Chronic recurrent multifocal osteomyelitis in association with pyoderma gangraenosum

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Abstract

Background: Chronic recurrent multifocal osteomyelitis (CRMO) is a rare acquired inflammatory skeletal disorder of unknown origin. CRMO was first described by Gideon in 1972 [1] and mainly affects children and young adults of female gender [2–4]. Very similar to CRMO is the SAPHO-Syndrome, whose hallmarks are the presence of synovitis, acne, pustulosis, hyperostosis and osteitis (SAPHO). SAPHO-Syndrome is considered to affect predominantly the adult population. The age of the patient is, however, not diaforo-diagnostic [5], since adults with CRMO and children with SAPHO-Syndrome have been described so far [6, 7].

The CRMO is part of the clinical picture of non-bacterial Osteomyelitis (NBO) and typically presents a relapsing recurring course with both remission and spontaneous exacerbation. CRMO is typically encountered in the limbs and the metaphysis of long bones in particular. Usually the clinical symptoms include painful swellings of the affected regions. This case report describes the rare case of a CRMO of the mandible in association with pyoderma gangraenosum.

Case presentation: A 14-year old female caucasian patient, residing in the south of Germany, presented in the oncological outpatient clinic of our Department of Paediatrics and Adolescent Medicine in June 2014 complaining of increasing neck pain and progressive swelling at her left cheek ongoing for about 6 weeks. These symptoms had been occurring quarterly for 4 years, but had never been as pronounced. Blood biochemistry showed a moderately elevated CRP (35 mg/l) and a significantly increased blood sedimentation rate (BSR 48/120 mm). The panoramic radiograph, however, revealed a bone alteration in the left mandibular region. Further investigations confirmed the diagnosis of CRMO.

Conclusion: The present case underlines the fact that rare diseases might occasionally present with even more rare symptoms. These occasions can obviously be considered to present a considerable diagnostic challenge.

Keywords: Chronic recurrent multifocal osteomyelitis, CRMO, Non-bacterial osteomyelitis, NBO, Pyoderma gangraenosum

Abbreviations: BSR/ESR, Blood sedimentation rate; CMFS, Cranio-maxillo-facial surgeon; CRMO, Chronic recurrent multifocal osteomyelitis; CRP, C-reactive protein; CT, Computed tomography; ENT, Ear nose throat specialist; FD, Fibrous dysplasia; IBD, Chronic inflammatory bowel disease; LCH, Langerhans cell histiocytosis; MR(II), Magnetic resonance imaging; NBO, Non-bacterial osteomyelitis; PG, Pyoderma gangraenosum; PR, Panoramic radiograph; SAPHO, Synovitis, acne, pustulosis, hyperostosis and osteitis

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**Case presentation**

A written consent for this case report has been obtained by the parents. A 14-year-old female caucasian patient, presented in the oncological outpatient clinic of our Department of Paediatrics and Adolescent Medicine in June 2014 complaining of increasing neck pain and progressive swelling at her left cheek about 6 weeks ago (Fig. 1a). These symptoms had been occurring quarterly for 4 years, but had never been as pronounced. In all previous occasions the symptoms had improved with the use of ibuprofen. Furthermore skin lesions were noticed on both lower legs. The patient told us that these lesions had persisted for 3 years and had usually improved during the summer (Fig. 1b).

Suffering from ongoing trismus, the patient consulted an ear nose throat (ENT) specialist 4 months ago. The initial work diagnosis was that of a parotitis and an antibiotic therapy was implemented. Due to the persistence of the symptoms the advice of a cranio-maxillo-facial surgeon (CMFS) was sought, so as to investigate a possible relation of the symptoms to the wisdom teeth. Pericoronitis or any other disease related to wisdom teeth could indeed be excluded with the help of clinical examination and radiological investigation (PR - panoramic radiograph).

The PR, however, revealed an alteration of the trabecular structure of the left mandibular angle (Fig. 2). A Magnetic Resonance Imaging (MR) was performed for further investigation. Suspicion for Langerhans Cell histiocytosis (LCH), Ewing sarcoma, fibrous dysplasia, lymphoma and osteomyelitis was risen (Fig. 3). The young patient was referred to our department of paediatric oncology.

