Role of hypogonadism in development of bone alterations in thalassemic patients

Nicola Napoli Enrico Carmina

Department of Clinical Medicine, University of Palermo

Address for correspondence: Prof. Enrico Carmina, M.D. Department of Clinical Medicine Via del Vespro 143, 90127 Palermo Ph. +39 091 6552953 Fax +39 091 6555995 E-mail: enricocarmina@libero.it

Summary

Although transfusions and chelation therapy have improved survival of patients afflicted with Thalassemia Major (TM), endocrine alterations are still common complications. Particularly, hypogonadism plays an important role in the aetiology of osteoporosis in these patients. It is clear that in most patients the gonadal failure is a consequence of a pituitary damage and that a hypogonadotropic hypogonadism occurs; nevertheless, also a gonadal damage because of iron deposition may be a further cause of hypogonadism. Prevalence or he pubertal failure ranges between 50 to 80% of the calles. I. some studies ferritin levels have been correlated with hyp. gonadism, suggesting that improvement of chelation . eau... may prevent or reduce the appearance of hyperana 'ism in TM patients. Treatment with HRT has show n cont, cting results, but patients who started the treatment is young age present better results than those who starre ' late. 'Jowever we may conclude that HRT is an impor'. It tre. 'ment option but should be prescribed in early age and issociated with biphosphonates.

KEY WORDS: thalassemia, hypoconadism. osteoporosis, iron overload.

Beta thalassemia $\ln n^{1/2}$ (TM) is an important health concern in many countries, most in the Mediterranean area, in the Middle East, $\ln n$ at Asia. In 1988 a report from the WHO documented that almost 50000 children per year were born with TM.

In the List decades, improved programs of transfusional and relation corrapy have permitted to TM patients to survive until heir forties or fifties and to get a better quality of life (1). However, so veral complications progressively arise and in particular endocrine alterations are common with diabetes, hypothyroidism, hypoparathyroidism, and hypogonadism being well known complications of adult thalassemia major (2-4).

More recently, in adult TM patients, bone mass deficiency has been recognised as a major problem that may cause pathologic fractures and limb deformities and greatly worsen the quality of life of these patients (5, 6). The etiology of bone mass deficiency in thalassemia is still unclear and many factors have been suggested as possible causes, including IGF-I deficiency, low vitamin D levels, alterations of genes related to collagen synthesis, bone cortical thinning because of bone ma row pansion and altered levels of some oligoelements while a zinc and copper (7). However, a main role is probable player by hypogonadism that is a hallmark of adult TM patier. s.

In this brief review, we will examine the cheracte. The soft hypogonadism in TM patients and the point ible link between hypogonadism and bone mass deficiently. Finally, we will discuss the role of the hormonal substitutive ther, py in the prevention and treatment of bone mass deficiency in True.

Prevalence and charac. The first of gonadal failure in thalassemia major

Most TM patients provent a delayed or absent puberty. In the patients who present puberty (at normal or late age), a gonadal failure offer provensished by a cours with appearance of menstrual disorder and an valation in women and spermatic abnormalities and reviced sexual activity in males. While it was initially suggented that a primary gonadal deficiency (a hypergunautrichic hypogonadism) occurs in TM patients, it is now clean that in most patients the gonadal failure is a consequence of a putuitary damage and that a hypogonadotropic hypogona hism occurs. Primitive gonadal damage may also occur but appears generally later (sometimes also in patients with hypogonadotropic hypogonadism). Therefore, in some adult TM patients (generally after the third decade) a mixed hypogonadism (central and gonadal) may be present (8).

All studies have documented that the prevalence of hypogonadism in adult TM patients is very high. In an Italian multicentric study, failure of the puberty was observed in 51% of the male and in 47% of female subjects. Between the female patients who reached a normal menarche, 23% presented a secondary amenorrhea and 11.6% oligomenorrhea (9).

In an Iranian study, puberty was delayed or absent in 72.6% of females and 80.8 % of the males and a disturbance of gonadal function was the most common endocrinopathy.

