Protein Kinase G Activation Reverses Oxidative Stress and Restores Osteoblast Function and Bone Formation in Male Mice With Type 1 Diabetes

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Bone loss and fractures are underrecognized complications of type 1 diabetes and are primarily due to impaired bone formation by osteoblasts. The mechanisms leading to osteoblast dysfunction in diabetes are incompletely understood, but insulin deficiency, poor glycemic control, and hyperglycemia-induced oxidative stress likely contribute. Here we show that insulin promotes osteoblast proliferation and survival via the nitric oxide (NO)/cyclic guanosine monophosphate (cGMP)/protein kinase G (PKG) signal transduction pathway and that PKG stimulation of Akt provides a positive feedback loop. In osteoblasts exposed to high glucose, NO/cGMP/PKG signaling was reduced due in part to the addition of O-linked N-acetylglucosamine to NO synthase-3, oxidative inhibition of guanylate cyclase activity, and suppression of PKG transcription. Cinaciguat—an NO-independent activator of oxidized guanylate cyclase—increased cGMP synthesis under diabetic conditions and restored proliferation, differentiation, and survival of osteoblasts. Cinaciguat increased trabecular and cortical bone in mice with type 1 diabetes by improving bone formation and osteocyte survival. In bones from diabetic mice and in osteoblasts exposed to high glucose, cinaciguat reduced oxidative stress via PKG-dependent induction of antioxidant genes and downregulation of excess NADPH oxidase-4–dependent H2O2 production. These results suggest that cGMP-elevating agents could be used as an adjunct treatment for diabetes-associated osteoporosis.

Bone loss and fractures are frequent and underappreciated complications of type 1 diabetes. Despite improvements in insulin replacement therapy, low bone mineral density and reduced bone mass are observed in most children and adults with type 1 diabetes (1–4). Multiple population-based studies have shown that patients with type 1 diabetes have a two- to fivefold increased risk of fractures, extending across the entire life span; moreover, fractures in patients with diabetes heal slowly, leading to significant morbidity and mortality (3–5). Bone loss in type 1 diabetes is mainly from reduced osteoblast activity; thus, current osteoporosis therapies aimed at inhibiting osteoclasts are inadequate (2,5,6). Patients with type 2 diabetes also have decreased bone formation and increased fracture risk, but bone mineral density may be normal or increased, while bone quality is impaired (1). Mouse models of type 1 diabetes more faithfully recapitulate the human bone disease than models of type 2 diabetes (2,7–9).

Poor glycemic control and diabetes-related vascular complications have emerged as predictors of fractures in most but not all studies of type 1 diabetes (1,3). In a group of patients who had type 1 diabetes for >50 years with excellent glycemic control and a low prevalence of cardiovascular disease, fracture rates were low, suggesting that optimal insulin therapy may protect from both complications (10). Insulin and IGF-1 have important anabolic effects in osteoblasts, as demonstrated in genetic mouse models (11,12). Thus, defective bone remodeling in type 1 diabetes is partly due to insulin deficiency, but the effect of hyperglycemia and hyperglycemia-induced oxidative stress is incompletely defined (13,14).

High glucose concentrations induce oxidative stress by several mechanisms: glucose auto-oxidation, nonenzymatic...
formulation of advanced glycation end products, increased mitochondrial superoxide \( (O_2^-) \) production, NADPH depletion from increased polypeptide flux, and increased expression or activation of NADPH oxidases (15). Growing evidence indicates increased reactive oxygen species (ROS) and decreased antioxidant defense mechanisms (e.g., thioredoxin) contribute to diabetic tissue damage and bone loss (8, 9, 16). High glucose also increases flux through the hexosamine pathway, leading to increased uridine diphosphate -N-acetylglucosamine (UDP-GlcNAc) concentrations, and posttranslational O-linked (O-)GlcNAc modification of proteins, such as NO synthase (NOS3) and SP1 transcription factors (15, 17, 18).

O-GlcNAc modification of NOS3 and oxidation of NOS3 cofactors reduce NOS3 activity in endothelial cells of patients with diabetes, causing impaired endothelium-dependent vasodilation, referred to as “endothelial dysfunction” (19). Levels of stable NO metabolites are low in serum and urine of patients with diabetes, suggesting a systemic NO deficiency (20). NO regulates bone formation, as illustrated by osteoblast defects in mice lacking NOS3, and NO donors preserve bone in estrogen-deficient rodents and postmenopausal women (21, 22). The major intracellular NO target is soluble guanylate cyclase (sGC), which produces cyclic guanosine monophosphate (cGMP), thereby activating cGMP-dependent protein kinase G (PKG1 and 2). We previously defined proproliferative, antiapoptotic pathways involving cGMP/PKG2-mediated activation of Src, extracellular signal–regulated kinase (ERK)-1/2, Akt, and \( \beta \)-catenin in osteoblasts and osteocytes (23–25). We now show that each step of the NO/cGMP/PKG pathway is impaired in osteoblasts and bones of mice with type 1 diabetes and that restoring cGMP synthesis reduces oxidative stress, recovers osteoblast functions, decreases osteocyte apoptosis, and prevents bone loss in the insulin-deficient mice.

RESEARCH DESIGN AND METHODS

Animal Experiments

All mouse procedures were approved by the University of California, San Diego Institutional Animal Care and Use Committee. Male C57Bl/6 mice (11 weeks old) from Envigo/Harlan were housed at three to four animals per cage in a temperature-controlled environment with a 12-h light/dark cycle and fed the standard Teklad Rodent Diet with ad libitum access to food and water. Male mice were used to avoid the confounding effects of estrogens on NO/cGMP signaling (22, 26). After 1 week of acclimatization, mice weighing 19–22 g were randomly divided into three groups. Groups 1 and 2 (12 mice each) received streptozotocin at 40 mg/kg/day intraperitoneally for 5 days, and group 3 (controls, 9 mice) received vehicle injections (7). Glucose was measured in tail vein blood 13 days after the start of the injections. Mice with a blood glucose concentration of >270 mg/mL were considered diabetic. Six streptozotocin-administered mice with blood glucose <270 mg/mL were excluded. Mice in groups 1 and 2 (8 and 10 mice, respectively) received vehicle or cinaciguat at 10 \( \mu \)g/kg/day intraperitoneally 6 days/week from day 21 through day 48. This dose of cinaciguat did not significantly reduce systolic blood pressure (<10 mmHg) and is sufficient to prevent bone loss in ovarioctomized mice (26). Double calcein labeling was performed by intraperitoneal injection of calcein (25 mg/kg) at 7 and 4 days before euthanasia. Mice were euthanized 1 h after the last drug or vehicle injection by CO2 intoxication and exsanguination.

