Twelve months of voluntary heavy alcohol consumption in male rhesus macaques suppresses intracortical bone remodeling


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Twelve months of voluntary heavy alcohol consumption in male rhesus macaques suppresses intracortical bone remodeling

Gino W. Gaddini, Kathleen A. Grant, Andrew Woodall, Cara Stull, Gianni F. Maddalozzo, Bo Zhang, Russell T. Turner, Urszula T. Iwaniec

Abstract

Chronic heavy alcohol consumption is a risk factor for cortical bone fractures in males. The increase in fracture risk may be due, in part, to reduced bone quality. Intracortical (osteonal) bone remodeling is the principle mechanism for maintaining cortical bone quality. However, it is not clear how alcohol abuse impacts intracortical bone remodeling. This study investigated the effects of long-duration heavy alcohol consumption on intracortical bone remodeling in a non-human primate model. Following a 4-month induction period, male rhesus macaques (Macaca mulatta, n = 21) were allowed to voluntarily self-administer water or alcohol (4% ethanol w/v) for 22 h/d, 7 d/wk for 12 months. Control monkeys (n = 13) received water and an isocaloric maltose-dextrin solution. Tetracycline hydrochloride was administered orally 17 and 3 days prior to sacrifice. Control monkeys and monkeys consuming alcohol demonstrated similar body weights and serum levels of ethanol (1.7 ± 0.1 ng/ml, versus 1.8 ± 0.5 ng/ml, p = 0.021) and labeled osteon density was lower (0.41 ± 0.2/mm², p = 0.003) in alcohol-consuming monkeys compared to controls, indicating a reduced rate of intracortical bone remodeling. In concordance, plasma CTx was lower (2.5 ± 0.3 ng/ml versus 1.7 ± 0.1 ng/ml, p = 0.028) in the alcohol group. These results suggest that chronic heavy alcohol consumption may negatively impact bone health, in part, by suppressing intracortical bone remodeling.

Introduction

Chronic heavy alcohol consumption is associated with an increase in all cause fracture risk [1–5]. A low bone mineral density (BMD) is often observed in chronic heavy alcohol consumers and when detected is generally associated with decreased bone formation [6–14]. In contrast, the precise role of bone resorption in the etiology of this condition is less clear [6,7,11,12,15–18]. Because alcohol consumption often results in an overall reduction in the rate of bone remodeling, bone loss may occur when the reduction in bone formation exceeds the reduction in bone resorption [19–21]. Although alcoholics often exhibit reduced BMD, the specific effects of alcohol on the human skeleton are poorly defined. This is due, in part, to complications associated with co-morbidities, such as smoking, poor diet and alcohol-induced disease [22]. In addition, alcohol intervention studies are exceedingly difficult to perform in humans. As a result, most intervention studies have been conducted using experimental animal models, with rodents being the mainstay for assessing the specific effects of alcohol on bone metabolism [23–25]. To date, rodent studies have focused on the effects of alcohol on cortical bone accrual and cancellous bone turnover. However, because intracortical (osteonal) bone remodeling is absent in small rodents, these animals are not ideal for evaluation of the effects of alcohol on cortical bone turnover [26].

Disclosure summary: The authors have nothing to disclose.

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In humans, cortical bone comprises the great majority (≥80%) of bone mass and plays a fundamental role in the mechanical function of the skeleton. Cortical bone fractures are common in young adult and middle aged males and are more prevalent in chronic heavy alcohol consumers [26,27]. Interestingly, heavy alcohol consumers are more likely to sustain low trauma fractures than non-heavy consumers [27]. Because the ability of a bone to resist fracture depends not only on bone mass, but also on factors such as architecture and intrinsic properties of the bone material [28], a deficit in cortical bone quality caused by detrimental changes in architecture and material properties can increase fracture risk independent of reduced BMD [29–31]. In this regard, intracortical remodeling is essential for maintenance of cortical bone quality [32].

