Acute myeloid leukaemia with t(8;21) associated with ‘‘occult’’ mastocytosis. Report of an unusual case and review of the literature

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CASE REPORT

Approximately 20% of patients with systemic mastocytosis (SM) have an associated haematological, clonal, non-mast cell lineage disease, and most exhibit an associated myelogenous neoplasm. This report describes a 48 year old man with acute myeloid leukaemia (AML) and a type t(8;21) cytogenetic abnormality. Associated bone marrow mastocytosis (a defined subtype of SM) was only detected after successful polychemotherapy in the state of bone marrow aplasia, and persisted after complete remission of AML. The diagnosis of mastocytosis was based on the demonstration of a multifocal dense mastocytic infiltrate. The atypical mast cells showed prominent spindling and an aberrant immunophenotype, with coexpression of tryptase, chymase, KIT, and CD25—which is expressed only on neoplastic (not normal) mast cells. In addition, the transforming somatic mutation D816V of the c-kit gene was detected. Re-examination of the pretherapeutic (initial) bone marrow revealed a slight diffuse increase in partially spindle shaped mast cell clusters, was seen in this second biopsy specimen. Strongly metachromatic mast cells, sometimes forming dense clusters, was seen in this second biopsy specimen. Accordingly, the diagnosis of bone marrow (systemic) mastocytosis was established. Reviewing the pretherapeutic bone marrow biopsy, the final diagnosis of SM-AML could be established retrospectively. Follow up revealed a stable complete remission of the AML could be detected in a bone marrow biopsy specimen. However, an increased number of strongly metachromatic mast cells, sometimes forming dense clusters, was seen in this second biopsy specimen. Accordingly, the diagnosis of bone marrow (systemic) mastocytosis was established. Reviewing the pretherapeutic bone marrow biopsy, the final diagnosis of SM-AML could be established retrospectively. Follow up revealed a stable complete morphological remission of AML, although mastocytosis persisted morphologically for about 18 months. About 20 months after initial biopsy, however, compact mast cell infiltrates were no longer detectable, and there was only a slight increase in loosely scattered mast cells. Only very few of these mast cells expressed CD25. Thus, according to the WHO criteria, a definitive diagnosis of SM was not longer possible.

SYSTEMIC mastocytosis (SM) is a neoplasm of multi-lineage, myelomastocytic, or mast cell committed haemopoietic progenitor cells. In a large proportion of cases, these progenitor cells have the capacity to transform not only into SM, but also into a haemopoietic clonal non-mast cell lineage disease (AHNMD). In approximately 20% of all SM cases, an associated AHNMD is diagnosed (SM-AHNMD). In most patients, myelogenous neoplasms such as myelodysplastic syndromes or acute myeloid leukaemia (AML) occur. It is particularly important for the patient that full diagnosis is established by applying reliable disease criteria. The World Health Organisation (WHO) has recently proposed a new classification for AML and for mastocytosis, which includes disease criteria that may be very helpful in this regard. Here, we present a patient with AML and t(8;21), in whom a coexisting mastocytosis was not detectable at initial diagnosis because of an excessive accumulation of AML blasts obviously masking the compact mast cell infiltrates. After successful chemotherapy, however, a few compact (diagnostic) mast cell infiltrates were detected in the aplastic bone marrow. Applying the full range of diagnostic criteria, a final diagnosis of AML with coexisting systemic (bone marrow) mastocytosis (SM-AML) could be established.

‘‘It is particularly important for the patient that full diagnosis is established by applying reliable disease criteria’’

CASE REPORT AND METHODS

Case report

A 48 year old man presented with AML, cytologically classified on bone marrow smears as AML M2 according to FAB criteria, and AML with t(8;21) according to the WHO classification. The complete karyotype was 46, XY, t(8;21;12)(q22; q22; q24), suggesting a secondary aberration in the setting of a t(8;21) translocation. His previous medical history was unremarkable. He received three cycles of chemotherapy with idarubicin and cytosinarabinosid and one cycle of high dose chemotherapy with cytosinarabinosid, administered within a period of six months. The high dose chemotherapy cycle was accompanied by a pseudomembranous colitis, with subsequent surgical resection of the caecum. At the end of the first cycle of chemotherapy, complete remission of the AML could be detected in a bone marrow biopsy specimen. However, an increased number of strongly metachromatic mast cells, sometimes forming dense clusters, was seen in this second biopsy specimen. Accordingly, the diagnosis of bone marrow (systemic) mastocytosis was established. Reviewing the pretherapeutic bone marrow biopsy, the final diagnosis of SM-AML could be established retrospectively. Follow up revealed a stable complete morphological remission of AML, although mastocytosis persisted morphologically for about 18 months. About 20 months after initial biopsy, however, compact mast cell infiltrates were no longer detectable, and there was only a slight increase in loosely scattered mast cells. Only very few of these mast cells expressed CD25. Thus, according to the WHO criteria, a definitive diagnosis of SM was not longer possible.

