Metabolic Acidosis Increases Intracellular Calcium in Bone Cells Through Activation of the Proton Receptor OGR1

Kevin K Frick, Nancy S Krieger, Keith Nehrke, and David A Bushinsky

ABSTRACT: Metabolic acidosis increases urine Ca without increasing intestinal absorption, leading to bone Ca loss. It is unclear how bone cells detect the increase in proton concentration. To determine which G protein–coupled proton sensing receptors are expressed in bone, PCR was performed, and products were detected for OGR1, TDAG8, G2A, and GPR4. We tested the hypothesis that the G protein–coupled proton sensor, OGR1, is an H⁺-sensing receptor in bone. To determine whether acid-induced bone resorption involves OGR1, we incubated mouse calvariae in neutral pH (NTL) or acidic (MET) medium ± the OGR1 inhibitor CuCl₂. CuCl₂ decreased MET-induced Ca efflux. We used fluorescent imaging of perfused bone cells to determine whether MET increases Caᵢ. Perfusion with MET induced a rapid, flow-independent, increase in Caᵢ in individual bone cells. To determine whether transfection of OGR1 into a heterologous cell type would increase Caᵢ in response to H⁺, we perfused Chinese hamster ovary (CHO) cells transfected with mouse OGR1 cDNA. Perfusion with MET induced a rapid increase in Caᵢ in OGR1-transfected CHO cells. These data indicate that OGR1 induces an increase in Caᵢ in response to MET and is a prime candidate for an osteoblast proton sensor.

J Bone Miner Res 2009;24:305–313. Published online on October 13, 2008; doi: 10.1359/JBMR.081015

Key words: bone, acidosis, proton, intracellular calcium, bone resorption, osteoblast

INTRODUCTION

CHRONIC METABOLIC ACIDOSIS alters calcium (Ca) homeostasis by increasing urine Ca excretion without a corresponding increase in intestinal Ca absorption, leading to depletion of the primary Ca reservoir in the body, the skeleton. In vivo, mild metabolic acidosis, induced by ample dietary protein intake and an age-related decline in acid excretion, may deplete bone Ca stores. This process can be modeled in vitro by culturing neonatal mouse bone in physiologically acidic medium. We have found that, in the earliest time periods (≤24 h), bone calcium efflux is directly related to the magnitude of the fall in pH and caused by non–cell-mediated physicochemical mineral dissolution. In contrast, at later time periods (>24 h), the continued calcium efflux is caused by cell-mediated bone resorption. Metabolic acidosis regulates both osteoblastic and osteoclastic activity. Acidosis decreases expression of osteoblastic extracellular matrix proteins, including type 1 collagen, osteopontin, and matrix gla protein, and increases expression of COX-2 and RANKL. Acidosis also increases secretion of osteoclastic β-glucuronidase.

How the bone cells sense the changes in extracellular pH (proton concentration, [H⁺]) is not clear. Various families of acid-sensitive ion channels including the ASICs, TRPV1, and TASK are known to be expressed in pain sensing, skeletal muscle, and taste cells. In addition, there is a family of G protein–coupled proton-sensing receptors. This small family (40–50% shared homology) of G protein–coupled receptors includes the ovarian cancer G protein–coupled receptor 1, OGR1 (also called GPR68), GPR4, T-cell death–associated gene 8 (TDAG8, also called GPR65), and G2A (also called GPR132). These receptors are coupled either to phosphoinositide metabolism and increased Caᵢ (OGR1 and G2A) or alteration in adenylate cyclase activity (GPR4 and TDAG8). Ludwig et al. reported that OGR1 is expressed in mouse osteoblasts and that reduction in pH (increases in [H⁺]) leads to an accumulation of phosphoinositide metabolites, making OGR1 a prime candidate for an osteoblastic proton sensor.

To test the hypothesis that OGR1 acts as an H⁺-sensing receptor in bone cells, we studied whether an inhibitor of OGR1 (CuCl₂) would diminish acidosis-induced Ca efflux from bone, whether metabolic acidosis would increase Caᵢ in cultured bone cells, and whether transfection of OGR1 into a heterologous cell type would permit cells to mimic the Caᵢ response to acidosis of primary bone cells. We found that CuCl₂ inhibits the acidosis-induced increase in Ca efflux from bone cells and that primary murine calvarial bone cells respond to a decrease in extracellular pH with an increase in Caᵢ. We found that Chinese hamster ovary fibroblasts (CHO cells) transfected with murine OGR1...
cDNA now respond to acidosis with increased Ca$. These data are consistent with a primary role for OGR1 as a sensor for H$ in bone cells.

MATERIALS AND METHODS

Reagents

All reagents, unless otherwise specified, were obtained from Sigma-Aldrich (St Louis, MO, USA).

Animals and cells

CD-1 mice were obtained from Charles River (Wilmington, MA, USA). CHO fibroblasts were a gift from Patricia Hinkle (University of Rochester, Rochester, NY, USA) and grown in αMEM (Invitrogen, Carlsbad, CA, USA) containing 5% heat-inactivated fetal bovine serum (FBS; Invitrogen) and penicillin (100 U/ml; Sigma-Aldrich).