Several years ago, we have shown that about 70% of adult TM patients (both males and females) present some degree of hypogonadism (19). More recently, studying 30 adult TM women (mean age 28.5 \pm 1.3 years), we found that TM patients had lower serum levels of LH, FSH and estradiol (Figure 1) than controls of similar age (10). A 80% prevalence of hypogonadotropic hypogonadism was found and 20% of patients with hypogonadotropic hypogonadism presented also some evidence of primary gonadal failure.

A subset of adult TM patients present normal gonadal function. In our study, six adult TM women had normal ovulatory cycles with normal circulating levels of LH, FSH, estradiol and progesterone. Two of these TM patients previously had spontaneous successful pregnancies. Therefore, while most TM patients progressively develop a sexual hormone deficiency, some remain eugonadal. It is unclear what mechanism may determine these differences. In fact, while it is possible that differences in chelation or transfusional programs may explain it, we did not find any evidence of it and serum ferritin levels were not significantly different in eugonadal compared to hypogonadic adult TM women.

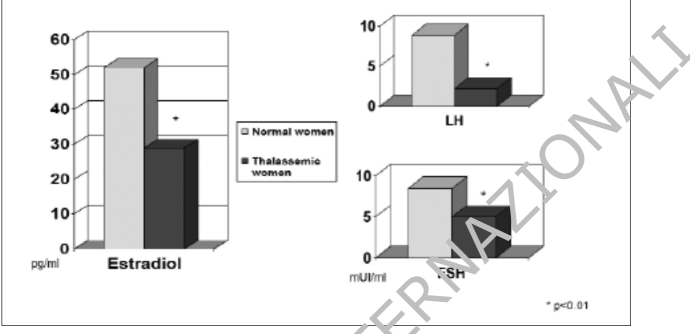


Figure 1 - Serum FSH, LH and estradiol in normal and in thalassemic adult patients

Causes of hypogonadism in adult TM patients

It has been suggested that in TM patients the hypoth lamus and the pituitary are damaged by the iron overload (1). In fact the pituitary gland is very sensitive to iron and along modest deposition may impair its functionalit. "List ogical studies have confirmed the damage of the pitul ory grand by iron overload in TM patients (12) and MR' has thow has significantly smaller anterior pituitary volume (13). It ther studies have shown that iron overload may iso therefore, it may be assumed that the hypogonadism of adult TM patients is intin inly a timesquence of the iron deposit.

Some authors have reported higher forritin levels in subjects who had hypogonadism compared to those with normal gonadal function (15). Ferritin 3 the protein that stores iron intracellularly and when it capat lity is exceeded an excess of active iron is released and crualyses the formation of free radicals. Free radicalr may camage membrane lipids, leading to mitochondrial and 'vsr somal damage and finally to cell death. Becaus of these findings it has been suggested that improvement f chelation treatments may prevent or reduce the appearaine of h/pogonadism in TM patients (16). While it may be, robasis, other studies did not find such correlations and our numogonadic patients had the same transfusion and chelation treatment than hypogonadic patients. Moreover, while hypogonadic patients had slightly higher ferritin circulatmg levels, the difference with normogonadic patients was not significant.

