Decreased osteoclastogenesis and high bone mass in mice with impaired insulin clearance due to liver-specific inactivation of CEACAM1

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ABSTRACT

Type 2 diabetes is associated with normal-to-higher bone mineral density (BMD) and increased rate of fracture. Hyperinsulinemia and hyperglycemia may affect bone mass and quality in the diabetic skeleton. In order to dissect the effect of hyperinsulinemia from the hyperglycemic impact on bone homeostasis, we have analyzed L-SACC1 mice, a murine model of impaired insulin clearance in liver causing hyperinsulinemia and insulin resistance without fasting hyperglycemia. Adult L-SACC1 mice exhibit significantly higher trabecular and cortical bone mass, attenuated bone formation as measured by dynamic histomorphometry, and reduced number of osteoclasts. Serum levels of bone formation (BALP) and bone resorption markers (TRAP5b and CTX) are decreased by approximately 50%. The L-SACC1 mutation in the liver affects myeloid cell lineage allocation in the bone marrow: the (CD3−CD11b−CD45R−) population of osteoclast progenitors is decreased by 40% and the number of (CD3−CD11b−CD45R+) B-cell progenitors is increased by 60%. L-SACC1 osteoclasts express lower levels of c-fos and RANK and their differentiation is impaired. In vitro analysis corroborated a negative effect of insulin on osteoclast recruitment, maturation and the expression levels of c-fos and RANK transcripts. Although bone formation is decreased in L-SACC1 mice, the differentiation potential and expression of the osteblast-specific gene markers in L-SACC1-derived mesenchymal stem cells (MSC) remain unchanged as compared to the WT. Interestingly, however, MSC from L-SACC1 mice exhibit increased PPARγ2 and decreased IGF-1 transcript levels. These data suggest that high bone mass in L-SACC1 animals results, at least in part, from a negative regulatory effect of insulin on bone resorption and formation, which leads to decreased bone turnover. Because low bone turnover contributes to decreased bone quality and an increased incidence of fractures, studies on L-SACC1 mice may advance our understanding of altered bone homeostasis in type 2 diabetes.

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Introduction

Patients with type 2 diabetes mellitus (T2DM) exhibit normal-to-higher bone mineral density (BMD) than nondiabetics. Despite higher BMD, T2DM patients are at a 2-fold higher risk for fracture [8]. This suggests that instead of bone mass, diabetes-related factors are essential determinants of bone quality and increased fracture risk.

Hyperglycemia and hyperinsulinemia are cardinal features of insulin resistance in T2DM and insulin appears to have an anabolic effect on bone [21]. In vitro analyses have shown that insulin decreases the resorptive activity in osteoclasts, but stimulates proliferation and differentiation while inhibiting apoptosis in osteoblasts [4,20,21]. Insulin action on bone may involve direct signaling through the insulin receptor, activation of bone anabolic IGF-1 signaling by binding to IGF-1 receptor, or synergistic effects with other anabolic agents such as parathyroid hormone (PTH) [21,24].

Most of the studies on the effect of insulin on bone are based on in vitro models. Available in vivo models of altered insulin metabolism are usually in the context of either high glucose levels or changes in the growth hormone/IGF-1 signaling axes, which may confound the effect of insulin on bone.

The carcinoembryonic antigen-related cell adhesion molecule 1 (CEACAM1) is a transmembrane glycoprotein belonging to the immunoglobulin superfamily [10]. The CEACAM1 regulates insulin action by promoting insulin clearance [5,15,25]. The CEACAM1
C-terminal serine (SS03) is a substrate for the insulin receptor tyrosine kinase and a regulator of receptor-mediated insulin endocytosis and its degradation. The dominant negative SS03A mutation prevents CEACAM1 phosphorylation and affects insulin clearance. Transgenic L-SACC1 animals overexpress CEACAM1 (SS03A) specifically in the liver [17]. The L-SACC1 mouse exhibits hyperinsulinaemia and glucose intolerance, due to impairment of hepatic insulin clearance [16]. Moreover, L-SACC1 mice have increased visceral adiposity with associated macrophage infiltration. The mice do not develop overt diabetes, as assessed by normal fasting glucose levels. Thus, we, herein, used the L-SACC1 mouse to investigate whether elevations in insulin levels cause an increase in bone mass and alter bone cell phenotype in the absence of altered glucose levels.