RNA isolation and PCR

RNA was isolated from primary calvarial cells using a Qiagen RNeasy kit (Valencia, CA, USA) as previously described. Total RNA (1 μg) was reverse transcribed to first-strand cDNA using an iScript cDNA synthesis kit (Biorad, Hercules, CA, USA) and amplified by PCR using iQ SYBR-green in an iCycler thermocycler (Biorad).

PCR primers were synthesized by Integrated DNA Technologies (Coralville, IA, USA) and were as follows: OGR1 (forward, 5′-CCTCAACCTGTTCGGACGTGC; reverse, 5′-CCACATATCACTCTCCGGGTCTC); G2A (forward, 5′-GACATGGATGCGGTGTGTC; reverse, 5′-CCCAAGCGTGACTCAAGGAC); GPR4 (forward, 5′-ACCGAGGGCCAGGAAAGT; reverse, 5′-GGAGGACCTGCCCACAGGC); and TDAG8 (forward, 5′-CTTCAACCTGTTCGGACGTGC; reverse, 5′-CCACATATCACTCTCCGGGTCTC). PCR products were resolved by electrophoresis in agarose (Invitrogen) and migration rate compared with a size standard (1 kb Plus DNA Ladder; Invitrogen).

Effect of CuCl$_2$ on bone calcium efflux

Calvariae from 4- to 6-day-old CD-1 mice were dissected and incubated in 2.8 ml DMEM (Lonza, Walkersville, MD, USA) containing 15% heat-inactivated horse serum (Invitrogen), heparin (10 USP units/ml; Baxter), and penicillin (100 U/ml) in 35-mm dishes at neutral pH (NTL, pH ≈ 7.4) or at a physiologically acidic pH (MET, pH ≈ 7.1) produced by a primary reduction in [HCO$_3^-$] as a model of metabolic acidosis as previously reported. The partial pressure of medium CO$_2$ (PCO$_2$) was maintained at the physiologic normal of ~40 mmHg. To closely replicate physiological conditions, only the HCO$_3^-$/CO$_2$ buffer system was used; no other buffers were added to the medium. All medium was preincubated at PCO$_2$ ~40 mmHg, 37°C, for at least 3 h before use. The number of pairs of bones chosen in each group was determined by power analysis using actual means and SD from initial experiments with an n = 4. We assumed probability of a type I error (α) = 0.05 and the probability of a type II error (β) = 0.10. We used Statistica (version 6.0; StatSoft, Tulsa, OK, USA). For a statistical power (1 – β) goal of 0.9, we needed eight pairs of bones in each group. In the protocols above, we used 10–12 pairs of bones in each group.

Primary bone cell culture

Primary bone cells, which are almost exclusively osteoblasts, were isolated from neonatal CD-1 mouse calvariae immediately after dissection as described previously. Briefly, bones were washed in PBS containing 4 mM EDTA for 10 min at 37°C and incubated in HEPES (25 mM), pH = 7.4, containing 2 mg/ml collagenase (Wako Pure Chemicals, Dallas, TX, USA) and 90 μM Nα-tosyl-l-lysyl chloromethyl ketone for three sequential 20-min digestion periods at 37°C in a shaking water bath. At the end of each digestion, released cells were collected and resuspended in HEPES buffer containing 1 mM MgSO$_4$ and all three digests were pooled in DMEM (Lonza, Walkersville, MD, USA) containing 15% heat-inactivated horse serum (Invitrogen), heparin (10 USP units/ml; Baxter), and penicillin (100 U/ml) for plating on 8-mm glass coverslips (Warner Instruments, Hamden, CT, USA). Coverslips were used for imaging 1–5 days after initial plating.

Cell imaging

To reduce background, DMEM without phenol red (Invitrogen) was used for perfusion while imaging. To closely replicate physiological conditions, only the HCO$_3^-$/CO$_2$ buffer system was used; no other buffers were added to the medium. The acidity of the medium was adjusted by a primary reduction in [HCO$_3^-$] as a model of metabolic acidosis as previously reported. The partial pressure of medium CO$_2$ (PCO$_2$) was maintained by gentle bubbling with 5% CO$_2$ in air (AirGas, Rochester, NY, USA) for at least 15 min before perfusion. A continuous flow of equilibrated medium was established through an open microscope chamber (RC-26; Warner Instruments; ~200-μl volume) by use of a peristaltic pump (~1 ml/min) and aspiration. To reduce gas loss, where possible, Viton (DuPont, Wilmington, DE, USA) or fluorinated ethylene-propylene (FEP) tubing was used. Flow control was accomplished with a manually operated perfusion valve system (V-6; Warner Instruments). To load cells on coverslips with fura-PE3 (TefLabs, Austin, TX, USA), coverslips were washed with ice-cold PBS and incubated for 45 min at 37°C in DMEM containing 4.4 μM fura-PE3 and 48 μM pluronic acid F127 (Invitrogen). Coverslips were mounted in the chamber with flow of neutral medium. For stimulation with acidic
Intracellular Ca (Ca_i) levels were measured from fluo-}

