Ghrelin and unacylated ghrelin stimulate human osteoblast growth via mitogen-activated protein kinase (MAPK)/phosphoinositide 3-kinase (PI3K) pathways in the absence of GHS-R1a

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Abstract

Recent studies demonstrate widespread expression of ghrelin among tissues and have uncovered its pleiotropic nature. We have examined gene expression of ghrelin and its two receptor splice variants, growth hormone secretagogue receptors (GHS-R) 1a and 1b, in human bone biopsies and in the human pre-osteoblastic SV-HFO cell line during differentiation. Additionally, we examined proliferative effects of ghrelin and unacylated ghrelin (UAG) in differentiating and non-differentiating cells. We detected GHS-R1b mRNA in human bone and osteoblasts but not ghrelin’s cognate receptor GHS-R1a, using two different real-time PCR assays and both total RNA and mRNA. In osteoblasts GHS-R1b mRNA expression remained low during the first 14 days of culture, but increased 300% in differentiating cells by day 21. Both human bone biopsies and osteoblasts expressed ghrelin mRNA, and osteoblasts were found to secrete ghrelin.

Overall, ghrelin gene expression was greater in differentiating than non-differentiating osteoblasts, but was not increased during culture in either group. Ghrelin and UAG induced thymidine uptake dose-dependently, peaking at 1 and 10 nM respectively, at day 6 of culture in both non-differentiating and differentiating osteoblasts. The proliferative response to ghrelin and UAG declined with culture time and state of differentiation. The proliferative effects of ghrelin and UAG were suppressed by inhibitors of extracellular-signal-regulated kinase (ERK) and phosphoinositide-3 kinase, and both peptides rapidly induced ERK phosphorylation. Overall, our data suggest new roles for ghrelin and UAG in modulating human osteoblast proliferation via a novel signal transduction pathway.


Introduction

Ghrelin is an acylated 28-residue peptide hormone and is the natural ligand for the growth hormone (GH) secretagogue receptor (GHS-R1a; Kojima et al. 1999, van der Lely et al. 2004, Kojima & Kangawa 2005). The predominant form of ghrelin in serum is unacylated ghrelin (UAG). The majority of circulating ghrelin is derived from X/A-like cells in the oxyntic mucosa of the stomach with greatest expression in the fundus (Kojima et al. 1999). Ghrelin gene expression is widespread among tissues in humans, although the presence of its mRNA in human bone has not been determined (Gnanapavan et al. 2002). The GHS-R mRNA is as widely expressed as ghrelin, whereas GHS–R1a gene expression is limited to the pituitary and a few other peripheral tissues (Guan et al. 1997, Gnanapavan et al. 2002). Ghrelin stimulates the release of growth hormone, as well as prolactin and corticotrophin (ACTH) from the anterior pituitary. However, much recent evidence suggests numerous other functions for ghrelin at the periphery, including effects on cell proliferation (Murata et al. 2002, Baldanze et al. 2002, Broglio et al. 2002, Andreis et al. 2003, Jeffery et al. 2003, Kim et al. 2004, Nanzer et al. 2004, Zhang et al. 2004). Interestingly, although only the acylated form of ghrelin activates the GHS–R1a receptor, recent evidence suggests that UAG may also modulate cellular function and body growth, presumably by acting through an alternative receptor. For example,
transgenic mice that overexpress UAG are smaller than their wild-type counterparts (Ariyasu et al. 2004), direct treatment with UAG alters adipogenesis in the bone marrow of rats (Thompson et al. 2003), and in vitro UAG inhibits apoptosis of human cardiomyocytes and endothelial cells (Baldanzi et al. 2002). To the best of our knowledge nothing has been reported on the proliferative effects of UAG in osteoblasts.

