Bone health in the prostate cancer patient receiving androgen deprivation therapy: a review of present and future management options

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Abstract

Osteoporosis and bone fractures are frequently overlooked complications of androgen deprivation therapy in men with nonmetastatic prostate cancer. All such patients should have their bone mineral density (BMD) monitored and be offered preventive measures, such as calcium and vitamin D supplementation; patients with low BMD should be offered treatment. Several agents, including bisphosphonates, are available (although this use is currently off-label), and upcoming treatments, such as denosumab and toremifene, have shown promise in reducing fracture risk in these patients.

Why should bone loss concern urologists?

Androgen deprivation therapy (ADT) is increasingly being prescribed both for men with locally advanced or high-risk nonmetastatic prostate cancer and for those with recurrent disease.1,2 With this increased exposure to ADT, clinicians have seen the emergence of longer-term treatment complications, including osteoporosis and osteopenia. Although osteoporosis is generally less frequent in men, it is increasingly recognized as a source of substantial morbidity and even mortality in the aging male. Men suffer one third of all hip fractures. In a Canadian study of 3981 hip fracture patients, the 1-year mortality rate among men over 60 after a hip fracture was 37.5%, exceeding that among women (28.3%; \(p < 0.001\)).3 Osteoporotic vertebral fractures have a radiological prevalence of up to 50% in both sexes; they often cause chronic pain, and even clinically silent fractures are associated with increased risks of future fracture (both vertebral and hip), kyphosis, restricted lung function, impaired activities of daily living and even increased mortality.4 A study of Canadian prostate cancer patients who were orchiectomized found that their 5-year risks of vertebral and hip fractures were 2.2-fold higher than those of patients who had not been orchiectomized (\(p < 0.001\) for both).5 Further, a claims-based study carried out in Ontario compared 19,079 men aged at least 66 years with prostate cancer who had used ADT for at least 6 months or who had undergone orchiectomy with matched controls who had not received ADT; the former group had a significantly increased risk of fragility fractures \((HR 1.65; 95\% CI 1.53–1.77)\).6 Among prostate cancer patients receiving ADT, induced hypogonadism is a major risk factor for osteoporosis and, hence, of fracture. Fractures also independently predict diminished survival in prostate cancer patients on ADT. In one retrospective study, a history of fracture since the diagnosis of prostate cancer decreased median overall survival from 160 months to 121 months \((p = 0.04)\).7

Despite these potentially serious complications, osteoporosis and osteopenia are greatly underdiagnosed and undertreated in this population. A recent retrospective chart review involving 174 veterans with prostate cancer in New Mexico who were receiving ADT found that only 34% of those with nonmetastatic disease had received any recommended screening, prophylaxis, or therapy for osteoporosis, and only 13% had received a dual-energy x-ray absorptiometry (DXA) scan. Many of these patients had additional risk factors for osteoporosis including diabetes, smoking, alcohol use and treatment with corticosteroids.8

This overview focuses on cancer-treatment-induced bone loss (CTIBL) in the patient with nonmetastatic prostate cancer. Reflecting the latest research in CTIBL, this article discusses...
mechanisms of bone loss, risk factors, treatment options and offers practical advice for the community urologist regarding the evaluation and management of their prostate cancer patient’s bone health.

The RANK/RANKL pathway and the effects of hormonal therapy on bone

Bones are composed of 2 main types of tissues: cortical bone and trabecular bone. Cortical (compact) bone is 80% to 90% calcified and has mainly mechanical and protective functions. Trabecular (cancellous or spongy) bone is only 15% to 25% calcified and constitutes only 20% of the total bone mass, but carries out most of the bone’s metabolic function. Bone strength is a function of bone mass and of other parameters including geometry (e.g., the diameter of the cortical bone), material properties (e.g., the quality of the bone matrix and inorganic crystals) and microstructure (e.g., the diameter and interconnectivity of the trabeculae). The bone mass of a normal adult is the outcome of a dynamic equilibrium between bone formation (mediated by osteoblasts) and bone resorption (mediated by osteoclasts). The function of the latter cells is regulated by a family of proteins that include receptor activator of nuclear factor κ-B (RANK), RANK ligand (RANKL) and osteoprotegerin (OPG). Binding of RANKL to RANK on the surfaces of osteoclast precursors will trigger maturation, activation, and prolonged survival of these cells. Thus, RANKL promotes bone resorption. In contrast, OPG is a “decoy receptor” that binds and neutralizes RANKL, thus inhibiting bone resorption. There is interplay between RANKL and OPG (Fig. 1a).

