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Linkage of Decreased Bone Mass with Impaired Osteoblastogenesis in a Murine Model of Accelerated Senescence

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Abstract

Bone marrow is the principal site for osteoclastogenesis and osteoblastogenesis; and an increase in the former has been linked with bone loss caused by acute loss of gonadal steroids. We have now used an established murine model of accelerated senescence and osteopenia (SAMP6) to test the hypothesis that reduced osteoblastogenesis is linked with decreased bone mass. At 1 mo of age, the number of osteoblast progenitors in SAMP6 marrow was indistinguishable from controls; however a threefold decrease was found at 3–4 mo of age. Impaired osteoblast formation was temporally associated with decreased bone formation and decreased bone mineral density, as determined by histomorphometric analysis of tetracycline-labeled cancellous bone and dual-energy x-ray absorptiometry, respectively. Osteoblastogenesis determined in ex vivo bone marrow cultures was also decreased in these mice, as was the number of osteoclasts in histologic sections. Moreover, unlike controls, senescence-accelerated mice failed to increase osteoclast development after gonadectomy. The osteoclastogenesis defect was secondary to impaired osteoblast formation as evidenced by the fact that osteoclastogenesis could be restored by addition of osteoblastic cells from normal mice. These findings provide the first demonstration of a link between low bone mineral density and decreased osteoblastogenesis in the bone marrow and validate the senescence-accelerated mouse as a model of involutional osteopenia. (J. Clin. Invest. 1996. 97:1732–1740.) Key words: osteoporosis • aging • remodeling • bone formation • osteoclast formation

Introduction

The skeleton is continuously remodeled during adult life with the resorption of old bone by osteoclasts and its subsequent replacement by osteoblasts. This process is carried out in a highly coordinated fashion with a specific temporal and spatial pattern. Disturbances in the process result in an imbalance between resorption and formation and are responsible for most metabolic bone diseases including osteoporosis. Many of the clinical and epidemiological features of osteoporosis can be accounted for by a two-stage model of bone loss (1, 2). In this model, sex hormone deficiency results in an increase in both osteoclast and osteoblast activity with the former exceeding the latter, resulting in rapid bone loss. This phase is followed by a period of slow bone loss due to aging (3–9). It is generally agreed that the phase of rapid bone loss does not continue indefinitely, but its duration varies considerably among individuals. Confounding factors such as age-associated changes in vitamin D status or age-related development of subclinical hyperparathyroidism may result in continued elevated bone remodeling in some individuals (10–13). Nevertheless, the existence of an age-induced mechanism of bone loss, distinct from age-related alterations in hormone levels, is indicated by the inability of hormone replacement therapy to prevent bone loss indefinitely and the occurrence of bone loss with advancing age in sex hormone-replete men (14, 15).

A consistent histologic feature of the osteopenia that occurs during aging is a decrease in wall thickness, especially in trabecular bone (16, 17). Wall thickness is a measure of the amount of bone formed in a remodeling packet of cells and is determined by the number and activity of osteoblasts at the remodeling site. Considerable evidence suggests that the decline in wall thickness is due to the aging process itself, as opposed to hormone changes that occur with advancing age, such as alterations in the level of sex steroids or parathyroid hormone (18, 19). Hence, the osteopenia due to aging may be due to a deficit in the number of osteoblasts recruited to the remodeling site relative to the demand for them (17).

The bone-forming osteoblast and the bone-resorbing osteoclast originate from mesenchymal and hematopoietic progenitors of the marrow, respectively. The development of osteoblasts and osteoclasts is governed by locally produced cytokines and growth factors that influence the differentiation of osteoclasts from colony-forming unit–granulocyte/macrophage (CFU-GM) and/or the differentiation of osteoblasts from colony-forming unit–fibroblast (CFU-F). Based on extensive histomorphometric data from humans, as well as the results of experimentation in animal models, we recently proposed the hypothesis that inappropriate production of osteoclasts or inadequate production of osteoblasts, as opposed to changes in the activity of these cells, is responsible for the mismatch between formation and resorption of bone in postmenopausal and age-related osteopenia, respectively (20).

