Erythromelalgic Thrombotic Thrombocythemia (ETT) and Hemorrhagic Trombocythemia (HT) in Patients with Essential Thrombocythaemia (ET) and Polycythaemia Vera (PV)

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Abstract
We analysed between 1908 and 1985 of two hundred cases of thrombocythemia from the literature between 1908 and 1985. Erythromelalgic, acrocyanotic ischemia and digital gangrene, cerebral or coronary ischemic events (erythromelalgic thrombotic thrombocythemia: ETT) in 99 thrombocythemia patients (essential thrombocythaemia: ET N=67 and polycythaemia vera: PV N=32) occurred at platelet counts in excess of 400 × 10^9/L. Hemorrhagic complications at time of presentation of hemorrhagic thrombocythemia (HT) in 100 HT patients included melana and/or hematemesis, skin bleedings (bruises, subcutaneous hematomas, painful hematomas after trauma, ecchymoses, but not petechiae, epistaxis and gum bleeding and secondary bleeding after trauma or surgery. ETT occurred at an early stage of thrombocythemia with a mean platelet count of 1110+477 × 10^9/L, whereas HT occurred at significantly higher platelet counts in excess of 1000 × 10^9/L with a mean platelet count of 2016±1070 × 10^9/L. HT patients frequently had a previous history of ETT or paradoxical occurrences of both thrombosis and bleeding (ETT/HT). The degree of thrombocythemia determined the sequential occurrence of ETT, HT in ET and PV patients. Hypersensitive thrombocythemic platelets (sticky platelets) are involved in the etiology of platelet-mediated thrombosis at platelet counts above 400 × 10^9/L whereas bleedings at high platelet count is related to an acquired von Willebrand syndrome processes on top of a blood clot retraction disturbance with erythrocyte fall out, that causes painful subcutaneous hematomas with a central swelling (clot) after a blow, trauma and secondary bleeding after surgery in thrombocythemia patients.

Keywords:
Erythromelalgia and Arterial Thrombosis in ET and PV 1878-1980

Mitchel [1] reported on arare vaso-motor neurosis of the extremities and labeled it as erythromelalgia (erythros=red, melos=extremity, and algos=pain) in the footnote of his article. Mitchell demonstrated that typical erythromelalgia in one or more toes may progress into unbearable intense burning, throbbing, and aching pain, complicated by neurologic ischemic attacks and visual disturbances in one and the same patient. The occurrence of erythromelalgia in polycythaemia is very well known in the literature since 1908 [2-5], (Norman and Allen). Oder in 1908 noted that there has been cases of PV in which pain in the hands and feet with the extreme congestion is suggestive for the erythromelalgia of Weir Mitchell [1]. In 1929, Oppenheimer recognized that the erythromelalgic symptom complex was often the earliest symptom of PV and even may precede PV for more than 10 years. He distinguished two types of peripheral circulatory symptoms at presentation of PV:1) the erythromelalgic syndrome, and 2) occlusion of peripheral arteries by thrombi (thromboangiitis obliterans) in later stages of the disease. Two of his patients suffered from erythromelalgia and recurrent attacks of dizziness and had platelet counts of 560 to 620 × 10^9/L. The peripheral microvascular symptoms at diagnosis of PV in 100 cases from the study of Brown and Griffin [4] were erythromelalgia in 6, acroparesthesias of burning type in 13, peripheral digital gangrene or ischemia in 8 (erythromelalgic ischemic complications 21%). Similar observations of erythromelalgic microvascular ischemia, amorosiosis fugax, and transient ischemic attacks and major arterial and venous thrombotic events were observed in the studies of Edwards and Cooley [6], and of Barabas et al. [7] (Table 1).

In 1940 Damasek and Henstell described recurrent burning pain and cyanosis of several toes of the right foot with gangrene of the third toe diagnosed as thrombo-angiitis obliterans in PV at platelet count above 1000 × 10^9/L and erythrocyte count above 6 × 10^12/L [8]. These observations prompted Damasek in 1940 to suggest the possibility of "platelet thrombophilia" with multiple small peripheral vascular thromboses (thromboangiitis obliterans). In 1937 Nygaard and Brown [9] described a 47-year-old man with recurrent burning, painful, red toes during a 10-year follow-up ultimately complicated by blister formation and gangrene of the right fourth and fifth toes in a case with persistent increase of peripheral blood platelet count in myeloproliferative thrombocythemia. Annets and Tracy [10] and Vreeken and van Aken [11] described microvascular painful ischemia of the toes as first manifestation of ET. Alarcon-Segovia et al. [12] recognized aspirin-sensitive erythromelalgia as a clue to the early diagnosis of ET and PV, but phlebotomy did not improve the erythromelalgic burning pain (cases 5, 6, 7 in Table 2).

