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Adults with germline CBL mutation complicated with juvenile myelomonocytic leukemia at infancy

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Abstract

Juvenile myelomonocytic leukemia (JMML) appears to be a life threatening disease and showed poor prognosis even after hematopoietic stem cell transplantation (HSCT) because of high relapse rate. On the other hand, recent molecular analysis revealed the heterogeneity of JMML. Here we report that 2 JMML patients survived more than 20 years without HSCT and both patients had uniparental disomy (UPD) of 11q23 where CBL is located without the phenomenon found in Noonan syndrome nor Noonan syndrome-like disorder. We think that some JMML patients with CBL mutation might show the good prognosis in later life after remission of JMML.
Introduction

Juvenile myelomonocytic leukemia (JMML) was estimated to show the poor prognosis.\textsuperscript{1} Hematopoietic stem cell transplantation (HSCT) was strongly recommended for JMML patients, however, about 30\% of patients showed relapse even after HSCT.\textsuperscript{2} Recent study revealed that the prognosis might be different from the genetic alterations, \textit{NRAS, KRAS, PTPN11, NF1} and \textit{CBL}.\textsuperscript{3}

Perez et al identified a heterozygous germline mutation in the \textit{CBL} gene (Y371H) in 3 unrelated JMML patients with a Noonan syndrome-like disorder.\textsuperscript{4} The mutation occurred de novo in 2 patients and was inherited from an unaffected father in 1 patient. Leukemic cells of all patients showed somatic loss of heterozygosity (LOH) at chromosome 11q23, including the \textit{CBL} gene. These findings indicate that heterozygous mutation in the \textit{CBL} gene is associated with predisposition for the development of JMML.

We found two JMML patients survived with good health longer than 20 years without HSCT (20 years old female and 31 years old male). Molecular analysis revealed that these 2 patients had the germline \textit{CBL} mutation with different frequency in each organ. Uniparental disomy (UPD) of chromosome
11q23 was also found in the recent peripheral blood, both. Furthermore, there are no typical features of Noonan syndrome nor Noonan syndrome like disorders in either patients. We will discuss about JMML with the CBL mutation.

**Patients**

Diagnostic criterion of JMML was depended on WHO 2008 classification. Case 1 : A six-month-old girl was admitted to our hospital because of abdominal swelling. She did not have the characteristics of Noonan syndrome (Supplemental Figure 1A). She was diagnosed as having JMML. We repeated chemotherapy using low-dose cytarabine (Ara-C) (30mg/m² for 10 days) and 6-mercaptoprine (6-MP) (40mg/m² for 7 days) for 126 times (about 10.5 years). Leukocytosis and hepatosplenomegaly diminished within one year after starting chemotherapy. She is now 20 years old and healthy without short stature, hearing loss, optic atrophy, congenital heart defects, malformations of certain blood and lymph vessels, hypertension or cardiomyopathy. Her intelligence quotient (IQ) was borderline (IQ test was 70 to 79). On her recent laboratory test, anti-nuclear antigen has been positive, however, she has no symptom of collagen disease.
Case 2: A nine-month-old boy was admitted to our hospital because of hepatosplenomegaly and leukocytosis. He did not have the characteristics of Noonan syndrome (Supplemental Figure 1B). He was diagnosed as having JMML. We continued chemotherapy using Ara-C and 6-MP for about 9 years. Hepatosplenomegaly and leukocytosis diminished within 2 years after starting chemotherapy. He is now 31 years old and healthy without short stature, intellectual disability, hearing loss, optic atrophy, congenital heart defects, malformations of certain blood and lymph vessels, hypertension or cardiomyopathy.

