Structural basis of growth-related gain and age-related loss of bone strength

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If bone strength was the only requirement of skeleton, it could be achieved with bulk, but bone must also be light. During growth, bone modelling and remodelling optimize strength, by depositing bone where it is needed, and minimize mass, by removing it from where it is not. The population variance in bone traits is established before puberty and the position of an individual’s bone size and mass tracks in the percentile of origin. Larger cross-sections have a comparably larger marrow cavity, which results in a lower volumetric BMD (vBMD), thereby avoiding bulk. Excavation of a marrow cavity thus minimizes mass and shifts the cortex radially, increasing rigidity. Smaller cross-sections are assembled by excavating a smaller marrow cavity leaving a relatively thicker cortex producing a higher vBMD, avoiding the fragility of slenderness. Variation in cellular activity around the periosteal and endocortical envelopes fashions the diverse shapes of adjacent cross-sections. Advancing age is associated with a decline in periosteal bone formation, a decline in the volume of bone formed by each basic multicellular unit (BMU), continued resorption by each BMU, and high remodelling after menopause. Bone loss in young adulthood has modest structural and biomechanical consequences because the negative BMU balance is driven by reduced bone formation, remodelling is slow and periosteal apposition continues shifting the thinned cortex radially. But after the menopause, increased remodelling, worsening negative BMU balance and a decline in periosteal apposition accelerate cortical thinning and porosity, trabecular thinning and loss of connectivity. Interstitial bone, unexposed to surface remodelling becomes more densely mineralized, has few osteocytes and greater collagen cross-linking, and accumulates microdamage. These changes produce the material and structural abnormalities responsible for bone fragility.

Key words: Ageing, Bone, Growth, Modelling, Remodelling, Fragility, Strength.

Introduction

Propulsion against gravity requires rigidity for leverage, so long bones must be stiff. Impact loading imparts energy that cannot be destroyed so bone must also be flexible to absorb energy by changing shape [1]. The elastic properties of bone allow it to absorb energy by deforming reversibly when loaded [2, 3]. If the load imposed exceeds the bone’s ability to deform elastically, irreversible plastic deformation is accompanied by permanent shape change with accumulation of microcracks that allow energy release [1]. The ability to develop microdamage is defence against the alternative—complete fracture—but microcracking compromises strength as it accumulates [4]. If both elastic and plastic zones of deformation are exceeded, structural failure—fracture—occurs. Bone must also be light to allow mobility.

Bone achieves the paradoxical properties of stiffness yet flexibility and strength yet lightness through its material composition and its structural design. Type 1 collagen is tough and distensible, but lacks resistance to bending so is stiffened by creating a composite of collagen plus mineral. Greater mineral content produces greater material stiffness, but the ability to deform and so absorb and store energy decreases [1]. Ossicles in the ear are over 80% mineral, sacrificing the ability to deform in favour of stiffness. The antlers of deer are less densely mineralized to facilitate deformation. Greater energy-absorbing capacity of antlers is favoured over stiffness, which is not needed as they are not load-bearing [5].

Growth and the attainment of peak bone strength and minimal mass

During growth, bone material is fashioned into three-dimensional masterpieces of biomechanical engineering by bone modelling, the formation of bone by osteoblasts without prior bone resorption. This process is vigorous during growth and changes bone size and shape. During remodelling, bone is refashioned first by resorption by osteoclasts, which remove bone, and then osteoblasts deposit bone in the same location. These cells form the basic multicellular unit (BMU), which reconstructs bone in distinct locations on the three (endocortical, intracortical and trabecular) components of its inner (endocortical) envelope and to a much lesser extent on the periosteal (outer) envelope [6, 7].