Cell Culture

Primary osteoblasts (POBs) and bone marrow stromal cells (BMSCs) were isolated from long bones (22, 24) of control mice (Con), mice administered streptozotocin plus vehicle (DM), and mice administered streptozotocin plus cinaciguat (DM+Cin). Cells were cultured in 10% FBS-containing DMEM with 5 mmol/L glucose (Con), 25 mmol/L glucose (DM), or 25 mmol/L glucose plus 100 nmol/L cinaciguat (DM+Cin). POBs were also isolated from mice carrying floxed prkg2 alleles (prkg2\textsuperscript{fl/fl}) and were infected with an adenovirus encoding CRE recombinase, as previously described (22). Human POBs were explant cultures from trabecular bone fragments obtained from knee replacement operations for osteoarthritis, according to an institutionally approved protocol (24); none of the patients had diabetes. Cells were grown in DMEM with 10% FBS and 25 mmol/L glucose, unless noted otherwise. All POBs were used at passages 1–5 and characterized after differentiation in medium supplemented with ascorbate (0.3 mmol/L) and \( \beta \)-glycerol phosphate (10 mmol/L), as previously described (24).

Quantitative RT-PCR

Femurs were cleaned of surrounding soft tissue, including the periosteum, ~1 mm was cut off the ends, and the bone marrow was flushed out before samples were snap frozen in liquid nitrogen. RNA was purified from pulverized bone shafts and reverse transcribed; PCR was performed using a MX3005P real-time PCR detection system with Brilliant II SYBR Green Mix (Agilent Technologies) (22). Primer sequences are in Supplementary Tables 3 and 4; all primers were tested with serial cDNA dilutions. Relative changes in mRNA expression were analyzed using the 2\(^{-\Delta\Delta{Ct}}\) method, with 18S and tuba1 serving as internal references. Mean \( \Delta{Ct} \) values (gene of interest minus reference gene) measured in vehicle-treated control groups were assigned a value of 1 (22).

Amplex Red Assay for \( \text{H}_2\text{O}_2 \)

Human POBs were seeded at 4 \( \times \) \( 10^4 \) cells/well in 96-well plates and incubated in 0.1 mL of reaction mixture containing 50 \( \mu \)mol/L Amplex Red and 0.1 units/mL horseradish peroxidase. Fluorescence was measured every 2 min using 540 nm excitation and 590 nm emission wavelengths.

MitoSOX Red Detection of Mitochondrial \( \text{O}_2^- \)

Human POBs seeded on glass cover slips were loaded with 1 \( \mu \)mol/L MitoSOX Red for 10 min at 37°C. After washing, fluorescence microscopy was performed on live cells using 515 nm excitation and 580 nm emission wavelengths.
Assay of Lipid Peroxidation Products
POBs (5 x 10^5 cells/well) were lysed in 0.15 mL ice-cold radioimmunoprecipitation assay buffer and sonicated three times for 5 s. To 0.1 mL of cell lysate or serum, 0.2 mL 10% trichloroacetic acid and 0.3 mL 0.67% thiobarbituric acid was added, and samples were boiled for 45 min. The thiobarbituric acid adducts were extracted in butanol and measured using 515 and 553 nm as excitation and emission wavelengths, respectively. Malondialdehyde standards were from Cayman.

Quantitation of NOx and cGMP
NO production was measured based on nitrite and nitrate accumulation in the medium, using a two-step colorimetric assay: nitrate was first converted to nitrite by nitrate reductase, and then nitrite was measured spectrophotometrically based on the Griess reaction (22). cGMP concentrations were measured using an ELISA kit (Biomedical Technologies) according to the manufacturer’s protocol.

Western Blotting and Immunofluorescence Staining
Western blotting was performed using the antibodies described in Supplementary Table 2. Protein carbamylation was detected using the OxyBlot Protein Oxidation Detection Kit (Millipore EMD). Immunofluorescence staining with antibodies specific for BrdU, cleaved caspase-3, or 8-OH-deoxyguanosine (1:100 dilution) was done as previously described (26). Images were analyzed with a Keyence BZ-X700 fluorescence microscope.

Bone Histomorphometry and Staining of Bone Sections
Tibiae were fixed in 70% ethanol, dehydrated, embedded in methyl-methacrylate, and sectioned at the University of Alabama at Birmingham Center for Metabolic Bone Disease. An investigator blinded to the treatment of the mice identified trabecular osteoblasts and osteoclasts on Masson’s trichrome–stained sections and assessed fluorochrome labeling on unstained sections (27). Trabecular and cortical bone were analyzed between 0.25 and 2.25 mm or between 0.25 and 4.25 mm, respectively, distal to the growth plate. TUNEL staining of deplasticized sections was performed as previously described (26). For 8-OH-deoxyguanosine staining, sections were treated with proteinase K (10 mg/mL) for 7 min, followed by treatment with 0.2% Triton X-100 for 15 min. Anti–8-OH-deoxyguanosine antibody (1:50 dilution) was followed by goat anti-mouse horseradish peroxidase antibody (1:200), and incubation with 3,3-diaminobenzidine (Vector Laboratories). Blood vessel density was measured based on staining with anti-CD31 antibody (1:100). Clusters of at least two endothelial cells with anti-CD31 membrane staining, separate from other microvessels, were counted as individual vessels. Slides were scanned with a Hamamatsu NanoZoomer 2.0 HT System and analyzed using Digital Pathology NDP.view2 software (22).

Micro-Computed Tomography
Ethanol-fixed tibiae were scanned using a SkyScan 1076 (Bruker, Kontich, Belgium) micro-CT scanner at a voxel size of 9 μm, applying an electrical potential of 50 kV and current of 200 μA, and using a 0.5-mm aluminum filter. A beam-hardening correction algorithm was applied before image reconstruction with DataViewer and CTAn software (Bruker). Cortical bone was analyzed by automatic contouring 3.6–4.5 mm distal to the proximal growth plate, using a global threshold to identify cortical bone and eroding 1 pixel to eliminate partial volume effects. Trabecular bone was analyzed by automatic contouring of the proximal tibial metaphysis 0.36–2.1 mm distal to the growth plate. An adaptive threshold (using the mean maximum and minimum pixel intensity values of the surrounding 10 pixels) was used to identify trabecular bone (28).

Statistical Analyses
Graph Pad Prism 5 software was used for two-tailed Student t test to compare two groups or one-way ANOVA with Bonferroni posttest analysis to compare more than 2 groups. P < 0.05 was considered significant.

RESULTS
Insulin Enhances Osteoblast Proliferation and Survival via NO/cGMP/PKG2
Insulin treatment of human or murine POBs activated Akt, induced NOS3 Ser1177 phosphorylation (a stimulatory site targeted by Akt), and enhanced NO production; these effects were prevented when Akt activation was blocked by LY294002 or MK2206 (Fig. 1A and B and Supplementary Fig. 1A; the NOS inhibitor L-NAME served as a control). These results suggest insulin increases NO synthesis in osteoblasts via Akt phosphorylation of NOS3, similar to the effects of insulin’s in endothelial cells (29). Insulin-induced Akt and ERK activation in POBs was reduced by >50% when NOS, sGC, or PKG were inhibited pharmacologically or when PKG2 expression was knocked out by CRE-mediated recombination (Fig. 1C–G). Correspondingly, insulin stimulation of osteoblast proliferation and survival was largely PKG2-dependent (Fig. 1H and I and Supplementary Fig. 1B).