MATERIAL AND METHODS

Bone marrow smears were routinely stained with Pappenheim stain, myeloperoxidase, non-specific esterase, and naphthol AS-D chloroacetate esterase. The bone marrow specimens were fixed in 5% neutral formalin, mildly decalcified overnight in edetic acid, and embedded in paraffin wax. All slides were routinely stained with haematoxylin and...
most remarkable finding, now enabling the diagnosis of bone an increased and focal accumulation of atypical metachro-
cytopoiesis. Blast cells were virtually absent now. However,
showed bone marrow aplasia, with pronounced stromal
gained after successful induction polychemotherapy 
with the diagnosis of AML. The second bone marrow biopsy 
were nearly absent. The findings were regarded as consistent 
negative (fig 1). Normal blood cell precursors and fat cells 
CD68, and terminal deoxynucleotidyl transferase were 
and myeloperoxidase, whereas antibodies against CD34, 
infiltration by blast cells expressing chloroacetate esterase 
treatment showed extreme hypercellularity, with diffuse 
hypercellularity, with subtotal depletion of the neutrophilic granulo-
cytogenetic point configuration of c-kit was found. Moreover, the last 
two consecutive trephine biopsies contained no compact mast cell aggregates, but exhibited a slight increase in loosely scattered mast cells, which were mostly negative for CD25. Hence, the diagnosis of morpho-
logic remission of both AML and mastocytosis was then established.

Bone marrow histology
The initial bone marrow trephine biopsy specimen before 
treatment showed extreme hypercellularity, with diffuse 
infiltration by blast cells expressing chloroacetate esterase 
and myeloperoxidase, whereas antibodies against CD34, 
and terminal deoxynucleotidyl transferase were negative (fig 1). Normal blood cell precursors and fat cells 
were nearly absent. The findings were regarded as consistent 
with the diagnosis of AML. The second bone marrow biopsy obtained after successful induction polychemotherapy 
showed bone marrow aplasia, with pronounced stromal oedema and subtotal depletion of the neutrophilic granulo-
cytogenesis. Blast cells were virtually absent now. However, 
an increased and focal accumulation of atypical metachro-
matic cells coexpressing tryptase, CD117, and CD25 was a most remarkable finding, now enabling the diagnosis of bone 
marrow mastocytosis (fig 2). On re-examination of the 
pretherapeutic bone marrow biopsy specimen, some incon-
spicuous, loosely scattered mast cells coexpressing tryptase 
and KIT (CD117) were detected. Dense (diagnostic) mast cell 
infiltrates were not present. Furthermore, immunohisto-
chemical analysis revealed an additional strong expression of 
CD25 on these mast cells, which emphasised their neoplastic nature. Obviously, mastocytosis had already been 
present in the initial biopsy, but was obscured by the dense, 
diffuse, blastic infiltrates of AML. The diagnosis of SM-
AHNMD, namely SM-AML M2 with t(8;21), could be 
established retrospectively, even in the initial bone marrow 
sample. Table 1 depicts the complete immunophenotypic 
characteristics of mast cells and myeloid blasts. In total, 
seven subsequent bone marrow trephine biopsy specimens 
were analysed during a follow up period of 20 months. All 
these bone marrow biopsies showed a continuing complete 
morphologic remission of AML, although some compact 
dense aggregates of atypical mast cells expressing CD25 were 
detectable in the first five follow up specimens. In contrast to 
AML, bone marrow mastocytosis persisted for about 18 
months. However, the last two consecutive trephine biopsies 
contained no compact mast cell aggregates, but exhibited a slight increase in loosely scattered mast cells, which were mostly negative for CD25. Hence, the diagnosis of morpho-
logic remission of both AML and mastocytosis was then established.