Achieving strength by modifying mass distribution rather than mass

If bone had only to be strong it could achieve this with bulk, but bone must also be light to facilitate mobility. Longer tubular bones need more mass to construct their greater length than shorter bones, but wider and narrower cross-sections do not necessarily differ in the amount of material needed to construct them [8]. Wider and narrower bone cross-sections are assembled using a similar amount of material, so larger cross-sections are assembled with less material relative to their size, producing a lower apparent volumetric BMD (vBMD) avoiding bulk. Smaller cross-sections are assembled with more material relative to their size, producing a higher vBMD and avoiding the fragility of slenderness (Fig. 1). Wider tubular bones are assembled with a thinner cortex (producing the same bone area because the thinner ‘ribbon’ of cortex is distributed around a larger perimeter).

The diversity in bone size, shape and mass distribution is the result of differing degrees of focal modelling around the periosteal perimeter and remodelling at the corresponding point on the endocortical surface during growth. Bone strength is optimized using the minimum net amount of bone needed. For example, total cross-sectional area (CSA) of the femoral neck is greatest adjacent to the shaft of the femur and smaller nearer the femoral head, but the amount of bone in each cross-section is no different (Fig. 2).

Adjacent to the femoral shaft, the femoral neck cross-section is elliptical with a long axis in the supero-inferior direction. Greater periosteal apposition superiorly and inferiorly than...
medio-laterally produces the elliptical shape. Greater periosteal apposition and perhaps less endocortical resorption inferiorly produces a thicker cortex than superiorly [8]. The bone in the cross-section at the junction of the femoral neck with the femoral shaft is largely cortical. Moving proximally, femoral neck shape becomes more circular, reflecting similar degrees of periosteal apposition around the perimeter, and the bone mass is distributed progressively more as trabecular and less as cortical bone, while cortical thickness is similar around the perimeter.

Early establishment of differences in bone size, shape and mass

Differences in bone size are established early in life, before puberty and perhaps even in utero. In a 3-yr prospective study of growth in 40 boys and girls, Loro et al. [9] report that the variance at Tanner Stage 2 (pre-puberty), in vertebral CSA and vBMD, femoral shaft CSA and cortical area, was no less than at Tanner Stage 5 (maturity); 60–90% of the variance at maturity was accounted for by the variance present before puberty. Thus, the magnitude of trait variances is established before puberty [9]. The ranking of individual values at Tanner Stage 2 was unchanged during 3 yrs in girls (Fig. 3).

These traits were tracked, and an individual with a large vertebral or femoral shaft cross-section, or higher vertebral vBMD or femoral cortical area, before puberty had these traits at maturity.

The deposition of the same amount of bone on the periosteal surface of an already larger cross-section confers more bending resistance than deposition of the same amount of bone on
a smaller cross-section, because resistance to bending is proportional to the fourth power of the distance from the neutral axis [10]. In this way, larger cross-sections are assembled with less mass (relative to their size) avoiding bulk, while smaller cross-sections are assembled with more mass (relative to their size), offsetting the fragility of slenderness.

Sexual dimorphism in bone structure rather than mass

The vertebral body is wider in males than in females [11]. Trabecular number per unit area is constant during growth; therefore, individuals with a low trabecular number in young adulthood are likely to have lower trabecular numbers in childhood [12]. The age-related increase in trabecular density is the result of increased thickness of existing trabeculae. Before puberty, there is no difference in trabecular density in boys and girls of either Caucasian or African American origin [13]. At puberty, trabecular density increases, but within a race there is no sex difference in trabecular density.

Growth does not build a ‘denser’ vertebral body in males than females, it builds a bigger vertebral body in males. Strength of the vertebral body is greater in young males than females because of size differences. Within a sex, African Americans have a higher iliac crest trabecular density than whites due to a greater increase in trabecular thickness [14].

As tubular bones increase in length by endochondral apposition, periosteal apposition widens the bone while concurrent endocortical resorption excavates the marrow cavity. As periosteal apposition is greater than endocortical resorption, the cortex thickens. In females, earlier completion of longitudinal growth with epiphyseal fusion and earlier inhibition of periosteal apposition produces a smaller bone. Cortical thickness is similar in males and females because endocortical apposition in females contributes to final cortical thickness [15]. Cortical thickness is similar by race and sex; what differs is the position of the cortex relative to the long axis of the long bone [16]. Racial differences in trabecular vBMD are also reported, but the morphological basis for these differences is yet to be defined [17].