Together with our previous data showing NO/cGMP regulation of osteoblast proliferation and survival via ERK and Akt (24,25), these results indicate that the anabolic effects of insulin on osteoblasts require NO/cGMP/PKG2 signaling and point toward a positive feedback loop, where insulin-induced NOS3 stimulation is enhanced by PKG2 activation of Akt (Fig. 1J). Proproliferative and antiapoptotic insulin signals are reduced when this feedback loop is interrupted by NOS, sGC, or PKG inhibition, or prkg2 gene knockout.

NOS Activity Is Reduced in POBs From Diabetic Mice and Human POBs Exposed to High Glucose
We isolated POBs from mice with streptozotocin-induced diabetes and control mice, and cultured them in medium containing “high” (25 mmol/L) or physiological “low” (5 mmol/L) glucose concentrations, respectively, to mimic in vivo conditions. Basal NO production was reduced by ~50% in osteoblasts from diabetic compared with control
Figure 1—Insulin signaling via NO/cGMP/PKG2 in POBs. A–D: Human (h)POBs were treated with pharmacological inhibitors of PI3K (LY294002, 10 μmol/L), Akt (MK2206, 10 μmol/L), NOS (l-NAME, 4 mmol/L), sGC (ODQ, 10 μmol/L), PKG (Rp-CPT-PET-cGMPS, 50 μmol/L), or vehicle (Veh) for 1 h before stimulation with insulin for 20 min. ODQ, 1H-(1,2,4)oxadiazolo[4,3-a]quinoxalin-1-one. A: Stable NOx metabolites (nitrite + nitrate) were measured in the medium by Griess reaction. Phosphorylation (p) of Akt (Ser473), NOS3 (Ser1177), and ERK-1/2 (Thr202/Tyr204) was assessed with phospho-specific antibodies (B and C) and densitometry scanning of Western blots (D). E–G: Murine (m)POBs containing “floxed” prkg2f/f alleles were infected with control virus (LacZ) or virus expressing CRE-recombinase (CRE). E and F: After 48 h, cells were stimulated with insulin for 10 min to assess ERK and Akt activation. G: prkg2 mRNA (normalized to 18S rRNA) was measured by quantitative RT-PCR, and PKG2 protein (in POB membranes) by Western blotting. H and I: prkg2f/f POBs infected with LacZ or CRE virus were transferred to low serum–containing medium with some cells receiving insulin. H: Proliferation was measured by 3H-thymidine incorporation into DNA. I: The effect of insulin on apoptosis was assessed by determining the percentage of cells staining for cleaved caspase-3 in the absence of serum (green, DNA counterstained blue with Hoechst 33342). Scale bar = 15 μm. Data represent mean ± SD of three to five independent experiments. *P < 0.05, **P < 0.01, and ***P < 0.001 for the indicated comparisons; in panel I ***P < 0.001 for comparison with serum starvation without insulin and ###P < 0.01 for the indicated comparison (by ANOVA, except t test in G). J: Insulin activates Akt and ERK downstream of the insulin receptor, and Akt phosphorylation of NOS3 activates the NO/cGMP/PKG2 pathway in osteoblasts to regulate proliferation and survival. Akt phosphorylation of NOS3 also occurs in a positive feedback loop. When NO/cGMP/PKG2 signaling is inhibited, this feedback loop is interrupted, and insulin signaling via ERK and Akt is reduced.
Figure 2—Reduced NOS activity in POBs exposed to high glucose. A: Mouse (m)POBs isolated from control (Con) and diabetic mice (DM) were cultured in 5 mmol/L or 25 mmol/L glucose, respectively, and NOx was measured 2 h after transfer to fresh medium, as in Fig. 1A. NOS3 (Ser1177) phosphorylation (p) was analyzed with a phospho-specific antibody (B), and NOS3 association with HSP-90 (C) or caveolin-1 (D) was determined by coimmunoprecipitation (IP) (10% input lysates are shown below). E: Human (h)POBs from individuals without diabetes were cultured for 3 weeks in medium with 5 mmol/L (low) glucose (LG) or 25 mmol/L (high) glucose (HG), with some cells in LG receiving 20 mmol/L mannitol; NOx was measured 2 h after insulin stimulation. F: NOS3 was isolated from human POBs cultured in LG or HG by ADP-agarose pull-down, and O-GlcNAc modification of the protein was determined on Western blots using the anti–O-GlcNAc antibody RL-2. The graph summarizes densitometry scans from three independent experiments. G and H: Human POBs were maintained in LG or HG medium or were switched from LG to HG and vice versa. NO production (G) and NOS3 (Ser1177) phosphorylation (H) were measured at 0 and 72 h and then weekly; for each time point, cells were plated at a density of 2 × 10⁴ cells/cm². I and J: Human POBs cultured for 3 weeks in LG or HG were infected with control virus or virus encoding OGT. I: Cells received vehicle or insulin for 20 min, and NOx was measured as in Fig. 1A. J: NOS3 phosphorylation and OGT expression were assessed in insulin-treated cells. Data represent mean ± SD of three independent experiments (except n = 2 in D). *P < 0.05, **P < 0.01, ***P < 0.001 for the indicated comparisons, and ###P < 0.001 for comparison between the same treatment in HG vs. LG (A–C, by t test; E–I, by ANOVA).
animals, as was NOS3 Ser\textsuperscript{1177} phosphorylation and NOS3 association with HSP90, a positive enzyme regulator (Fig. 2A–C) (30). In contrast, NOS3 association with its negative regulator caveolin-1 (30) was increased in diabetic osteoblasts (Fig. 2D).

To concentrate on hyperglycemic effects on NO synthesis and simulate diabetic conditions in human POBs, cells from donors without diabetes were cultured for 3 weeks in medium containing high or low glucose. Basal and insulin-stimulated NO production were both reduced in POBs exposed to high glucose compared with cells cultured in low glucose. The differences were not from increased osmolarity, because adding 20 mmol/L mannitol to the 5 mmol/L glucose medium neither affected NO production nor insulin-induced Akt or ERK activation (Fig. 2E and Supplementary Fig. 2A–C). To determine whether reduced NO production in human POBs cultured in high versus low glucose may be partly related to O-GlcNac modification of NOS3, we isolated the enzyme via ADP-agarose pull-down and probed Western blots with the anti-O-GlcNac antibody RL-2. We found a 2.3-fold increase in NOS3 O-GlcNac-acytelylation in cells cultured in high versus low glucose (Fig. 2F and Supplementary Fig. 2D). These data are in keeping with findings in endothelial cells exposed to high glucose and in vascular tissue from diabetic rats, where O-GlcNac modification of NOS3 on Ser\textsuperscript{1177} reduces NOS activity by interfering with Akt phosphorylation of Ser\textsuperscript{1177} (17,31).

