Skeletal Involution by Age-associated Oxidative Stress and Its Acceleration by Loss of Sex Steroids*

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Both aging and loss of sex steroids have adverse effects on skeletal homeostasis, but whether and how they may influence each other's negative impact on bone remains unknown. We report herein that both female and male C57BL/6 mice progressively lost strength (as determined by load-to-failure measurements) and bone mineral density in the spine and femur between the ages of 4 and 31 months. These changes were temporally associated with decreased rate of remodeling as evidenced by decreased osteoblast and osteoclast numbers and decreased bone formation rate; as well as increased osteoblast and osteocyte apoptosis, increased reactive oxygen species levels, and decreased glutathione reductase activity and a corresponding increase in the phosphorylation of p53 and p66shc, two key components of a signaling cascade that are activated by reactive oxygen species and influences apoptosis and lifespan. Exactly the same changes in oxidative stress were acutely reproduced by gonadectomy in 5-month-old females or males and reversed by estrogens or androgens in vivo as well as in vitro. We conclude that the oxidative stress that underlies physiologic organismal aging in mice may be a pivotal pathogenic mechanism of the age-related bone loss and strength. Loss of estrogens or androgens accelerates the effects of aging on bone by decreasing defense against oxidative stress.

Age-related loss of bone mass and strength is an invariable feature of human biology, affecting women and men alike. Moreover, population-based studies demonstrate that substantial bone loss begins as early as the 20s in young adult women and men, long before any hormonal changes (1). The extent to which estrogen deficiency contributes to age-related bone loss and the slower rate of decline of bone mass and strength during the late postmenopausal years, and the molecular and cellular mechanisms of such putative interactions, are unknown.

The universality of age-associated bone loss irrespective of sex steroid status notwithstanding, age is by far a more critical determinant of fracture risk than bone mass in humans indicating that age-related increase in fracture risk reflects a loss of bone strength that is only partly accounted for by loss of bone mass (2). Whereas an increased propensity to fall due to age-related decline in neuromuscular function is a factor, there are also age-related changes in the bone itself. Such changes include disrupted architecture, altered composition of the bone mineral and matrix, delayed repair of fatigue microdamage, excessive turnover, and inadequate bone size (3–7). The most recently appreciated qualitative factor is loss of osteocytes (8, 9), former osteoblasts entombed into the mineralized matrix. Osteocyte death may influence the signals necessary for mechanical adaptation and repair and also lead to long term changes in bone hydration. The anti-apoptotic effect of sex steroids on osteocytes, which has been well documented in mice, rats, and humans (10–12), may contribute to their anti-fracture efficacy independently of their effect on bone mineral density (BMD) (8).

We and others had shown earlier that estrogens and androgens protect the adult skeleton against bone loss by suppressing the rate of bone turnover and maintaining a focal balance between bone formation and resorption (13–15). Suppression of bone turnover results from attenuating effects of sex steroids on the birth rate of osteoblast and osteoclast progenitors. Maintenance of a focal balance between formation and resorption results from opposite effects on the lifespan of osteoblasts/osteocytes and osteoclasts: an anti-apoptotic effect on the former and a pro-apoptotic effect on the latter cell type. Conversely, loss of sex steroids increases the rate of remodeling by up-regulating osteoblastogenesis and osteoclastogenesis. Specifically, we have shown that estrogen loss rapidly up-regulates osteo-

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4 The abbreviations used are: BMD, bone mineral density; AFU, arbitrary fluorescence units/mg of protein; BSO, l-buthionine-(S,R)-sulfoximine; DEM, diethyl maleate; DHT, dihydrotestosterone; E2, estradiol; ER, estrogen receptor; NAC, N-acetyl-L-cysteine; ORX, orchidectomy; OVX, ovariectomy; ROS, reactive oxygen species; TNF, tumor necrosis factor; N, newton; ERK, extracellular signal-regulated kinase; AR, androgen receptor; ANOVA, analysis of variance; MSC, mesenchymal stem cell; FOXO, Forkhead box O; MEK, mitogen-activated protein kinase/extracellular signal-regulated kinase kinase.
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**FIGURE 1.** BMD and strength decrease with age in sex steroid sufficient female or male C57BL/6 mice. A, BMD at the spine and femur was assessed by dual energy x-ray absorptiometry in two experiments with female and one experiment with male mice. The 

n was 12 animals per age group in each experiment with females, and 10–12 in the experiment with males. In the two experiments with females the age-dependent changes were statistically indistinguishable, hence the data were combined. B, load-to-failure, a measure of strength, was determined by compression testing of the 6th lumbar vertebra (L6) and by 3-point bending of the left femur. Colored horizontal lines in A and B indicate the mean values for each sex. * and † indicate the age at which a time-dependent decline began in females and males, respectively.

