Liver tumor concurrent with chronic myelocytic leukemia and extreme thrombocytosis: A rare case report

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Case Report

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Abstract

Background Chronic myelocytic leukemia (CML) may occasionally occur after organ transplantation or long-term chemotherapy for solid tumors; Solid tumors secondary to long-term chemotherapy and biotherapy in CML have also been reported. However, the concurrence of solid tumor with CML and extreme thrombocytosis in an untreated patient is seldom reported.

Case presentation We describe a 61-year-old woman transferred to liver surgery department for the discovery of a large mass in the liver and an elevated plasmic AFP. She was initially diagnosed with liver cancer. Blood tests indicated a marked increase of platelets(2464x10^9/L). The chromosome examination of bone marrow biopsy indicated the existence of t(9;22) translocation, fluorescence in situ hybridisation (FISH) and PCR were both positive for the Bcr-Abl rearrangement. The diagnosis of CML was made. She received hydroxyurea and imatinib to treat CML, and platelet-lowering therapy, and then underwent a liver biopsy, which suggested a moderately-poorly differentiated adenocarcinoma, or might be a hepatic metastatic carcinoma. However, the patient refused further pathological examination and primary site screening for the tumor. She died six and a half months after discharge.

Conclusion: Here, we describe a rare case of liver cancer concurrent with CML and extreme thrombocytosis in an old patient. However, the exact relationship between the two tumors is still unclear, and more cases are desired.

Background

Chronic myeloid leukemia (CML) is a myeloproliferative disease caused by the reciprocal translocation of chromosome 9 and 22 which leads to a chimeric gene product known as Bcr-Abl[1]. Bcr-Abl fusion protein possesses constitutively activated Abl tyrosine kinase activity and activates downstream signalling pathways, including Ras/MAPK, JAK-STAT and PI3K/AKT, which are responsible for the pathogenesis of CML[2]. Clinically, CML is customarily considered a triphasic disease with an initial chronic phase (CP), an intermediate accelerated phase (AP) and a final, fatal blastic phase (BP)[3]. Patients with CML are usually characterised by extreme leucocytosis, myelocyte bulge, basophilia, eosinophilia in peripheral blood[4]. Some patients of CML can have mild to moderate thrombocytosis, there are even individual reports that platelets in CML patients can reach more than 10,000 × 10^9/L[5].

The relation between CML and solid tumors are rarely reported in four conditions: 1) Myeloid sarcoma (MS), commonly known as extramedullary acute myeloid leukemia (EM-AML), granulocytic sarcoma (GS) or chloroma, is a tumor composed of myeloid cells occurring at an extramedullary site[6]. The most common sites of involvement include small intestine, bone, skin, and lymphnode[7]. It can occur in different clinical settings: in patients with acute myeloid leukemia (AML) as a localized tissue mass, as blast crisis in patients with chronic myeloid leukemia (CML) or leukemic transformation in myelodysplastic syndrome (MDS), before AML and as an isolated neoplasm without evidence of AML[8]; 2) Secondary solid tumors after treatment for CML, the solid tumors reported after CML treatment
including hepatocellular carcinoma (HCC)[9], pulmonary carcinoid[10], and thyroid adenocarcinoma[11]. The underlying mechanisms is unclear, but may be relevant to the chemotherapy induced DNA damage, multiple genetic alterations and increasing host susceptibility to oncogenic viruses due to immunosuppression[9]; 3) Secondary CML after treatment for solid tumors, the formation of CML after treatment for solid tumors have been reported in non-small cell lung cancer[12], thyroid carcinoma[13, 14], breast cancer[15], and gastric adenocarcinoma[16, 17]. Up to now, evidence is lacking as to the frequency of therapy-related CML complicating cytotoxic therapy. 4) CML after organ transplant, more than 30 cases of CML associated with organ transplants have been reported up to now, most of them developed after kidney transplants[18, 19]. The development of CML in this situation may be associated with the postorgan transplant immunosuppressed state. However, it remains controversial whether the incidence of CML in such patients is truly higher than the general population owing to the limited number of reported cases.

In this case report, we reported the concurrent of liver cancer with CML and extreme thrombocytosis in an untreated old woman.