To examine the relationship between osteoblast progenitor development in the bone marrow, bone remodeling, and bone mass, we have used the SAMP6 mouse, a model of early senes-

1. Abbreviations used in this paper: BMC, bone mineral content; BMD, bone mineral density; CFU-F, colony-forming unit–fibroblast; CFU-GM, colony-forming unit–granulocyte/macrophage; CFU-OB, colony-forming unit–osteoblast; DEXA, dual-energy x-ray absorptiometry; LDH, lactate dehydrogenase; TRAPase, tartrate-resistant acid phosphatase.
cence that exhibits osteopenia (21, 22). The SAMP6 strain as well as the “normal” control strain SAMR1 were developed over the past 20 yr by investigators at Kyoto University by brother–sister mating of AKR/J mice for > 50 generations. When reared under specific pathogen-free conditions, SAMP6 exhibit normal incidence of tumors and other specific abnormalities. Thus, the confounding effects of age-related diseases of inbred mice do not account for either the rapid aging or the osteopenia of SAMP6 mice. The osteopenic phenotype of this strain may have arisen from unfavorable combinations of alleles at multiple loci of the genome affecting bone mass. Alternatively, retroviruses from the AKR/J parental strain, or another aberrantly introduced strain, may have become activated when placed in a heterologous genetic background after a presumed (accidental) outcrossing at the initial stages of the development of this strain (23).

In this report we present evidence for a tight linkage among decreased osteoblastogenesis in the bone marrow, diminished bone formation in remodeling cancellous bone, and low bone mineral density (BMD).

**Methods**

**Animals.** SAMP6 mice and SAMR1 mice were obtained from a colony established from breeders kindly provided to us by Dr. Toshio Takeda of Kyoto University (Kyoto, Japan). Individual animals were electronically tagged (BioMedic Data System Inc., Maywood, NJ) at weaning. For some experiments, 3–4-mo-old mice were gonadectomized as described previously (24, 25). Neonatal C57Bl/6J mice were obtained from Harlan Sprague Dawley, Inc. (Indianapolis, IN). All animals were maintained and used in accordance with National Institutes of Health guidelines on the care and use of laboratory animals.

**Detection and quantification of osteoblast progenitors.** CFU-F were detected by the formation of a colony of adherent fibroblastoid cells in ex vivo marrow cell cultures (26). The presence of osteoblasts in these colonies was assayed by determining the ability of cells present in the colony to form a mineralized matrix (27). Mineralizing colonies were designated as colony-forming unit–osteoblast (CFU-OB). The number of CFU-F and CFU-OB in murine bone marrow was determined by limiting dilution assay (27). Marrow cells were obtained from the femur (24), and replicate cultures (n = 16–24) of serially diluted cells were established from each animal in 0.3-cm² wells of a 96-well plate at 6–8 seeding densities that ranged from ~300,000 to 10,000 cells per well. Cells were cultured in phenol red–free αMEM containing 15% preselected FBS, 50 μM ascorbic acid, and 10 mM β-glycerophosphate. One-half of the medium was replaced every 5 d.

For the determination of CFU-F, cultures were maintained for 10 d, fixed with neutral buffered formalin, stained for alkaline phosphatase (kit 180; Sigma Immunochemicals, St. Louis, MO), and counted. The number of wells that contained CFU-F or CFU-OB was determined as colony-forming unit–osteoblast (CFU-OB). The number of CFU-F and CFU-OB in murine bone marrow was determined by limiting dilution assay (27). Marrow cells were obtained from the femur (24), and replicate cultures (n = 16–24) of serially diluted cells were established from each animal in 0.3-cm² wells of a 96-well plate at 6–8 seeding densities that ranged from ~300,000 to 10,000 cells per well. Cells were cultured in phenol red–free αMEM containing 15% preselected FBS, 50 μM ascorbic acid, and 10 mM β-glycerophosphate. One-half of the medium was replaced every 5 d. For the determination of CFU-F, cultures were maintained for 10 d, fixed with neutral buffered formalin, stained for alkaline phosphatase (kit 180; Sigma Immunochemicals, St. Louis, MO), and counterstained with hematoxylin. Colonies containing > 50 cells were designated as CFU-F colonies. For the determination of CFU-OB, cells were cultured for 25 d, fixed in 50% ethanol, 18% formaldehyde, and stained for calcium deposits using either Von Kossa stain or alizarin red. The percentage of wells that contained CFU-F or CFU-OB was determined for each cell concentration and the number of progenitors per 10⁶ marrow cells was graphically determined as described previously (27).