The ratio of RANKL to OPG is a critical factor determining the balance between bone resorption and bone formation. Vitamin D₃, parathyroid hormone, tumour necrosis factor-α (TNF-α), activated T-cells, and glucocorticoid therapy all increase this ratio, promoting bone resorption. Estrogen deficiency states (including menopause) also produce osteoporosis because normal levels of 17β-estradiol inhibit RANKL production and stimulate OPG. Testosterone stimulates osteoblasts, inhibits the apoptosis of both osteoblasts and osteoclasts, and is a precursor of estrogen via aromatization; its net effect is to stimulate bone formation. In males with hypogonadism (whether induced by orchietomy, ADT, hyperparathyroidism, or other causes), both testosterone and estrogen levels fall, shifting the balance of bone turnover toward resorption (Fig. 1b). It has been hypothesized that several malignancies including prostate and breast cancer and multiple myeloma also promote bone resorption by expressing or stimulating RANKL.

Who is at risk? Who should be screened?

In its clinical practice guidelines on the diagnosis and management of osteoporosis, the Osteoporosis Society of Canada (OSC) identified 4 robust, independent risk factors for osteoporotic fracture: low bone mineral density (BMD), a prior fragility fracture, age ≥65 and a family history of osteoporosis (Box 1). In turn, there are also other risk factors for osteoporosis, including lifestyle and dietary factors, and diseases and treatments associated with bone loss (Table 1).

**Fig. 1a.** Interplay between RANKL and OPG in the control of osteoclast activity. RANKL is expressed by osteoblasts and binds to its specific receptor RANK on the surface of osteoclast precursors, triggering pathways that promote the differentiation, activation and survival of osteoclasts. OPG binds to RANK and neutralizes it, blocking the differentiation and activation of new osteoclasts and shortening the survival time of existing osteoclasts. Adapted from Boyle WJ, et al. *Nature* 2003;423:337.

**Fig. 1b.** Hormonal effects on bone. Estrogen inhibits bone resorption, while androgens promote bone formation. When either androgen or estrogen is deficient (for example, in postmenopausal patients, those taking aromatase inhibitors or men with prostate cancer taking ADT), bone resorption predominates over bone formation, and the net effect is loss of bone. Adapted from Perez EA, et al. *Oncology, Williston Park*, 2006;1029.
Prostate cancer itself is associated with osteoporosis, even among ADT-naive patients without metastatic disease. In a cross-sectional study, 45.2% of such patients had osteopenia and 35.4% had osteoporosis even before starting ADT. In addition, prevalence increased with duration of treatment until after 10 years no patient on ADT had a BMD within the normal range (Fig. 2). Numerous prospective longitudinal studies have confirmed that BMD generally decreases significantly at the spine and hip, particularly during the first year of ADT; reported BMD losses after only 1 year of ADT range up to 4.8% at the lumbar spine and 3.8% at total hip.

Systematic retrospective reviews have also shown the association between ADT and increased fracture risk. For example, a large study of Medicare records from 50613 prostate cancer patients in the United States found overall fracture rates of 19.4% after 5 years among those who received ADT within 6 months of diagnosis and 12.6% among those who did not (p < 0.001; Fig. 3). The authors estimated that about 3000 fractures a year were attributable to gonadotropin-releasing hormone (GnRH) agonist use.

Because age and hypogonadism are both considered major risk factors for osteoporosis, all prostate cancer patients beginning ADT should be screened with DXA scans at baseline; anyone aged ≥65 and anyone with kyphosis, back pain, substantial height loss, or other symptoms suggesting vertebral fractures should also be screened with thoracic and lumbar spine x-rays. These recommendations are consistent with those of a recent Canadian review of the diagnosis and management of osteoporosis in men.

### Lifestyle measures, calcium and vitamin D supplementation

Lifestyle modifications to address osteoporosis include exercise, smoking cessation and moderating alcohol and caffeine intake. In addition, men over 50 should have a total of 1500 mg daily of calcium and 800 IU daily of vitamin D (D₂ being preferable to D₃). However, the OSC guidelines state that while adequate calcium and vitamin D (whether dietary or supplemented) are essential adjuncts to prevent and treat osteoporosis, they are insufficient by themselves as treatments.

