Essential thrombocythemia: Biology, clinical features, thrombotic risk, therapeutic options and outcome

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Abstract

Essential Thrombocythemia (ET) is currently classified as a Philadelphia negative myeloproliferative neoplasm (MPN) together with polycythemia vera (PV) and primary myelofibrosis (PMF); the latter can be further divided in pre-fibrotic primary myelofibrosis (pre-PMF) and overt myelofibrosis, as listed in the revised 2016 World Health Organization classification of myeloid malignancies (WHO 2016). Overall, respect to the others MPNs, ET is characterized by favorable prognosis, lower life expectancy if compared to the control population, increased risk of thrombohemorrhagic complications along with possible evolution in myelofibrosis and leukemic transformation. In this review the authors will review current knowledge on biology, clinical aspects, prognosis and stratification of thrombotic risk, therapeutic options and outcome in ET patients.

Introduction

The revised 2016 World Health Organization classification for hematological malignancies and acute leukemia defines the criteria for the diagnosis of essential thrombocythemia. 4 major criteria and 1 minor criterion have been identified. Respectively, a platelet value greater than or equal to 459,000x10⁹ L⁻¹; bone marrow biopsy showing proliferation mainly of the megakaryocyte lineage with increased number of enlarged, mature megakaryocytes with hyperlobulated nuclei; no significant increase or left shift in neutrophil granulopoiesis or erythropoiesis and very rarely (grade 1) increase in reticulin fibers; exclusion of WHO criteria for BCR-ABL, CML, PV, PMF, myelodysplastic syndromes, or other myeloid neoplasms; presence of JAK2, CALR, or MPL mutations and a clonal marker or absence of evidence for reactive thrombocytosis. Diagnosis of ET requires meeting all major criteria or the first three major criteria and the minor criterion [1]. The estimated ET annual incidence in the United States is 2.5 cases per 100,000 whereas the prevalence is estimated to be 24 cases per 100,000. Furthermore, other studies estimated the annual ET incidence in Western countries in 0.2–2.5 cases per 100,000 with a prevalence of 38-57 cases per 100,000 [2,3]. Moreover, the incidence is higher in women than men population with an approximate ratio of 2:1 [4], the median age at diagnosis is 60 years with a 20% of patients receiving a diagnosis under 40 years of age. According to some studies over the last few years there has been a decrease in the age of ET diagnosis (56 years versus 60 years.) [5,6]. In our experience, in 234 consecutive patients diagnosed with ET between 1997 and 2019, 36.75% were diagnosed under 61 years, 26.07% under 51 years and 16.67% under 41 years. Interestingly, ET is characterized by overall favorable prognosis if compared to the other MPNs (but life-expectancy in ET is inferior to the control population) with an expected survival of 19.8 years (compared to 13.5 years in PV and 5.9 years in PMF). Cumulative incidence of blast transformation is lower in ET (3.8%) than in PV (6.8%) and PMF (14.2%). Moreover, similar cumulative incidence of fibrotic transformations is lower in ET (9.2%) than PV (21%) [7].
Diagnosis and mutational status