An initial assessment of the patient revealed a good general but a deteriorated nutritional status. The skin of the lower legs showed multiple ulcerous blueish skin lesions (Fig. 1b) and the skin of the face and the upper part of the body presented several acne scars. A 7x5cm, hardened, tender on palpation mass was detected at her left cheek. No clear local signs of inflammation were present. The lymph node status was unremarkable.

Further clinical examination revealed an abnormal posture, characterized by inclination of the head to the left, elevated shoulders and scoliosis. Pain was elicited on percussion of the cervical spine. Blood biochemistry showed a moderately elevated CRP (35 mg/l) and a significantly increased blood sedimentation rate (BSR 48/120 mm).

Ultrasound of the head and neck showed submandibular cervical lymphadenopathy at the left side and a diffuse swelling affecting the soft tissues of the left cheek (Fig. 4).

On chest x-ray (pa), no suspicious pulmonary nodules or consolidations were found.

Further imaging with contrast-enhanced CT was performed which revealed an extensive thickened left corpus and ramus of the mandible with a soft tissue mass (Fig. 5a). Destruction of the cortex of the mandible and a massive periosteal reaction (Fig. 5b), a left-sided cervical lymphadenopathy and diffuse contrast-enhancement in the soft tissue surrounding the mandible were identified. Similar osseous and soft tissue lesions were detected in the cervical vertebrae 2-5 including the vertebral arches and the transverse processes.

Biopsies were taken from both the lower jaw and the skin lesions. The histological examination of the specimen from the lower jaw showed features of a benign

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Fig. 1 a Distinct swelling of the left cheek (circle). b Skin lesions on both lower limbs (circles)
fibro-osseous lesion, consistent with fibrous dysplasia or secondary changes after osteomyelitis (Fig. 6). GNAS mutational analysis was performed to define it to exclude FD. The dermatopathological examination showed an abscess-forming neutrophil inflammation; taking into consideration the clinical appearance, the findings were consistent with pyoderma gangraenosum (PG).

Treatment with Ibuprofen and Sulbactam/Amoxicillin led to significant decrease of the mandibular swelling. Furthermore, the pain in the mandible and the back subsided, resulting in regular posture. An obvious reduction of the lower limb skin lesions and of acne vulgaris as well as increasing appetite were observed (Fig. 7).

After taking into consideration the clinical findings, the results of the performed investigation and the response to the provided treatment, the diagnosis of a chronic recurrent multifocal osteomyelitis (CRMO) was made.

The CRMO is part of the clinical picture of non-bacterial Osteomyelitis (NBO) which can be classified into three progression forms - acute NBO, persistent chronic NBO or chronic recurrent multifocal osteomyelitis [8]. CRMO typically presents a relapsing recurring course.
with both remission and spontaneous exacerbation [9, 10]. Intermission could last from months to years [11]. Jansson proposed major and minor diagnostic criteria of NBO (Table 1) [8]. NBO is quite likely if two major or one major and three minor criteria are met. Jannson also showed that the fewer criteria are met the more likely is an acute NBO. Furthermore the more criteria are met the more likely is a CRMO.

CRMO is a diagnosis that is made by exclusion. Medical history, radiological diagnostics and a bone biopsy are the cornerstones for establishing the diagnosis of CRMO. According to the current AWMF-guideline [12] numerous diagnostic methods might be used to exclude malignant or inflammatory conditions. The exclusion of malignant processes has the highest priority.

CRMO is typically encountered in the limbs and the metaphysis of long bones in particular. Usually the clinical symptoms include painful swellings of the affected regions. None of the typically affected sites were involved in our patient; two rare sites – the lower jaw and vertebral bodies – were affected instead. Radiological signs are hyperostosis and osteitis with lightening of the affected areas and surrounding sclerosis. As in children multifocal bone lesions could be consistent with a good

**Fig. 4** Ultrasound of the soft tissue of the cheeks (a, b). Inhomogenous widening on the left side (b) (circle)

**Fig. 5** a, c The CT reveals an extensive thickening of the left mandible with contrast agent enhancing soft tissue mass (circles). b, d Destruction of the cortex of the mandible and massive periostal reaction. Similar changes can also be found in the posterior aspect of the cervical spine (circles)
number of differential diagnoses—e.g. Langerhans cell histiocytosis or leukemia [13], a biopsy is recommended [14]. Ewing sarcoma could not radiologically be excluded since it might look identically. An exclusion of this diagnosis could only be achieved through the following biopsy.