**Materials and methods**

**Animals**

Transgenic L-SACC1 mice were generated as previously described [17]. Animals were housed on 12-h dark/light cycle and fed standard chow and water ad libitum. All procedures were approved by the University of Toledo Health Science Campus Institutional Animal Care and Utilization Committee. Only male animals were used in presented experiments. To permit dynamic bone histomorphometry, mice were injected ip with 30 μg/g body weight of tetracycline 7 and 2 days before sacrifice. Animals were euthanized by CO2 inhalation and cervical dislocation. Weights of epididymal fat corresponding to white adipose tissue (WAT) and interscapular fat corresponding to brown adipose tissue (BAT) were recorded. Bone marrow was harvested from both femora. Right tibia and L5 vertebrae were analyzed using micro-computed tomography (mCT), whereas left tibia was decalcified and embedded in paraffin for histological assessment. After mCT, the right tibiae were embedded undecalcified in methyl methacrylate and analyzed for dynamic histomorphometry at the Center for Orthopaedic Research, University of Arkansas for Medical Sciences (Little Rock, AR), as previously described [11].

**Measurements of metabolic serum parameters**

Animals were fasted for 18 h and serum samples were prepared from blood collected by cardiac puncture immediately after euthanasia. Serum insulin levels and IGF-1 levels were measured using immunoassays provided by ALPCO Diagnostics (Salem, NH) and MECORE Laboratory (Bangor, ME), respectively. Bone-specific alkaline phosphatase was measured colorimetrically using the Alkaline Phosphatase Diagnostic Kit (Sigma). In the coculture experiments, nonadherent PBMC were plated over U-33/γ2 cells, which represent marrow cells of mesenchymal lineage naturally producing RANKL and M-CSF and able to support osteoclastogenesis [11]. U-33/γ2 cells were plated at the density of 1 × 105/cm2 on 48-well/plate 1 day before nonadherent PBMC were added (at 2 × 105/cm2). Cultures were grown in the media supplemented with 10−8 M 1,25-dihydroxy vitamin D3 for 8 days with media changed on day 4.

**Fluorescence-activated cell sorting (FACS)**

Freshly isolated bone marrow was subjected to FACS using FACS-Calibur (BD Pharmingen, Franklin Lakes, NJ) according to the standard protocol. Sorting was performed using following antibodies, CD3-PE (T-cell lineage) (cat #555275), CD11b-FITC (macrophage lineage) (cat #553310), CD45R-APC (B-cell lineage) (cat #553092), and CD16/CD32 (blocking nonspecific binding) (cat #553142), which were purchased from BD Pharmingen. Initial sorting, which excluded cell aggregates and dead cells, was followed by sorting for CD11b. The cell population negative for this antigen was subsequently sorted for both CD3 and CD45R antigens. The population of cells negative for all three antigens (CD11b−CD3−CD45R−) was considered osteoclast precursors [7].

**Analysis of gene expression in mature osteoclasts**

Nonadherent marrow cells harvested after 24 h growth of PBMC, as described above, were plated at the density of 2.5 × 105 cells/cm2 in the presence of growth medium (α-MEM and 15% FBS) supplemented with 50 ng/ml M-CSF. Cells were incubated for 3 days, then received medium supplemented with 50 ng/ml M-CSF and 50 ng/ml RANKL. After additional 3 days of growth, cells were harvested and total RNA isolated using TRIzol reagent (Invitrogen, Carlsbad, CA). Gene expression analysis was performed using real-time PCR (StepOnePlus, Applied Biosysytem, Foster City, CA), as previously described [11]. Relative expression was measured by the comparative C规章制度 method after each sample was normalized to the quantity of 18S RNA. Real-time PCR analysis was performed using previously described primers sets: c-fos, RANK, and M-CSF-R [23]. IGF-1 (exon 3 and exon 6) and IGF-1r [13]. PPARγ (both isoforms), PPARγ2, Runx2, Dlx5, RANKL, osteocalcin, collagen I and RANKL [11].

