Osteoporosis and type 2 diabetes mellitus: what do we know, and what we can do?

Abstract: Diabetes mellitus (DM) is a pandemic and chronic metabolic disorder with substantial morbidity and mortality. In addition, osteoporosis (OP) is a silent disease with a harmful impact on morbidity and mortality. Therefore, this systematic review focuses on the relationship between OP and type 2 diabetes mellitus (T2DM). Systematic reviews of full-length articles published in English from January 1950 to October 2010 were identified in PubMed and other available electronic databases on the Universiti Sains Malaysia Library Database. The following keywords were used for the search: T2DM, OP, bone mass, skeletal. Studies of more than 50 patients with T2DM were included. Forty-seven studies were identified. The majority of articles (26) showed increased bone mineral density (BMD), while 13 articles revealed decreased BMD; moreover, eight articles revealed normal or no difference in bone mass. There were conflicting results concerning the influence of T2DM on BMD in association with gender, glycemic control, and body mass index. However, patients with T2DM display an increased fracture risk despite a higher BMD, which is mainly attributable to the increased risk of falling. As a conclusion, screening, identification, and prevention of potential risk factors for OP in T2DM patients are crucial and important in terms of preserving a good quality of life in diabetic patients and decreasing the risk of fracture. Patients with T2DM may additionally benefit from early visual assessment, regular exercise to improve muscle strength and balance, and specific measures for preventing falls. Patient education about an adequate calcium and vitamin D intake and regular exercise is important for improving muscle strength and balance. Furthermore, adequate glycemic control and the prevention of diabetic complications are the starting point of therapy in diabetic patients.

Keywords: bone, diabetes, osteopenia, osteoporosis, skeletal

Introduction

Diabetes mellitus (DM) is a chronic metabolic disorder with substantial morbidity and mortality, characterized by the presence of hyperglycemia. Pharmacotherapy, continuing medical care, and education are crucial for preventing acute and chronic complications. On the other hand, osteoporosis (OP) is a painless weakening of the bones that constitutes an enormous socioeconomic crisis, with a harmful impact on morbidity and mortality. It leads to increased skeletal fragility and microarchitectural deterioration of bone tissue, causing a decrease in bone mineral density (BMD), bone quality, and strength. Osteoporosis can result in height loss, severe back pain, deformity, impairments in a person’s ability to walk, disability, and even death.

Moreover, OP has been considered to be a disorder of postmenopausal white women; several studies investigating BMD and fracture risk in women of different ethnicities
have revealed conflicting evidence regarding osteoporosis risk. In addition, white women have higher hip fracture rates than black, Asian, and Hispanic women. Similarly, many results have indicated that Hispanic women are at higher risk for developing OP than non–Hispanic white women.

Along with an increased risk of complications, including retinopathy, nephropathy, neuropathy, and cardiovascular events, there is strong evidence for reduced BMD in children, adolescents, and adults with type 1 diabetes mellitus (T1DM), which might reduce the peak bone mass attained and increase the risk of osteoporosis and its related complications in later life. Though the relationship between T2DM and osteoporosis has been widely investigated, it remains controversial. Diabetes could influence bone through several mechanisms, some of which may have contradictory effects. Obesity, widespread in T2DM, is strongly associated with higher BMD, probably through mechanical loading and hormonal factors, including insulin, estrogen, and leptin. Hyperinsulinemia may promote bone formation. Therefore, low levels of insulin and the progression of T2DM may cause reductions in BMD. Higher glucose levels in the blood are known to interact with several proteins to generate a higher concentration of advanced glycation end-products (AGEs) in collagen that may reduce bone strength. Yamagishi et al hypothesized that AGEs in collagen may interact with bone to reduce bone strength, resulting in osteoporosis in patients with diabetes. Accumulated AGEs in the body may stimulate apoptosis of osteoblasts, thereby contributing to deficient bone formation. Another indirect effect of hyperglycemia is glycosuria, which causes hypercalcicuria, leading to decreased levels of calcium in the body and poor bone quality, hastening bone loss. There is established evidence that low levels of vitamin D are not only associated with the incidence of DM but also that altered vitamin D metabolism leads to diabetic osteopenia.

In addition, microvascular complications of diabetes lead to reduced blood flow to bone and may contribute to bone loss and fragility. Additional studies are required to determine whether DM is a leading cause of the development of osteoporosis or whether osteoporosis is aggravated by the presence of DM and should be considered as one of the long-term complications of diabetes. Thus, identifying and evaluating populations at increased risk of developing osteoporosis is critical in disease prevention and management.

