Primary myelofibrosis is not primary anymore since the discovery of MPL<sup>515</sup> and CALR mutations as driver causes of mono-linear megakaryocytic and dual megakaryocytic granulocytic myeloproliferation and secondary myelofibrosis

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Abstract

Primary myelofibrosis (PMF) is a distinct clinicopathological myeloproliferative disease (MPD) not preceded by any other MPD ET, PV, CML, or MDS. With the advent of the MPL<sup>515</sup>, CALR and JAK2<sup>V617F</sup> mutations as the driver causes of three variants of ET (thrombocythemias) the term ET is not essential and PMF is not primary anymore simple because myelofibrosis is a secondary event in all three molecular variants of thrombocythemia and PV. CALR-mutated hypercellular thrombocythemia and MPL<sup>515</sup>-mutated normocellular thrombocythemia are two novel distinct myeloproliferative neoplasms (MPN) disease entities without features of PV and complicated by splenomegaly and secondary myelofibrosis during life-long follow-up. The life expectancy of JAK2<sup>V617F</sup>, MPL<sup>515</sup> and CALR mutated MPNs are related to allele burden, bone marrow cellularity and the degree of anemia, splenomegaly and myelofibrosis during life-long follow-up.

Primary myelofibrosis (PMF) according to the criteria of the Polycythemia Vera Study Group (1975 PVSG) is a clinicopathological entity not preceded by any other MPD PV, CML, or MDS, and characterized by various degrees of anemia, splenomegaly, leuko-erythroblastosis, with tear drop-shaped erythrocytes, and dry tap on BM aspiration due to various degrees of myelofibrosis (MF) or osteosclerosis [1]. In 1977 Silverstein reviewed the characteristic findings in PMF [1,2]. All PMF patients, are usually above the age of 50 years, have large spleens, a leuko-erythroblastic blood reaction, striking teardrop poikilocytosis and dry tap on bone marrow aspiration. Under low-power magnification, a typical marrow section shows stranding, considerable fibrosis, and a few scattered megakaryocytes: at higher power, a dense fibrotic reaction is usually apparent. The marrow tends to be hypocellular in about 85%, normocellular in about 5% and hypercellular in about 10% of PMF patients. Anemia of ineffective erythropoiesis develops in about 60% of PMF patients within 5 to 10 years. Thrombocytopenia and leucopenia related to hypersplenism is seen in 30% and 14% of PMF patients. The diagnostic PVSG criteria for PMF have not been changed in the 2001, 2008 and 2016 WHO criteria for PMF [3-10].
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Personal observations Rotterdam MPD study group 1975-2015

Case 1 in table 1 and related figure shows a typical case of primary pre-fibrotic PMF (prePMF) followed by progressive myelofibrosis, splenomegaly during 20 years follow-up. Sequential bone marrow aspirates and biopsies in 1971 and 1978 showed normal cellularity, no increase of erythropoiesis, selective increase of large megakaryocytes in a bone marrow smears and fine reticulin fibers in a bone marrow biopsy. In 1978, early stage PMF with reticulin fibrosis grade 2 was associated with significant splenomegaly but normal platelet counts and no anemia. Since 1985 full blown primary myelofibrosis (PMF) with progressive anemia and splenomegaly and leuko-erythroblastosis developed and the patient died in 1989 (Table 1). PMF was complicated by a transient episode of erythromelalgic thrombocytopenia at platelet counts above 400x10^9/L at time of progressive PMF during the sequential fibrotic stages of increased reticulin fibrosis followed by collagen fibrosis. Pre-PMF with reticulin fibers grade 1 was diagnosed in 1971 in a 61-year-old man. The spleen had progressed to 7 cm in 1978 and to 13 cm below the costal margin in 1988. Since 1985 he suffered from aspirin responsive burning pain and red swelling of the right hand, fingers and left foot toes (erythromelalgia) at platelet counts of 400 to 500x10^9/L. After correction of increased platelet counts (800x10^9/L) by hydroxyurea to 200x10^9/L, erythromelalgia did not recur because anemia, splenomegaly and thrombocytopenia developed (Table 1).