### Bisphosphonates

Bisphosphonates (BPs) are often used for patients with bone metastases to prevent pathologic fractures, reduce bone pain or control hypercalcaemia, but they are not specifically indicated for the prevention and treatment of CTIBL. Numerous randomized controlled trials have explored the effects of BPs on BMD in the setting of ADT for nonmetastatic prostate...
Table 2. Published randomized controlled trials in the management of cancer-treatment-induced bone loss in prostate cancer patients receiving androgen deprivation therapy

<table>
<thead>
<tr>
<th>ADT</th>
<th>Author (year)</th>
<th>N</th>
<th>Treatment arms</th>
<th>Duration of study follow-up</th>
<th>Key end point</th>
<th>Main results*</th>
<th>Adverse events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bisphosphonates — alendronate</td>
<td>Greenspan et al. (2007)</td>
<td>112</td>
<td>Alendronate 70 mg po q 1 week vs. placebo</td>
<td>1 year</td>
<td>BMD</td>
<td>Alendronate &gt; placebo (spine, FN)</td>
<td>NSS</td>
</tr>
<tr>
<td>Bisphosphonates — alendronate</td>
<td>Greenspan et al. (2008)</td>
<td>112</td>
<td>Continue alendronate 70 mg po q 1 week vs. crossover to placebo vs. crossover to alendronate</td>
<td>2 years (re-randomization after 1 year)</td>
<td>BMD</td>
<td>Alendronate/ alendronate &gt; alendronate/ placebo, placebo/ alendronate (spine, TH, FN)</td>
<td>NSS</td>
</tr>
<tr>
<td>Bisphosphonates — neridronate</td>
<td>Morabito et al. (2004)</td>
<td>48</td>
<td>Neridronate 25 mg IM q 1 month vs. none</td>
<td>1 year</td>
<td>BMD</td>
<td>Neridronate &gt; none (spine, TH)</td>
<td>None relevant</td>
</tr>
<tr>
<td>Bisphosphonates — neridronate</td>
<td>Magno et al. (2005)</td>
<td>60</td>
<td>Maximal androgen blockade (A) or bicalutamide (B), with neridronate 25 mg IM q 1 month (A2, B2) or without (A1, B1) (n = 15 each)</td>
<td>1 year</td>
<td>BMD</td>
<td>Arms with neridronate &gt; arms without neridronate</td>
<td>None relevant</td>
</tr>
<tr>
<td>Bisphosphonates — pamidronate</td>
<td>Diamond et al. (2001)</td>
<td>21</td>
<td>Pamidronate 90 mg IV q 6 months vs. placebo (crossover at 6 months)</td>
<td>1 year (crossover at 6 months)</td>
<td>BMD</td>
<td>Pamidronate &gt; placebo (spine, FN)</td>
<td>Not reported</td>
</tr>
<tr>
<td>Bisphosphonates — pamidronate</td>
<td>Smith et al. (2001)</td>
<td>47</td>
<td>Pamidronate 60 mg IV q 3 months vs. none</td>
<td>48 weeks open-label</td>
<td>BMD</td>
<td>Pamidronate &gt; none (spine, TH, trochanter)</td>
<td>2 withdrawals (angiosarcoma, memory disorder) in pamidronate group 3 acute phase reactions in pamidronate group</td>