In 1948, Albright and Reifenstein reported a loss of bone mass leading to osteoporosis in diabetic patients with poor glycemic control. Many clinical studies have reported that osteoporosis is one of the chronic complications associated with DM. These findings have received a great deal of attention and have been investigated by a number of researchers. Osteoporosis is a prevalent metabolic bone disease worldwide, and its occurrence in patients who have diabetes further increases their burden of disease. BMD is often used as an indication of susceptibility to osteoporosis. Bone mass is determined by the measurement of peak bone mass, which attains a maximum in the second decade of life and starts to decrease after the third decade of life. Therefore, early BMD tests are very important. In order to define osteoporosis, a group of experts convened by the World Health Organization divided BMD T-scores into the three following categories: “normal (T-score of −1.0 or higher), osteopenia (T-score between −1.0 and −2.5), or osteoporosis (T-score ≤ −2.5).” In addition to the diagnostic tools for osteoporosis (BMD test), Cummings et al suggested that it would be more sensible to focus on the risk of fracture based on a combination of factors, rather than solely relying on the diagnostic labels obtained from a T-score, since fracture risk factors are independent of BMD. For example, a woman who has a vertebral fracture has a fourfold increase in risk of another vertebral fracture regardless (or independent) of her BMD.

Risk factors that predict hip fracture independently of BMD include age, personal history of fracture, parental history of hip fracture, current cigarette smoking, low body weight, poor health, low calcium intake, low level of vitamin D, alcoholism, inadequate physical activity, and use of (or plans to use) oral corticosteroids for longer than 3 months, or serious long-term conditions thought to increase fracture risk and the risk of falling, such as hyperthyroidism, hypogonadism, diabetes mellitus, rheumatoid arthritis, malabsorption, increased levels of markers of bone resorption, and very low serum levels of estradiol.

**Methods**

Systematic reviews of full-length articles published in English from January 1950 to October 2010 were identified in PubMed, Medline, Inside Web, ISI Web of Knowledge, Science Direct, Springer Link, Ebsco Host, and other available electronic databases at Universiti Sains Malaysia Library. The following keywords were used in the search: type 2 diabetes mellitus, osteoporosis, bone, skeletal, BMD, dual X-ray absorptiometry (DXA), and quantitative ultrasound scan (QUS). More than 70 articles were identified, and those judged to be relevant by the authors were further evaluated. Clinical studies that included BMD measurements.
in patients with T2DM were extended to those using DXA and other methods. For studies that investigated BMD in T2DM, only studies that included more than 50 patients were reviewed.

**Results and discussion**

**Key findings**

Forty-seven studies were identified: seven were from the US, six from Japan, four each from Turkey, Iran, and Italy, three from China, two each from Spain, the UK, Saudi Arabia, Kuwait, and Netherlands, and one each from Denmark, Greece, Sweden, Croatia, Canada, Norway, Egypt, Korea, and Finland. In general, most of the studies showed controversy over the effect of diabetes on bone mass in T2DM. The majority of articles (26) showed increased BMD, while 13 articles revealed decreased BMD; moreover, eight articles revealed normal or no difference bone mass, as shown in Table 1.

**Osteoporosis in diabetes mellitus type 2**

Diabetes has burdened the health-care system, which is already under strain due to other chronic diseases. Uncontrolled diabetes has led to an increase in the rate of complications, and thus has doubled the cost of treating these patients. It has long been known that the alterations in bone and mineral metabolism are clinically complicated in patients with DM. Although osteopenia is an established complication of T1DM, particularly in patients with poor control who have been treated with large doses of insulin, contradictory results have been found for patients with T2DM. In addition, there have been conflicting observations concerning the incidence of osteoporosis depending on differences in sex, age, and race or the methods used to detect the increase or decrease in BMD. In fact, a reduction in bone mineral content has been observed in both T1DM and T2DM patients.

For patients with T2DM, some authors have reported an elevated BMD, other studies have reported a decreased BMD, and some have reported unaltered bone density; some cross-sectional studies have even found normal BMD. Several mechanisms have been proposed for diabetes-related osteoporosis. These include both the comorbidities of diabetes and more direct pathophysiological effects of the disease itself.

**Fractures and diabetes mellitus type 2**

The incidence of fracture was reported to be lower in T2DM patients compared to nondiabetic controls in some but not all studies. In addition, the relationship between fracture and T2DM is less clear, because many factors have an effect, such as an increase in BMD and different study designs, ages, body mass index (BMI), race, and gender, which all contribute to the conflicting results. In a study of nearly 1000 diabetic subjects, Heath and co-workers found decreased fracture rates in men and women with diabetes compared to the fracture incidence in the nondiabetic population, but they were unable to adjust for obesity, which is positively associated with T2DM and inversely associated with osteopenia. This was in contrast to the results of other studies, which was surprising given the increased fracture risk associated with T2DM. Similar study found that whilst women with T2DM had a significant risk of hip fractures, this was much lower than for the women with T1DM.

