Spontaneous splenic rupture in a patient with acute promyelocytic leukaemia during induction chemotherapy

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Background. Acute promyelocytic leukaemia (APL) is a subtype of acute myeloid leukaemia with high curability rates. However, it is often accompanied by severe coagulopathy and bleeding risk and thus represents a potentially fatal haematological emergency requiring immediate treatment. Spontaneous splenic rupture is a rare event in all haematological malignancies. Only two clinical cases have been described so far in a setting of APL.

Case report. We report a patient with APL without preceding splenomegaly who underwent urgent splenectomy for spontaneously occurring splenic rupture during induction chemotherapy. After surgery the patient completed induction chemotherapy and achieved complete remission.

Conclusion. This is the second case of spontaneous splenic rupture without preceding splenomegaly in a patient with APL during induction chemotherapy described so far. Our case demonstrates that emergent splenectomy can be lifesaving even in the unfavourable condition of patient with severe immune deficiency.

Key words: acute promyelocytic leukaemia, splenic rupture, splenomegaly, abdominal pain

INTRODUCTION

APL is a subtype of acute myeloid leukaemia (AML) with specific biological and clinical features and high curability rates due to current treatment possibilities. From a cytogenetic and molecular point of view, it is well characterized by the presence of a balanced reciprocal translocation between chromosomes 15 and 17 (ref.1) giving rise to a fusion gene PML-RAR-α whose transcript detection by reverse transcriptase polymerase chain reaction (RT-PCR) has become the golden standard in APL diagnostics and minimal residual disease monitoring2,3. Administration of ATRA (all-trans retinoic acid) at a daily dose of 45 mg/m2 in combination with anthracycline-based chemotherapy for induction and consolidation followed by maintenance according to disease risk stratification has become a standard therapy4-8. APL is generally considered a curable disease with excellent long-term prognosis in cases where the initial disease specific and treatment related complications are managed. Cure rates from 80% to 90% can be expected in patients who survive induction and achieve complete remission9. Because of the high early mortality rate, APL represents a medical emergency. Immediate ATRA administration is essential to decrease the high early mortality rate, mostly resulting from bleeding into the central nervous system, lungs and gastrointestinal tract10. Spontaneous splenic rupture (SSR) is a rare but often life threatening complication most often seen in neoplastic disorders11. To our knowledge, only two clinical cases of SSR during induction chemotherapy of APL have been documented in the literature so far12,13. Here we report a case of spontaneously occurring splenic rupture during induction of APL.

CASE REPORT

A 57-year-old man was referred to our centre from a county hospital where he had been admitted for spontaneously occurring hematomas and febrile periods in combination with newly detected anaemia, thrombocytopenia and the presence of leukemic promyelocytes in the peripheral blood smear. On admission, the patient was in sepsis, complicated by hypotension. No spleen size enlargement was detected at physical examination. The initial blood count and laboratory evaluation showed severe anemia 71 g/L, thrombocytopenia 30x109/L, mild leukocytosis 13x109/L and significant CRP and renal function parameter elevation (Table 1). Pronounced coagulopathy was in accordance with general APL manifestation. The patient was assigned to high-risk disease according to Sanz14. Abundant intravenous hydration together with antibiotics were started with improvement in renal parameters and significant decrease in CRP. ATRA was administered immediately after suspicion of APL that was confirmed by conventional cytogenetic exam and RT-PCR on the second day after admission. The bone marrow leukemic immune phenotype examination revealed positiv-
In myeloid markers CD33, CD13, CD65 and CD15. Conventional bone marrow cytogenetic investigation confirmed t(15;17) and PML-RAR-α positivity was detected by RT-PCR in both bone marrow and peripheral blood (Fig. 1). Specific reciprocal translocation between chromosomes 15 and 17 was confirmed by the fluorescence in situ hybridisation (green arrows), where green signal corresponds to the promyelocytic leukaemia gene (PML) on chromosome 15 and the red signal to the retinoic acid receptor-alpha gene (RAR-α) on chromosome 17.

During the first 3 days of therapy, the patient developed signs and symptoms of manifested ATRA-syndrome with typical fluid retention and dyspnoea requiring intensive diuretic support and oxygen therapy. Chest X-ray showed signs of mild pulmonary congestion, discrete right-sided fluidothorax and confluent opacities in the basal region of the right lung suggestive of incipient pneumonia. The overall condition corresponded to rapidly increasing leucocytosis reaching almost 70x10^9/L on day 3 (Fig. 2). ATRA had to be stopped and hydroxyurea was added instead to the patient’s medications as cytoreduction together with corticosteroids. On day 4 of chemotherapy, the patient started complaining about abdominal discomfort with increased flatulence. The abdominal physical exami-

Fig. 1. Bone marrow aspirate showing diagnostic promyelocytes (left) and chromosomal translocation t(15;17)(q22;q21) diagnostic for APL (right).

Diagnostic bone marrow aspirate showed abundant promyelocytes with typical Auer rods in the cytoplasm (black arrow). Specific reciprocal translocation between chromosomes 15 and 17 was confirmed by the fluorescence in situ hybridisation (green arrows), where green signal corresponds to the promyelocytic leukaemia gene (PML) on chromosome 15 and the red signal to the retinoic acid receptor-alpha gene (RAR-α) on chromosome 17.

Fig. 2. WBC counts before and after splenectomy.