Varicance in bone mass at completion of growth is an order of magnitude greater than variance in rates of bone loss during ageing (1 s.d. = 10% vs 1%, respectively), so bone size, architecture and mass attained during growth determine the relevance of bone loss during advancing age. For example, in children with larger tibial cross-sections, the advantage of avoiding bulk by assembling the larger bone with a relatively thinner cortex may be a disadvantage when age-related bone loss occurs. Women with hip fractures and their daughters have larger femoral neck diameters and reduced vBMD [18].

Adulthood and the emergence of bone fragility

The purpose of modelling and remodelling during adulthood is to maintain bone strength by removing damaged bone. Bone, like roads, buildings and bridges, develops fatigue damage during repeated loading, but only bone has a mechanism enabling it to detect the location and magnitude of the damage, to remove it, replace it with new bone and so to restore the bone’s material composition, micro- and macro-architecture [19, 20]. Resorption is not necessarily bad. The resorptive phase of the remodelling cycle removes damaged bone and is essential to bone health. The formation phase of the remodelling cycle restores the bone’s structure.

Microcracks damage the canalicular system causing osteocyte apoptosis [21]. Osteocytic death from many causes, such as corticosteroid therapy or oestrogen deficiency, is associated with loss of bone strength and so may be a form of damage itself [22, 23]. Osteocyte death provides the topographical information needed to identify the location and extent of damage, initiate osteoclastogenesis and provide an appropriately sized work force of osteoclasts for targeted remodelling. Apoptosis precedes osteoclastogenesis [24, 25]. Death of the osteocyte-like cell line MLO-Y4, induced by scratching, results in the formation of tartrate-resistant acid phosphatase-positive cells along the scratching path. Osteocyte apoptosis occurs within 3 days of immobilization and is followed by osteoclastogenesis and bone resorption [26].

Damage may be signalled via the osteocytic network to flattened bone lining cells, which digest unmineralized osteoid creating a cavity beneath which becomes a bone remodelling compartment (BRC). Osteoclast precursors may be delivered from the marrow via the circulation for both cortical osteonal remodelling and trabecular hemiosteonal remodelling [27]. Osteoblast precursors may arise from local marrow stromals, from the circulation or from the canopy of the BRC. Whatever the mechanism, bone formation follows resorption partly or completely refilling the excavation site. Most osteoblasts die, others become lining cells, while others are entombed in osteoid leading to ‘rewiring’ of the osteocytic canalicular communicating system for subsequent mechanotransduction, damage detection and repair [28] (Fig. 4).
Abnormalities in bone remodelling

Four age-related changes in bone modelling and remodelling compromise bone’s material properties and structural design [29].

(i) A reduction in bone formation at the tissue level. Periosteal apposition slows precipitously after completion of longitudinal growth and continues in adulthood modestly during the next 60 yrs [30–32].

(ii) A reduction in bone formation at the cellular level within each BMU [33, 34].

(iii) Continued resorption in the BMU. The volume of bone resorbed in each BMU does not increase, but decreases or remains unchanged [23, 35, 36].

(iv) An increase in the rate of bone remodelling after the menopause accompanied by worsening of the negative bone balance in each BMU as the volume of bone resorbed increases and the volume of bone formed decreases in the many more BMUs now remodelling bone [23] (Fig. 5).