On the other hand, T2DM was previously believed to be associated with normal to increased BMD, which may be considered as an osteoprotective effect. These reports were based on the concept of BMD alone and not from prospective controlled large trials. Moreover, BMD measurement in predicting osteoporotic fractures may be limited by two main factors: decreased bone quality (which cannot be measured by DXA) and a higher risk of falls. Bone quality changes may also be affected by microvascular events common in diabetes. Patients with T2DM generally have an increased risk of falling because of peripheral neuropathy, possible hypoglycemia, nocturia, and visual impairment. In addition, many type 2 diabetic patients are obese and sedentary; coordination and balance factors that are protective in falls may be absent. Thus, patients with generally larger body size and relatively high bone mass may have higher fracture rates. Therefore, normal BMD values may be misleading. A large prospective study of older women obtained from the Study of Osteoporotic Fractures confirmed that women with type 2 diabetes experience higher fracture rates in regions of the hip, humerus, and foot than do nondiabetic women.

**Osteoporosis and gender**

Several studies found that people with T2DM have increased, normal, decreased, or no difference in bone mass compared to healthy control subjects. On the other hand, the problem of osteoporosis in men has been overlooked in the past. Although not as common as in women, hip and spine fractures in men are associated with higher morbidity and mortality than in women, and the prevalence of vertebral fractures in men is similar to or even higher than that in women.
<table>
<thead>
<tr>
<th>Country</th>
<th>Sample size/sex (F/M)*</th>
<th>Age (years)</th>
<th>Duration (years)</th>
<th>Method of BMD measurement/study design</th>
<th>Control</th>
<th>Major finding</th>
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<td>Duration</td>
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<td>BMD</td>
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<tr>
<td>Canada²⁶</td>
<td>n, 63 F nc, 99</td>
<td>78</td>
<td>8–40</td>
<td>Roentgenographic</td>
<td>Age-matched control</td>
<td>Radius</td>
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<td>77</td>
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<td></td>
<td>NR NR NR ↑</td>
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<tr>
<td>USA²⁹</td>
<td>n, 79 POM nc, 59</td>
<td>50–70</td>
<td>8.6 ± 7.2</td>
<td>Photon absorptiometry</td>
<td>Age-matched control</td>
<td>Radius</td>
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<tr>
<td>Japan²⁶</td>
<td>n, 168 (84/84) nc, 78 (27/51)³⁰</td>
<td>30–80</td>
<td>1–31</td>
<td>X-ray film/microdensitometer</td>
<td>Age-matched control</td>
<td>Metacarpal bone</td>
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<tr>
<td>Italy²⁴</td>
<td>n, 40 F nc, 35</td>
<td>42–83</td>
<td>NR</td>
<td>DPA</td>
<td>Age-matched control</td>
<td>LS</td>
</tr>
<tr>
<td>Italy²⁵</td>
<td>n, 53 (26/27)²⁹ DM1 nc, 47 (29/18)²⁹ DM2 nc, 500 (300/200)</td>
<td>40.9 ± 2.3 DM1 15.8 ± 1.3 DM1</td>
<td>15.8 ± 1.3 DM1 14.7 ± 1.1 DM2</td>
<td>SPA CS</td>
<td>Age-matched control</td>
<td>Forearm/DR</td>
</tr>
<tr>
<td>USA²⁵</td>
<td>n, 28 F nc, 207</td>