blastoegenesis in mice, and this effect is at least in part cell autonomous, and independent of the increased osteoclastic resorption, as it is manifested in ovariectomized (OVX) mice treated with bisphosphonates, known inhibitors of osteoclastic bone resorption (16). Moreover, the adverse effects of estrogen or androgen withdrawal on bone seem to be mediated, at least in part, by cells of the osteoblastic lineage and are obviated by constraints on osteoblastogenesis, whether genetic (17) or acquired (18), most likely because of the critical dependence of osteoblastogenesis on osteoblast aging (19, 20). Nonetheless, the effect of the loss of estrogens on the rate of remodeling and the up-regulation of osteoblastogenesis and bone formation in animals and humans wanes with time (5–10 years in women and 6–8 weeks in mice) to about the same rate as in elderly eugonadal males, raising the possibility that aging might be overriding the acute effects of estrogen deficiency.

Increased levels of reactive oxygen species (ROS) influence numerous cellular processes, including the timing of death by apoptosis; and have been linked to aging and the development of age-related diseases. Moreover, oxidative stress has been strongly correlated with longevity in flies, nematodes, and mammals (21, 22). The p53 tumor suppressor and the adapter protein p66shc represent key components of a signal transduction pathway that not only is activated by increased intracellular ROS and converts oxidative signals into apoptosis but also generates ROS in the mitochondria (23–25). Strikingly, an activating mutation of p53 causes early onset of aging-associated phenotypes in mice (26). Conversely, deletion of p66shc increases resistance to oxidative stress, as well as lifespan, by as much as 30% (23). One of the mechanisms used by cells to defend against oxidative damage involves the reduction of peroxides to harmless alcohols in a reaction in which glutathione peroxidase oxidizes glutathione (GSH) to the disulfide GSSH, and glutathione reductase (GSR) converts it back into GSH (27).

The above lines of evidence have strongly suggested to us the possibility that organismal aging per se, rather than an age-associated failure of other organs and tissues, may be the predominant mechanism of the bone fragility disease, which has become synonymous with osteoporosis, just one of the many features and risk factors underlying the problem of fractures. And, that loss of estrogens or androgens may exaggerate the adverse effects of organismal aging on bone. We report several previously unappreciated age-related changes in both female and male C57BL/6 mice that may provide critical clues into the mechanisms of the age-related decline of bone strength and mass, and the influence of sex steroid deficiency in the process.

**EXPERIMENTAL PROCEDURES**

**Chemicals and Reagents**—L-Buthionine-(S,R)-sulfoximine (BSO), diethyl maleate (DEM), N-acetyl-L-cysteine (NAC), etoposide, fluotamide, H$_2$O$_2$, dihydrotestosterone (DHT), and E$_2$ were purchased from Sigma. PD98059 was purchased from Cell Signaling Technology, Inc. (Danvers, MA). ICI 182,780 was purchased from Tocris Cookson Inc. (Ellisville, MO), PP1 from Biomol International P.A. (Plymouth Meeting, PA), and U0126 from Promega (Madison, WI). Tumor necrosis factor (TNF) α recombinant protein was purchased from R&D Systems (Minneapolis, MN). Sixty-day slow-release pellets containing DHT were purchased from Innovative Research of America (Sarasota, FL). Estradiol was assayed with a kit from DiaSorin (Stillwater, MN) and testosterone with a kit from MP Biomedicals (Costa Mesa, CA). Glutathione reductase was assayed with a kit from Cayman Chemical Co. (Ann Arbor, MI). Intracellular ROS were quantified using dichlorodihydrofluorescein diacetate dye (28), using bone marrow cells flushed from femurs and washed with phosphate-buffered saline.

**Animal Experimentation**—Male and female C57BL/6 mice 4–31 months old were purchased from Harlan Inc. from a cohort maintained with support from the National Institute of...
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Aging. The age-associated changes of intact animals were studied in three separate experiments: two with females (the first at 8, 16, 25, and 31 months old; and the second at 4, 8, 16, and 25 months old) and one with males (4, 8, 16, 25, and 31 months old). For the studies examining the effects of sex steroid deficiency, 5 month-old C57BL/6 mice were purchased from Harlan Sprague-Dawley Inc. Two to 3 days before surgery, all animals were electronically tagged (Biomedic Data System Inc., Maywood, NJ), BMD measurements were performed on each, and then sham-operated, OVX, or orchidectomized (ORX). Sham-operated animals were administered vehicle or BSO (2 mmol/kg intraperitoneally) twice a day (n = 12 per group). BSO was also included in the drinking water (20 mm). OVX and ORX animals were subcutaneously injected with vehicle or with replacement doses of E2 (30 ng/g) or NAC (100 mg/kg/day) twice a day or were implanted with 60-day slow-release pellets containing DHT (10 mg) (n = 12 per group). After 6 weeks of treatment, animals were sacrificed and the tissues dissected for further analyses. BMD, bone geometry measurements, histomorphometry, and osteoblast/osteocyte apoptosis were performed as previously described (29–31).