When human POBs were transferred from low to high glucose, NO production and NOS3 Ser\textsuperscript{1177} phosphorylation decreased gradually over ~3 weeks. NO production and NOS3 phosphorylation both recovered with similar kinetics when cells were transferred from high to low glucose (Fig. 2G and H). Similar results were obtained in POBs isolated from mice with type 1 diabetes (Supplementary Fig. 2E). Thus, decreased basal NO synthesis in diabetic POBs is reversible when hyperglycemia is corrected.

To test the effect of O-GlcNac modification on osteoblast NOS activity, we infected human POBs with a viral vector encoding O-GlcNac transferase (OGT), which adds a single GlcNac in O-glycosidic linkage to Ser and Thr residues. OGT expression inhibited insulin-induced NO production and NOS3 Ser\textsuperscript{1177} phosphorylation in low and high glucose medium, and there was a trend toward OGT reducing basal NO synthesis (Fig. 2I and J). In murine POBs in high glucose, OGT expression decreased basal NO production significantly, whereas expression of N-acetylglucosaminidase (GCA), which removes O-GlcNac, increased NO production (Supplementary Fig. 2F and G). These results are consistent with O-GlcNac modification of NOS3 on Ser\textsuperscript{1177} interfering with NOS3 phosphorylation on this site, thereby reducing NOS activity (17).

### Guanylate Cyclase Is Oxidized and NO-Induced cGMP Synthesis Is Impaired in POBs Exposed to High Glucose

The NO donor PAPA-NONOate [(Z)-1-[N-(3-aminopropyl)-N-(propyl)amino]diazen-1-ium-1,2-diolate] elicited a dose-dependent increase in the intracellular cGMP concentration in osteoblasts from normal mice in low glucose, but in cells from diabetic mice in high glucose, only the highest PAPA-NONOate concentration elicited a small increase in cGMP (Fig. 3A). sGC protein concentrations were similar in both conditions (Fig. 3B). To determine whether the decreased NO responsiveness of sGC was due to oxidation of the heme-bound Fe\textsuperscript{2+}, we compared cinaciguat—a NO- and heme-independent sGC activator that can activate oxidized (Fe\textsuperscript{3+}) or heme-free sGC in addition to the reduced (Fe\textsuperscript{2+})-heme-bound enzyme to riociguat, which stimulates only the (Fe\textsuperscript{2+})-heme-containing enzyme (32,33). Cinaciguat increased the intracellular cGMP concentration similarly in osteoblasts from control and diabetic mice, whereas comparable concentrations of riociguat had minimal effect in cells from diabetic mice (Fig. 3C). Similarly, PAPA-NONOate and riociguat induced ERK and Akt activation much less effectively in cells from diabetic mice compared with control cells, whereas cinaciguat was equally effective under both conditions (Fig. 3D and E). The cell permeable ROS scavenger tempol partly restored sGC responsiveness to PAPA-NONOate and riociguat in cells from diabetic mice, providing further support that the sGC heme group is oxidized under diabetic conditions (Fig. 3F). Thus, cinaciguat, the prototype of a new class of NO-independent sGC activators under clinical development (32), effectively raises osteoblast cGMP under hyperglycemic conditions.

### Expression of prkg1 and prkg2 Is Downregulated in POBs Exposed to High Glucose

The main cGMP targets in osteoblasts are cytosolic PKG1 and membrane-bound PKG2, encoded by prkg1 and prkg2, respectively (24). Both enzymes have some overlapping but also distinct functions in promoting osteoblast proliferation, differentiation, and survival (24,25). We found decreased PKG1 and 2 protein and prkg1 and 2 mRNA in osteoblasts from diabetic mice compared with control mice, whereas nos3 and gucy1 mRNAs (encoding NOS3 and the common sGC-B1 subunit) were unchanged (Fig. 3G–I). Switching human osteoblasts from low to high glucose medium decreased prkg1 and 2 mRNAs over 2–3 weeks, and changing cells from high to low glucose increased both mRNAs over a similar time course (Fig. 3J). The stability of both mRNAs was not altered in high versus low glucose medium (Supplementary Fig. 3A and B).

To determine whether glucose-induced oxidative stress contributed to transcriptional downregulation of prkg1 and 2, we treated osteoblasts in high glucose medium with tempol. Tempol increased prkg1 and 2 mRNAs in human and murine POBs in high glucose to concentrations found in low glucose–containing medium; concomitantly, the drug decreased the intracellular concentration of malondialdehyde, a lipid oxidation marker (Fig. 3K and Supplementary Fig. 3C and D).

We have shown that high glucose–induced O-GlcNac modification of the transcription factor SP1 leads to
Figure 3—Oxidized guanylate cyclase and reduced prkg1 and prkg2 expression in POBs exposed to high glucose. A–E and G–J: POBs were isolated from control (Con) or diabetic mice (DM) and cultured as in Fig. 2A. A: Cells were stimulated with the NO donor PAPA-NONOate (0.3–10 μmol/L) for 30 min, and cGMP was measured by ELISA. B: Expression of sGC was assessed by Western blotting using an antibody for the common β1 subunit, with α-tubulin (Tub) as the loading control. C: Cells were stimulated with vehicle (Veh), cinaciguat (Cin, 0.1 and 1 μmol/L) or riociguat (Rio, 1 and 10 μmol/L) for 30 min, and cGMP was measured by ELISA. D and E: Cells were treated with PAPA-NONOate, Cin, or Rio for 10 min; ERK and Akt phosphorylation (p) were assessed as in Fig. 1C. F: Human (h)POBs cultured in 5 or 25 mmol/L glucose for 3 weeks were treated with 200 μmol/L tempol for 48 h before stimulation with 3 μmol/L PAPA-NONOate or 10 μmol/L Rio for 30 min; cGMP was measured as in panel A. G: Cytosolic and membrane fractions from control and diabetic POBs were analyzed by Western blotting with antibodies for PKG1, tubulin, PKG2, and caveolin-1. H: Blots were quantified by densitometry. I: nos3, gucy1 (sGC common β1 subunit), and prkg1 and 2 mRNAs were quantified by RT-PCR and normalized to 18S rRNA in control and diabetic POBs; expression in control POBs was assigned a value of 1. J: After 3 weeks in medium with 5 mmol/L (low) glucose (LG) or 25 mmol/L (high) glucose (HG), human POBs were maintained in the same medium or were switched from LG to HG and vice versa, and prkg1 and 2 mRNAs were measured as in panel I. K and L: Human POBs cultured in LG or HG for 3 weeks received vehicle vs. tempol (Tem, 200 μmol/L) or were infected with control virus encoding green fluorescent protein (GFP).
decreased promoter activity and expression of target genes and that SP1 positively regulates prkg1 expression (18). Expressing GCA in POBs in high glucose medium restored prkg1 and 2 mRNAs to concentrations found in low glucose (Fig. 3I). Thus, decreased prkg1 and 2 transcription under diabetic conditions is likely caused by oxidative stress and O-GlcNAc modification of transcriptional regulators.