Relationships between hypogonadism and bone mass deficiency in TM

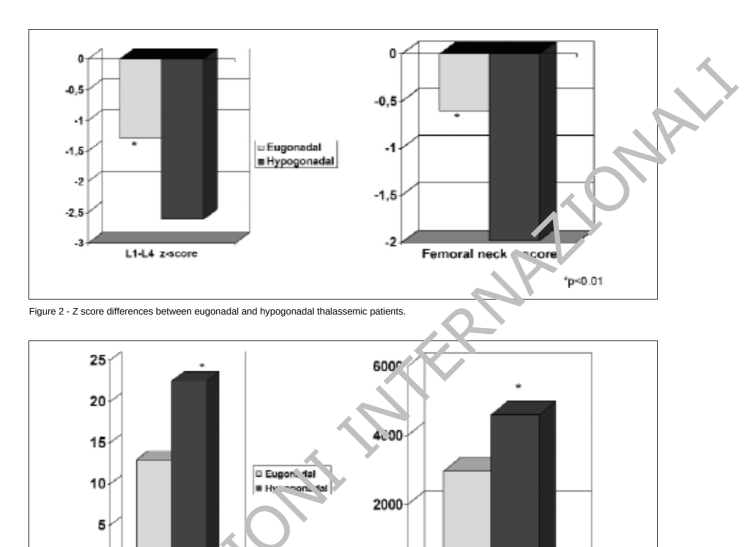
It is well known that sexual hormone deficiency determines an alteration of bone metabolism with increased bone resorption and progressive evolution towards osteoporosis. While nost studies have been conducted in menopausal women, it is clear, from clinical and experimental data, that sexual hormone deficiency may determine osteopenia or osteoporosis at any age and in both sexes. Because of it and although many other possible causes are present in thalassemia, hypogonadism may have a main role in determining bone mass deficiency.

Consistent with this hypothesis, we have shown that in adult TM patients a direct correlation between low bone mass and reduced estradiol exists, suggesting that reduced bone mass in many TM patients is mostly dependent on sex hormone deficiency (10). Moreover, hypogonadic TM patients had lower L1-L4 z score (Figure 2) and higher levels of bone markers (Figure 3) than eugonadal TM patients. All these data confirm that gonadal failure plays a main role in inducing bone mass deficiency of adult TM patients.

HRT effects on bone mass deficiency

Because this important role of hypogonadism, we should expect that hormonal replacement therapy (HRT) corrects or prevents bone mass deficiency in adult TM patients. Conflicting results have been presented with some studies indicating a clear improvement and other giving doubtful results. In our recent study, we found that female TM patients treated with HRT still have reduced bone mass and increased bone turnover compared to normal women of similar age. However, patients who started the HRT in younger age had better results than patients who started the treatment later. Maybe, it is important to start the treatment early, at the pubertal age. On the other hand, it cannot be excluded that HRT is not sufficient because other factors contribute to the bone mass deficiency of TM patients. Consistent with our results, Lasco et al. found that bone mass, measured by DEXA, was lower in all TM patients than in controls, but the difference was more marked in patients who didn't receive HRT (17). We can conclude that HRT is an important

Role of hypogonadism in development of bone alterations in thalassemic patients



 Bone alkalyni
 CTX

 phosphalase
 proliference is portials, fir e phosphatase and CTX between eugonadal and hypogonadal thalassemic patients.

part of the treatment of bone mass deficiency in adult TM patients ' ut , at the reatment has to be started early and that often alor, a is not sufficient to normalize bone mass.

Fecause C^t the disappointing results of HRT on bone mass in dult TM patients, it has been suggested to associate HRT and b rohor phonates (18). Limited experience is available but the results of our group and of other authors suggest that the results may be better than with HRT alone.

Conclusions

Hypogonadism is very common in adult TM patients and may be present in about 80% of the patients. It is important to assess hypothalamic-pituitary-gonadal function in young women with beta thalassemia major, so that those with gonadal fail-

Clinical Cases in Mineral and Bone Metabolism 2005; 2(1): 21-24

ure may start as soon as possible the replacement therapy. Patients with normal gonadal function should re-evaluate periodically their sexual hormone function because hypogonadism may develop after puberty and involve both pituitary and the gonads. In patients who do not respond adequately to HRT, other possible causes of osteoporosis should be screened and bisphosphonates could be added to the substitution therapy.

References

- Modell B, Letsky EA, Flynn DM, Peto R, Weatherall DJ. Survival and desferrioxamine in thalassaemia major. Br Med J. (Clin Res. Ed) 1982;284:1081-1084.
- 2. Vullo C, De SV, Katz M, Wonke B, Hoffbrand AV, Bagni B, Torre-

N. Napoli et al.

sani T, Tolis G, Masiero M, Di Palma A et al. Endocrine abnormalities in thalassemia. Ann N Y Acad Sci. 1990;612:293-310.