**Analysis of individual CFU-F.** Marrow cells were seeded in a 96-well plate at 10,000, 20,000 or 30,000 per well (n = 48 wells for each seeding density) and cultured as described above for the determination of CFU-F. Cells were dried and then fixed with neutral buffered formalin (10%) at 4°C for 10 min. The activity of lactate dehydrogenase (LDH) was used as an index of the number of cells present in each well and was determined by coupling the enzyme reaction to the reduction of a tetrazolium salt to a formazan as described previously (28). Each well contained 100 μl of substrate solution consisting of 0.2 M Tris buffer, pH 8.2, 27 mM sodium (+)-lactate, 0.65 mM β-nicotinamide adenine dinucleotide, 1.3 U/ml diaphorase, and 0.3 mg/ml 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfo-pheny)-2H-tetrazolium, inner salt (MTS). The rate of MTS reduction was determined by measuring the absorbance at 490 nm over a 12–16-h period, using a microtiter plate reader. LDH activity was expressed as the change in OD₉⁰₀ per minute. Preliminary experiments indicated that as few as 50 MCT3-E1 osteoblastic cells could reliably be detected using this assay (data not shown). After determination of LDH activity, wells were rinsed with distilled water, and the alkaline phosphatase activity in each well was quantitated as follows. Each well contained 100 μl of 0.1 M 2-amino-2-methyl-1-propanol buffer, pH 10.4, and 7 mM p-nitrophenyl phosphate (kit 104; Sigma Immunochemicals). The production of p-nitrophenol was determined by measuring the absorbance at 402 nm over a 12–16-h period. One micromol of activity was defined as that amount of enzyme capable of hydrolyzing 1 pmol p-nitrophenolphosphate per minute at room temperature. Data on cell number (LDH activity) and alkaline phosphatase content were analyzed only in the case of wells that contained a single colony, determined by microscopic examination.

**Bone histomorphometry.** Mice were injected (intraperitoneally) with tetracycline hydrochloride (15 mg/kg body weight) at 8 and 3 d before killing. The distal femur and lumbar vertebrae were fixed at 4°C in phosphate-buffered 10% formalin, pH 7.4, containing 0.5% sucrose, and embedded in methyl methacrylate as described previously (29, 30). Uniform positioning of embedded bones was assured by placing them into a carved groove in a preformed, sticky methacrylate wafer at the bottom of each embedding vial. 5-μm-thick longitudinal sections taken from halfway through the specimens were cut on Zeiss 350 heavy-duty microtome (Carl Zeiss, Inc., Thornwood, NY) with a D profile knife (Delaware Diamond Knives Corp., Wilmington, DE). Osteoid and osteoblasts were best distinguished when stained with 0.25–1% toluidine blue, pH 2.8–3.2. Murine osteoclasts were identified by staining for tartrate-resistant acid phosphatase (TRAPase) (Sigma Immunochemicals). Adjacent 8-μm-thick sections were left unstained for epifluorescence observations.