</tr>
<tr>
<td>Bisphosphonates — zoledronic acid</td>
<td>Smith et al. (2003)</td>
<td>106</td>
<td>Zoledronic acid 4 mg IV q 3 months vs. placebo</td>
<td>1 year</td>
<td>BMD</td>
<td>Zoledronic acid &gt; placebo (spine, FN, TH)</td>
<td>NSS</td>
</tr>
<tr>
<td>Bisphosphonates — zoledronic acid</td>
<td>Ryan et al. (2006)</td>
<td>122</td>
<td>Zoledronic acid 4 mg IV q 3 months vs. placebo</td>
<td>1 year, double-blind</td>
<td>BMD</td>
<td>Zoledronic acid &gt; placebo (spine, FN, TH)</td>
<td>Nausea more common in zoledronic acid group Otherwise, No mandibular osteonecrosis seen</td>
</tr>
<tr>
<td>Bisphosphonates — zoledronic acid</td>
<td>Ryan et al. (2007)</td>
<td>42</td>
<td>Zoledronic acid 4 mg IV q 3 months vs. placebo</td>
<td>1 year, double-blind</td>
<td>BMD</td>
<td>Zoledronic acid &gt; placebo (spine, FN)</td>
<td>NSS (1 severe AE in placebo group)</td>
</tr>
<tr>
<td>Bisphosphonates — zoledronic acid</td>
<td>Michaelson et al. (2007)</td>
<td>40</td>
<td>Zoledronic acid 4 mg IV (single dose) vs. placebo</td>
<td>1 year</td>
<td>BMD</td>
<td>Zoledronic acid &gt; placebo (spine, TH)</td>
<td>No serious treatment-related adverse events</td>
</tr>
<tr>
<td>Bisphosphonates — zoledronic acid</td>
<td>Israeli et al. (2007)</td>
<td>215</td>
<td>Zoledronic acid 4 mg IV q 3 months vs. placebo</td>
<td>1 year</td>
<td>BMD</td>
<td>Zoledronic acid &gt; placebo (spine, TH)</td>
<td>NSS Traumatic fractures for 2 in zoledronic acid group and 3 in placebo group No ONJ</td>
</tr>
<tr>
<td>SERMs — raloxifene</td>
<td>Smith (2004)</td>
<td>48</td>
<td>Raloxifene 60 mg daily vs. none</td>
<td>12 months, open-label</td>
<td>BMD</td>
<td>Raloxifene &gt; none (TH)</td>
<td>Well tolerated</td>
</tr>
<tr>
<td>SERMs — toremifene citrate</td>
<td>Steiner (2004)</td>
<td>46</td>
<td>Toremifene citrate vs. placebo</td>
<td>6 months</td>
<td>BMD</td>
<td>Toremifene &gt; placebo</td>
<td>Also decreased hot flushes</td>
</tr>
<tr>
<td>Nonsteroidal antiandrogens — bicalutamide</td>
<td>Smith (2004)</td>
<td>52</td>
<td>Bicalutamide vs. leuprolide</td>
<td>12 months, open-label</td>
<td>BMD</td>
<td>Bicalutamide &gt; leuprolide (spine, etc.)</td>
<td>Also decreased fat mass, fatigue, loss of libido, hot flushes</td>
</tr>
<tr>
<td>Antibodies directed against RANKL — denosumab</td>
<td>Smith et al. (2009)</td>
<td>1468</td>
<td>Denosumab 60 mg s.c. q 6 months vs. placebo</td>
<td>36 months, double-blind</td>
<td>BMD</td>
<td>Denosumab &gt; placebo (lumbar spine, total hip, femoral neck, distal radius)</td>
<td>NSS</td>
</tr>
</tbody>
</table>