At some stage in young adulthood, the volume of bone formed in the formation phase of a remodelling cycle is less than the volume of bone resorbed in the resorptive phase of that cycle producing a net negative BMU balance, bone loss, structural decay and bone fragility. Bone mass decreases in young adulthood due to the decline in bone formation [37–39]. About 40% of the trabecular bone lost across life is lost before the age of 50 yrs in women and men, but cortical loss is minimal before the age of 50 yrs. The consequences are likely to be less than bone loss later because remodelling rate is slow, bone loss proceeds by reduced bone formation rather than increased bone resorption in the BMU, bone loss proceeds by trabecular thinning rather than by loss of connectivity, producing less loss of strength [40]. Periosteal apposition partly offsets endocortical bone loss and shifts the cortices radially [32].

At the menopause, the steady state is perturbed by an increase in the birth rate of new BMUs on bone’s endosteal envelope. The many BMUs remove bone while the fewer BMUs created before the menopause complete remodelling by depositing bone. This perturbation produces accelerated bone loss and a rapid decline in BMD. This is a partly reversible loss of bone mass and bone mineral is produced by the normal delay in onset and slower progression of the formation phase of the remodelling cycle [41]. The temporary deficit in bone mass and mineral has three components, the excavation site, the osteoid that lacks mineral and bone that has undergone primary but not secondary mineralization, the slow enlargement of crystals of calcium hydroxyapatite-like mineral whose completion takes months to years [42].

Bone loss slows in the 3–5 yrs after the menopause because the steady state is restored at a new higher remodelling rate. Now the large numbers of BMUs created producing resorption of bone are matched by completion of bone formation of their remodelling cycle and by the large numbers of BMUs created in the perimenopause. Bone loss continues at a faster rate than before the menopause, but at a slower rate than immediately after, because BMU balance is perhaps more negative than before the menopause. Now there are many more sites being created and completed, but the negative BMU balance produces a permanent deficit in bone mass and mineral mass driven by the high remodelling rate.

Structural changes resulting from abnormalities in remodelling

Remodelling occurs on bone surfaces. As trabecular bone has more surface per unit bone volume than cortical bone, accelerated bone remodelling initially produces more trabecular than cortical bone loss. Complete loss of trabeculae reduces the surface available for remodelling, but remodelling on endocortical and intracortical surfaces increases the surface available for remodelling in cortical bone; cortical bone is trabecularized. Cortical thinning occurs and increased porosity results in an increase in the surface/volume ratio in cortical bone. The total bone surface available for resorption does not change (increasing in cortical bone, decreasing in trabecular bone) or increases (in regions of cortical bone only),
so that late in life bone loss is more cortical than trabecular in origin. The continued remodelling at a similar intensity with its negative BMU balance on the same amount or more surface removes the same amount of bone from an ever decreasing amount of bone accelerating bone loss and structural decay.

Rapid remodelling is associated with an increased risk of fracture because more densely mineralized bone is removed and replaced with younger, less densely mineralized bone, reducing stiffness, and excavated resorption sites create stress 'concentrators', which predispose to microdamage, and increased remodelling impairs isomerization of collagen [43–45]. Interstitial bone, deep bone, is not exposed to remodelling and becomes more densely mineralized, and more highly cross-linked with advanced glycation products like pentosidine, both of which reduce bone toughness. It is easier for microcracks to travel through a more homogeneously mineralized bone. Interstitial bone, which has fewer osteocytes, accumulates microdamage [46–50]. Cortical thinning and increased porosity reduce resistance to crack propagation. Pores coalesce so that the number of pores in cortical bone decreases, but the total area of porosity increases late in life. The ability of bone to limit crack propagation declines so that bone cannot absorb the energy imparted by a fall and this energy is released in the worst possible way—fracture.

Reduced periosteal apposition

Periosteal apposition during ageing is slow [30]. It is believed to increase as an adaptive response to compensate for the loss of strength produced by endocortical bone loss. In a 7-yr prospective study of over 600 women, Szule et al. [32] report that endocortical bone loss occurred in pre-menopausal women with concurrent periosteal apposition. Periosteal apposition was less than endocortical resorption so the cortices thinned, but there was no net bone loss because the thinner cortex was distributed around a larger perimeter, conserving total mass, and resistance to bending increased because this same amount of bone was distributed further from the neutral axis (Fig. 6).