Static and dynamic histomorphometric measurements were made as described previously using a computer and digitizer tablet (OsteoMeasure; Osteometrics, Inc., Atlanta, GA) interfaced to a Zeiss Axioscope (Carl Zeiss, Inc.) with a drawing tube attachment (29, 30). The tetracycline labels were measured at a magnification of 400 (numerical aperture 0.75). The terminology used is that recommended by the Histomorphometry Nomenclature Committee of the American Society for Bone and Mineral Research (31). To ensure that only remodeling bone of the secondary spongiosa was measured, measurements were restricted to the area between 1 and 5 mm distal to the growth plate–metaphyseal junction of the distal femur, and the central cancellous bone of the vertebrae. We avoided areas of bone undergoing modeling, specifically the endocortical margins of the diaphysis, and the cancellous bone furthest away from the growth plate. To estimate bone turnover, the rate of bone formation per unit of bone area was calculated as the bone formation rate per bone perimeter multiplied by the perimeter to area ratio, and expressed as the percentage of bone area remodeled per day. Values are expressed as means ± SD.

**Bone densitometry.** Dual-energy x-ray absorptiometry (DEXA) was performed on a QDR-2000 Plus densiometer (Hologic, Waltham, MA) that was calibrated daily with a hydroxyapatite phantom of the human lumbar spine. Each set of animal scans was preceded by scanning a plastic embedded murine phantom. Before scanning, animals were sedated (intramuscularly) with 120 μg/10 grams body weight sodium pentobarbital (Sigma Immunochemicals). Scans were done with the animals positioned prone. Measurements were made with a customized mouse whole body software package (Hologic) that allowed specified region of interest windows and subtraction of the animal identification transponder device. The global window was defined as the whole body image minus the calvarium, mandible, and teeth.
Duplicate BMD determinations with repositioning between the scans (n = 32 mice) showed that the mean difference in BMD between the two measurements was not significantly different from zero, thus proving that repositioning did not influence the measurement. The mean coefficients of variation for the short-term reproducibility studies were 2.17% for the global measurement, 2.19% for the spine, and 3.10% for the hindquarters. Long-term reproducibility was assessed by repeated measurement of the murine phantom. In a 6-mo period, 55 measurements were made with a coefficient of variation of 4.39%.

The accuracy of DEXA measurements was demonstrated by determining bone mineral content (BMC) and ash weight on the same murine bone samples as follows. Mice (n = 47) that were 23–378 d old and weighing 9.0–69.8 g were killed by CO2 inhalation. After determination of BMC by DEXA, the entire skeleton (excluding skull and tail) was harvested for ashing. In 24 of these animals, the spine, hindquarters, and remaining bones were each ashed separately. Ashing was done in a muffle furnace at 800°C for 24 h in covered quartz crucibles (Fisher Scientific, Pittsburgh, PA). After determining ash weight, data were analyzed by linear regression with ash weight as the independent variable and BMC as the dependent variable (Fig. 1). We observed a strong linear relationship between ash weight and global BMC (r = 0.96, P < 0.00001). The relationship between the ash weight and BMC of the spine and hindquarter subregions was also strong (r = 0.97 and 0.95, respectively, P < 0.00001). For the three relationships, the slope ranged from 0.928 to 1.194 and the intercept from −0.127 to 0.048. The relationships were not affected by sex, weight, or age of the mice.

Assay of osteoclast formation. Marrow cells (four to six replicate cultures) from each animal were cultured at 1.5 × 10^6 per 2-cm² well and maintained for 9 d in the presence of 10⁻⁸ M 1,25(OH)₂D₃ and 10% FBS in αMEM as described previously (24, 25, 32). We also used a coculture assay to quantify osteoclast progenitors (24). In these experiments, exogenous stromal/osteoblastic cells from neonatal murine calvaria cells were provided as support for osteoclast progenitors. Stromal/osteoblastic cells were prepared by sequential collagenase digestion of calvaria dissected from 3–5-d-old C57Bl/6J mice as described previously (24, 32, 33). Marrow cells (200,000 per 2-cm² well) were cocultured with 40,000 calvaria cells and maintained for 8 d in the presence of 10 nM 1,25(OH)₂D₃ and 10% FBS in αMEM. The number of osteoclasts formed was determined by counting cells that both bound 55Ca and expressed TRAPase. The number of osteoclasts formed per femur was calculated using the marrow cell yield from the animal.