The PMF case in table 1 could be diagnosed as primary megakaryocytic granulocytic myeloproliferation (PMGM) when the Hannover Bone Marrow classification of the MPDs are applied [2]. Sequential bone marrow biopsies in 1985 and 1989 showed normal cellularity, coarse reticulin fibers, collagen fibrosis (with dry tap on aspiration), and increase of clustered large megakaryocytes. PMGM is misclassified as chronic idiopathic myelofibrosis (CIMF) by Thiele et al. in 1996, as CIMF in the 2001 WHO classification by Thiele & Vardiman and as PMF by Tefferi in the 2008 and 2016 WHO classifications. According to the 2006-2008 ECMP classification Michiels & De Raeve re-defined CIMF and PMF as JAK2 wild type PMGM (Table 2). PMGM has been recognized by Michiels in 2006 as JAK2/MPL wild type Thrombocythemia associated with PMGM since 2006 and as CALR mutated hypercellular thrombocytopenia associated with PMGM since 2014 [11-18].

JAK2^{V617F}, MPL^{515} mutated and JAK2/MPL wild type MPN as the cause of PMF

MPL is the thrombopoeitin receptor (TPOR) on hematopoietic bone marrow cells discovered by William Vainchenker, who also discovered the JAK2^{V617F} as the cause of trilinear myeloproliferative disorders: polycythemia vera (PV) and essential thrombocythemia (ET). About 40% of ET patients are JAK2 wild type. There is a small subgroup of 3% to 8.5% within JAK2 wild type ET patients, who carry an acquired gain

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of function MPL\textsuperscript{515} mutation as the cause of monolinear ET [8-10]. Tefferi from the division of Hematology, Mayo Clinic, Rochester, MN reviewed in January 2012, 1000 consecutive patients with primary myelofibrosis (PMF) seen at Mayo Clinic between November 4, 1977, and September 1, 2011 [3]. The International Prognostic Scoring System (IPSS), dynamic IPSS (DIPSS), and DIPSS-plus were applied for risk stratification. Separate analyses were performed for PMF patients seen at time of referral (N=1000), at initial diagnosis (N=340), and within or after 1 year of diagnosis (N=660). Anno 2012 there were 592 documented deaths and 68 leukemic transformations. Parameters at initial diagnosis vs time of referral included median age (66 vs 65 years), male sex (61% vs 62%), red cell transfusion need (24% vs 38%), hemoglobin level less than 10 g/dL (38% vs 54%), platelet count less than 100 \times 10^9/L (18% vs 26%), leukocyte count more than 25 \times 10^9/L (13% vs 16%), marked splenomegaly (21% vs 31%), constitutional symptoms (29% vs 34%), and abnormal karyotype (31% vs 41%). Mutational frequencies were 61% for JAK2\textsuperscript{V617F}, 8% for MPL\textsuperscript{515}, and 4% for IDH1/2. DIPSS-plus risk distributions at time of referral were low in 10%, intermediate-1 in 15%, intermediate-2 in 37%, and high in 37%. At that time CALR mutation as the driver cause of ET and PMF was not yet discovered. The corresponding median survivals were 17.5, 7.8, 3.6, and 1.8 years vs 20.0, 14.3, 5.3, and 1.7 years for patients younger than 60 years of age. Compared with both DIPSS and IPSS, DIPSS-plus showed better discrimination among high risk groups. Five-year leukemic transformation rates were 6% and 21% in low- and high-risk patients, respectively. Anno 2012 Dr. Tefferi concluded that prognosis in PMF should be assessed after a period of observation after diagnosis rather than at diagnosis. In addition, it seems that about 50% are treated in this manner whereas about 25% are candidates for allogenic stem cell transplant. Roughly 20% can be treated with a JAK2 inhibitor Jakafi/Ruxolitinib.