In vivo, ghrelin may have an indirect effect on bone through its stimulation of GH release. However, the GHS-R1a agonist hexarelin inhibits markers of bone resorption in the rat, an effect not observed with GH treatment (Sibilia et al. 1999). This suggests that GH secretagogues may have GH-independent effects on bone in rodents. Interestingly, in rodents and in humans gastrectomy often results in osteopenia, although the mechanism for this is unclear (Bernstein et al. 2003). Furthermore, extracts from the oxyntic mucosa mobilize Ca^{2+} intracellular stores in osteoblast cell lines (Larsson et al. 2001), suggesting the presence of an osteotropic hormone(s) in the gastric mucosa, with ghrelin a possible candidate. Despite this, a recent study found that ghrelin did not induce a Ca^{2+} signal in a rat osteosarcoma cell line (Larsson et al. 2002). However, this does not exclude the possibility that ghrelin can signal via alternative intracellular pathways, and perhaps other receptors (Papotti et al. 2000, Baldanzi et al. 2002). In fact, three recent studies demonstrate that ghrelin has proliferative effects in rodent osteoblasts (Fukushima et al. 2005, Kim et al. 2005, Maccarinelli et al. 2005).

Since there are many physiological differences between rodents and humans we undertook to examine the role of ghrelin in regulating human osteoblast growth. In this study we have assessed the gene expression of ghrelin and its receptor subtypes in human bone cells. To further characterize the functional activity of ghrelin and its unacylated form, we have examined the effects of ghrelin and UAG on the growth of the well-characterized human osteoblast cell-line, SV-HFO (Weyts et al. 2003, Jansen et al. 2004, Eijken et al. 2005), and the possible intracellular signalling pathways involved in this process.

Materials and Methods

Human femoral head biopsy

Human bone material was obtained from femoral head biopsies of two osteoarthritic patients (bones 1 and 2, derived from separate patients). Bone marrow was removed and the inner cortical bone surfaces were carefully rinsed and homogenized using a Mikro Dismembrator S (Sartorius, Goettingen, Germany). RNA was isolated from the resulting homogenate as described below. Collection of human tissue was approved by the Erasmus MC Medical Ethical Commission (MEC No. 204-287).

Cell culture

A simian virus 40 (SV40)-immortalized human fetal osteoblast (SV-HFO) cell line (Weyts et al. 2003, Jansen et al. 2004, Eijken et al. 2005) was seeded at a density of 5500 cells/cm^2. The cells were cultured in Phenol Red-free α-minimal essential medium (αMEM; Gibco BRL, Paisley, UK), supplemented with 100 IU/ml penicillin, 100 µg/ml streptomycin (Life Technologies, Breda, The Netherlands), 20 mM HEPES, 1·8 mM CaCl_{2} and 10% heat-inactivated fetal calf serum (Gibco BRL) at 37°C in 5% CO_{2} in a humidified atmosphere. Medium was changed every 2–3 days and after 1 week cells were trypsinized for further culture under different experimental conditions. For the differentiation study, the cells were seeded at a density of 10 000 cells/cm^2. The cells were cultured in αMEM with 2% heat-inactivated charcoal-treated fetal calf serum at 37°C in 5% CO_{2}. From day 2 onward cells were cultured in the presence of 10 mM β-glycerophosphate (Sigma, St Louis, MO, USA) and in the absence (non-differentiating conditions) or presence (differentiating conditions) of 10^{-6} M dexamethasone (9α-fluoro-16α-methylprednisolone; Sigma). The medium was changed every 2–3 days. Human ghrelin was obtained from Neosystems (Strasbourg, France) and human UAG was kindly provided by Theratechnologies (Montreal, Canada).

DNA content

For DNA measurements 100 µl SV-HFO cell lysates were treated with 200 µl heparin (8 IU/ml in PBS) and 100 µl RNase A (50 µg/ml in PBS) for 30 min at 37°C. This was followed by adding 100 µl ethidium bromide solution (25 µg/ml in PBS). Samples were analyzed on the Wallac 1420 Victor2 (Perkin Elmer, Wellesley, MA, USA) using a 340 nm excitation filter and a 590 nm emission filter. Calf thymus DNA (Sigma) was used for standards.

Alkaline phosphatase activity

Alkaline phosphatase activity was assayed by determining the release of p-nitrophenol from p-nitrophenylphosphate (20 mM in 1 M diethanolamine buffer supplemented with 1 mM MgCl_{2} at pH 9·8) in the SV-HFO cell lysates for 10 min at 37°C. The reaction was stopped by adding 0·06 M NaOH. Absorption was measured at 405 nm. Results were adjusted for DNA content of the corresponding cell lysates.