Tokyo, Japan) equipped with a CCD camera (Cooke,}

Stable transfection of CHO cells

Expression of H^+-sensing receptors in bone cells

Effect of CuCl2 on bone resorption

Measurement of Ca_i in primary bone cells

Because OGR1 has been shown to be coupled to phospholipase C, leading to increased IP_3 production and subsequent increases in intracellular Ca (Ca_i), we next determined whether Ca_i was altered in response to a physiologic reduction in pH. Changes in Ca_i were measured by loading primary bone cells with Fura-PE3 and...
measuring the ratio of fluorescent emissions at 510 nm after excitation at 340 and 380 nm. The bone cells were continuously perfused with neutral (NTL, pH = 7.40, PCO2 = 47 mmHg, [HCO3-] = 29 mM) medium, which did not lead to a significant alteration of Ca_i (Fig. 3A).

Cells were perfused with NTL followed by MET. Again, perfusion of the cells with NTL (pH = 7.42 ± 0.02, PCO2 = 44.4 ± 1.2 mmHg, [HCO3-] = 28.2 ± 1.0 mM) did not change Ca_i (Fig. 3B, 0–180 s). In contrast, when the cells were perfused with MET (pH = 6.90 ± 0.03, PCO2 = 41.8 ± 1.7 mmHg, [HCO3-] = 8.2 ± 0.8 mM), there was a substantial, rapid increase in Ca_i in multiple cells (Fig. 3B, 182–600 s). The increase in Ca_i in response to MET was asynchronous, occurred in individual cells at various times after the initial exposure to MET, and varied in magnitude and duration. A pseudocolor representation of selected fields among those analyzed for Fig. 3B is shown in Fig. 3C. The first frame (C1) indicates the initial time (time = 0 s), and the latter three frames show representative responses to acidification at 198, 221, and 288 s for C2, C3, and C4, respectively. Arrows indicate cells whose relative fluorescence intensity is shown in the corresponding traces in D1–D4. Cells shown in C1–C4 and corresponding traces D1–D4 show various patterns of response, with some cells displaying single peaks, whereas others exhibit more complex responses.

Changes in Ca_i were quantified by two methods: the distribution of cell number by the peak to nadir ratio and the percentage of cells with an increase in Ca_i above specific threshold values.

The ratio of the maximum to minimum Ca_i for each cell during the initial NTL and MET phases of perfusion was determined for all 300 time points for each cell and the distribution of values plotted. The distribution of the peak to nadir ratio for cells during the MET phase of perfusion was significantly different than that for the NTL phase of perfusion with more cells in MET having a greater peak to nadir ratio (n = 2306 cells in each group; Fig. 4A) and the mean value of the ratio (inset) was significantly increased with MET.

The percentage of cells with an increase in Ca_i above a specific threshold (1.1) was determined. A peak in cell Ca_i was defined as a >1.1 increase in area under the curve of Ca_i for each 2-s time interval compared with the mean area under the curve for the previous five time intervals for that cell. This calculation was repeated for threshold values of 1.2, 1.3, 1.4, and 1.5. Compared with the initial NTL phase of imaging, there was a significant increase in the percentage of cells with Ca_i peaks >1.1, >1.2, >1.3, and >1.4 during the MET phase of imaging (n = 2306 cells individually analyzed; Fig. 4B).

Measurement of Ca_i in OGR1-transfected CHO cells

We next tested the hypothesis that expression of OGR1 in CHO cells would lead to a MET-induced increase in Ca_i. Nontransfected CHO cells do not increase Ca_i in response to MET (pH = 7.02, PCO2 = 52 mmHg, [HCO3-] = 13 mM) compared with NTL (pH = 7.37, PCO2 = 51 mmHg,
[HCO$_3^-$] = 29 mM; $n = 1234$ cells in each group, % cells with peaks $>1.2$: NTL = 4.1 ± 2 versus MET = 2.7 ± 0.7, $p$ = not significant; data not shown).

CHO cells were transfected with a plasmid encoding full-length mouse OGR1 cDNA and selected for antibiotic resistance. Stable lines were screened for expression of a V5 epitope encoded by the expression plasmid; up to $\sim$30% of cells were positive for V5 expression in various lines, and the three maximally expressing lines were chosen for further study. The responses of these three lines were not different, and data for these lines were combined.

To determine whether Ca$_i$ was altered in response to a physiologic reduction in pH, OGR1-transfected CHO cells were loaded with fura-PE3, and Ca$_i$ was determined during continuous perfusion of NTL followed by MET medium.