**CALR mutations as the driver cause in JAK2/MPL wild type ET and PMF**

The molecular etiology of JAK2/MPL wild type ET and PMF remained elusive until Kralovics discovered the calreticulin (CALR) mutations as the driver cause in JAK2/ MPL wild type ET and PMF patients negative for JAK2\textsuperscript{V617F} and MPL\textsuperscript{515} mutation [4]. On behalf of Austrian/Italian hematologists Dr Kralovics detected the CALR mutation in 78 of 311 (25%) ET patients and in 72 of 203 (35%) MF patients and in none of 382 PV patients. The CALR, JAK2\textsuperscript{V617F} and MPL\textsuperscript{515} mutations mutually exclude each other
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since all ET and MF patients with CALR in exon 9 are JAK2 and MPL wild type [4,5]. Out of a large cohort of 289 JAK2/MPL wild type ET 195 (67%) carried one of the CALR mutations. Out of a cohort of 120 JAK2/MPL wild type MF a CALR mutation was detected in 105 (80%). Green and his team in the UK confirmed the high frequency of somatic CALR mutations in 70 to 84% of 120 ET or PMF patients negative for the JAK2/MPL mutation [6]. CALR mutations were present in 110 of 158 JAK2/MPL wild type MPN patients including 80 of 112 (70%) ET patients, 18 of 32 (56%) MF patients and in 12 of 13 patients with advanced PMF. CALR mutations were identified in 10 of 120 (8%) MDS patients (RA in 5 of 53, RA-S in 3 of 27 and RAEB-T in 2 of 27), and in one patient each with chronic myelomonocytic leukemia (CMML) and atypical CML [6].

In a recent large study of Tefferi et al., 254 WHO-defined PMF patients carried the JAK2$^{V617F}$, MPL$^{515}$- and CALR-mutation in 58%, 8.3% and 25% respectively and 8.7% were triple negative [7]. The median overall survival (OS) among 253 WHO-defined PMF patients in 83 CALR-, 21 MPL-, and 147 JAK2-mutated cases and in 22 triple negative cases was 8.2, 4.1, 4.3 and 2.5 years. As compared to CALR wild type MF, CALR-mutated MF patients were younger, had higher platelet count, lower leukocyte count, higher hemoglobin (less anemic) and lower DIPSS-plus score. Tefferi confirmed the findings of Kralovics that CALR-mutated MF patients had a favorable impact on median survival as compared to CALR-negative MF patients whether ASXL1-negative or positive. Among 181 WHO-defined CALR-negative MF patients, the median overall survival was 2.3 years in 55 CALR-negative/ASXL1-positive as compared to 5.6 years in 126 CALR-negative/ASXL1-negative MF patients. Among 146 cases of WHO-defined CALR-positive MF the median survival was 7 years in 20 CALR-positive/ASXL1-positive MF patients as compared to 9.6 years in 126 CALR-positive/ASXL1-negative MF patients.

By including bone marrow histology for ET, PV and PMF since 1980 Michiels & De Raeve changed the concept of PVSG-WHO defined Myeloproliferative Disorders (MPD) into the improved ECMP and WHO-CMP classifications of Myeloproliferative Neoplasms (MPN) [8-10]. The broad spectrum of JAK2$^{V617F}$ mutated trilinear MPN varied from ET, prodromal PV, masked PV, erythrocythemic PV, classical PV, and PV complicated by splenomegaly and myeloﬁbrosis (MF). ET heterozygous for the JAK2$^{V617F}$ mutation is associated with low MPN disease burden and normal life expectancy. JAK2$^{V617F}$ mutation load increases from less than 50% in early stage PV to 100% in combined heterozygous/homozygous or homozygous JAK2$^{V617F}$ mutated advanced PV and post-PV MF. Pre-treatment bone marrow morphology and cellularity distinguish JAK2$^{V617F}$ mutated trilinear MPN from CALR and MPL mutated MPN. The morphology of clustered large pleomorphic megakaryocytes with hyperlobulated nuclei are similar in JAK2$^{V617F}$ ET and PV patients (Figure 1). CALR mutated thrombocythemia shows bone marrow characteristics of PMGM (Table 2). CALR mutated thrombocythemia lacks features of PV and present with hypercellular thrombocythemia featured by dense clusters of immature large to giant megakaryocytes with hyperchromatic bulbous (cloudy) nuclei, which are characteristic for PMGM and not seen in JAK2$^{V617F}$ and MPL$^{515}$ mutated MPN [8,9]. MPL$^{515}$ mutated thrombocythemia typically shows mono-linear proliferation of large to giant megakaryocytes with hyperlobulated staghorn like nuclei in a normocellular bone marrow without features of PV or PMGM. JAK2$^{V617F}$ mutation load in heterozygous mutated JAK2$^{V617F}$ normocellular ET ranges from low to 25%, in prodromal PV from 25% to 50% and increases to above 50% up to 100% in hypercellular heterozygous/homozygous or homozygous JAK2$^{V617F}$ mutated PV and advanced PV with myelofibrosis. The CALR and MPL$^{515}$ mutation load is low in pre-fibrotic stages and rises from values below 25% to maximal values of around 50% in fibrotic stages of MPN [8,9]. JAK2$^{V617F}$, CALR and MPL$^{515}$ allele burden slowly increases together with the degree of splenomegaly, myelofibrosis and constitutional symptoms during life-long follow-up.
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Bone marrow histology in MPL<sup>515</sup> mutated normocellular ET

