed in
cortical BMC, BMD or thickness (Figures 19E and F).

Example 12. Increased osteoclast formation and activity in critically ill patients occurs
spontaneously and is potentiated with autologous patient serum
To further examine the cause of bone loss during critical illness in a clinically relevant model,
PBMCs were isolated and pooled from an age- sex- and BMI-matched set of critically ill
patients and healthy controls, and analyzed for osteoclast formation and activity.
To establish whether cells from critically ill patients are programmed to differentiate into
osteoclasts before any in vitro manipulation, the number of osteoclast precursors present in
critically ill patient or healthy control peripheral blood was assessed by Fluorescence
Activated Cell Sorting (FACS) analysis. Double positive CD14+/CD1 lb+ cells are
considered to be early osteoclast precursors, and triple positive CD14+/CD1 lb+/VNR+ cells
are considered to be more mature circulating osteoclasts (D'Amelio P, Grimaldi A,
Pescarmona GP, Tamone C, Roato I, Isaia G. Spontaneous osteoclast formation from
peripheral blood mononuclear cells in postmenopausal osteoporosis. FASEB J.
Mechanisms of spontaneous osteoclastogenesis in cancer with bone involvement. FASEB J.
2005;19(2):228-30). There were more early osteoclast precursors (CD14+, CD1 lb+) in blood
from critically ill patients than in blood from healthy controls (Figure 20A and 20B) with only a trend for more mature circulating osteoclasts (CD14+, CD1 lb+, VNR+).

To confirm that these precursor cells indeed form active osteoclasts, the isolated PBMCs were cultured for 14 days, with and without RANKL and M-CSF, and the number and activity of osteoclasts in patients were compared with healthy matched controls. After 14 days of culture with RANKL and M-CSF, the formation of mature, multi-nuclear (>3 nuclei, tartrate-resistant acid phosphatase (TRAP) positive) osteoclasts was 16.8-fold higher in cultures from critically ill patients than those from healthy controls (Figure 21, A and B). Analysis of the activity of the osteoclasts was done by quantifying the degree of hydroxyapatite-resorption after culturing the cells for 21 days in hydroxyapatite-coated wells. This revealed a 36.2-fold higher resorption in patient PBMC cultures with RANKL and M-CSF as compared with controls (Figure 21, A and C). PBMCs from critically ill patients also displayed spontaneous differentiation into osteoclasts (without RANKL and M-CSF; 23.8-fold increase vs. healthy cells), but the degree of hydroxyapatite resorption in spontaneously-formed osteoclast cultures from patients and controls was similar (Figure 21, A and C).

To assess the impact of humoral factors in the patient's blood on the differentiation potential of PBMCs into osteoclasts, PBMCs from patients and controls were cultured in the presence of 10% autologous patient serum (PS) or 10% autologous healthy serum (HS). In the presence of RANKL and M-CSF, patient PBMCs cultured with 10% HS did not form significantly more mature, multi-nuclear TRAP positive osteoclasts per well than cells from healthy control grown under the same conditions (Figure 22, A and B). In addition, also without RANKL and M-CSF, in the presence of 10% HS, minimal spontaneous osteoclast differentiation was observed, similar in patients and controls. However, in the presence of RANKL and M-CSF, the addition of 10% PS resulted in a 3.7-fold increase in osteoclast formation in patient PBMCs compared to cells grown with 10% HS. A similar increase was observed in 'unstimulated' patient PBMC cultures (without RANKL and M-CSF), as when cultured in 10% PS, spontaneous osteoclast formation was increased by 6.1-fold compared to spontaneous osteoclast formation in patient PBMC cultures with 10% HS (Figure 22, A and B). However, the addition of 10% PS to cultures of healthy control PBMCs did not increase osteoclast formation in the presence or absence of RANKL and M-CSF. These data suggest a unique interaction between patient cells and PS that is not present in healthy control cultures and PS.

The formation of F-actin rings displayed a similar pattern to that observed with the formation of TRAP positive multi-nuclear cells. A significant increase in F-actin ring formation was
detected both with and without RANKL and M-CSF in patient cells cultured with 10% HS (30-fold and 6.25-fold, respectively) (Figure 22, A and C), compared to healthy control cells cultured with 10% HS. A further 5.9-fold increase in F-actin ring formation was observed in patient cells grown in 10% PS, compared to 10% HS, and when grown without RANKL and M-CSF, an increase of 4.72-fold was still observed.

Resorption of hydroxyapatite was also significantly increased when 10% PS was added to patient PBMC cultures, but not when added to healthy control cultures (Figure 22, A and D). With PS added to patients cells cultured in the presence of RANKL and M-CSF, 62.7% of the total surface area was resorbed as compared with 15.8% with HS (3.9-fold increase). A 1.63 fold increase in resorption was maintained in the absence of RANKL and M-CSF.

In order to assess whether the humoral factors that seem to interact with the primed PBMCs from critically ill patients could be the elevated levels of the major pro-inflammatory cytokines (Van den Berghe G, Van Roosbroeck D, Vanhove P, Wouters PJ, De Pourcq L, Bouillon R. Bone turnover in prolonged critical illness: effect of vitamin D. J Clin Endocrinol Metab. 2003;88(10):4623-32), cells were grown in 10% PS with or without neutralizing antibodies for IL-6 or TNF-a. Unexpectedly, however, adding anti-IL6 or anti-TNF-a further increased the formation of multi-nuclear TRAP positive osteoclasts in a dose-dependent manner. Strikingly, adding both antibodies in combination resulted in the formation of giant multi-nuclear osteoclasts (Figure 23, A, B and C).

Example 13. Critically ill patient serum does not affect osteoblast differentiation but reduces expression of angiogenesis markers in vitro. In order to assess the impact of circulating factors during critical illness on bone formation, an in vitro model of osteogenesis during critical illness was set up. For this purpose, we used hPDCs, a pool of mesenchymal cells isolated from the human periosteum, that have shown to proliferate, migrate and differentiate into chondrogenic and osteogenic lineages upon stimuli such as trauma, fracture or infection (De Ban C, Dell’Accio F, Vanlauwe J, Eyckmans J, Khan IM, Archer CW, et al. Mesenchymal multipotency of adult human periosteal cells demonstrated by single-cell lineage analysis. Arthritis Rheum. 2006;54(4):1209-21. Roberts SJ, Chen Y, Moesen M, Schrooten J, Luyten FP. Enhancement of osteogenic gene expression for the differentiation of human periosteal derived cells. Stem Cell Res. 7(2):137-44.). Cells were cultured for 21 days in osteogenic medium, as previously described (Roberts SJ, Chen Y, Moesen M, Schrooten J, Luyten FP. Enhancement of osteogenic gene expression for the differentiation of human periosteal derived cells. Stem Cell Res. 7(2):137-44, De Bari C, DelVAccio F, Luyten FP. 27
Human periosteum-derived cells maintain phenotypic stability and chondrogenic potential throughout expansion regardless of donor age. Arthritis Rheum. 2001;44(1):85-95. As a late marker of hPDC osteoblast differentiation, calcium deposition was measured quantitatively using Alizarin Red staining on differentiated hPDCs.

Addition of fetal bovine serum (FBS), HS or PS to the osteogenic medium did not affect differentiation nor mineralization of the cells (Figure 24, A and B). Also gene expression analysis of osteogenic markers RUNX2 and COL1A1 (type I collagen) revealed no difference between PS and HS (despite a reduction in expression of these genes in both these groups as compared with cells grown in FBS) (Figure 24, C and D). Expression of ALP (bone specific alkaline phosphatase) was also not different between PS and HS (Figure 24E). These data suggest that osteogenic differentiation is unaffected by circulating factors during critical illness, assessed in vitro. Interestingly, expression of the angiogenesis factor vascular endothelial growth factor receptor (VEGF) alpha (VEGFA) was not significantly different between PS and HS groups (Figure 24F), although a significant reduction in VEGF-receptor 1 (VEGFR1) expression with PS as compared with HS was detected at the gene and protein level, suggesting that angiogenesis may be compromised during bone formation and skeletal healing in critical illness (Figures 24G and H).

Example 14. In an in vivo murine model of steletogenesis, bone formation by critically ill patient cells is reduced, coinciding with increased osteoclast activity and reduced angiogenesis

Bone formation is a multi-factorial process involving a tightly-regulated cascade of events which include the recruitment, proliferation, and differentiation of osteoprogenitor cells, along with the formation of a well-defined vascular compartment. Therefore although no differences in osteogenesis in the presence of PS were observed in hPDCs in vitro, it was plausible that aberrant bone formation might still occur in vivo through deficient angiogenesis. In order to test this hypothesis, hPDCs were incubated with FBS, HS or PS for 7 days in basal medium, before being seeded onto calcium phosphate (CaP) NuOssD scaffolds overnight, transplanted into NMRI-nu/nu mice, and incubated for 8 weeks (Roberts SJ, Geris L, Kerckhofs G, Desmet E, Schrooten J, Luyten FP. The combined bone forming capacity of human periosteal derived cells and calcium phosphates. Biomaterials.32(19):4393-405). Upon explanation and analysis of bone formation by µCT, scaffolds containing PS-treated hPDCs revealed significantly less mature bone than those containing HS-treated cells (Figure 25A). No significant differences in calcium phosphate
granules or fibrous tissue were observed. In view of the increased osteoclast activity observed in previous experiments, possibly contributing to bone hyperresorption in critical illness, the trend for an increase in osteoclastic activity in PS scaffolds, as detected by TRAP staining, was unsurprising (Figure 25B). The local vasculature is vital for the formation of new bone during normal bone maintenance and following fracture. Therefore, the formation of blood vessels was measured with CD31 immunohistochemistry, and revealed that angiogenesis was significantly reduced in PS scaffolds compared to HS scaffolds (Figure 24C).

The present invention is predicated on the discovery by the inventors that the disorder of critical illness [MeSH Descriptor C23.550.291.625] correlates with increased osteoclast formation from circulating precursors in the blood, osteoclast maturation and osteoclast bone resorption activity in comparison to healthy patients, resulting in complications such as "Osteopenia" (ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851) or leading to the indication Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease, Osteoporosis Secondary to ICU Admission with more extreme bone loss which increases the risk of critically ill patients for fractures such as stress fractures (ICD-10-CM 733.93 Stress fracture of tibia or fibula, 733.94 Stress fracture of the metatarsals, 733.95 Stress fracture of other bone, 733.96 Stress fracture of femoral neck, 733.97 Stress fracture of shaft of femur and/or 733.98 Stress fracture of pelvis) and that usually needs that usually needs post-ICU ambulatory care. The applicants found that trabecular bone mineral content and density were reduced in critically ill rabbits, coinciding with low serum levels of ionized calcium and osteocalcin. Moreover it was found that increased number of PBMC osteoclast precursors in the blood of critically ill humans resulted in increased in vitro osteoclast formation, which was further potentiated by the addition of serum from critically ill patients. Unexpectedly, neutralizing the inflammatory cytokines TNF-a and IL-6 in patient serum further increased osteoclast formation in patient PBMC cultures.

In an in vitro model of osteoblast formation, hPDC differentiation was unaffected by critically ill patient serum, although a reduction in VEGF-R1 gene and protein expression was observed. In vivo, bone formation was reduced in scaffolds containing patient serum-treated hPDCs, possibly due to an increase in osteoclastic activity together with a reduction in vascularization. As previously reported, the rabbit model of critical illness mirrors many of the metabolic and endocrine changes observed in critically ill patients (Weekers F, Giulietti
AP, Michalaki M, Coopmans W, Van Herck E, Mathieu C, et al. Metabolic, endocrine, and immune effects of stress hyperglycemia in a rabbit model of prolonged critical illness. Endocrinology. 2003;144(12):5329-38.). In the current study, critically ill rabbits displayed low levels of ionized calcium throughout the 7-day period of illness. During critical illness in humans, abnormalities in ionized calcium levels are common, with 90% of patients displaying mild hypocalcaemia, although hypercalcemia has also been reported in the protracted phase of illness (Egi M, Kim I, Nichol A, Stachowski E, French CJ, Hart GK, et al. Ionized calcium concentration and outcome in critical illness. Crit Care Med. 2011; 39(6): 1295-300). Ostensibly, these alterations may be linked with decreased levels of vitamin D, which have previously been reported during critical illness (Van den Berghe G, Van Roosbroeck D, Vanhove P, Wouters PJ, De Pourcq L, Bouillon R. Bone turnover in prolonged critical illness: effect of vitamin D. J Clin Endocrinol Metab. 2003;88(10):4623-32). Also, the observed reduction in serum osteocalcin in critically ill rabbits after 4 days is in line with findings in critically ill humans (Van den Berghe G, Van Roosbroeck D, Vanhove P, Wouters PJ, De Pourcq L, Bouillon R. Bone turnover in prolonged critical illness: effect of vitamin D. J Clin Endocrinol Metab. 2003;88(10):4623-32), which suggests that bone formation may also be reduced during critical illness. This was confirmed by the applicant’s study, in which trabecular BMC and BMD were found reduced by 30.6% and 28.8%, respectively. In another rabbit model of osteoporosis evoked by methylprednisolone, trabecular BMD was reduced by 36% only after 10 weeks of treatment (Baofeng L, Zhi Y, Bei C, Guolin M, Qingshui Y, Jian L. Characterization of a rabbit osteoporosis model induced by ovariectomy and glucocorticoid. Acta Orthop.81(3):396-401. PMCID: 2876847), and therefore the observation of similarly reduced trabecular bone after only 7 days of critical illness is remarkable. A severe catabolic state is observed during critical illness (Weekers F, Giulietti AP, Michalaki M, Coopmans W, Van Herck E, Mathieu C, et al. Metabolic, endocrine, and immune effects of stress hyperglycemia in a rabbit model of prolonged critical illness. Endocrinology. 2003;144(12):5329-38)

There was observed a direct association between the extreme catabolic state experienced during critical illness and subsequent skeletal morbidity. This is confirmed by a recent retrospective longitudinal case-cohort study by Orford and colleagues revealing an increased risk of fragility fractures over an 8-year period in elderly female survivors of critical illness (Orford NR, Saunders K, Merriman E, Henry M, Pasco J, Stow P, et al. Skeletal morbidity among survivors of critical illness. Crit Care Med. 2011; 39(6): 1295-300).

This increase in osteoclast precursors is often characterized by an increase in osteoclast formation and activity, which can be recapitulated *in vitro* through the isolation and differentiation of PBMCs into osteoclasts, with the addition of osteoclastogenic factors such as RANKL and M-CSF. The applicants found that, PBMCs isolated from critically ill patients displayed a significant increase in differentiation into osteoclasts compared to healthy controls.

In cultures of PBMCs from critically ill patients, we observed osteoclast formation both in the presence and absence of RANKL and M-CSF. However, these mature, multi-nuclear osteoclasts were only able to actively resorb hydroxyapatite in the presence of osteoclastogenic factors.