**Analysis of marrow mesenchymal cells**

To assess the potential of MSC to differentiate toward osteoblasts, PBMC were seeded as a colony forming units type of culture and analyzed as described previously [11]. Gene expression analysis was performed on total RNA isolated from PBMC cultured for 10 days, as described [12]. To assess the effect of insulin on MSC proliferation and alkaline phosphatase activity, PBMC from the wild-type strain were plated on 96-well plates at the density 1 × 105 cells/well. Cultures were grown for 10 days either in the absence or presence of insulin (0.1 μM or 1 μM concentration). Cell proliferation was assayed by MTT assay (Promega) and alkaline phosphatase activity was measured and normalized to the cell number, as previously described [12]. Each experiment was repeated three times.

**Micro-computed tomography analysis**

mCT analysis was performed using a SCANCO μCT35 (SCANCO Medical AG, Bassersdorf, Switzerland) equipped with a 10-mm focal spot microfocus x-ray tube. Scans were performed at the following instrument settings: E = 70 KVP, I = 110 μA, increment 7 μm, threshold value = 289 [6,11]. Two hundred slices of the proximal tibia was decalcified and analyzed using micro-computed tomography (mCT), whereas left tibia was decalcified and embedded in paraffin for histological assessment. After mCT, the right tibiae were embedded undecalcified in methyl methacrylate and analyzed for dynamic histomorphometry at the Center for Orthopaedic Research, University of Arkansas for Medical Sciences (Little Rock, AR), as previously described [11].

**Osteoclastogenesis assay**

Primary bone marrow cells (PBMC) were harvested as described previously [12], and seeded at the density of 2.5×105/cm2 in the presence of α-MEM (Invitrogen, Carlsbad, CA) supplemented with 15% FBS (HyClone, Waltham, MA). Floating, nonadherent cells were harvested after 24 h and seeded at the density of 2 × 105/cm2 into 48-well plate with the medium supplemented with M-CSF (50 ng/ml) and RANKL (50 ng/ml) (R&D System, Minneapolis, MN). After 4 days of growth, cultures were stained for TRAP5b using the Leukocyte Acid Phosphatase (TRAP5b) Kit (Sigma). In the coculture experiments, nonadherent PBMC were plated over U-33/γ2 cells, which represent marrow cells of mesenchymal lineage naturally producing RANKL and M-CSF and able to support osteoclastogenesis [11]. U-33/γ2 cells were plated at the density of 1 × 105/cm2 on 48-well/plate 1 day before nonadherent PBMC were added (at 2 × 105/cm2). Cultures were grown in the media supplemented with 10−8 M 1,25-dihydroxy vitamin D3 for 8 days with media changed on day 4.
Statistical analysis was performed on three independent experiments, each of them consisted of age-matched groups of L-SACC1 and WT animals (n=4 animals per group). Statistically significant differences between groups in each experiment were detected using one-way ANOVA within the SigmaStat software (SPSS, Inc., Chicago, IL). In all cases, \( p < 0.05 \) was considered significant.

Dynamic histomorphometric data were collected from one experiment, 4 animals per group, and differences between parameters determined using ANOVA within the SigmaStat software (SPSS Science, Chicago, IL). All values are presented as the mean \( \pm \) SD. Differences were considered significant if \( p < 0.05 \).

### Results

**L-SACC1 mice are hyperinsulinemic, obese and have changes in fat distribution**

Consistent with previous reports [17], 5-month-old L-SACC1 males exhibited elevation in body weight (30.3 \( \pm \) 2.5 g vs. 23.5 \( \pm \) 2.7 g; \( p < 0.05 \)) with increased visceral fat weight (0.874 \( \pm \) 0.069 g vs. 0.19 \( \pm \) 0.05 g).