**Case Presentation**

A 61-year-old woman was transferred to our hospital with complaints of upper abdominal pain accompanied by anorexia for four months. Doctors in local hospital found a large mass in her right hepatic lobe by magnetic resonance imaging (MRI) and an elevated plasma level of AFP to 216.5 ng/ml, and she was initially diagnosed with liver cancer and admitted to our liver surgery department. She had a 5-year history of hepatitis B, but has never been treated; she also had a 5-year history of hypertension, and treated with indapamide tablet.

The physical examination on admission showed no jaundice, ascites, pedal edema or spider naevi, the liver and spleen are not touched below the costal margin. There were no obvious enlargement of superficial lymph nodes, and the remaining physical examination findings were unremarkable. Laboratory tests at admission (Table 1) showed a hemoglobin level of 111 g/L, white blood cell count of 7.01 × 10^9/L, and platelet count of 2464 × 10^9/L. The liver function test results were serum alanine aminotransferase of 55 U/L, serum aspartate aminotransferase of 82 U/L, and globulin of 69 g/L. HBV-DNA was 2.11 × 10^7 IU/ml. AFP was 308.8 ng/ml. Abdominal color Doppler ultrasonography showed a large hypoechoic area of about 9.5 cm x 5.7 cm in the right lobe of the liver, which was irregular in shape, unclear in boundary and uneven in internal echo, and the distal part of the right hepatic vein was not clearly displayed due to the compression of the mass(Fig. 1A). Abdominal contrast-enhanced CT showed a slightly low-density lesion in the right lobe of the liver with a maximum diameter of about 9.1 × 5.5 cm, and was slightly uneven enhancement and delayed enhancement(Fig. 1B,C). Hilar and retroperitoneal lymph nodes were increased and enlarged. These imaging and biochemical examination results suggested a neoplastic lesion in the liver, particularly HCC.
Table 1
The main laboratory test results at admission

|                         | Our values   | Normal range          |
|-------------------------|--------------|-----------------------|
| **WBC**                 | 7.01 × 10⁹ /L| 3.50–9.50 × 10⁹ /L    |
| **Neutrophils**         | 3.28 × 10⁹ /L| 1.80–6.30 × 10⁹ /L    |
| **RBC**                 | 3.71 × 10¹² /L| 3.80–5.10 × 10¹² /L   |
| **Hb**                  | 111 g/L      | 115–150 g/L           |
| **Platelet**            | 2.46 × 10⁹ /L| 125–350 × 10⁹ /L      |
| **AST**                 | 82 U/L       | 0–32 U/L              |
| **ALT**                 | 55 U/L       | 0–33 U/L              |
| **Total bilirubin(TBil)**| 8.9 µmol/L | 3.4–20.5 µmol/L       |
| **LDH**                 | 387 U/L      | 135–214 U/L           |
| **Total protein**       | 92.4 g/L     | 64–83 g/L             |
| **globulin**            | 64.4 g/L     | 20–35 g/L             |
| **Creatinine**          | 67 µmol/L    | 45–84 µmol/L          |
| **AFP**                 | 308.8 ng/ml  | 0–7 ng/ml             |
| **CEA**                 | 4.69 ng/mL   | 0–5 ng/mL             |
| **CA19-9**              | 0.6 U/mL     | 0.3–4 U/mL            |
| **ESR**                 | 95 mm/H      | 0–22 mm/H             |
| **ANA**                 | 1: 320       | -                     |
| **Urine protein**       | ++           | -                     |
| **HBV-DNA**             | 2.11 × 10⁷ IU/ml | -              |
| **PT**                  | 17.3 s       | 11.5–14.5             |
| **APTT**                | 46.8 s       | 29–42 s               |