**CFU-GM assay.** Marrow cells were cultured separately to establish three replicate cultures. Cells were plated at 100,000 cells per 10-cm² culture dish containing 1.5 ml of semisolid methylcellulose medium with pokeweed mitogen–stimulated spleen cell conditioned medium (No. M3430; Stem Cell Technologies, Inc., Vancouver, British Columbia, Canada). After 7 d, CFU-GM colonies (>50 cells) were enumerated. The number of CFU-GM formed per femur was calculated using the marrow cell yield from the animal.

Assessment of individual CFU-F colonies for ability to support osteoclast formation. Marrow cells were seeded at 20,000 per well in each well of a 96-well plate. After 5 d of culture spleen cells from normal Swiss/Webster mice were added as a source of osteoclast progenitors (34). Cells were incubated for 8 d with 10 nM 1,25(OH)₂D₃ to stimulate osteoclast formation. At the end of the experiment, each well was examined microscopically for the presence of CFU-F colonies and osteoclastic cells, identified by staining for TRAPase.

**Statistics.** Student’s t test was used to test for significant differences between means of two groups, after testing for equivalence of variances and normal distribution of data. Data which did not meet this criteria were analyzed by the Mann-Whitney rank sum test. Correlation coefficients were calculated to test for an association between two independently measured variables. Fisher’s exact test was used to test for significant differences in the proportion of positive elements in two populations.

**Results**

**Analysis of osteoblast progenitors of the bone marrow.** Growing SAMR1 and SAMP6 mice (~1 mo old) had identical numbers of CFU-F and CFU-OB in the bone marrow (Fig. 2 A). On the other hand, marrow cells from 3–4-mo-old adult SAMP6 mice contained approximately threefold fewer CFU-OB than SAMR1 (Fig. 2 B); but the number of CFU-F present in the marrow of SAMP6 and SAMR1 mice remained identical. Analysis of individual CFU-F colonies of 3-mo-old mice revealed that a larger proportion of the SAMP6 colonies was smaller and contained less alkaline phosphatase than colonies of SAMR1 mice (Fig. 3, A and B). The median cell number index for SAMR1 colonies was 0.44 × 10⁻³ OD/min vs. 0.11 × 10⁻³ OD/min in the case of SAMP6 colonies (P < 0.01 by Mann-Whitney rank sum test). The median alkaline phos-
phatase activity of SAMR1 colonies was 2.49 vs. 0.96 μU/col-

Bone histomorphometry. The cancellous bone of the distal femur and lumbar vertebra displayed numerous resorption and formation surfaces (Fig. 4). Cement lines or reversal lines, which are deposited after the osteoclasts resorb bone, demarcate the old bone from the new. The scalloped appearance of the reversal line reflects previous episodes of osteoclast action at the remodeling site. We used TRAPase staining to identify osteoclasts because we found that murine osteoclasts were dif-

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Concurrent determination of CFU-OB, by limiting dilution assay, was also performed on marrow cells from the mice used for histomorphometric studies shown in Table I. The number of CFU-OB was directly correlated with the vertebral mineral appositional rate \((r = 0.91, P = 0.032)\), to the vertebral cancellous bone area \((r = 0.88, P = 0.047)\), to the femoral mineralizing perimeter \((r = 0.88, P = 0.047)\), and to the femoral osteo-

The bone formation rates in the lumbar vertebrae of 3–4-wk-

Bone densitometry. The global, spine, and hindquarters BMD were measured in weaned growing animals (36–38 d old), in sexually mature young adults (93–128 d old) and in very old animals (436–486 d old). As shown in the data of Table II, the BMD of the three skeletal sites in the 36–38-d-old animals was similar in SAMP6 and SAMR1 mice. In 3–4-mo-old adults, there was a significant decrease in global and spine BMD of SAMP6 mice; but there was no significant difference in the hindquarters BMD. Very old SAMP6 mice showed signif-

