Multiple Hereditary Osteochondromatosis: A Case Report

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ABSTRACT
Objectives: Common carious lesions owing to vomiting are not widespread in children. In this case, we aimed to report an 11-years-old male patient with common carious lesions due to repeated vomitings, chewing and eating difficulty and retarded growth with Multiple Hereditary Osteochondromatosis (MHO).
Case Report: An 11-years-old boy was referred to Department of Pediatric Dentistry in Faculty of Dentistry because of eating difficulty owing to common carious lesions. It was seen that the patient growth was generally retarded in extra-oral examination. Some exostoses were also present on the extremities. It was learned that he was previously diagnosed as MHO in Faculty of Medicine. Nausea and vomiting have been commonly occurring after taking of Didronat. Chewing and eating difficulty and inadequate nutrition were present because of bad oral hygiene, carious lesions and remained roots. Growth was negatively affected by malnutrition and MHO.
Results: Diet recommendations were given and oral hygiene behaviors were rearranged. Preventive, surgical, restorative and prosthetical dental applications were applied for dental treatments. (Eur J Dent 2007;1:183-187)
Key Words: Bisphosphonates; Bone tumor; Chondrosarcoma; Multiple hereditary exostoses; Hereditary multiple osteochondromatosis.

INTRODUCTION
Multiple Hereditary Osteochondromatosis (MHO) is an autosomal dominant developmental disorder characterized by the presence of multiple osseous prominences with cartilage caps, arising most commonly from the metaphysis of long bones.1-4 However, these exostoses have also been found on the diaphysis of long bones, on flat bones, and/or on vertebrae.1,5,7 Osteochondromas are the most common benign osseous tumors. Sarcomatous changes have been documented to occur in approximately 1-5% of affected patients1-5,8 and defective endochondral ossification is likely to be involved in the formation of osteochondrosarcomas. Data indicated that most chondrocytes involved in the growth of osteochondromas can proliferate, and that some of them exhibit bone-forming cell characteristics.9
Cervical spinal cord compression resulting from osteochondroma is a rare and extremely serious complication of MHO.7,10 Some symptoms as myelopathy and paralysis may develop. Neurosurgical approach should be recommended in order to
achieve a spinal cord decompression, which usually results in excellent functional recovery and it usually has a favorable outcome provided surgical decompression is performed before major neurological damage develops.

Osteochondromas may contribute to altered osseous growth and growth plate of long bones. This altered discrepancy of limb-length or angular deformities and may lead to decreased range of motion, impaired function and possibly to premature osteoarthritis. Also, local muscle, tendon or nerve irritations can cause symptoms of secondary pain.

For local irritations and/or esthetic reasons, corrective or reconstructive surgery and excision of the exostosis may be performed. Treatment should aim not only at surgical resection of the masses but also at prevention of deformities. But, it was also reported that the risk of an abnormal scarring with keloid formation after osteochondroma excision in MHO patients after surgery. However if a malignancy suspected, complete surgical excision is the preferred treatment. Otherwise, if a spinal cord compression is present, patient may get an excellent recovery without neurologic defects after surgery.

Rarely, multiple osteocartilaginous nodules in temporomandibular joint space and associated joint dysfunction because of synovial osteochondromatosis are reported.

In this case report, an 11-years-old male patient with generalized carious lesions caused by vomiting, bad oral hygiene and his chewing and eating difficulty, and retarded growth affected by inadequate nutrition and MHO are presented.

**CASE REPORT**

An 11-years-old male patient was referred to Department of Pediatric Dentistry of Faculty of Dentistry due to severe carious lesions and eating difficulty. The patient growth was generally retarded. Some exostoses and shape anomalies were present on the extremities and vertebral column. In anamnesis, his mother reported that, he was diagnosed as MHO at early childhood at one of Faculty of Medicine. Mother had not got any serious problem during pregnancy and there was not kinship between mother and father. The baby had a normal body shape after birth, but the several deformities and outgrowth of the bones had begun to occur at the age of 2. When the MHO was diagnosed, the medical treatment was started immediately. The patient has been treating with Vitamin D (Devit-3 oral drop, 400 IU/day, daily use), Calcium (Calcimed Fort effervescent tablet, 500 mg/day, daily use) and Etidronate disodium (Didronat-PMO tablet, 400 mg/day, to be used in a scheme once a day for 15 days following an interruption duration of 2.5 months) to prevent from osteoporosis. His mother reported that, he had severe gastrointestinal symptoms such as nausea and vomiting with every intaking periods of Didronat. He had also more carious lesions. The chewing and eating difficulties and an inadequate nutrition were present. There was a general physical weakness. Growth was negatively affected by MHO and malnutrition.

His height was 124 centimeters and his weight was 21 kilograms. The growth of body was seen retarded according to other children in the same age. Several skeletal deformities were present on extremities and vertebras. The right and left forearms, wrists and shoulders, right shank and patella and left side of thorax were affected by disease (Figures 1 and 2). Mental retardation was not present.

Oral hygiene was very poor. In clinical and radiographical examinations, there were severe carious lesions on 11, 21, 12, 22, 31, 41, 32, 42, 73, 83 numbered teeth and remained roots were present from 74, 75, 36, 46 numbered teeth. 16 and 26 numbered teeth were not present. It was learned that, they were more previously extracted caused by severe carious lesions and pain. It was seen that, the rotation of the 23, 24 and 25 numbered teeth was abnormal in panoramic radiography and they were not shown in oral cavity yet. Wide plaque layers were present on the surfaces of the teeth and roots (Figures 3 and 4).