Intracortical bone remodeling is initiated by osteoclast-mediated resorption on a quiescent bone surface to form a cutting cone, which is followed spatially and temporally by a closing cone lined by osteoblasts. In a completed osteon, bone fills most of the cavity that was created by the action of osteoclasts, leaving only a Haversian canal [33]. An increase in the overall rate of bone remodeling, or a remodeling imbalance where bone formation fails to keep pace with bone resorption, can result in increased cortical bone porosity and decreased BMD [34]. The effect of heavy alcohol consumption on intracortical bone remodeling rate and balance is unknown. The purpose of this study was to evaluate the specific impact of alcohol on intracortical bone remodeling and cortical porosity in a non-human primate model for chronic heavy alcohol consumption. The study focused on late adolescent to young adult male monkeys to model the human population that is most prone to chronic heavy alcohol consumption [35].

Materials and methods

Animals

The study population was comprised of a total of 34 late adolescent/young adult (5.4 ± 0.1 years old at initiation of alcohol protocol) male rhesus macaques (Macaca mulatta). Epiphyseal closure in rhesus macaque males is complete by 6.5 years of age [36]. Animals were pooled from 3 cohorts of monkeys selected from a colony born and reared in captivity at the Oregon National Primate Research Center at Oregon Health and Sciences University. The first cohort was sent to necropsy in May 2010 (n = 5 control, 8 alcohol), the second in March 2012 (n = 4 control, 8 alcohol), and the third in July 2012 (n = 4 control, 5 alcohol). All of the animals from each cohort were subjected to the same experimental design (described below). Throughout the study, monkeys were housed individually under constant temperature (20–22 °C) and humidity (65%) and an 11-h light cycle (light 0700–1800 h) in a room allowing visual, auditory, and olfactory contact with other monkeys. Body weights were recorded weekly throughout the study. Alcohol intake and blood alcohol concentration data for some of these subjects have been published elsewhere [37,38].

Experimental design

Induction phase

The experimental design is described in detail elsewhere [39]. Briefly, monkeys were trained to self-administer food and beverage (using an operant panel integrated into the side of their cage) and to submit their leg for blood sampling. Monkeys in the treatment group were then induced to drink increasing volumes of an alcohol solution (4% w/v ethanol mixed in deionized water) in a step-wise fashion over 4 consecutive 30-day periods for a total of 120 days. To induce drinking, a 1 g flavored food pellet was delivered every 5 min and water was the only fluid available. After water consumption became associated with the delivery of the pellet, the monkeys underwent a 30-day session where water was the only drinking fluid provided. During the second 30-day interval, animals drank a predetermined volume of alcohol solution corresponding to 0.5 g/kg/d alcohol, followed by volumes of alcohol corresponding to 1.0 and 1.5 g/kg/d during the third and fourth 30-day intervals, respectively. Monkeys were allowed to drink only 4% w/v alcohol until the required dose of alcohol was reached (e.g., 0.5 g/kg/d), at which point animals were allowed to drink only water. This step-wise increase in alcohol induction was done to circumvent alcohol taste aversion and increase the opportunity to associate drinking alcohol with its intoxicating effects.

Voluntary drinking phase

Following the 120-day induction, monkeys in the alcohol group (n = 21) were given simultaneous access to both water and alcohol (4% w/v) and allowed to voluntarily self-administer alcohol and/or water for 22 h/d (1100–0900 h each day), 7 d/wk for 12 months. Control animals (n = 13) were allowed to self-administer a volume of maltose-dextrin solution isocaloric to the mean volume of alcohol consumed by the alcohol group. Consumption was recorded daily by using weighing scales (Ohaus Corp., Parsippany, NJ) to measure the change in the mass of containers dispensing the solutions.