Cinaciguat Restores Proliferation and Differentiation of Preosteoblasts Exposed to High Glucose

POBs isolated from diabetic mice and cultured in high glucose showed lower BrdU incorporation into DNA, and, when differentiated in vitro, produced less alkaline phosphatase (ALP) and mineralized less well than POBs isolated from control mice cultured in low glucose (Fig. 4A–D; POBs in Fig. 4C and D were postconfluent when they were switched to differentiation medium). Treating diabetic mice with cinaciguat (see Fig. 7A, below) and culturing POBs from these mice in high glucose in the presence of 100 nmol/L cinaciguat returned BrdU incorporation, ALP production, and cell mineralization to values found in cells from control mice in low glucose (Fig. 4A–D). In addition, cinaciguat restored mRNA expression of osteoblast marker genes and c-fos in osteoblasts from diabetic mice to that of control mice (Fig. 4E and F). BMSCs isolated from diabetic mice and cultured in high glucose generated fewer ALP-producing colonies and expressed lower amounts of osteoblast-specific genes compared with cells from control mice, and once again, cinaciguat restored these functions (Fig. 4G–I and Supplementary Fig. 4).

Cinaciguat Reduces Oxidative Stress in POBs Exposed to High Glucose

We assessed oxidative stress by measuring whole-cell H2O2 production, and mitochondrial O2− production (15). In POBs isolated from diabetic mice and in human POBs cultured in high glucose, H2O2 production was higher than in control POBs in low glucose, and cinaciguat reduced H2O2 production significantly (Fig. 5A and B). The cGMP analog 8-p-chlorophenylthio–cGMP (8-CPT–cGMP) and the antioxidant tempol decreased H2O2 production comparable to cinaciguat (Fig. 5B). The effect of cinaciguat was mediated by PKG2, because the drug was ineffective in prkg2-deficient osteoblasts (Fig. 5C). Protein, lipid, and DNA oxidation and p38 and JNK phosphorylation were increased in cells exposed to high glucose, and cinaciguat largely reversed these changes (Fig. 5D–J and Supplementary Fig. 5A–C). Again, 8-CPT–cGMP and tempol had similar effects as cinaciguat, and the effects of cinaciguat on lipid peroxidation were absent in prkg2-deficient osteoblasts (Fig. 5E and F).

In contrast to findings in endothelial cells (19), we found no evidence that NOS uncoupling contributed to H2O2 production in osteoblasts in high glucose (Supplementary Fig. 5D). Using the mitochondrial O2− indicator MitoSOX Red, we found increased mitochondrial O2− production in human POBs exposed to high glucose compared with POBs cultured in low glucose. Cinaciguat reduced mitochondrial O2− production toward values seen in low glucose (Supplementary Fig. 5E; antimycin A was used as a positive control). Thus, cinaciguat reduces oxidative stress in osteoblasts under high glucose conditions.

Cinaciguat Regulates Thioredoxin-Related Genes and NADPH Oxidase 4, Which Is a Major Source of ROS in POBs Exposed to High Glucose

Because transgenic expression of thioredoxin (txn1) reduces oxidative stress and partly prevents diabetes-induced bone loss in mice with type 1 diabetes (16), we determined the effect of hyperglycemia and cinaciguat on txn1 and thioredoxin-related genes. Compared with normoglycemic conditions, we found reduced expression of the antioxidant genes txn1, txnrd1 (thioredoxin reductase), and prdx1 (thioredoxin peroxidase) in osteoblasts under hyperglycemic conditions, with almost full recovery after cinaciguat treatment (Fig. 6A). The pro-oxidant gene txnip (thioredoxin-interacting protein) was regulated in the opposite direction (Fig. 6A). Thioredoxin-interacting protein has pro-oxidant properties because it inhibits thioredoxin and enhances nox4 expression under diabetic conditions (34).

NADPH oxidase 4 (NOX4) is a constitutively-active NADPH oxidase that generates predominantly H2O2, and is transcriptionally upregulated by a variety of cellular stresses, including high glucose (35,36). High glucose increased nox4 mRNA and protein in mouse and human POBs, with cinaciguat preventing the upregulation; nox2 expression was not altered under these conditions (Fig. 6A–C and Supplementary Fig. 6A and B). Cinaciguat’s suppression of nox4 mRNA was cGMP and PKG2 mediated and was mimicked by tempol, suggesting a mechanism linked to ROS (Fig. 6D and E). Txr1 mRNA regulation was reciprocal to that of nox4: trxl was downregulated in high glucose, cinaciguat and tempol increased its expression, and cinaciguat’s effect required PKG2 (Fig. 6F and G). Cinaciguat, cGMP, and tempol did not affect nox4 or trxl mRNA under low glucose conditions, suggesting regulation occurred only under high oxidative stress (Supplementary Fig. 6C and D).

To evaluate whether NOX4 upregulation could account for the increased H2O2 production in POBs exposed to high glucose, H2O2 production was measured in POBs exposed to high glucose with and without cinaciguat. In control POBs, the addition of cinaciguat to high glucose decreased H2O2 production (Fig. 6H). In diabetic POBs, cinaciguat had a significantly smaller effect on H2O2 production, consistent with the larger basal H2O2 production in diabetic versus control POBs. These findings confirm that NOX4 is a major source of oxidative stress in diabetic POBs. NOS uncoupling did not affect H2O2 production under high glucose conditions (Supplementary Fig. 6D).