In the current study, osteoclast formation and activity was further increased in patient PBMC cultures upon the addition of 10% PS (patient serum), whereas no increase was observed with HS (Healthy serum), or in healthy PBMC cultures treated with PS. This suggested that factors present in the PS were reacting with unique factors present in patient PBMCs to promote osteoclastogenesis. Inflammatory cytokines have previously been shown to promote the formation of osteoclasts *in vitro* and *in vivo* (*D’Amelio P, Roato I, D’Amico L, Veneziano L, ...
Suman E, Sassi F, et al. Bone and bone marrow pro-osteoclastogenic cytokines are upregulated in osteoporosis fragility fractures. Osteoporos Int., Roato I, D'Amelio P, Gorassini E, Grimaldi A, Bonello L, Fiori C, et al. Osteoclasts are active in bone forming metastases of prostate cancer patients. PLoS One. 2008;3(11):e3627. PMCID: 2574033, Boyce BF, Li P, Yao Z, Zhang Q, Badell IR, Schwarz EM, et al. TNF-alpha and pathologic bone resorption. Keio J Med. 2005;54(3):127-31), and have been directly related to fragility fractures in postmenopausal females (D'Amelio P, Roato I, D'Amico L, Veneziano L, Suman E, Sassi F, et al. Bone and bone marrow pro-osteoclastogenic cytokines are up-regulated in osteoporosis fragility fractures. Osteoporos Int.). However, neutralizing antibodies for both TNF-a and IL-6 further increased osteoclast formation in patient PBMC cultures with 10% PS. Resorption is thought to be mediated largely by the increased local production of pro-inflammatory cytokines, such as TNF-a, which is thought to induce resorption indirectly by affecting the production of RANKL and/or its soluble decoy receptor, osteoprotegerin, by osteoblasts or directly by enhancing the activity of osteoclasts (Boyce BF, Xing L. Functions of RANKL/RANK/OPG in bone modeling and remodeling. Archives of Biochemistry and Biophysics. 2008;473(2):139-46, Otero JE, Dai S, Alhawagri MA, Darwech I, Abu-Amer Y. IKKbeta activation is sufficient for RANK-independent osteoclast differentiation and osteolysis. J Bone Miner Res. 25(6):1282-94. PMCID: 3153134.).

Furthermore the impact of prolonged critically ill patient serum on the differentiation capacity of hPDCs was evaluated by mineralization of the extracellular matrix with calcium deposition. Although an increase in mineralization was observed in all differentiated conditions compared to non-differentiated cells, no differences were observed between HS or PS. Similarly, gene expression of RUNX2, COL1A1 and ALP revealed no differences in expression between HS and PS conditions, although a reduction in RUNX2 and COL1A1 expression was observed in both human serums compared to the standard FBS condition. Under specific conditions, hPDCs are known to be capable of differentiation into the chondrocyte, osteoblast, adipocyte, and skeletal myocyte lineages in vitro and in vivo.

However, in the current study carried out by applicants, hPDCs were directed towards the osteoblast lineage in order to focus directly on the effect of critical illness during bone formation. Although these findings suggest that in vitro, factors such as inflammatory cytokines present in critically ill PS do not have an effect on osteogenic differentiation, the fact that the expression of VEGF-R1 at both the gene and protein level was significantly reduced with PS suggests that vascularization may be inhibited during skeletal healing and bone formation in critical illness.

This hypothesis was corroborated by the in vivo model of bone formation during critical illness, where a reduction in bone formation in PS-coated NuOss™ scaffolds implanted in NMRI-nu/nu mice correlated with a significant reduction in vascularization, along with an increase in osteoclast activity. This finding directly supports the hypothesis that hypoxia is a major risk factor for impaired fracture healing, such as evidenced by sternal healing problems after internal mammary artery harvesting for coronary bypass surgery. Due to the fact that the majority of studies aimed at enhancing sternal closure utilize mechanical interventions such as wire stabilization (Iwakura A, Tabata Y, Nishimura K, Nakamura T, Shimizu Y, Fujita M, et al. Basic fibroblast growth factor may improve devascularized sternal healing. Ann Thorac Surg. 2000;70(3):824-8, Iwakura A, Tabata Y, Koyama T, Doi K, Nishimura K, Kataoka K, et al. Gelatin sheet incorporating basic fibroblast growth factor enhances sternal healing after harvesting bilateral internal thoracic arteries. J Thorac Cardiovasc Surg. 2003;126(4):1113-20), the finding that vascularization and bone formation in the model was inhibited by critically ill PS suggests that a biological approach to enhance fracture healing aimed at increasing vascularisation also warrants further investigation.

An autophagy activator compound for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject can have different structures. The autophagy
activator compound can be a compound of the group consisting of Rapamycin, Nigericin, Wiskostatin, Fluspirilene, Niguldipine, Trifluoperazine, Nicardipine and Penitrem A (Tremortin). An autophagy activator compound for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject can have different structures.

The autophagy activator compound can be a compound of the group consisting of Loperamide, Amiodarone, Niguldipine, Pimozide, Nicardipine, Penitrem A, Fluspirilene, Trifluoperazine, and pharmaceutically acceptable salts thereof. Trehalose, a non-reducing disaccharide present in many non-mammalian species, including bacteria, yeast, fungi, insects, invertebrates (natural hemolymph sugar of invertebrates), and plants is an autophagy inducer according present invention for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Trehalose is also known as mycose or tremalose, is a natural alpha-linked disaccharide formed by an α-1,1-glucoside bond between two α-glucose units.

Aspects of present invention are the following. Loperamide for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Amiodarone for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Niguldipine for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Pimozide for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Nicardipine for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Penitrem A for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Fluspirilene for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Trifluoperazine for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Trehalose for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. A particular aspect of present invention features a kit which includes: (i) a pharmaceutical composition comprising an autophagy inducing compound and (ii) instructions for administering the composition to a subject for the treatment of the bone degenerative disorder. Moreover in a particular embodiment of this aspect, the autophagy inducing compound is administered to reach plasma concentrations of (about) 0.1 nM to (about) 150 nM. In a preferred embodiment of this aspect, the autophagy inducing compound is administered at a dosis to reach plasma concentration of (about) 3.0 nM to (about) 9.0 nM.
Present invention also concerns a pharmaceutical composition comprising an autophagy inducing compound in an amount effective for treating a bone degenerative disorder, wherein said compound is selected from the group consisting of: (a) Loperamide; (b) Amiodarone; (c) Niguldipine; (d) Pimozide; (e) Nicardipine; (f) Penitrem A; (g) Fluspirilene, (h) Trifluoperazine and (i) trehalose or present invention concerns a pharmaceutical composition comprising an autophagy inducing compound, wherein said compound is selected from the group consisting of: (a) Loperamide; (b) Amiodarone; (c) Niguldipine; (d) Pimozide; (e) Nicardipine; (f) Penitrem A; (g) Fluspirilene, (h) Trifluoperazine and (i) trehalose for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Such pharmaceutical composition can be as described here above, wherein said composition further comprises a pharmaceutically acceptable carrier. The bone degenerative disorder can be to inhibit increased bone resorption disorder, to inhibit osteoclast activity, to inhibit formation of osteoclasts (TRAP-positive multinuclear cells) from blood progenitor cells, or to activate bone formation. The bone degenerative disorders, can be bone degenerative disorder caused by acute critical illness or prolonged critical illness, or the bone degenerative disorder is caused by (parenteral) nutrition-induced suppression of autophagy, or the bone degenerative disorder is caused by mTOR activation, or the bone degenerative disorders is hyperresorption of bone, or the bone degenerative disorders is an imbalance in the regulation of bone resorption and bone formation resulting in metabolic bone diseases such as osteoporosis, or the bone degenerative disorders is elderly osteoporosis, or the bone degenerative disorders is osteoporosis or the bone degenerative disorders is pathological increased bone resorption and pathological decreased bone formation, or the bone degenerative disorders is disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness, or the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness, or the bone degenerative disorders is increased osteoclast formation from circulating precursors leading to osteoporosis, or the bone degenerative disorders is increased osteoclast formation due to a increased cytokine production disorder, or the bone degenerative disorders is increased osteoclast formation due to hormonal disorder, or the bone degenerative disorders is increased osteoclast formation due to VitaminD deficiency, or the bone degenerative disorders is increased osteoclast formation due to glucocorticoids (pre)treatment, or the bone degenerative disorders is increased osteoclast formation due to heparinoids (pre)treatment, or the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder, or the bone degenerative disorders is osteoclast activity increase due to acute critical illness and/or prolonged critical
illness, or the bone degenerative disorders is osteoclast activity increase due to increased cytokine production disorder or the bone degenerative disorders is osteoclast activity increase due to hormonal disorder.

Present invention also concerns a pharmaceutical composition comprising an autophagy inducing compound in an amount effective for treating a bone degenerative disorder, wherein said compound is selected from the group consisting of: (a) Loperamide; (b) Amiodarone; (c) Niguldipine; (d) Pimozide; (e) Nicardipine; (f) Penitrem A; (g) Fluspirilene, (h) Trifluoperazine and (i) trehalose or present invention concerns a pharmaceutical composition comprising an autophagy inducing compound, wherein said compound is selected from the group consisting of: (a) Loperamide; (b) Amiodarone; (c) Niguldipine; (d) Pimozide; (e) Nicardipine; (f) Penitrem A; (g) Fluspirilene, (h) Trifluoperazine and (i) trehalose for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject. Such pharmaceutical composition can be as described here above, wherein said composition further comprises a pharmaceutically acceptable carrier. The bone degenerative disorder can be to inhibit increased bone resorption disorder, to inhibit osteoclast activity, to inhibit formation of osteoclasts (TRAP-positive multinuclear cells) from blood progenitor cells, to activate bone formation. The bone degenerative disorders is bone metabolism disorder, can be bone degenerative disorder caused by acute critical illness or prolonged critical illness, or the bone degenerative disorder is caused by (parenteral) nutrition-induced suppression of autophagy or the bone degenerative disorder is caused by mTOR activation, or the bone degenerative disorders is hyperresorption of bone or the bone degenerative disorders is an imbalance in the regulation of bone resorption and bone formation resulting in metabolic bone diseases such as osteoporosis or the bone degenerative disorders is elderly osteoporosis or the bone degenerative disorders is osteoporosis or the bone degenerative disorders is pathological increased bone resorption and pathological decreased bone formation, or the bone degenerative disorders is disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness, or the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness, or the bone degenerative disorders is increased osteoclast formation from circulating precursors leading to osteoporosis, or the bone degenerative disorders is increased osteoclast formation due to a increased cytokine production disorder, or the bone degenerative disorders is increased osteoclast formation due to hormonal disorder, or the bone degenerative disorders is increased osteoclast formation due to VitaminD deficiency, or the bone degenerative
disorders is increased osteoclast formation due to glucocorticoids (pre)treatment, or the bone degenerative disorders is increased osteoclast formation due to heparinoids (pre)treatment, or the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder, or the bone degenerative disorders is osteoclast activity increase due to acute critical illness and/or prolonged critical illness, or the bone degenerative disorders is osteoclast activity increase due to increased cytokine production disorder or the bone degenerative disorders is osteoclast activity increase due to hormonal disorder.

Present invention also concerns a pharmaceutical composition for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject said pharmaceutical composition comprising an autophagy inducing compound in an amount effective for treating a bone degenerative disorder, wherein said the compound is at least one compound selected from the group consisting of:

(a) compounds of formula (I):

wherein X is selected from CR4R5 and NR6; R1 is selected from hydrogen, C1-6 alkyl and phenyl, wherein the alkyl and phenyl are substituted with 0 or 1 halogen; R2 is selected from hydrogen, C1-6 alkyl and phenyl, wherein the alkyl and phenyl are substituted with 0 or 1 halogen; R3 is selected from hydrogen, hydroxyl, C1-6 alkyl and phenyl; R4 is selected from hydrogen,
R5 is selected from CI-6 alkyl and phenyl, halophenyl, benzimidazole, dihydrobenzimidazole, benzimidazolone; optionally R4 and R5 are taken together to form a 5 or 6 membered heterocycloalkyl comprising two nitrogen atoms, wherein the heterocycloalkyl is substituted with 1, 2 or 3 substituents selected from the group consisting of CI-6 alkyl, phenyl, and =0; R6 is selected from hydrogen and CI-6 alkyl; R7a, R8a, R9a, RIOa, R11a, R7b, R8b, R9b, RIOb, and R11b are each independently selected from hydrogen, hydroxyl, halogen and CI-6 haloalkyl; optionally R11a and R11b are taken together to form a heterocycle of the following structure:

wherein R11 is selected from CH2, NH, O and S; R12 and R13 are each independently selected from hydrogen and CI-6 alkyl; R14a and R14b are each independently selected from hydrogen and CI-6 alkyl; R15 is selected from phenyl substituted with 0 or 1 halogen or nitro; R16 is selected from hydrogen and CI-6 alkyl; Y is N or CH; and pharmaceutically acceptable salts thereof; (b) compounds of formula (II):

wherein R17 is selected from hydrogen and CI-6 alkyl; R18a and R18b, are each independently selected from hydrogen and CI-6 alkyl; R19a, R19b, R20a, R20b, and R21 are each independently selected from hydrogen, halogen and nitro; R22 is selected from hydrogen and CI-6 alkyl; R23 is selected from —(CH2)nNR24aR24b and —(CH2)nR24a; R24a and R24b are each independently selected from CI-6 alkyl and phenyl, wherein the alkyl is substituted with 0 or 1 phenyl substituents; optionally R24a and R24b are taken together with
the nitrogen to which they are attached to form a piperidine which is substituted with 0, 1 or 2 phenyl substituents; n is a positive integer from 2 to 4; and pharmaceutically acceptable salts thereof.

(c) compounds of formula (III):

wherein R25 is selected from hydrogen and C1-6 alkyl; R26a, R26b, R27a, and R27b are each independently selected from hydrogen, halogen and C1-6 alkyl; R28 is selected from —O(CH2)mNR29aR29b and —NH(CH2)mNR29aR29b; R29a and R29b are each independently selected from hydrogen and C1-6 alkyl; Z is O, S or NH; m is a positive integer from 1 to 3; and pharmaceutically acceptable salts thereof.

(d) compounds of formula (IV):

wherein R30 is selected from hydrogen, C1-6 alkyl and halogen; R31a and R31b are each independently selected from hydrogen, hydroxyl and C1-6 alkyl; R32 is selected from hydrogen, hydroxyl and C1-6 alkyl; R33 and R34 are each independently selected from hydrogen and C1-6 alkyl; R35a and R35b are each independently selected from hydrogen, hydroxyl and C1-6 alkyl; R36a and R36b are each independently selected from hydrogen, hydroxyl and C1-6 alkyl; R37a and R37b are each independently selected from hydrogen, hydroxyl and C1-6 alkyl; R38 is selected from hydrogen, hydroxyl and C1-6 alkyl; optionally
R37a and R38 are taken together to form a three membered heterocycle of the formula:

wherein R38' is O, S or NH; R39a is selected from hydrogen, hydroxyl and C1-6 alkyl; R39b is selected from hydrogen, hydroxyl, C1-6 alkyl and C2-6 alkenyl; U, V and W are each independently selected from O, S, and NH; and pharmaceutically acceptable salts thereof.