### Table 1

<table>
<thead>
<tr>
<th></th>
<th>BV (mm³)</th>
<th>BV/TV (%)</th>
<th>Tb. N. (1/mm²)</th>
<th>Tb. Th. (mm)</th>
<th>Tb. Sp. (mm)</th>
<th>ConnD (1/mm³)</th>
<th>SMI</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Prox. tibia</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>0.20 ( \pm ) 0.04</td>
<td>8.1 ( \pm ) 0.02</td>
<td>3.43 ( \pm ) 0.75</td>
<td>0.022 ( \pm ) 0.002</td>
<td>0.279 ( \pm ) 0.062</td>
<td>361.7 ( \pm ) 105.1</td>
<td>2.6 ( \pm ) 0.2</td>
</tr>
<tr>
<td>LS</td>
<td>0.42 ( \pm ) 0.05</td>
<td>14.1 ( \pm ) 0.02</td>
<td>4.94 ( \pm ) 0.54**</td>
<td>0.028 ( \pm ) 0.002**</td>
<td>0.176 ( \pm ) 0.022**</td>
<td>667.6 ( \pm ) 181.2**</td>
<td>1.8 ( \pm ) 0.2*</td>
</tr>
<tr>
<td><strong>Vertebrae</strong></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>0.12 ( \pm ) 0.04</td>
<td>7.9 ( \pm ) 0.04</td>
<td>2.83 ( \pm ) 1.14</td>
<td>0.021 ( \pm ) 0.004</td>
<td>0.366 ( \pm ) 0.116</td>
<td>161.8 ( \pm ) 167.7</td>
<td>3.2 ( \pm ) 0.6</td>
</tr>
<tr>
<td>LS</td>
<td>0.19 ( \pm ) 0.05**</td>
<td>10.6 ( \pm ) 0.02</td>
<td>3.78 ( \pm ) 0.41</td>
<td>0.024 ( \pm ) 0.001</td>
<td>0.243 ( \pm ) 0.034</td>
<td>297.6 ( \pm ) 51.0</td>
<td>2.7 ( \pm ) 0.2</td>
</tr>
</tbody>
</table>

All values are expressed as means \( \pm \) SD.

* \( n=4 \) animals per group.

* \( p<0.01 \) vs. WT.

** \( p<0.05 \) vs. WT.

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0.321 ± 0.091 g; p < 0.05) and decreased interscapular fat mass 
(0.059 ± 0.024 g vs. 0.093 ± 0.010 g; p < 0.05). They also had higher 
fasting levels of serum insulin (168 ± 51.8 PM vs. 48.5 ± 2.1 PM; 
< 0.05) and there is no difference in fasting levels of glucose between 
L-SACC1 and WT animals (103.2 ± 5.3 mg/dL vs. 109.7 ± 7.7 mg/dL). 
The L-SACC1 animals used in these studies were hyperinsulinemic 
and obese, but normoglycemic, with changes in fat distribution 
indicating changes in the energy metabolism.

L-SACC1 mice have increased trabecular and cortical bone mass, 
decreased bone formation, and decreased number of osteoclasts

L-SACC1 had higher content of trabecular bone in the tibia and 
vertebrae as compared to wild-type (WT) mice (Figs. 1A and B, Table 1). 
The trabecular bone volume (BV/TV) in the tibia was increased by 
74% (p = 0.0004), connectivity density (ConnD) by 84% (p = 0.006), 
number of trabeculae (TbN) by 44% (p = 0.018), and trabecular 
thickness (TbTh) by 26% (p = 0.008). Consistently, the trabecular 
separation (TbSp) was decreased by 37% (p = 0.02). The measure of 
structure model index (SMI) indicated that L-SACC1 trabecular bone 
architecture was more platelike, in contrast to more rodlike architecture 
seen in control animals (Fig. 1A and Table 1). Due to large 
variations between animals, the vertebral measurements did not 
achieve statistical significance as compared to the control group 
(Table 1).

Measurements of cortical bone geometry from the midshaft of the 
tibia showed that L-SACC1 bone had a larger cortical perimeter due to 
increased cortical area (22.5%; p = 0.02) and cortical thickness (11.4%; 
p = 0.02) (Fig. 1C and Table 2). Although being larger and thicker, 
L-SACC1 bone was not osteopetrotic and had a slightly increased 
medullary area (8.3%, NS).