Mineralization

SV-HFO cell lysates were incubated overnight in 0·24 M HCl at 4°C. Calcium content was determined colorimetrically with a calcium assay kit (Sigma) according to
the manufacturer’s description. Results were adjusted for DNA content of the corresponding cell lysates.

RNA isolation, cDNA synthesis and real-time PCR

Total RNA from bone biopsies and cultured osteoblasts was isolated using RNA-Beo solution (Tel-Test, Friendswood, TX, USA). mRNA was subsequently isolated from total RNA using the Oligotex Direct mRNA mini kit (Qiagen). RNA was quantified using a Ribogreen assay (Molecular Probes, Eugene, OR, USA). One µg total RNA was reverse transcribed into cDNA, using 0.5 µg oligo(dT)18, 0.2 µg random hexamer primers and Moloney murine leukemia virus reverse transcriptase according to the protocol of the manufacturer (MBI Fermentas, St Leon-Rot, Germany). Five ng (biopsies) or 20 ng (differentiating and non-differentiating SV-HFO cells at 7, 14 and 21 days of culture) of cDNA was then assayed for ghrelin, GHS-R1a and GHS-R1b genes using one-step TaqMan real-time PCR with an ABI Prism 7700 sequence-detection system (PE Biosystems, Rotkreuz, Switzerland). The strategy for distinguishing GHS-R1a and GHS-R1b splice variants and PCR assay conditions were similar to those described by Gnanapavan et al. (2002). The presence of GHS-R1b mRNA in osteoblasts was also confirmed using cDNA derived from poly(A)+ RNA samples from all culture time points with and without dexamethasone treatment. Reactions without reverse transcriptase were included to check for genomic DNA contamination. Since GHS-R1a was not detected in total RNA, SV-HFO mRNA was reverse transcribed and assayed for GHS-R1a gene expression by real-time PCR using alternative primers and a locked nucleic acid (LNA) probe (Exiqon A/S, Vedbaek, Denmark). Samples were also processed without reverse transcriptase as negative controls. Samples were assayed in duplicate and corrected for GAPDH mRNA. The data presented for ghrelin and GHS-R gene expression in SV-HFO cells represent analyses of RNA from four different, independent experiments of two cultures each. Primer and probe sequences were as follows: ghrelin, forward 5'-GGGCAGAGATGAACTGGGAA-3', reverse 5'-CCCTGGCTGCTGCTGTA-3', probe 5'-FAM-TCCGGTTCACCCGAC-TAMRA-3'; GHS-R1a, forward 5'-ACACGACACCAACACCC-3', reverse 5'-AGGTTTCGAATCCCAGAAGT-3', probe 5'-FAM-CTGCTGCTCCCTTCCAG-TAMRA-3'; GHS-R1b, forward 5'-CTGCTGGTGGCTGCTGTA-3', reverse 5'-GCTGAGACCCACCCAGCA-3', probe 5'-AGGGA CCAGAACCACAAGCAACCCG-3'; GAPDH, forward 5'-ATGGGGATGGTAGTGAC-3', reverse 5'-TAA AAGCGGCCCTGGTACC-3', probe 5'-FAM-CGCC CAATACGACACACATCCGGTACC-TAMRA-3' (FAM: carboxyfluorescein; TAMRA: carboxytetramethylrhodamine). Sequences of LNA probe and primers: GHS-R1a LNA probe, 5'-CTGGCTCCTCC-3'; GHS-R1a, forward 5'-GCA AACCGTGAAATGCTG-3'; reverse 5'-AAATATCG CCCTACTGGGAA-3'. Spleen total RNA was used as a positive control for ghrelin and GHS-R1b expression, since it is known to express these genes (Gnanapavan et al. 2002). Pituitary total RNA was used as a positive control for GHS-R1a gene expression. Data presented are relative mRNA levels calculated as $2^{-\Delta Ct}$ where $\Delta Ct = Ct_{\text{gene of interest}} - Ct_{\text{GAPDH}}$ (Bustin & Nolan 2004). Samples in which no amplified product could be detected by cycle 40 were considered not to express the gene being assayed.