<td>26–84, mean, 59.2 ± 2.9</td>
<td>8.9 ± 1.2</td>
<td>DPA NR</td>
<td>Age- and WT-matched control</td>
<td>LS</td>
</tr>
<tr>
<td>USA²⁵</td>
<td>n, 80 (39/41)³ì nc, 381 (242/139)³ì</td>
<td>72 (55–88)</td>
<td>1–5</td>
<td>SPA/DXA CS</td>
<td>IGT and NGT</td>
<td>WRist, midradius/LS, FN</td>
</tr>
<tr>
<td>Japan²⁸</td>
<td>n, 78 (40/38)³ì</td>
<td>62/63</td>
<td>12</td>
<td>DXA NR</td>
<td>Age- and sex-matched control</td>
<td>LV</td>
</tr>
<tr>
<td>Italy²³</td>
<td>n, 110 DM2 [60(35/25)]³ì nc, 50 (26/24)³ì μl</td>
<td>60–75</td>
<td>&gt;5–15</td>
<td>DPA CS</td>
<td>Age-, sex- and BMI-matched control</td>
<td>BMC</td>
</tr>
<tr>
<td>Netherlands³³</td>
<td>n, 578 (335/243)³ì nc, 5353 (3115/2238)³ì</td>
<td>≥55</td>
<td>NR</td>
<td>DXA CPS/PS population-based study</td>
<td>Age-matched control</td>
<td>LS</td>
</tr>
<tr>
<td>Norway³⁵</td>
<td>n, 36 (15 POM/21)</td>
<td>40–65</td>
<td>3–15</td>
<td>DXA/US CS</td>
<td>Age- and sex-matched control</td>
<td>BMCL/FN, calcaneus LVL, FN, trochanter, intertrochanter, WT</td>
</tr>
<tr>
<td>Spain³⁶</td>
<td>n, 47 F nc, 252</td>
<td>61.3 ± 7</td>
<td>NR</td>
<td>DXA/QCT CS</td>
<td>Age-matched control</td>
<td>BMCL/FN, calcaneus LVL, FN, trochanter, intertrochanter, WT</td>
</tr>
<tr>
<td>Country</td>
<td>n, gender</td>
<td>Age range</td>
<td>Gender distribution</td>
<td>DXA method</td>
<td>Control group</td>
<td>T score</td>
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</tbody>
</table>
| Korea     | 185 M     | 35–74     | 0–16                | DXA CS      | Age-matched control LV Hip | NR ↑ (ve R) | NR ↑ | ↓ by 10% in M.  
> 15 y duration of DM  
↓ after 55 y age  
↓ at YSM  
↑ at FN |
| Italy     | 66 F (POM) | 35–74     | 0–16                | DXA CS      | Age-matched control LS FN | NR NR NR NR | NR ↑ | ↑ in F |
| Egypt     | 81 M      | 63.2 ± 7.4 | ≥2                  | DXA CS      | Age-matched control LS FN | NR NR NR NR | NR ↑ | ↓ in POM – T1 DM  
↑ in POM – DM2 |
| Denmark   | 56 M      | 52.9 ± 6.16 | 2                  | DXA CS      | Age-matched control LS PF | NR NR NR NR | NR ↑ | ↓ in DM1  
↑ in DM2 |
| Finland   | 56 M      | (61.7 ± 6.3) | dx after 30 y       | DXA CS      | Age-matched control PF NR | NR /DM2 ↑ in DM1 NR | NR ↑ | ↓ in DM1  
↑ in DM2 |
| Japan     | 104 M     | 54.0 ± 1.0 | 5.6 ± 0.6           | CXD NR      | Age-matched control Metacarpal NR ↔ ↑ R P | NR ↑ | ↑ | ↓ |
| Turkey    | 161 F (POM) | 54.3 ± 1.3 | ≥2                  | DXA NR      | Age-matched control LS FN | NR NR NR NR | NR ↑ | ↑ in F |
| USA       | 80 M      | (64/61) | At least 3          | DXA/QCT NR  | Age-matched control Z-score NR | NR NR NR NR | NR ↑ | ↓ in M  
↑ in FMD1 6.8% in woman; 7.6% lower in men |
| USA       | 600 M     | 54.8 ± 12.5 F  | New dx and older   | DXA CS      | Age-matched control LS Hip FN | ↔ ↔ ↔ ↔ | NR ↑ | ↑ by MN 3.7%  
FN in F |
| Turkey    | 277 M      | 6.5 ± 5.3  | 2                  | DXA PS/CS   | Age-matched control LS FN Trochanter WT | ↔ ↔ ↔ ↔ | NR ↑ | ↑ in F/M (51–60 y)  
↓ LS in M  
↔ at WT/tronchanter (more in LS than FN)  
ON 43.6%  
OP 46.8%  
14% low vit D |
| Saudi Arabia | 104 POM | 55–70     | 6–30                | DXA PS      | Age-matched control LS FN | NR ↔ ↔ NR | NR ↑ | ↓ by 10% in M.  
> 15 y duration of DM  
↓ after 55 y age  
↓ at YSM  
↑ at FN |