The remarkable knowledge acquired over the last 15 years in the field of molecular biology has improved the diagnostic ability of Philadelphia negative myeloproliferative neoplasms. V617F, janus kinase 2, located on chromosome 9p24, was first reported in 2005 and represents the most frequent mutation with an estimated frequency, in ET patients, of 50–60% [8-11]. Furthermore, in 2006 and 2013 further driver mutations have been identified, respectively, in MPL and CALR genes (calreticulin). MPL, gene, myeloproliferative leukemia virus oncogene, is located on chromosome 1p34, also known as the thrombopoietin (TPO) receptor. Mutations on MPL are present in about 5% of patients with ET [12,13]. Finally, CALR gene, is located on chromosome 19p13.2 [14,15] and according to some authors, CALR mutations are closely related to platelets production. Unlike JAK2 and MPL mutations, CALR mutations are present in patients with thrombocytosis other than ET, such as PMF or refractory anemia with ring sideroblasts and marked thrombocytosis. The most frequent CALR mutations are, respectively, a 52 bp deletion (type 1) and a 5 bp insertion (type 2) [15,16]. In our series of 234 patients with ET, 71.79% showed V617F-JAK2 mutation, 8.9% CALR mutations, 1.71% MPL mutations [17], contrariwise, 52% of our ET patients are triple negative. Driver mutations were considered to be mutually exclusive. However, Mansier and colleagues have recently reported that additional mutations in CALR or MPL co-exist in approximately 10% of patients with low JAK2 burden. Unfortunately, the clinical significance of this double mutation is still unclear [17]. JAK2 mutated ET patients tend to be older, show higher hemoglobin levels and white blood cell counts as well as a lower platelet counts and serum EPO levels than patients without mutation [18]. Moreover, JAK2 mutated ET patients are more likely to develop thrombosis [19,20]. Furthermore, patients with CALR mutations have a different phenotype than V617F-JAK2 or MPL mutated patients. Indeed, they have a greater platelet value, a lower hemoglobin values and low leukocyte levels along with a lower thrombotic risk [21,22]. Several CALR variants, related to different phenotypes, have been identified. In particular, the type1-like variant is associated with a greater risk of myelofibrosis transformation, while the type2-like variant is associated with a more indolent clinical course [23]. The mutational landscape of MPNs is actually very complex. Indeed, alongside the aforementioned driver or phenotypic mutations (sufficient to determine the clinical phenotype of the disease), other sub-clonal mutations have been also identified. (TET2, ASXL1, CBL, IDH and IKZF1). Sub-clonal mutations can occur more often in conjunction with phenotypic mutations but can temporally either precede or follow them. To date, sub-clonal mutations do not have a clear diagnostic value, in fact they are present in myelodysplastic syndromes as well as in other hematological neoplasms such as acute leukemia. Interestingly, these mutations have been reported to have an important negative prognostic value in primary myelofibrosis (PMF). Their presence is associated with lower survival and a greater risk of acute leukemia evolution. Such mutations are identified in lower frequencies in ET and PV if compared to MF [24-27]. In 2013, the study reported by Nangalia et al. highlighted a median of 6.5 mutations in patients with ET if compared to 13 mutations found in patients with PMF. The most frequent sub-clonal mutations were found in DNMT3A, TET2, and ASXL1 [28]. Moreover, in a cohort of 181 ET patients, 46% were found to have somatic mutations including TET2 (13%), ASXL1 (11%), DNMT3A (6%), SF3B1 (5%), CEBPA (4%), along with mutations in TP53, SH2B3, EZH2, and CSF3R (each 2%). The impact on prognosis is not clear and today the use of next-generation sequencing in ET is still not routine neither recommended by guidelines [29].

Clinic feature

In MPN symptomatic burden is often severe and affect the majority of patients with the disease. Some authors have conducted an online survey on 1179 patients with MPN to quantify the burden of illness; results indicate that constitutional and splenomegaly associated symptoms were predominant (70% of patients) and compromised quality of life. Additional signs included fatigue (81%), pruritus (52%), night sweats (49%), bone pain (44%), fever (14%), and weight loss (13%). Most often patient’s experienced profound fatigue in excess of age matched controls and required assistance for daily activities or severe disabilities (34.5%) and reported MPN-associated disability (11.2%). Subsequently, the same authors attempted to validate a widely applicable 18-item tool (evaluation of the symptoms of myeloproliferative neoplasias Module [MPN-SAF]), coordinated with the Brief Fatigue Inventory, to evaluate the symptoms of myelofibrosis, essential thrombocytemia and polycythemia vera between potential cohorts in the United States, Sweden and Italy. Patients reported that symptoms associated with MPN disease were severe and frequent among all three MPNs. In a cohort of 161 ET patients the most frequently reported symptoms were fatigue (90.3%), numbness (58.8%), insomnia (58%), sad mood (57.3%), vertigo (56.1%), concentration problem (55.8%), inactivity (53.7%), early satiety (53.2%), night sweats (51.3%), sexual problem (51.0%), headache (47.1%), abdominal discomfort (45.3%), bone pain (45.2%), cough (41.4%), itching (40.6%), abdominal pain (38.2%), weight loss (23.4%) and fever (17%). Patients with ET such symptoms present with a slightly lower frequency compared to PV and PMF and are significantly less severe [30]. In figure 1, we reported the symptoms and the relative frequency in a cohort of 234 ET patients enrolled at our institution.

Thrombotic risk and bleeding complication

An increased risk of vascular complications over time is the main clinical feature of ET. In a study conducted on
Today, the current classification of thrombotic risk in patients and Barbui adding the negative effect of MPL mutation [40].

