Clinical Correlates Involving Deficiency of Vitamin D and Calcium

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Abstract: The effects of vitamin D and calcium deficiency are evident in several patients suffering from various diseases including osteoporosis and rickets. Other hormones such as glucocorticoids also play a role in bone homeostasis leading to bone growth promotion or retardation. Consequently, hormone imbalances in patients should be considered with suspected vitamin D and calcium deficiency. The deficiency of vitamin D and calcium lead to non-bone related abnormalities for instance promotion of prostate cancer and retardation of blood clotting, respectively. However, this study will be focusing on the detrimental effects of osteoporosis and rickets on patients. Vitamin D and calcium deficiency may be prevented by adequate sun exposure as well as diet supplementation. Vitamin D and calcium deficiency result in lack of intracellular mechanisms, loss of bone matrix and density hence, weaker bones. This leads to an increased probability of bone fractures.

Key words: Vitamin D, calcium, osteoporosis, rickets, diet, deficiency

INTRODUCTION

Several diseases such as osteoporosis, rickets, osteomalacia and rheumatoid arthritis have been linked with problems due to bone metabolism, often involving deficiency of vitamin D and calcium. The recommended daily dietary intake of vitamin D is 600 IU, ideally increased to 800 IU in the elderly, apart from having adequate exposure to UV light (http://ods.od.nih.gov/factsheets/vitaminD). Vitamin D rich foods include oily fish, milk and eggs as mentioned previously. Calcium rich sources include milk, cheese and yoghurt and the recommended daily intake is shown in the Table 1. According to Dowd and Stafford (2008), ideal vitamin D levels should be between 50 and 70 and additional calcium supplementation is unnecessary if the diet is acid-base balanced. For instance in osteoarthritis, the disease in bone remodelling occurs due to vitamin deficiency and acid excess in the diet. Therefore, calcium supplementation would be unlikely to improve the prognosis of the patient.

Calcium is absorbed in the small intestine via proteins such as TRPV6, calbindin 9K, FMCA1 and NCX1 and a small percentage remains in the intestines to be egested in the stools. Since, vitamin D is lipid soluble when obtained through the diet it diffuses through intestinal epithelium. Until recently, it was thought that people suffering from Crohn’s disease and coeliac disease have impaired calcium and vitamin D uptake (Douard et al., 2010). This would increase the risks having of bone fractures, rickets in growing children and osteoporosis. However, in a study shown by Kumari et al. (2010), it was found that young men having stable Crohn’s disease do not have impaired calcium uptake. It was only in patients who had inflammatory Crohn’s disease that calcium uptake was impaired and this problem was solved by administering vitamin D as it has been suggested to increase calcium transporting proteins and decrease the inflammatory response. This treatment was also found to reduce bone turnover markers and TNF-α levels.

THE ROLE OF OTHER HORMONES IN MAINTAINING BONE HOMEOSTASIS

Disorders in Growth Hormone (GH), glucocorticoids, Insulin-like Growth Factors (IGFs), thyroid hormones and steroids (such as oestrogens) also affect bone growth. For instance, glucocorticoids given to children may stunt bone growth. Receptors for glucocorticoids are found on osteoblasts and so it seems to be important in bone formation and mineral deposition. However, it was discovered that when given to mice in pharmacological

Table 1: A table showing the recommended daily amount of calcium in the diet

<table>
<thead>
<tr>
<th>Age</th>
<th>Male (mg)</th>
<th>Female (mg)</th>
<th>Pregnant (mg)</th>
<th>Lactating (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-6 months</td>
<td>200</td>
<td>200</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>7-12 months</td>
<td>200</td>
<td>200</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>1-3 years</td>
<td>700</td>
<td>700</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>4-8 years</td>
<td>1,000</td>
<td>1,000</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>9-11 years</td>
<td>1,300</td>
<td>1,300</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>14-18 years</td>
<td>1,300</td>
<td>1,300</td>
<td>1,300</td>
<td>1,300</td>
</tr>
<tr>
<td>19-50 years</td>
<td>1,000</td>
<td>1,000</td>
<td>1,000</td>
<td>1,000</td>
</tr>
<tr>
<td>51-70 years</td>
<td>1,000</td>
<td>1,200</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>71+ years</td>
<td>1,200</td>
<td>1,200</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

* Adequate Intake (AI)
most likely deficient in calcium and vitamin D. Effective calcium and vitamin D supplementation and weight-bearing exercises can help prevent osteoporosis. However, in certain cases, osteoporosis can begin in utero if the mother is vitamin D deficient. Osteoporosis affects the inorganic component of bone and results in thinning of bone.

Consequently, bone loses its strength and so is easily fractured. The typical clinical scenario would be an old lady falling and fracturing her hip and neck of femur. More often than not, patients would be unaware of their bone loss (in post-menopausal women it is about 1% bone loss per year) which is why osteoporosis is referred to as a silent disease.

It develops due to a decrease in oestrogen levels in women. However, it also occurs in men due to a decline in bone density as the body ages (Dowd and Stafford, 2008; Sunyecz, 2008).

In order to minimize risks of developing osteoporosis, it is recommended that exercise, calcium and vitamin D supplementation is taken before the age of 30 as that time would be peak bone mass after which a decline in bone density is observed. Vitamin D deficiency enhances osteoclast proliferation and so increased calcium mobilisation from bone.