BMD = bone mineral density; FN = femoral neck; NSS = not statistically significant; ONJ = osteonecrosis of the jaw; TH = total hip; SERMs = selective estrogen receptor modulators; RANKL = receptor activator for nuclear factor k B ligand. *A > B means A produced a significantly better result than B. Printed with permission. Saad F, et al. J Clin Oncol 2008;26:1.
Bone health in prostate cancer

Fig. 3. Prevalence of fractures in men with prostate cancer: effect of ADT type and dose. Unadjusted fracture-free survival among patients with prostate cancer, according to androgen-deprivation therapy. The survival curves start at 12 months after diagnosis and androgen deprivation was initiated within 6 months after diagnosis. GnRH denotes gonadotropin-releasing hormone. The number of doses is the number administered within 12 months after diagnosis. Printed with permission. Shahinian VB, et al. N Engl J Med 2005;352:154.

Table 3. A simple algorithm for the management of cancer-treatment-induced bone loss in patients with nonmetastatic prostate cancer receiving androgen deprivation therapy

<table>
<thead>
<tr>
<th>DXA scans</th>
<th>Obtain baseline BMD</th>
<th>Repeat every 1 to 2 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thoracic and lumbar spine x-rays</td>
<td>If patient is ≥65 years old or has kyphosis, height loss ≥6 cm, or acute severe back pain: Rule out vertebral fracture</td>
<td></td>
</tr>
<tr>
<td>Lifestyle modifications</td>
<td>Calcium and vitamin D intake</td>
<td>Smoking cessation</td>
</tr>
<tr>
<td>Treatment</td>
<td>Consider medical therapy to increase BMD and/or reduce fracture risk</td>
<td></td>
</tr>
</tbody>
</table>


Raloxifene

Raloxifene is a selective estrogen receptor modulator (SERM) often used to treat osteoporosis in women. In a 12-month open-label study enrolling 48 men with nonmetastatic prostate cancer receiving ADT, the addition of raloxifene 60 mg daily significantly improved BMD at the total hip and spine.32

Bicalutamide

Bicalutamide is a nonsteroidal anti-androgen, which increases estradiol levels when given as monotherapy. A 12-month, open-label comparison of leuprolide versus bicalutamide (150 mg daily) in 52 men with nonmetastatic prostate cancer showed that bicalutamide increased BMD at several sites (e.g., lumbar spine BMD +2.5% vs. −2.5%; p < 0.001), as well as decreasing fat mass, fatigue, loss of libido and hot flushes compared with leuprolide. Breast tenderness and enlargement were seen more frequently in the bicalutamide group.34 Finally, a recent prospective study whose population included 253 prostate cancer patients with osteoporosis found that bicalutamide treatment maintained BMD over 6 years.41

Upcoming agents: toremifene citrate and denosumab

The effects on BMD of toremifene citrate, a new SERM, were tested in a 6-month, placebo-controlled dose-finding study with 46 men with prostate cancer receiving ADT. An oral dose of 60 mg daily significantly improved BMD and decreased hot flushes.33 A 2-year, double-blind, placebo-controlled phase-III multicentre study of oral toremifene 80 mg has been completed in 1389 ADT patients with advanced prostate cancer; this compound reduced new morphometric vertebral fractures (the primary endpoint).
by 53% ($p = 0.034$). Bone mineral density at lumbar spine, hip, and femur was also increased significantly ($p < 0.0001$), and lipid profiles were improved compared with placebo.42,43

Denosumab is a fully human monoclonal antibody that specifically targets RANKL and is delivered by subcutaneous injection twice a year.44 This therapy was found to be very effective in reducing fractures and was well-tolerated in the clinical settings of osteoporotic postmenopausal women45,46 and protecting BMD in osteopenic postmenopausal women receiving adjuvant aromatase inhibitors for breast cancer.47 More recently, denosumab (60 mg subcutaneously, every 6 months) was evaluated in a 36-month, phase-III, placebo-controlled randomized clinical trial involving 1468 men with nonmetastatic prostate cancer who were receiving ADT.48 Compared with placebo, denosumab significantly improved BMD at all sites measured, including lumbar spine (the primary endpoint) by 6.7% ($p < 0.001$), total hip by 4.8% ($p < 0.001$), femoral neck by 3.9% ($p < 0.001$), and distal radius by 5.5% ($p < 0.001$) at 24 months; by the end of the trial (36 months), denosumab dramatically reduced the risk of new vertebral fractures (a secondary endpoint) by 62% ($p = 0.006$). Due to the entrance criteria of the study, the patients enrolled in the study were healthier than average prostate cancer patients in terms of BMD. Based on current knowledge in the field of osteoporosis, the impressive reduction in fracture risk reported in the pivotal denosumab study will likely be at least as impressive if one were to begin therapy in patients with low BMD and at higher risk of fractures. Denosumab was well-tolerated, with rates of serious adverse events similar to placebo.

**Conclusion**

Among patients with nonmetastatic prostate cancer who are receiving long-term ADT, CTIBL is a frequent, serious, and often overlooked treatment complication. Cancer-treatment-induced bone loss can readily be detected with DXA scans, and essential measures such as calcium and vitamin D supplementation can be offered to the patient at the outset. Although, at this point, evidence is sparse for fracture reduction in this setting, BPs are most often recommended to treat CTIBL in prostate cancer. Upcoming treatments, such as denosumab and toremifene citrate, have demonstrated promise in this setting. A simple 4-step algorithm for the management of CTIBL is shown in Table 3.14,17

With the increased use of agents affecting bone metabolism and an aging population susceptible to the complications associated with CTIBL, it behooves all urologists to become more familiar with the biology, diagnosis, and treatment of ADT-induced bone disease.

References

Bone health in prostate cancer


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