vs. virus encoding GCA for 48 h before measuring prkg1 and 2 mRNAs. Graphs show mean ± SD of three to four independent experiments. By ANOVA (or t test in H and I); *P < 0.05, **P < 0.01, ***P < 0.001 for the indicated comparisons or comparison with vehicle (C and K) or GFP (L); and #P < 0.05, ##P < 0.01, ###P < 0.001 for comparison between control and diabetic osteoblasts treated with the same drug concentration (C and D) or between absence and presence of tempol (F).
glucose, we used a gene silencing approach. Short hairpin (sh)RNA–mediated knockdown of nox4 decreased H2O2 production and lipid peroxidation in human POBs in high glucose to values found in control POBs in low glucose; in contrast, the nox4 shRNA had little effect on basal H2O2 production and lipid peroxidation in cells in low glucose (Fig. 6H and I; Fig. 6J shows efficient NOX4 depletion by the shRNA). These results indicate nox4 upregulation is
Figure 5—Reduction of oxidative stress by cinaciguat. A and D: POBs were isolated from control mice (Con), diabetic mice (DM), or diabetic mice treated with cinaciguat (DM+Cin) and were cultured as described in Fig. 4. B, E, and G–J: Human (h)POBs were cultured in 5 mmol/L (low) glucose (LG) or 25 mmol/L (high) glucose (HG) for 3 weeks; some cells received cinaciguat (Cin, 0.1 μmol/L), 8-CPT-cGMP (cGMP, 100 μmol/L), or tempol (Tem, 200 μmol/L) for the last 48 h, as indicated. C and F: POBs were isolated from “flanked” prkg2−/− mice. Cells were cultured in LG or HG for 3 weeks, infected with control virus (LacZ) or virus expressing CRE-recombinase (CRE), and treated with cinaciguat for 48 h. A–C: H₂O₂ production was measured by Amplex Red fluorescence and normalized to cellular protein content, with control cells assigned a value of 100%. D: Protein carbamylation was detected using OxyBlot after derivatization with 2,4-dinitrophenylhydrazine (+DNPH); nonderivatized extracts served as controls (**). E and F: Malondialdehyde (MDA) concentrations were measured to estimate lipid peroxidation products. G: Oxidative DNA damage was assessed by immunofluorescence using an 8-OH-deoxyguanosine–specific antibody (red; DNA counterstained blue with Hoechst 33342); the percentage of red-fluorescent nuclei is shown below (n = 3). **P < 0.01 compared with LG, #P < 0.05 and ##P < 0.01 compared with HG+vehicle. H–J: JNK and p38 activation was evaluated using phospho (p)-specific antibodies and densitometry scanning of Western blots. Graphs represent mean ± SD of three to six independent experiments. By ANOVA *P < 0.05, **P < 0.01, ***P < 0.001 for the indicated comparisons, and ###P < 0.001 for drug-treated compared with vehicle-treated cells in HG.
largely responsible for increased H₂O₂ production in POBs in high glucose, and cinaciguat reduces oxidative stress under hyperglycemic conditions by suppressing \( \text{nox4} \) and \( \text{txnip} \) and inducing antioxidant genes.

**Cinaciguat Treatment of Diabetic Mice Improves Bone Microarchitecture and Bone Formation, While Decreasing Resorption**

To induce type 1 diabetes, we administered streptozotocin to mice (7), and after 3 weeks—when the mice had been hyperglycemic for at least 1 week—we began administering a 4-week course of daily cinaciguat (Fig. 7A and Supplementary Table 1). In the diabetic mice, serum cGMP concentrations were decreased by \( \sim 60\% \) compared with nondiabetic animals, similar to patients with type 1 diabetes (37), and cinaciguat largely restored concentrations to normal (Fig. 7B).

As observed in osteoblasts from diabetic mice in vitro, \( \text{prkg1} \) and \( \text{prkg2} \) mRNAs were reduced in bones of diabetic mice, whereas \( \text{nos3} \) mRNA was unchanged. Cinaciguat restored \( \text{prkg1} \) expression but had little effect on \( \text{prkg2} \) mRNA (Fig. 7C).

Micro-CT analysis of tibial bone showed decreased trabecular and cortical bone volume fraction, trabecular number, and cortical cross-sectional bone area in diabetic mice, with cinaciguat treatment improving all three parameters (Fig. 7D–F). Mineral apposition rate, mineralizing surface per bone surface, and bone formation rate (BFR) were dramatically decreased in the diabetic mice, as described before (9), but cinaciguat largely restored these parameters, indicating improved osteoblastic activity (Fig. 7G–J).

Consistent with previous reports, trabecular and cortical osteoblast numbers were reduced and osteocyte apoptosis
was increased in the diabetic mice (9,13); cinaciguat almost completely reversed these changes (Fig. 7K and L and Supplementary Fig. 7A and B). Osteoclast number and serum concentrations of the osteoclast activity marker COOH-terminal telopeptide of type 1 collagen (CTX) were increased in the diabetic mice (Fig. 7M and N), as observed by some but not all investigators (9,13,38). Cinaciguat-treated diabetic animals showed reduced serum CTX concentrations and a trend toward reduced osteoclast numbers (Fig. 7M and N).
Histological examination of femur sections revealed an increase in bone marrow adiposity in the diabetic mice (Supplementary Fig. 7C and D), which is thought to be due to increased adipogenic differentiation of mesenchymal stem cells (2). Cinaciguat treatment slightly decreased the percentage of bone marrow occupied by fat cells, but this trend did not reach significance (Supplementary Fig. 7C and D). cGMP/PKG can stimulate vasculogenesis via induction of vascular endothelial growth factor-A (VEGF-A), and VEGF-A is expressed by osteoblasts (39,40). We found decreased vegfa mRNA and a trend toward reduced microvessel density in the bones of mice with streptozotocin-induced diabetes. Cinaciguat restored vegfa mRNA to levels found in control mice and caused a trend toward improved bone marrow microvascular density in diabetic mice, but the latter change did not reach statistical significance (Supplementary Fig. 8A–C).

These data indicate that cinaciguat largely reverses the negative effects of diabetes on trabecular and cortical bone by increasing osteoblast number and bone formation parameters while reducing osteoclast activity; however, the drug had only minor effects on bone marrow adiposity and microvessel density.

**Cinaciguat Restores Osteoblastic and Osteoclastic Gene Expression to Control Levels and Reduces Oxidative Stress in Diabetic Bone**

Concomitant with the decrease in osteoblast number and bone formation, expression of osteoblast differentiation-associated genes (dlx1, runx2, sp7, bglap, lrp5) was reduced in the bones of diabetic mice, but cinaciguat restored expression to levels found in control mice (Fig. 8A), analogous to its effect in POBs in vitro (Fig. 4E). The Wnt antagonist sclerostin (sost) is produced predominantly by osteocytes. Its expression was increased in diabetic animals, as previously reported (13), but cinaciguat reduced sost mRNA to control levels (Fig. 8A; tubulin mRNA was unchanged under all conditions). Consistent with the increase in osteoclast numbers, osteoclast-specific gene expression (apc5, ctsk) was increased in the bones of diabetic mice, with cinaciguat reducing expression to control levels (Fig. 8B).