(Continued)
Table 1 (Continued)

<table>
<thead>
<tr>
<th>Country</th>
<th>Sample size/sex (F/M)*</th>
<th>Age (years)</th>
<th>Duration (years)</th>
<th>Method of BMD measurement/study design</th>
<th>Control</th>
<th>Major finding</th>
</tr>
</thead>
<tbody>
<tr>
<td>USA</td>
<td>n, 566 (243/323)*</td>
<td>70–79</td>
<td>11 F/10 M (&lt;5–20)</td>
<td>DXA/spine QCT CS/PS cohort</td>
<td>Age-matched control</td>
<td>LS ↔ ↓ NR ↑ 4%–5% regardless of sex and race</td>
</tr>
<tr>
<td>Spain</td>
<td>n, 92 (56/36)*</td>
<td>40–81</td>
<td>1–40 (10 ± 8 y)</td>
<td>DXA NR CS</td>
<td>Age-matched control</td>
<td>Calcaneus ↔ NR NR ↑ 8% in F</td>
</tr>
<tr>
<td>UK</td>
<td>n, 32/33 DM2</td>
<td>59–71</td>
<td>New dx</td>
<td>DXA CS</td>
<td>Age-matched control</td>
<td>FN ↔ ↔ ↔ ↑ in F</td>
</tr>
<tr>
<td>Iran</td>
<td>n, 146 (20 PRM/124 POM)</td>
<td>40–81</td>
<td>DXA historical cohort study</td>
<td>Age- and BMI-matched control</td>
<td>Age- and BMI-matched control</td>
<td>LS NR NR NR ↔ in PRM ↑ LS in POM</td>
</tr>
<tr>
<td>Netherlands</td>
<td>n, 792 (483/309)*</td>
<td>73.8 ± 9.2</td>
<td>dx &gt; 30</td>
<td>DXA PS</td>
<td>Age- and gender-matched control</td>
<td>LS NR NR NR ↑ fracture risk by 1.69-fold</td>
</tr>
<tr>
<td>Japan</td>
<td>n, 145 (81/64)*</td>
<td>(67/63)*</td>
<td>&gt;2 y</td>
<td>DXA CS</td>
<td>Age-matched control</td>
<td>LS ↑ NR NR ↑ DR</td>
</tr>
<tr>
<td>Sweden</td>
<td>n, 67 F</td>
<td>75</td>
<td>9.8 (0.04–61)</td>
<td>DXA/US NR</td>
<td>Age- and gender-matched control</td>
<td>LS NR ↔ ↓ NP</td>
</tr>
<tr>
<td>Greece</td>
<td>n, 40 D-EMPDM2</td>
<td>58.7 ± 5</td>
<td>13.9 ± 3.9</td>
<td>DXA NR</td>
<td>Age- and BMI-matched control</td>
<td>LV ↑ ↔ NR ↑ LV BMD in D-EMP</td>
</tr>
<tr>
<td>UK</td>
<td>n, 35 DM1 M</td>
<td>49.3 DM1</td>
<td>19.7 DM1</td>
<td>DXA CS/controlled study</td>
<td>Age-matched control</td>
<td>DR ↔ ↔ ↔ ↑ in EMP</td>
</tr>
<tr>
<td>Iran</td>
<td>n, 40 MW</td>
<td>POM</td>
<td>38.5 control</td>
<td>DXA Case-control study</td>
<td>Age-, BMI-, and MPL-matched control</td>
<td>LS FN ↑ NR ↑ FNL Normal</td>
</tr>
<tr>
<td>Saudi Arabia</td>
<td>n, 154 M</td>
<td>50–76</td>
<td>DXA NR</td>
<td>Age-matched control</td>
<td>Age-matched control</td>
<td>LS Hip ↔ NR NR</td>
</tr>
</tbody>
</table>