Ghrelin enzyme immunoassay

Human osteoblasts were cultured for 24 and 72 h in serum-free medium. Ghrelin was measured in the conditioned media using an enzyme immunoassay kit that measures both acylated and unacylated forms of human ghrelin (Total Ghrelin kit; Phoenix Europe, GMBH, Karlsruhe, Germany). Medium that had not been conditioned was also assayed using this kit to check the specificity of the assay in our hands, and the presence of exogenous ghrelin in the medium. Total protein was determined using the BioRad protein assay reagent (BioRad, Hercules, CA, USA).

Cell-growth experiments

Proliferation was examined by [3H]thymidine uptake measurements. SV-HFO cells were seeded into 96-well
plates as described above, in differentiating and non-differentiating conditions. After culture for 5, 12 or 19 days the cells were treated for 20 h with ghrelin or UAG at various concentrations in 10–27 replicate wells per treatment at day 5 (thymidine assay on day 6), and 6–10 replicate wells at days 12 and 19 (thymidine assays on days 13 and 20).

Experiments with extracellular-signal-regulated kinase (ERK) and phosphoinositide-3 kinase (PI3K) inhibitors were also started on day 5 (n=20). Cells were pre-incubated for 40 min with vehicle (0.1% DMSO; Sigma), 10 µM U0126 (Promega, Madison, WI, USA), 35 µM PD98059 or 100 nM wortmannin (Sigma). These concentrations were found to have no effect on basal thymidine uptake and are similar to those used in other studies involving osteoblasts (e.g. Grey et al. 2002, Jansen et al. 2004, Kim et al. 2005, Osyczka & Leboy 2005). These initial treatments were replaced with media containing vehicle or inhibitor, as appropriate, with or without ghrelin or UAG. Cells were then incubated for 20 h. Subsequently (culture day 6), [3H]thymidine (Amersham, Little Chalfont, Bucks, UK) was added to a concentration of 1 µCi/ml and incubated for a further 7 h. Cells were then harvested onto glass-fibre mats (Tomtec-Harvester 96; Hamden, CT, USA) and radioactivity measured with a Wallac MicroBeta scintillation counter.

Figure 2 GHS-R1b mRNA expression in human bone. (A) Relative quantitation of GHS-R1b mRNA (see the Materials and Methods section) in human osteoarthritic bone biopsy samples compared with that in human spleen. (B) Relative expression of GHS-R1b mRNAs in non-differentiating and differentiating SV-HFO osteoblasts. (a, P<0.005 versus differentiating cells at day 7 (d7)). (C) Relative quantitation of ghrelin mRNA in human osteoarthritic bone biopsy samples compared with that in human spleen. (D) Relative expression of ghrelin mRNAs in non-differentiating and differentiating SV-HFO osteoblasts. Bone 1 and bone 2 refer to samples from two different subjects. SV-HFO data are derived from four separate experiments and are corrected for GAPDH mRNA expression (see the Materials and Methods section). GHS-R1a mRNA was undetectable in these samples. Functionality of the GHS-R1a TaqMan primer/probe set was confirmed using human pituitary RNA (see the Results section).

ERK phosphorylation assay

ERK activation was determined using a fast activated cell enzyme-linked immunosorbent assay kit (FACE; Active-Motif, Carlsbad, CA, USA). Briefly, osteoblasts were seeded into 96-well plates at a density of 10 000 cells/cm² as described above, in non-differentiating medium. At day 6, cells were treated with medium lacking or containing 10 nM ghrelin or 100 nM UAG for 5 min. The cells were then fixed and incubated with specific antiphosphorylated ERK antibodies followed by incubation with a secondary horseradish peroxidase-conjugated antibody. The levels of phosphorylated ERK were quantified using a colorimetric readout and are expressed as the absorbance at 450 nm measured in each well.

