Earlier this year the Canadian Panel of the International Society for Clinical Densitometry (whose members are listed at the end of this article) issued standards and guidelines for the practice of densitometry in Canada.1 The guidelines were based on a review of the literature and reflected the consensus of the panel. This article summarizes the key messages from those guidelines. It focuses on the use of bone densitometry by means of central dual-energy x-ray absorptiometry (DXA) in adults. It does not address densitometry in the pediatric population, nor does it cover technologies that evaluate bone mineral density (BMD) at peripheral skeletal sites such as heel, tibia or phalanx.

Bone densitometry is invaluable for the diagnosis of osteoporosis. Osteoporosis is associated with increases in the risk of fractures, morbidity and mortality.2 It is important to identify people with osteoporosis before the onset of fractures, because the occurrence of both vertebral and appendicular fractures is associated with greater risk of subsequent fractures.3 Bone densitometry allows for accurate and precise skeletal assessment and enables the detection of osteoporosis before the development of clinical fractures.

How does bone densitometry help in defining risk of fracture?

Several technologies are available for the measurement of BMD. Central DXA is currently the technology of choice.4 It measures BMD at the lumbar spine and the hip, and, with the appropriate software, it can also be used to measure BMD at other sites.

The relation between BMD and fracture risk in untreated patients has been evaluated prospectively in a number of large, well-designed studies.4–10 A meta-analysis of these studies confirmed that a decrease in BMD is associated with an increased risk of fracture.5 The predictive power of BMD for hip fracture is similar to the predictive power of blood pressure for stroke and better than the predictive power of serum cholesterol level for cardiovascular disease.6 In postmenopausal women the risk for hip fracture increases by a factor of 2.6 for each age-adjusted standard deviation (SD) decline in the BMD of the femoral neck.4,5 BMD at other sites (radius, calcaneus, hip and spine) has also been shown to correlate with the fracture risk at all sites.4 The best validation of diagnostic thresholds determined by DXA has been obtained for the hip.11

A working group of the World Health Organization (WHO) defined osteoporosis on the basis of the relation between BMD and fracture risk in postmenopausal white women.12 According to the WHO definitions, people with BMD more than 2.5 SDs below the mean for young adult women have osteoporosis.12 The T-score is the number of SDs that the patient’s BMD is above or below the mean reference value for young adults (the age of peak bone mass). The T-score is thus a comparison of the patient’s BMD with mean BMD for the young adult population. The Z-score is the number of SDs that the patient’s BMD is above or below the mean reference value for people of the same age. The Z-score is thus a comparison of the patient’s BMD with that of people of the same age.12 There are currently insufficient data regarding the relation between BMD and fracture risk to allow specific definitions of osteoporosis in premenopausal women, non-white women or men.

For example, the relation between BMD and fracture risk may be different in men and women.13 Current data are contradictory, and large prospective studies are needed to clarify the relation. Racial differences in skeletal size, skeletal geometry and hip axis length may contribute to racial differences in hip fracture rates.14 Therefore, it may be inappropriate to apply the WHO criteria to other groups.
without modification.

In evaluating fracture risk, BMD should be considered in conjunction with other clinical risk factors for fracture.\(^1\)\(^{16}\) Important independent risk factors include age 65 years or older, history of fracture as an adult, family history of osteoporotic fracture (especially of the hip) and poor neuromuscular function.\(^6\) Whether and how to intervene should be decided on the basis of a combined assessment of BMD and clinical risk factors for fracture.

**What are the indications and contraindications for bone densitometry?**

National guidelines\(^1\)\(^{16}\)\(^{17}\) suggest that BMD testing be targeted at people who have clinical risk factors for osteoporosis and those with conditions or disorders associated with bone loss (Tables 1 and 2). The Study of Osteoporotic Fractures provided a detailed analysis of risk factors for hip fractures.\(^6\) The same risk factors have been shown to predict fractures at other sites, including the spine.\(^1\)\(^4\)

BMD testing should be completed only if the results of the test will affect patient management. For example, a woman experiencing menopause without risk factors for osteoporosis (Table 2) does not require routine densitometry, whereas a woman with a personal history of fragility fracture after age 40 does require such testing. Fragility fractures are those that occur spontaneously or after minor trauma, such as a fall from standing height (e.g., from roller skates or ice skates), a fall from the sitting position or the lying position (e.g., from a chair or bed less than 1 m high), a fall after having missed 1 to 3 steps in a staircase or a fall while coughing.