Enhancement of IPSET-Thrombosis was proposed by Tefferi of ET diagnosis, were reported in 231 cases (17.8%) [31]. Risk factors (10/59) compared to patients with only one cardiovascular risk factor (18/92) and compared to patients with more than one cardiovascular risk factors (34/83) (p=0.0009). In table 1 we listed the cardiovascular risk factors in a cohort of 234 ET patients. The frequency of bleeding complications in patients with ET varies in different studies. In a 2012 international study conducted on 891 ET patients, 55 major bleeding events occurred (6.17%) [41], previously, several reports indicated incidence of bleeding events in 36-37% of patients with ET. In 2005, Elliott and Teffery reported an incidence of bleeding events equal to 0.33% per person per year [42]. The most frequent hemorrhages concern the gastrointestinal and urogenital apparatus but may occur in several sites among which intracranial hemorrhages are particularly dangerous. Bleeding episodes are correlated with extreme thrombocytosis (platelet >1000-1500*10^9/L) and may be associated with acquired Von Willebrand disease (AVWS). The mechanism that determines the AVWS consists in the absorption of large Von Willebrand multimer by the platelet’s membranes. For these reasons, extreme thrombocytosis is an indication for cytoreductive therapy [43,44].

Disease progression

In ET patient’s transformation to AML and/or post-ET myelofibrosis (post-ET MF) are rare events. Risk of transformation to acute leukemia was reported at 2–3% at 10 years and 5% at 15 years [45]. Subsequently, following the review of the case definition carried out by the WHO in 2016, the distinction between myelofibrosis in the pre-fibrotic phase and ET significantly reduced the incidence of transformation in post-ET myelofibrosis and acute leukemia [46]. When ET is confirmed on the basis of bone marrow morphology, the risk of leukemic transformation is estimated at 0.7% at 10 years and 2.1% at 15 years [47]. As for the evolution in myelofibrosis one of the largest longitudinal cohort studies reported the cumulative probability of myeloid disease transformation to be 3.8% at 10 years, 19.9% at 20 years, and 28.9% at 30 years [48]. But even in this case, the distinction between ET and pre-PMF could have an important impact on the incidence of transformation into myelofibrosis. Some risk factors seem to be related to leukemic transformation in patients with ET; the presence of anemia, platelets count <1000,000, advanced age and leukocytosis as well as reticulin grading and bone marrow cellularity [47,49] seem to play a pivotal role. Mutation subtype may play a role in the risk of transformation to AML or post-ET MF, but more data are still needed. For AML, while one study highlighted that JAK2-mutated patients have

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a higher risk of transformation to AML than CALR-mutated patients [50], these differences did not prove any statistical significance.

**Therapeutic option**

In 2018, the panel of experts from the European LeukemiaNet consortium (ELN) published the therapy recommendations for patients with MPNs. Based on the results of retrospective studies, the use of low-dose aspirin is recommended for high-risk patients according to the IPSET thrombosis classification. In low-risk or intermediate-risk patients aspirin is recommended at low doses if age > 60 years and with uncontrolled cardiovascular risk factors or in the presence of JAK2 mutation [51]. Of particular interest is a study carried out on a cohort of 433 patients with low-risk ET in which the use of aspirin at low doses, in patients with bleeding or a platelet count greater than 1,500 x 10^9 L. Cytoreduction or following a major or hemorrhagic thrombotic event and/or a platelet count greater than 1,500 x 10^9 L is recommended. The LeukemiaNet panel also recommends the use of anagrelide or rINFalfa as a second-line therapy is intolerance or insufficient response to hydroxyurea therapy, or in patients with low-risk ET in which the use of aspirin was of insufficient quality, and the risk-benefit ratio unfavorable. Contrariwise, in patients with low-risk ET, cytoreductive therapy is not recommended. In case of intolerance or insufficient response to hydroxyurea, the use of anagrelide or rINFalfa as a second-line therapy is recommended. The LeukemiaNet panel also recommends the early initiation of cytoreductive therapy in patients who switch to the high-risk category following the age of 60 years or following a major or hemorrhagic thrombotic event and/or a platelet count greater than 1,500 x 10^9 L. Cytoreduction may be also required for progressive myeloproliferative (e.g., increasing splenomegaly) or uncontrolled ET-related systemic symptoms [51].

Other drugs have been experimentally used in patients resistant or tolerant to hydroxyurea. A total of 18 patients in two sequential cohorts received an initial dose of 7.5mg or 9.4mg of imetelstat (Telomerase inhibitor) per kilogram of body weight intravenously once a week until attainment of a platelet count of approximately 250,000 - 300,000 per cubic millimeter. The primary end point was the best hematologic response. Imetelstat induced hematologic responses in all 18 patients, and 16 patients (89%) had a complete hematologic response. Molecular responses were seen in 7 of 8 patients who were positive for the JAK2-V617F mutation (88%; 95% confidence interval, 47 to 100). CALR and MPL mutant allele burden was also reduced by 15 to 66%. However, imetelstat therapy showed a high risk of grade 3-4 neutropenia, thrombotic and abnormal liver function test [54]. The histone deacetylase inhibitors vorinostat and gavinostat were able to induce some signals of spleen reduction, symptoms relief and hematologic response in a substantial proportion of patient [55,56]. Verstovsek et al., used ruxolitinib in patients with ET resistant or intolerant to hydroxyurea. The results of this study demonstrated that patients can achieve clinically meaningful and durable reduction in platelet and WBC counts and improvements in ET-related symptoms with ruxolitinib treatment [57]. According to other authors, since the ET, especially after the revision of the case definition by the WHO in 2016, has a relatively benign course and the evolution in MF PET is quite a rare event, the use of Ruxolitinib appears questionable [58].