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In view of the evidence that osteoblastogenesis is defective in marrow of adult SAMP6 mice, we examined the possibility that the marrow of SAMP6 mice was also deficient in the stromal/osteoblastic cells that are required for osteoclast formation. Marrow cells from SAMP6 mice formed significantly fewer osteoclasts than cells from SAMR1 mice (Fig. 5A). This finding could be due to a decrease in the number of osteoclast progenitors and/or a decrease in stromal/
osteoblastic support for osteoclast formation in ex vivo marrow cultures from SAMP6 mice. However, the number of CFU-GM was identical in marrow of SAMR1 and SAMP6 mice as determined in an ex vivo colony assay (Fig. 5B). Furthermore, osteoclast formation in ex vivo cultures of marrow cells from SAMP6 mice was slightly higher than SAMR1 mice when marrow cells were cultured together with exogenous stromal/osteoblastic cells prepared from calvaria of normal neonatal C57Bl/6J mice (Fig. 5C). In these cocultures, there was a four- to eightfold increase in the number of osteoclasts, compared with those seen in marrow cultures alone.

To directly establish whether marrow of SAMP6 mice had reduced numbers of osteoclastogenic support cells, we compared osteoclast formation in CFU-F colonies of SAMP6 or SAMR1 mice that were cocultured with spleen cells as a source of osteoclast progenitors, as described in Methods. We found that whereas 40% of the CFU-F colonies (total n = 60) from SAMR1 mice contained osteoclast-like cells, only 14% of the CFU-F colonies (total n = 35) derived from SAMP6 mice contained osteoclast-like cells. Thus, significantly fewer CFU-F colonies of SAMP6 mice contained stromal/osteoblastic cells capable of supporting osteoclast formation ($P < 0.02$ by Fisher’s exact test).

We and others have shown previously that the stromal/osteoblastic cells of the marrow are a target of the action of sex steroids to suppress IL-6 production and that IL-6 mediates the increase in osteoclast formation and the bone loss caused by loss of sex steroids (6, 24, 25, 32). In view of the evidence that marrow of SAMP6 mice is deficient in cells of the osteoblast lineage, we studied the effect of loss of sex steroids on osteoclast progenitors in SAMP6 mice. As shown in Fig. 6, loss of estrogens or androgens had no effect on osteoclast progenitors of the marrow of SAMP6 mice, regardless of whether osteoclast formation was assessed in marrow cells cultured alone or in marrow cells cocultured with stromal/osteoblastic cells from normal neonatal mice. Nevertheless, sex steroid deficiency caused the expected increase in osteoclast progenitors in SAMR1 mice.

### Table II. DEXA Determination of BMD

<table>
<thead>
<tr>
<th>Age (d)</th>
<th>Number analyzed</th>
<th>BMD (mg/cm²)</th>
<th>Global</th>
<th>Spine</th>
<th>Hindquarters</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SAMR1</td>
<td>SAMP6</td>
<td>SAMR1</td>
<td>SAMP6</td>
<td>SAMR1</td>
</tr>
<tr>
<td>33–38</td>
<td>13</td>
<td>15</td>
<td>37.5±2.6</td>
<td>33.6±3.2</td>
<td>41.5±3.3</td>
</tr>
<tr>
<td>91–149</td>
<td>46</td>
<td>47</td>
<td>50.6±3.2</td>
<td>48.0±1.5</td>
<td>56.9±3.3</td>
</tr>
<tr>
<td>463–486</td>
<td>6</td>
<td>6</td>
<td>51.7±4.6</td>
<td>39.2±2.0*</td>
<td>57.9±6.3</td>
</tr>
</tbody>
</table>

Data shown are the mean±SD. *$P < 0.05$ vs. SAMR1 of the same age; †$P < 0.05$ vs. 91–149-d-old SAMP6.
Discussion

Bone loss occurs when the amount of bone formed during each remodeling cycle does not exactly replace the bone removed during the preceding resorption phase. Theoretically, this could arise from (a) a reduction in the number of osteoblasts that differentiate from progenitors in the marrow; (b) a defect in the processes that cause the newly recruited team of differentiated osteoblasts to arrive at the remodeling site at the correct place and time; or (c) an increase in bone resorption.

