



DXA parameters, Trabecular Bone Score (TBS) and Bone Mineral Density (BMD), in fracture risk prediction in endocrine-mediated secondary osteoporosis

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Abstract

Osteoporosis, a disease characterized by low bone mass and alterations of bone microarchitecture, leading to an increased risk for fragility fractures and, eventually, to fracture; is associated with an excess of mortality, a decrease in quality of life, and co-morbidities. Bone mineral density (BMD), measured by dual X-ray absorptiometry (DXA), has been the gold standard for the diagnosis of osteoporosis. Trabecular bone score (TBS), a textural analysis of the lumbar spine DXA images, is an index of bone microarchitecture. TBS has been robustly shown to predict fractures independently of BMD. In this review, while reporting also results on BMD, we mainly focus on the TBS role in the assessment of bone health in endocrine disorders known to be reflected in bone.

Keywords Bone mineral density · Dual-energy X-ray absorptiometry · Trabecular bone score · Secondary osteoporosis · Diabetes mellitus · Endocrine disorders

Background

Osteoporosis is a common disease characterized by low bone mass (i.e. quantity) and altered bone microarchitecture (i.e. quality), resulting in decreased bone strength with an increased risk of fractures [1]. It is associated with an excess of mortality, a decrease in quality of life, and co-morbidities; and has a high social and economic burden, representing a major public health problem. Osteoporosis affects mostly postmenopausal women and fewer men. It is mainly related to the normal aging process— primary osteoporosis.

Nevertheless, it can also occur due to secondary causes such as bone-affecting treatments, clinical disorders, or lifestyle habits—secondary osteoporosis [2]. Nine million osteoporotic fractures occur worldwide yearly [3]. The accurate identification of fracture risk, followed by relevant management, would reduce these numbers and the associated costs [4].

Bone mineral density (BMD), measured by dual X-ray absorptiometry (DXA), has been the gold standard for osteoporosis diagnosis in the absence of established fragility fractures [5]. BMD, a bone quantity parameter, is a major determinant of bone strength and fracture risk. Nevertheless, a considerable overlap (up to 45%) exists in BMD values between individuals who develop fractures and those who do not, suggesting that fracture risk prediction based solely on BMD is suboptimal [6]. Thus, the development of bone quality assessments has gained special interest to address this gap. Trabecular bone score (TBS) is one of the most widely used assessments of bone quality [7]. Both BMD and TBS are independent predictors of fragility fractures and are the two pillars of the World Health Organization's (WHO) clinical definition of osteoporosis [1]. In this article, we review the role of DXA-derived parameters, BMD and TBS, with a special focus on TBS, in

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fracture risk prediction in endocrine-mediated secondary osteoporosis.

DXA: the gold standard for BMD assessment

DXA is the most widely used technique and the gold standard for osteoporosis diagnosis and management [8–10]. It is a simple, quick, painless, and safe examination, which uses very low doses of dual X-rays (considerably lower than those of an X-ray and a CT scan), hence it can be repeated safely over time. DXA acquisition can provide images for total body, hip, posterior–anterior (PA) lumbar spine (LS), and/or forearm. BMD (expressed in g/m^2) is measured in well-defined regions of interest in these images. LS, total hip, femoral neck, and/or 1/3 radius BMD values are considered for the osteoporosis diagnosis and/or management. The decision on the use of each of them depends on factors, such as sex and age of the patient. For example, the PA LS is the preferred region for the BMD assessment in women up to 60 years and in men up to 65; the femoral neck and total hip are the chosen sites for BMD assessment in older individuals and/or in the presence of discrepancies or degenerations in the LS. Forearm (1/3 radius) is a backup site taken into consideration when the main sites (PA LS, total hip, and femoral neck) are impacted by certain health conditions such as hyperparathyroidism or obesity (exceeding the weight limit of the DXA device's table).