Biomechanical Testing—The load bearing properties of L6 were measured using a single column material testing machine and a calibrated tension/compression load cell (model 5542, Instron Corp., Canton, MA). Load cell calibration was verified in accordance with American Society for Testing and Materials E74-02 standards and traceable to the National Institute of Standards and Technology. Data were recorded and analyzed using the Merlin IX software package (Instron Corp.). The L6 specimens were cleaned of surrounding soft tissue, wrapped in gauze soaked in 37 ± 0.5 °C normal saline, and tested on the day of sacrifice. The length, width, and depth of the bones were recorded with a digital caliper at a resolution of 0.01 mm (Mitutoyo number 500-196, Ace Tools, Ft. Smith, AR). The cross-sectional area was assumed to be an ellipse and calculated as A = 0.25 π (width)(depth). Articular and spinous processes that would interfere with compression were excised using an iris scissors. After pre-seating with less than 0.5 newtons (N) of applied load, vertebrae were compressed between screw-driven loading platens using a lower-platen, customized miniature spherical seat that minimizes shear by adjusting to irregularities in the end plates of the specimens. Best seating was obtained with the load applied along the caudocephalad axis at a speed of 0.5 mm/min until failure. Standard materials for compression were run before each set of determinations. Three-point bending of the femur was also performed at 37 ± 0.5 °C using a miniature bending apparatus with the posterior femoral surface lying on lower supports (7 mm apart) and the left support immediately proximal to the distal condyles. Load was applied to the anterior femoral surface by an actuator mid-way between the two supports moving at a constant rate of 3 mm/min to produce a physiological in vivo stain rate of 1% for the average murine femur. The external measurements (length, width, and thickness) of the femora were recorded with a digital caliper. Measurements of the internal marrow cavity (greater and lesser diameters) were obtained with a hand-held micr scope at ×100 magnification using a calibrated linear reticule eyepiece (Klarmann Rulings, Manchester, NH). Maximum load (N) and displacement (mm) were recorded. The mechanical properties were normalized for bone size and ultimate strength or stress (N/mm²; in megapascals) was calculated. Standard precision steel piano wire with stiffness in the same range as murine femoral bone was evaluated before each set of determinations.

Western Blot Analysis—The phosphorylation status of p53, p66shc, and ERK1/2 was analyzed by immunoblotting in fifth lumbar vertebra lysates, as previously described (32). The antibodies used were: a rabbit polyclonal antibody recognizing Ser15-phosphorylated p53 (Cell Signaling Technology, Inc.,
Danvers, MA), a mouse monoclonal antibody recognizing Ser36-phosphorylated p66shc (Calbiochem, San Diego, CA), and a mouse monoclonal antibody recognizing tyrosine-phosphorylated ERK1/2 (Santa Cruz Biotechnology Inc., Santa Cruz, CA). Protein levels of p53, p66shc, and ERK1/2 were analyzed using a mouse monoclonal antibody recognizing p53 (Cell Signaling), a rabbit polyclonal antibody recognizing p66shc (BD Biosciences, Palo Alto, CA), and a rabbit polyclonal antibody recognizing total ERK1/2 (Santa Cruz).

Cell Culture—OB-6 cells, an osteoblastic cell line derived in our laboratory from the murine bone marrow (33), were cultured in α-minimal essential medium supplemented with 10% fetal bovine serum, antibiotics as above, and 1% sodium pyruvate. Osteoclasts were derived from bone marrow cells cultured in α-minimal essential medium supplemented with 30 ng/ml macrophage-colony stimulations factor and 30 ng/ml soluble RANK ligand. For the quantification of osteoblast apoptosis, OB-6 cells were treated for 1 h with BSO or DEM followed by the steroids for 1 and 6 h with the pro-apoptotic agent etoposide (5 × 10^{-5} M), TNFα (10^{-9} M), or H2O2 (5 × 10^{-5} M). Osteoclast apoptosis was assayed in cells treated with BSO or DEM for 1 h followed by the steroids for 24 h. Apoptotic cells were quantified by measuring caspase 3 activity as described previously (32). For quantification of osteoclasts, after 4 days in culture, cells were treated for 1 h with BSO or DEM followed by the steroids for 24 h. Osteoclasts were enumerated after staining for TRAPase; both mononucleated and multinucleated cells were counted.