The initial NTL (pH = 7.37 ± 0.02, PCO$_2$ = 46.6 ± 0.7 mmHg, [HCO$_3^-$] = 26.6 ± 1.6 mM) phase of perfusion again did not lead to alteration of Ca$_i$ (Fig. 5A, 0–180 s). Perfusion with MET (pH = 6.90 ± 0.03, PCO$_2$ = 43.2 ± 1.3 mmHg, [HCO$_3^-$] = 8.2 ± 1.0 mM) led to a rapid increase in Ca$_i$ in multiple cells (Fig. 5A, 182–600 s). The increase in Ca$_i$ in response to MET was again asynchronous; the increase in Ca$_i$ in individual cells occurred at various times after the initial exposure to MET and varied in magnitude and duration. Pseudocolor representations of selected fields among those analyzed for Fig. 5A are shown in Fig. 5B. The first frame (B1) indicates the initial time and the latter three frames show representative responses to acidification at 250, 280, and 300 s for B2, B3, and B4, respectively. Arrows indicate cells whose relative fluorescence intensity are shown in the corresponding traces in C1–C4. The cell indicated in B1 and C1 did not respond to acid. Cells shown in B2–B4 and corresponding traces C2–C4 illustrate various patterns of response, with some cells displaying single peaks, whereas others exhibit more complex responses. The period of time before changes in Ca$_i$ were observed after MET reached the perfusion chamber was shorter than with primary bone cells (compare time to initial peaks in Fig. 5A with Fig. 3B).

Again, changes in Ca$_i$ were quantified by two methods: the distribution of cell number by the peak to nadir ratio and the percentage of cells with an increase in Ca$_i$ above specific threshold values.

The ratio of the maximum to minimum Ca$_i$ for each cell during the initial NTL and MET phases of perfusion was determined and the distribution of values plotted. The distributions were significantly different, $p < 0.01$. Inset, mean ± SE for all NTL and all MET cells. (B) The percentage of cells with an increase in Ca$_i$ above each specified threshold. The peak in cell Ca$_i$ was defined as an increase in Ca$_i$ of greater than the specified threshold (1.1–1.5) over the mean of the previous five time points (time point taken every 2 s) for that cell. $n = 2306$ cells individually analyzed for A and B; *$p < 0.05$. 

**FIG. 4.** Quantitation of Ca$_i$ in primary bone cells. Changes in Ca$_i$ were quantified by two methods: (A) the distribution of cell number as a function of the peak to nadir ratio and (B) the percentage of cells with an increase in Ca$_i$ above specific threshold values. (A) The ratio of the maximum to minimum Ca$_i$ for each cell during the NTL and MET phases of perfusion was determined and the distribution of values plotted. The distributions were significantly different, $p < 0.01$. Inset, mean ± SE for all NTL and all MET cells. (B) The percentage of cells with an increase in Ca$_i$ above each specified threshold. The peak in cell Ca$_i$ was defined as an increase in Ca$_i$ of greater than the specified threshold (1.1–1.5) over the mean of the previous five time points (time point taken every 2 s) for that cell. $n = 2306$ cells individually analyzed for A and B; *$p < 0.05$. 

[ACIDOSIS INCREASES INTRACELLULAR CA THROUGH OGR1 309]
point taken every 2 s) for that cell. This calculation was repeated for threshold values of 1.2, 1.3, 1.4, and 1.5. Compared with the initial NTL phase of imaging, there was a significant increase in the percentage of cells with $Cai > 1.1, > 1.2, > 1.3, > 1.4,$ and $> 1.5$ in the MET phase of imaging ($n = 4758$ cells individually analyzed; Fig. 6B).

**DISCUSSION**

An acidic extracellular pH increases osteoclastic bone resorption through an increase in osteoblastic COX-2, leading to an increase in RANKL and subsequent increase in net Ca efflux from bone. A class of $H^+$-sensing G protein–coupled receptors, whose activity is mediated by release of the second messengers $Ca_i$ or cAMP, has now been identified. Stimulation of these $H^+$ receptors is an attractive mechanism by which bone cells could sense and initiate the response to acidosis. In this study, we confirmed the expression of each of four known proton receptors OGR1, TDAG8, GPR4, and G2A by RT-PCR of RNA from primary bone cells (Fig. 1).

Because CuCl$_2$ binds to essential histidine residues in OGR1, we next showed that this compound significantly decreased acid-induced bone net Ca efflux, a marker of bone resorption, in cultured neonatal mouse calvariae. To our knowledge, the specificity of CuCl$_2$ for proton receptors other than OGR1 has not be reported. The four histidine residues in OGR1 determined by Ludwig et al. to be essential for acid induction of inositol phosphate formation are conserved in GPR4. On the basis of primary structure, the positioning of histidine residues in TDAG8 and G2A differs from OGR1. These structural features suggest that GPR4 might also be sensitive to CuCl$_2$, whereas TDAG8 and G2A would be less susceptible; however, such predictions would require experimental testing.

The observed suppression of Ca release by CuCl$_2$ in neutral conditions may indicate that OGR1 or similar receptors also have a role in establishing the basal rate of Ca release.

