AB0829
OSTEOGENESIS IMPERFECTA: EVOLUTION OF THE BONE MINERAL DENSITY AND EVALUATION OF FRACTURE RISK UNDER BISPHOSPHONATES

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Background: Osteogenesis imperfecta (OI), or Lobstein's disease, also known as 'Glass bone disease' is a rare genetic disorder predisposing to low bone mass with impaired bone microarchitecture and abnormal quality of bone material, causing fracture susceptibility and bone deformities.

Objectives: The main objective of our work was to study the clinical and para-clinical characteristics, and global management, of adult patients and children with OI in a rheumatology department by comparing them with data from the literature.

The secondary objectives were to study the evolution of the bone mineral density and to evaluate the fracture incidence under treatment by Bisphosphonates.

Methods: This is a cross-sectional, observational, descriptive, analytical and monocentric study of subject with OI followed at Fattouma Bourguiba University Hospital in Monastir, Tunisia.

Results: We collected 13 patients with OI, including 5 children and 8 adults. Seven patients were female and 6 were male, including 12 patients with type IB and one with type IIIB. Three adult patients had a family history of OI. For children, it was sporadic and the genetic survey did not reveal similar cases in the family. The mean age of clinical onset of the disease was 3 years and 3 months [0-33 years] with a mean age at diagnosis of 15 years and 7 months [14 months-40 years]. The clinical manifestations were dominated by osteo-articular symptoms. Chronic mechanical bone pain related to fractures was present in eight patients.

Twelve patients had a history of fracture. These were repeated fractures without trauma or minimal trauma in the majority of cases mainly in the lower limbs. The first fracture site was the femur associated with other fracture sites (there were no fractures in the pelvis, hands and skull).

Deformities were observed in 7 patients, mainly in the limbs (6 patients), in the spine (2 patients) and in the thorax (1 patient).

In our study the frequency of osteoporosis and/or osteopenia regardless of the measured site and using the T-score value for women -2.5 and the Z-scores for men was 84.62%. All patients received Aredia protocol with calcium and vitamin D supplementation. The treatment was also multidisciplinary, Bisphosphonate treatment was well tolerated for the majority of patients. There was a decrease in the number of fractures and also a densitometric gain objective for the majority of patients.

Conclusion: Our results confirm the efficacy of BP in patients with OI regardless of age group. Short-term adverse are rare and transient.

REFERENCE

Disclosure of Interests: None declared


AB0830
CG AND DMARDS INDUCED BMD CHANGES IN PRE-, POSTMENOPAUSAL WOMEN AND MAN WITH RHEUMATOID ARTHRITIS (RA)

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Background: Risk factors for GC-induced fracture include low bone strength at the beginning of GC treatment and the rate of decline in bone mass during treatment, which is largely determined by the dose and duration of GC use [1]. The absolute risk of future fracture in an individual is substantially influenced by demographic and other characteristics (age, race, sex, and concomitant OP risk factors). For these reasons, it is important to identify differences in BMD changing depends on sex and reproductive status.

Objectives: To compare the BMD changes in man, pre- and postmenopausal women with RA depends on different treatment regiments.

Methods: The study was performed on 145 patients: 117 women (mean age 45.4±13.9 years, mean disease duration 9.7±7.7 years, 41% (n=48) postmenopausal) and 28 man (mean age 45.5±17.5 years, mean disease duration 5.7±4.8 years) with RA. Female patients were divided in two groups by menopause: premenopausal (Prem) in mean age 36.9±9.3 years and postmenopausal (PM) in the mean age 57.6±5.9 years. 68.4% women and 54.3% man received prednisolone ≤10 mg/day during 5.67±4.82 and 3.5±5.9 years respectively. 60.7% of men received MTX, 50% in combination with CGs. Among the women group 87% Prem patients received MTX, 58.3% in combination with GCs and 46.4% with biologics. 85.4% PM women took MTX, 68.8% in combination with GCs and 33.3% with biologics. BMD was measured in 3 part of the skeleton: hip (total and neck), lumbar spine, distal part of forearm.

Results: BMD was decreased in 44.5% of women and 42.9% of man. BMD of hip, lumbar spine, distal part of forearm were respectively decreased in 26.1%, 26.1%, 18.8% Prem women and 66.7%, 70.8%, 79.2% PM women. 39.3% of man had decrease BMD in the hip and 42.8% - in the lumbar spine. According to logistic regression in all women group the fact of GC intake was strongly associated with BMD decrease in the hip (total and neck) and lumbar spine (p<0.25, p<0.05), in man no association with the fact of CG intake was found. The effect of CGs on BMD in man was cumulative dose dependent, negative correlation between treatment duration and low hip neck (r=-0.44, p<0.05) and lumbar spine BMD (r=-0.5, p<0.05) was detected. In Prem women only association between CG therapy and decreased total and neck hip BMD (r=-0.33, p<0.05) was detected, any relationship with cumulative dose and treatment duration wasn’t found. Only in PM patients BMD decreasing depends on average cumulative dose and treatment duration. MTX intake was negatively correlated with BMD only in PM women in total hip and lumbar spine (r=-0.37, p<0.05) and in men in the distal part of forearm (r=2.48, p<0.05). In PM women GCs and in total hip and distal part of forearm was significantly higher in MTX-GCs-biologics group than without biologics (r=0.38, p<0.05). The level of PTG was significantly higher in PM patients on GCs and was negatively correlation with cumulative dose (r=-0.77, p<0.05). In Prem women GCs intake was negatively correlated with C-TP level (r=-0.37, p<0.05) and positively with RANKL level (r=0.35, p<0.05) and with the difference between C-TP and RANKL level (r=0.31, p<0.05).

Conclusion: A sexual differences in BMD changes during different treatment regiments was observed in different parts of the skeleton. In man BMD lost in hip and lumbar spine depends on GCs treatment duration and on in the distal part of forearm on MTX intake. In PM women BMD decreasing was strong associated with GC and MTX cumulative dose and treatment duration.

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AB0831
PERSPECTIVE ON THE CLINICAL PRACTICE MANAGEMENT OF HYPOVITAMINOSIS D FROM A MEETING OF ITALIAN EXPERTS

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