Malondialdehyde concentrations, marking lipid peroxidation end products, were elevated in the serum of mice with type 1 diabetes compared with control mice (Fig. 8C), similar to results in patients with type 1 diabetes (37). However, cinaciguat treatment returned malondialdehyde to control levels (Fig. 8C). Osteocyte nuclear 8-OH-deoxyguanosine content, marking DNA oxidation, was increased in the diabetic mice, but cinaciguat restored this marker back to control levels (Fig. 8D). Consistent with depletion of antioxidant defense mechanisms in diabetic tissues (15), thioredoxin-related antioxidant genes were reduced in diabetic bones, while the pro-oxidant gene txnip1 was increased (Fig. 8E). nox4 mRNA was dramatically induced in diabetic bones, whereas nox2 was unchanged (Fig. 8F). Cinaciguat reversed these gene expression changes in diabetic bones (Fig. 8E and F), similar to its effects in POBs (Fig. 6A).

Thus, cinaciguat reduced oxidative stress markers, improved bone formation, and prevented bone loss in diabetic mice, suggesting that activators of oxidized guanylate cyclase may represent a new class of bone anabolic agents for diabetic osteoporosis.

**DISCUSSION**

Compared with other diabetic complications, diabetic bone disease has received relatively little attention, and treatment of diabetic osteoporosis remains unsatisfactory (5). Bisphosphonates, which inhibit bone resorption, can improve bone mineral density but may worsen bone quality by repressing bone formation (1,38). Parathyroid hormone analogs, the only U.S. Food and Drug Administration-approved drugs that stimulate bone formation, have not been studied in patients with diabetes (1,41). A better understanding of the molecular mechanisms underlying diabetic bone loss is needed to design safe and effective therapies, especially in patients with suboptimal glycemic control. We now describe several findings that could lead to a novel adjunct therapy for diabetic osteoporosis (Fig. 8G): 1) insulin stimulation of osteoblast proliferation and survival requires NO/cGMP/PKG2 signaling; 2) each step of the NO/cGMP/PKG2 pathway is impaired in osteoblasts under hyperglycemic conditions; 3) cinaciguat, via cGMP/PKG2, exerts potent antioxidant effects in diabetic osteoblasts; and 4) cinaciguat improves bone formation and reduces bone loss in mice with type 1 diabetes by restoring impaired osteoblast proliferation, differentiation, and survival, constraining osteoclast activity, and reducing oxidative stress.

**Insulin Signaling in Osteoblasts**

Insulin promotes proliferation, survival, and differentiation of osteoblasts via insulin receptor activation of the Ras/Raf/mitogen-activated protein kinase (MEK)/ERK and phosphatidylinositol 3-kinase (PI3K)/Akt pathways, and mice lacking the insulin receptor in osteoblasts are osteopenic (12). In mice with type 1 diabetes, insulin therapy improves bone formation and partly preserves bone volume (12,42). We found that insulin requires NO/cGMP/PKG2 signaling for its prosurvival and proproliferative effects in osteoblasts. Although we did not study insulin signaling in vivo, we observed decreased insulin activation of the NO/cGMP/PKG2 pathway in osteoblasts exposed to high glucose, suggesting the bone-anabolic effects of insulin are diminished under hyperglycemic conditions. Consistent with this hypothesis, high-dose continuous insulin infusion only partly prevented bone loss in mice with type 1 diabetes, in which blood glucose remained above nondiabetic levels (42).

FOXO transcription factors are important insulin targets, with insulin restraining FOXO activation and nuclear translocation via PI3K/Akt. FOXOs 1, 3, and 4 influence osteoblast proliferation, survival and resistance to oxidative stress (43). Mice with a deficiency of FOXO1 in Col1A-CRE–expressing osteoblastic cells have low bone volume with decreased osteoblast number and BFR, without change in
Cinaciguat effects in diabetic mice: regulation of osteoblast- and osteoclast-related gene expression and reduction of oxidative stress. Type 1 diabetes was induced, and diabetic mice were treated with vehicle (DM) or cinaciguat (DM+Cin) as described in Fig. 7A. A and B: Gene expression was measured in tibial bone and normalized to 18S rRNA as described in Fig. 7C. The means of the control group were assigned a value of 1 (dlx1, distal-less homeobox-1; sp7, osterix; bglap, osteocalcin; alpl, ALP; ltp5, LDL receptor-related protein-5; sost, sclerostin; tuba1, tubulin-a1; acp5, tartrate-resistant acid phosphatase; csk, cathepsin K; hprt, hypoxanthine phosphoribosyltransferase). C: Serum malondialdehyde concentrations were measured spectrophotometrically to assess lipid peroxidation. D: Osteocyte oxidative DNA damage was assessed by immunohistochemical staining of femoral cortex with an 8-OH-deoxyguanosine–specific antibody (black). The percentage of black cells is shown below. E and F: Gene expression was measured as in panel A. A–F: n = 5–7 mice/group. *P < 0.05, **P < 0.01, ***P < 0.001 for the indicated comparisons, except in panel D, *P < 0.05 for the comparison with diabetic, vehicle-treated mice (by ANOVA). G: Insulin activates NO/cGMP/PKG signaling via Akt, and insulin stimulation of osteoblast proliferation and survival requires a positive feedback loop involving cGMP/PKG activation of Akt. Under hyperglycemic conditions, each step of the NO/cGMP/PKG pathway is impaired: NOS activity is reduced by O-GlcNAc modification, sGC activity and NO responsiveness is reduced by oxidation of the enzyme’s heme group, and PKG1 and 2 transcription is downregulated by ROS- and O-GlcNAc-sensitive transcriptional regulators. Cinaciguat activates oxidized sGC in an NO-independent fashion and exerts antioxidant effects via cGMP activation of PKG2, by downregulating NOX4 and upregulating thioredoxin family antioxidant proteins. Reduction of ROS improves proliferation and survival of osteoblasts and their precursors.
osteoblast apoptosis (44). In contrast, deletion of FOXOs 1/3/4 in osteoblast progenitors via Osx1-CRE increases osteoblast number, BFR, and cortical bone volume, and attenuates diabetes-induced trabecular bone loss, without correcting low BFR or cortical bone loss in mice with type 1 diabetes (43). Because we found cGMP-dependent Akt activation in osteoblasts, some of the bone-protective effects of cinaciguat in diabetic mice could be explained by inhibition of FOXO transcription. However, the roles of FOXOs in bone are complex, and more work is required to define the role of each FOXO family member.

**Defective NO/cGMP Signaling in Diabetes**

Endothelial NO generation and serum nitrite/nitrate and cGMP concentrations are reduced in patients with diabetes, similar to our finding of reduced serum cGMP in diabetic mice (20,37); however, few studies have addressed cGMP/PKG signaling downstream of NO under diabetic conditions (19). Consistent with our results in osteoblasts, O-GlcNAC modification of NOS3 is increased approximately twofold in endothelial cells exposed to high glucose or isolated from diabetic mice compared with control cells (17,45). O-GlcNAC modification of NOS3 occurs predominantly on Ser1177 and blocks phosphorylation and activation of NOS3 by Akt, but NOS association with HSP90 and caveolin-1 was not studied previously under diabetic conditions (17,30). We saw no evidence of uncoupled NOS activity in diabetic osteoblasts, in contrast to what occurs in diabetic endothelium (19).