*Note: *F/M = female/male; DM = diabetes mellitus; PRM = prediabetes risk; POM = prediabetes; D-EMP = diabetes; D-NMP = prediabetes risk; DM2 = diabetes mellitus type II; iFG = impaired fasting glucose; NS = not significant; CS = case-control study; PS = population study; TBD = treatment breakdown data; NL = Netherlands; DM1 = diabetes mellitus type I; LS = lumbar spine; FN = femur neck; PF = proximal femur; DP = distal phosphorous; DR = distal radius; NP = normal; HbA1c = hemoglobin A1c; ↑ = increase; ↓ = decrease.
<table>
<thead>
<tr>
<th>Country</th>
<th>n</th>
<th>Gender</th>
<th>Age (years)</th>
<th>Sex</th>
<th>BMI (kg/m²)</th>
<th>Method</th>
<th>Control</th>
<th>Trochanter</th>
<th>Metacarpal</th>
<th>Femoral Shaft</th>
<th>Radius</th>
<th>Hip</th>
<th>Calcaneus</th>
<th>Total Hip</th>
<th>OP/ON (%)</th>
<th>OP/ON (%)</th>
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<tbody>
<tr>
<td>Turkey</td>
<td>52 (37/15)*</td>
<td>n, 48 (34/14)*</td>
<td>41–64</td>
<td>0–20</td>
<td>Age-, sex-, and BMI-matched control</td>
<td>Forearm</td>
<td>NR</td>
<td>NR</td>
<td>NR</td>
<td>↑ FN</td>
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<tr>
<td>China</td>
<td>131 M</td>
<td>n, 131 M</td>
<td>73.12 ± 5.54</td>
<td>5.28 ± 3.56</td>
<td>DXA</td>
<td>NR</td>
<td>FN</td>
<td>LS</td>
<td>FT</td>
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<tr>
<td>Iran</td>
<td>146 F</td>
<td>n, 146 F</td>
<td>59.3/48.3</td>
<td>6.18/0.68</td>
<td>DXA</td>
<td>Age- and BMI-matched control</td>
<td>LS</td>
<td>NR</td>
<td>NR</td>
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<tr>
<td>Japan</td>
<td>151 M</td>
<td>n, 151 M</td>
<td>55.4 ± 1.2</td>
<td>5.8 ± 1.2</td>
<td>DXA</td>
<td>Age- and BMI-matched control</td>
<td>Trochanter</td>
<td>↔</td>
<td>NR</td>
<td>NR</td>
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<tr>
<td>USA</td>
<td>4929 (2505/2424)*</td>
<td>n, 3967 (1993/1974)*</td>
<td>50–79</td>
<td>NR</td>
<td>DXA</td>
<td>Age-matched control</td>
<td>Trochanter</td>
<td>↔</td>
<td>NR</td>
<td>NR</td>
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<tr>
<td>Kuwait</td>
<td>210 F</td>
<td>n, 210 F</td>
<td>59</td>
<td>11.99 ± 8.6</td>
<td>DXA</td>
<td>Age-matched control</td>
<td>LS</td>
<td>NR</td>
<td>←ve R</td>
<td>NR</td>
<td>↔</td>
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<tr>
<td>Iran</td>
<td>242 POM</td>
<td>n, 242 POM</td>
<td>53.6 ± 10.59</td>
<td>5.9 ± 3.7</td>
<td>DXA</td>
<td>Age- and BMI-matched control</td>
<td>LS</td>
<td>NR</td>
<td>NR</td>
<td>NR</td>
<td>↑</td>
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</tr>
<tr>
<td>Croatia</td>
<td>130 POM</td>
<td>n, 130 POM</td>
<td>67 (45–80)</td>
<td>NR</td>
<td>DXA</td>
<td>Age-matched control</td>
<td>LS</td>
<td>NR</td>
<td>NR</td>
<td>NR</td>
<td>↑</td>
<td></td>
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<tr>
<td>Japan</td>
<td>164</td>
<td>n, 196 POM</td>
<td>69.7 ± 7.1</td>
<td>14.6 ± 10.7</td>
<td>QUS</td>
<td>Age-matched control</td>
<td>Calcaneus</td>
<td>↔</td>
<td>↔</td>
<td>↓ with NP</td>
<td>↑</td>
<td></td>
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<tr>
<td>Kuwait</td>
<td>122 PRM</td>
<td>DM2 n, 122 PRM</td>
<td>26–50</td>
<td>NR</td>
<td>DXA</td>
<td>Age-, gender-, and BMI-matched control</td>
<td>LS</td>
<td>NR</td>
<td>NR</td>
<td>NR</td>
<td>↑ LS</td>
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<tr>
<td>China</td>
<td>1042 POM</td>
<td>n, 919</td>
<td>62.2 ± 7.15</td>
<td>6.6 ± 6.09</td>
<td>DXA</td>
<td>Age-matched control</td>
<td>LS/FN</td>
<td>NR</td>
<td>NR</td>
<td>NR</td>
<td>↑ LS</td>
<td></td>
<td></td>
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<tr>
<td>Turkey</td>
<td>206 POM</td>
<td>n, 61</td>
<td>44–88</td>
<td>10.3 ± 8.2</td>
<td>DXA</td>
<td>Age-matched control</td>
<td>LS</td>
<td>NR</td>
<td>↑</td>
<td>←ve R</td>
<td>–</td>
<td>↔</td>
<td></td>
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</tr>
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</table>

(Continued)
Table 1 (Continued)

<table>
<thead>
<tr>
<th>Country</th>
<th>Sample size/sex (F/M)*</th>
<th>Age (years)</th>
<th>Duration (years)</th>
<th>Method of BMD measurement/study design</th>
<th>Control</th>
<th>Method of BMD</th>
<th>Major finding</th>
</tr>
</thead>
<tbody>
<tr>
<td>China</td>
<td>n=190 (42M, 148F)</td>
<td>&gt;50</td>
<td>NR</td>
<td>DXA</td>
<td>NR</td>
<td>CN</td>
<td>↑ in non-obese DM2</td>
</tr>
</tbody>
</table>
|         | (58.5/56.1)            |             | NR               | CS                                    | FN      | LS            | ↔ in women | ±
|         | (1837)                 |             |                  | PS                                    | Fn      | FN            | ↓ in men   |

Abbreviations: BMC, body mineral content; BMD, bone mass density; BMi, body mass index; CS, cross-sectional study; CN, comparative study; DXA, dual X-ray absorptiometry; LS, lumbar spine; PN, premenopausal women; FN, postmenopausal women; DM1, diabetes mellitus type 1; DM2, diabetes mellitus type 2; D-EMP, diabetic women with premature menopause; D-NMP, diabetic women with normal menopause; DXA, dual X-ray absorptiometry; DR, distal radius; dx, diagnosis; DPA, dual photon absorptiometry; EMP, prematurely menopausal women; FT, femoral trochanter; F/M, female/male; FS, prospective study; FN, femoral neck; FT, femoral trochanter; FF, femoral fracture risk; F, female; iFG, impaired fasting glucose; iGT, impaired glucose tolerance; LS, lumbar vertebrae; M, male; Mw, menopausal women; OPP, osteoporosis; ON, osteopenia; PRM, premenopausal women; POM, postmenopausal women; PS, prospective study; PF, proximal femoral; QCT, quantitative computerized tomography; RP, retinopathy; SPA, single photon absorptiometry; TBD, total bone density; wT, weight; US, ultrasound; y, years; YSM, years since menopause; →, positive correlation with BMD; ↔, no association.