Quantitative PCR—Total RNA was extracted from tibia or calvaria using Ultraspec RNA (Biotecx Laboratories, Houston, TX) and reverse-transcribed using the High Capacity cDNA Archive Kit (Applied Biosystems, Foster City, CA). Primers and probes for estrogen receptor (ER) α, ERβ, androgen receptor (AR), and glyceraldehyde-3-phosphate dehydrogenase were manufactured by the TaqMan® Gene Expression Assays service (Applied Biosystems).

Transient Transfections—C2C12 cells were transfected with 0.1 μg of green fluorescent protein, 0.1 μg of empty vector or wild-type p66shc plasmid, and 0.2 μg of pcDNA using Lipofectamine Plus (Invitrogen).

Statistical Analysis—Effects of age and gender on BMD and strength of the spine and the femur were evaluated by two-way ANOVA. The time of onset of the changes was determined by linear regression followed by a test for significant lack of fit or departure from linearity (34). Whenever there was significant departure from linearity for the entire data set, data from one or two earlier time points (4 month or both 4 and 8 months) were excluded from the analysis so that linearity could be established within the remaining set of data. Subsequently, the earliest time point that was used to establish linearity, i.e. 8 or 16 months, was compared with 4 or 4 and 8 months by
Student’s t test or ANOVA. If no different from the preceding time points, the earliest time point of the linearity plot was declared the time after which a particular change was manifested and the change was continuous from that time point on; if, on the other hand, differences were detected, we concluded that the change began at the earliest time point, but it was discontinuous.

ANOVA was used to detect other age-related changes, and effects of various in vivo and in vitro treatments, after establishing that the data were normally distributed and equivalency of variances. Bonferroni’s method was used to perform appropriate pairwise comparisons of treatment groups. In cases when one or both of the requirements for performing ANOVA were not met, Kruskal-Wallis ANOVA on Ranks test was used followed by Dunn’s method to perform pairwise comparisons of treatment groups.

RESULTS

Similar Age-associated Changes in BMD and Strength in Female and Male Mice—Both female and male C57BL/6 mice exhibited loss of bone mass, as determined by the surrogate measurement of BMD, and loss of strength in the spine and femur between ages of 4 and 31 months (Fig. 1, A and B). The fall in spinal BMD and femoral strength occurred earlier in females than in males.

Age-associated Changes in Osteoblast and Osteoclast Number, Bone Formation, and Osteoblast and Osteocyte Apoptosis—As evidenced from histomorphometric analysis (Fig. 2) of the vertebrae of the first of the two experiments with female C57BL/6 mice, advancing age from 8 to 31 months was associated with decreased cancellous bone area and trabecular width and increased trabecular separation (Fig. 2A). Furthermore, advancing age was associated with decreased wall width, decreased osteoblast and osteoclast number, as well as a decrease in bone formation rate. All these changes are unmistakable indications of a low turnover state, with resorption exceeding formation; and are clearly distinct from the high turnover state caused by loss of estrogens or androgens in rodents and humans. Furthermore, advancing age was associated with an increase in osteoblast and osteocyte apoptosis (Fig. 2B). The increase in osteoblast and osteocyte apoptosis with age in females was confirmed in the experiment with the males between the ages of 4 and 31 months. As is the case in humans (35, 36), both female and male mice exhibited an age-dependent increase in both the external and internal diameter of the femoral diaphysis, indicating ongoing periosteal apposition and endocortical resorption, respectively (Fig. 3).

Age-related Increase of Oxidative Stress in the Bone of Females and Males—Consistent with the evidence for a critical role of oxidative damage in age-related diseases in general, and of the p53/p66* signaling cascade in apoptosis, the changes in bone mass, strength, osteoblast/osteocyte apoptosis, osteoblast number, and bone formation rate with age in the aging murine model were associated with age-dependent decreased activity of GSR (Fig. 4A) and increased levels of ROS in the bone marrow (Fig. 4B). Furthermore, the pattern of all these changes was temporally associated with a progressive increase in the phos-
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FIGURE 5. Uterine or seminal vesicle weight or the estrogen and androgen receptor mRNA levels do not change with age in C57BL/6 mice. A, wet uterine or seminal vesicle weight; total body weight (B) and uterine or seminal vesicle (C) corrected for body weight of mice from the second experiment with females and the single experiment with males are shown. Bars represent mean ± S.D.; and n = 10–12 animals per group. * indicates p < 0.05 versus 4-month-old animals. D, mRNA levels of ERα, ERβ, or AR were determined by quantitative PCR in calvaria and tibia obtained from the second experiment with females (n = 5–9), as described under “Experimental Procedures.”