The quantitative assessment of BMD is compared with the average BMD value of a population of healthy young adults in the age of peak bone mass (young adult reference population). The number of standard deviations that the BMD differs from the BMD of the young adult reference population is indicated by the term BMD T-score. The WHO classification of osteoporosis is based on the lowest BMD T-score value from PA LS, total hip or femoral neck (or 1/3 radius for the specified scenarios): normal: BMD T-score greater or equal than -1 ; osteopenia: BMD T-score between -1 and -2.5 ; osteoporosis: BMD T-score equal or less than -2.5 . This classification applies only to the BMD assessed by DXA. Extensive evidence supports the use of DXA BMD for osteoporosis diagnosis, prognosis, and follow-up, for fracture risk stratification and treatment initiation decision-making [11]. The importance of BMD testing, the skeletal sites to measure it, its report and interpretation, and the follow-up intervals between measurements, have been well-defined in several guidelines and taught worldwide by a combined course put together (Osteoporosis Essentials[®]) by the International Osteoporosis Foundation (IOF) and the International Society of Clinical Densitometry (ISCD) [12, 13].

BMD is a measurement of high reliability and precision, able to predict osteoporotic fractures well in advance, and

useful in osteoporotic patients' follow-up. Black et al. demonstrated that BMD and history of nonvertebral fracture could predict fractures in a period as long as 20–25 years in a large cohort of postmenopausal women [14]. The most widely used tool for fracture prediction, FRAX, provides the fracture probability for a period of 10 years [15]. An extended DXA measurement of the hip diaphysis is recommended in patients on long-term treatment with bisphosphonates in order to detect early signs of atypical femur fractures [16]. ISCD recommends the use of BMD as assessed by DXA for antiosteoporotic treatment follow-up [17].

An acknowledged drawback of DXA is its limited capability to assess bone health in type 2 diabetes mellitus (T2DM) patients [18–20]. Although bone fragility is a known complication of diabetes, T2DM patients have normal or higher BMD levels as compared to non-diabetics. Napoli et al. [21] thoroughly describe the possible mechanisms that could explain this association: *in vivo* and *ex vivo* studies have shown the anabolic effect of insulin on osteoblasts, thus hyperinsulinemia in patients with T2DM might explain the high BMD levels; also, sclerostin levels are higher among T2DM and they are positively associated with BMD. Despite the normal or higher BMD values in diabetics as compared to non-diabetics, several studies [22–24] have shown that low BMD values are associated with a higher risk of fracture in diabetics as compared to non-diabetics. Moreover, BMD is useful at antiosteoporotic treatment follow-up in diabetic patients [22–25]. Other limitations of DXA are the overestimation of LS BMD as a consequence of degenerative changes in the spine or aortic calcification presence, and the influence of BMD by BMI or regional soft tissue presence [26, 27]. Armameto-Villareal et al. [28] has recently shown that weight loss among obese individuals is associated with bone loss as indicated by BMD values. However, BMD was negatively associated with the presence of fat tissue, indicating that the increase in body fat may increase systemic inflammation leading to frailty and poor bone quality. To address the increase in bone loss and eventual fracture risk in obese undergoing weight loss therapy, Armameto-Villareal et al. suggest a combination of aerobic and resistance exercise and calcium and vitamin D supplements. Other bone health assessments using bone density derived from high-resolution peripheral quantitative computed tomography (HR-pQCT) or other bone strength surrogates (such as bone turnover markers or advanced glycation end-products, etc.) in obese and diabetics have been shown useful at their overall bone health evaluation [29–31].

Major improvements in DXA technologies over the years have enabled the assessment of bone health parameters other than BMD. For example, the improved spatial resolution (as low as 250 micrometers for some of DXA