Statistical analysis

Statistical analyses were performed with StatView5 for Macintosh (SAS Institute, Cary, NC, USA). Results were tested for statistical significance using analysis of variance (ANOVA), followed by Fisher’s protected least-significant difference post-hoc test. Data are presented as means ± S.E.M.

Results

Osteoblast differentiation

During SV-HFO cell culture, osteoblast differentiation was monitored by measuring DNA levels, alkaline phosphatase activity and mineralization. In the absence of dexamethasone alkaline phosphatase activity remained very low throughout the whole 23-day culture period. In contrast, cultures that were treated continuously with dexamethasone showed an increase of alkaline phosphatase activity, which peaked around day 14 (Fig. 1). In these dexamethasone-treated cultures mineralization was initiated around day 14 of culture and increased further up to day 19, after which it levelled off (Fig. 1). In the absence of dexamethasone, cultures showed no evidence of osteoblast differentiation and mineralization. Dexamethasone-treated and non-dexamethasone-treated cell cultures will be referred to as differentiating and non-differentiating osteoblasts, respectively.

Expression of the GHS-R splice variants and ghrelin in human bone cells

Like an earlier study of human bone marrow using RNase protection assay with poly(A)⁺ RNA (Guan et al. 1997), GHS-R1a mRNA was not detected in the bone samples, although identical TaqMan assays of human pituitary RNA produced positive amplification data (GAPDH ratio of 1200–1500) from 20 ng cDNA; see Materials and Methods section). However, GHS-R1b mRNA was detected in femoral bone (Fig. 2A) at 200–450% of its level of expression in spleen.

Bone biopsies probably contain a mixed population of cell types including osteoblasts. Therefore, to clarify whether the GHS-R subtype mRNAs are expressed
in osteoblasts, we measured their expression by real-time PCR in cultured human osteoblasts. Like the biopsies, no GHS-R1a mRNA was detectable in total RNA from human osteoblast cultures at any of the time points examined. To examine this negative observation further we ran real-time PCRs of cDNAs derived from mRNA using an alternative primer set and an LNA probe. Again, no GHS-R1a mRNA expression was detected. In contrast, human osteoblasts expressed GHS-R1b mRNA and no GHS-R1a mRNA expression was detected. In controls, reaching maxima at 1 nM in both non-differentiating or non-differentiating conditions and then thymidine uptake was measured to determine their rate of proliferation.

Thymidine uptake was significantly and dose-dependently stimulated by ghrelin relative to untreated controls, reaching maxima at 1 nM in both non-differentiating (170% of untreated controls, \( P<0.001 \); Fig. 3A) and differentiating (155% of untreated controls, \( P<0.001 \); Fig. 3B) cultures. Ghrelin (10 nM) was also confirmed to increase cellular DNA content to 144% (\( P<0.007 \)) and 119% (\( P<0.02 \)) of control levels in non-differentiating and differentiating cultures after 72 h.

We next examined whether the proliferative effect of ghrelin on the SV-HFO cells was dependent on culture time and whether or not the cells had been induced to differentiate. In non-differentiating cells the proliferative response declined with time in culture, with a decline in response at day 13 and eventually leading to loss of a significant effect of ghrelin detectable during the third week. In differentiating osteoblasts, the response to ghrelin disappears within 2 weeks of culture, with no significant effect of ghrelin detectable during the third week. (B) In differentiating osteoblasts the response to ghrelin is lost by day 13.
UAG also had a marked dose-dependent inductive effect on thymidine uptake in both non-differentiating and differentiating cells at day 6 of culture (Fig. 5). UAG had maximal effect at 10 nM (180% of controls in non-differentiating cells \(P=0.0003\)) and 195% of controls in differentiating cells \(P=0.0003\)). Furthermore, the response to UAG declines with time in culture and with differentiation in a way that is comparable to the ghrelin response, suggesting that the mechanism of response is similar, if not identical (Fig. 6). Studies on alkaline phosphatase activity and mineralization during the 3-week culture period demonstrated that neither ghrelin nor UAG affect these two processes (data not shown).