**Pregnancy**

Out of 234 patients with ET visited in our center, 33 of 161 women (20.49%) are fertile at the time of diagnosis, this implies the possibility to manage a pregnancy in this setting of ET patients. In this patients the risk of first-trimester fetal losses (about 3.5-fold) and placental complications (e.g. abruption, pre-eclampsia) is increased if compared to healthy women [59-61]. These events are possibly mediated by decidual thrombosis. Risk factors include previous pregnancy complications and possibly the presence of a JAK2-V617F mutation. Venous thrombosis may occur, particularly in the postpartum period, and the risk is higher in patients with a history of vascular events [62]. Furthermore, the incidence of postpartum hemorrhage is estimated at 9%. With regard to fetal complications, the perinatal mortality rate was 17 per 1000 live and stillbirths, and 22% were below the 10th percentile in weight, while 13% required admission to the neonatalcare unit [63]. In 2011 the panel of LeukemiaNet consortium defined treatment approach during pregnancy, although evidence about the impact on outcome is weak. For low risk patient low dose aspirin is recommended; moreover, prophylactic low molecular weight heparin dose after delivery, until 6 weeks postpartum, is recommended [64]. For high risk patient with previous major thrombosis events or severe pregnancy complication a low molecular weight heparin dose throughout pregnancy is recommended (stop aspirin in case of bleeding complications; if platelet count > 1,500x10^9/L: consider rINFalfa , if previous major bleeding; avoid aspirin and consider interferon alfa to reduce thrombocytosis) [64]. In 2014, consensus statement of the Haemostasis Working Party of the German Society of Hematology and Oncology (DGHO), the Austrian Society of Hematology and Oncology (ÖGHO) and Society of Thrombosis and Haemostasis Research, offer similar recommendations [65].
Survival

Overall, ET is considered as an indolent disease. The 15-year survival is 75%, with risk factors for poor survival identified as lower hemoglobin level than normal, age 60, leucocyte count ≥15x10^9/L, smoking, diabetes, thrombosis, thrombocythemia, and male gender [66]. The International Prognostic Score for ET (IPSET) is a prognostic model that was developed to predict survival at diagnosis. Patients identified as intermediate risk (total score 1–2) had a median survival of 24.5 years, while patients identified as high risk (total score 3–4) had a median survival of 13.8 years [67].

Familial ET

Rare cases of ET in the same household are reported. Predisposition haplotypes, including 46/1, [68] and predisposition alleles have been identified and seem to confer an increased risk of developing not only MPN but also JAK2-V617F clonal hematopoiesis [69-71]. Maffioli, Mora et al. analyzed 22 cases of familial MPNs, for a total of 45 patients. Within this cohort of 22 familial MPN pedigrees disease phenotype and genotype are heterogeneously distributed and occurrence is more frequently vertical [72]. Among our patients we have identified a unique case of familial ET. The case concerns a brother and a sister who have in common only one parent (the father); in both cases the diagnosis of ET has been done according to the WHO 2016 criteria, of particular interest is the presence of a CALR mutation in the male and a JAK2-V617F mutation in the sister (unpublished data). Unfortunately, it has not been possible to perform further molecular testing to the parent as he has been dead for several years. We recommend that physicians interview all patients with ET to determine whether there is a family history of thrombocytosis or MPN [73].

Conclusion

ET is a relatively rare chronic myeloproliferative disease, characterized by a favorable prognosis with similar or slightly lower survival than the general population. The distinction made in 2016 by the WHO in pre-PMF and myelofibrosis overt has further improved the prognosis as several cases of ET have been reclassified to pre-PMF which has a slightly worse prognosis. The mutational state and the thrombotic risk may influence the therapeutic choice. The knowledge acquired in the last two decades, especially in terms of molecular biology, have greatly improved understanding of the pathogenesis and the ability to diagnose negative Philadelphia myeloproliferative diseases. To date, however, still several cases of ET are diagnosed after the thrombotic event. Therefore, early diagnosis could further minimize the risk of thrombotic even and improve the outcome.

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