It is important to note that calcium supplements may interfere with drugs the patient is taking for instance calcium will increase digoxin to toxicity levels causing arrhythmia and palpitations. All people above the age of 50 should take a bone density test (dual-energy X-ray absorptiometry implementing the T-score or the Z-score) so as to prevent the risk of bone fractures by starting treatment early on. Pharmacological treatment may be given by using bisphosphonates such as reclus and actonel which act by killing off osteoclasts and effectively inhibiting further bone resorption (Sunyecz, 2008).

Osteoporosis causes the spaces between one vertebra and the next to decrease, often resulting in compressional fractures. The patient usually suffers from kyphosis and appears shorter in size because of the compressed vertebrae as shown in Fig. 3 (http://www.true-beauty-tips.com/osteoporosis-alternatives.html).

Osteoporosis is a condition in which lack of calcium and vitamin D and age, amongst other factors induce a higher rate of bone resorption than bone formation. Consequently, the bone matrix is destroyed, cortical bone thins and larger spaces are formed within the bone architecture as shown in Fig. 4. Bones also appear thinner on X-Rays of osteoporotic patients (http://www.chiropractic-help.com/CAuses-of-Osteoporosis.html).
RICKETS

This disease manifests in children who are vitamin D deficient and is known as osteomalacia when it occurs in adults often leading to osteoporosis which is a decrease of total bone mass including collagenous tissue in the bone matrix. In the advanced stages, rickets is characterized by growth retardation, epiphysis enlargement in long bones, abnormal projections from the rimeage and bowing or bending of the legs and spine with weak, toneless muscles. This can be reversed by supplementation of high doses vitamin D or calcitriol and ensuring a calcium and phosphate rich diet to replenish the bone reservoir of ions required for homeostasis (Holick, 2006).

Since, vitamin D is deficient in most cases, calcium cannot be absorbed from the intestines to be utilized by bone. The decrease in calcium plasma levels causes an increase in PTH secretion which in turn enhances RANKL on osteoblasts resulting in increased resorption by osteoclasts. Also, phosphorous reabsorption is inhibited in the kidneys by PTH causing increased loss of phosphate in the urine. Since, serum calcium concentrations are also decreased, there is not enough calcium and phosphorous to ensure laying down of bone by osteoblasts.

Also, 1, 25-(OH),D[subscript]3 levels are usually normal or high in these children since, excess PTH stimulates its production in the kidneys and so it may appear that the child is not deficient for vitamin D. Coloured people were found more likely to be deficient in vitamin D when compared to their white counterparts. Since, rickets causes poor mineralization of the whole skeleton, bones which grow rapidly such as long bones show the first signs of abnormality especially in infants <18 months of age (Holick, 2006).

It is essential that people are exposed to enough sunlight as many people are deficient for vitamin D even though they live in a country with plenty of sunlight. However, this together with an appropriate diet is still not enough for some people because they are genetically vitamin D resistant which leads to rickets. This occurs because a point mutation can occur causing a variation in spliceing of VDR by leaving out exon 8, rendering VDR non-functional (Ma et al., 2009) or due to a mutation in an enzyme converting 25-(OH),D[subscript]3 to 1, 25-(OH),D[subscript]3. The signs for this condition are aggravated when compared to vitamin D deficiency. The treatment for this condition is pharmacological doses of calcitriol, calcium and phosphorous (Pettifor, 2005).

This X-ray was taken after a 10 months old, coloured boy sustained minor trauma to his left femur. The Fig. 5 shows bowing of the shaft of femur and an incomplete fracture. He was clearly suffering from vitamin D deficiency either due to a primary cause (inadequate diet and lack of sun exposure) or a secondary cause such as chronic renal failure, hypophosphataemia or vitamin-independent rickets.

The bowing and fracture occurred because of lack of calcium in the bones causing the bones to be weak and unable to support the patient’s body. As a result, minor trauma will cause bones to fracture easily since little resistance is offered to stress (http://www.radpod.org/2007/01/08/rickets/).

The Fig. 6 shows the classical appearance of a severe case of rickets including bowed legs, pigeon chest, wide joints, ribs and spinal deformities (http://lancastria.net/blog/tag/starvation).
The effects of rickets

- Odd shaped skull
- Pigeon chest
- Wide elbow joints
- Wide wrist joints
- Bowlegs
- Wide ankle joints
- Wide knee joints
- Pelvic deformities
- Odd shaped ribs
- Spine deformities
- Short stature (Stunted growth)
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Fig. 5: A radiograph showing left femur bowing and an incomplete fracture

Fig. 6: The clinical signs and symptoms of rickets in a child

CONCLUSION

It is remarkable how many people throughout the world are deficient for both vitamin D and calcium without themselves knowing. The best way to reach out to people is by educating the required daily amounts for vitamin D and calcium and showing them the reasons why they are an essential part of the diet especially in regions of the world lacking sunlight. Research is continually unfolding mysteries of the etiology of major diseases for instance prostate cancer and colorectal cancer, several of which seeming to involve a deficiency in vitamin D (Dowd and Stafford, 2008). This further stresses the importance of vitamin D and calcium in not only bone but the entire body’s physiology.

REFERENCES


