Case report: Double L611S/V617L JAK2 mutation in a patient with polycythemia vera originally diagnosed with essential thrombocythemia

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Double JAK2 mutations have rarely been described in myeloproliferative neoplasms (MPNs) and are demonstrated to be associated with the polycythemia vera (PV) phenotype. Here, we first report a case of a PV patient with a de novo double L611S/V617L in cis mutation of JAK2. A 40-year-old woman was admitted to the hospital with massive splenomegaly, multiple splenic infarcts, and abdominal pain. She had a 4-year history of erythrocytosis with an antecedent 10-year history of thrombocytosis before coming to our hospital. She was diagnosed with JAK2 L611S/V617L double-mutant PV after a detailed medical examination in 2017. According to the literature, IFNα therapy can induce clinical, hematological, histopathological, and occasionally molecular remission in individuals with MPNs. Our report demonstrates that combination therapy with ruxolitinib and IFNα can lead to a substantial reduction in JAK2 L611S/V617L double-mutant allele burden.

Keywords: polycythemia vera, essential thrombocythemia, myeloproliferative neoplasm, JAK2 L611S/V617L mutation, CSF1R mutation, phenotype transformation

Introduction

Somatic JAK2 mutations are the most common disease-causing events in patients with myeloproliferative neoplasms (MPNs). Double JAK2 mutations have rarely been described. A JAK2 L611V missense mutation that presented in cis with V617F has been reported in three adult polycythemia vera (PV) patients, which resulted in the JAK2 L611V/V617F double mutation (1). The coexistence of JAK2 L611S and JAK2 V617F variants was described in one adult PV patient who carried the JAK2 L611S mutation in cis with the JAK2 V617F mutation (2). A double L611S/V617F JAK2 mutation was detected in a pediatric patient with PV (3). These double JAK2 mutations were associated
with a PV phenotype. Acquisition of additional mutations in hematopoietic stem cells (HSCs), which creates either compound-mutant JAK2 alleles or a compound-heterozygous state (4), could influence the subclones and affect the disease phenotype in MPNs.

Here, we first report a case of a PV patient with a de novo JAK2 L611S/V617L double mutation in cis. A 4-year history of erythrocytosis with an antecedent 10-year history of thrombocytosis in this case suggests that essential thrombocytemia (ET) and PV are a continuum of one basic disease. Sørensen et al. had reported promising 2-year end-of-study results of the clinical trial investigating combination treatment with ruxolitinib and low-dose pegylated interferon-α2 (PEG-IFNα2), with improved cell counts, reduced bone marrow cellularity and fibrosis, decreased JAK2 V617F burden, and reduced symptom burden with acceptable toxicity in several patients with PV or myelofibrosis (5). In our case, the combination of ruxolitinib and IFNα could improve treatment efficacy and produced a good outcome.

Case description

In January 2003, a woman suffered her first seizure. The initial hemogram showed a high platelet count (517 × 10⁹/L) and a normal hemoglobin concentration, hematocrit level, and white blood cell count. She had no special medical, family, or psychosocial history. A diagnosis of epilepsy was made, and carbamazepine was prescribed by the doctor at the local hospital as seizure treatment. While the diagnosis of epilepsy was made, and carbamazepine was prescribed, the patient was taking carbamazepine for epilepsy from January 2003 to May 2009, her hemogram showed a moderately high platelet count (maximum value: 659 × 10⁹/L). However, she did not have another treatment for abdominal pain. The hemogram showed a high hemoglobin concentration (182 g/L), hematocrit level (57%), platelet count (347 × 10⁹/L), and white blood cell count (12.97 × 10⁹/L). Abdominal MRI revealed giant splenomegaly (thickness, 8.6 cm; length, 23.5 cm), multifocal hemosiderin deposition, and multiple infarcts in the spleen. The patient had a 4-year history of erythrocytosis with an antecedent 10-year history of thrombocytosis before coming to our hospital. Her medical, family, or psychosocial histories were otherwise unremarkable. As such, she was directly transferred from the outpatient clinic to the inpatient department for further medical examination.