**Signaling pathways for ghrelin and UAG proliferative activity**

Ghrelin in particular, but also UAG, have been shown to activate mitogen-activated protein kinase (MAPK) and PI3K pathways in other cell types (e.g. Baldanzi et al. 2002, Kim et al. 2004, Nanzer et al. 2004). ERK and PI3K are important components of signalling pathways that are often directly involved in proliferative responses of cells to growth factors. Therefore, the possible involvement of these pathways was investigated by utilizing specific inhibitors of ERK activation (U0126 and PD98059) and PI3K activity (wortmannin) at concentrations that had no significant effect on basal cell proliferation. Since the most potent effects of ghrelin and UAG on proliferation occurred in non-differentiating osteoblasts we tested the inhibitors under these conditions using the peptides at concentrations that had been confirmed to induce proliferation. All three inhibitors blocked or suppressed the proliferative response of human osteoblasts to ghrelin (Fig. 7A) and UAG (Fig. 7B).

To further confirm the involvement of the MAPK signalling pathway in the proliferative effects of ghrelin and UAG we examined the acute effects of these peptides on the phosphorylation of ERK by ELISA. We found that ghrelin and UAG (10 and 100 nM, respectively) significantly stimulated the phosphorylation of ERK relative to their untreated controls \(123\%_p\ 8.6\) and \(124\%_p\ 9.1\%\, \text{respectively; } P<0.05, \text{by ANOVA}\) in serum-starved human osteoblasts within 5 min of treatment (Fig. 8).

**Discussion**

The current study demonstrates ghrelin gene expression in human bone tissue, suggesting that, like rodents, ghrelin may have auto-/paracrine activity in human bone, in addition to the endocrine activity of gastric-derived hormone. However, the absence of GHS-R1a suggests that an auto-/paracrine effect might be mediated by an alternative receptor in human bone. The presence of a non-type-1a GHS-R is also suggested by the response of the cells to UAG. In rat osteoblasts, GHS-R1a protein was detected and thus may mediate the proliferative effects of ghrelin observed in these cells (Fukushima et al. 2005, Maccarinelli et al. 2005). This observation suggests species
Ghrelin and unacylated ghrelin induce osteoblast growth

The proliferative effect of UAG, like ghrelin, is dependent on time in culture and state of differentiation. Cells were treated for 20 h with UAG, starting on days 5, 6, 13 and 20. (A) In non-differentiating cells the effect of UAG on thymidine uptake generally declines with time in culture having only approximately 50% of its effect at all doses at week two, and with no significant effect detectable during the third week. (B) In differentiating osteoblasts the response to UAG disappears after the first week of culture (a, P<0·05; b, P<0·005; c, P<0·0005 versus 0 nM controls on the same day).

![Graph A](Image)

![Graph B](Image)

Figure 6 The proliferative effect of UAG, like ghrelin, is dependent on time in culture and state of differentiation. Cells were treated for 20 h with UAG, starting on days 5 (n=10–27 replicates per treatment), 12 or 19 (n=6–10 replicates per treatment), then thymidine uptake was measured following 7 h incubation on days (d) 6, 13 and 20. (A) In non-differentiating cells the effect of UAG on thymidine uptake generally declines with time in culture having only approximately 50% of its effect at all doses at week two, and with no significant effect detectable during the third week. (B) In differentiating osteoblasts the response to UAG disappears after the first week of culture (a, P<0·05; b, P<0·005; c, P<0·0005 versus 0 nM controls on the same day).

Differences between rodent and human bone in the expression of the cognate receptor.

The expression in human bone biopsies and the changes in GHS-R1b mRNA are intriguing. GHS-R1b is almost identical to GHS-R1a but is truncated after transmembrane domain 5, and current evidence demonstrates that this receptor does not stimulate an intracellular signalling pathway (Howard et al. 1996, Feighner et al. 1998). Mutational analysis of the GHS-R suggests that the ligand-binding site (at least of synthetic GHS) involves residues D99, C116 and E124, which are located in transmembrane domains 2 and 3 and extracellular loop 1 (Feighner et al. 1998). So it is conceivable that GHS-R1b can either bind ligand or modify interaction of the ligand with the cognate receptor. There is some evidence for the latter where overexpression of GHS-R1b appears to antagonize the response of GHS-R1a to synthetic GH secretagogues (Chan & Cheng 2004).