**Can I trust a change in the bone density measurement?**

Current methods for measuring BMD typically demonstrate precision errors of the same order as natural short-term changes in BMD, which underscores the need for careful quality control of the instrument, the scanning technique and the analysis. The site measured is also important. In early postmenopausal women, bone loss from the spine exceeds that from the hip because of the more rapid turnover of trabecular bone, which predominates in the vertebral bodies.\(^1\)\(^9\) Similarly, an increase in skeletal BMD related to antiresorptive treatment (bisphosphonates, calcitonin, hormone replacement therapy or raloxifene) is usually most evident in the spine.\(^2\)\(^0\) For older subjects who are not receiving treatment, the decline in BMD in the hip generally exceeds that in the spine because of age-related degenerative sclerosis in the spine, which increases the BMD measurement but does not contribute to bone strength.\(^2\)\(^1\)

The precision error provides a measure of the reproducibility of the result in a repeated measurement. It is influenced by the instrument used, as well as by technologist- and patient-dependent factors (the last 2 of these factors tending to be more important than the first one). Therefore, it is strongly recommended that, whenever possible, follow-up measurements for a given patient be obtained by the same technologist using the same scanning procedure and the same instrument as for the original measurement. In addition, each laboratory should determine the precision of its measurements by evaluating a variety of clinical subjects under conditions that reflect “real-life” situations\(^2\)\(^2\) and should use this information in judging the significance of any change observed in a single patient. In routine clinical settings the following reproducibilities have been reported: for lumbar spine 1.8% to 2.3%, for femoral neck 2.3% to 3.6% and for total hip 1.7% to 2.5%.\(^2\)\(^3\)\(^{2}\) Each laboratory should include on the BMD report its own DXA reproducibilities for each measurement site. If the clinician is to conclude (with 95% confidence) that a change is not related to measurement error, the change must be at least 2.7 times the site-specific precision error at that centre. For example, a BMD of 2% at the lumbar spine would not be sta-

---

**Table 2: Factors identifying people who should be assessed for osteoporosis**

<table>
<thead>
<tr>
<th>Major risk factors</th>
<th>Minor risk factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age ≥ 65 yr</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Vertebral compression fracture</td>
<td>History of clinical hyperthyroidism</td>
</tr>
<tr>
<td>Fragility fracture after age 40</td>
<td>Long-term anticonvulsant therapy</td>
</tr>
<tr>
<td>Family history of osteoporotic fracture (especially hip fracture in mother)</td>
<td>Weight loss of &gt; 10% of weight at age 25</td>
</tr>
<tr>
<td>Systemic glucocorticoid therapy of at least 3 months’ duration</td>
<td>Weight &lt; 57 kg</td>
</tr>
<tr>
<td>Malabsorption syndrome</td>
<td>Smoking</td>
</tr>
<tr>
<td>Primary hyperparathyroidism</td>
<td>Excess alcohol intake</td>
</tr>
<tr>
<td>Propensity to fall</td>
<td>Excess caffeine intake</td>
</tr>
<tr>
<td>Appearance of osteopenia on radiograph</td>
<td>Low dietary calcium intake</td>
</tr>
<tr>
<td>Hypogonadism</td>
<td>Long-term heparin therapy</td>
</tr>
<tr>
<td>Early menopause (before age 45)</td>
<td></td>
</tr>
</tbody>
</table>

**Table 1: Common indications and contraindications for bone densitometry**

<table>
<thead>
<tr>
<th>Indications</th>
<th>Minor risk factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>At least 1 major or 2 minor risk factors for osteoporosis (see Table 2)</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Age ≥ 65 years, even without other risk factors</td>
<td>History of clinical hyperthyroidism</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>Long-term anticonvulsant therapy</td>
</tr>
<tr>
<td>Recent gastrointestinal contrast studies and nuclear medicine tests; suggested wait of at least 72 hours before a central bone densitometry scan (7 days for long-lived isotopes such as gallium)</td>
<td>Weight loss of &gt; 10% of weight at age 25</td>
</tr>
</tbody>
</table>