Suitable Rapamycin analogues for present invention are of the group of the Rapamycin derivatives such as Everolimus, Temsirolimus, 40-O-(2-hydroxyethyl)-rapamycin, and/or 32-deoxorapamycin, and/or 16-pent-2-ynyloxy-32-deoxorapamycin, and/or 16-pent-2-ynyloxy-32 (S or R) -dihydro-rapamycin, and/or 16-pent-2-ynyloxy-32 (S or R)-dihydro-40-O-(2-hydroxyethyl)-rapamycin, and/or 40-[3-hydroxy-2-(hydroxy-methyl)-2-methylpropanoate]-rapamycin (also known as CCI779) and/or 40-epi-(tetrazolyl)-rapamycin (also known as ABT578), and/or the so-called rapalogs, e.g. as disclosed in WO9802441, WO014387 and WO0364383, AP23573, AP23464; AP23675 or AP23841, e.g. AP23573, and/or compounds disclosed under the name Tafa-93, and/or compounds disclosed under the name biolimus. Other for present invention preferred rapamycin derivative is selected from the group consisting of 40-O-(2-hydroxyethyl)-rapamycin (also known as everolimus), and/or 32-deoxorapamycin, and/or 16-pent-2-ynyloxy-32-deoxorapamycin, and/or 16-pent-2-ynyloxy-32 (S or R) -dihydro-rapamycin, and/or 16-pent-2-ynyloxy-32 (S or R)-dihydro-40-O-(2-hydroxyethyl)-rapamycin, and/or 40-[3-hydroxy-2-(hydroxy-methyl)-2-methylpropanoate]-rapamycin (also known as CCI779 or temsirolimus) and/or 40-epi-(tetrazolyl)-rapamycin (also known as ABT578), and/or AP23573, such as 40-O-(2-hydroxyethyl)-rapamycin.

Suitable phenothiazine derivatives for present invention are for example compounds chosen from quinacrine, imipramine, carbamazepine, phenazine, phenothiazine, promazine,
Another mTOR dependent autophagy inducer suitable for the medicament and treatment of present invention is Resveratrol and derivatives thereof as described in WO2005102298 or WO2007096078. Resveratrol is an autophagy induced via the mTOR-Rictor survival pathway. Resveratrol at lower doses induces autophagy while higher doses are known to attenuate autophagy. The activation of mammalian target of rapamycin (mTOR) is differentially regulated by low-dose resveratrol, i.e. the phosphorylation of mTOR at serine 2448 is inhibited, whereas the phosphorylation of mTOR at serine 2481 is increased, which is attenuated with a higher dose of resveratrol. Low-dose resveratrol significantly induces the expression of Rictor, a component of mTOR complex 2, and activated its downstream survival kinase Akt (Ser 473).

The term "resveratrol, a derivative, metabolite or analogue thereof concerns compounds encompassed by the general formula I

![Chemical Structure][1]

wherein A denotes a carbon-carbon single or double bound, and the latter hereby may be trans or cis, and R1, R2, R3, R4, R5 and R6, independently from each other denote hydrogen, hydroxyl, etherified hydroxyl or esterified hydroxy groups.

Another mTOR dependent autophagy inducer suitable for the medicament and treatment of present invention is fenofibrate. Fenofibrate functions as autophagy activator by activating AMPK. AMPK switches on p53-dependent cell cycle metabolic check point and autophagy. AMPK antagonizes this Akt-induced mTOR activation. AMPK-dependent protein kinase (AMPK) plays an integral role in the response to starvation by sensing the rise in AMP/ATP ratio and switching off the ATP-consuming anabolic processes, such as protein and lipid synthesis or DNA replication. AMPK can induce several rescue pathways, which enhance cell survival during glucose deprivation (Figures 1 and 2). One of them includes p53-dependent check point, which promotes autophagy.

chloropromazine, haloperidol, clozapine, 2-chlorophenothiazine, promethazine (10-(2dimethylaminopropyl)phenothiazine), chlorprothixen, and acepromazine.
Other compounds suitable for the medicament and treatment of present invention are DNA methylation inhibitors and in particular the DNA methylation inhibitor, 5-azacytidine.

Present invention also concerns an autophagy inducing compound for use in a treatment of treating or preventing bone degenerative disorder in a mammalian, wherein said the compound is at least one compound selected from the group consisting of:

(a) compounds of formula (I):

![Chemical Structure](image)

wherein X is selected from CR4R5 and NR6; R1 is selected from hydrogen, CI-6 alkyl and phenyl, wherein the alkyl and phenyl are substituted with 0 or 1 halogen; R2 is selected from hydrogen, CI-6 alkyl and phenyl, wherein the alkyl and phenyl are substituted with 0 or 1 halogen; R3 is selected from hydrogen, hydroxyl, CI-6 alkyl and phenyl; R4 is selected from hydrogen, hydroxyl, CI-6 alkyl and phenyl; R5 is selected from CI-6 alkyl and phenyl, halophenyl, benzimidazole, dihydrobenzimidazole, benzimidazolone; optionally R4 and R5 are taken together to form a 5
or 6 membered heterocycloalkyl comprising two nitrogen atoms, wherein the heterocycloalkyl is substituted with 1, 2 or 3 substituents selected from the group consisting of C1-6 alkyl, phenyl, and =0;

R6 is selected from hydrogen and Cl-6 alkyl; R7a, R8a, R9a, R10a, R11a, R7b, R8b, R9b, R10b, and R11b are each independently selected from hydrogen, hydroxyl, halogen and Cl-6 haloalkyl; optionally R11a and R11b are taken together to form a heterocycle of the following structure:

wherein R11 is selected from CH2, NH, O and S; R12 and R13 are each independently selected from hydrogen and Cl-6 alkyl; R14a and R14b are each independently selected from hydrogen and Cl-6 alkyl; R15 is selected from phenyl substituted with 0 or 1 halogen or nitro; R16 is selected from hydrogen and Cl-6 alkyl; Y is N or CH; and pharmaceutically acceptable salts thereof;

(b) compounds of formula (II):
wherein R17 is selected from hydrogen and Cl-6 alkyl; R18a and R18b, are each independently selected from hydrogen and Cl-6 alkyl; R19a, R19b, R20a, R20b, and R21 are each independently selected from hydrogen, halogen and nitro; R22 is selected from hydrogen and Cl-6 alkyl; R23 is selected from -(CH2)nNR24aR24b and -(CH2)nR24a; R24a and R24b are each independently selected from Cl-6 alkyl and phenyl, wherein the alkyl is substituted with 0 or 1 phenyl substituents; optionally R24a and R24b are taken together with the nitrogen to which they are attached to form a piperidine which is substituted with 0, 1 or 2 phenyl substituents; n is a positive integer from 2 to 4; and pharmaceutically acceptable salts thereof

(c) compounds of formula (III):

wherein R25 is selected from hydrogen and Cl-6 alkyl; R26a, R26b, R27a, and R27b are each independently selected from hydrogen, halogen and Cl-6 alkyl; R28 is selected from -(CH2)mNR29aR29b and -NH(CH2)mNR29aR29b; R29a and R29b are each independently selected from hydrogen and Cl-6 alkyl; Z is O, S or NH; m is a positive integer from 1 to 3; and pharmaceutically acceptable salts thereof

(d) compounds of formula (IV):

wherein R30 is selected from hydrogen, Cl-6 alkyl and halogen; R31a and R31b are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R32 is selected from hydrogen, hydroxyl and Cl-6 alkyl; R33 and R34 are each independently selected from hydrogen and Cl-6 alkyl; R35a and R35b are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R36a and R36b are each independently selected from hydrogen,
hydroxyl and CI-6 alkyl; R37a and R37b are each independently selected from hydrogen, hydroxyl and CI-6 alkyl; R38 is selected from hydrogen, hydroxyl and CI-6 alkyl; optionally R37a and R38 are taken together to form a three membered heterocycle of the formula:

wherein R38' is O, S or NH; R39a is selected from hydrogen, hydroxyl and CI-6 alkyl; R39b is selected from hydrogen, hydroxyl, Cl-6 alkyl and C2-6 alkenyl; U, V and W are each independently selected from O, S, and NH; and pharmaceutically acceptable salts thereof.

The bone degenerative disorder can be to inhibit increased bone resorption disorder, to inhibit osteoclast activity, to inhibit formation of osteoclasts (TRAP-positive multinuclear cells) from blood progenitor cells, to activate bone formation. The bone degenerative disorders is bone metabolism disorder, can be bone degenerative disorder caused by acute critical illness or prolonged critical illness, or the bone degenerative disorder is caused by (parenteral) nutrition-induced suppression of autophagy or the bone degenerative disorder is caused by mTOR activation, or the bone degenerative disorders is hyperresorption of bone or the bone degenerative disorders is an imbalance in the regulation of bone resorption and bone formation results in metabolic bone diseases such as osteoporosis or the bone degenerative disorders is elderly osteoporosis or the bone degenerative disorders is osteoporosis or the bone degenerative disorders is pathological increased bone resorption and pathological decreased bone formation, or the bone degenerative disorders is disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness, or the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness, or the bone degenerative disorders is increased osteoclast formation from circulating precursors leading to osteoporosis, or the bone degenerative disorders is increased osteoclast formation due to a increased cytokine production disorder, or the bone degenerative disorders is increased osteoclast formation due to hormonal disorder, or the bone degenerative disorders is increased osteoclast formation due to VitaminD deficiency, or the bone degenerative disorders is increased osteoclast formation due to glucocortoids
(pre)treatment, or the bone degenerative disorders is increased osteoclast formation due to heparinoids (pre)treatment, or the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder, or the bone degenerative disorders is osteoclast activity increase due to acute critical illness and/or prolonged critical illness, or the bone degenerative disorders is osteoclast activity increase due to increased cytokine production disorder or the bone degenerative disorders is osteoclast activity increase due to hormonal disorder.

The invention also concerns such compound of present invention, described in the embodiments above, for use in a treatment to activate bone formation in a subject, for use in a treatment to treat a bone metabolism disorder, for use in a treatment to treat a bone metabolism disorder caused by acute critical illness or prolonged critical illness, for use in a treatment to treat a bone metabolism disorder caused by (parenteral) nutrition-induced suppression of autophagy, for use in a treatment to treat a bone metabolism disorder caused by mTOR activation, for use in a treatment to treat hyperresorption of bone, for use in a treatment to treat an imbalance in the regulation of bone resorption and bone formation resulting in metabolic bone diseases such as osteoporosis, for use in a treatment to treat elderly osteoporosis, for use in a treatment to treat osteoporosis, for use in a treatment of a patient with pathological increased bone resorption and pathological decreased bone formation, for use in a treatment of disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness, for use in a treatment of disorders of bone metabolism of enhanced osteoclast formation from circulating precursors in the blood during critical illness, for use in a treatment of increased osteoclast formation from circulating precursors leading to osteoporosis, for use in a treatment of increased osteoclast formation due to an increased cytokine production disorder, for use in a treatment of increased osteoclast formation due to hormonal disorder, for use in a treatment of increased osteoclast formation due to VitaminD deficiency, for use in a treatment of increased osteoclast formation due to glucocorticoids (pre)treatment, for use in a treatment of increased osteoclast formation due to heparinoids (pre)treatment, for use in a treatment of increased osteoclast formation due to autophagy deficiency disorder, for use in a treatment to inhibit osteoclast activity increase due to acute critical illness and/or prolonged critical illness, for use in a treatment to inhibit osteoclast activity increase due to increased cytokine production disorder, for use in a treatment to inhibit osteoclast activity increase due to hormonal disorder, for use in a treatment to inhibit osteoclast activity increase due to VitaminD deficiency, for use in a
treatment to inhibit osteoclast activity increase due to glucocorticoids (pre)treatment, for use in a treatment to inhibit osteoclast activity increase due to heparinoids (pre)treatment, for use in a treatment to inhibit osteoclast activity increase due to autophagy deficiency disorder or for use in a treatment to inhibit bone hyperresorption.

The present invention also relates to a pharmaceutical composition comprising a pharmacologically acceptable amount of an autophagy inducing or autophagy inducing compound or an autophagy inducing or autophagy inducing compound analog as described above, a pharmaceutically acceptable salt, solvate or isomer thereof, or combinations thereof. The invention also concerns the pharmaceutical for use in a treatment to activate bone formation in a subject, for use in a treatment to treat a bone metabolism disorder, for use in a treatment to treat a bone metabolism disorder caused by acute critical illness or prolonged critical illness, for use in a treatment to treat a bone metabolism disorder caused by (parenteral) nutrition-induced suppression of autophagy, for use in a treatment to treat a bone metabolism disorder caused by mTOR activation, for use in a treatment to treat hyperresorption of bone, for use in a treatment to treat an imbalance in the regulation of bone resorption and bone formation resulting in metabolic bone diseases such as osteoporosis, for use in a treatment to treat elderly osteoporosis, for use in a treatment to treat osteoporosis, for use in a treatment of a patient with pathological increased bone resorption and pathological decreased bone formation, for use in a treatment of disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness, for use in a treatment of disorders of bone metabolism of enhanced osteoclast formation from circulating precursors in the blood during critical illness, for use in a treatment of increased osteoclast formation from circulating precursors leading to osteoporosis, for use in a treatment of increased osteoclast formation due to a increased cytokine production disorder, for use in a treatment of increased osteoclast formation due to hormonal disorder, for use in a treatment of increased osteoclast formation due to Vitamin D deficiency, for use in a treatment of increased osteoclast formation due to glucocorticoids (pre)treatment, for use in a treatment of increased osteoclast formation due to heparinoids (pre)treatment, for use in a treatment of increased osteoclast formation due to autophagy deficiency disorder, for use in a treatment to inhibit osteoclast activity increase due to acute critical illness and/or prolonged critical illness, for use in a treatment to inhibit osteoclast activity increase due to increased cytokine production disorder, for use in a treatment to inhibit osteoclast activity increase due to hormonal disorder, for use in a treatment to inhibit osteoclast activity increase due to Vitamin D deficiency, for use in a
treatment to inhibit osteoclast activity increase due to glucocorticoids (pre)treatment, for use in a treatment to inhibit osteoclast activity increase due to heparinoids (pre)treatment, for use in a treatment to inhibit osteoclast activity increase due to autophagy deficiency disorder or for use in a treatment to inhibit bone hyperresorption.

5 A further embodiment of present invention concerns a DNA methylation inhibitor for use in a method of inhibiting for suppressing osteoclastogenesis or osteoclast differentiation in a subject in need for a treatment of a disorders of bone density (ICD-10-CM M80-M85). This DNA methylation inhibitor can be for use in a method of inhibiting for suppressing enhanced osteoclastogenesis or increased osteoclast differentiation in a subject in need for a treatment of a disorders of bone density (ICD-10-CM M80-M85). Moreover this DNA methylation inhibitor can be for use in a method of inhibiting for suppressing critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis or increased osteoclast differentiation in a subject in need for a treatment of Critical Illness Related Metabolic Bone Disease or of critical illness induced Osteopenia [ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851] secondary to ICU Admission. Suitable methylation inhibitor according to this invention is for instance a DNA methylation inhibitor selected from the group consisting of decitabine (or 5-aza-2’-deoxycytidine or ) 5-azadC, azacitidine (or 5-azacytidine), vorinostat (Zolinza™), procainamide and derivatives thereof. It was surprisingly found that DNA methylation inhibitor according to present invention can be delivered or administrated to subject in need thereof to reach very low nanomolar plasma concentration of a value of the range of 1 to 10 nM to suppress osteoclastogenesis or osteoclast differentiation.

25 When administered to a patient, an autophagy inducing or autophagy inducing compound or a methylation inhibitor is preferably administered as a component of a composition that optionally comprises a pharmaceutically acceptable carrier or vehicle. In one embodiment, these compositions are administered orally. In a preferred embodiment, the autophagy inducing or autophagy inducing compound of present invention is a component of a pharmaceutical composition that is administered intravenously.