Blood samples were collected every 5 days from the saphenous vein of monkeys in the alcohol group just before the lights were turned off (between 1800 and 1900 h), which corresponded to 7 h into the 22 h sessions. Blood samples were sealed in airtight vials containing 0.5 ml of distilled water and 0.02 ml of 10% isopropanol (internal standard), and stored at −4 °C until analysis.

Food consisted of 1 g banana-flavored pellets (carbohydrate, 63%; fat, 4%; protein, 22%; PJ Noyes, Lancaster, NH). Over the 12-month duration of the experiment, the monkeys were required to eat their daily allotment of food in no fewer than 3 “meals,” with at least 2 hours between each meal. A meal was defined by the proportion of daily food allotted to each monkey and the pace of the animal to obtain the food. The meal ended if one-third of the daily food allotment was obtained at a time, or if the monkey took longer than 2 minutes to obtain a pellet.

Fluorochrome tetracycline hydrochloride (20 mg/kg) was administered orally (17 and 3 days) prior to sacrifice for determination of active mineralization sites and rates of bone formation. At necropsy, the right tibia with attached fibula was harvested from each animal, placed into 70% alcohol, and stored at 4 °C until analysis.

Blood alcohol concentrations

Blood alcohol concentrations were assayed in plasma using a gas chromatograph (Hewlett-Packard 5890 Series II, Avondale, PA) equipped with a headspace autosampler, flame ionization detector, and a Hewlett Packard 3392A integrator.

Blood marker of bone resorption

Carboxyterminal cross-linking telopeptide of type 1 collagen (CTX), a marker of bone resorption, was measured in samples collected at necropsy using Serum CrossLaps ELISA (Immunodiagnostic Systems, Fountain Hills, AZ).

Blood vitamin D

25-Hydroxyvitamin D was measured in plasma samples collected at necropsy using an electrochemiluminescence binding assay (Roche Diagnostics, Indianapolis, IN).

Dual-energy X-ray absorptiometry

Bone mineral content (BMC, g), bone area (cm²), and areal BMD (g/cm²) in tibia/fibula (the tibia and fibula of each animal were analyzed together) were determined post mortem using a dual-energy X-ray absorptiometry.
absorptiometry (DXA) scanner (Hologic Discovery A, Waltham, MA) and Hologic APEX System Software, Version 3.1.1. In addition to analysis of total tibia/fibula, a subregion in the distal tibia diaphysis (~2.5 mm in length, enclosing the region evaluated by µCT and histomorphometry; please see below) was evaluated. Quality control check was performed against the Anthropomorphic Spine Phantom and Small Animal Step Phantom provided by the manufacturer. The coefficient of variation evaluating test-retest reliability for DXA scans in our laboratory is 1.0% for BMC, area, and areal BMD. The least significant difference is 0.003–0.006 g/cm², depending on skeletal site, at the 95% confidence level.

Microcomputed tomography

Microcomputed tomography (µCT) was used for nondestructive 3-dimensional evaluation of cortical bone architecture and porosity in tibial diaphysis. Tibial length was measured as the distance between the proximal tip of the intercondylar eminence and the distal tip of the medial malleolus. The distal third of the tibia was then excised using an Isomet® Low Speed Saw (Buehler, Lake Bluff, IL) and scanned in 70% ethanol at a voxel size of 30 x 30 x 30 µm (55 kVp, 145 µA, and 200 ms, 500 projections/rotation) on a Scanco µCT40 scanner (Scanco Medical AG, Basserdorf, Switzerland). Evaluations were conducted with filtering parameters sigma and support set to 0.8 and 1, respectively. Thirty-three consecutive slices (1.0 mm) of cortical bone (at proximal end of the distal third of the tibia) were analyzed at a threshold of 245 (gray scale of 0–1000) determined empirically. This threshold corresponds to 374 mg hydroxyapatite/cm². Cortical measurements included (1) cross-sectional tissue volume (cortical and marrow volume, mm³), (2) cortical volume (mm³), (3) marrow volume (mm³), (4) cortical thickness (µm), and (5) polar moment of inertia (an estimate of bone strength in torsion, mm⁴).