An effective tooth-brushing method was shown. The oral hygiene behaviours were rearranged. Essential diet recommendations were given and a topical fluoride gel was applied. The patient was prescribed with a chlorhexidine gluconate solution to reduce the levels of pathogen microorganisms before dental treatment. All remained roots with excessive carious lesions were then extracted (74,75,36,46 numbered teeth).

The carious lesions on 11,21,12,22,31,41,32,42,73,83 numbered teeth were cleaned with a high speed, water-cooled dental torque (KaVo Super
Torque, Germany) and teeth were then restored with a composite resin and a total-etch adhesive system [Charisma composite resin, Heraeus-Kulzer, Germany, Single Bond adhesive system, 3M, USA]. A consultation with Department of Orthodontics was made for patient. With two clinical considerations together, to make removable partial dentures for providing a sufficient chewing function immediately and observation of oral condition with frequently dental visits were approved for treatment.

**DISCUSSION**

Osteochondromatosis, also known as osteocartilaginous exostosis is the most frequent benign bone tumor of the skeletal system.1,5,22 Osteochondroma may occur in some short bones and may develop endochondral ossification despite its preference for long bones.22 Affected individuals exhibit bilateral deformities and multiple osteochondromas in the hands, wrists, legs and feet. The forearms and wrists are the most common influenced sites. These deformities include relative shortening of the ulna, bowing of the radius, ulnar deviation of the distal aspect of the radius, wrist and hand.5,11-14 In this case, all these deformities and multiple osteochondromas were present in defined locations. However, rarely reported mul-

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**Figure 1.** The clinical view of the shoulders and left side of the thorax.

**Figure 2.** The clinical view of the subjacent extremities.

**Figure 3.** The intra-oral clinical view of the patient.

**Figure 4.** The intra-oral radiographical view of the patient.

**Figure 5.** The intra-oral clinical view of the patient after prosthetic dental treatment.
tiple osteocartilaginous nodules in temporoman-
dibular joint space at Synovial Osteochondroma-
tosis17-20 were not present in our case.

Patil et al23 reported that the males were pre-
dominantly affected with Osteochondromatosis
such as in our case.

Some inhibitor drugs which were called as
Bisphosphonates are necessary in the treat-
ment of MHO for reducing bone resorption and improve
the bone mineral density (BMD).24-26 Bisphos-
phonates are powerful inhibitors of osteoclastic
activity25 and they are synthetic analogs of inor-
ganic pyrophosphate that have a high affinity for
calcium.24,25 They are excreted through the kid-
neys without metabolic alteration.26 During bone
resorption, they are released from the bone
surface and may be reincorporated into newly
formed bone or osteoclasts.24 So, they can play
an important role in the treatment of loss of min-
eral density and significantly increase BMD.24,27,28
Otherwise, bisphosphonates can cause some
gastrointestinal system (GIS) reactions, mucosal
ulcerations, exposed devitalized bone etc.24 They
led to compromised oral hygiene of the dentition
and local infections. This impaired oral hygiene,
in turn, may have facilitated local infection lead-
ing to advanced bone necrosis [osteonecrosis],24,25
and subsequent increased tooth mobility and loss.
Past dental history for many of the patients re-
vealed recent tooth extraction.24

Didronat [Etidronate disodium] which was
used medicament in this case is one of drugs of
Bisphosphonates.29,30 It increases the BMD and
bone turnover but may also give rise to some GIS
symptoms such as diarrhea, nausea, and vomit-
ing29 and/or esophageal lesions such as oesophag-
itis or oesophageal ulceration.29 Repeating vom-
itings cause to acidic levels of oral pH in oral cavity
and acidic pH helps to starting and developing of
the carious lesions on the teeth. If patient does
not adequately brushes his/her teeth daily, bad
oral hygiene rapidly boosts this condition. Carious
lesions cause to loss of the dental structures of
the teeth. The chewing and eating difficulties may
occur owing to the structural lost in time. Insuf-
ficient nutrition may affect growth negatively and
quality of life may also decrease.

In our case, adverse GIS effects of Didronat
was also caused to bad oral hygiene and some
GIS reactions. Therefore, several carious lesions,
loss of teeth and malnutrition were occurred and
all these symptoms have been negatively affected
the growth and life quality of the child. The pre-
ventive and other dental treatments are very im-
portant and necessary to obtain a good oral condi-
tion and sufficient chewing and eating functions.
The improved nutrition may provide a better life
quality for the patient. So, necessary dental treat-
ments were made for obtaining these better con-
ditions and patient was also called for regularly
dental controls and possible necessary dental
treatments in the future.

CONCLUSION

In this case report, a male patient in 11-years-
old was presented with common severe carious
lesions caused by repeated vomitings due to the
mandatory using of a medicine (Didronat) for MHO
and also bad oral hygiene. A multidisciplinary den-
tal treatment had applied to get improvements in
the oral and systemic conditions of the patient
near medical treatment.

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medical info of the third author for assessment of
the prognosis of the disease and the the effects of
the medicines on disease and body.

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