**FIG. 5.** Imaging of $Cai$ in OGR1-transfected CHO cells. CHO cells transfected with mouse OGR1 on coverslips were loaded with fura-PE3 and perfused with physiologically neutral (NTL) or acid (MET) medium. $Cai$ levels were determined from fluorescent emissions at 510 nm after excitation at 340 and 380 nm using a dual wavelength excitation system. (A) Perfusion with NTL (0–180 s) followed by MET (182–600 s). Arrow, time of chamber acidification. Relative fluorescence as a function of time for all cells is shown in B ($n = 255$ individual cells). (B) Pseudocolor images of relative fluorescence intensity; B1, time = 0; B2, time = 250 s; B3, time = 280 s; B4, time = 300 s. Arrows indicate selected cells whose change in fluorescence as a function of time is shown in C1–C4, respectively.

**FIG. 6.** Quantitation of $Cai$ in OGR1-transfected CHO cells. Changes in $Cai$ were quantified by two methods: (A) the distribution of cell number as a function of the peak to nadir ratio and (B) the percentage of cells with an increase in $Cai$ above each specified threshold. The peak in cell $Cai$ was defined as an increase in $Cai$ of greater than the specified threshold ($1.1–1.5$) over the mean of the previous five time points (time point taken every 2 s) for that cell ($n = 4758$ cells individually analyzed for A and B; $^{*}p < 0.05$).
resorption. OGR1 has been previously detected in osteoblasts and is coupled to phospholipase C and increases in diacylglycerol and inositol phosphate (IP) metabolism, leading to an increase in Ca$_i$. We found that a subset of cultured primary calvarial cells respond to acidosis pH with an increase in Ca$_i$. To further support OGR1 as a prime candidate as an osteoblastic H$^+$ sensor, we asked whether heterologous cells transfected with OGR1 would mount a similar Ca$_i$ response to acidosis as do primary bone cells. CHO cells were stably transfected with mouse OGR1 cDNA. When cultures of these transfecteds were perfused with acidic medium, there was a prompt increase in Ca$_i$ that was not present in nontransfected CHO cells. These data indicate that OGR1 is capable of transducing an increase in Ca$_i$ in response to MET and is a prime candidate as the osteoblast proton sensor.

To replicate, to the extent possible, physiologic conditions, in this study, only the HCO$_3^-/CO_2$ buffer system was used to adjust [H$^+$]; medium pH was adjusted by varying the [HCO$_3^-$] while maintaining a constant physiologic PCO$_2$. The open chamber used in the perfusion experiments permits adequate fluid flow while limiting fluid shear stress, which has been shown to alter Ca$_i$. We did not observe significant changes in Ca$_i$ during perfusion with neutral pH medium, indicating that there was no response to either flow or the neutral pH medium (Fig. 3A).

With primary bone cells, the time from when the cells were acidified to their maximal increase in Ca$_i$ and the magnitude of that increase were heterogeneous among the cells. Not every cell in the field showed a measurable increase in Ca$_i$ to acidosis. By visual examination, the responding cells were morphologically indistinguishable from cells that did not respond. It is not clear why some cells responded to an increase in [H$^+$] and others did not. Whereas the primary bone cells used in this study are almost exclusively osteoblasts, they are clearly a heterogeneous population at different levels of maturation. The ability of the cells to respond to acidosis with increased Ca$_i$ may be related to their level of maturation. While the use of more acidic medium (greater [H$^+$]) might have elicited a more robust response, we chose to use a perfusion pH within the pathophysiologic range (i.e., conditions that are compatible with life). With the OGR1-transfected CHO cells, cell-to-cell variations in the Ca$_i$ response to H$^+$ may be the result of differences in expression of OGR1.

This study does not exclude that other G protein–coupled H$^+$ sensors are involved in the response of bone to acidosis. The function of G2A as an H$^+$ receptor is controversial and may be related to specific cell type, although its activation would also lead to an increase in Ca$_i$. However, there seemed to be little expression of G2A in cultured bone cells (Fig. 1). In contrast, although apparently abundant in cultured bone cells, TDAG8 and GPR4 are coupled to adenylate cyclase and should not directly alter Ca$_i$. Further studies will be needed to elucidate the potential roles, if any, of other proton receptors such as TDAG8 and GPR4, whose effect is mediated through cAMP in the response of bone to acid.

Other proteins that modulate biological responses to changes in extracellular [H$^+$] include the pH-sensitive leak K$^+$ channel TASK, the vanilloid receptor TRPV1, and the proton-gated, voltage-insensitive sodium channels, the ASICs. Expression of TASK and TRPV1 by osteoblasts has not been reported, although osteoclasts express TRPV1. Human pre-osteoblasts express ASIC 1–4 and retain expression during mineralization. Whereas the ASICs are responsive to pH increments in the range of 0.05–0.06 units, their pH of half-maximal activation ranges from 6.5 to 4.4. In the experiments reported here, pH was reduced from 7.4 to 6.9, well above the region of maximal responsiveness. However, with ASIC3, significant changes in current can be induced by reducing the bath pH from 8.0 to 7.0, and it is possible that the pH range of the ASICs could be modulated by accessory proteins. Thus, we cannot exclude a role for ASICs in acid-induced bone resorption.