...phorylation of p53 (Fig. 4C) and p66<sup>shc</sup> (Fig. 4D) in lysates from vertebrae. In agreement with published studies by others showing no change in circulating estrogen or androgen levels with age in C57BL/6 mice (37, 38), we found no evidence of a decline in estrogen or androgen status between 4 and 31 months of age, as measured by the sensitive estrogen and androgen status indicators, uterine and seminal vesicle weight, respectively (Fig. 5, A–C). Likewise, transcript levels of the ERα or -β or the androgen receptor in bone were not altered with age (Fig. 5D). In addition, using commercially available radioimmunoassays, we attempted to measure serum levels of testosterone and 17β-estradiol. Detectable levels of testosterone were found in all males from which we have available serum and at all ages, and there was no difference among ages (4 months = 0.09 ng/ml ± 0.11, 8 months = 0.04 ng/ml ± 0.03, 16 months = 0.06 ng/ml ± 0.04, 25 months = 0.17 ng/ml ± 0.34, and 31 months = 0.10 ng/ml ± 0.06; n = 3–9 per group). On the other hand, serum estradiol was at or below the detection limit of the radioimmunoassay (2 pg/ml) in all the animals we assayed (n = 45) at all ages (4, 8, 16, 25, and 31; n = 9 per group).

Gonadectomy Acutely Recapitulates the Effect of Aging on Oxidative Stress, Osteoblast and Osteocyte Apoptosis, and BMD: Protection by an Antioxidant—Prompted by evidence that ROS, such as H<sub>2</sub>O<sub>2</sub>, may be responsible for the increased loss of bone caused by OVX in mice (39), we proceeded to test the hypothesis that sex steroid deficiency accelerates the effects of aging on bone, by examining whether sex steroid deficiency in younger mice will acutely reproduce the effects of advancing age on oxidative stress (Fig. 6). To this end, we sham-operated or gonadectomized 5-month-old C57BL/6 females or males. Immediately after surgery, the sham-operated animals were treated with either saline or BSO, a specific inhibitor of GSH synthesis. Gonadectomized animals received vehicle, E<sub>2</sub>, DHT, or the antioxidant NAC. Six weeks later, animals were sacrificed and uterine or seminal vesicle weight, ROS, and GSR activity in the bone marrow, as well as the phosphorylation of p53 and p66<sup>shc</sup> in vertebral lysates was determined. In either sex of animals, E<sub>2</sub>, DHT, or NAC prevented the effects of gonadectomy on all measures of oxidative stress tested here, including GSR activity, ROS levels, as well as p53 and p66<sup>shc</sup> phosphorylation (Fig. 6, A–D). Moreover, NAC was as effective as E<sub>2</sub> or DHT in preventing the decrease of spinal BMD caused by either OVX or ORX (Fig. 7A). Furthermore, NAC prevented the increase in osteoblast and osteocyte apoptosis induced by OVX (Fig. 7B); and, in agreement with Lean et al. (39), NAC prevented OVX-induced loss of cancellous bone (Fig. 7C). As expected, OVX decreased the uterine weight, and estrogen replacement reversed this effect (Fig. 7D). DHT also restored uterine weight as noted previously by us and others. Likewise, ORX decreased seminal vesicle weight, and DHT replacement prevented it. BSO or NAC had no effect on uterine weight; and BSO, E<sub>2</sub>, or NAC had no effect on seminal vesicle weight.

The Effects of Both Estrogens and Androgens on Osteoclasts and Osteoblasts Result from Antioxidant Actions—Finally, we investigated the possibility that the bone protective effects of estrogens or androgens result from direct antioxidant actions on bone cells. BSO or the electrophilic agent DEM abrogated the suppressive effects of E<sub>2</sub> or DHT on osteoclastogenesis (Fig. 8A). BSO or DEM also abrogated E<sub>2</sub>- or DHT-induced osteoclast apoptosis (Fig. 8B). In addition, E<sub>2</sub> or DHT stimulated the activity of GSR in osteoclasts (Fig. 8C); and the effect of E<sub>2</sub> or DHT on GSR was abrogated by the ER antagonist ICI 182,780 or the AR antagonist flutamide, respectively. Furthermore, the effects of E<sub>2</sub> or DHT on GSR were abrogated by the specific inhibitors of Src kinase PP1 and MEK kinase U0126. Consistent with the idea that low doses of H<sub>2</sub>O<sub>2</sub> promote osteoclast survival, H<sub>2</sub>O<sub>2</sub> at concentrations reported to cause NF-κB activation (1.5 to 12.5 μM), attenuated osteoclast apoptosis; however, at higher concentrations (25 to 50 μM) it was ineffective (Fig. 8D).

BSO or DEM also abrogated the anti-apoptotic effects of E<sub>2</sub> or DHT on bone marrow-derived osteoblastic cells, OB-6 (Fig.
9A), as well as in primary cultures of calvaria cells and the C2C12 osteoblast progenitor cell line (data not shown), irrespective of whether apoptosis was induced by the topoisomerase inhibitor etoposide, TNFα, or H$_2$O$_2$. BSO or DEM, at higher concentrations than those that blocked the anti-apoptotic effect of E$_2$, induced apoptosis of calvaria cells, presumably by depleting GSH (data not shown). This observation raises the possibility that a threshold level of GSH production by osteoblasts may be an inhibitor of endogenous ROS production and, thereby, apoptosis.