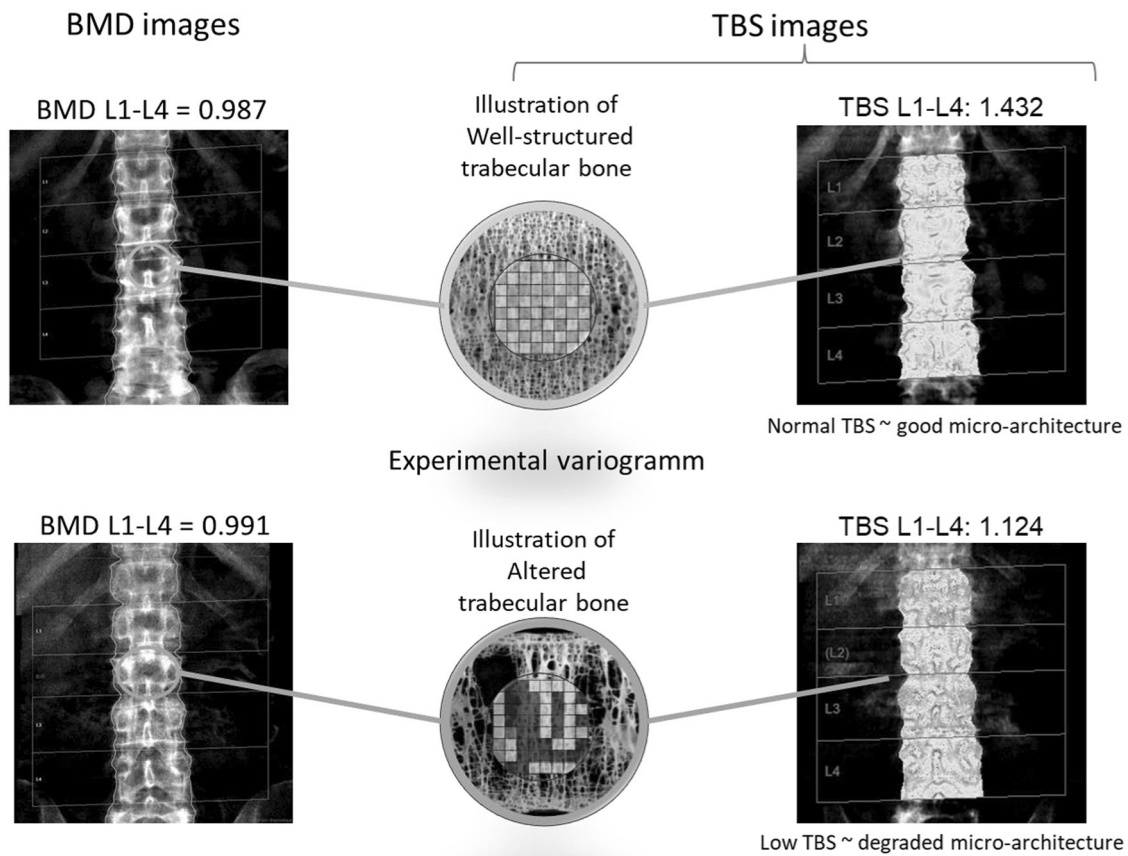


Fig. 1 Concept of TBS. Two different patients with equivalent bone mineral density (BMD) but different trabecular bone score (TBS)

devices) allows the search for vertebral fractures (vertebral fracture assessment—VFA) and the assessment of trabecular bone texture (TBS) (further elaborated below). Different algorithms can also estimate structural parameters of hip geometry, such as femoral strength index, axis length, cross-sectional area, etc. (hip structural analysis—HSA) [32–35].

DXA: beyond bone health assessment

Besides bone, DXA assesses parameters of muscle and fat [36]. Several authors reported the direct and indirect role of DXA in the follow-up of malabsorption conditions [37–40]. In addition, recently, emerged more and more the key concept of the “modified functional muscle–bone unit”, in which BMD is strongly associated with either mass and quality of muscles, which are further correlated with fracture risk [41]. In this sense, DXA has been demonstrated useful in providing a precise assessment of the skeletal muscle mass, which is crucial in disorders such as