Bone marrow histology from a patient with JAK2 wild type ET carrying the MPL<sup>515</sup>L mutation displayed clusters large megakaryocytes with a greater number of giant megakaryocytes with hyper-lobulated stag-horn nuclei in a normal cellular bone marrow and no increase of erythropoiesis (Figure 2) [10]. Michiels & De Raeve described the essential differences in bone marrow histopathology features of differential diagnostic significance between patients with MPL<sup>515</sup> mutated (N=12) versus JAK2<sup>V617F</sup> mutated MPN (N=12) (Figure 2) [8-12]. The presence of clustered small and giant megakaryocytes with deeply lobulated stag-horn like nuclei (Figure 2) in ET carrying the MPL<sup>515</sup> mutation are not seen in JAK2<sup>V617F</sup> positive prodromal PV, and classical PV (Figure 1). The pleomorphic medium to large megakaryocytes in JAK2<sup>V617F</sup> mutated ET and PV in bone marrow smears and bone marrow biopsy were similar in size and pleomorphy (Figure 1). There is local increase of erythropoiesis in areas of loose clustered pleiomorphic megakaryocytes in prodromal PV (Figure 1), which is not seen in MPL<sup>515</sup> mutated ET (Figure 2). JAK2 wild type MPL<sup>515</sup> mutated ET have no clinical, laboratory and bone marrow features of prodromal PV at diagnosis, do not evolve into PV during follow-up, and have normal LAP score, serum EPO and ferritin levels [1,5]. Laboratory and bone marrow histology evaluations have the diagnostic potential to separate the JAK2<sup>V617F</sup> mutated ET and prodromal PV with increased LAP score, low serum EPO and pleomorphic megakaryocyte morphology from MPL<sup>515</sup> mutated ET with normal LAP score and serum EPO and giant mature megakaryocytes with staghorn-like nuclei [1,3,4]. The natural history of MPL<sup>515</sup> patients clearly differ from JAK2<sup>V617F</sup> mutated ET and is typically featured by progressive anemia with increased reticuline fibrosis in a normocellular or may progress into refractory anemia in a hypocellular bone marrow and a high risk of MDS or leukemic transformation [10,11]. Pich et al., performed in 2012 a comparative bone marrow histology study in 103 WHO defined ET patients and demonstrated that the values for hemoglobin, hematocrit, and erythrocyte count, and bone marrow erythropoiesis were significantly lower in 44 JAK2<sup>V617F</sup> wild type ET as compared to 59 JAK2<sup>V617F</sup> mutated ET [13].
The megakaryocytes in JAK2 V617F wild type patients were large to giant with more pronounced hyperlobulation of nuclei, whereas the megakaryocytes in JAK2 V617F mutated ET and PV patients are medium sized to large (pleomorphic) with only a few giant forms [13].

