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The Clinical Picture

Anemia, leukocytosis, abdominal pain, flushing, and bone and skin lesions



FIGURE 1. Confluent erythematous maculopapular lesions involving the trunk (maculopapular cutaneous mastocytosis).



FIGURE 2. Radiologic evaluation showed diffuse osteosclerosis together with lytic and blastic areas (arrows).

72-YEAR-OLD MAN presented with abdominal cramping, diarrhea, intermittent flushing, asthenia, and a weight loss of 10 kg (22 lb) in the past 6 months. Physical

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examination revealed hepatosplenomegaly and an erythematous, maculopapular, confluent rash on the trunk (FIGURE 1) that displayed the Darier sign (redness, swelling, and itching in response to stroking in the involved area).

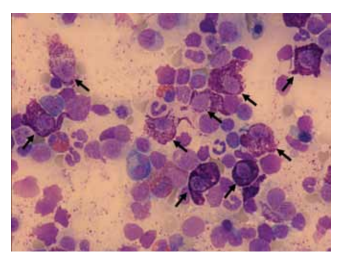


FIGURE3. Bone marrow smear demonstrating increased numbers of abnormal mast cells (May-Grünwald-Giemsa stain, x 600).

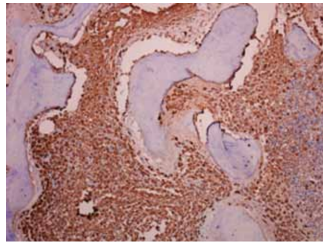


FIGURE 4. Bone marrow study demonstrating a massive infiltrate of abnormal mast cells (tryptase stain, x 200).

Laboratory analyses

- Hemoglobin 9.8 g/dL (normal 13–17 g/dL)
- White blood cell count 22.9 × 10⁹/L (3.8– 10)
- Vitamin B₁₂ 1,730 pg/mL (220–900)
- Serum tryptase 516 μg/L (5.5–13.5)
- Beta-2 microglobulin 4.14 mg/L (1.39–2.11).

Radiologic evaluation

Radiologic evaluation showed diffuse osteosclerosis with lytic and blastic areas (FIGURE 2).

- Q: Which is the most likely diagnosis?
- ☐ Carcinoid syndrome
- ☐ Histiocytosis
- ☐ Acute myeloblastic leukemia
- ☐ Systemic mastocytosis
- ☐ Chronic myeloblastic leukemia

A: The correct answer is systemic mastocytosis. The diagnosis was made according to the World Health Organization (WHO) diagnostic criteria for mastocytosis on the basis of the following findings in the bone marrow:

- Morphologically abnormal mast cells characterized by large size, spindle shape and poorly granulated cytoplasm (FIGURE 3) together with criteria for refractory cytopenia and multilineage dysplasia
- Diffuse infiltration by tryptase-positive mast cells as assessed by immunohistochemical study (FIGURE 4)

- One percent of mast cells that are immunophenotypically aberrant (CD25bright+), all of them showing an immature profile,¹ associated with features of multilineage dysplasia² as assessed by flow cytometry
- The activating D816V KIT mutation, detected by peptide nucleic acid-mediated polymerase chain reaction clamping technique.³
 The release of mast cell

MASTOCYTOSIS HAS SEVEN VARIANTS

Mastocytosis is a rare heterogeneous group of disorders characterized by proliferation and accumulation of abnormal mast cells in diverse organs and tissues, such as the skin, bone marrow, gastrointestinal tract, liver, spleen, or lymph nodes. ⁴⁻⁶ The release of mast cell mediators causes a wide variety of symptoms, ranging from pruritus, flushing, abdominal cramping, and diarrhea to severe anaphylaxis with vascular collapse. ^{7,8}

The WHO defines seven variants⁶:

- Cutaneous mastocytosis
- Indolent systemic mastocytosis
- Systemic mastocytosis with an associated (clonal) hematologic non-mast-cell disease (SM-AHNMD)
- Aggressive systemic mastocytosis
- Mast cell leukemia
- Mast cell sarcoma
- Extracutaneous mastocytoma.⁶

The release of mast cell mediators causes a wide variety of symptoms

KIT mutation as a diagnostic criterion and prognostic factor

In most cases of systemic involvement, the clonal nature of the disease can be established by finding activating mutations of *KIT*, usually D816V, in lesions in the skin, bone marrow cells, or both.⁹ Apart from its value as a diagnostic criterion for systemic mastocytosis, *KIT* mutation has been reported to be strongly associated with progression of indolent systemic mastocytosis, including the development of myeloid malignancies, when the mutation is detected not only in mast cells but in all hematopoietic lineages.¹⁰

In cases of SM-AHNMD, a possible pathophysiologic relationship between the disorder in the mast cells and the disorder in other cells could be explained by a *KIT* mutation in early hematopoietic progenitor cells, which further evolve into phenotypically different subclones.

A rational management plan for mastocytosis must include carefully counselling the patient and care providers, avoiding factors that trigger acute release of mast cell mediators, and giving antimediator therapy such as oral cromolyn sodium (Gastrocrom), antihistamines, and leukotriene antagonists to relieve the symptoms caused by mast-cell-mediator

The patient was alive with stable disease 14 months after starting therapy

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release.¹¹ In cases of SM-AHNMD, the clinical course and long-term prognosis are usually dominated by the concomitant hematologic malignancy, which should be treated as a separate entity.

CASE CONTINUED

Our patient's bone marrow was analyzed for the *KIT* mutation in highly purified bone marrow cell subpopulations sorted by fluorescence-activated cell sorting. The mutation was detected in his mast cells, CD34+ cells, eosinophils, monocytes, neutrophils, lymphocytes, and nucleated erythroid precursors. According to the WHO recommendations, he had SM-AHNMD, the associated hematologic disease being a myelodysplastic syndrome.

In view of his advanced age and concomitant myelodysplastic syndrome presenting with leukocytosis, we gave him hydroxyurea (Droxia; available in Spain as Hydrea) rather than other cytoreductive drugs as the first-line therapy. Additionally, we gave him corticosteroids in low doses, sodium cromolyn, and antihistamines to treat mastocytosis-related gastrointestinal symptoms. The patient was alive with stable disease 14 months after starting therapy.

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