30 A pharmaceutical composition comprising a autophagy inducing or autophagy inducing compound of present invention can be administered via one or more routes such as, but not limited to, oral, intravenous infusion, subcutaneous injection, intramuscular, topical, depo injection, implantation, time-release mode, and intracavitary. The
pharmaceutical composition is formulated to be compatible with its intended route of administration. Examples of routes of administration include parenteral, e.g., intravenous, intramuscular, intraperitoneal, intracapsular, intraspinal, intrasternal, intratumor, intranasal, epidural, intra-arterial, intraocular, intraorbital, intradermal, subcutaneous, oral (e.g., inhalation), transdermal (topical-particularly to the ears, nose, eyes, or skin), transmucosal (e.g., oral) nasal, rectal, intracerebral, intravaginal, sublingual, submucosal, and transdermal administration.

Administration can be via any route known to be effective by a physician of ordinary skill. Parenteral administration, i.e., not through the alimentary canal, can be performed by subcutaneous, intramuscular, intra-peritoneal, intratumoral, intradermal, intracapsular, intra-adipose, or intravenous injection of a dosage form into the body by means of a sterile syringe, optionally a pen-like syringe, or some other mechanical device such as an infusion pump. A further option is a composition that can be a powder or a liquid for the administration in the form of a nasal or pulmonary spray. As a still further option, the administration can be transdermally, e.g., from a patch. Compositions suitable for oral, buccal, rectal, or vaginal administration can also be provided. In a preferred embodiment, administration of the autophagy inducing or autophagy inducing compound compound of present invention is via an intravenous injection, e.g. an intravenous bolus injection or by gradual perfusion over time.

The autophagy inducing or autophagy inducing compound compound and the pharmaceutical composition of present invention can also be administered by a small bolus injection followed by a continuous infusion. One protocol for treatment with autophagy inducing or autophagy inducing compound or a autophagy inducing or autophagy inducing compound analog is as follows: (i) initial bolus injection over a period of 1-2 minutes; (ii) high level infusion for 1 hour; (2) low level maintenance infusion for 2-3 hours.

The whole of the dose of autophagy inducing or autophagy inducing compound required to achieve a protective effect could also be administered as one or more bolus injections e.g. ranging between 1-100percent of the estimated required 24h dose, or administered with a 50cc syringe at a rate of 2 ml per hour.

The autophagy inducing or autophagy inducing compound and the pharmaceutical composition of present invention can also be administered by a small bolus injection followed by a continuous infusion. One protocol for treatment with autophagy inducing or autophagy inducing compound or a autophagy inducing or autophagy inducing compound analog is as follows: (i) initial bolus injection over a period of 1-2 minutes; (ii) high level infusion for 1 hour; (2) low level maintenance infusion for 2-3 hours.
The whole of the dose of autophagy inducing or autophagy inducing compound required to achieve a protective effect could also be administered as one or more bolus injections, e.g. administered with a 50cc syringe at a rate of 2 ml over 1 hour.


In yet another embodiment, a controlled release system can be placed in proximity of the target. For example, a micropump can deliver controlled doses directly into bone or adipose tissue, thereby requiring only a fraction of the systemic dose (See e.g., Goodson, 1984, in Medical Applications of Controlled Release, vol. 2, pp. 115-138). In another example, a pharmaceutical composition of the invention can be formulated with a hydrogel (See, e.g., U.S. Pat. Nos. 5,702,717; 6,1 17,949; 6,201,072).

In one embodiment, it may be desirable to administer the pharmaceutical composition of the invention locally, i.e., to the area in need of treatment. Local administration can be achieved, for example, by local infusion during surgery, topical application (e.g., in conjunction with a wound dressing after surgery), injection, catheter, suppository, or implant. An implant can be of a porous, non-porous, or gelatinous material, including membranes, such as sialastic membranes, or fibres.

In certain embodiments, it may be desirable to introduce the autophagy inducing or autophagy inducing compound into the central nervous system by any suitable route, including intraventricular, intrathecal, and epidural injection. Intraventricular injection may be
facilitated by an intraventricular catheter, for example, attached to a reservoir, such as an Ommaya reservoir.

Pulmonary administration can also be employed, e.g., by use of an inhaler or nebulizer, and formulation with an aerosolizing agent, or via perfusion in a fluorocarbon or synthetic pulmonary surfactant.

The skilled artisan can appreciate the specific advantages and disadvantages to be considered in choosing a mode of administration. Multiple modes of administration are encompassed by the invention. For example, a autophagy inducing or autophagy inducing compound of the invention can be administered by subcutaneous injection, whereas another therapeutic agent can be administered by intravenous infusion. Moreover, administration of one or more species of autophagy inducing or autophagy inducing compound compounds, with or without other therapeutic agents, can occur simultaneously (i.e., co-administration) or sequentially. In another embodiment, the periods of administration of a autophagy inducing or autophagy inducing compound, with or without other therapeutic agents can overlap. For example a autophagy inducing or autophagy inducing compound can be administered for 7 days and another therapeutic agent can be introduced beginning on the fifth day of autophagy inducing or autophagy inducing compound treatment. Treatment with the other therapeutic agent can continue beyond the 7-day autophagy inducing or autophagy inducing compound treatment.

A pharmaceutical composition of a autophagy inducing or autophagy inducing compound can be administered before, during, and/or after the administration of one or more therapeutic agents. In one embodiment, autophagy inducing or autophagy inducing compound can first be administered to stimulate the expression of insulin, which increases sensitivity to subsequent challenge with a therapeutic agent. In another embodiment, autophagy inducing or autophagy inducing compound can be administered after administration of a therapeutic agent. In yet another embodiment, there can be a period of overlap between the administration of the autophagy inducing or autophagy inducing compound and the administration of one or more therapeutic agents.

A pharmaceutical composition of the invention can be administered in the morning, afternoon, evening, or diurnally. In one embodiment, the pharmaceutical composition is administered at particular phases of the circadian rhythm. In a specific embodiment, the pharmaceutical composition is administered in the morning. In another specific embodiment, the pharmaceutical composition is administered at an artificially induced circadian state.

The present pharmaceutical compositions can take the form of solutions, suspensions, emulsion, tablets, pills, pellets, capsules, capsules containing liquids, powders, sustained-
release formulations, suppositories, emulsions, aerosols, sprays, suspensions, or any other form suitable for use. In one embodiment, the pharmaceutically acceptable vehicle is a capsule (See e.g., U.S. 5,698,155).

Pharmaceutical compositions adapted for parenteral administration include, but are not limited to, aqueous and non-aqueous sterile injectable solutions or suspensions, which can contain antioxidants, buffers, bacteriostats and solutes. Other components that can be present in such pharmaceutical compositions include water, alcohols, polyols, glycerine and vegetable oils, for example. Compositions adapted for parenteral administration can be presented in unit-dose or multi-dose containers (e.g., sealed ampoules and vials), and can be stored in a freeze-dried (i.e., lyophilized) condition requiring the addition of a sterile liquid carrier (e.g., sterile saline solution for injections) immediately prior to use. Extemporaneous injection solutions and suspensions can be prepared from sterile powders, granules and tablets.

Pharmaceutical compositions adapted for transdermal administration can be provided as discrete patches intended to remain in intimate contact with the epidermis for a prolonged period of time. Pharmaceutical compositions adapted for topical administration can be provided as, for example, ointments, creams, suspensions, lotions, powders, solutions, pastes, gels, sprays, aerosols or oils. A topical ointment or cream is preferably used for topical administration to the skin, mouth, eye or other external tissues. When formulated in an ointment, the active ingredient can be employed with either a paraffinic or a water-miscible ointment base. Alternatively, the active ingredient can be formulated in a cream with an oil-in-water base or a water-in-oil base.

Pharmaceutical compositions adapted for topical administration to the eye include, for example, eye drops or injectable pharmaceutical compositions. In these pharmaceutical compositions, the active ingredient can be dissolved or suspended in a suitable carrier, which includes, for example, an aqueous solvent with or without carboxymethylcellulose.

Pharmaceutical compositions adapted for topical administration in the mouth include, for example, lozenges, pastilles and mouthwashes.

Pharmaceutical compositions adapted for nasal administration can comprise solid carriers such as powders (preferably having a particle size in the range of 20 to 500 microns). Powders can be administered in the manner in which snuff is taken, i.e., by rapid inhalation through the nose from a container of powder held close to the nose. Alternatively, pharmaceutical compositions adapted for nasal administration can comprise liquid carriers such as, for example, nasal sprays or nasal drops. These pharmaceutical compositions can comprise aqueous or oil solutions of an autophagy inducing or autophagy inducing compound.
Compositions for administration by inhalation can be supplied in specially adapted devices including, but not limited to, pressurized aerosols, nebulizers or insufflators, which can be constructed so as to provide predetermined dosages of the autophagy inducing or autophagy inducing compound.

Typically, pharmaceutical compositions for injection or intravenous administration are solutions in sterile aqueous buffers. Where necessary, the composition can also include a solubilizing agent and a local anesthetic such as lidocaine to ease pain at the site of the injection. Generally, the ingredients are supplied either separately or mixed together in unit dosage form, for example, as a dry lyophilized powder or water-free concentrate in a hermetically sealed container such as an ampoule or sachet indicating the quantity of active agent.

Where the composition is to be administered by infusion, it can be dispensed with an infusion bottle, bag, or other acceptable container, containing sterile pharmaceutical grade water, saline, or other acceptable diluents. Where the composition is administered by injection, an ampoule of sterile water for injection or saline can be provided so that the ingredients may be mixed prior to administration.

For a patient who cannot orally ingest a nutrient, it is essential to supply all nutrients such as an amino acid, a saccharide and an electrolyte through a vein. This way is called the total parenteral nutrition therapy, (TPN therapy) which can be provided by a TPN solution. Such TPN solutions are particularly suitable for critically ill patients for a therapy in the Intensive Care Unit. As a TPN solution employed in the TPN therapy, there has been known (1) a TPN solution containing a saccharide, an amino acid, a fat and an electrolyte (Japanese Unexamined Patent Publications No. 186822/1989, WO08503002 and EP-A-0 399 341), (2) an emulsion for injection comprising an amino acid and a fat (Japanese Unexamined Patent Publication No. 74637/1986), (3) a TPN solution comprising two separate infusions, one of which contains glucose and an electrolyte and the other of which contains an amino acid (Japanese Unexamined Patent Publications No. 52455/1982 and No. 103823/1986) and the like. In the TPN therapy, an infusion containing a high concentration of saccharide is usually administered to a patient.

As indicated the high nutritional content of such TPN solutions may lead to hyperglycemia and has been found to have a detrimental effect on the repair processes in critically ill patients by inhibition the autophagy process, which contributes to the removal of damaged organelles. Accordingly, a further aspect of the present invention relates to a TPN solution combined with a autophagy inducing or autophagy inducing compound of the present invention. This
combined composition is used to improve the condition of a critically ill patient or to reduce or treat multiple organ dysfunction syndrome in a critically ill patient.

Compositions for parenteral nutrition, in particular for intravenous administration are isotonic or hypertonic solutions (e.g. prepared by NaCl and/or dextrose or lactated Ringers) further comprising a saccharide such as glucose in a range between 10 and 20 % (w/v) to obtain a high nutritional content, and further comprising lipids and/or amino acids and/or added vitamins.

Compositions for parenteral administration comprise further to autophagy inducing or autophagy inducing compound of the present invention a saccharide such as glucose. Final glucose concentrations in a composition for administration are typically in the range from 10 to 20 % (w/v) e.g. 12.5 or 16 %.

Compositions for parenteral administration typically further comprise saturated, mono-unsaturated and essential poly-unsaturated fatty acids such as refined olive oil and/or soybean oil. Final lipid concentrations in a composition for administration are typically in the range of 2 to 6 % (w/v) e.g 4 %.

Compositions for parenteral administration typically further comprise one or more amino acids. Final amino acid concentrations are typically in the range from 2 to 6 % (w/v) e.g. 4 %.

Compositions for parenteral administration optionally further comprise trace elements such as one or more of Fe, Zn, Cu, Mn, F, Co, I, Se, Mo, Cr e.g. under the form of respectively the following salts ferrous gluconate, copper gluconate, manganese gluconate,zinc gluconate, sodium fluoride,cobalt II gluconate, sodium iodide,sodium selenite, ammonium molybdate and chromic chloride.

Compositions for parenteral administration optionally further comprise one or more vitamins such as Vitamin A (Retinol), Vitamin D3,Vitamin E (a tocopherol), Vitamin C, Vitamin B1 (thiamine), Vitamin B2 (riboflavin), Vitamin B6 (pyridoxine), Vitamin B12, Folic Acid, Pantothenic acid, Biotin, and Vitamin PP (niacin), e.g. under the form of Retinol palmitate, Colecalciferol, DL-a-tocopherol, Ascorbic acid, Cocarboxylase tetrahydrate, Riboflavin dihydrated sodium phosphate, Pyridoxine hydrochloride, Cyanocobalamin, Folic acid, Dexpantenol, D-Biotin and Nicotinamide.

Compositions for parenteral administration prior to administration can be isotonic solutions, or more particularly hypertonic solutions e.g. solutions with osmolality between 1000 and 1500, or between 1200 and 1500 mOsm/liter, eg. 1250 or 1500 mOsm/liter.
Compositions for parenteral administration can be provided as one solution comprising all constituents or as a kit of parts wherein different constituents are provided separately (saccharide, lipids, amino acids) and wherein the autophagy inducing or autophagy inducing compound is dissolved in one of the constituents or is provided separately. One or more of the different constituents may be provided in a dried form, which is redissolved prior to use. The compositions for parenteral nutrition in accordance with the present invention further comprise a autophagy inducing or autophagy inducing compound such as spermidine or spermine, autophagy inducing or autophagy inducing compound or putrescine in a concentration between 0.05, 0.1, 0.2 or 0.5 to 1, 2, 3 or 4 % (w/v).

The compositions for intravenous administration are typically packed in plastic bags with spike ports for delivery by intravenous drips.

In a specific embodiment, the present compositions contain spermine or autophagy inducing or autophagy inducing compound.

For patients which do not rely on parenteral food pharmaceutical compositions herein described can be provided in the form of oral tablets, capsules, elixirs, syrups and the like. Compositions for oral administration might require an enteric coating to protect the composition(s) from degradation within the gastrointestinal tract. In another example, the composition(s) can be administered in a liposomal formulation to shield the autophagy inducing or autophagy inducing compound compound disclosed herein from degradative enzymes, facilitate the molecule's transport in the circulatory system, and affect delivery of the molecule across cell membranes to intracellular sites.

A autophagy inducing or autophagy inducing compound compound intended for oral administration can be coated with or admixed with a material (e.g., glycercly monostearate or glycercly distearate) that delays disintegration or affects absorption of the autophagy inducing or autophagy inducing compound compound in the gastrointestinal tract. Thus, for example, the sustained release of a autophagy inducing or autophagy inducing compound compound can be achieved over many hours and, if necessary, the autophagy inducing or autophagy inducing compound compound can be protected from being degraded within the gastrointestinal tract. Taking advantage of the various pH and enzymatic conditions along the gastrointestinal tract, pharmaceutical compositions for oral administration can be formulated to facilitate release of a autophagy inducing or autophagy inducing compound compound at a particular gastrointestinal location.
Selectively permeable membranes surrounding an osmotically active driving compound are also suitable for orally administered compositions. Fluid from the environment surrounding the capsule is imbibed by the driving compound, which swells to displace the autophagy inducing or autophagy inducing compound through an aperture, can provide an essentially zero order delivery profile instead of the spiked profiles of immediate release formulations. A time delay material such as, but not limited to, glycerol monostearate or glycerol stearate can also be used.