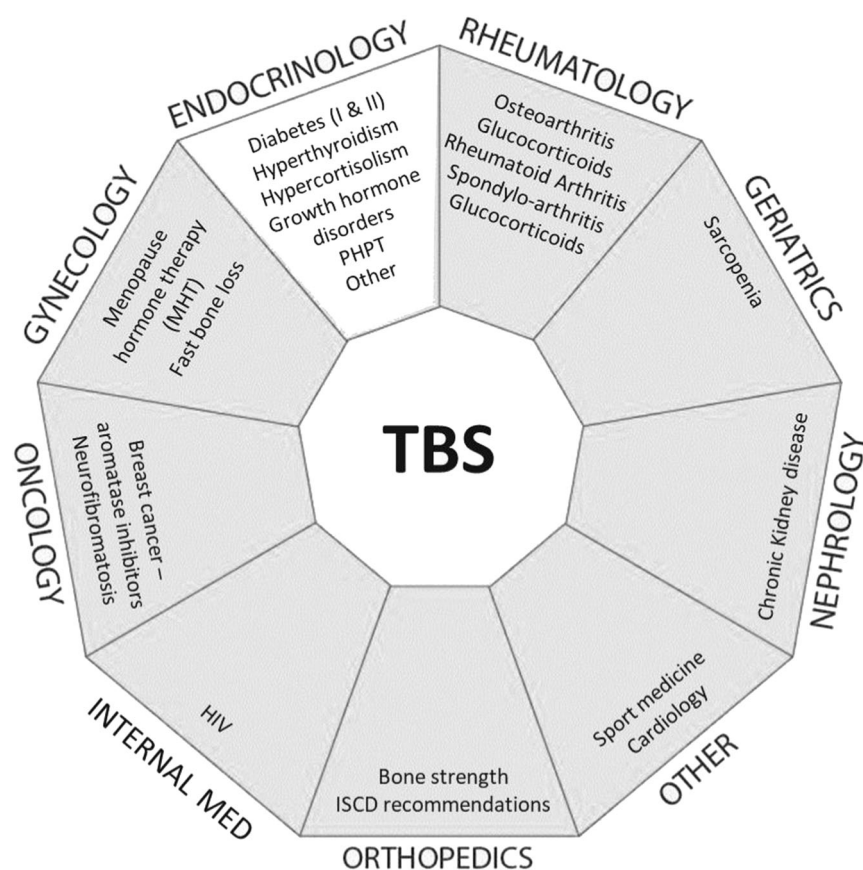
sarcopenia or geriatric syndrome [42, 43]. Other parameters of body composition, indicating the presence and distribution of fat mass in different regions of the body are essential in the assessment and follow-up of health disorders associated with fat presence [44–48].

An interesting recent study demonstrated that during weight-loss programs, BMI can not specify the fat distribution, with limited use of this datum; in contrast with DXA report, which is able to quantify the metabolic redistribution of total and regional fat mass and visceral adipose tissue [49]. Another study compared the accuracy of six different osteoporosis risk assessment tools in a restricted group of women in Kuala Lumpur, further demonstrating the importance of DXA in such context [50].

Definition of TBS and its clinical validation/implication

TBS is a textural analysis resulting from a computed evaluation of pixel gray-level variations in previously obtained

Fig. 2 Summary of pathologies (per medical specialty) in which TBS has demonstrated added clinical value (published studies only)—adapted to this paper needs—Courtesy of Medimaps Group SA (Switzerland)



LS DXA images. It is an index of bone microarchitecture correlated with parameters of bone strength [51–53]. Higher values of TBS indicate a better microarchitecture, whereas lower values indicate a degraded microarchitecture (Fig. 1). Studies have robustly shown that TBS predicts fractures independently of BMD and other clinical risk factors for fracture. The added value of the TBS to BMD in fracture risk assessment has been extensively documented in cross-sectional, prospective, and longitudinal studies [54] and endorsed by medical societies of bone field (IOF, the European Society for Clinical and Economic Aspects of Osteoporosis and Osteoarthritis, and the ISCD) [55]. More recently, for regions where intervention guidelines are based solely on BMD T-score, an alternative approach for using TBS in clinical practice based upon a “risk-equivalent” offset adjustment to BMD T-score has been developed [56]. The early validations of this approach illustrate how TBS contributes to vertebral fracture risk assessment by increasing the specificity of the model (by about 20%) without compromising the sensitivity.

Several studies have also demonstrated the ability of TBS to predict fragility fractures in secondary osteoporosis caused by diabetes, PHPT, rheumatoid arthritis, adrenal incidentaloma, chronic kidney disease, long-term glucocorticoid therapy, HIV, or oncological conditions (Fig. 2) [2].

