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| **Original Article** |

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| **Low Bone Mineral Density in Inflammatory Bowel Disease** |

**Incidence and Risk Factors for Low Bone Mineral Density in Inflammatory Bowel Disease**

**Imran Arshad, Shaista Zeb, Ehsan Rahim Memon and Prem Kumar**

**ABSTRACT**

**Objective:** Inflammatory bowel disease often causes osteoporosis. Inflammatory bowel disease patients' BMD falls due to clinical causes (IBD). Despite little data, BMD prevalence and risk factors are poorly known. Thus, this research investigated IBD's low BMD prevalence and causes.

**Study Design:** Cross-sectional study

**Place and Duration of Study:** This study was conducted at the Isra University Hospital's Gastroenterology Department included 65 adult ulcerative colitis patients from August 2021 to July 2022.

**Materials and Methods:** Patients' ages, BMIs, illnesses, sex, sickness durations, vitamin D levels, and steroid use histories were recorded. Dual-energy X-ray absorptiometry evaluated lumbar and femur bone mineral density (DEXA). Bone metabolism biochemical markers included deoxypyridinoline, serum calcium, osteocalcin, and phosphorus. Low bone mineral density was compared to medications, steroid usage, disease duration, age, and body mass index. SPSS 25 analyzed the data.

**Results:** The diagnosis of ulcerative colitis was made in 30 women (46.2% of the total) and 35 males (53.8% of the total). 38.6 3.54 years. 68.9% (n=45) had abnormal bone mineral density. 35.4% and 33.8% of 45 individuals with abnormal BMD had osteoporosis and osteopenia. Steroid use and illness duration substantially correlated with low bone mineral density in univariate analysis. Poor bone mineral density predicted disease duration in multivariate studies. Poor BMD was unrelated to age, body mass index, gender, vitamin D status, or steroid usage.

**Conclusion:** High levels of osteoporosis and osteopenia were seen in this study of people with inflammatory bowel disease. Poor bone mineral density was strongly associated with disease progression. Illness seems to be the biggest risk factor for low bone mineral density. Early detection of low MBD allows for prevention.

**Key Words:** Unknowns Inflammatory colitis, Bone deficiency

**Citation of article: Arshad I, Zeb S, Memon ER, Kumar P. Incidence and Risk Factors for Low Bone Mineral Density in Inflammatory Bowel Disease. Med Forum 2022;33(11):32-35.**

**INTRODUCTION**

Inflammatory bowel disease patients had 2% to 42% osteoporosis [1,2]. In numerous studies [2,3], inflammatory bowel disease patients had lower bone mineral density. Age, gender, body mass index, sickness duration, smoking, steroid history, and reduced food intake were negatively correlated with bone mineral density [4]. Many processes link inflammatory bowel disease to osteoporosis. Genetics, low body mass index, small intestinal resection, malabsorption, corticosteroid treatment, hypogonadism, and vitamin D deficiency may cause low bone mineral density [5-7].

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Received: August, 2022

Accepted: September, 2022

Printed: November, 2022

Crohn's disease increases bone mineral density risk. Crohn's disease may be diagnosed with low bone density. Crohn's disease patients with low bone mineral density demonstrated a greater connection with treatment than ulcerative colitis patients [8]. Crohn's disease is a major cause of osteopenia/osteoporosis, since azathioprine-induced remission increased BMD in patients. Steroids for low BMD may not work. Corticosteroids may lower BMD, however studies disagree [9].

Research on low bone mineral density has shown conflicting results (BMD). Jahsen et al. [10] observed no association between vitamin D and BMD in 120 IBD patients. Maldonado et al. [11] observed no association between vitamin D and calcium intake and bone mineral density in premenopausal inflammatory bowel

disease patients. Khadgawat [12] found a positive connection between poor BMD and calcium intake in Indian patients with inflammatory bowel disease. Despite their assumptions, steroid usage, age, and sickness duration did not affect BMD. Bishop et al. [13] found that corticosteroid usage, male gender, and inadequate vitamin D intake increased the risk for low BMD in 166 IBD patients, while age and disease site did not. Hip and vertebral fractures are prevalent in IBD patients. Breaks may be caused by osteoporosis factors. To monitor and treat low bone mineral density, this study investigated its prevalence and causes.

**MATERIALS AND METHODS**

This cross-sectional research at Hyderabad's Isra University Hospital's Gastroenterology Department included 65 adult ulcerative colitis patients from August 2021 to July 2022. Ages, BMIs, diagnoses, illness durations, vitamin D levels, and steroid use were documented. Dual-energy X-ray absorptiometry measured lumbar and femur BMD (DEXA). Deoxypyridinoline, serum calcium, osteocalcin, and phosphorus measured bone metabolism. Low bone mineral density was compared to drugs, steroids, illness duration, age, and BMI. DEXA eliminated isolated proctitis individuals without steroid history. Medical records and questionnaires provided demographic and clinical data. Endoscopic, clinical, histological, and radiographic tests identified IBD. Cancer, diabetes, chronic liver disease, pregnancy, and blood creatinine >1.5 mg/dL were excluded from this research.