Molecular findings
Table 2 summarises the results of the molecular analyses. It is 
of major importance that the initial bone marrow biopsy contained the transforming point mutation at codon 816 (Asp816→Val) of the c-kit protooncogene, a finding that 
supports the diagnosis of mastocytosis. Although amplifiable 
DNA was not available from the second and fifth bone 
marrow biopsy specimens, the specific mutation could be 
demonstrated in biopsy specimens numbers 3 and 4. Although bone marrow biopsy specimens 6 and 7 both 
contained compact (diagnostic) mast cell infiltrates, the wild-
type configuration of c-kit was found. Moreover, the last 
trephine biopsy specimen again contained the specific 
mutation of c-kit (fig 3C), but morphologically diagnostic 
compact mast cell infiltrates were not detected, although a few mast cells were found to express CD25. The persistent

<table>
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<th>Blast cells</th>
<th>Mast cells</th>
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<tr>
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<td>117</td>
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<tr>
<td>LFA-2</td>
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AML, acute myeloid leukaemia; NC, not clustered; SM, systemic mastocytosis.

Figure
Acute myeloid leukaemia (AML) with “occult” mastocytosis (SM- 
AML). (A) Bone marrow section with extreme hypercellularity and subtotal 
depletion of normal blood cell precursors and fat cells. Sheets of blast 
cells dominate the picture. Infiltrates of mastocytosis cannot be detected 
(haematoxylin and eosin stain). (B) Bone marrow smear with a large cluster of 
medium sized to large blast cells exhibiting pleomorphic, sometimes 
slightly indented, nuclei and prominent nucleoli (Pappenheim stain). (C) Bone 
marrow section with chloroacetate esterase staining showing strong 
expression of this enzyme by blast cells. Note the bright granular cytoplasmic 
reactivity of almost all the blast cells (naphthol AS-D chloroacetate esterase). 
(D) Immunostaining with an antibody against tryptase reveals loosely 
scattered mast cells in the immediate vicinity of a trabeculum, possibly 
remnants of a distorted dense infiltrate. Note that the blast cells were non-
reactive with anti-tryptase (AA1; avidin–biotin complex method).

Table 1  SM-AML with t(8;21). Immunhistochemical phenotype of mast cells and AML blast cells

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associated clonal haemopoietic disorder is diagnosed. In about 20% of patients with SM, an extracutaneous organs. Urticaria pigmentosa-like skin lesions by an abnormal accumulation of mast cells in one or more general, systemic variants of mastocytosis are characterised with respect to clinical course, extent of organ involvement, and clinical outcome. In most major variants of systemic mast cell proliferative diseases. The revised WHO classification on mastocytosis defines four major subtypes of myeloid malignancies having been described. The AHNMD should be strictly classified according to the WHO or FAB criteria. Even pure cutaneous forms of mastocytosis (urticaria pigmentosa) have been reported to be associated with haematological malignancies, in particular AML. To the best of our knowledge, approximately 20 cases of SM-AML have been published. In six of these cases, a t(8;21) translocation is reported.

Our present case validates the usefulness of the diagnostic guidelines included in the WHO system of classification for mastocytosis, even to detect and define a so called occult mastocytosis”

Here, we present an unusual case in which the final diagnosis of SM-AML could be established only in the second (control) bone marrow biopsy specimen after chemotherapy for AML with t(8;21). Apparently, the complete diagnosis was obscured by a diffuse, compact infiltration of an extremely hypercellular bone marrow by myeloblasts. The cytoreductive treatment disclosed multifocal dense infiltrates of mastocytosis in an aplastic bone marrow. The morphological diagnosis in this trephine specimen is that of an isolated

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**Table 2** SM-AML with t(8;21). Follow up at 20 months, with histological and molecular/cytogenetic findings

<table>
<thead>
<tr>
<th>Sample (bone marrow trephine)</th>
<th>AML</th>
<th>Mastocytosis</th>
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<td>(pretherapeutic)</td>
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<td>2 (after induction therapy)</td>
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<td>3 (follow up)</td>
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<td>8 (follow up)</td>
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<tr>
<td>9 (follow up)</td>
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**AMN** acute myeloid leukaemia; **NT** not tested; **WT** wild type; **SM-AHNMD** systemic mastocytosis with haemopoietic clonal non-mast cell lineage disease; +, present; –, absent.