TBS in endocrine-mediated secondary osteoporosis

TBS and diabetes mellitus

It is well established that the skeleton is one of the organs affected by diabetes mellitus type 1 and 2, listing diabetes as one of the risk factors for fragility fractures [57]. Despite the high fracture risk in diabetics, their BMD is generally higher compared to non-diabetics. Thus, it is hypothesized that diabetes may be associated with a reduction of bone strength that is not reflected in the measurement of BMD alone. Robust evidence has shown that—differently than BMD—TBS is lower in diabetics than non-diabetics [16, 17, 19, 57]. In 2013, Leslie et al. were the first to report this finding in a study of 29,000 women, of which 2,356 had diabetes [58]. Moreover, it has been confirmed that TBS is negatively related to levels of HbA1c (e.g. bone structure remained normal when HbA1c was lower than 7.5), fasting plasma glucose, and fasting insulin [59, 60]. Currently, several studies conducted in more than 40,508 (4,269 diabetics) individuals altogether have robustly shown that TBS is lower in diabetics than in controls, whereas BMD is higher in diabetics than in controls [58–70]. Furthermore, TBS outperforms BMD in fracture

discrimination and prediction in diabetics. A recent meta-analysis of 7,819 women and men also showed that type 2 diabetes was associated with decreased TBS in fully adjusted models; and that compared to controls, also pre-diabetics had significantly lower TBS [71]. Overall, combining TBS and BMD incrementally improves fracture prediction. Nevertheless, the increased fracture risk is not entirely explained by the TBS difference between the diabetics and non-diabetics: diabetes is associated with a 32% increase in the fracture risk; whereas one standard deviation decrease in TBS is associated with 1.27-fold (95% CI 1.10–1.46) increase in the risk of fracture in diabetics [72]. The mechanism remains unclear. A hypothesis to explain the association of TBS with diabetes is that advanced glycation end products could mediate this association [61]. Nevertheless, it necessitates further investigation to be proven true.

TBS and growth hormone (GH) disorders

GH is an important regulator of bone growth; its effects are important to maintain bone mass [73]. *Growth hormone deficiency* (GHD) is related to reduced bone strength and the GH long-term replacement therapy can successfully revert this condition [68]. The effect of GH treatment on bone quality is unclear. Kuřma et al. showed that bone quality, as assessed by TBS, improved only in the GHD cases with sufficient vitamin D levels. However, an insignificant decrease in TBS was seen after 7 years of GH treatment [74]. *Acromegaly* is a rare disease characterized by excessive GH and insulin-like growth factor 1 (IGF-1), caused mainly by GH-producing pituitary adenoma [75]. Evidence shows that patients with acromegaly are predisposed to develop fractures regardless of their BMD values. To explain the high incidence of fractures in these patients, TBS may be useful in assessing skeletal fragility, given BMD is unable to accurately assess bone strength. Previous studies suggested that GH excess may positively affect cortical bone and negatively affect trabecular bone [74]. Currently, there are conflicting BMD results in acromegaly, showing mainly no effect or increase. Further, acromegaly treatment had different effects on TBS and BMD: TBS decreased significantly, whereas BMD increased [74].

TBS in hyperparathyroidism

Primary hyperparathyroidism (PHPT) is an endocrine disorder characterized by elevated parathyroid hormone levels with hypercalcemia. Patients with PHPT exhibit increased fracture risk [76]. Several studies have reported that PHPT cases have decreased TBS, indicating deteriorated bone microarchitecture. TBS is shown to be a predictor of fracture independent of BMD in PHPT patients. Also, the

fractured patients with PHPT had lower TBS than the non-fractured ones [77–80]. Moreover, the TBS values significantly improved after parathyroidectomy compared to after conservative management of PHPT [81].

TBS in hyperthyroidism

Thyroid hormone stimulates bone resorption. Hyperthyroidism is associated with bone loss and increased risk for fracture. A negative association between bone parameters and the fT4 level has been reported by Hwangbo et al. [82]. Moreover, TBS was studied in thyroid carcinoma patients receiving long-term *thyroid-stimulating hormone suppressive therapy*, and resulted to be lower than in patients in shorter-term therapy [83, 84]. *Grave's disease*, an autoimmune disorder that causes hyperthyroidism, has been shown to have a strong correlation with decreased TBS, whereas BMD did not differ between the cases and control groups [85]. The findings of the studies on TBS and health conditions exhibited with hyperthyroidism are in line with each other and bring evidence that TBS is a bone parameter that aids in the management of patients suffering from this thyroid disorder.