We found oxidative stress reduced sGC activation and downregulated PKG1 and 2 transcription under hyperglycemic conditions in vitro and in vivo. The reduced NO-responsiveness of sGC in POBs exposed to high glucose was due to oxidative stress of the enzyme’s heme iron rather than to reduced enzyme levels, because 1) cinaciguat, which can activate oxidized (Fe3+) and reduced (Fe2+) sGC, stimulated cGMP synthesis similarly in POBs in high and low glucose, whereas riociguat and NO, which activate only reduced (Fe2+) sGC, were much more effective in low than in high glucose, despite similar levels of the heme-containing β-subunit of sGC; and 2) the ROS-quenching agent tempol increased cGMP synthesis more than fivefold in response to riociguat or NO in POBs in high glucose, whereas the antioxidant had no effect in low glucose (32,33). The NO/cGMP/PKG signaling defects observed in high glucose were reversible, albeit slowly: 1) basal and insulin-induced NOS activity were improved by removing O-GlcNAC modifications; 2) NO-induced cGMP synthesis was improved by tempol, and cGMP synthesis was restored by cinaciguat; and 3) prkg1/2 transcriptional downregulation was reversed by tempol or glycine. After osteoblasts were transferred from high to low glucose medium, prkg1/2 transcription gradually increased, compatible with loss of O-GlcNAC modifications due to slow protein turnover. SP-1 is a ROS-sensitive transcriptional regulator modified by O-GlcNAC and required for prkg1 promoter activity (18). Other candidate transcription factors include Nrf-2, and we are presently characterizing oxidative stress–responsive transcriptional regulators and high glucose–induced changes in the osteoblast transcriptome. Thus, hyperglycemia alters NO/cGMP/PKG signaling at every step, leading to long-term cellular dysfunction even after correction of hyperglycemia (hyperglycemic memory).

**Oxidative Stress in Diabetes**

Biomarkers of oxidative stress are elevated in patients with type 1 and 2 diabetes, and increased oxidative stress contributes to the development of diabetic cardiovascular and kidney disease (15,46). Oxidative stress also appears to contribute to diabetic bone loss, because 1) transgenic over-expression of thioredoxin attenuates osteoporosis in mice with streptozotocin-induced diabetes (16); 2) antioxidant flavonoids improve bone volume in rats with type 1 diabetes (47); and 3) oxidative stress inhibits osteoblast differentiation and induces osteoblast death in vitro (48).

Increased ROS production in diabetic tissues has been attributed to increased NADPH oxidase activity, NOS uncoupling, and enhanced mitochondrial superoxide (O2−) production (15,19,34,49,50). We identified nox4 upregulation as the major source of increased H2O2 and oxidative stress in diabetic osteoblasts and bones: shRNA-mediated nox4 depletion reduced H2O2 production and lipid peroxidation in osteoblasts in high glucose to values found in low glucose, and cinaciguat reduced nox4 mRNA expression and DNA oxidation in femurs of diabetic mice to values in control mice. Increased nox4 expression has been observed in diabetic heart and kidney, and global nox4 deletion alleviates diabetes-induced kidney injury (51). How NOX4 contributes to diabetic bone loss will require generating osteoblast-specific nox4 knockout mice, because global nox4−/− mice have a high bone mass due to reduced osteoclast numbers and activity (52).

Mitochondrial O2− and H2O2 production contribute to age-related bone loss, based on mice lacking mitochondrial superoxide dismutase or expressing mitochondrial-targeted catalase in osteoblastic cells, respectively (53,54). We found increased mitochondrial O2− production in POBs exposed to high glucose. NOX4 localizes to both plasma membranes and mitochondria and may contribute to mitochondrial O2− and H2O2 production (36,51).

**Antioxidant Effects of cGMP/PKG2**

We uncovered novel antioxidant functions of cGMP in osteoblasts cultured in high glucose: cinaciguat and 8-CPT-cGMP increased the antioxidant genes txn1, txnrd1, and prdx1, while simultaneously downregulating the pro-oxidant genes nox4 and txnip. Cinaciguat induced the same changes in bone of diabetic mice and reduced markers of DNA oxidation under hyperglycemic conditions in vivo. Cinaciguat, 8-CPT–cGMP, and tempol did not change nox4 or txn1 mRNA under normoglycemic conditions, suggesting nox4 and txn1 are regulated by ROS-responsive transcription factors in hyperglycemic cells. We found that the antioxidant effects of cGMP in osteoblasts were mediated by
PKG2, whereas antioxidant effects of cGMP in endothelial cells are attributed to PKG1 posttranscriptionally upregulating catalase and glutathione peroxidase (55).

**Anabolic Effects of Cinaciguat in Diabetic Bone**

We identified reduced NO/cGMP/PKG signaling as a major mechanism of diabetic bone loss and showed that restoring cGMP synthesis with cinaciguat in type 1 diabetic mice reduced oxidative stress, improved bone formation, and attenuated trabecular and cortical bone loss. We showed previously that cinaciguat protects mice from ovariectomy-induced trabecular bone loss, but the drug had no significant effect on bone microarchitecture in control mice (26). Cinaciguat reduced osteocyte apoptosis in ovariectomized and diabetic mice. This antiapoptotic effect likely contributes to bone protection in both disease models because osteocyte apoptosis has been causally linked to trabecular and cortical bone loss (54,56). We found that cinaciguat reduced osteocyte apoptosis in ovariectomized mice, and in diabetic mice. This antiapoptotic effect likely contributed to bone protection in both disease models because osteocyte apoptosis has been causally linked to trabecular and cortical bone loss (54,56). We found that cinaciguat increased expression of runx2 and bglap mRNA in vivo and in POBs and BMSCs in vitro, suggesting a stimulatory effect on osteoblastic differentiation. In contrast, Jafari et al. (57) found that the nonspecific protein kinase inhibitor H-8 promoted osteoblast differentiation of telomerase-overexpressing human BMSCs and suggested the effect of H-8 was via PKG1 inhibition because it was partly mimicked by small interfering RNA knockdown of PKG1. The differences in our results and those obtained by Jafari et al. (57) could be due to differences in cell type and differentiation protocol but could also relate to off-target effects of a non-specific protein kinase inhibitor.

Cinaciguat is a prototype of recently developed NO- and heme-independent sGC activators, and, other than hypotension at high doses, was well tolerated in clinical trials of cardiovascular diseases (32). The cinaciguat dose we used does not affect systolic blood pressure (26). We conclude that hyperglycemia-induced disruption of NO/cGMP/PKG signaling contributes to oxidative stress and bone loss in mice with type 1 diabetes and that NO/cGMP/PKG pathway activators present a novel paradigm for enhancing bone formation under conditions of hyperglycemia and oxidative stress.

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