This patient was transferred to our hematology department, further biochemical examination results showed that ESR was 95 mm/H, beta-2 microglobulin was 4.62 mg/L and anti-nuclear antibody was 1:320. No M protein was found in the test of monoclonal gamma globulin disease. A bone marrow aspiration and biopsy was performed to examine bone marrow, and results showed a hypercellular marrow with marked granulocytic and megakaryocytic hyperplasia (Fig. 2A,B). FISH analysis with Bcr/Abl DF probe was used to detect Bcr/Abl loci, when counting 200 cells in interphase, about 20% of the cells...
expressed two fusion signals, one red signal and one green signal (Fig. 3A), suggesting that Bcr/Abl fusion was positive. Chromosome examination also indicated the existence of t(9;22) translocation (Fig. 3B). Quantitative PCR for Bcr/Abl (p210) fusion transcript showed that the copy number of Bcr/Abl transcript was 421662, the ABL transcript copy was 384566, the ratio of Bcr/Abl to Abl transcript was 109.65%, and the international standardisation (IS) value was 109.65%. The diagnosis of chronic-phase CML was made. She received hydroxyurea (1 g po tid) and imatinib (0.4 g po qd) to treat CML, and platelet-lowering therapy with anagrelide (0.5 mg po bid), bayaspirin (100 mg po qd) and sodium bicarbonate (1 g po tid). She also received anti-hepatitis B virus therapy with entecavir (0.5 mg po qd). She developed agranulocytosis on the 7th day of treatment with hydroxyurea (Fig. 4A), so we reduced the dosage of hydroxyurea and gradually stopped it. She was treated with anti-infection and recombinant human granulocyte stimulating factor injection. Her hemoglobin and platelets gradually decreased (Fig. 4B, C). On the 14th day of hydroxyurea treatment, platelets returned to normal range (317 × 10^9/L), three days later, it continued to drop to 21 × 10^9/L, but gradually returned to normal. The main therapeutic drugs during this course are shown in Fig. 4D. The second bone marrow aspiration and biopsy results both suggested the erythrocyte and granulocyte series of bone marrow were suppressed, quantitative PCR for Bcr/Abl showed that the copy number of Bcr/Abl transcript was 79692, the Abl transcript copy was 1282604, the ratio of Bcr/Abl to Abl transcript was 6.21%, and the international standardisation (IS) value was 2.36%. Flow cytology of bone marrow suggested increased proportion of lymphocyte, no abnormal expression of T lymphocytes, NK cells and monoclonal B lymphocytes. The proportion of myeloid primordial cells was significantly decreased, and the neutrophils and monocytes were mainly in the mature stage (complete results is available in Additional file 1).

She underwent a liver biopsy when her blood routine results returned to near normal, HE staining suggested heterotopic adenoid and solid cell nests with necrosis in the chronic inflammatory liver tissue (Fig. 5A). The Ki-67 index was about 30–40% (Fig. 5B). Immunohistochemical studies showed positive staining of VILLIN, CK7, and EMA, while negative for Hepatocyte, Glypican-3, Arginasc1, AFP, CK19, CK20, CDX2, TTF-1, NapsinA, PAX8, ER, PR, and GATA3 (Fig. 5C-R). Pathologists concluded a moderately-poorly differentiated adenocarcinoma, and hepatic metastatic carcinoma can't be excluded. However, the patient refused further pathological examination and primary site screening for tumor, and asked to be discharged from hospital. She continued to take imatinib after discharge with a dosage of 0.4 g per day. However, she took the drug by herself at a reduced dose of 0.1 g per day due to a severe gastrointestinal reaction. At the fourth month of discharge, her platelet count was in normal range (272 × 10^9/L), while a color Doppler ultrasonography showed that the liver tumor was enlarged (9.8 cm x 7.3 cm). She died six and a half months after discharge, and the exact cause of her death was not clear.

**Discussion And Conclusions**

CML is a clonal disorder of hematopoietic stem cell characterised by the presence of Philadelphia chromosome t(9;22), it is manifested with increased number of myeloid cell count, and marked leucocytosis, myelocyte bulge, basophilia, eosinophilia and normal to mild thrombocytosis in peripheral
blood[4]. Rare cases of CML may present with an isolated, marked thrombocytosis, defined as a platelet count more than \(1000 \times 10^9/L\)[20]. Sora et al evaluated the marked thrombocytosis of CML in a large series of patients from 16 different Italian haematological centres from January 2002 to December 2015[21], and 87 of 1591 CML patients with extreme thrombocytosis were identified, with platelet count ranging from \(1054 \times 10^9/L\) to \(4720 \times 10^9/L\). CML patients with atypical clinical features and severe thrombocytosis were often mimicked with essential thrombocytosis (ET) as reported previously[22]. In our case, the patient's platelet count was as high as \(3622 \times 10^9/L\), but her erythrocyte and granulocyte series in peripheral blood were generally normal. However, FISH, chromosome examination and RT-PCR confirmed the diagnosis of CML. Moreover, bone marrow aspiration and biopsy revealed megakaryocytes that were smaller than normal cells and had a typical hyperlobated round nuclei in morphology. Therefore, we and other researchers propose that every case of ET should be tested for the Philadelphia chromosome to avoid missed diagnosis of CML[23].