The evidence presented in this report demonstrates that in SAMP6 mice the first scenario is responsible for a reduction in the rate of bone formation in remodeling trabecular bone, and the reduced bone mass. Thus, at \( \sim 1 \) mo of age, SAMR1 and SAMP6 mice have identical numbers of osteoblast progenitors in the marrow, identical rates of bone formation in remodeling trabecular bone, and identical BMD. However, by 3–4 mo of age, the number of osteoblast progenitors in the marrow of SAMP6 mice is reduced, and this is accompanied by a reduction in the rate of bone formation and diminished bone mass. Furthermore, a close relationship between osteoblastogenesis and both bone mass and formation is strongly suggested by the direct correlations between CFU-OB and spinal cancellous bone area, mineralizing perimeter, and mineral appositional rate. These observations are thus consistent with the accelerated aging that characterize these mice and establish that the osteopenic phenotype becomes apparent after sexual maturity.

It is important to note that the bone formation we measured in the secondary spongiosa of both growing juvenile mice and adult mice is due to remodeling activity. Indeed, we noted discrete erosion bays, as well as sites of bone formation associated with reversal lines in the cancellous bone we examined. Matsushita et al. (22) have reported previously that lon-
nitudinal growth of the femur and the age at which peak bone mass is attained are identical in SAMR1 and SAMP6 mice. Thus, at this stage, there is no evidence that alterations in growth or modeling in bone of SAMP6 mice contribute to the development of their osteopenia.

The osteopenic phenotype of SAMP6 was originally noted by the occurrence of “spontaneous” tibial fractures. Additional studies revealed a reduction in the cortical thickness of the diaphysis of the femur that became apparent at 3-4 mo of age (22). Although our histomorphometric measurements revealed significant differences in bone architecture, cell number, and the kinetics of bone formation in the cancellous bone of SAMR1 and SAMP6 mice, this approach can provide only limited information on overall bone mass due to the small size of the bone specimen available for study and the microheterogeneity of cancellous bone. Thus, although a significant reduction in cancellous bone was apparent in vertebral bone, the reduction in the distal femur was less impressive. However, a significant reduction in trabecular width was noted in the distal femur, indicating that the decreased number of osteoblast progenitors at this site was accompanied by trabecular bone atrophy. With further remodeling of this bone in the face of insufficient osteoblasts, additional bone loss will ensue.

Using a Hologic QDR-2000 Plus densitometer adapted for the measurement of bone mass by DEXA in live mice, we noted a reduction in bone mass in vertebral bone of SAMP6 mice at 3-4 mo of age. In addition, we observed that very old (15 mo) animals exhibited osteopenia in all skeletal sites examined. Thus, SAMP6 mice exhibit progressive loss of bone between 3 and 15 mo of age, a phenomenon which apparently did not occur in SAMR1 mice. However, as they approach 15 mo of age, a larger percentage of SAMP6 mice die, creating a group of unique survivors. Therefore, additional longitudinal studies are required to fully characterize age-related bone loss in these mice. In any event, the development of DEXA methodology for use in live mice, described here for the first time, makes such studies feasible and permits concurrent histomorphometric and BMD determinations. We believe that this methodologic advance will also allow exploitation of the wealth of information on the genetics and strain diversity in the mouse for studies on the development of osteopenia (35).

The marrow of adult SAMP6 mice has few osteoblast progenitors, but normal numbers of multipotential mesenchymal progenitors, CFU-F. Analysis of the properties of individual CFU-F colonies formed in ex vivo marrow of cultures indicated a considerable degree of heterogeneity with respect to the number of cells in each colony and the level of alkaline phosphatase expression, reflecting differences in the properties of the progenitors that gave rise to each colony. Comparison of individual colonies formed in cultures from SAMP6 and SAMR1 mice indicated that the mesenchymal progenitors of SAMP6 mice gave rise to smaller colonies than marrow of SAMR1 mice. These findings suggest that the marrow of SAMP6 mice is deficient in a population of rapidly dividing progenitors committed to the osteoblast lineage. Previous studies have indicated increased adipogenesis in marrow of SAMP6 mice (36). In addition, we have reported elsewhere evidence for an increase in hematopoietic activity in marrow of adult SAMP6 mice (37). These findings are consistent with the notion that there is an alteration in the pattern of differentiation of the progenitors of the mesenchymal cell lineage in the bone marrow of SAMP6 mice.