*People with at least 1 major or 2 minor risk factors should be considered candidates for bone densitometry. Reproduced from Brown and colleagues with permission.*

---

*Adapted from Baran and associates\(^1\)\(^4\) and Brown and colleagues.\(^1\)\(^6\)
tistically significant at the 95% confidence level if the precision error at the lumbar spine is 1% (since $2.77 \times 1% = 2.77\%$, which is greater than the observed 2% change). In summary, it is important not to over-interpret small changes.

**When should I repeat the bone density measurement?**

Average rates of bone loss are variable. The rate of loss is greater in untreated early postmenopausal women (approximately 1% to 2% per year) than in older women (less than 1% per year).

Follow-up BMD measurements in patients who are not receiving active treatment can help in identifying the subset of patients with rapid bone loss (“fast losers”). Repeat testing may also be useful in confirming a positive response to treatment. However, interpretation of results from testing undertaken to confirm a positive treatment effect is not completely straightforward. Although large increases in BMD are associated with large decreases in fracture risk, even small increases in BMD can dramatically reduce the risk of fractures. For example, raloxifene therapy has been associated with minimal increases of BMD at the lumbar spine and hip but significant decreases in the incidence of vertebral fractures. This pattern may indicate that some of the benefit from current antiresorptive therapies is mediated through mechanisms other than an increase in BMD. Conversely, absence of measurable change in BMD does not necessarily imply therapeutic failure and lack of antifracture benefit. In general, repeat BMD measurement can be considered 1 year after the original measurement if there is concern about rapid progressive bone loss (e.g., in glucocorticoid-induced osteoporosis, immobilization, acute gonadal insufficiency or primary hyperparathyroidism) or if the patient has started a new intervention (e.g., bisphosphonates). Less frequent repeat scanning (every 2 to 3 years) is appropriate in patients who have started therapies that increase BMD only minimally, such as nasal calcitonin and raloxifene, and those whose condition is already known to be stable or improving with drug therapy.

**How significant is exposure to radiation with DXA?**

Radiation safety practice requires that all radiation doses be considered harmful. However, the dose delivered by DXA is extremely small, of the order of everyday doses from background radiation (the environmental radiation to which everyone is exposed). Studies have not shown any ill effects (either long- or short-term) from such small doses.

In explaining radiation risk to the patient, the clinician can relate the dose from the procedure (an effective dose equivalent between 0.0005 and 0.0060 mSv [millisievert, the unit for dose equivalence]) to the dose from background radiation (effective dose equivalent 2.5 mSv/yr, although this value varies with altitude and geologic substrate) or the dose from everyday occurrences, such as flying from coast to coast (effective dose equivalent 0.5 mSv) or undergoing chest radiography (effective dose equivalent 0.1 mSv). For example, the radiation dose from DXA is 1/1000th the dose received when a person flies from coast to coast and 1/2000th that received from chest radiography. Because radiation risk relates not only to dose, but also to dose rate, such analogies are qualitative, not quantitative.

It should be noted that a fetus is more susceptible than an adult to radiation damage. Thus, for patients who may be pregnant, the indications and benefits of the assessment should be reviewed and the BMD measurement deferred until immediately after the patient’s next menstrual period, which might not be until after delivery (if the patient is actually pregnant). However, if the measurement has already been performed in a woman who might be pregnant, the additional risk associated with the radiation dose is so small that therapeutic abortion cannot be justified.

**What information does the bone density laboratory need?**

Clinical information is important in the interpretation of bone densitometry results. Referral information or the patient’s responses to a questionnaire completed at the time of the procedure are used as background information. The pertinent information includes age, menopausal status, prior history of atraumatic fractures, loss of height and any specific treatment for osteoporosis. Other medications such as thyroid replacement therapy, anticonvulsants and corticosteroids should be mentioned. Also important are smoking history, intake of vitamin D and calcium, family history of osteoporosis, coexisting illnesses and conditions, and history of alcohol use.