**Bone marrow histology in CALR mutated JAK2 wild type ET and MF**

Michiels & De Raeve studied between 2014 and 2018 twelve consecutive newly diagnosed CALR positive ET cases. We found a spectrum of two main stages of normocellular (40%-60%) ET and hypercellular bone marrow (60-80%) ET characteristics as the presenting feature of pre-fibrotic ET and hypercellular ET associated with a PMGM bone marrow picture (Table 2). Bone marrow histology in pre-fibrotic CALR normocellular ET (Figure 3) and in early fibrotic CALR myelofibrosis (MF, Figure 4) showed ‘dysmorphic’ large to giant megakaryocytes with definite abnormalities of maturation with bulky (bulbous) hyperchromatic nuclei and some disturbances of the nuclear cytoplasmic ratio consistent with CALR mutated PMGM, which are not seen in MPL515 mutated ET and also not in JAK2 V617F mutated ET, prodromal PV and classical PV. Pich performed in 2018 a comparative analysis of bone marrow cellularity, reticulin fibrosis, and the number, clustering and morphology, size of megakaryocytes and nuclei in 90 cases of WHO defined ET caused by the driver mutation JAK2 V617F, CALR or MPL515 at time of diagnosis [14]. The JAK2 V617F mutation was found in 58.9%, CALR in 28.9%, and MPL in 4.4% of the cases, and 7.8% were triple-negative. JAK2 V617F mutated ET showed increased bone marrow cellularity due to significant increase of erythropoiesis (Figure 3). MPL515 mutated ET showed the highest number of clustered large mature megakaryocytes with hyperlobulated staghorn-like nuclei in a normocellular or reduced cellular bone marrow (Figure 2). The bone marrow in CALR-mutated ET is normocellular or hypercellular due to dual megakaryocytic granulocytic myeloproliferation with no increase or reduced erythropoiesis CALR mutated ET in the Pich study showed dense clusters of large to giant dysmorphic megakaryocytes with hypo/hyperlobulated hyperchromatic bulbous or cloud-like nuclei similar as observed by Michiels & De Raeve (Figures 4-6) [10,11,15-18].
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Figure 3: Bone marrow biopsy showing increased cellularity (70%) due to increased erythropoiesis and megakaryocytic (EM) proliferation with loose clusters of large megakaryocytes with hyper-lobulated nuclei in WHO defined JAK2V617F ET (a) in two studies of Piche 2008 and 2018. Dense clusters of large to giant megakaryocytes with hyperchromatic hypohyper-lobulated bulbous (cloudy) nuclei and relative reduction of erythropoiesis in WHO defined CALR mutated ET in the study of Pich 2018. Bone marrow biopsy with fields of normocellular and reduced cellularity (h) and the presence of dense clustered large to giant megakaryocytes with hyper-lobulated staghorn-like nuclei (i and j) in WHO defined MPL515 mutated ET in the study of Pich 2018.

Figure 4: Bone marrow biopsy showing a normocellular bone marrow with loose and dense clusters of immature large to giant megakaryocytes with hypo/hyper-lobulated hyperchromatic cloudy (bulbous) nuclei in CALR mutated ET at platelet counts around 1000x10^9/L and no increase of reticulin fibers grade 0/1. Source Michiels & De Raeve.

Figure 5: Bone marrow biopsy showing some increase of cellularity (60%) due to increase of primary megakaryocytic and granulocytic myeloproliferation (PMGM) with loose and dense clusters of immature large to giant megakaryocytes with hyperchromatic cloudy (bulbous) nuclei in CALR thrombocythemia (platelets 1296x10^9/L) and splenomegaly (spleen size 16 cm on echogram) complicated by acquired von Willebrand syndrome in the study of Michiels 2017.
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Conclusion

With the advent of the MPL515, CALR and JAK2V617F mutations as the driver causes of three distinct variant of ET (thrombocythemia) the term ET is not essential and PMF is not primary anymore simple because myelofibrosis is a secondary event in all three molecular variants of thrombocythemia and PV [15-18]. CALR-mutated hypercellular thrombocythemia and MPL515-mutated normocellular thrombocythemia are two distinct JAK2V617F wild MPN disease entities without features of PV and complicated by splenomegaly and secondary myelofibrosis during life-long follow-up. In conclusion, the term PMF is not primary anymore and should be replaced by CALR-thrombocythemia, MPL515-thrombocythemia and secondary MF similar as in JAK2V617F mutated post ET-MF and post-PV-MF as a secondary event [15-18].

References


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