Conventional DEXA assessed femur and L2-4 spine BMD. BMD produced T or Z-scores. Z-scores measure standard deviations from age- and gender-specific mean values, whereas T-scores relate to a young adult's bone mass peak. The WHO distinguishes osteoporosis from osteopenia using standard deviation. Osteoporosis is a WHO-defined T score of -2.5 SD, whereas osteopenia is -1 SD but not >2.5 SD [14]. Normal, insufficient, or deficient vitamin D 25-hydoxy levels were 20–30 ng/mL.

Comparing healthy and abnormal BMD patients (osteoporosis and osteopenia). SPSS 25 analyzed data. Low BMD was linked to illness duration, age, BMI, steroid use, gender, and vitamin D. Chi-square and Student's tests analyzed category and numeric variables. Univariate odds ratios have 95% confidence intervals.

**RESULTS**

Thirty females (46.2% of the total) and thirty-five boys (53.8% of the total) were diagnosed with ulcerative colitis. 38.6 years. 68.9% (n=45) had abnormal BMD. 35.4% and 33.8% of 45 abnormal BMD patients had osteoporosis and osteopenia. Steroid usage and illness duration were linked with poor bone mineral density in univariate analysis. Multivariate analyses predicted illness duration with low bone mineral density. Poor BMD was unrelated to age, BMI, gender, vitamin D status, or steroid use. Gender ratios. Figure 2 shows BMD abnormality. Table-I shows baseline features of DEXA-tested inflammatory bowel disease patients. Mann-Whitney.

**DISCUSSION**

The study's major findings showed that inflammatory bowel disease patients had a higher risk of osteoporosis and osteopenia. Bowel resection and corticosteroid medication most typically caused osteopenia or osteoporosis in IBD patients. Disease-related activities were independent risk factors. Similar to tertiary care referral studies, this study found higher incidence of osteopenia and osteoporosis [15]. In our study, inflammatory bowel disease patients had a higher rate of osteoporosis [16,17]. Inflammatory bowel disease patients with osteopenia and osteoporosis benefit from bisphosphonates in addition to vitamin D and calcium.

Pathological BMD in IBD patients showed that the femoral neck was less effective than the lumbar spine. Most studies show femoral neck osteoporosis is more common [18,19], yet this may not be true. Corticosteroids deplete trabecular bone. Bokemeyer et al. found reduced BMD in Chinese IBD patients [20]. In a study of 50 patients, 26 of whom had ulcerative colitis, inflammatory bowel disease patients had inadequate bone mineral density 63% more often than healthy young adults of the same age and gender. The recent study found that low bone mineral density prolonged sickness. Azuma et al. [21] found similar findings in 41 ulcerative colitis patients. Low BMI has long been linked to poor BMD. Poor BMD was not linked to BMI in this study. Asians may eat better than Westerners due to the BMI-nutrition link. In a cross-sectional study of 1250 postmenopausal women, reduced bone mineral density (BMD) was associated with higher BMI and worse socioeconomic status [22]. In this study, vitamin D levels did not affect BMD. The patient's limited capability may relate. Inflammatory bowel illness may cause vitamin D insufficiency due to poor diet, malabsorption, sun exposure, or circulation [23]. Hilmi et al. [24] observed no association between vitamin D and low BMD in 74 inflammatory bowel disease patients. Mouli et al. found that inflammatory bowel disease patients with vitamin D deficiency had low BMD [25].

In IBD patients, corticosteroids are typically utilized as first-line treatment for active BMD, which decreases intestinal calcium absorption, impairs osteoblast function, increases renal calcium excretion, and induces osteoblast death [26]. This study found no link between steroid usage and BMD. Alireza et al. [27] found a significant positive connection between poor BMD and steroid usage in 122 people with inflammatory bowel illness. Steroids cause osteopenia and osteoporosis, Abraham said. Corticosteroids have been linked to osteoporosis in Crohn's disease, and patients with abnormal BMD receive more steroids. 110 g of steroids may decrease bone mineral density (BMD) [28].

**CONCLUSION**

Osteoporosis and Osteopenia were shown to be prevalent in this study's population of individuals with inflammatory bowel disease. Furthermore, a strong correlation between disease progression and poor bone mineral density was found. The greatest risk factor for poor bone mineral density seems to be the illness itself. If patients with low MBD can be identified in the early stages, an appropriate preventative approach may be devised.

**Author’s Contribution:**

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| Concept & Design of Study: | Imran Arshad |
| Drafting: | Shaista Zeb |
| Data Analysis: | Ehsan Rahim Memon |
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**Conflict of Interest:** The study has no conflict of interest to declare by any author.

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