In this study we chose to focus on acid-induced changes in Ca$_i$. In addition to its effects on inositol phosphate and Ca$_i$, OGR1 in human aortic smooth muscle cells also activates cAMP formation. However, the response to H$^+$, as a function of the magnitude of the acidosis, is quite different for cAMP. Whereas IP accumulation is half-maximal at pH ~ 7.1, the cAMP effect is half-maximal at pH ~ 6.6, which is well outside the pH range generally thought to be compatible with life. We cannot, without further study, exclude a small contribution of increased cAMP to the OGR1-mediated increase in Ca$_i$ shown in this study to acid-induced bone resorption. We have never found any measurable change in cAMP in response to MET in either intact calvariae or primary calvarial cells (unpublished data). A recent publication has studied acid induction of COX-2 in the human osteoblastic cell line NHOst. These cells express OGR1 as the sole G protein–coupled proton–sensing receptor. A reduction of the incubation pH to 6.7 caused a substantial increase in Ca$_i$, which was sensitive to the G$_o$ inhibitor YM-254890. Acid induction of Ca$_i$ was also inhibited by siRNA directed against OGR1.

OGR1 may have roles in multiple bone cell types. The primary cells used in this study are almost exclusively osteoblasts, and our results support the role of OGR1 in the osteoblastic response to acidosis. However, the importance of OGR1 in osteoclastogenesis has also been established. A decrease in medium pH from 7.4 to 7.0 induces Ca$_i$ transients in rat osteoclasts, and these transients are suppressed by the phospholipase C inhibitor U73122. In its proliferative state, the monocytic cell line RAW264.7 expresses low levels of OGR1 and acidification causes only a minor change in Ca$_i$. Osteoclastic differentiation of RAW264.7 with RANKL increases OGR1 levels and the Ca$_i$ response to acidification. A short hairpin RNA construct directed against OGR1 decreased the Ca$_i$ response to acidification in transfected RAW264.7 cells. Whereas acidification of osteoclastic cells stimulates NFATc1 translocation to the nucleus, acidification also promotes osteoclast survival through a Ca- and ERK-dependent pathway that is NFAT independent.

The expression of mouse OGR1 in CHO cells models the acid-induced increase in Ca$_i$ in primary bone cells, suggesting that OGR1 acts as the primary mediator of the response of osteoblasts to increased proton concentration. The data presented in this study and that of Tomura...
et al. are consistent with a model in which proton activation of OGR1 leads to increased Ca levels. This increase acts as a second messenger to mediate the effects of acidosis on osteoblasts, including increased COX-2 (14) and RANKL (13) expression, which result in increased osteoclastic bone resorption.

ACKNOWLEDGMENTS

The authors thank Theresa Sherman for construction of the OGR1 expression plasmid and Anne Michalenga and Kelly LaPlante Strutz for expert technical assistance. This work was supported in part by Grants AR 46289 and DK 75462 from the National Institutes of Health and a grant from the Renal Research Institute.