Suitable pharmaceutical carriers also include starch, glucose, lactose, sucrose, gelatin, saline, gum acacia, talc, keratin, urea, malt, rice, flour, chalk, silica gel, sodium stearate, glycerol monostearate, talc, sodium chloride, dried skim milk, glycerol, propylene, glycol, water, and ethanol. If desired, the carrier, can also contain minor amounts of wetting or emulsifying agents, or pH buffering agents. In addition, auxiliary, stabilizing, thickening, lubricating, and colouring agents may be used. The composition can be formulated as a suppository, with traditional binders and carriers such as triglycerides.

For oral administration in the form of a tablet or capsule, the active drug component can be combined with an oral, non-toxic, pharmaceutically acceptable, inert carrier such as, but not limited to, lactose, starch, sucrose, glucose, methyl cellulose, magnesium stearate, dicalcium phosphate, calcium sulfate, mannitol, and sorbitol. For oral administration in liquid form, the oral drug components can be combined with any oral, non-toxic, pharmaceutically acceptable carrier such as, but not limited to, ethanol, glycerol, and water. Moreover, suitable binders, lubricants, disintegrating agents and colouring agents can also be incorporated into the mixture. Suitable binders include, but are not limited to, starch, gelatin, natural sugars (e.g., glucose, beta-lactose), corn sweeteners, natural and synthetic gums (e.g., acacia, tragacanth, sodium alginate), carboxymethylcellulose, polyethylene glycol, and waxes. Lubricants useful for an orally administered drug, include, but are not limited to, sodium oleate, sodium stearate, magnesium stearate, sodium benzoate, sodium acetate, and sodium chloride. Disintegrators include, but are not limited to, starch, methyl cellulose, agar, bentonite, and xanthan gum.

Pharmaceutical compositions adapted for oral administration can be provided, for example, as capsules or tablets; as powders or granules; as solutions, syrups or suspensions (in aqueous or non-aqueous liquids); as edible foams or whips; or as emulsions. For oral administration in the form of a tablet or capsule, the active drug component can be combined with an oral, non-toxic, pharmaceutically acceptable, inert carrier such as, but not limited to, lactose, starch, sucrose, glucose, methyl cellulose, magnesium stearate, dicalcium phosphate, magnesium
carbonate, stearic acid or salts thereof, calcium sulfate, mannitol, and sorbitol. For oral administration in the form of a soft gelatin capsule, the active drug component can be combined with an oral, non-toxic, pharmaceutically acceptable, inert carrier such as, but not limited to, vegetable oils, waxes, fats, semi-solid, and liquid polyols. For oral administration in liquid form, the oral drug components can be combined with any oral, non-toxic, pharmaceutically acceptable carrier such as, but not limited to, ethanol, glycerol, polyols, and water. Moreover, suitable binders, lubricants, disintegrating agents and colouring agents can also be incorporated into the mixture. Suitable binders include, but are not limited to, starch, gelatin, natural sugars (e.g. glucose, beta-lactose), corn sweeteners, natural and synthetic gums (e.g., acacia, tragacanth, sodium alginate), carboxymethylcellulose, polyethylene glycol, and waxes. Lubricants useful for an orally administered drug, include, but are not limited to, sodium oleate, sodium stearate, magnesium stearate, sodium benzoate, sodium acetate, and sodium chloride. Disintegrators include, but are not limited to, starch, methyl cellulose, agar, bentonite, and xanthan gum.

Orally administered compositions may contain one or more agents, for example, sweetening agents such as, but not limited to, fructose, aspartame and saccharin. Orally administered compositions may also contain flavouring agents such as, but not limited to, peppermint, oil of wintergreen, and cherry. Orally administered compositions may also contain colouring agents and/or preserving agents.

The autophagy inducing or autophagy inducing compound compounds of present invention can also be administered in the form of liposome delivery systems, such as small unilamellar vesicles, large unilamellar vesicles and multilamellar vesicles. Liposomes can be formed from a variety of phospholipids, such as cholesterol, stearylamine or phosphatidylcholines. A variety of cationic lipids can be used in accordance with the invention including, but not limited to, N-((2,3-dioleyloxy)propyl)-N,N,N-trimethylammonium chloride ("DOTMA") and diolesylphosphotidylethanolamine ("DOPE"). Such compositions suit the mode of administration.

The autophagy inducing or autophagy inducing compounds of the present invention can also be delivered by the use of monoclonal antibodies as individual carriers to which the compounds can be coupled. The compounds can also be coupled with soluble polymers as targetable drug carriers. Such polymers can include polyvinylpyrrolidone, pyran copolymer, polyhydroxypropylmethacrylamide-phenol, polyhydroxyethylaspartamid-phenol, or polyethyleneoxide-polylysine substituted with palmitoyl residues. Furthermore, the autophagy inducing or autophagy inducing compounds can be coupled to a class of biodegradable
polymers useful in achieving controlled release of a drug, for example, polylactic acid, polyglycolic acid, copolymers of polylactic and polyglycolic acid, poly epsilon caprolactone, polyhydroxy butyric acid, polyorthoesters, polyacetals, polydihydopyrans, polycyanoacrylates and cross linked or amphipathic block copolymers of hydrogels.

5 Pharmaceutical compositions adapted for rectal administration can be provided as suppositories or enemas. Pharmaceutical compositions adapted for vaginal administration can be provided, for example, as pessaries, tampons, creams, gels, pastes, foams or spray formulations.

Suppositories generally contain active ingredients in the range of 0.5% to 10% by weight. Oral formulations preferably contain 10% to 95% active ingredient by weight. In a preferred embodiment, the composition is formulated in accordance with routine procedures as a pharmaceutical composition adapted for intratumoral injection, implantation, subcutaneous injection, or intravenous administration to humans.

Conveniently, the blood autophagy inducing or autophagy inducing compound level is kept within the ranges mentioned in connection with the present invention for as long a period of time as the patient is critically ill. Hence, as a general rule, the blood autophagy inducing or autophagy inducing compound level is kept within the ranges mentioned in connection with the present invention as long as the patient is critically ill. Consequently, the blood autophagy inducing or autophagy inducing compound level is usually kept within the ranges mentioned in connection with the present invention for a period of time of more than about 8 hours, preferably more than about 24 hours, even more preferred more than about 2 days, especially more than about 4 days, and even more than about 7 days. In certain cases, it may even be preferred that the blood autophagy inducing or autophagy inducing compound level is kept within the ranges mentioned in connection with the present invention after the patient (previously) considered as being critically ill has been transferred from the Intensive Care Unit to another part of the hospital or even after the patient has left the hospital.

A critically ill patient, optionally entering an ICU, may be fed continuously, on admission with mainly intravenous glucose (for example, about 200 g to about 300 g per 24 hours) and from the next day onward with a standardized feeding schedule aiming for a caloric content up to between about 10 and about 40, preferably between about 20 and about 30, non-protein Calories/kg/24 hours and a balanced composition (for example, between about 0.05 and about 0.4, preferably between about 0.13 and about 0.26, g nitrogen/kg/24 hours and between about 20% and about 40% of non-protein Calories as lipids) of either total parenteral, combined
parenteral/enteral or full enteral feeding, the latter mode attempted as early as possible. Other concomitant ICU therapy can be left to the discretion of attending physicians. Alternatively, the following procedure can be used or it is possible to use a combination or variant of these procedures, as the physician considers advantageous for the patient:

A critically ill patient may be fed, on the admission day, using, for example, a 20% glucose infusion and from day 2 onward by using a standardized feeding schedule consisting of normal caloric intake (for example, about 25-35 Calories/kgBW/24 h) and balanced composition (for example, about 20%-40% of the non-protein Calories as lipids and about 1-2 g/kgBW/24h protein and about 0.01-100 mg/kg BW/24h autophagy inducing or autophagy inducing compound) of either total parenteral, combined parenteral/enteral or full enteral feeding, the route of administration of feeding depending on assessment of feasibility of early enteral feeding by the attending physician. All other treatments, including feeding regimens, were according to standing orders currently applied within the ICU.

The autophagy inducing or autophagy inducing compound and optionally another therapeutic agent are administered at an effective dose. The dosing and regimen most appropriate for patient treatment will vary with the disease or condition to be treated, and in accordance with the patient's weight and with other parameters.

An effective dosage and treatment protocol can be determined by conventional means, comprising the steps of starting with a low dose in laboratory animals, increasing the dosage while monitoring the effects (e.g., histology, disease activity scores), and systematically varying the dosage regimen. Several factors may be taken into consideration by a clinician when determining an optimal dosage for a given patient. Additional factors include, but are not limited to, the size of the patient, the age of the patient, the general condition of the patient, the particular disease being treated, the severity of the disease, the presence of other drugs in the patient, and the in vivo activity of the autophagy inducing or autophagy inducing compound.

A typical effective human dose of a autophagy inducing or autophagy inducing compound would be from about 1 µg/kg body weight/day to about 100 mg/kg/day, preferably from about 5 µg/kg/day to about 50 mg/kg/day, and most preferably about 10 µg/kg/day to 20 mg/kg/day.

As analogues of the autophagy inducing or autophagy inducing compound disclosed herein can be 2 to 100 times more potent than naturally occurring counterparts, a typical effective dose of such an analog can be lower, for example, from about 10 ng/kg body weight/day to 1 mg/kg/day, preferably 1 µg/kg/day to 900 µg/kg/day, and even more preferably 2 µg/kg/day to 250 µg/kg/day.
In another embodiment, the effective dose of a autophagy inducing or autophagy inducing compound of present is less than 1 μg/kg/day. In yet another embodiment the effective dose of a autophagy inducing or autophagy inducing compound of present is greater than 100 mg/kg/day.

The specific dosage for a particular patient, of course, has to be adjusted to the degree of response, the route of administration, the patient's weight, and the patient's general condition, and is finally dependent upon the judgment of the treating physician. Especially the highly critical condition of ICU patients requires a specific dosage and dosage regime.

It is understandable that the ideal dosage per serving to have the health effect will have to vary according the body weight of the subject who consumes the oral ingestible dosage form which comprises the autophagy inducing or autophagy inducing compound of present invention. A beneficial effect can be obtained in a subject with about 50 kg body weight by an orally ingestible dosage form comprising between 0.05 mg and 5 gram, preferably 0.25 mg to 2 gram, more preferably between 0.5 mg and 1.5 gram, more preferably between 1 mg and 750 mg of the autophagy inducing or autophagy inducing compound of present invention per administration (as demonstrated in Table 1).

Table 1. Possible amount of the autophagy inducing or autophagy inducing compound active ingredient of present invention per serving by a subject (BW: body weight).

<table>
<thead>
<tr>
<th>Dose mg/kg</th>
<th>50 mg</th>
<th>60 mg</th>
<th>70 mg</th>
<th>80 mg</th>
<th>90 mg</th>
<th>100 mg</th>
<th>110 mg</th>
<th>120 mg</th>
<th>130 mg</th>
<th>140 mg</th>
</tr>
</thead>
<tbody>
<tr>
<td>0,1</td>
<td>5</td>
<td>6</td>
<td>7</td>
<td>8</td>
<td>9</td>
<td>10</td>
<td>11</td>
<td>12</td>
<td>13</td>
<td>14</td>
</tr>
<tr>
<td>0,2</td>
<td>10</td>
<td>12</td>
<td>14</td>
<td>16</td>
<td>18</td>
<td>20</td>
<td>22</td>
<td>24</td>
<td>26</td>
<td>28</td>
</tr>
<tr>
<td>0,3</td>
<td>15</td>
<td>18</td>
<td>21</td>
<td>24</td>
<td>27</td>
<td>30</td>
<td>33</td>
<td>36</td>
<td>39</td>
<td>42</td>
</tr>
<tr>
<td>0,4</td>
<td>20</td>
<td>24</td>
<td>28</td>
<td>32</td>
<td>36</td>
<td>40</td>
<td>44</td>
<td>48</td>
<td>52</td>
<td>56</td>
</tr>
<tr>
<td>0,5</td>
<td>25</td>
<td>30</td>
<td>35</td>
<td>40</td>
<td>45</td>
<td>50</td>
<td>55</td>
<td>60</td>
<td>65</td>
<td>70</td>
</tr>
<tr>
<td>1</td>
<td>50</td>
<td>60</td>
<td>70</td>
<td>80</td>
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<td>100</td>
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<td>120</td>
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<td>5</td>
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<td>300</td>
<td>350</td>
<td>400</td>
<td>450</td>
<td>500</td>
<td>550</td>
<td>600</td>
<td>650</td>
<td>700</td>
</tr>
</tbody>
</table>
A beneficial effect can also be obtained in a subject with about 50 kg body weight as part of a TPN therapy comprising between 0,05 mg and 2,5 gram, preferably 0,5 mg to 2 gram, more preferably between 1 mg and 1,5 gram, more preferably between 2 mg and 750 mg of the autophagy inducing or autophagy inducing compound of present invention per administration.

Another aspect of present invention concerns to treat by an autophagy activator or enhancer a disorder of bone metabolism in a subject for instance to reverse a disorder of decrease in bone formation and an increase in bone resorption. Such disorder can be caused by critical illness. Particular aspects and embodiments of present invention are the following. An autophagy activator or autophagy inducing compound for use as a medicine to inhibit increased bone resorption. An autophagy activator or autophagy inducing compound for use as a medicine to inhibit osteoclast activity. An autophagy activator or autophagy inducing compound for use as a medicine to inhibit the formation of osteoclast (TRAP positive multinuclear cells) from blood progenitor cells. An autophagy activator or autophagy inducing compound for use as a medicine to activate bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is a bone metabolism disorder. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is caused by acute critical illness or prolonged critical illness. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is caused by (parenteral) nutrition induced suppression of autophagy. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder which is caused by mTOR activation. And/or an autophagy activator or autophagy inducing...
compound for use as a medicine to treat a bone degenerative disorder which is hyperresorption of bone.

An embodiment of present invention concerns an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is pathological increased bone resorption and pathological decreased bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is a disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness. Another embodiment of present invention concerns an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder caused by enhanced osteoclast formation from circular precursors in the blood during critical illness. Yet an embodiment of present invention concerns an autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is pathological increased bone resorption and pathological decreased bone formation. An autophagy activator or autophagy inducing compound for use as a medicine to treat a bone degenerative disorder that is increased osteoclast formation from circulating precursor leading to osteoporosis.