Subsequently, a 1-mm thick cross-section of bone (for further µCT evaluation) and a 50-µm thick cross-section of bone (for histomorphometric evaluation) were removed from the proximal end of the scanned portion of the tibia using the Isomet® Low Speed Saw. The 1-mm thick section was cut into 4 quadrants (sectioned at 45° angles to the cranial-caudal and medial-lateral axes to generate cranial, lateral, caudal, and medial quadrants) and each quadrant was scanned at a voxel size of 6 x 6 x 6 µm to provide adequate resolution for assessment of intracortical porosity. Cutting the cross-section into quadrants was essential due to constraints on specimen size for scanning at 6 µm voxel size. Sixty-eight consecutive slices (408 µm) were evaluated in each quadrant. Inverse thresholding was used to create an image of the canal size and distribution throughout the bone specimen. Direct measurements included cortical porosity (cortical volume/bone volume, %), canal thickness (µm), canal number (mm⁻¹), and canal spacing (µm). Results are reported for each quadrant and as a mean for the 4 quadrants.

Quantitative bone histomorphometry

The 50-µm thick cross-sections were ground on a roughened glass surface (using 220-grit aluminum oxide powder) to an approximate thickness of 25 µm for histomorphometric evaluation. Fluorochrome-based measurements of intracortical bone remodeling included (1) labeled osteone density (number of single- and double-labeled osteons/bone area, number/mm²), (2) mineral apposition rate (the distance between two fluorochrome markers that comprise a double label in an osteon divided by the 14-d interval between consecutive labels, µm/d), (3) bone formation rate (osteonal double + half single labeled perimeter multiplied by mineral apposition rate and expressed per bone area, %/year), and (4) formation period (osteonal wall thickness/mineral apposition rate, d). Activation frequency is not reported because it approached 0 in the alcohol group. In addition, the number and size (area bounded by a cement line) of incomplete osteons (i.e., actively remodeling osteons where the formation phase of the remodeling cycle has not been completed) was determined. The osteon count was normalized to bone area (number/mm²).

Fluorochrome-based measurements of endocortical and periosteal bone formation included (1) mineralizing perimeter (mineralizing perimeter/bone perimeter: bone perimeter covered with double plus half single label normalized to bone perimeter, %), (2) mineral apposition rate (the distance between two fluorochrome markers that comprise a double label divided by the 14-d interlabel interval, µm/day), and (3) bone formation rate (bone formation rate/bone perimeter: calculated by multiplying mineralizing perimeter by mineral apposition rate normalized to bone perimeter, µm²/µm²/µm/day).

Table 1

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<th>Study population.</th>
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<td>Age (years)</td>
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<td>67 ± 0.2</td>
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<td>Body weight (kg)</td>
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<td>87 ± 0.2</td>
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<td>Blood alcohol concentration (mg/dL)</td>
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Data are mean ± SE.
Fig. 1. Variation in daily alcohol consumption (A) and corresponding blood alcohol concentrations taken every fifth day at seven hours following onset of session (B) over the 12-month duration of study in four representative male rhesus macaques.

Fig. 2. Effects of 12 months of chronic alcohol consumption on tibial length (A), total tibia/fibula bone area (B), bone mineral content (C), and bone mineral density (D) and on bone area (E), bone mineral content (F), and bone mineral density (G) in a subregion of the distal tibia enclosing the region evaluated by μCT and histomorphometry in male rhesus macaques. Data are mean ± SE.
Fig. 3. Effects of 12 months of chronic alcohol consumption on tibial diaphysis cross-sectional volume (A), cortical volume (B), marrow volume (C), cortical thickness (D), and polar moment of inertia (E) in male rhesus macaques. Data are mean ± SE.