After detailed hematologic examination in November 2017, the patient’s bone marrow morphological examination revealed hyperplasia of all three lineages with megakaryocyte hyperplasia with micromegakaryocytes. Reticular fiber staining showed mild increase in reticular fibers (grade 0 myelofibrosis). The bone marrow saMPLs were investigated by flow cytometry. Flow cytometric analysis of HLA-DR, CD33, CD34, CD117, CD10, CD56, CD19, CD5, CD7, CD2, CD11b, CD15, CD13, CD16, CD71, CD41, and CD45 revealed no obvious expression disorder. There was no detection of abnormal blasts and the immunophenotypic characteristics analyzed by flow cytometry were normal. Chromosomal analyses of the bone marrow showed a normal karyotype. BCR-ABL1 fusion gene variants (p190, p210, p230, and rare variants) were not detected by reverse transcription-polymerase chain reaction (RT-PCR). Rearrangements of the PDGFRα, PDGFRβ, or FGFR1 genes were not detected by FISH analysis. There was no detection of a JAK2 V617F mutation, JAK2 exon 12 mutation, MPL W515L mutation, MPL W515K mutation, or a CALR Reticulin (CALR) exon 9 deletion or insertion by PCR with Sanger sequencing. We then performed next-generation sequencing (NGS) for whole exome sequencing (WES) of her bone marrow cells. This revealed de novo double mutations in JAK2, including the JAK2 V617L mutation (c.1849G>C; p.V617L), with a variant allele frequency (VAF) of 44.0%, and the JAK2 L611S mutation (c.1832T>C; p.L611S), with a VAF of 44.7%. She also had a colony-stimulating factor 1 receptor (CSF1R) mutation (c.1460C>T; p.A487V; VAF 46.2%). Neither the double L611S/V617L JAK2 mutation nor other MPN-associated mutations were detected in her buccal mucosal cells by NGS-WES, but the CSF1R A487V mutation was detected with a VAF of 46.2%, and thus proven to be germline. The Integrative Genomics Viewer visualization tool showed that both JAK2 variants were located in the same reads (Figure 1). Cranial MRI revealed ischemic degeneration in the bilateral frontal cortex. Gene mutations associated with epilepsy were not detected.

The patient was diagnosed with JAK2 L611S/V617L double-mutant PV after a detailed medical examination,
and she received continuous treatment with ruxolitinib from November 2017 and combination therapy with ruxolitinib and IFNα from January 2021 (Table 1). During the follow-up time from November 2017, the hemoglobin concentration, hematocrit level, platelet count, and white blood cell count returned to normal 10 weeks after treatment initiation. Significant reductions of 45% and 56% in the cross-sectional area of the spleen were achieved 12 weeks (spleen: thickness, 6.3 cm; length, 17.5 cm) and 2 years later (spleen: thickness, 5.7 cm; length, 15.7 cm), respectively. Alleviation of the constitutional symptoms including excessive and disabling fatigue, early satiety, anorexia, night sweats, abdominal pain and discomfort, and cognitive coMPLiments had improved her quality of life 6 months later. VAFs of both the JAK2 L611S and JAK2 V617L mutations were 44%, 15%, and 11% in November 2017, November 2019, and November 2020, respectively, which suggested that these mutations had occurred in the same clone. The addition of IFNα to ruxolitinib therapy substantially decreased JAK2 L611S/V617L allele burden. After the patient received the combined treatment of IFNα and ruxolitinib for 6 months, the VAF of the double L611S/V617L JAK2 mutation was reduced from 11% to 5% (Table 2). During the follow-up period from November 2017 to November 2020 (ruxolitinib alone), ruxolitinib improved the patient’s splenomegaly and other symptoms, including early satiety, bloating, portal hypertension, fatigue, and undesired weight loss, which improved her overall quality of life.