Phosphorylation of serine 36 (Ser36) of p66$^{shc}$ is required for transduction of oxidant stress signals leading to apoptosis. Consistent with this, overexpression of p66$^{shc}$ in C2C12 cells induced apoptosis both under basal conditions and in the presence of H$_2$O$_2$ (Fig. 9B). Therefore, we examined whether E$_2$ or DHT affects p66$^{shc}$ phosphorylation. H$_2$O$_2$ stimulated the phosphorylation of p66$^{shc}$ as early as 2 min and for at least 1 h following its addition to cultures of OB-6 osteoblastic cells but had no effect on the expression of the protein (Fig. 9C). E$_2$ or DHT suppressed the H$_2$O$_2$-induced p66$^{shc}$ phosphorylation as early as 15 min following treatment of OB-6 cells; and the effect of either steroid was reversed in the presence of the MEK inhibitor PD98059. These results suggest that, not only attenuation of osteoclastogenesis, but also the stimulation of osteoclast and attenuation of osteoblast apoptosis by estrogens involve non-protein thiol metabolism and are the result of actions mediated via cytoplasmic kinases and probably downstream transcriptional control (32).

**DISCUSSION**

The results reported herein demonstrate that both female and male C57BL/6 mice exhibit an age-related decrease in strength, BMD, bone remodeling, and bone formation rate together with an increase in osteoblast and osteocyte apoptosis. These changes are temporally associated with increased ROS levels, decreased GSR activity, and increased phosphorylation of p53 and p66$^{shc}$. The exact same changes in ROS, GSR, and the phosphorylation of p53 and p66$^{shc}$ were acutely reproduced by gonadectomy in 5-month-old females or males; and prevented by the antioxidant N-acetyl-L-cysteine. Moreover, estradiol or DHT acted directly in vitro to attenuate osteoclastogenesis and osteoblast apoptosis, stimulate osteoclast apoptosis, and suppress p66$^{shc}$ phosphorylation by a kinase-dependent mechanism that involves non-protein thiol metabolism. This evidence strongly suggests that increased ROS is a pivotal mechanism of age-related bone loss and strength; and that estrogen or androgen deficiency accelerates the adverse effects of aging on bone by decreasing defense against ROS. To the best of our knowledge, the in-depth characterization of the mice used in our studies, including the demonstration of increased ROS and p66$^{shc}$ phosphorylation with physiologic aging and acute loss of estrogens or androgens, demonstrates for the first time that aging and sex steroid deficiency exert their adverse effects by similar mechanisms in an animal model at the molecular, cellular, and tissue level. A model summarizing the results of the present report
and bone formation rate in aging C57BL/6 mice, in studies by Chambers and colleagues (41) both osteoblast numbers and bone formation were decreased in 2-month-old B6;129SF2 mice treated with the glutathione inhibitor BSO. Likewise, in agreement with our findings with the 5-month-old ovariectomized C57BL/6 mice, these workers had shown that the antioxidants, NAC and ascorbate, or inhibition of H₂O₂ by pegylated catalase prevents the increased osteoclastogenesis and loss of bone caused by acute loss of estrogens in 2-month-old MF-1 mice (39, 41).

Beneficial effects of estrogens in several other tissues, such as lens epithelial cells, arteries, central nervous system, fat, liver, and oviducts, are also shown to result from improved defense against oxidative stress (42–53). Importantly, in view of the fact that E₂ inhibits oxidized LDL (54) inhibition of 15-lipoxygenase, the enzyme responsible for the generation of oxidized low density lipoprotein, partially prevents OVX-induced bone loss (55). Moreover, bone loss and vascular calcification progress in parallel with advancing age, indicating an age-dependent relationship between atherosclerosis and osteoporosis (56). On the other hand, adverse effects of estrogens on breast cancer, the uterus, and spermatogenesis may be due to increased ROS production or decreased antioxidant defense (57–59).