The main pathogenesis of CML is the reciprocal translocation between chromosomes 9 and 22, which leads to the tyrosine kinase coding gene ABL of chromosome 9 translocated into BCR region of chromosome 22. An important result of this chromosome translocation is the product of fusion gene Abl-Bcr that encodes a deregulated tyrosine kinase resulting in manifestations of CML. Inhibition of Bcr-Abl oncogene expression and protein function are the primary treatment of CML based on its pathogenesis[24]. Bcr-Abl tyrosine-kinase inhibitors (TKI) are the first-line therapy for CML, among them, imatinib is one of the most classic, first-line drug[25]. In the recent study of Sora et al as mentioned previously, the majority of patients patients received a pre-treatment with hydroxycarbamide, and then treated with TKIs(63 patients received imatinib, 16 received dasatinib, and 8 were given nilotinib), 87% patients obtained a complete cytogenetic response (CCyR) or a major molecular response (MMR) after 12 months of treatment[21]. In our case, the patient received a pre-treatment with hydroxycarbamide and platelet-lowering therapy with anagrelide, but it did not achieve a relevant reduction in platelet count. However, when imatinib therapy was started, the platelet count rapidly decreased to the normal range. At the four-month follow-up after discharge, her platelet count remained in normal range.

Another issue in our report is the solid tumor complicating with CML in one patient. The abnormal elevated AFP, liver masses in CT imaging, and past history of hepatitis B infection all support the clinical diagnosis of liver tumor, particularly HCC. However, continuous monitoring of tumor markers revealed a downward trend in AFP(FigureS1). The pathological diagnosis of liver biopsy suggested a moderately-poorly differentiated adenocarcinoma, and it may also be metastatic liver tumor. However, neither abdominal enhanced CT, chest CT nor gastroscopy(FigureS2) showed tumors in other parts of the abdomen. In view of the particularity of liver tumors concurrent with hematological tumors, we consulted the literature and found some reports about liver tumors coexisted with hematological tumors. Katoh reported a 55-year-old Japanese man with CML, who had received an alkylating agent for 16 years, was diagnosed as HCC with clinically evident splenic metastases[9]. This is the first report in the literature on HCC in association with CML. The authors analyzed the long-term chemotherapy induced DNA damage may have endowed the HCC with an aggressive proliferative potential[9]. Ansari presented a case of a 72-
year-old man with a diagnosis of CML, showing subsequently with a liver mass which was pathological
diagnosed as Histiocytic sarcoma (HS)[26]. The liver mass showed a retained BCR-ABL1 translocation
suggesting clonality between the CML and HS. Therefore, the author proposed that the concurrent
expression of immunoglobulin heavy (IGH)/light-chain rearrangements or cytogenetic markers common
to the primary CML suggested an evolutionary mechanism involving lineage switching that could
potentially be affected by genetic or epigenetic factors which may occur at the level of a progenitor or the
malignant cell itself[26]. MS is a rare extramedullary presentation of neoplastic myeloid cells. They may
occur before, concurrent with, or after the diagnosis of myeloproliferative disorders, such as AML or
CML[27, 28]. Norsworthy reviewed 51 cases of biopsy-proven MS involving the liver, biliary tree or
pancreas[29], most cases presented with systemic disease, mainly in association with AML and CML.
The main symptom includes jaundice, right upperquadrant abdominal pain, fatigue, anorexia, nausea and
vomiting. Many cases were initially diagnosed as cholecystitis, pancreatitis or carcinoma, especially in
those with unknown hematological diseases, and underwent surgery. As immunohistochemistry
confirmed, the most commonly expressed cell surface markers of MS in these cases were
myeloperoxidase (63%), CD68 (40%), CD45 (30%), CD34 (30%), CD43 (30%), CD117 (28%), CD33 (15%),
lysozyme (15%) and CD13(8%)[29]. Unfortunately, for our case, there are no additional samples for further
immunohistochemical detection of MS markers. Although the platelet count was reduced to a normal
range, imatinib therapy did not reduce the size of the liver tumor.