Graph showing evolution of WBC counts (x10^9/L) with a marked time point (day 6) at which splenectomy was performed.
nation showed no signs of peritoneal irritation. However, there was extreme drop in haemoglobin level from 76 g/L down to 59 g/L. The patient became severely hypotensive, showing signs of manifested circulatory failure requiring intensive blood and vasopressor support in terms of continual infusion of noradrenaline. Despite intensive packed red cell transfusions there was continuous decrease in haemoglobin level to 42 g/L and the patient’s overall condition was deteriorating. Computerized tomographic angiography (CTA) of the abdomen showed massive fluid haemorrhagic liquid and SSR with large intraparenchymal hematoma with no signs of active bleeding (Fig. 3). The consultant surgeon indicated splenectomy as a lifesaving emergency intervention. The patient was given 3 transfusion units (TU) of packed red cells, 2 TU of thrombocyte concentrates, 600 units of Prothromplex (® Baxter) and 4 g of fibrinogen as preoperative preparation. Blood count and coagulation parameters before substitution are shown in Table 2. Splenectomy was performed without complications. Pathological examination showed a spleen of 120x75x40 mm in size with subcapsular haemorrhagic focus of 65 mm and parenchymal rupture.

Cytoreduction with hydroxyurea was continued for 3 days and was stopped at the point where white cell count decreased to 3.76x10^9/L. Resumption the ATRA treatment followed immediately as the oral intake had been restored. The patient completed induction chemotherapy after the transfer to our department on the 4th

Table 1. Laboratory values at diagnosis.

<table>
<thead>
<tr>
<th>Blood count</th>
<th>WBC (x10^9/L)</th>
<th>HGB (g/L)</th>
<th>PLT (x10^9/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual differential count</td>
<td>neutrophil segments 5%, neutrophil bands 11%, neutrophil metamyelocytes 6%, neutrophil myelocytes 10%, neutrophil promyelocytes 56%, lymphocytes 11%, blasts 1%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coagulation</td>
<td>PT (s)</td>
<td>Fibrinogen (g/L)</td>
<td>D-dimers (ng/L)</td>
</tr>
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</tr>
<tr>
<td></td>
<td>17.8</td>
<td>1.70</td>
<td>20287</td>
</tr>
<tr>
<td>Biochemistry</td>
<td>Urea (mmol/L)</td>
<td>Creatinine (μmol/L)</td>
<td>CRP (mg/L)</td>
</tr>
<tr>
<td></td>
<td>13.1</td>
<td>162</td>
<td>150</td>
</tr>
</tbody>
</table>

Patient’s pathological blood count, coagulation and biochemical values at admission showing pronounced renal insufficiency and CRP elevation due to sepsis.

Table 2. Blood count and coagulation parameters before preoperative substitution.

<table>
<thead>
<tr>
<th>Blood count</th>
<th>WBC (x10^9/L)</th>
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<th>PLT (x10^9/L)</th>
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<tr>
<td></td>
<td>15</td>
<td>0.9</td>
<td>9633</td>
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Coagulopathy, severe anaemia and thrombocytopenia requiring thorough preoperative haematological substitution was revealed.
postoperative day complicated by febrile neutropenia followed by an episode of hypotension needing vasopressor support. The patient completed the first consolidation chemotherapy with idarubicin 5 mg/m² and Ara-C.

1 g/m² without any serious complications and is about to start the second consolidation with mitoxantrone 10 mg/m² and etoposide 100 mg/m² according to GIMEMA protocol. The patient has reached PML-RAR-α negative status in bone marrow currently.

**DISCUSSION**

There are very limited data available concerning SSR in APL. Only two clinical cases with SSR in APL were described so far. One of them had a moderate splenomegaly at ultrasound examination and the other one had no splenomegaly detected. Therefore, our case report presenting SSR in an APL patient during induction chemotherapy without previous splenomegaly is the second one published so far. SSR is a very rare event. However, according to a systematic review by Renzulli et al., haematological neoplastic disorders represent the largest etiologic group (16.4%) followed by infectious diseases (14.8%) (ref.11). Non-Hodgkin’s lymphomas are reported among the most often to be prone to SSR followed by various types of leukemia and chronic myeloproliferative disorders. Two cases from our centre with SSR occurring in a patient with diffuse large B-cell lymphoma and a patient with hairy cell leukemia were published a few years ago. AML represents the third most often aetiological subgroup of SSR of all haematological neoplastic disorders. The highest SSR-related mortality rate is associated with neoplastic disorders and was proven to be significantly related to the presence of splenomegaly and age over 40 years, whereas male sex, primary surgical treatment and multiple aetiological factors were not associated with mortality. Nevertheless, SSR can occur without previous splenomegaly as happened in our APL patient. Tachycardia, hypotension, abdominal pain, nausea, vomiting and rarely occurring Kehr’s sign are general symptoms of intra-abdominal haemorrhage. However, their manifestation can be blurred by supportive and concomitant therapy. Abdominal CT and CTA are the diagnostic methods of choice with high sensitivity and specificity as in our case.

**CONCLUSION**

Our clinical case demonstrates that emergent splenectomy can be lifesaving even in the severely unfavourable condition in a patient with SSR, severe immune deficiency and manifested coagulopathy after thorough haematological preoperative preparation. It requires an experienced surgical team with excellent multidisciplinary postoperative care.

**ABBREVIATIONS**

APL, Acute promyelocytic leukaemia; AML, Acute myeloid leukaemia; RT-PCR, Reverse transcriptase polymerase chain reaction; ATRA, All-trans retinoic acid; SSR, Spontaneous splenic; WBC, White blood cell count; HGB, Haemoglobin (HGB); PLT, Platelets.

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**REFERENCES**