Endocortical resorption increased during the perimenopausal period, but periosteal apposition decreased—it did not increase in compensation so the cortices thinned. Nevertheless, bending strength remained unchanged because periosteal apposition was still sufficient to shift the thinning cortex outwards. Bone fragility emerged after the menopause when acceleration in endocortical bone resorption and deceleration in periosteal apposition produce further cortical thinning with little outward displacement of the thinning cortex, so the cortical area now declined, as did resistance to bending.

The periosteal envelope is not only a bone-forming surface [30]. Blizotes et al. [7] report that bone resorption occurs in adult non-human primates. Femurs from 16 intact adult male and female non-human primates showed that periosteal remodelling of the femoral neck in intact animals was slower than in cancellous bone, but more rapid than at the femoral shaft. Gonadectomized females showed an increase in osteoclast number on the periosteal surface compared with intact controls. If these findings are
correct, adult skeletal dimensions may decrease in size as age advances.

Fuller Albright suggested over 65 yrs ago that osteoporosis was a disorder of reduced bone formation [51]. During ageing, both increasing endocortical bone resorption and reduced periosteal apposition cause net bone loss, alterations in the distribution of the remaining bone and the emergence of bone fragility. The cellular basis of the vigour of bone formation during growth and progressive decline in vigour during ageing on the periosteal surface and within each BMU is yet to be defined.

Sex differences in trabecular and cortical bone loss

A greater proportion of women than men sustain fragility fractures because (i) men’s skeletons are larger than women’s, and therefore more resistant to bending; (ii) men do not have a mid-life decline in sex hormones and increase in remodelling rate; (iii) bone loss in most men is the result of a negative BMU balance produced by reduced formation not increased resorption, so trabecular bone loss occurs by thinning rather than loss of connectivity [52]; the loss of strength is less than produced by loss of connectivity [40] even though the amount of trabecular bone loss across age is only slightly greater in women than men [53], or is similar [16, 52, 54–59]; (iv) cortical porosity increases less in men than in women because remodelling rate is lower in men; thus, crack propagation in cortical bone is probably better resisted in men than in women; and (v) periosteal apposition is purported to be greater in men than in women in some [16, 58–59], but not all studies [53].

The absolute risk for fracture in women and men of the same age and BMD is similar [60, 61]. Thus, the lower fracture incidence in men than in women is likely to be the result of lower proportion of elderly men than elderly women with material and structural properties below the level at which the loads on the bone are greater than the bone’s net ability to tolerate them.

Structural failure occurs less in men because the relationship between load and bone strength is better maintained in men than in women [62].

**Heterogeneous material and structural basis of bone fragility in patients with fractures**

Patients with fractures do not share the same pathogenesis and structural basis of bone fragility. Patients with vertebral fractures may have high, normal or low remodelling rates [63], while others have a negative BMU balance due to reduced formation, increased resorption, or both, or no negative BMU balance at all [64]. Some patients with vertebral fractures have increased, others reduced, tissue mineral density [65]. Some patients have reduced osteocyte density, while others do not [66]. Whether anti-fracture efficacy can be improved by defining the pathogenesis and structural basis in an individual remains uncertain, but it is worthy of consideration.

**Conclusion**

Modelling and remodelling are successful during growth, but not ageing. Longevity is accompanied by reduced bone formation on the periosteal envelope and abnormalities in remodelling balance and rate on the endosteal envelope that compromise the material and structural properties of bone. Understanding of why or how bones fail at the material and structural level is essential if we are to provide targeted approaches to drug therapy.

**Rheumatology key messages**

- Bone fragility is the result of age-related abnormalities in bone remodelling and the osteocoenic system which prevents damage accumulation and removes it when it occurs.
- Each remodelling event, whether adaptive or reparative leaves a deficit in bone volume producing structural decay.

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