Histologically CRMO shows characteristics of inflammation and/or fibrosis without a detectable pathogen [13, 14]. In our patient the pattern of a benign fibro-osseous lesion was present consistent with an expired inflammatory process. Fibrous dysplasia could be excluded from the differential diagnosis since a negative GNAS1-mutational status (GNAS 1 wild type) was identified. The histomorphological pattern was consistent with changes after osteomyelitis or previous surgical procedure. As there were no earlier interventions CRMO became more likely.

Jansson determined that in 31 % of CRMO patients exoskeletal findings were present [8]. An association with autoimmune disorders as IBD (chronic inflammatory bowel disease) [2] respectively Crohn’s disease [15] or Takayasu arteritis [16] is described. Far more frequent - incidence varies from 23 to 80 % [7]- associated skin lesions are present – as in our patient. Not only psoriasis and palmoplantar pustulosis [13] but also pyoderma gangrenosum [2] have been described so far. However various autoimmune diseases are considered as associated skin lesions but as they also occur alone, it can easily happen that these lesions are not included in the diagnosis -making [17–20]. Our patient also presented dubious skin lesions consistent with pyoderma gangrenosum. Dagan et al. wrote a case report of a CRMO in the ninth and tenth ribs as well as the tibia with associated PG. Nurre et al. presented a case of CRMO with associated PG. However CRMO presented initially at the left hand as well as other bones except the mandible. After biopsy a PG presented at the site of biopsy. Innis described a similar case with involvement of several bones. In his case, PG was present at the scalp and the limbs. In an article of Edwards et al. CRMO manifested at both tibiae and associated PG. To our knowledge this is the first of such case – involvement of the mandible with associated PG- described in the literature.

Due to the fact that its pathogenesis remains unclear, its treatment is mainly symptomatic. Anti-inflammatory drugs form the basis of the current treatment concept; occasionally the use of cortisone therapy might be

![Fig. 6 Biopsy from the lower jaw: Features of a benign fibro-osseous lesion, consistent with fibrous dysplasia or secondary changes after osteomyelitis. HE 100x](image)

![Fig. 7 MRI (a) and the patient (b) 6 months after therapy. The MR shows a distinct decrease of the thickening of the left mandible and the soft tissue mass in the posterior aspect of the cervical spine (circles)](image)
radiologic differentiation and classification of a patient sample of 86 cases.

In addition, an accurate medical history is of paramount importance when uncertain changes of the cranial skeleton are present. The physician should always enquire about new osseous changes, pain or new lesions and additional questions e.g. SCHOLAR [26]. If uncertain skin lesions or rheumatic disorders are present, CRMO should be in mind. NBO criteria might be helpful for diagnosis. As there are various manifestations, an interdisciplinary approach can be useful.

Conclusion

The present case underlines the fact that rare diseases might occasionally present with even more rare symptoms. CRMO in the mandible and CRMO with associated PG have already been described. For the first time we described an involvement of the mandible with associated PG. These occasions and combinations can obviously be considered to present a considerable diagnostic challenge. An in depth knowledge about the disease and its symptoms is a matter of extreme value in such cases.

In addition, an accurate medical history is of paramount importance when uncertain changes of the cranial skeleton are present. The physician should always enquire about new osseous changes, pain or new lesions and additional questions e.g. SCHOLAR [26]. If uncertain skin lesions or rheumatic disorders are present, CRMO should be in mind. NBO criteria might be helpful for diagnosis. As there are various manifestations, an interdisciplinary approach can be useful.

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Authors’ contributions

MCW, IB, ML, KB, MC, JJ, FWN, CW examined and treated the patient and collected the data. MCW, IB, ML, KB, KM, MC, JJ, FWN, CW discussed the case and data. MCW, OW, KM, KB wrote the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent

Written informed consent was obtained from the parents for publication of this case report and any accompanying images. A copy of the written consent is available for review by the editor of this journal.

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