There are autophagy-inducing compounds for the above embodiments of present invention. For instance the macrolide autophagy inducing compounds. The macrolides are a group of drugs (typically antibiotics) whose activity stems from the presence of a macrolide ring, a large macrocyclic lactone ring to which one or more deoxy sugars, usually cladinose and desosamine, may be attached. The lactone rings are usually 14-, 15-, or 16-membered. Macrolides belong to the polyketide class of natural products. There are several macrolides autophagy inducers for instance the Azithromycin macrolide (an azalide, a subclass of macrolide antibiotics, IUPAC name: 2R,3S,4R,5R,8R,10R,1 IR,12S,13S,14R)-2-ethyl-3,4,10-trihydroxy-3,5,6,8,10,12,14-heptamethyl-15-oxo-11-([3,4,6-trideoxy-3-(dimethylamino)-P-D-xylo-[oxy]-l-oxa-6-azacyclopentadec-13-yl 2,6-dideoxy-3-C-methyl-3-0-methyl-a-L-ribo-hexopyranoside) is an autophagy-inducing compound in a dose-dependent manner in a reversible manner and this effect is removable by an autophagy inhibitor such as 3-methyladenine or LY294002 (Stamatiou R et al. EUROPEAN RESPIRATORY JOURNAL Volume: 34 Issue: 3 Pages: 721-730 Published: SEP 2009 )
Other suitable autophagy inducing compounds that are available for use in a medicament as described above are the calpain inhibitors (calpain 1 or calpain 2 inhibitors) for instance of the group consisting of calpastatin, ALLM, calpeptin, leupeptin, α-dicarbonyls, quinolinecarboxamides, sulfonium methyl ketones, diazomethyl ketones, Leu-Abu-CONHEt (AK275), 27-mer calpastatin peptide, Cbz-Val-Phe-H (MDL28170), calpeptin (Z-Leu-Nle-H), α-mercaptoacrylic acids, phosphorus derivatives, epoxysuccinates, acyloxymethyl ketones, halomethylketones and E64 have been demonstrated to induce autophagy (WO2007003941) and are suitable in the manufacture of a medicament for increasing autophagy in an individual. IMPase inhibitors for instance of the group consisting of L-690330, lithium, valproate, carbamazepine and salts, analogues and derivatives thereof have been demonstrated to induce autophagy (WO2006079792) and are suitable in the manufacture of a medicament for increasing autophagy in an individual. Compounds or agents that inhibit or reduce the activity of the cAMP/EPAC/PLC for instance of the group consisting of clonidine, rilmenidine, tyramine, morphine, baclofen, mastoparan, propanolol, bupivacain, N-dodecyl lysinamide, Gsa and/or PACAP are suramin, NF449, NF503 minoxidil, pinacidil, cromakalim or an analog or derivative thereof and have been demonstrated to induce autophagy (WO2008099175) and are suitable in the manufacture of a medicament for increasing autophagy in an individual. A dose of glutamine (Gin) such that it induces low millimolar concentrations (for instance, 1-10 mM, preferably 2-4 mM) of ammonium (Gln-derived ammonia) after Gin deamination in mitochondria stimulates autophagy. Christina H. Eng et al; Autophagy 6:7, 968-970; October 1, 2010; © 2010 Landes Bioscience. This can for instance be reached by feeding a patient parenterally with a parenteral nutrition comprising water having dissolved therein from about 1 to 150 mMoles/l of glutamine.

Trehalose (α,α-Trehalose; α-D-glucopyranosyl-(1→1)-α-D-glucopyranoside with Molar mass 342.296 g/mol (anhydrous) and 378.33 g/mol (dihydrate)) for instance D-(+)-trehalose dehydrate is an autophagy inducing agent (Casarejos MJ et al Conference Information: 14th International Congress of Parkinsons Disease and Movement Disorders Buenos Aires, ARGENTINA, JUN 13-17, 2010 Casarejos yr:2010 vol:25 iss:7 pg:S417 -S417; Catarina Gomes et al. Mutant superoxide dismutase 1 overexpression in NSC-34 cells: Effect of trehalose on aggregation, TDP-43 localization and levels of co-expressed glycoproteins Neuroscience Letters 475 (2010) 145-149 and Jos). This effects can be reached in vivo (e A. Rodriguez-Navarro et al. Neurobiology of Disease 39 (2010) 423-438). For the use of
present invention as described in the embodiments here above the dose is such to reach 1 to 100 mM, preferably 2 - 50 mM or more preferably 5 to 20 mM at cells. This can for instance be reached by delivery trehalose in a parenteral delivery form and replacing a fraction of the carbohydrates so that 0.1 - 50 mg/kg body weight / day, preferably 0.2 - 25 mg/kg body weight / day, yet more preferably 0.4 - 5 mg/kg body weight / day, yet more preferably 0.5 - 1.5 mg/kg body weight / day and most preferably 1 mg/kg body weight / day is provided. For oral delivery daily dose of Trehalose can be higher for instance about 0.5 to about 100 g/adult/day or 0.5 to 100 g/adult/day, and preferably about 1 to about 50 g/adult/day with respect to the amount of Trehalose or its derivates.

Compounds for medical use as described in the above embodiments are for instance the autophagy inducers Silibinin or Curcumin

Compounds for medical use as described in the above embodiments a can also be one or more compounds of the mTOR-dependent autophagy inducers are compounds such as Rapamycin and analogs (CCI-779, RAD001, AP23573), Perhexiline, Amiodaronel, Niclosamide, Rottlerin, Tori-1 (with structure) PI103 and structurally related compounds, Phenethyl isothiocyanate (PEITC) or Dexamethasone

Suitable Rapamycin analogues for present invention are of the group of the Rapamycin derivatives such as Everolimus, Temsirolimus, 40-O-(2-hydroxyethyl)-rapamycin, and/or 32-deoxorapamycin, and/or 16-pent-2-ynloxy-32-deoxorapamycin, and/or 16-pent-2-ynloxy-32 (S or R)-dihydro-rapamycin, and/or 16-pent-2-ynloxy-32 (S or R)-dihydro-40-O-(2-hydroxyethyl)-rapamycin, and/or 40-[3-hydroxy-2-(hydroxy-methyl)-2-methylpropanoate]-rapamycin (also known as CCI779) and/or 40-epi-(tetrazolyl)-rapamycin (also known as ABT578), and/or the so-called rapalogs, e.g. as disclosed in WO9802441, WO0114387 and WO0364383, AP23573, AP23464; AP23675 or AP23841, e.g. AP23573, and/or compounds disclosed under the name TAF-A-93, and/or compounds disclosed under the name biolimus. Other for present invention preferred rapamycin derivative is selected from the group consisting of 40-O-(2-hydroxyethyl)-rapamycin (also known as everolimus), and/or 32-deoxorapamycin, and/or 16-pent-2-ynloxy-32-deoxorapamycin, and/or 16-pent-2-ynloxy-32 (S or R)-dihydro-rapamycin, and/or 16-pent-2-ynloxy-32 (S or R)-dihydro-40-O-(2-hydroxyethyl)-rapamycin, and/or 40-[3-hydroxy-2-(hydroxy-methyl)-2-methylpropanoate]-
rapamycin (also known as CC1779 or temsirolimus) and/or 40-epi-(tetrazolyl)-rapamycin (also known as ABT578), and/or 40-O-(2-hydroxyethyl)-rapamycin.

Suitable phenothiazine derivatives for present invention are for example compounds chosen from quinacrine, imipramine, carbamazepine, phenazine, phenothiazine, promazine, chloropromazine, haloperidol, clozapine, 2-chlorophenothiazine, promethazine (10-(2dimethylaminopropyl)phenothiazine), chloroprothixen, and acepromazine.

Compounds for medical use as described in the above embodiments a can also be one or more compounds of the mTOR-independent autophagy inducers are compounds such as Lithium, L-690,330, Carbamazepine, sodium valproate, Verapamil, loperamide, Amiodarone, nimodipine, Nitrendipine, niguldipine, Nicardipine, pimozide, Calpastatin, calpeptin, Clonidine, rilmenidine, 2',5'-Dideoxyadenosine, NF449, Minoxidil, Penitrem A, Trehalose, Spermidine, Resveratrol, Fluspirilene, trifluoperazine, SMER10, SMER18, SMER28 OR Compound C (dorsomorphin)

A method of inhibiting critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis in a patient in need thereof, comprising administering to the patient an autophagy activating compound, wherein the patient suffers from a condition selected from the group consisting of Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease and Osteoporosis Secondary to ICU Admission and whereby the autophagy activating compound is administered in an amount sufficient to critical illness-induced osteoclastogenesis in the patient. The present invention provides in an embodiment that autophagy inducing compounds are administered in a dose such that a such that plasma levels are reached sufficient prevent or inhibit PBMCs in critically ill patients of differentiating into osteoclasts. The autophagy inducing compound can be any one of the group of glutamine, resveratrol, fenofibrate, Rapamycin, Spermidine, Everolimus, Promethazine or a derivative thereof or any of the autophagy inducing compounds of present invention for instance disclosed in this application.

A autophagy inducing compound of present invention for use in a method of inhibiting critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis in a patient in need thereof, comprising administering to the patient said autophagy activating compound such that plasma levels reached sufficient prevent or inhibit PBMCs in critically
ill patients of differentiating into osteoclasts, wherein the patient suffers from a condition selected from the group consisting of Critical Illness Bone Atrophy, Metabolic Bone Disease Following ICU Admission, Critical Illness Related Metabolic Bone Disease and Osteoporosis Secondary to ICU Admission and whereby the autophagy activating compound is administered in an amount sufficient to critical illness-induced osteoclastogenesis in the patient. The present invention provides in an embodiment that autophagy inducing compounds are administered in a dose such that plasma levels are reached sufficient prevent or inhibit PBMCs in critically ill patients of differentiating into osteoclasts. The autophagy inducing compound can be any one of the group of glutamine, resveratrol, fenofibrate, Rapamycin, Spermidine, Everolimus, Promethazine or a derivative thereof or any of the autophagy inducing compounds of present invention for instance disclosed in this application.

Some of the other embodiments of the invention are set forth in claim format directly below:

1. An autophagy activator compound for use as a medicine to cure or prevent bone degenerative disorder in a mammalian subject whereby the autophagy activator compound is glutamine and whereby the bone degenerative disorder is caused by acute critical illness or prolonged critical illness.

2. The autophagy activator compound according to embodiment 1, whereby the bone degenerative disorders is osteoclast activity increased due to acute critical illness and/or prolonged critical illness.

3. The autophagy activator compound according to any one of the previous embodiments, whereby the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder of critically ill patients.

4. The autophagy activator compound according to any one of the previous embodiments, whereby the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness.

5. The autophagy activator compound according to any one of the previous embodiments, whereby the bone degenerative disorders is enhanced osteoclast formation from circulating precursors due to autophagy deficiency disorder.

6. An autophagy activator compound for use as a medicine to cure or prevent bone degenerative disorder in a mammalian subject whereby the autophagy activator compound
is glutamine and whereby the bone degenerative disorder is caused by (parenteral) nutrition-induced suppression of autophagy.

7. The autophagy activator compound according to any one of the previous embodiment 6, whereby the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder by (parenteral) nutrition-induced suppression of autophagy in critically ill patients..

8. The autophagy activator compound according to any one of the previous embodiment 6, whereby the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness (parenteral) nutrition-induced suppression of autophagy.

9. The autophagy activator compound according to any one of the previous embodiments 6 to 8, whereby the bone degenerative disorders is enhanced osteoclast formation from circulating precursors due to autophagy deficiency disorder.

10. The autophagy activator compound according to any one of the previous embodiments, whereby the autophagy activator compound that is glutamine inhibits osteoclast activity.

11. The autophagy activator compound according to any one of the previous embodiments 1 to 10, whereby the glutamine compound is administered to said patient or mammalian subject such to such that it induces low millimolar concentrations (for instance, 1 - 10 mM, preferably 2-4 mM) of ammonium (glutamine-derived ammonia) after glutamine deamination in mitochondria which stimulates autophagy.

12. The autophagy activator compound according to any one of the previous embodiments 1 to 10, whereby the compound is administered to said patient or mammalian subject by delivering said compound in a watery fluid delivery (for instance enteral or parenteral) comprising the watery fluid having dissolved therein from 1 to 150 mMoles/1 of glutamine.

Some of the other embodiments of the invention are set forth in claim format directly below:

1. An autophagy activator compound for use as a medicine to cure or prevent bone degenerative disorder in a mammalian subject whereby the autophagy activator compound is a trehalose, for instance of the group consisting of a-D-glucopyranosyl-(l→l)-a-D-glucopyranoside, a-D-glucopyranosyl-(l→1)-β-D-glucopyranoside and β-D-
glucopyranosyl-(1→1)-β-D-glucopyranoside or a trehalose-based polymer such as poly(6-Vinyladipoyl-Trehalose) and whereby the bone degenerative disorder is caused by acute critical illness or prolonged critical illness.

2. The autophagy activator compound according to embodiment 1, whereby the bone degenerative disorders is osteoclast activity increased due to acute critical illness and/or prolonged critical illness.

3. The autophagy activator compound according to any one of the previous embodiments, whereby the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder of critically ill patients.

4. The autophagy activator compound according to any one of the previous embodiments, whereby the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness.

5. The autophagy activator compound according to any one of the previous embodiments, whereby the bone degenerative disorders is enhanced osteoclast formation from circulating precursors due to autophagy deficiency disorder.

6. An autophagy activator compound for use as a medicine to cure or prevent bone degenerative disorder in a mammalian subject whereby the autophagy activator compound is a trehalose, for instance of the group consisting of α-D-glucopyranosyl-(1→1)-α-D-glucopyranoside, α-D-glucopyranosyl-(1→1)-β-D-glucopyranoside and β-D-glucopyranosyl-(1→1)-β-D-glucopyranoside or a trehalose-based polymer such as poly(6-Vinyladipoyl-Trehalose) and whereby the bone degenerative disorder is caused by (parenteral) nutrition-induced suppression of autophagy.

7. The autophagy activator compound according to any one of the previous embodiment 6, whereby the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder by (parenteral) nutrition-induced suppression of autophagy in critically ill patients.

8. The autophagy activator compound according to any one of the previous embodiment 6, whereby the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness (parenteral) nutrition-induced suppression of autophagy.

9. The autophagy activator compound according to any one of the previous embodiments 6 to 8, whereby the bone degenerative disorders is enhanced osteoclast formation from circulating precursors due to autophagy deficiency disorder.
10. The autophagy activator compound according to any one of the previous embodiments, whereby the autophagy activator compound that is a trehalose, for instance of the group consisting of \( \alpha-D\)-glucopyranosyl-(1→1)-\( \alpha-D\)-glucopyranoside, \( \alpha-D\)-glucopyranosyl-(1→1)-\( \beta-D\)-glucopyranoside and \( \beta-D\)-glucopyranosyl-(1→1)-\( \beta-D\)-glucopyranoside or a trehalose-based polymer such as poly(6-Vinyladipoyl-Trehalose) inhibits osteoclast activity.

11. The autophagy activator compound according to any one of the previous embodiments 1 to 10, whereby the compound is administered to said patient or mammalian subject such to reach 1 to 100 mM, preferably 2 - 50 mM or more preferably 5 to 20 mM at cells.

12. The autophagy activator compound according to any one of the previous embodiments 1 to 10, whereby the compound is administered to said patient or mammalian subject by delivering said compound in a parenteral delivery form and replacing a fraction of the carbohydrates of having a concentration so that 0.1 - 50 mg/kg body weight / day, preferably 0.2 - 25 mg/kg body weight / day, yet more preferably 0.4 - 5 mg/kg body weight / day, yet more preferably 0.5 - 1.5 mg/kg body weight / day and most preferably 1 mg/kg body weight / day of said compound is provided.