The results of the present study in mice also suggest that loss of bone strength with age is only partially accounted for by a reduction in bone mass. Moreover, even though we did not study fractures in mice, our observations are in line with clinical evidence from humans that the age-related increase in fracture risk reflects a loss of bone strength that is only partly accounted for by loss of bone mass (2). Indeed, consistent with this contention, at the lowest values for bone mass for humans, a 20-year increase in age is accompanied by a 4-fold increase in fracture risk. Whereas an increased propensity to fall due to age-related decline in neuromuscular function is clearly a factor in humans, there are also age-related changes in the bone itself. The importance of non-mass factors is demonstrated by several lines of evidence. First, a fracture at any site increases the risk of a subsequent fracture at any other site (60). Second, only a small part of the reduction in fracture incidence in response to anti-catabolic therapy can be accounted for by the increase in bone mass (61). Third, many of the genetic effects on bone strength are mediated by factors other than bone mass (62, 63). The results reported here support the idea that loss of osteocytes, which may be an additional factor (8, 64). Importantly, the age-dependent decline in murine bone strength in the present study occurred despite the expansion in bone size, exactly as is the case in humans. Therefore, our findings also support the contention that the C57BL/6 mouse is a good model for the study of age-related geometric changes in human bone.

In line with the results of the present report suggesting that sex steroid status was not altered appreciably, at least as far as the skeletal changes concerned, between the age of 4 and 31 months in C57BL/6 mice, Mobbs and Finch (37) had found no difference in estradiol levels in females up to the age of 18 months or androgen in male C57BL/6 mice up to 31 months (38). Nonetheless, in more detailed studies of the estrus cycle in the same mice these workers subsequently reported that estro-

and findings from earlier studies by us and others on the effects of ROS and sex steroids on bone cells is provided in Fig. 10.

In agreement with the findings of the present report, osteoporosis has been noted by us and others in mouse models of premature aging (17, 26, 40). Moreover, consistent with our observations temporally linking increased oxidative stress with increased osteoblast apoptosis and decreased osteoblast number
gen levels on day 3 and the preovulatory rise beginning on day 4 were decreased at 10–12.5 months as compared with 5.5–7.5 months of age (65). Moreover, whereas Nelson et al. (38) did not show declines in testosterone levels in healthy mice, mice with other diseases of aging did have markedly reduced testosterone levels. In addition, in a very recent paper (66) the authors found marked decreases in testosterone levels in male C57BL/6 mice at age 15 months compared with 4 months. We and others have shown here and elsewhere that seminal vesicle weight has less than half the variance of the testosterone measurements and indicates cumulative androgen status in mice better than a single serum sample (18, 67, 68). Be that as it may, we cannot categorically exclude the possibility that the age-related findings we have demonstrated herein may be accentuated by superimposed sex steroid deficiency in aging mice, as they are in humans. That having been said, however, acute loss of sex steroids causes an increase in the rate of bone remodeling, underlined by an increase in osteoclastogenesis and osteoblastogenesis and a corresponding increase in both bone resorption and formation, with the former exceeding the latter. In contrast, our aging C57BL/6 mice (and elderly individuals without vitamin D deficiency and secondary hyperthyroidism) exhibit a low rate of bone remodeling, as well as a decrease in osteoblast number and bone formation, a phenotype reminiscent also of the osteoporosis associated with glucocorticoid excess. In fact, work, from our group and others in several animal models, has strongly suggested that decreased wall width, the hallmark of age-associated osteoporosis and decreased bone formation results from decreased osteoblastogenesis (13, 17). If both aging and loss of sex steroids exert their adverse effects on bone by oxidative damage, how can osteoblast and osteoclast number be low in the former and high in the latter?

In studies reported in the accompanying manuscript by Almeida et al. (82) we have determined that ROS antagonize the skeletal effects of Wnt/β-catenin in vitro by diverting β-catenin from Tcf- to FoxO-mediated transcription. Moreover, consistent with the notion that increased ROS production with age attenuates Wnt/β-catenin signaling, Axin2 and Opg mRNA were decreased in old as compared with young C57BL/6 mice. Because activation of Wnt signaling enables osteoblastogenesis, suppresses osteoblastic cell apoptosis, inhibits osteoclastogenesis and increases bone mass, the cellular changes seen with increased aging and the accompanying oxidative damage may be due in part to attenuation of Wnt signaling.

In work reported elsewhere (69, 70), we have found that the estrogen-liganded ERα inhibits BMP-2-induced osteoblast progenitor differentiation, whereas the unliganded ERα or ERβ, but not the AR, promote the pro-differentiating actions of BMP-2 by enhancing Smad-mediated transcription. In followed by E2 or DHT (10⁻⁷ M) for 24 h. Osteoclasts were enumerated after staining for TRAPase and apoptosis was quantified by determining caspase 3 activity. C, GSR activity in osteoclasts treated with ICI 182,780 (10⁻⁷ M), flutamide (10⁻⁷ M), PP1 (10⁻⁶ M), or U0126 (10⁻⁶ M) for 1 h followed by the indicated steroids for 24 h. D, apoptosis was quantified by determining caspase 3 activity in bone marrow-derived osteoclasts treated for 24 h with 10⁻⁷ M E2 or the indicated doses of H₂O₂. Bars indicate mean ± S.D. of triplicate determinations; *p < 0.05 versus vehicle.
addition, we have developed methods of expanding the self-renewal of mesenchymal stem cell (MSC) progenitors of osteoblasts, while preserving stemness, and found that their replication capacity is significantly lower in old versus young mice (71).