Materials and Methods

We analyzed several genes affecting the JMML leukemogenesis as follows, NRAS, KRAS, PTPN11, CBL, SETBP1 and JAK3. Written informed consent including picture presentation was obtained from each patient and institutional review board of Okayama University Hospital approved this project. Total DNA was extracted from stored BM mononuclear cells (BM-MNCs) sample, stored or recent peripheral blood MNCs (PB-MNCs) sample, buccal smear cells, nail or hair using QIAamp DNA Mini or Investigator kit.
Polymerase chain reaction (PCR) was performed using primer pair as
described previously.\textsuperscript{6,11,12,13,14} PCR product was directly sequenced using ABI
310 or 3130 sequencer (Applied Biosystems, Tokyo).

We used pyrosequencing to quantify the fraction of mutated alleles in DNA
samples from the different somatic tissues. DNA extracted from samples was
analyzed using the PyroMarkQ24 Gold Reagents according to the
manufacturer’s recommendation (QIAGEN, Hilden, Germany). Data analysis
was performed using the allelic quantitation software of the PyroMark Q24
system.

Genome-wide analysis for genetic lesions of mutated \textit{CBL} was performed
by single nucleotide polymorphism (SNP) array analysis. DNA extracted from
samples was analyzed using the GeneChip Human Mapping 250K \textit{NspI} array
(Affymetrix, Santa Clara, California). The data thus obtained were processed
using CNAG/AsCNAR software.

\textbf{Results}

The same \textit{CBL} Tyr(Y)371His(H) mutation was found in the recent PB-MNCs
in both patients without other gene mutations. Furthermore, the same \textit{CBL
mutation was found in the diagnostic BM-MNCs of case 1. Unfortunately we could not check the diagnostic sample of case 2 because his sample was not available. We identified the same \textit{CBL} Y371H mutation in DNA derived from buccal smear cells, nails of hands and hairs in two patients (Figure 1). The mutated frequency by pyrosequencing was different in each sample (Figure 1). In DNA from buccal smear cells of the both patients’ parents, no mutation was detected (Figure 1). We found that both cases are de novo mutation. SNP array data suggested that \textit{CBL} mutations were related to the LOH of chromosome 11q which included the \textit{CBL} gene, both (Figure 2).
Discussion

JMML was estimated to be a life-threatening disease and HSCT was strongly recommended as soon as possible, however, high relapse rate was still observed and resulted in the poor prognosis. Recent study revealed JMML prognosis might be quite different from the genetic alterations.\textsuperscript{6,7,8,9,10}

Our two cases survived for more than 20 years without HSCT and had the same sporadic germline \textit{CBL} mutation with 11qUPD. Recently germline \textit{CBL} mutation syndrome was presented.\textsuperscript{3,4,15,16} Interestingly, \textit{CBL} mutation in our two patients was found with different frequency in the different organ even after JMML remission. \textit{CBL} mutations are generally associated with LOH of the 11q23 chromosomal region resulting in apparent homozygosity for a \textit{CBL} mutation in JMML,\textsuperscript{17} but our cases still showed the LOH of the 11q23 in healthy PB samples, especially case 1. Previous reports suggested that germline heterozygous missense \textit{CBL} mutations were detected in 4 sporadic and 2 familial cases (total of 7 cases).\textsuperscript{4,15} None of the 7 individuals with a \textit{CBL} mutation had any hematological or solid tissue malignancy; however, the authors proposed the hypothesis that carriers of a germline \textit{CBL} mutation could be at increased risk for both, analogous to the predisposition to
malignancies seen in NF1, another disorder involving the RAS-MAPK pathway.\textsuperscript{4,15} These studies suggest that germline heterozygous CBL mutation carriers are susceptible to malignancy if reduction to homozygosity in somatic tissues occurs due to acquired UPD. For example, Kato et al reported duplication of KRAS due to acquired UPD caused JMML aggressive transformation.\textsuperscript{18} However, now our two cases have no malignancy except for JMML at infancy, although they had the germline CBL mutation with 11qUPD. We think the relationship between CBL mutation and 11qUPD in both cases (Supplemental figure 2). There seems to be four types which were shown in supplemental figure 2, they would exist as mixed status in each patients, however, type A would be dominant in case 1, and Type B would be dominant in case 2. Further large study about CBL mutation with 11q23UPD in adult cases will be needed in future.