13. The autophagy activator compound according to any one of the previous embodiments 1 to 10, whereby the compound is administered to said patient or mammalian subject by delivering said compound in a parenteral delivery form an enteral dose of about 0.5 to about 100 g/adult/day or 0.5 to 100 g/adult/day, and preferably about 1 to about 50 g/adult/day with respect to the amount of compound is provided.

14. The autophagy activator compound according to any one of the previous embodiments 1 to 13 in combination with glutamine.

15. The autophagy activator compound according to any one of the previous embodiments 1 to 13 in combination with glutamine whereby the glutamine compound is a dose such that it induces low millimolar concentrations (for instance, 1 - 10 mM, preferably 2-4 mM) of ammonium (glutamine-derived ammonia) after glutamine deamination in mitochondria which stimulates autophagy when administered to said patient or mammalian subject.

16. The autophagy activator compound according to any one of the previous embodiments 1 to 13 in combination with glutamine whereby the compounds are in a watery fluid delivery form comprising the watery fluid having dissolved therein from 1 to 150 mMoles/1 of glutamine for enteral or parenteral administration to said patient or mammalian subject.
Some of the other embodiments of the invention are set forth in claim format directly below:

1. An autophagy activator compound for use as a medicine to cure or prevent bone degenerative disorder caused by acute critical illness or prolonged critical illness in a mammalian subject.

2. An autophagy activator compound for use as a medicine to inhibit osteoclast activity increase due to autophagy deficiency disorder caused by parenteral or enteral nutrition-induced suppression of autophagy.

3. An autophagy activator compound for use as a medicine according to any one of the embodiments 1 to 2, whereby bone degenerative disorder is increased osteoclast activity.

4. Embodiment 1, 2 or 3, whereby the autophagy activator compound is selected from the group consisting of: (a) Loperamide; (c) Niguldipine; (d) Pimozide; (e) Nicardipine; (f) Penitrem A; (g) Fluspirilene; (h) Trifluoperazine.
5. Embodiment 1, 2 or 3, whereby the autophagy activator compound is selected from the group consisting of: (a) 10-NCP, (b) Promazine whereby $R = H$, (c) Chlorpromazine whereby $R = Cl$

\[
\text{COT}
\]

, (d) Triflupromazine whereby $R = CF_3$, (e) Prometazine

, (f) Mesoridazine and (g) Thioridazine.

6. Embodiment 1, 2 or 3, whereby the autophagy activator compound is at least one compound selected from the group consisting of:

(a) compounds of formula (I):

wherein $X$ is selected from CR4R5 and NR6; $R1$ is selected from hydrogen, Cl-6 alkyl and phenyl, wherein the alkyl and phenyl are substituted with 0 or 1 halogen; $R2$ is selected from hydrogen, Cl-6 alkyl and phenyl, wherein the alkyl and phenyl are substituted with 0
or 1 halogen; R3 is selected from hydrogen, hydroxyl, CI-6 alkyl and phenyl; R5 is selected from CI-6 alkyl and phenyl, halophenyl, benzimidazole, dihydrobenzimidazole, benzimidazolone; optionally R4 and R5 are taken together to form a 5 or 6 membered heterocycloalkyl comprising two nitrogen atoms, wherein the heterocycloalkyl is substituted with 1, 2 or 3 substituents selected from the group consisting of CI-6 alkyl, phenyl, and =O; R6 is selected from hydrogen and CI-6 alkyl; R7a, R8a, R9a, R10a, R11a, R7b, R8b, R9b, R10b, and R11b are each independently selected from hydrogen, hydroxyl, halogen and CI-6 haloalkyl; optionally R11a and R11b are taken together to form a heterocycle of the following structure:

wherein R11 is selected from CH2, NH, O and S; R12 and R13 are each independently selected from hydrogen and CI-6 alkyl; R14a and R14b are each independently selected from hydrogen and CI-6 alkyl; R15 is selected from phenyl substituted with 0 or 1 halogen or nitro; R16 is selected from hydrogen and CI-6 alkyl; Y is N or CH; and
pharmaceutically acceptable salts thereof; (b) compounds of formula (II):

![Chemical Structure](image)

wherein R17 is selected from hydrogen and Cl-6 alkyl; R18a and R18b, are each independently selected from hydrogen and Cl-6 alkyl; R19a, R19b, R20a, R20b, and R21 are each independently selected from hydrogen, halogen and nitro; R22 is selected from hydrogen and Cl-6 alkyl; R23 is selected from \( -(CH_2)_nNR_{24a}R_{24b} \) and \( -<CH_2>_nR_{24a} \); R24a and R24b are each independently selected from Cl-6 alkyl and phenyl, wherein the alkyl is substituted with 0 or 1 phenyl substituents; optionally R24a and R24b are taken together with the nitrogen to which they are attached to form a piperidine which is substituted with 0, 1 or 2 phenyl substituents; n is a positive integer from 2 to 4; and pharmaceutically acceptable salts thereof

![Chemical Structure](image)

(c) compounds of formula (III):

wherein R25 is selected from hydrogen and Cl-6 alkyl; R26a, R26b, R27a, and R27b are each independently selected from hydrogen, halogen and Cl-6 alkyl; R28 is selected from \( -0(CH_2)_mNR_{29a}R_{29b} \) and \( -NH(CH_2)_mNR_{29a}R_{29b} \); R29a and R29b are each independently selected from hydrogen and Cl-6 alkyl; Z is O, S or NH; m is a positive integer from 1 to 3; and pharmaceutically acceptable salts thereof

![Chemical Structure](image)

(d) compounds of formula (IV):
wherein R30 is selected from hydrogen, Cl-6 alkyl and halogen; R31a and R31b are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R32 is selected from hydrogen and Cl-6 alkyl; R33 and R34 are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R35a and R35b are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R36a and R36b are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R37a and R37b are each independently selected from hydrogen, hydroxyl and Cl-6 alkyl; R38 is selected from hydrogen, hydroxyl and Cl-6 alkyl; optionally R37a and R38 are taken together to form a three membered heterocycle of the formula:

wherein R38' is O, S or NH; R39a is selected from hydrogen, hydroxyl and Cl-6 alkyl; R39b is selected from hydrogen, hydroxyl, Cl-6 alkyl and C2-6 alkenyl; U, V and W are each independently selected from O, S, and NH; and pharmaceutically acceptable salts thereof.

7. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Azithromycin macrolide (an azalide, a subclass of macrolide antibiotics, IUPAC name: 2R,3S,4R,5R,8R,10R,1 IR,12S,13S,14R)-2-ethyl-3,4,10-trihydroxy-3,5,6,8,10,12,14-heptamethyl-15-oxo-11-[[3,4,6-trideoxy-3-(dimethylamino)-β-D-xylo]-oxy]-1-oxa-6-azacyclopentadec-13-yl 2,6-dideoxy-3-C-methyl-3-0-methyl-a-L-ribo-hexopyranoside).

8. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Loperamide.

9. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Silibinin.

10. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Niguldipine.

11. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Pimozide.
12. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Nicardipine.

13. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Penitrem A.

14. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Fluspirilene.

15. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Trifluoperazine.

16. Any one of the previous embodiments, whereby the autophagy activator compound is 10-NCP.

17. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Promazine.

18. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Chlorpromazine.

19. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Triflupromazine.

20. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Promazine.

21. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Mesoridazine.

22. Any one of the previous embodiments 1 to 3, whereby the autophagy activator compound is Thioridazine.

23. Any one of the previous embodiments 1 to 22, whereby the treatment is to inhibit increased bone resorption disorder.

24. Any one of the previous embodiments 1 to 22, whereby the treatment is to inhibit osteoclast activity.

25. Any one of the previous embodiments 1 to 22, whereby the treatment is to inhibit formation of osteoclasts (TRAP-positive multinuclear cells) from blood progenitor cells.

26. Any one of the previous embodiments 1 to 22, whereby the treatment is to activate bone formation.

27. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is bone metabolism disorder.

28. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorder is caused by acute critical illness or prolonged critical illness.
29. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorder is caused by (parenteral) nutrition-induced suppression of autophagy.

30. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorder is caused by mTOR activation.

31. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is hyperresorption of bone.

32. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is an imbalance in the regulation of bone resorption and bone formation results in metabolic bone diseases such as osteoporosis.

33. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is elderly osteoporosis.

34. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is osteoporosis.

35. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is pathological increased bone resorption and pathological decreased bone formation.

36. Any one of the previous embodiments, whereby the bone degenerative disorders is disrupted calcium and bone metabolism by acute critical illness and/or prolonged critical illness.

37. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is caused by enhanced osteoclast formation from circulating precursors in the blood during critical illness.

38. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation from circulating precursors leading to osteoporosis.

39. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation due to a increased cytokine production disorder.

40. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation due to hormonal disorder.

41. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation due to VitaminD deficiency.

42. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation due to glucocorticoids (pre)treatment.

43. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation due to heparinoids (pre)treatment.
44. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is increased osteoclast formation due to autophagy deficiency disorder.

45. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is osteoclast activity increase due to acute critical illness and/or prolonged critical illness.

46. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is osteoclast activity increase due to increased cytokine production disorder.

47. Any one of the previous embodiments 1 to 22, whereby the bone degenerative disorders is osteoclast activity increase due to hormonal disorder.

48. A therapeutic agent or a pharmaceutical composition comprising a autophagy activator compound according to any one of the previous embodiments or an analogue thereof for use in a treatment of treating or preventing bone degenerative disorder in a mammalian subject.

49. Embodiment 48, whereby the bone degenerative disorders is increased bone resorption disorder or bone hyperresorption.

50. Embodiment 48, whereby the treatment is to prevent, inhibit or reduce increased bone resorption or bone hyperresorption.

Particular and preferred aspects of the invention are set out in the accompanying independent and dependent claims. Features from the dependent claims may be combined with features of the independent claims and with features of other dependent claims as appropriate and not merely as explicitly set out in the claims.

Thus, the claims following the detailed description are hereby expressly incorporated into this detailed description, with each claim standing on its own as a separate embodiment of this invention.

**Drawing Description**

**BRIEF DESCRIPTION OF THE DRAWINGS**

The present invention will become more fully understood from the detailed description given herein below and the accompanying drawings which are given by way of illustration only, and thus are not limitative of the present invention, and wherein:
Figure 1 concerns graphics (B) showing spermidine dose response - osteoclast activity (Resorption) in stimulated healthy Peripheral Blood Mononuclear Cells (PBMCs) and shows the photos (A) on the osteoclast PBMCs with spermidine. Healthy PBMCs were differentiated for 14 days in the presence of RANKL and M-CSF, with or without spermidine (2 x 10⁻⁸ M - 2 x 10⁻⁹ M), on a hydroxyapatite film layer. At 14 days, cells were removed with sodium hypochlorite, and the hydroxyapatite layer was stained with Von Kossa. Osteoclast resorption was analysed using Image J software. (A) Von Kossa staining of the hydroxyapatite film layer. (B) Percentage of total area resorbed. (**p<0.01 vs. Control; *p<0.05 vs. Control). As the lowest concentration still affecting osteoclast resorption, 2x10⁻⁸ M spermidine was selected for use in future experiments.

Figure 2 demonstrates the effect of spermidine osteoclast formation in stimulated (+RANKL/MCSF) PBMCs. Fig. 2A is a photodisplay and Fig 2B a graphic on mature osteoclast formation in healthy and sick stimulated PBMCs with spermidine. Healthy and sick PBMCs were differentiated for 14 days in the presence of RANKL and M-CSF, with or without spermidine (2x10⁻⁸M). At 14 days, cells were stained for tartrate-resistant acid phosphatase (TRAP), and nuclei visualised with DAPI staining. Mature osteoclasts were identified as cells staining positively for TRAP, with 3 or more nuclei. (A) Micrograph images of PBMCs in culture at day 7, and at day 14 following TRAP/DAPI staining (B) Number of TRAP positive multinuclear cells per 5x10⁵ cells plated (** p<0.001 vs. Healthy cells; ap <0.05 vs. sick cells; n=4)

Figure 3 concerns the effects of spermidine on spontaneous osteoclast formation in unstimulated PBMCs. Figure3 demonstrates spontaneous mature osteoclast formation in healthy and sick unstimulated PBMCs with spermidine. Healthy and sick PBMCs were cultured for 14 days with or without spermidine (2x10⁻⁸M). At 14 days, cells were stained for tartrate-resistant acid phosphatase (TRAP), and nuclei visualised with DAPI staining. Mature osteoclasts were identified as cells staining positively for TRAP, with 3 or more nuclei. (A) Micrograph images of PBMCs at day 14 following TRAP/DAPI staining (B) Number of TRAP positive multinuclear cells per 5x10⁵ cells plated (** p<0.001 vs. Healthy cells; ap ≤ 0.05 vs. sick cells; n=4)

Figure 4. concerns the effect of spermidine on osteoclast activity in stimulated (+RANKL/MCSF) PBMCs. Figure4. Osteoclast activity in stimulated healthy and sick
PBMCs with spermidine. Healthy and sick PBMCs were cultured for 14 days in the presence of RANKL and M-CSF with or without spermidine (2x10^-8 M) on a hydroxyl apatite film layer. At 14 days, cells were removed with sodium hypochlorite, and the hydroxyapatite layer was stained with VonKossa. Osteoclast resorption was analysed using Image J software. (A) Von Kossa staining of the hydroxyapatite film layer. (B) Percentage of total area resorbed. (** p < 0.01 vs. Healthy; * p ≤ 0.05 vs. sick; n=4)

**Figure 5.** concerns the effect of spermidine on spontaneous osteoclast activity in unstimulated PBMCs. Healthy and sick PBMCs were cultured for 14 days with or without spermidine (2x10^-8 M) on a hydroxyl apatite film layer. At 14 days, cells were removed with sodium hypochlorite, and the hydroxyapatite layer was stained with VonKossa. Osteoclast resorption was analysed using Image J software. (A) Von Kossa staining of the hydroxyapatite film layer. (B) Percentage of total area resorbed.

**Figure 6.** concerns autophagy marker p62 protein expression in healthy and sick PBMCs/osteoclasts (RANKL/MCSF stimulated) and the effect of spermidine on healthy PBMCs/osteoclasts. Healthy and sick PBMCs were cultured for 14 days with or without spermidine (2x10^-8 M) in 6 well plates (5 x 10^6 cells/well). After 14 days, protein was extracted using the Ambion PARIS extraction kit (AppliedBiosystems), and proteins quantified using the BC Protein Assay (ThermoFischerScientific). (A) 5ug of protein was subjected to 15% Bis-tris gelelectrophoresis, transferred to nitrocellulose membrane and incubated at 4°C overnight in Anti-p62 (NovusBiologicals;1:1000). p62 protein was detected by ECL following incubation with AntiMouseIgG (DAKO;1:1000). GAPDH was used as a loading control. (B) p62 relative expression was quantified with Image QuantTL (Amersham Biosciences)and normalised to GAPDH (n=3; *** p ≤ 0.05 vs. Healthy).

**Figure 7:** Demonstrates that Osteoblast markers such as bone specific alkaline phosphatase also decrease over time in ICU. bsALP = Serum Bone Specific Alkaline phosphatase (bsALP).

**Figure 8** demonstrates effects on Osteoclast markers such as Crosslaps increase over time in ICU. Serum Crosslaps (collagen crosslinks).
Figure 9 provides a graphic overview of provides a review on osteoclast formation from circulating precursors in the blood, osteoclast maturation, osteoclast bone resorption activity, and the current marks with are indicative for such.