Fig. 4. Effects of 12 months of chronic alcohol consumption on cortical porosity in the tibial diaphysis in male rhesus macaques. Tibial cross-sections were divided into four quadrants and porosity assessed in the cranial (A), lateral (B), caudal (C), and medial quadrants (D) and as a mean for the 4 quadrants (E). Representative μCT images (6 x 6 μm voxel size) of cranial quadrants depicting cortical porosity are shown for a control monkey and an alcohol-consuming monkey (F). Data are mean ± SE. *Different from Control, p < 0.05; †p < 0.1.
ally, labeled osteon density was 98% lower (0.41 ± 0.2/mm² vs. 0.01 ± 0.01/mm², p = 0.002) in the alcohol group compared to the control group.

In the tibial diaphysis are shown in Fig. 5. The density of incomplete osteons was 91% lower (0.47 ± 0.2/mm² vs. 0.04 ± 0.01/mm², p = 0.001, D) in the alcohol-consuming monkeys compared to control monkeys. The lower cortical porosity in the cranial quadrant was associated with lower canal thickness and number and higher canal spacing (Table 2). Canal thickness was also lower in the alcohol-consuming monkeys compared to control monkeys in the lateral and caudal quadrants and in total cross-section (four quadrants combined). XCT images illustrating differences in cortical porosity in a representative cranial quadrant from a control and an alcohol-consuming monkey are shown in Fig. 4F.

The effects of alcohol consumption on intracortical bone remodeling in the tibial diaphysis are shown in Fig. 5. The density of incomplete osteons was 91% lower (0.47 ± 0.2/mm² versus 0.04 ± 0.01/mm², p < 0.001) in the alcohol group than in the control group (A). Additionally, labeled osteon density was 98% lower (0.41 ± 0.2/mm² versus 0.01 ± 0.01/mm², p < 0.003) in alcohol-consuming monkeys relative to controls (B). Differences in mineral apposition rate were not detected with treatment (C). Bone formation rate was 98% lower (5.6 ± 2.8%/year versus 0.10 ± 0.00%/year, p = 0.019) in the alcohol group than the control group (D). Significant differences between control and alcohol-consuming monkeys were not detected for osteon size (E) or formation period (F). Photomicrographs of histological sections illustrating differences in intracortical remodeling in a control and alcohol monkey are shown in Figs. 5 (G-H).

The effects of alcohol on bone formation in periosteal and endocortical envelopes in the tibial diaphysis are shown in Fig. 6. Periosteal mineralizing perimeter tended to be lower (p = 0.074) in alcohol-consuming in comparison to control monkeys (A). Differences in periosteal mineral apposition rate (B) and bone formation rate (C) were not detected with treatment. Alcohol resulted in 91% lower endocortical mineralizing perimeter (19.6 ± 7.0%/month versus 1.7 ± 1.4%/month, p < 0.001, D), 47% lower mineral apposition rate (0.86 ± 0.07/mm²/month versus 0.46 ± 0.06/mm²/month, p = 0.004; E), and 95% lower bone formation rate (6.3 ± 2.4/mm²/month versus 0.3 ± 0.3/mm²/month; p = 0.001, F) in the alcohol group compared to the control group.

The effects of alcohol consumption on plasma CTx and 25-hydroxyvitamin D are shown in Fig. 7. Alcohol-consuming monkeys had 31% lower plasma CTx levels compared to control monkeys (2.5 ± 0.3 ng/ml versus 1.7 ± 0.1 ng/ml, p = 0.028) (A). Significant differences in 25-hydroxyvitamin D were not detected with treatment (B).