Recent barium studies (within the previous week) may leave residues that will attenuate the densitometry x-ray beam and falsely elevate BMD measurements. Ingestion of radiopaque medications may have the same affect. Recently injected radioisotopes for nuclear medicine procedures may emit radiation, which will falsely reduce measured BMD. This effect may be significant for approximately 72 hours after the radioisotope has been given, longer (up to a week) for long-lived radioisotopes such as gallium.

**What information should I expect to find in the bone density report?**

The T-score and the WHO Working Group definitions of normal (T-score greater than or equal to –1), osteopenia (T-score less than –1 but greater than –2.5) and...
osteoporosis (T-score equal to or below –2.5) will be given. When relevant, the Z-score will also be reported. A Z-score below –2 may indicate a need for detailed evaluation of the patient for secondary causes of osteoporosis (e.g., multiple myeloma, a malabsorption syndrome or glucocorticoid-induced osteoporosis). The report should outline any technical problems that might compromise the validity of the examination, such as degenerative changes (e.g., facet joint sclerosis or osteophytes), anatomic abnormalities (e.g., severe scoliosis) or compression fractures. The report should also include a qualitative assessment of fracture risk, as well as comments on serial BMD changes for patients who have previously undergone densitometry (which should take into account the precision error of the testing densitometer in assessing whether any change occurring since the previous study is in fact significant).

The results of the BMD assessment should be considered in conjunction with other clinical risk factors for fracture and are helpful in determining the patient’s future risk of fragility fractures. Table 4 of the Canadian osteoporosis guidelines provides information about assessing the patient’s risk in light of the BMD results and factors such as sex and age.

The case revisited

Mrs. B’s personal history of radial fracture after age 40 may indicate that she has osteoporosis, and further evaluation is appropriate. A BMD assessment will be helpful in assessing fracture risk. Because of the radial fracture, her risk of future fracture is greater than it would be otherwise.

On densitometry, the patient’s BMD for the lumbar spine (L1 to L4) was 1.04 g/cm², which corresponds to a T-score of –1.3. Total hip BMD was within the normal range (T-score –0.6). The patient is reassured by these results. Her current risk of fracture can be considered low, but it will increase as she ages. Some of the age-related increase in fracture susceptibility relates to bone loss, but much of it reflects other factors such as an increase in the risk of falling. This patient would benefit from appropriate calcium intake (1500 mg/day from all sources) and vitamin D supplementation (20 µg or 800 IU/day), as well as regular weight-bearing exercise.

The interpretation of repeat assessments should take into consideration the precision error at the testing centre. Serial assessments are of value in ensuring that rapid progressive bone loss does not occur and in monitoring the effectiveness of therapy.

Conclusions

Central DXA, a proven technology for the diagnosis and management of bone mineral loss, is now widely used across Canada. Optimum benefit of the technology requires maintenance of high standards in technical application, medical supervision and interpretation of results.

References


Correspondence to: Dr. Aliya A. Khan, McMaster University, 331-209 Sheldon Ave., Oakville ON L6J 1X8; fax 905 844-8966; Avkhan@aol.com


---

**A CMAJ Call for Medical Images: Clinical Vistas**

Send us your interesting clinical images!

Through scopes and scanners, on film and computer screens, with ultrasonography and microscopy, clinicians capture stunning images of illness and healing. *CMAJ* invites you to share your normally privy visual perspectives on anatomy, pathology, diagnostic procedures and therapeutic techniques. Let colleagues outside your specialty take a close look at the characteristic signs of rare conditions (Kayser-Fleischer rings in Wilson’s disease) or the interior marvels of your clinical terrain (colonoscopic view of an adenomatous polyp). We’re also interested in images that take a wider angle on the context of care (a recently cord-clamped newborn on a cold steel scale). If you have original, unpublished images that are beautiful or informative, rare or classic, we’d like to include them in *CMAJ’s* Clinical Vistas.

Send your images or queries to:

**Editorial Fellow • Canadian Medical Association Journal**

1867 Alta Vista Drive • Ottawa ON K1G 3Y6 Canada

or email pubs@cma.ca