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**DISCUSSION**

The revised WHO classification on mastocytosis defines four major variants of systemic mast cell proliferative diseases. These entities strongly differ with respect to clinical course, extent of organ involvement, and clinical outcome. In general, systemic variants of mastocytosis are characterised by an abnormal accumulation of mast cells in one or more extracutaneous organs. Urticaria pigmentosa-like skin lesions may be present and are often associated with an indolent course of the disease. In about 20% of patients with SM, an associated clonal haemopoietic disorder is diagnosed. This observation led to the definition of a new entity termed systemic mastocytosis with associated clonal haemopoietic non-mast cell lineage disorder (SM-AHNMD). Quantitatively, SM-AHNMD assumes an intermediate position between the relatively common indolent SM and the rare aggressive SM. The clinical presentation and outcome in the setting of SM-AHNMD is often more related to the associated haematological neoplasm than to the mastocytosis. In most patients with SM-AHNMD, a myeloid neoplasm can be diagnosed. All major subtypes of myeloid malignancies including myelodysplastic and myeloproliferative syndromes have been described. The AHNMD should be strictly classified according to the WHO or FAB criteria. Even pure cutaneous forms of mastocytosis (urticaria pigmentosa) have been reported to be associated with haematological malignancies, in particular AML. To the best of our knowledge, approximately 20 cases of SM-AML have been published. In six of these cases, a t(8;21) translocation is reported.

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**Figure 2** Acute myeloid leukaemia (AML) with “overt” mastocytosis (SM-AML). (A) Bone marrow section after first cycle of chemotherapy shows overall pronounced stromal oedema, with subtotal aplasia of blood cell precursors. Note a giant erythron in the right lower quadrant (haematoxylin and eosin stain). (B) At this time a few dense infiltrates of strongly metachromatic mast cells intermingled with lymphocytes and macrophages were easily detected (Giemsa stain). (C) One year later, the bone marrow still showed complete remission of the AML, but contained some dense mast cell infiltrates, which were immunopositive using the antibody against tryptase (AA1; avidin–biotin complex (ABC) method). (D) The neoplastic nature of the mast cells is underlined by the demonstration of aberrant coexpression of the antigen CD25, which is not seen on normal/reactive mast cells, even in states of extreme hyperplasia. Note the specific anular (membrane associated) staining of the mast cells (anti-CD25, ABC method).

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**Table 2 SM-AML with t(8;21). Follow up at 20 months, with histological and molecular/cytogenetic findings**

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<td>9 (follow up)</td>
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**Ratio of AML1-ETO/G6PDH (housekeeping gene); †according to minor World Health Organisation criteria; ‡not sufficient for definite SM diagnosis.**
bone marrow mastocytosis, which, by definition, belongs to the SM group.

Ideally, the correct diagnosis of SM-AML would have been possible even in the initial biopsy diffusely infiltrated by AML, although compact mast cell infiltrates as the one and only major diagnostic criterion for mastocytosis were not detected. However, two minor diagnostic criteria were fulfilled morphologically, namely spindling in more than 25% of the loosely scattered mast cells and the proof of CD25 expression by mast cells, suggesting their neoplastic nature. Because the transforming point mutation D816V was also detected, three of a total of four minor diagnostic criteria were met, thus enabling the definitive diagnosis of mastocytosis. Our present case validates the usefulness of the diagnostic guidelines included in the WHO system of classification for mastocytosis, even to detect and define a so-called occult mastocytosis.

According to Puillart et al, patients with associated SM and AML t(8;21) have a worse prognosis with standard chemotherapy protocols than patients with AML t(8;21). However, our patient is still in complete haematological remission after two years of follow up, although the polychemotherapy was much more effective (and gave a much faster response) on the AML than the mastocytosis, which persisted morphologically for a period of at least 18 months, whereas the AML showed immediate complete remission after induction phase chemotherapy. In this respect, our case confirms the experiences reported by others, and underlines the resistance of neoplastic mast cells to conventional chemotherapy. The delayed decrease of mast cell infiltration in response to chemotherapy may be explained by the fact that treatment resulted in the depletion of mast cell progenitors but not of mature mast cells. In fact, mast cells are long lived cells and any effect of chemotherapy on their immature progenitors must be expected to lead to a reduction in mast cell numbers only after several months. In this regard, it is also of interest that the last bone marrow biopsy specimen to be analysed contained the transforming D816V mutation, whereas the morphological findings were regarded as not sufficient for a diagnosis of persistent mastocytosis. In fact, this last bone marrow biopsy showed a nearly normal microarchitecture, with intact haemopoiesis and a diffuse moderate increase in loosely scattered mast cells, about 15% of which had a spindle shape appearance and less than 5% of which expressed CD25. These findings may best be interpreted as minimal residual mastocytosis.

## ACKNOWLEDGEMENTS

This work was supported in part by a grant to KS from the University of Tübingen (fortune 1067-0-0).

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REFERENCES