Three studies investigated the decrease in bone mass in men; two studies were performed using computed X-ray densitometry at the metacarpus bone in 151 and 104 male T2DM patients aged 55.84,102 The third study assessed BMD measured by DXA in 131 elderly T2DM males in China. The findings from this study also suggested that decreased BMD and poor glycemic control were correlated with osteoporosis.103 In contrast, Bridges et al assessed the BMD in 35 and 90 men with T1DM and T2DM, respectively, compared to 50 control subjects. This study found no difference between bone mass and diabetic status. In addition, no correlation was found between BMD and the glycosylated hemoglobin concentration, disease duration or the presence of microvascular or macrovascular complications in either of the diabetic groups.104

On the other hand, three studies reported a lower incidence of osteopenia and increased bone mass in older women with diabetes. In 1967, Meema and Meema reported a significantly increased cortical thickness of the radial bone in 63 elderly women with diabetes compared to 133 women without diabetes. This difference persisted after adjusting for body weight, and the authors postulated that diabetes was an “antiosteoporotic condition.”79

In 1985, Johnston and colleagues reported a significantly greater mid-radial bone mass in 79 postmenopausal women with T2DM (68% of whom were being treated with insulin) compared to 59 in the control group. The calculated rate of bone loss was about half that expected and was not explained by obesity.79 In 1989, Weinstock et al reported a nonsignificantly higher BMD in 28 T2DM women with mean age of 59 years compared to 207 age-matched volunteers.73 Thus, women with T2DM are not at an increased risk of diminished BMD, and may even be protected against bone loss. Relatively little attention has been paid to these remarkable findings. Since these studies were limited to women, sex differences were not described, and also the sample size was small. A similar finding was reported for 47 elderly T2DM women where the BMD was measured using two different methods (DXA and quantitative computed tomography), where no evidence was found to show that T2DM produces any change in bone metabolism or mass.84 Moreover, a population-based family study of Mexican Americans (600 subjects from 34 families) confirmed that diabetic women aged 55 years have a higher BMD compared to their nondiabetic counterparts.46 In addition, two studies assessed the association of T2DM with BMD in older patients (aged ≥55 years), and both studies found an increase in BMD.75,89 In one population-based Rotterdam
Study with a sample size of 5931, it was shown that both men and women with T2DM had an increased BMD, and that in women this was associated with a lower frequency of non-vertebral fractures. A similar finding related to increased BMD found in white diabetic women, whereas no differences in bone density according to diabetic status were observed in men, and the sex differences were explained by the greater androgenicity reported in women with hyperglycemic and hyperinsulinemic conditions.

On the other hand, menopause is one of the most important risk factors for osteoporosis in women and is characterized by rapid bone loss in newly postmenopausal women. The loss of BMD is accelerated after the cessation of ovarian secretions and estrogen deficiency is an important factor causing osteoclast activation. In our survey of the scientific literature related to the association between T2DM and bone mass, we identified 16 studies out of 47 that assessed bone mass in postmenopausal T2DM; the majority (13) of studies showed increased BMD and lower osteoporosis risk, whereas two studies showed decreased BMD and one study showed no difference in BMD when compared with normal subjects, as shown in Table 1.

**Osteoporosis and glycemic control**

Most of the studies were performed on T2DM patients under fairly controlled and stable conditions; in one of these studies, the mean glycosylated hemoglobin was 6.7%. Therefore, it is as yet unknown whether changes in glycemic control influence bone turnover in T2DM patients, although some papers have reported that BMD decreased more severely in patients with poorly controlled type 2 diabetes. In 1997, one study assessed BMD before and after glycemic control for 3 weeks in 78 patients with poorly controlled T2DM (age 28–73 years) with an initial glycosylated hemoglobin level of more than 8%, and found that metabolic improvements in poorly controlled T2DM decreased bone loss within a short period. Thus, glycemic control might protect T2DM patients from bone loss. On the other hand, some studies have reported that insulin level has a vital effect on bone mass. Japanese T2DMs have the clinical feature of low insulin secretion, which may predispose them to the risk of lower BMD. In 2005, Majima et al assessed the association between T2DM, BMD, metabolic control, and insulin-secretion capacity in 145 elderly Japanese diabetic patients compared to 95 subjects in the control group who were of a similar age. This study indicated that there was loss in cortical bone and positive correlation between the levels of insulin secretion and BMD at different sites (lumbar spine, femoral neck, distal radius); in addition, maintaining good metabolic control was key in preventing bone loss in T2DM. In contrast, Oz et al, in 2006, found an association between T2DM and higher BMD in 52 diabetic men and women aged 41–64 years compared to 48 nondiabetic control subjects. These findings suggest that although bone mass formation is lower in T2DM, diabetic patients are not susceptible to bone resorption. This low bone turnover can slow the rate of bone loss and cause a higher BMD than would be expected for a certain age.