Because we found decreased osteoblast progenitors and reduced bone formation rate in bone of adult SAMP6 mice, we expected that the number of osteoblasts in bone would also be reduced. Although the mean number of osteoblasts was lower in cancellous bone of SAMP6 mice, the reduction was not statistically significant. This result may be due to high variation in histomorphometric measurements of osteoblasts, as noted previously (38). Compared with osteoblasts of rat or human bone, murine osteoblasts exhibited reduced size, were columnar rather than cuboidal, and exhibited less granularity. Due to their less distinctive features, therefore, we may have overestimated the number of mature osteoblasts that are synthesizing bone matrix using current histological criteria. Some of these cells could have been osteoblast progenitors or inactive osteoblasts. Development of new methods to identify matrix-synthesizing osteoblasts should allow more precise quantification of these cells.

The cancellous bone of adult SAMP6 mice exhibited a dramatic reduction in the number of osteoclasts, compared with bone of SAMR1 mice. Further, the marrow of SAMP6 mice, in contrast to the control SAMR1 mice, did not exhibit increased osteoclastogenesis when challenged by the removal of sex steroids. This phenomenon was due to a defect in the ability of stromal/osteoblastic cells in the marrow SAMP6 mice to support osteoclast formation and therefore was secondary to the defective osteoblastogenesis. In view of the extensive evidence that cells of the stromal/osteoblastic cell lineage are required for osteoclast formation, this finding was not unexpected. However, this is the first in vivo demonstration of this dependency. More important, the evidence that loss of sex steroids failed to increase osteoclastogenesis in the marrow of SAMP6 mice demonstrates the essential role of stromal/osteoblastic cells in the increased osteoclastogenesis caused by loss of sex steroids. In fact, this observation is consistent with the view that cells of the stromal/osteoblastic cell lineage are the principal targets of estrogens and androgens and therefore the essential mediators of the increased osteoclastogenesis caused by sex steroid deficiency (24, 25).

The trabecular bone of SAMP6 mice exhibits many features that are analogous to that of aging humans. In addition to osteopenia, the trabeculae were infrequent and widely separated in the spine, suggesting increased perforation by osteoclasts. Although osteoclast numbers were reduced in SAMP6 mice, the rate of bone turnover was still sufficiently rapid to explain the osteopenia in these mice. Based on the bone turnover measured at 3 mo of age, 0.5–4% of the cancellous bone is remodeled every 10 d. Thus, the osteoblast defect in SAMP6 was severe enough to preempt the bone formation required to reconstitute the previously eroded cavities. The reduced remodeling and the altered architecture of bone of SAMP6 mice are cardinal features of bone in aging humans. Thus, many cellular and architectural features of aging bone in humans are duplicated in this mouse model in a way that permits investigation of the cause of this osteopenia in animals that are the same age. Hence, the SAMP6 model should be useful for the exploration of the molecular and cellular basis of the relationship among osteoblastogenesis, bone formation in remodeling bone, and bone mass in a context that is similar to the age-related osteopenia that occurs in humans.

In conclusion, the demonstration of a close association between the development of a defect in osteoblastogenesis in a murine model of accelerated aging with resulting decreases in
osteoclast formation, bone formation, and bone mass strongly suggests the critical role of the bone marrow in the development of the osteopenia in this model. Because a decreased supply of osteoblasts may explain the diminished amount of bone formed during each remodeling cycle during aging in humans, our findings establish the validity of the SAMP6 mouse as an animal model of involutional osteoporosis. Finally, they pave the way for chromosome mapping studies aiming to elucidate genes that contribute to osteopenia (35).

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