**Discussion**

We investigated the effects of 12 months of daily voluntary alcohol consumption on bone density, cortical bone architecture, and intracortical bone remodeling in tibiae of young adult male rhesus macaques. To our knowledge, this is the first study to examine the effects of chronic alcohol consumption on indices of intracortical bone remodeling using an animal model. An average daily alcohol intake of 2.7 ± 0.2 g/kg, resulting in average daily blood alcohol concentration of 77 ± 9 mg/dl, had no effect on tibial BMC and areal BMD nor on cortical bone mass or architecture in tibial diaphysis. In contrast, alcohol consumption resulted in reduced density of fluorochrome-labeled osteons and incomplete osteons, reduced intracortical porosity, and reduced endocortical bone formation. Finally, alcohol consumption resulted in reduced plasma level of the bone resorption marker CTx.

Macaques have greater than 95% gene homology with humans and share many similarities in physiology [35], including bone physiology. Furthermore, attributes of alcohol consumption in macaques closely resemble attributes of alcohol consumption in humans [35]. This includes similar absorption and metabolism of alcohol and the propensity to voluntarily consume large quantities of alcohol [35]. Based on the aforementioned features, macaques should be an excellent model to evaluate the effects of alcohol on the skeleton.

Consistent with prior studies in cynomolgus macaques [39], rhesus macaques in the current study frequently engaged in episodes of binge drinking. Although average daily blood alcohol concentrations were only moderately high, the levels were highly variable, with day-to-day levels ranging from 0 to over 250 mg/dl (a blood alcohol concentration which can result in severe motor impairment and unconsciousness). The patterns of alcohol consumption and the daily blood alcohol concentrations exhibited by the monkeys in this study are similar to drinking patterns and blood alcohol concentrations of alcoholic men given 24 h/day free access to alcohol for 10–60 consecutive days in a hospital ward [40–42].

Bone fracture risk is influenced by a number of parameters including bone mass, architecture, and quality. The findings of the present cross-sectional analysis demonstrate that cortical bone mass and architecture in young adult male rhesus macaques were not significantly affected by 15 months (3 month induction phase plus 12 month voluntary drinking phase) of chronic heavy alcohol consumption. Many studies in human alcoholics have associated chronic heavy alcohol consumption with low BMD [3,4,43–50]. In the majority, subjects had alcoholism-related comorbidities, such as liver or kidney disease, pancreatitis, smoking, impaired ability to carry out activities of daily living, or poor nutrition, which could have alcohol-independent detrimental skeletal effects [3, 47,51–53]. Additionally, not all studies, particularly ones that excluded individuals with alcohol-related secondary diseases, have demonstrated an association between chronic heavy alcohol intake and low BMD. Laitinen et al. (1993), for example, reported that an average consumption of 186 ± 85 g/d (~13 standard drinks/d) in women with low BMD is a predictor of fracture risk in the general population, the increase in low trauma, as well as all cause fractures in heavy alcohol consumers exceeds values predicted by the extent of bone loss [46, 54]. These findings suggest that chronic heavy alcohol consumption...
may reduce bone strength by a mechanism that does not necessarily lead to a reduction in BMD.

Intracortical bone formation is normally coupled to bone resorption, suggesting that the reduced intracortical bone formation in alcohol-consuming monkeys may be caused by near-complete suppression of initiation of new secondary osteons. The observed reductions in intracortical porosity, incomplete osteon density, and blood level of CTx, with normal osteonal mineral apposition rate in the alcohol-consuming monkeys support this interpretation as does μCT analysis indicating that the reduction in intracortical porosity was due to reduced canal size; alcohol consumption decreased (by 52%) overall canal size with only a tendency to lower (by 3%) canal number. Alternatively, resorption may have been initiated and terminated prematurely.

However, incomplete osteons in the alcohol-consuming monkeys were not smaller nor was the formation interval reduced when compared to controls. Thus, the tissue level reduction in intracortical bone formation observed in alcohol-consuming monkeys was largely due to decreased activation of osteonal bone remodeling.

