Granulocytic Sarcoma with Meningeal Leukemia. A Case Report

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ABSTRACT

The case of a 63-year-old woman is presented who developed 18 years after undergoing allogenic bone marrow transplantation for chronic myeloid leukaemia a granulocytic sarcoma localised in the presacral region. Complete radiologic remission was obtained with the administration of nilotinib. She developed a metachron granulocytic sarcoma in the right frontal region associated to leptomeningeal infiltration. Complete clinical and near complete radiological remission was obtained with the administration of nilotinib. The patient was finally lost of progression of the frontal and meningeal localisations 45 months after the presentation of the first presacral lesion. Throughout the follow-up no sign of systemic leukaemia was detected.

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Introduction

Granulocytic sarcoma is a rarely occurring extramedullar form of myelocytic malignancy called also myeloid sarcoma or chloroma. Usually it is a manifestation of acute myeloid leukaemia or chronic myeloid leukaemia in accelerated or blastic phase [1, 2]. A retrospective analysis found 8 granulocytic sarcomas in 615 chronic myeloid leukaemia patients [2]. Median overall survival was 14 months with tyrosine kinase inhibitors (high dose imatinib and nilotinib).

The localisation can be highly variable with soft tissues in predilection but also lymph nodes, bone, epidural region and viscera are possible [3, 4]. The diagnosis is challenging. The uncommon irregular mass can be mistaken for haematoma. The histologic examination needs special expertise.

Case Report

A 63-year-old woman was hospitalised for lumbar pain. Tonsillectomy, thymectomy, cholecystectomy, hysterectomy and bilateral ovaricectomy can be revealed in her medical history. She underwent allogenic bone marrow transplantation for chronic myeloid leukaemia 18 years earlier and has been followed for a while for mild graft versus host disease. MRI showed a presacral mass. Whole body CT scan did not show any other tumoral lesion. The histologic examination of the presacral mass described immature round or ovoid cells with small cytoplasm. Immunohistochemistry showed CD117+, CD99+, CKAE1/AE3-, EMA-, CD56-, chromogranine-, MPO+, CD34 focal+, vimentin focal+, desmine-, CD3-, CD20-, CD138-, Ki67: 50%. The finding was compatible with the diagnosis of granulosarcoma (Figure 1). The diagnosis was confirmed by a second expert opinion and a positivity for Philadelphia chromosome detected with FISH.

The patient received major opioid analgesics and palliative irradiation to the sacral region. Imatinib 400 mg treatment was started and allowed a rapid clinical improvement with complete disappearance of the low back pain. The poor tolerance necessitated a dose reduction in the first step and sixteen months after treatment initiation the patient completely stopped to take the drug although, the efficacy was supported by PET CT showing no viable malignancy a year after the instauration of imatinib. Six months after having stopped to take imatinib she was complaining of frontal headache and blurred vision. Brain MRI showed a right frontal basicranial mass infiltrating the ethmoidal cellules associated to small meningeal lesions in several localisations. Liquor cytology showed elevated cell count, predominantly of lymphocytic and granulocytic type. Whole brain irradiation was discussed, but finally

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A treatment with nilotinib was started. Complete clinical and near complete radiological regression were obtained.

Eighteen months later, she presented with recurrent headache and vomiting. Brain MRI showed expansion of the cerebral and meningeal lesions. Liquor cytology showed again elevated cell count with 40% of blasts. Whole brain irradiation and four cycles of intrathecal cytarabine chemotherapy were given and allowed only a short clinical improvement. The patient was lost of progressing intracranial hypertension caused by uncontrolled meningeal sarcomatosis forty-five months after the diagnosis of the presacral granulosarcoma. Throughout the follow-up no sign of systemic leukaemia was detected (Figure 2).

**Discussion**

The unique feature of this case is that granulocytic sarcoma developed independently of active leukaemia. The relatively long survival observed in this pathology of poor prognosis is probably related to the lack of systemic manifestation. The omission of cerebral irradiation upon the detection of the intracranial lesions may be discussed. We decided not to perform it because the patient’s complaints were mild and the evidences of concomitant use with tyrosine kinase inhibitors are lacking. Nilotinib allowed a significant improvement in some days. It supports the good penetration of the drug through the blood-brain barrier. At the time of progression radiotherapy was finally realised and followed by intrathecal chemotherapy without real efficacy.

**Conclusion**

Granulocytic sarcoma may present without associated leukaemia. The response to imatinib and nilotinib is excellent at the onset of the treatment but the long-term efficacy is shorter as compared to chronic myeloid leukaemia.

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