REFERENCES

1. Lemann J Jr, Bushinsky DA, Hamm LL 2003 Bone buffering of acid and base in humans. Am J Physiol Renal Physiol 285: F811–F832.
2. Jehle S, Zanetti A, Muser J, Hulter HN, Krapf R 2006 Partial neutralization of the acidogenic western diet with potassium citrate increases bone mass in postmenopausal women with osteopenia. J Am Soc Nephrol 17:3213–3222.
3. Bushinsky DA 1995 The contribution of acidosis to renal osteodystrophy. Kidney Int 47:1816–1832.
4. Krieger NS, Frick KK, Bushinsky DA 2004 Mechanism of acid-induced bone resorption. Curr Opin Nephrol Hypertens 13:423–436.
5. Bushinsky DA, Wolbach W, Sessler NE, Mogilevsky R, Levi-Setti R 1993 Physicochemical effects of acidosis on bone calcium flux and surface ion composition. J Bone Miner Res 8: 93–102.
6. Bushinsky DA, Krieger NS, Geisser DI, Grossman EB, Coe FL 1983 Effects of pH on bone calcium and proton fluxes in vitro. Am J Physiol (Renal Fluid Electrolyte Physiol 14) 245: F204–F209.
7. Frick KK, Jiang L, Bushinsky DA 1997 Acute metabolic acidosis inhibits the induction of osteoblastic egr-1 and type 1 collagen. Am J Physiol (Cell Physiol 41) 272: C1450–C1456.
8. Krieger NS, Sessler NE, Mogilevsky R, Levi-Setti R 1993 Physicochemical effects of acidosis on bone calcium flux and surface ion composition. J Bone Miner Res 8: 93–102.
9. Bushinsky DA, Krieger NS, Geisser DI, Grossman EB, Coe FL 1983 Effects of pH on bone calcium and proton fluxes in vitro. Am J Physiol (Renal Fluid Electrolyte Physiol 14) 245: F204–F209.
10. Frick KK, Jiang L, Bushinsky DA 1997 Acute metabolic acidosis inhibits the induction of osteoblastic egr-1 and type 1 collagen. Am J Physiol (Cell Physiol 41) 272: C1450–C1456.
11. Krieger NS, Sessler NE, Mogilevsky R, Levi-Setti R 1993 Physicochemical effects of acidosis on bone calcium flux and surface ion composition. J Bone Miner Res 8: 93–102.
12. Frick KK, Jiang L, Bushinsky DA 1997 Acute metabolic acidosis inhibits the induction of osteoblastic egr-1 and type 1 collagen. Am J Physiol (Cell Physiol 41) 272: C1450–C1456.
13. Frick KK, Jiang L, Bushinsky DA 1997 Acute metabolic acidosis inhibits the induction of osteoblastic egr-1 and type 1 collagen. Am J Physiol (Cell Physiol 41) 272: C1450–C1456.
14. Frick KK, Jiang L, Bushinsky DA 1997 Acute metabolic acidosis inhibits the induction of osteoblastic egr-1 and type 1 collagen. Am J Physiol (Cell Physiol 41) 272: C1450–C1456.
15. Gitterman DP, Wilson J, Randall AD 2005 Functional properties and pharmacological inhibition of ASIC channels in the human SJ-RH30 skeletal muscle cell line. J Physiol 562:759–769.
16. Ugawa S 2003 Identification of sour-taste receptor genes. Anat Sci Int 78:205–210.
17. Lin W, Burks CA, Hansen DR, Kinnamon SC, Gilbertson TA 2004 Taste receptor cells express pH-sensitive leak K+ channels. J Neurophysiol 92:2909–2919.
18. Tomura H, Mogi C, Sato K, Okajima F 2005 Proton-sensing and lysolipid-sensitive G-protein-coupled receptors: A novel type of multi-functional receptors. Cell Signal 17:1466–1476.
19. Ludwig MG, Vanek M, Guerine D, Gasser JA, Jones CE, Junker U, Hofstetter H, Wolf RM, Seuwen K 2003 Proton-sensing G-protein-coupled receptors. Nature 425:93–98.
20. Wang JQ, Kon J, Mogi C, Tobo M, Damirin A, Sato K, Komachi M, Malcinkhenu E, Murata N, Kimura T, Kawabara A, Wakamatsu K, Koizumi U, Ueda T, Tsujimoto G, Kuruse H, Sato T, Harada A, Misawa N, Tomura H, Okajima F 2004 TDAG8 is a proton-sensing and psychosine-sensitive G-protein-coupled receptor. J Biol Chem 279:45626–45633.
21. Ishii S, Kihara Y, Shimizu T 2005 Identification of T cell death-associated gene 8 (TDAG8) as a novel acid sensing G-Protein-coupled receptor. J Biol Chem 280: 9083–9087.
22. Murakami N, Yokomizo T, Okuno T, Shimizu T 2004 G2A is a proton-sensing G-protein-coupled receptor antagonized by lysophosphatidylcholine. J Biol Chem 279:42484–42491.
23. Krieger NS, Parker WR, Alexander KM, Bushinsky DA 2000 Prostaglandins regulate acid-induced cell-mediated bone resorption. Am J Physiol Renal Physiol 279:F1077–F1082.
24. Krieger NS, Heffley TJ, 1989 Differential effects of parathyroid hormone on protein phosphorylation in two osteoblast-like cell populations isolated from neonatal mouse calvaria. Calcif Tissue Int 44:192–199.
25. Krieger NS 1992 Demonstration of sodium/calcium exchange in rodent osteoblasts. J Bone Miner Res 7:1105–1111.
26. King-Smith C, Chen P, Garcia D, Rey H, Burnside B 1996 Calcium-independent regulation of pigment granule aggregation and dispersion in teleost retinal pigment epithelial cells. J Cell Sci 109:33–43.
27. Fowler CJ, Tiger G 1997 Calibration of Fura-2 signals introduces errors into measurement of thrombin-stimulated calcium mobilisation in human platelets. Clin Chim Acta 265: 247–261.