**TABLE 1** Clinical and biological characteristics of this patient with the double L611S/V617L JAK2 mutation.

| Time                  | January 2003 | January 2003 to May 2009 | November 2010 | November 2010 to September 2011 | October 2011 to October 2017 | November 2017 | November 2017 to December 2020 | January 2021 |
|-----------------------|--------------|--------------------------|---------------|---------------------------------|-------------------------------|---------------|-------------------------------|--------------|
| Diagnosis             | Epilepsy     | Epilepsy                 | ET            | Epilepsy                        | ET switching to PV in 2014? Epilepsy | PV            | PV                            | PV           |
| PLT (×10^9/L)         | 517          | 659                      | 798           | 370–593                         | October 2014: 393             | 347           | Normal                         | Normal       |
| HGB (g/L)             | 142          | Normal                   | 157           | Normal                          | October 2014: 172             | 182           | Normal                         | Normal       |
| HCT                   | 40.0%        | Normal                   | 47.2%         | Normal                          | October 2014: 55.5%           | 57.0%         | Normal                         | Normal       |
| Treatment             | CBZ          | CBZ                      | HU            | HU                              | CBZ                           | Ruxolitinib   | Ruxolitinib                    | IFNα +       |
|                       |              |                          | CRZ           | CRZ                             |                               | CRZ           | Ruxolitinib                    | Ruxolitinib  |

WBC, white blood cell; HGB, hemoglobin; HCT, hematocrit; PLT, platelet; ET, essential thrombocythemia; PV, polycythemia vera; HU, hydroxyurea. CRZ, carbamazepine.
subtypes (12). However, the specific role of the double JAK2 L611S/V617L mutation in our PV patient with a history of ET is unclear.

The presence of different gene alterations within the same gene often results in different biological properties that make it challenging for us to deeply understand the de novo double JAK2 L611S/V617L in cis mutation in PV. This patient experienced a decrease in the JAK2 L611S/V617L double mutation allele burden under the combination therapy of ruxolitinib and IFNα. The de novo JAK2 L611S/V617L double mutation in cis identified in this case might be an oncogenic mutation or driver mutation, as JAK2 V617F was discovered in 2005 as the driver mutation of the majority of non-BCR-ABL1 MPNs (13, 14).

The patient in this study had a germline CSF1R mutation (c.1460C>T; p.A487V). Unlike the reported pathogenic germline CSF1R mutations located within the PTK domain (15), the mutation identified in our patient was located within the immunoglobulin domain. The clinical significance of the CSF1R A487V mutation remains unknown. The dominant transmembrane receptor controlling the differentiation and survival of almost all macrophages regardless of their origins is the colony-stimulating factor 1 receptor. Although the CSFIR A487V germline mutation is known to be nonpathogenic, we speculate that the presence of JAK2 mutations in HSCs and endothelial cells, bone marrow-derived macrophages, or yolk sac-derived brain-specific macrophages with the CSF1R A487V mutation might cooperatively promote multiple splenic infarcts and ischemic degeneration in the bilateral frontal cortex (probably microthrombosis), which likely triggered her seizures. These seizures might be TIAs provoked by the increased number of more adhesive thrombocytes.

HSC heterogeneity underlies the disparate phenotypes of MPNs harboring the same initiating mutation, and malignant transformation of neoplasms might involve a specific subset of stem cells within a heterogeneous stem cell population. In ET, HSCs with JAK2 mutations are more sensitive to IFN signaling to megakaryocyte differentiation and show strong megakaryocyte lineage priming (9). Furthermore, hematopoietic stem and progenitor cells (HSPCs) with JAK2 mutations show a lineage bias towards the megakaryocyte-erythroid fate and contribute to the development of thrombosis (12). IFNα therapy can induce clinical, hematological, histopathological, and occasionally molecular remission in individuals with MPNs. Some results suggested that upon treatment (including long course therapeutic dose of IFNα), the mutant megakaryocyte-primed HSC population

### Table 2: Next-generation sequencing screening result of this patient.

| Gene    | Mutation                                      | Exon  | VAF (2017) | VAF (2019) | VAF (2020) | VAF (2021) |
|---------|-----------------------------------------------|-------|------------|------------|------------|------------|
| JAK2    | NM_004972.3:c.1460C>T(p.A487V) exon14         | 44.0% | 15.2%      | 11.0%      | 5.6%       |
| JAK2    | NM_004972.3:c.1832T>C(p.L611S) exon14          | 44.7% | 15.5%      | 11.4%      | 5.4%       |
| CSF1R   | NM_005211.3:c.1460C>T(p.A487V) exon10          | 46.2% | 47.0%      | 50.1%      | 49.5%      |

VAF, variant allele frequency.