Dynamic histomorphometry of the trabecular tibia bone showed 
atenuated bone formation in L-SACC1 mice (Fig 1D and Table 3). 
Assessment of bone formation based on incorporation of 
tetracycline into newly formed bone shows that only 5% of 
L-SACC1 bone is active as compared to approximately 70% of 
WT bone. Due to an insufficient number of double-labeled bone 
surfaces in L-SACC1 bone evaluations of the mineral apposition rate 
(MAR) and the bone formation rate (BFR) were not possible (Fig. 1D 
and Table 3).

The histologic assessment of trabecular bone in the tibia showed a 
decrease in the number of TRAP-positive osteoclasts in the L-SACC1 
bone, as compared to WT (Fig. 1E). The number of adipocytes in the 
L-SACC1 bone marrow was 3-fold higher than WT (Fig. 1F).

The effects of L-SACC1 mutation on serum bone turnover markers

Consistent with a lower number of osteoclasts, the levels of bone 
resorption markers, such as TRAP5b enzyme activity and CTX, a 
marker of collagen degradation, were significantly lower in serum of 
L-SACC1 animals (Table 4). The activity of bone-specific alkaline 
phosphatase was lower in L-SACC1 than in control mice, however 
the level of P1NP, a marker of collagen production, was higher in 
L-SACC1 animals (Table 4). Levels of circulating IGF-1 cytokine, which 
has an anabolic effect on bone, were not different between both 
strains.

### Table 2

Cortical parameters of midshaft of tibia.

<table>
<thead>
<tr>
<th></th>
<th>Cross-section area (mm²)</th>
<th>Cortical area (mm²)</th>
<th>Medullar area (mm²)</th>
<th>Cortical thickness (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>0.203 ± 0.017</td>
<td>0.127 ± 0.009</td>
<td>0.076 ± 0.009</td>
<td>0.184 ± 0.011</td>
</tr>
<tr>
<td>L-SACC1</td>
<td>0.238 ± 0.029</td>
<td>0.155 ± 0.016*</td>
<td>0.083 ± 0.015</td>
<td>0.205 ± 0.011*</td>
</tr>
</tbody>
</table>

All values are expressed as means ± SD.
* n = 4 animals per group.
* p < 0.05 vs. WT.

### Table 3

Dynamic histomorphometry of L-SACC1 (LS) and wild-type (WT) tibia trabecular bone.

<table>
<thead>
<tr>
<th></th>
<th>Single-labeled surface/BS (%)</th>
<th>Double-labeled surface/BS (%)</th>
<th>Nonlabeled surface/BS (%)</th>
<th>MAR (μm/d)</th>
<th>BFR/BS (μm²/mm²/d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>142 ± 11.1</td>
<td>61.4 ± 9.2</td>
<td>35.4 ± 7.7</td>
<td>0.75 ± 0.03</td>
<td>0.27 ± 0.05</td>
</tr>
<tr>
<td>LS</td>
<td>43.4 ± 3.5*</td>
<td>1.4 ± 1.0*</td>
<td>94.3 ± 2.2*</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

Values are expressed as means ± SD of n = 4 animals per group. BS indicates bone surface; MAR, mineral apposition rate; BFR, bone formation rate; ND, not detected.
* p = 0.001 L-SACC1 vs. WT.

Reduced number and impaired differentiation of osteoclast precursors from L-SACC1 mice

FACS analysis of freshly isolated bone marrow showed that the 
lineage distribution of myelopoietic cells in L-SACC1 and WT marrow 
was different. The number of osteoclast precursors, measured as a pool 
of cells negative for CD11b, CD3, and CD45R, was 40% lower in 
L-SACC1 than WT mice (Table 5). The number of CD45⁺ cells, which 
represent B-cell precursors, was increased by 60%. No differences in 
the number of cells positive for CD11b (macrophage marker) and CD3 
(T-cell marker) were observed (Table 5).