- Kwan EY, Lee AC, Li AM, Tam SC, Chan C., Lau YL, Low LC. A cross-sectional study of growth, puberty and endocrine function in patients with thalassaemia major in Hong Kong. J Paediatr Child Health. 1995;31:83-87.
- Costin G, Kogut MD, Hyman CB, Ortega JA. Endocrine abnormalities in thalassemia major. Am J Dis Child. 1979;133:497-502.
- Jensen CE, Tuck SM, Agnew JE, Koneru S, Morris RW, Yardumian A, Prescott E, Hoffbrand AV, Wonke B. High prevalence of low bone mass in thalassaemia major. Br J Haematol. 1998;103:911-915.
- Voskaridou E, Kyrtsonis MC, Terpos E, Skordili M, Theodoropoulos I, Bergele A, Diamanti E, Kalovidouris A, Loutradi A, Loukopoulos D. Bone resorption is increased in young adults with thalassaemia major. Br J Haematol. 2001;112:36-41.
- Bashir NA. Serum zinc and copper levels in sickle cell anaemia and beta-thalassaemia in North Jordan. Ann Trop Paediatr. 1995; 15:291-293.
- De SV, Vullo C, Katz M, Wonke B, Hoffbrand AV, Bagni B. Hypothalamic-pituitary-gonadal axis in thalassemic patients with secondary amenorrhea. Obstet Gynecol. 1988;72:643-647.
- Multicentre study on prevalence of endocrine complications in thalassaemia major. Italian Working Group on Endocrine Complications in Non-endocrine Diseases. Clin Endocrinol. (Oxf) 1995; 42:581-586.
- Carmina E, Di Fede G, Napoli N, Renda G, Vitale G, Lo PC, Bruno D, Malizia R, Rini GB. Hypogonadism and hormone replacement therapy on bone mass of adult women with thalassemia major. Calcif Tissue Int. 2004;74:68-71.

- 11. Iancu TC. Biological and ultrastructural aspects of iron overload: an overview. Pediatr Pathol. 1990;10:281-296.
- Bergeron C, Kovacs K. Pituitary siderosis. A histologic, immunocytologic, and ultrastructural study. Am J Pathol. 1978;93:295-309.
- Chatterjee R, Katz M, Oatridge A, Bydder GM, Porter JB. Selective loss of anterior pituitary volume with severe pituitary-gonadal insufficiency in poorly compliant male thalassemic patients with pubertal arrest. Ann N Y Acad Sci. 1998;850:479-482.
- 14. Canale VC, Steinherz P, New M, Erlandson M. Endocrino functor in thalassemia major. Ann N Y Acad Sci. 1974;232:333-34.
- Soliman AT, Nasr I, Thabet A, Rizk MM, El Mata, Yu, Hu, an chorionic gonadotropin therapy in adolescent bours with constitutional delayed puberty vs those with beta-the asseminmetor. Metabolism. 2005;54:15-23.
- Wang C, Tso SC, Todd D. Hypogona ... tropic hyper gonadism in severe beta-thalassemia: effect of thela ton and pulsatile gonadotropin-releasing hormone thera y. J Clin Endocrinol Metab. 1989;68:511-516.
- Lasco A, Morabito N, Gaudi A, Buen N, Wasniewska M, Frisina N. Effects of hormonal replacer therapy on bone metabolism in young adults with Lata-th Cassemia major. Osteoporos Int. 2001;12:570-575.
- Morabito N, Lasco Gaco A, Crisafulli A, Di Pietro C, Meo A, Frisina N. Bir A, Sphenates in the treatment of thalassemia-induced oster poror s. Osteoporos Int. 2002;13:644-649.
- Lo laco o F, in vizia R, Volpe FP, Carmina E, Galbo G. Esperienze cl'inche ui ma. trazione sessuale spontanea od indotta con gonr uotro une in talassemici politrasfusi. In: La maturazione sessi ale nei viseta-Thalassemia Major, Atti dell'Accademia delle Scio ze di Ferrara, 1986;225-228.