An explanation for loss of a proliferative response to both peptides with time in culture and state of differentiation is lacking. It could be related to differentiation-related changes in receptor signalling pathways. In this respect it is intriguing that there appears to be an inverse relationship between the loss of proliferative response to ghrelin and the increased expression of GHS-R1b. We found that GHS-R1b gene expression increased particularly in the last week of culture, and was lower in non-differentiating than differentiating cells at this time. A similar stimulatory effect of glucocorticoids on GHS-R gene expression has been demonstrated in the pituitaries of rats (Tamura et al. 2000). It is tempting to speculate that the increase in GHS-R1b during dexamethasone-induced differentiation may correspond with an inhibitory effect on ghrelin-induced cell growth.

From the current data it is clear that ghrelin is expressed by osteoblasts. Moreover, ghrelin and UAG are potent stimulators of proliferation in human osteoblasts, being effective in the nanomolar range. Inhibition of these proliferative effects by inhibitors of ERK and PI3K, and the activation of the MAPK pathway through phosphorylation of ERK, suggests the requirement for these pathways in both ghrelin and UAG signalling in human osteoblasts, and indicates a similar mechanism to that described for cardiomycocytes (Baldanzi et al. 2002) and hepatoma cells (Murata et al. 2002). However, the mechanism by which the proliferative effects of ghrelin and UAG on osteoblasts are initiated is unclear since GHS-R1a mRNA is not expressed at detectable levels in these cells, and GHS-R1b has been demonstrated to be non-functional, at least through activation of a Ca$^{2+}$ pathway (Howard et al. 1996). Recently, it has been demonstrated that UAG stimulates adipogenesis in vivo, and that ghrelin has similar effects that appear to be independent of GHS-R1a (Thompson et al. 2003). However, the mechanism, or receptor, which mediates the response to these peptides remains unknown. The involvement of both MAPK and PI3K pathways does not exclude the possibility that receptor types other than a G-protein-coupled receptor are involved in transducing the response. It is also possible that an indirect mechanism is involved, as has been suggested for prostate cells, where cell proliferation could be mediated via an auto-/paracrine pathway involving other growth factors (Jeffery et al. 2003). Even so, a receptor for ghrelin and UAG would still be necessary to stimulate such an intermediate mechanism. Therefore, it will be a challenge to identify a new receptor for ghrelin in peripheral tissues and to define its mode of action.

Indirect evidence for our hypothesis that a receptor other than GHS-R1a may be involved in bone metabolism is the finding that GHS-R1a homozygous knockout mice do not have a significantly altered bone phenotype.
(Sun et al. 2004), as assessed by bone mineral density (BMD) and bone mineral content (BMC). Our findings suggest that ghrelin only has significant proliferative effects early on in osteoblast cultures rather than direct effects on osteoblast differentiation. Therefore, it is possible that in vivo ghrelin acts primarily on the proliferation of osteoblast progenitors and/or preosteoblasts rather than mature osteoblasts. However, the lack of phenotype in the gene-deleted animals may be linked to redundancy in regulatory mechanisms that operate during this period of osteoblast development. Similar compensatory mechanisms may be at work in the ghrelin-knockout mice, which also have no obvious bone phenotype, again as assessed only by BMD and BMC (Sun et al. 2003). Compensation for the loss of ghrelin in these mice is evidenced by the lack of effect on appetite, food intake and body composition (Sun et al. 2003, 2004, Wortley et al. 2004). However, BMD and BMC are relatively crude markers of bone turnover, and it would be of interest to examine in greater detail bone morphology and markers of bone turnover in these animals.

The current study clearly reveals that ghrelin is produced by human bone cells and that ghrelin and UAG stimulate proliferation of human osteoblasts via a mechanism that involves activity of the MAPK and PI3K signalling pathways. These data uncover a new regulatory mechanism for bone metabolism in humans. The potent effects of UAG suggest a new, as yet undefined, receptor/signalling pathway in bone.

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