As evidenced by the decrease in cortical bone porosity, osteons that were in the process of formation prior to dietary intervention were completed by the end of study. This is not surprising because the formation interval (~2 months) of the bone remodeling cycle was much shorter than the treatment interval. Several classes of pharmaceuticals (e.g., bisphosphonates, estrogens, and selective estrogen receptor modulators) preserve bone mass and quality during high turnover states, such as following menopause, by reducing initiation of bone formation.
remodeling [55]. Consumption of moderate quantities of alcohol may confer a similar benefit in postmenopausal women [56]. However, highly suppressed intracortical bone remodeling, of the magnitude observed in the present study, is associated with failure to repair microdamage resulting in reduced bone quality [57,58].

In the present study, the dramatic decrease in cortical bone turnover in the femur was associated with site-specific reductions in porosity. The decrease in porosity was not associated with a change in areal BMD. Thus, prolonged suppression of intracortical bone remodeling is a plausible mechanism by which chronic heavy alcohol consumption could lead to increased fracture risk independent of BMD. If our presumption is correct, conventional screening with DXA may not detect an alcohol-induced loss of bone strength.

Chronic heavy alcohol consumption in humans is consistently associated with reduced biochemical markers of global bone formation [9, 17, 43, 45–47, 59, 60]. However, these markers provide little insight as to region- or compartment-specific skeletal effects. Histomorphometric analysis of biopsies allows for accurate assessment of local bone formation. Analyses of human transiliac biopsies in chronic heavy alcohol consumers show reduced cancellous bone formation [6, 16, 48, 61, 62]. Unfortunately, biopsy studies are uncommon and we are aware of only one study that has reported the effects of alcohol on indices of intracortical bone remodeling [61]. Although the alcoholics in the aforementioned study had reduced cortical thickness associated with increased endocortical bone resorption, all of the chronic alcohol abusers had pancreatitis and many had additional comorbidities that could influence bone metabolism, including poorly controlled type 2 diabetes, iron overload, liver disease and low serum magnesium and vitamin D levels [63]. Thus, the specific effect of chronic heavy alcohol consumption on cortical bone turnover in humans remains to be established.

Rodents have been extremely useful for modeling the effects of alcohol consumption on bone. High levels of alcohol, whether administered continuously in a liquid diet [64, 65], intermittently as a model for binge drinking [66], or by total enteral nutrition [67], have been shown to suppress bone growth in rats. Although small rodents do not normally exhibit intracortical bone remodeling, they do remodel cortical bone at the endocortical surface and several important bone regulating hormones (e.g., estrogen and parathyroid hormone) have been shown to modify marrow cavity volume and cortical bone thickness by regulating endocortical bone formation and resorption [68, 69]. Under certain circumstances, including chronically elevated parathyroid hormone levels,
intracortical bone turnover can be induced in rodents, resulting in greatly increased intracortical porosity [70]. Few studies have investigated the effects of alcohol on cortical bone architecture in skeletally mature rodents, but feeding high levels of alcohol (35% caloric intake) to 6-month-old male rats for 16 weeks was shown to have no effect on intracortical porosity [71] or on marrow volume [72]. These findings suggest that as in male rhesus macaques, chronic heavy alcohol consumption does not result in loss of cortical bone in adult male rats. It is likely that alcohol acts on bone through a combination of mechanisms, both direct and indirect [23–25], although the cellular and molecular mechanisms underlying the skeletal effects of alcohol are not fully understood. High concentrations of alcohol directly affect osteoblasts, osteoclasts, and osteocytes in culture [23]. Furthermore, alcohol may act as an endocrine disruptor by altering levels of and/or skeletal quality and lead to reduced bone mechanical properties independent of bone mass and density or cortical bone architecture. However, this pattern of reduced bone turnover in rodents does not fully correlate with findings in alcoholics. Bone Miner 1987:12:221–5.


Li J, Sato M, Jerome C, Turner CH, Fan Z, Burr DB. Microdamage accumulation in the monkey vertebra does not occur when bone turnover is suppressed by 50% or less with estrogen or raloxifene. J Bone Miner Metab 2005;23(Suppl.):48–54.