To further address the issue of decreased osteoclastogenesis 
observed in L-SACC1 mice, we compared the efficiency of osteoclast 
recruitment from the pool of hematopoietic cells and their differentiation 
toward osteoclasts in the presence of either an excessive amount of exogenously added RANKL and M-CSF cytokines, or a limited supply of these cytokines such as coculturing with U-33/ 
γ2 cells [11,14]. The number of TRAP⁺ osteoclastlike cells in L-SACC1 
was lower by 40% when cultured in the presence of exogenously 
added cytokines (Fig. 2A), and by 20-fold when the cytokines supply 
was limited (Fig. 2B). The more marked reduction in TRAP⁺ cells in 
the condition of limited cytokine supply indicates impairment in 
differentiation potential of existing osteoclast precursors. Moreover, 
there was a notable difference in the maturation of TRAP⁺ cells in both conditions, with those from WT nonadherent cells being more mature (appearing large and possessing multiple nuclei) than the majority of TRAP⁺ cells from L-SACC1 precursors which were smaller and possessed a low number of nuclei (Fig. 2C).

Analysis of the expression of genes essential for intrinsic 
regulation of osteoclastogenesis showed that osteoclast precursors 
from L-SACC1 animals expressed lower levels of c-fos and RANK gene 
transcripts (Fig. 2D). Products of these genes are essential for RANKL 
signaling and osteoclast differentiation [3]. The level of receptors 
for other osteoclast-supporting factors, M-CSF and IGF-1, was not 
different between L-SACC1 and WT animals. Similarly, transcript 
levels for PPARγ nuclear receptor, which has been recently identified as a transcriptional regulator of c-fos and RANK expression in 
osteoclasts [23], was not altered (data not shown).

Insulin decreased osteoclastogenesis and the expression of c-fos and RANK in nonadherent PBMC

Since L-SACC1 animals have elevated circulating levels of insulin, 
we tested the effect of insulin on WT-derived osteoclast differentiation

### Table 4

Serum bone turnover markers.

<table>
<thead>
<tr>
<th></th>
<th>WT*</th>
<th>L-SACC1*</th>
</tr>
</thead>
<tbody>
<tr>
<td>TRAP5b (U/L)</td>
<td>0.428 ± 0.073</td>
<td>0.216 ± 0.068</td>
</tr>
<tr>
<td>CTX (ng/ml)</td>
<td>15.60 ± 2.48</td>
<td>10.33 ± 3.50</td>
</tr>
<tr>
<td>BALP (μg/min)</td>
<td>0.026 ± 0.003</td>
<td>0.014 ± 0.002</td>
</tr>
<tr>
<td>P1NP (ng/ml)</td>
<td>1.44 ± 0.37</td>
<td>2.09 ± 0.25</td>
</tr>
<tr>
<td>IGF1 (ng/ml)</td>
<td>296.3 ± 30.9</td>
<td>297.3 ± 34.0</td>
</tr>
</tbody>
</table>

TRAP5b indicates tartrate-resistant acid phosphatase form 5b; CTX, C-terminal telopeptide of type I collagen; BALP, bone-specific alkaline phosphatase; P1NP, N-terminal propeptide of type I procollagen; IGF1, insulinlike growth factor 1. *
* n = 4 animals per group, 3 experiments.

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and osteoclast-specific gene expression in vitro. As shown in Fig. 3A, an addition of insulin to the ex vivo osteoclast cell cultures affected the number of TRAP$^+$ cells and their multinucleation. In the presence of 0.1 μM insulin the number of TRAP$^+$ cells was decreased by 40% (Fig. 3B) and the number of multinucleated TRAP$^+$ cells was reduced by 20% (Fig. 3C). Interestingly, in the presence of 1 μM insulin the number of TRAP$^+$ cells was decreased by 50% (Fig. 3B), while the number of multinucleated cells was reduced by more than 90% (Fig. 3C). These results are consistent with in vivo findings and indicate that insulin affects both, recruitment and osteoclast differentiation, however by different mechanisms which are determined by the dose of insulin in in vitro conditions.

An analysis of gene expression showed that insulin affects the expression of c-fos and RANK but not M-CSF and IGF-1 receptors (Fig. 3D). This change of gene expression by insulin in vitro is identical to that of osteoclasts derived from L-SACC1 animals.