TBS in hypercortisolism

Cushing's disease (CD) is characterized by an increased level of cortisol, namely hypercortisolism; and is an endocrine cause of obesity [86]. It has been shown that hypercortisolism has a negative effect on bone due to decreased bone formation that it causes. Nevertheless, only a mild decline in BMD has been reported; whereas the TBS decline was more pronounced [87, 88]. Moreover, TBS was lower in CD than in primary obesity patients, indicating the sensitivity of TBS to detect the variations along the hypercortisolism spectrum. Eller-Vainicher et al. investigated TBS in *subclinical hypercortisolism*; and demonstrated that the TBS decrease was associated with the cortisol excess and the severity of fractures and that TBS was more accurate than BMD in identifying patients at high risk for fracture [89]. Interestingly, Gonzalez Rodriguez et al. concluded in their study that high values of *evening cortisol* were associated with low TBS and increased prevalence of fracture in healthy postmenopausal women [90]. This evidence suggests the utility of TBS in bone health assessment in the spectrum of cortisol disorders.

TBS and other endocrine disorders

Few efforts have been devoted to investigating TBS in *hypogonadism* conditions, such as *Klinefelter Syndrome* or *estrogen deprivation*. TBS did not change significantly between men with the Klinefelter Syndrome than in

controls; whereas BMD was lower in cases [91]. In estrogen deprivation conditions, the decrease of TBS was more than the decrease of BMD, suggesting its independent role in bone health assessment and fracture prediction in these individuals [92].

Primary aldosteronism (PA), characterized by the hypersecretion of aldosterone, is associated with high fracture risk; a fact lacking support from studies assessing the association between PA and BMD. Kim et al. studied the values of TBS and BMD in cases of PA versus controls and showed that TBS was significantly lower among cases than controls, whereas BMD did not change significantly between the two groups [93]. PA might represent another endocrine disorder where TBS outperforms BMD in the assessment of skeletal fragility and the explanation of the high risk of fracture.

TBS in obesity

Inconsistent evidence exists on the presence of fracture risk and bone health condition in obesity. In general, BMD has shown to be higher among obese, while TBS lower. The difference in their associations with BMI has mainly been addressed to the possible technical limitation of TBS. However, the impact of overlying soft tissue on the measurement site is not only a matter of TBS. It is also problematic for BMD [94]. Nevertheless, to account for such limitation in obese, TBS is not applied in individuals with a BMI > 37 kg/m². Furthermore, for a deeper understanding and solution-seeking of this issue, a correction for direct regional soft tissue thickness, taking into account the morphotype of the patients had recently been developed to the TBS algorithm and has shown to surpass this limitation [95]. Further investigations of the updated—in regards to the soft tissue adjustment—TBS algorithm are necessary to support this initial evidence [96].

Conclusion

Alterations of bone strength in certain endocrine disorders are not always reflected in the BMD values, as their cause may lie in the microarchitecture of the bone. The TBS utility in fracture risk assessment in primary osteoporosis and diabetes has been robustly established. However, further larger studies could help for a deeper insight into the TBS behavior in other endocrine disorders where the presence of increased fracture risk is not (or only partially) explained by the BMD values, while TBS alone or in combination with BMD have proved promising at explaining the increased fracture risk.

In summary, knowledge of bone microarchitecture enriches our understanding of the pathophysiology of primary

and secondary osteoporosis. Nevertheless, bone quality cannot be comprehensively characterized by one sole parameter. Current noninvasive imaging techniques in combination with ex vivo mechanical and compositional techniques might provide a thorough understanding of bone quality. Currently, integrating the combined use of BMD, TBS, and clinical risk factors in clinical routine have proven efficient for osteoporosis and fracture risk management. This combination—within FRAX or not—enables us to fine-tune the risk of fracture stratification, treatment decision, and disease management.

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Compliance with ethical standards

Conflict of interest D.H. holds stock in Medimaps Group, the makers of the Trabecular Bone Score software. All other authors state that they have no competing interests.

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