28. Bushinsky DA 1988 Net proton influx into bone during metabolic but not respiratory, acidosis. Am J Physiol (Renal Fluid Electrolyte Physiol 23) 254:F306–F310.
29. Xu Y, Casey G 1996 Identification of Human OGR1, a Novel G Protein-Coupled Receptor That Maps to Chromosome 14. Genomics 35:397–402.
30. Radu CG, Nijagal A, McLaughlin J, Wang L, Witte ON 2005 Differential-proton sensitivity of related G protein-coupled receptors T cell death-associated gene 8 and G2A expressed in immune cells. Proc Natl Acad Sci USA 102:1632–1637.
31. Mogi C, Tomura H, Tobo M, Wang JQ, Damirin A, Kon J, Komachi M, Hashimoto K, Sato K, Okajima F 2005 Sphin-gosylphosphorylcholine antagonized proton-sensing ovarian cancer G-protein coupled receptor 1 (OGR1)-mediated inositol phosphate production and cAMP accumulation. J Pharmacol 1999:160–167.
32. Zhang J, Ryder KD, Bethel JA, Ramirez R, Duncan RL 2006 PTH-induced actin depolymerization increases mechanosensitive channel activity to enhance mechanically stimulated Ca2+ signaling in osteoblasts. J Bone Miner Res 21:1729–1737.
33. Komioka H, Sugawara Y, Murshid SA, Ishihara Y, Honjo T, Takano-Yamamoto T 2006 Fluid shear stress induces less calcium response in a single primary osteocyte than in a single osteoblast: Implication of different focal adhesion formation. J Bone Miner Res 21:1012–1021.
34. Batra NN, Li YJ, Yellowley CE, You L, Malone AM, Kim CH, Jacobs CR 2005 Effects of short-term recovery periods on leading acid sensors expressed in human nociceptors. J Clin Invest 110:1185–1190.
35. Gitterman DP, Wilson J, Randall AD 2005 Functional properties and pharmacological inhibition of ASIC channels in the human SJ-RH30 skeletal muscle cell line. J Physiol 562:759–769.
36. Centrella M, Casinghino S, McCarthy TL 1994 Differential actions of prostaglandins in separate cell populations from fetal rat bone. Endocrinology 135:1611–1620.
37. Tang LY, Kimmel DB, Jee WSS, Yee JA 1996 Functional characterization of prostaglandin E2 inducible osteogenic colony forming units in cultures of cells isolated from the neonatal rat calvarium. J Cell Physiol 166:76–83.
38. Bushinsky DA 1989 Internal exchanges of hydrogen ions: Bone. In: Seldin DW, Giebisch G (eds.) The Regulation of Acid-Base Balance. Raven Press, New York, NY, USA, pp. 69–88.
39. Bushinsky DA 1995 Metabolic acidosis. In: Jacobson HR, Striker GE, Klahr S (eds.) The Principles and Practice of Nephrology. Mosby, St. Louis, MO, USA, pp. 924–932.
40. Bushinsky DA, Krieger NS 1992 Role of the skeleton in calcium homeostasis. In: Seldin DW, Giebisch G (eds.) The Kidney: Physiology and Pathophysiology. Raven Press, New York, NY, USA, pp. 2395–2430.
41. Arnett TR 2008 Extracellular pH regulates bone cell function. J Nutr 138:415S–418S.
42. Jahr H, van Driel M, van Osch GJVM, Weinans H, van Leeuwen JPTM 2005 Identification of acid-sensing ion channels in bone. Biochem Biophys Res Commun 337:349–354.
43. Krishtal O 2003 The ASICs: Signaling molecules? Modulators? Trends Neurosci 26:477–483.
44. Xiong ZG, Zhu XM, Chu XP, Minami M, Hey J, Wei WL, MacDonald JF, Wemmie JA, Price MP, Welsh MJ, Simon RP 2004 Neuroprotection in Ischemia: Blocking Calcium-Permeable Acid-Sensing Ion Channels. Cell 118:687–698.
45. Naves LA, McCleskey EW 2005 An acid-sensing ion channel that detects ischemic pain. Braz J Med Biol Res 38:1561–1569.
46. Tomura H, Wang JQ, Komachi M, Damirin A, Mogi C, Tobo M, Kon J, Misawa N, Sato K, Okajima F 2005 Prostaglandin I2 production and cAMP accumulation in response to acidic extracellular pH through OGR1 in human aortic smooth muscle cells. J Biol Chem 280:34458–34464.
47. Tomura H, Wang JQ, Liu JP, Komachi M, Damirin A, Mogi C, Tobo M, Nochi H, Tamoto K, Im DS, Sato K, Okajima F 2008 Cyclooxygenase-2 expression and prostaglandin E2 production in response to acidic pH through OGR1 in a human osteoblastic cell line. J Bone Miner Res 23:1129–1139.
48. Komarova SV, Pereverzev A, Shum JW, Sims SM, Dixon SJ 2005 Convergent signaling by acidosis and receptor activator of NF-κB ligand (RANKL) on the calcium/calcineurin/NFAT pathway in osteoclasts. Proc Natl Acad Sci USA 102:2643–2648.
49. Pereverzev A, Komarova SV, Korcok J, Armstrong S, Tremblay GB, Dixon SJ, Sims SM 2008 Extracellular acidification enhances osteoclast survival through an NFAT-independent, protein kinase C-dependent pathway. Bone 42:150–161.

Address reprint requests to: Kevin K Frick, PhD
University of Rochester School of Medicine and Dentistry
Division of Nephrology
Department of Medicine
601 Elmwood Avenue, Box 675
Rochester, NY 14642, USA
E-mail: Kevin_Frick@URMC.Rochester.edu

Received in original form March 13, 2008; revised form July 1, 2008; accepted October 7, 2008.