**Osteoporosis and body mass index**

Another important factor that affects BMD and confuses findings is the BMI, in addition to heredity, nutritional dietary customs, height, and lean mass. A low BMI is associated with decreased BMD, the increased possibility of osteoporosis, and the risk of fracture. A meta-analysis demonstrated that BMI is also an important predictor of BMD in T2DM. Overweight and obesity are believed to be protective factors of BMD. Compared with Caucasian populations, diabetic women in Asia are relatively shorter, a lower percentage are overweight, and they have a lower insulin sensitivity. Furthermore, BMI is a powerful and modifiable risk factor for both DM and OP. However, the impact of BMI with racial/ethnic disparities in overweight and obesity (measured by BMI) is well documented in many studies. BMI disparities are more pronounced and consistent among women, age, education, existence of morbid conditions, and marital status. Seo and Torabi examined the effect of race/ethnicity on BMI among US adults by gender, adjusting the effects of age, education, serious morbidity, and marital status. There was no evidence of decrease in the prevalence of overweight or BMI and diabetes among US adults.

However, in Western countries, a BMI $\geq 27$ kg/m$^2$ is often used to define obesity. Using this definition, 33% of US adults are considered to be overweight, while in Hong Kong Chinese of working age, only 11.6% have a BMI $\geq 27$ kg/m$^2$. If BMI $\geq 30$ kg/m$^2$ is used to define obesity, 8%–15% of Caucasians but only 2.2%–4.8% of Hong Kong Chinese will be considered to be obese. Similarly, the mean BMI in UK subjects is 26.0 kg/m$^2$ for men and 26.3 kg/m$^2$ for women. This is compared to 23.4 kg/m$^2$ and 23.3 kg/m$^2$ in Hong Kong Chinese men and women, respectively. In addition, Asians, although mean BMI is lower, have a higher percentage body fat and more upper-body subcutaneous fat. A recent study of more than 13,000 Chinese men and women...
with significantly lower mean BMI (21 kg/m²) found the percentage body fat was inversely related to BMD as measured by DXA at the spine, hip, and total body.\textsuperscript{144} In a meta-analysis that included data for three Asian groups, Deurenberg et al found that the percentage body fat was higher than predicted at low BMI levels for Chinese. Body fat was underestimated across all BMI levels for Thais and Indonesians.\textsuperscript{145} In a similar study of women in Hawaii, Novotny et al found that Asian women had a greater percentage of body fat than did white women with the same BMI.\textsuperscript{146} The limited data concerning the correlation between BMI and adiposity suggest that health effects of BMI may differ as US-born Asian Americans are significantly more likely to be obese or overweight than the foreign-born Asian Americans.\textsuperscript{127} These conflicting results suggest a complex relationship between fat mass and bone mass is likely depending on the patient’s age, sex and ethnicity.\textsuperscript{147} To date, relatively few studies have examined this relationship.\textsuperscript{148}

**Conclusion**

With progressive aging of the population, there will be a huge increase in the prevalence of osteoporosis, which is considered to be one of the most common public health problems in the world. However, the impact of diabetes disease on osteoporosis has not yet been carefully considered. Therefore, there is a need for further longitudinal studies, including the incidence and risk factors for osteoporotic fractures. In clinical routine, the extent of diagnostic and therapeutic activities in patients with T2DM in respect to generalized bone disease or diabetic osteopenia should be based on individual conditions and risk profile for osteoporosis. In addition, osteoprotective behavior must be assessed, as low educational levels about OP may put those populations at high risk of fractures. Therefore, patient education needs to highlight that weight-bearing exercise and consuming calcium-rich foods not only prevent osteoporosis but also can decrease BMI, reduce blood pressure, and improve lipid and diabetic control, which are often significant risk comorbidities in diabetic patients. Therefore, the routine screening of risk factors and patient education about bone density evaluation are important and should be recommended to all diabetic patients. Also, all patients with diabetes, and particularly those with high risk of fractures, should be given general information regarding proper home-safety measures to reduce risk of falling, including the wearing of hip protectors, especially in the elderly. Future studies need to be prospective, controlled with large sample sizes, and include evaluations of bone quality at different sites (not just bone size), BMI, the duration of diabetes, race, glycemic control, fractures, complications, and BMD, in order to identify the possible relationship between diabetes and osteoporosis.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**

Osteoporosis and type 2 diabetes mellitus


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