### Discussion

To our knowledge, this is the first report of a double JAK2 L611S/V617L mutation in PV. The JAK2 L611S mutation alone was detected in a child with a germline mutation associated with thrombocytosis (7). The presence of the JAK2 L611S mutation leads to the constitutive activation of JAK2/STAT signaling, even in the absence of erythropoietin receptor expression (8). The JAK2 V617L mutation was detected in an individual with ET, and germline fibroblast testing confirmed the somatic origin of the mutation (9). JAK2 V617L and JAK2 V617I mutations have previously been shown to induce cytokine independence and constitutive downstream signaling in Ba/F3 cells randomly mutated at position 617 of JAK2 (10).

The double JAK2 L611V/V617F mutation increases the activation of JAK2, AKT, and ERK1/2 (but not of STAT5) and is associated with isolated erythrocytosis (1). We speculate that a similar mechanism could underlie the effects of double JAK2 L611S/V617L mutation.

The studies by Van Egeren et al. (9) and Williams et al. (11) showed that the latency between the acquisition of the driver gene mutation and the manifestation of MPN was much longer than that was generally assumed. This provides fascinating insights into the early steps in the pathogenesis of MPN, which might raise opportunities for earlier intervention and provide a new paradigm for cancer development.

Studies have revealed that the JAK2 V617F allele burden, differential signal transducer and activator of transcription activation, order of other somatic mutations, and interindividual genetic variation influence the predisposition to specific MPN subtypes (12). However, the specific role of the double JAK2 L611S/V617L mutation in our PV patient with a history of ET is unclear.
was reduced by promoting apoptosis or quiescence of the mutant cells (12). Clinical evaluation of pegylated IFNα-2a and ruxolitinib in a phase II study of PV and MF showed substantial reductions of JAK2 V617F allele burden with 41% of patients showing a molecular response along with improved cytosis and fibrosis as well as acceptable toxicity (5). These findings demonstrate that pegylated IFNα adds substantial clone suppression to ruxolitinib therapy and that the combination with ruxolitinib improves tolerability of IFNα therapy. These results suggested that our patient would benefit from sustained combination therapy with ruxolitinib and IFNα.

A recent study confirmed that therapeutic targeting of Y-Box Binding Protein 1 (YBX1)-dependent ERK signaling in combination with JAK2 inhibition could eradicate cells harboring mutations in JAK2 (16). The ruxolitinib/nilotinib/prednisone combination showed synergistic inhibitory effects on the JAK/STAT and MAPK signaling pathways in MPN cells (17, 18). Thus, IFNα combined with JAK2 or MEK inhibition might improve therapeutic efficacy.

Given the lifelong trajectories of MPNs in humans and considering the underlying mechanisms in MPNs, the heterogeneity of JAK2 mutant HSCs and HSPCs, and clonal expansion and evolution, studies to increase the understanding of MPNs and to improve our management of MPN patients are needed.

Data availability statement

The original contributions presented in the study are included in the article/supplementary materials. Further inquiries can be directed to the corresponding author.

Ethics statement

The studies involving human participants were reviewed and approved by The Third Affiliated Hospital of Sun Yat-Sen University. The patients/participants provided their written informed consent to participate in this study.

Author contributions

X-QL participated in the clinical care, analyzed the data, and was a major contributor in writing the manuscript. J-JL conducted the literature search and revised the manuscript. C-CL conducted the study, provided patient care, analyzed the data, and contributed with final manuscript drafting. All authors contributed to the article and approved the submitted version.

Funding

The project is funded by GuangZhou Basic and Applied Basic Research Foundation, the grant number is 202201010973.

Acknowledgments

We thank all the investigators, including the physicians, nurses, pathologists, and laboratory technicians in this study.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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