In summary, we presented a rare case of liver cancer concurrent with CML and extreme thrombocytosis in
an old patient. Although the exact pathological relationship between the liver cancer and CML in this case
is unknown, we wish there are more reports on the concurrent CML and solid tumors like our case to
explore the underline mechanisms.

Declarations

Abbreviations: PCR: polymerase chain reaction; CT: computed tomography; ESR: erythrocyte
sedimentation rate; NK cell: natural killer cell; AFP: alpha-fetoprotein

Author Contributions: Ping Han and Zhiqiang Han collected and analyzed the data, and wrote the first
draft of the manuscript; Xia Mao, Jin Wang, Min Xiao and Qinglu Li performed the tests of bone marrow
cytology, chromosome, FISH and flow cytometry. Li Meng and Dean Tian were in charge of the patient’s
management; Zhenya Hong supervised patient’s management, critically revised the text, and made
substantial scientific contributions. All authors approved the final version of the manuscript.

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Ethics approval and consent to participate: The study was conducted in accordance with the Declaration of Helsinki, and the protocol was approved by the Ethics Committee of Tongji Hospital, Tongji Medical College, Huazhong University of Science and Technology. The patient’s daughter gave her written informed consent in accordance with the Declaration of Helsinki.

Consent for publication: The authors have obtained consent to publish from the participants to report individual patient data.

Competing interests: We have no conflicts of interest to declare.

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Figures
Imaging features of the liver tumor in the patient. Abdominal color Doppler ultrasonography (A) showed a large hypoechoic area in the right lobe of the liver. It was irregular in shape, unclear in boundary and uneven in internal echo, and the distal part of the right hepatic vein was compressed; abdominal contrast-enhanced CT (B) showed a 9.1x5.5cm slightly low-density lesion in the right lobe of the liver, it was slightly uneven enhancement and delayed enhancement.
Figure 2

Bone marrow aspiration and biopsy during hospitalization. The cytology of bone marrow at admission(A) showed a large number of platelet aggregation on bone marrow cytology slide. Megakaryocytes can be easily found, and there were 16 granular megakaryocyte and 4 thromocytogenic megakaryocyte in 20 megakaryocytes. Excessive nuclear lobulation, abnormal nuclear lobulations were also displayed. The bone marrow biopsy at admission(B) suggested the megakaryocytes were clustered or scattered in
different cell sizes, with hypolobation (mononuclear) or multinucleation (two or more round separated nuclei). The second bone marrow cytology(C) and biopsy (D) after treatment both showed that the proliferation of bone marrow was severely inhibited, megakaryocytes were easily seen and hypolobations of megakaryocytes were also observed.

**Figure 3**

Bone marrow genetic testing results. A representative image of FISH analysis with Bcr/Abl DF probe(A). The red probe represents Abl (chromosome 9), the green probe Bcr (chromosome 22), and the yellow area shows where the two fluorophores are in close contact, indicating the gene fusion. Chromosome
examination (B) indicated the reciprocal translocation of long arm of chromosome 22 and the long arm of chromosome 9, arrows indicated the chromosome translocation regions.

![Graphs A, B, C](image)

**Figure 4**

Treatments and the response to treatments of the patient. A, B, C representing the trend of white blood cells, hemoglobin and platelets after admission respectively. The abscissa represents the days after admission, and the ordinate shows the value of WBC(x10^9/L), hemoglobin(g/L) and platelets(x10^9/L), respectively. D showing the main drugs and treatments during hospitalization. rhG-CSF, recombinant human granulocyte-colony stimulating factor. rHuGM-CSF, recombinant human granulocyte-macrophage colony-stimulating factor. rhIL-11, recombinant human interleukin 11. CRCs, red blood cells suspension.
Figure 5

HE and immunohistochemical stainings of puncture liver tissues. HE staining of liver biopsy specimen (A) showed heterotopic adenoid and solid cell nests with necrosis in the chronic inflammatory liver tissue (x100). The Ki-67 labeling index was about 30-40% (B). C-R indicating the immunohistochemical stainings of CK7, EMA, VILLIN, Hepatocyte, Glypican-3, Arginase1, AFP, CK19, CK20, CDX2, TTF-1, NapsinA, PAX8, ER, PR, and GATA3, respectively (x100).

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