Based on our own findings and parallel developments in the understanding of the role of FoxOs in hematopoietic stem cell resistance to physiologic oxidative stress (72) we hypothesize that estrogens influence osteoblastogenesis by two distinct but overlapping cell autonomous mechanisms: (a) antioxidant effects on MSC osteoblast progenitors and (b) transcriptional repression of BMP-induced differentiation of their progeny. Increased levels of ROS have deleterious effects on MSC self-renewal (by increasing apoptosis), but promote MSCs entry into the cell cycle and terminal differentiation. These effects are counteracted by ROS-enhanced binding of FOXO to β-catenin, leading to increased defense against ROS. Because of their ability to defend against ROS, estrogens favor MSC quiescence and survival and oppose terminal differentiation. Following acute loss of estrogens, the increased ROS levels in MSCs promote their exit from quiescence, replication, and development of transit amplifying osteoblast progenitors. These mechanisms, together with the unleashing of the transcriptional restraint of the liganded ERα on BMP, lead to up-regulation of osteoblast differentiation and thereby increased bone formation. However, this effect is reversed with time because the gradual buildup of ROS production with aging (accelerated by estrogen deficiency) antagonizes Wnt signaling and also leads to a decrease in the size of the MSC compartment by increasing MSC apoptosis.

In addition, we have found that as compared with 4-month-old mice, 25- and 31-month-old mice exhibited a 2-fold increase in serum corticosterone, adrenal weight, and bone mRNA for 11β-Hsd1, the enzyme that amplifies glucocorticoid action.5 The age-dependent decrease in bone formation rate and increased osteoblast and osteocyte apoptosis, cardinal features of the adverse effects of glucocorticoid excess on bone, along with the increase in 11β-Hsd1 mRNA, suggest that local amplification of endogenous glucocorticoids may also contribute to decreased bone formation, the decline in bone strength, and the disparity between bone quantity and quality with aging. Additional mechanisms could well include increased production of peroxisome proliferator-activated receptor γ-activating oxidized lipids derived from the lipoxygenase Alox15 (55, 73) and the skeletal unloading

5 R. S. Weinstein, R. L. Jilka, and S. C. Manolagas unpublished data.

FIGURE 9. Estrogens or androgens regulate the survival of osteoblasts via antioxidant actions. A, apoptosis was quantified by determining caspase 3 activity in OB-6 cells treated for 1 h with BSO or DEM followed by the steroids for 1 and 6 h with the pro-apoptotic agent etoposide (5 × 10⁻⁵ M), TNFα (10⁻⁹ M), or H₂O₂ (5 × 10⁻⁵ M). Bars indicate mean ± S.D. of triplicate determinations; * indicates p < 0.05 versus vehicle. B, C2C12 cells transfected with a vector control or wild-type p66shc plasmid, along with green fluorescent protein, were treated with or without H₂O₂ (5 × 10⁻⁵ M). The number of apoptotic cells was determined by examining the nuclear morphology of fluorescent cells 6 h later. Bars indicate mean ± S.D. of triplicate determinations; * p < 0.05 versus vector control untreated. C, OB-6 cells were treated for 1 h with the MEK inhibitor PD98059 (5 × 10⁻⁵ M), then the indicated steroids (10⁻⁸ M) were added, and 1 h later the cultures were exposed for 15 min to H₂O₂ (5 × 10⁻⁵ M). Phosphorylated p66shc was determined by Western blot analyses in cell lysates.
resulting from reduced physical activity with old age (74). Hence, the effects of aging must override the effects of sex steroid deficiency by several mechanisms, some of which result directly from increased oxidative stress. In agreement with this contention we have shown before that the expected changes in bone cell progenitor numbers, histomorphometry, and BMD that occur after orchidectomy were either absent or greatly attenuated in a mouse model of defective osteoblastogenesis or in Swiss Webster mice treated simultaneously with glucocorticoids (17, 18).

Although much has been learned about age-related bone loss, it remains unknown whether its mechanisms are the same in all aging persons, but recent evidence indicates that the cellular and molecular mechanisms responsible may vary from subject to subject (75–77). One reason for such variation could be that the increase in responsiveness differs among individuals, so that some would lose more bone than others at, for example, the same glucocorticoid level. In closing, we submit that the results presented in this report constitute a major paradigm shift in the understanding of the pathogenesis of osteoporosis from the “estrogen-centric” view that has dominated the last 50 years to a multifactorial one, in which oxidative damage is a pivotal mechanism for females and males alike.
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