**The effects of L-SACC1 mutation and insulin on mesenchymal stem cell (MSC) phenotype**

The differentiation potential of MSC toward osteoblasts and adipocytes was measured in CFU-OB and CFU-AD type of culture, respectively, and was not different between L-SACC1 and WT animals (data not shown). Similarly, mRNA levels of osteoblast-specific genes for Runx2, Dlx5, RANKL, osteocalcin and collagen I, were not different in MSCs isolated from L-SACC1 and WT animals (data not shown). Although the levels of circulating IGF-1 were not different (Table 4), however the expression of the bone-specific long transcript for IGF-1 was decreased and that of PPARγ2, an adipocyte-specific transcription factor, was increased in MSC of L-SACC1 animals (Fig. 4A). Increased expression of PPARγ2 correlates with the increased number of adipocytes in the marrow of L-SACC1 mice (Fig. 1G). Ex vivo analysis of insulin effects on PBMC showed that, at the tested doses, insulin neither affected MSC proliferation (Fig. 4B) nor MSC alkaline phosphatase activity (Fig. 4C).

**Discussion**

The current studies present evidence for an importance of efficient hepatic insulin clearance in the maintenance of bone homeostasis. In the L-SACC1 murine model we have shown that impaired insulin clearance in the liver, which results in high levels of circulating insulin and insulin resistance in peripheral tissues, leads to decreased bone turnover. It affects bone remodeling process by decreasing both, bone

<table>
<thead>
<tr>
<th>OCP (CD1−CD11b−CD45R−)</th>
<th>Macrophage (CD11b+)</th>
<th>B cell (CD3−CD11b−CD45R+)</th>
<th>T cell (CD3−CD11b−CD45R−)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT$^a$ 12.48 ± 3.09</td>
<td>33.23 ± 5.27</td>
<td>15.3 ± 5.04</td>
<td>2.3 ± 0.22</td>
</tr>
<tr>
<td>LS$^a$ 7.9 ± 1.1$^a$</td>
<td>32.23 ± 1.78</td>
<td>24.13 ± 1.83$^a$</td>
<td>2.0 ± 0.14</td>
</tr>
</tbody>
</table>

All values are expressed as means ± SD and represent a percent of sorted PBMC. OCP indicates osteoclast progenitors.

$^a$ n = 4 animals per group.

$^a$ p < 0.05 vs. WT.
formation and bone resorption. It also changes myelopoietic cell commitment toward the osteoclast and B-cell lineages. We have demonstrated in both, in vivo and in vitro systems, that high levels of insulin affect recruitment and differentiation of osteoclasts by impairing the RANKL signaling pathway. In osteoclasts, expression of the RANK receptor is under the control of the c-fos transcription factor [1], and insulin modulates the expression of both genes. Using PBMC derived from normoinsulinemic animals, we showed that exogenously added insulin affects osteoclast differentiation and the expression of RANK and c-fos in a manner identical to L-SACC1 animals. These findings provide evidence that insulin negatively regulates the phenotype of cells of the osteoclast lineage, and suggest that systemic hyperinsulinemia has a permanent effect on osteoclast progenitors.

The effect of insulin on cells of mesenchymal lineage is less evident. Differentiation potential of MSCs derived from L-SACC1 animals toward osteoblast lineage and the expression of osteoblast gene markers were not different from MSCs derived from WT animals, with exception to IGF-1 which expression was decreased. In contrast to studies of other’s [4,21], insulin neither increase alkaline phosphatase activity nor cell proliferation when added to the PBMC culture. However, serum parameters of bone formation in L-SACC1 were significantly different from that of WT animals. These data suggest that systemic changes, which result from CEACAM1 mutation in the liver, affect osteoblast function in the bone in an indirect manner, which does not involve intrinsic changes in MSC differentiation potential. In contrast to cells of hematopoietic lineage, in vitro effect of insulin on MSC osteoblast phenotype does not recapitulate attenuated bone formation in L-SACC1 animals, suggesting that CEACAM1 mutation in the liver affects marrow mesenchymal cells by different mechanism than cells of hematopoietic lineage.