Niemeyer et al suggested that germline CBL mutations have developmental, tumorigenic and functional consequences that resemble disorders that are caused by hyperactive Ras/Raf/MEK/ERK signaling and include NF-1, Noonan syndrome, Costello syndrome, cardiofaciocutaneous syndrome and Legius syndrome.\textsuperscript{16} Therefore, these germline mutated syndrome might
complicate JMML at infancy like transient abnormal myelopoiesis (TAM)/
transient myeloproliferative disorder (TMD) in Down syndrome.\textsuperscript{19}
Furthermore, \textit{CBL} mutation syndrome was reported with or without JMML
depended on the mutated site of \textit{CBL} gene.\textsuperscript{15,16} Interestingly, several reported
cases and our two patients with the same mutation Y371H does not have any
physiologic abnormalities such as hearing loss, optic atrophy, hypertension or
cardiomyopathy.\textsuperscript{4} Future study will enable to predict the prognosis of \textit{CBL}-
mutated JMML patients. Further study in long term survivor of JMML
patients will be needed in future.

In conclusion, JMML seems to show the heterogeneity due to the genetic
alterations. Some \textit{CBL}-mutated patients without typical phenomena like
Noonan syndrome might show the good clinical course after JMML remission.

\textbf{Acknowledgement}

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\textbf{Figure Legends}

Figure 1. Results of mutation analysis of \textit{CBL} by direct sequencing and
pyrosequencing in cases 1 and 2. The same mutation of Y371H, 1111 T>C was
observed in both patients. The frequencies of mutated allele by
pyrosequencing were different from the tissue type as follows, for Case 1, PB-
MNCs 87%, buccal smear cells 52%, nails 62%, hair 60%. For Case 2, PB-
MNCs 58%, buccal smear cells 48%, nails 42%, hair 51%. Their parents did
not have this CBL mutation in their buccal smear cells.

Figure 2. Results of SNP array analysis in cases 1 and 2. Figure 2A, and 2B:
In case 1, uniparental disomy (UPD) of chromosome 11q was observed in the
first diagnostic bone marrow sample (5 months old after birth) (Figure 2A)
and in the recent peripheral blood (20 years old) (Figure 2B). Loss of
heterogeneity (LOH) of chromosome 11q including the CBL gene was
observed and UPD was also observed in both samples. Figure 2C: Results of
the recent peripheral blood of case 2 (31 years old). The difference is very
small but UPD was found.

Supplemental Figure1. Patients’ pictures at present. A: Case 1, 20-year-old
female. B: Case 2, 31-year-old male. Informed consent was obtained from each
patients. No typical findings are observed of Noonan syndrome.

Supplemental Figure 2. Suspected relationship between \textit{CBL} mutation and 11qUPD according to the frequency of \textit{CBL} mutation and SNP array analysis in both cases. Four types about \textit{CBL} mutation and 11qUPD were considered, Type A: \textit{CBL} mutation with 11qUPD, Type B: \textit{CBL} mutation without 11qUPD, Type C: 11qUPD alone, Type D: no \textit{CBL} mutation and no 11qUPD. Star-shaped indicates \textit{CBL} mutation. We think they would exist as mixed status. Type A would be dominant in case 1, and Type B would be dominant in case 2.
References


Case 1: (CBL, Tyr(Y)371His(H) 1111T>C)

Figure 1. Direct sequencing and Pyrosequencing in Case 1 and Case 2.

Case 2: (CBL, Tyr(Y)371His(H) 1111T>C)
Figure 2. SNP array
Supplemental Figure 1. Pictures at present

A: Case 1. She is 20 years old.

B: Case 2. He is 31 years old.
Supplementary Figure 2

A: CBL mutation + UPD
B: CBL mutation
C: UPD
D: Normal

Case 1

Case 2