Figure 10a demonstrates osteoclast formation in PBMCs from Critically 111 Patients. Osteoclast formation is observed in Patient PBMCs after only 7 days in culture with RANKL and MCSF.

Figure 10b demonstrates the osteoclast formation in PBMCs from Critically 111 Patients. By Day 21 osteoclast formation is observed in Patient PBMCs with and without RANKL/MCSF.

Figure 10c shows the Day 21 Patient PBMCs with and without RANKL/MCSF are multinuclear.

Figure 11 shows mTOR-dependent autophagy inducers.

Figure 12 shows mTOR-independent autophagy inducers.

Figure 13 shows additional mTOR-independent autophagy inducers.

Figure 14 displays a dose response of spermidine on osteoclast formation.

Figure 15 displays a dose response of rapamycin on osteoclast formation.

Figure 16 displays a dose response of Everolimus on osteoclast formation.

Figure 17 displays a dose response of prometazine on osteoclast formation.

Figure 18 displays a dose response of 5-Azacytidine on osteoclast formation.

Figure 19. Rabbit in vivo study of prolonged critical illness (A) Serum ionized calcium levels were significantly lower in critically ill rabbits compared to healthy controls over the 7-day period (n = 30 critically ill (grey boxes) and 15 healthy (white boxes); **p<0.01; multiple
testing ANOVA). (B) Serum osteocalcin levels were significantly reduced (62.3%) in critically ill rabbits at day 4 of illness (n = 30 critically ill and 15 healthy; **p<0.01). (C) Trabecular BMC and (D) BMD were significantly reduced (30.6% and 28.8%, respectively) in the proximal tibiae of critically ill rabbits vs. healthy controls. (E) No significant difference in cortical BMC or (F) thickness was observed (n = 30 and 15, respectively; *p<0.05, **p<0.01).

**Figure 20. Circulating osteoclast precursors in healthy and critically ill patient peripheral blood** (A) Representative dot plots of CD14+/CD1 lb+ early osteoclast precursors and CD14+/CD1 lb+/VNR+ osteoclast precursors in healthy and patient peripheral blood. (B) FACS analysis revealed a significant increase in early osteoclast precursors (CD14/CD1 lb positive) from the peripheral blood of critically ill patients (99.8 ± 0.38% and 83.9 ± 7.09%, respectively). A trend towards an increase in mature circulating osteoclasts (CD14/CD1 lb/VNR positive) was also observed (n = 5; **p<0.01).

**Figure 21. Human in vitro study of osteoclast differentiation** (A) Formation of mature, multi-nuclear osteoclasts was visualized by TRAP positivity (scale bar = 100µm) (I) and resorption on hydroxyapatite (scale bar = 500µm) (II) in healthy or patient cultures, in the presence and absence of RANKL and M-CSF. (B) The number of TRAP positive multi-nuclear cells was significantly increased in critically ill patient PBMC cultures with and without RANKL and M-CSF (C) Resorption of hydroxyapatite was increased in patient cultures in the presence and absence of RANKL and M-CSF. (n = 8; ***p<0.05 versus healthy cells).

**Figure 22. Effect of critically ill patient serum on osteoclast differentiation and activity** (A) Formation of mature, multi-nuclear osteoclasts was visualized by TRAP positivity (top row), F-actin ring formation (second row), formation of multi-nuclear cells (third row, merged image of TRAP, DAPI and F-actin ring staining) and resorption on hydroxyapatite (bottom row; analysed with Von Kossa staining) in pooled PBMC cultures from healthy controls and critically ill patients. (B) The number of TRAP positive multi-nuclear cells was significantly increased in patient PBMC cultures with (white bars) and without (grey bars) osteoclastogenic factors, in the presence of PS (C) F-actin ring formation was also significantly increased in
the presence of patient serum in patient PBMC cultures in the presence (white bars) and absence (grey bars) of RANKL and MCSF. (D) Resorption of hydroxyapatite was increased in patient cultures cultured with 10% HS or 10% PS in the presence (white bars) and absence (grey bars) of RANKL and MCSF (n = 8; ***p<0.05 versus patient cells plus HS and healthy cells plus PS; ap<0.05 versus healthy cells plus HS).

**Figure 23.** Inhibition of inflammatory cytokines in patient serum increases in vitro osteoclast formation (A) Formation of mature, multinuclear osteoclasts was visualized by TRAP in pooled PBMC cultures from critically ill patients with 10% PS and varying concentrations of Anti-IL-6 and/or Anti-TNF-a neutralizing antibodies. (B) The number of TRAP positive multi-nuclear cells was significantly increased in patient PBMC cultures with µg/ml Anti-IL-6 or 4µg/ml and 8µg/ml of Anti-TNF-α, or a combination of Anti-IL-6 (1µg/ml) and Anti-TNF-a (8µg/ml) (n = 4; *p<0.05, ***p<0.001 vs. control cultures).

**Figure 24.** Effect of critically ill patient serum on osteogenic differentiation and angiogenesis in vitro (A) Alizarin Red staining of hPDC monolayers cultured for 21 days in 10% FBS, 10% HS or 10% PS (B) Alizarin Red staining quantified by assessment of optical density at 540 ran revealed no significant difference in mineralization. (C) qRT-PCR analysis of RUNX2; (D) COL1A1; (E) ALP; and (F) VEGFA normalised to β-actin revealed no differences in gene expression between HS and PS conditions, although a reduction in VEGF-R1 expression was observed in patient serum conditions (G), which was also observed at the protein level (H) (n = 8; ***p<0.001 versus FBS; **p<0.01 versus FBS; ap0.01 versus healthy serum).

**Figure 25.** Effect of critically ill patient serum on bone formation in vivo (A) Bone quantification of hPDC NuOSS™ implants was carried out using µCT analysis 8 weeks after implantation, and revealed a significant reduction in bone formation in patient-serum conditions compared to HS (mature bone = yellow; remaining CaP grains = blue). No significant differences in CaP grains or fibrous tissue were detected in the scaffold. (B) A trend towards an increase in the number of TRAP positive osteoclasts per scaffold in patient serum-treated hPDCs was observed, although this did not reach significance (p = 0.07). (C) Upon explant, scaffolds coated with PS treated hPDCs had visibly less vasculature than scaffolds coated with HS hPDCs (insets). This was confirmed by quantification of CD31 immunohistochemistry, with a significant reduction in the number of CD31 positive blood
vessels in scaffolds with PS treated hPDCs (n = 4; ***p<0.001 versus healthy serum; ap<0.01 versus FBS).
OSTEOCLAST ACTIVITY

Claims

1. An autophagy inducing compound for use in a method of inhibiting for suppressing critical illness [MeSH Descriptor: C23.550.29J.625] that enhanced osteoclastogenesis or increased osteoclast differentiation in a subject in need for a treatment of Critical Illness Related Metabolic Bone Disease or of critical illness induced Osteopenia [ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851] secondary to ICU Admission.

2. The autophagy activator compound according to claim 1, wherein the compound is a mTOR independent autophagy inducer.

3. The autophagy activator compound according to claim 1, wherein the compound is a mTOR independent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 10 g per day or less, or such that serum level of said autophagy inducing compound in the patient is 10 µmol/L or less.

4. The autophagy activator compound according to claim 1, wherein the compound is a mTOR independent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 1 g per day or less, or such that serum level of said autophagy inducing compound in the patient is 1000 nmol/L or less.

5. The autophagy activator compound according to any one of the claims 2 to 4, wherein the mTOR independent autophagy inducer is a spermidine derivative.

6. The autophagy activator compound according to claim 2, wherein the mTOR independent autophagy inducer is a phenothiazine derivative.

7. The autophagy activator compound according to claim 2, wherein the mTOR independent autophagy inducer is a phenothiazine derivative and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 1 g per day or less, or such that serum level of said autophagy inducing compound in the patient is 500 nmol/L or less.

8. The autophagy activator compound according to any one of the claims 2 to 4, wherein the mTOR independent autophagy inducer is spermidine.

9. The autophagy activator compound according to claim 2, wherein the mTOR independent autophagy inducer is promethazine.
10. The autophagy activator compound according to claim 2, wherein the mTOR independent autophagy inducer is promethazine and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 100 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 150 nmol/L or less.

11. The autophagy activator compound according to claim 1, wherein the compound is a mTOR dependent autophagy inducer.

12. The autophagy activator compound according to claim 1, wherein the compound is a mTOR dependent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 10g per day or less, or such that serum level of said autophagy inducing compound in the patient is 1000 nmol/L or less.

13. The autophagy activator compound according to claim 1, wherein the compound is a mTOR dependent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 1g per day or less, or such that serum level of said autophagy inducing compound in the patient is 1000 nmol/L or less.

14. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative.

15. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 5 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 100 nmol/L or less.

16. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 2 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 50 nmol/L or less.

17. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound
by the patient such that autophagy inducing compound intake is such that serum level of said autophagy inducing compound in the patient is 10 nmol/L or less.

18. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is Everolimus.

19. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is Everolimus and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 10 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 100 nmol/L or less.

20. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is Everolimus and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 1 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 50 nmol/L or less.

21. The autophagy activator compound according to claim 11, wherein the mTOR dependent autophagy inducer is Everolimus and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is such that serum level of said autophagy inducing compound in the patient is 5 nmol/L or less.

22. A method of inhibiting for suppressing critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis or increased osteoclast differentiation in a subject in need for a treatment of Critical Illness Related Metabolic Bone Disease or of critical illness induced Osteopenia [ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851] secondary to ICU Admission, the method comprising inhibiting the osteoclast formation or osteoclastogenesis by an autophagy inducing or activating compound.

23. The method according to claim 22, wherein the compound is a mTOR independent autophagy inducer.

24. The method according to claim 23, wherein the compound is a mTOR independent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 10g per day or less, or such that serum level of said autophagy inducing compound in the patient is 1000 nmol/L or less.
25. The method according to claim 23, wherein the compound is a mTOR independent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 1 g per day or less, or such that serum level of said autophagy inducing compound in the patient is 1000 nmol/L or less.

26. The method according to any one of the claims 23 to 25, wherein the mTOR independent autophagy inducer is a spermidine derivative.

27. The method according to claim 23, wherein the mTOR independent autophagy inducer is a phenothiazine derivative.

28. The method according to claim 23, wherein the mTOR independent autophagy inducer is a phenothiazine derivatives and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 1 g per day or less, or such that serum level of said autophagy inducing compound in the patient is 500 nmol/L or less.

29. The method according to claim 23, wherein the mTOR independent autophagy inducer is spermidine.

30. The method according to claim 23, wherein the mTOR independent autophagy inducer is promethazine.

31. The method according to claim 23, wherein the mTOR independent autophagy inducer is promethazine and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 100 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 150 nmol/L or less.

32. The method according to claim 22, wherein the compound is a mTOR dependent autophagy inducer.

33. The method according to claim 32, wherein the compound is a mTOR dependent autophagy inducer and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 1 g per day or less, or such that serum level of said autophagy inducing compound in the patient is 1000 nmol/L or less.

34. The method according to claim 32, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative.

35. The method according to claim 32, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative and wherein the autophagy inducing compound is to
be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 5 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 100 nmol/L or less.

36. The method according to claim 32, wherein the mTOR dependent autophagy inducer is rapamycin or a rapamycin derivative and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said compound by the patient such that autophagy inducing compound intake is 2 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 50 nmol/L or less.

37. The method according to claim 32, wherein the mTOR dependent autophagy inducer is Everolimus.

38. The method according to claim 32, wherein the mTOR dependent autophagy inducer is Everolimus and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 10 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 100 nmol/L or less.

39. The method according to claim 32, wherein the mTOR dependent autophagy inducer is Everolimus and wherein the autophagy inducing compound is to be used in conjunction with restricting intake of said by the patient such that autophagy inducing compound intake is 1 mg per day or less, or such that serum level of said autophagy inducing compound in the patient is 50 nmol/L or less.

40. Any one of the previous claims wherein the critically ill patient is subjected to total parenteral nutrition.


42. A DNA methylation inhibitor according to claim 41, for use in a method of inhibiting for suppressing enhanced osteoclastogenesis or increased osteoclast differentiation in a subject in need for a treatment of a disorders of bone density (ICD-10-CM M80-M85).

43. A DNA methylation inhibitor for use in a method of inhibiting for suppressing critical illness [MeSH Descriptor: C23.550.291.625] enhanced osteoclastogenesis or increased osteoclast differentiation in a subject in need for a treatment of Critical Illness Related Metabolic Bone Disease or of critical illness induced Osteopenia [ICD-10 M85.8, ICD-9 733.90, DiseasesDB 29870 or MeSH D001851] secondary to ICU Admission.
44. The DNA methylation inhibitor according to any one of the previous claims 41 to 43 wherein said DNA methylation inhibitor is selected from the group consisting of decitabine (or 5-aza-2'-deoxycytidine or 5-azadC, azacitidine (or 5-azacytidine), vorinostat (Zolinza™), procainamide and derivatives thereof.

45. The DNA methylation inhibitor according to any one of the previous claims 41 to 44, wherein said DNA methylation inhibitor is delivered or administrated to subject in need thereof to reach very low nanomolar plasma concentration of a value of the range of 1 to 10 nM to suppress osteoclastogenesis or osteoclast differentiation.
Figures

White spots are bone resorption (osteoclasts eating away bone)

Control

2 μM

200 nM

20 nM

2 nM

Fig. 1A
Healthy       Sick       Healthy + Spermidine       Sick + Spermidine

A

Healthy
Sick

Day 7 - In Culture

Day 14 - TRAP

Day 14 - TRAP/DAPI

These are red spots

Red spots surrounded by smaller blue spots

Fig 2A
Fig. 2B

TRAP+ve cells/5x10⁵ cells

Healthy  Sick  Healthy Spermidine  Sick Spermidine
Fig. 3A

Red spot surrounded by little blue dots

Fig. 3A (continued)
White spots are bone resorption

Fig. 4.
Bone resorption white spot

Fig 5
Fig 6
Osteoclasts

Mononuclear cell

Pre-osteoclast

Mature Osteoclast

++

RANKL/MCSF Cytokines

++

Fig 9
mTOR-dependent autophagy inducers

Rapamycin and analogs (CCI-779, RAD001, AP23573), Perhexiline, Amiodarone, Niclosamide, Rottlerin, Tori-1 (with structure) PI103

and structurally related compounds, Phenethyl isothiocyanate (PEITC) and Dexamethasone

Fig 11
mTOR-independent autophagy inducers

Lithium, L-690,330

Carbamazepine, sodium valproate, Verapamil, calpeptin

Sodium valproate, Amiodarone, nimodipine

Loperamide, Nitrendipine, niguldipine

Verapamil, Nicardipine, pimozide

Amiodarone, Nifedipine, Calpastatin

Nifedipine, Prunesin, Clonidine

Calpastatin, rilmenidine, Clonidine, 2',5'-Rilmenidine

Dideoxyadenosine, NF449

**Fig 12**
mTOR-independent autophagy inducers

- Minoxidil
- Penitrem A
- Trehalose
- Spermidine
- Resveratrol
- Fluspirilene
- Trifluoperazine
- SMER10
- SMER18
- SMER28
- Compound C (dorsomorphin)

**Fig 13**
Fig 21
Fig 22
Fig 23
**Fig 24**

Healthy Patient

151 kDa VEGF-R1

42 kDa β-actin