We have reported previously that PPARγ2, an adipocyte-specific transcription factor, suppresses the expression of IGF-1 in bone, especially its longer isoform which contains exon 6 [13]. Here, we showed that L-SACC1 mice are characterized by increased expression of PPARγ2 in marrow MSC. Thus, PPARγ2 may be responsible for decreased IGF-1 expression. Interestingly, although we did not observe an increase in CFU-AD formation in ex vivo conditions, L-SACC1 animals possess higher number of adipocytes in the bone marrow. The discrepancy between ex vivo and in vivo observation indicate that in in vivo conditions, an additional factor, perhaps insulin, which possesses proadipocytic activity [27], enhanced adipocytic differentiation of MSC. Taken together, these results suggest that prolonged exposure to high levels of insulin induces

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Fig. 3. In vitro analysis of insulin effects on osteoclast differentiation and gene expression. Nonadherent marrow cells were derived from WT animals (n = 4) and cultured separately as described in Materials and methods. The experiment was repeated 3 times. (A) An appearance of TRAP+ osteoclastlike cells after 4 days of culture in the presence of M-CSF and RANKL and insulin at different concentrations. (B) Calculated number of TRAP+ cells. (C) The ratio of multinucleated TRAP+ cells (more than 3 nuclei) to the total TRAP+ cells. (D) The effect of insulin on gene expression in osteoclastlike cells developed as in (A) and analyzed by real-time PCR. *p< 0.05 vs. control; #p< 0.05 vs. 0.1 μM insulin.

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formation, as assessed by low levels of alkaline phosphatase, patients and in diabetic Pima Indians showed reduced bone T2DM [2,9]. An analysis of serum bone turnover parameters in T2DM animals. contribute to the decreased bone formation rate observed in L-SACC1 proliferation. (C) The MSC were derived from bone marrow of WT animals (n=4) and assayed as 4 IGF-1, a positive regulator of osteoblast function[19], which may FIG. 4.

The PPARγ2 and IGF-1 gene expression in MSC derived from WT (black bars) and LS (gray bars) animals (n = 4 animals per group). (B) The in vitro effect of insulin on MSC proliferation. (C) The in vitro effect of insulin on alkaline phosphatase activity in MSC. The MSC were derived from bone marrow of WT animals (n = 4) and assayed as 4 separate cell isolates. The experiment was repeated 3 times. *p < 0.05; **p < 0.01.

the adipocyte phenotype in MSC and decreases the expression of

IGF-1, a positive regulator of osteoblast function [19], which may contribute to the decreased bone formation rate observed in L-SACC1 animals.

Several studies point toward a decreased bone turnover rate in T2DM [2,9]. An analysis of serum bone turnover parameters in T2DM patients and in diabetic Pima Indians showed reduced bone formation, as assessed by low levels of alkaline phosphatase, osteocalcin and IGF-1 [2]. Another study based on longitudinal observations of bone status showed attenuation of the rate of bone loss with aging and relatively higher bone mass in T2DM patients [9]. The higher bone mass in L-SACC1 animals is entirely consistent with low bone turnover, which results from attenuated bone resorption and decreased bone formation.

A paradox between normal or higher BMD and increased fracture risk in T2DM suggests altered quality in diabetic bone. In addition to hyperinsulinemia leading to low bone turnover, hyperglycemia may account for changes in bone biomaterial quality by modification of collagen fibers [18]. Highly reactive glucose metabolites (AGEs), of which circulating levels are increased in diabetic hyperglycemia, are implicated in forming cross-links between collagen fibers, which affect bone biomechanical properties by increasing its stiffness and fragility [22,26].

Using the L-SACC1 model of hyperinsulinemia caused by liver-specific impairment in insulin clearance, we have demonstrated that high insulin levels affect bone resorption and impair bone formation, leading to a decreased bone turnover rate. Because altered insulin action in L-SACC1 mice is primarily hepatic, our studies suggest that insulin metabolism in the liver, which regulates overall insulin action in peripheral tissues (muscle and adipose tissue) [16], regulates also the maintenance of bone mass. The similarity of the phenotype of L-SACC1 mice with hyperinsulinemia in humans emphasizes the relevance of the current studies in advancing our understanding of the altered bone homeostasis in diabetes.

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