Discovery of Aspirin-responsive Erythromelalgia in Thrombocythemia

Between 1970 and 1973 we discovered that aspirin responsive burning red extremities (erythromelalgia) is causally related to thrombocythemia in ET and PV patients (Figure 1). Acroparesthesias, for example tingling, "pins and needles" sensations, and numbness
in the toes or fingers, usually precede the disabling and burning distress, and aspirin promptly relieves pain for approximately three days [13-15]. A few cases of erythromelalgia associated with primary (essential) thrombocythemia has been recognized in the 1970s [16-20]. Between 1974 and 1977 Weatherley-Mein clearly documented that erythromelalgia was the presenting feature of primary (essential) thrombocythemia. Preston et al. [21] interpreted the burning painful and ischemic digital circulation disturbances as thrombotic thrombocythemia. Between 1975 and 1981 we demonstrated that aspirin responsive erythromelalgia is a pathognomonic manifestation of thrombocythemia in ET and PV patients is caused by platelet-mediated inflammation and thrombosis in the end-arterial circulation (Figure 1). Aspirin relieves the peripheral, cerebral and ocular ischemic disturbances by irreversible inhibition of platelet cyclooxygenase (COX) activity and aggregation ex vivo and vitamin K antagonist (Warfarin), dipiridamol, ticlopedine, sulfinpyrazone and sofium salicytate have no effect on platelet COX activity and are ineffective in the treatment thrombocythemia-specific manifestations of erythromelalgia and migraine-like ischemic attacks (MIAs) (Figure 2), [13-15]. Aspirin responsive erythromelalgia in thrombocythemia appeared to be utterly dissimilar from aspirin resisten incurable primary erythermalgia and secondary erythermalgia in SLE or RA patients [22-24]. Erythromelalgia disappeared by correction of platelet counts to normal (<350 × 10^9/L) in ET and PV, but not by correction of hemoglobin to normal by bloodletting in PV patients [13,15].

**Delineation of HT in Various Myeloproliferative Disorders 1923-1980**

Minot and Buckmann recognized in 1923 [25] a bleeding tendency of ecchymosis, nose bleeds, and secondary bleedings but absence of pectechiae in association with persistent increased platelet count in a case of polycythemia vera. Rosenthal [26] recognized a group of post-splenectomy patients with persistent increase of platelet counts in excess of 1000 × 10^9/L associated with hemorrhagic or thrombotic complications. Thrombocythemia may be associated with both a hemorrhagic diathesis as well as thrombotic complications in one and the same patient [27,28]. Epstein and Goedel [29] introduced the term hemorrhagic thrombocythemia (HT) for a disease entity featured by recurrent gums and nose bleeds associated with very high platelet count of 1735 × 10^9/L and atrophic spleen in a 56 year old man This case was previously described as a case with "hochgradigen Thrombocytenvermehrung=high grade platelet increase" by Epstein and Kretz [30]. Similar case histories are described in the literature under a diversity of terms including hemorrhagic thrombocthemia (Uotila, Reid, Mortensen, Woodrow and Lope, Hardisty and Wolf,
Binswanger et al., Koller and Bounnameux, Spaet and Bauer, Fountain, Gunz, Shaw, Fountain and Losowsky, Webb et al., Cronberg et al., Hall, Ohler et al., Bensiger et al., and the Polycythemia Vera study Group, PVSIG [31-45]) primary hemorrhagic thrombocythemia, [28,46-48]) primary thrombocythemia [49], essential thrombocythemia [50-53], persistent thrombocythemia [54], thrombocythemia [55-57], post-splenectomy thrombocythemia [58], thrombocytosis [59], hemorrhagic thrombocytosis [60,61], myelose hyperthrombocytaire [62] and hyperplaquettose [63].

According to Reid [34], Fanger et al. [55], Ozer et al. [28] and Silverstein [48] HT may be part of or clearly associated with polycythemia vera, chronic myeloid leukemia, or myelofibrotic myeloid metaplasia. Others labeled HT as a distinct disease even when associated with other myeloproliferative diseases like polycythemia vera and myelofibrotic myeloid metaplasia [31]. According to Dameshek in 1950 [64], the cause of the hemorrhagic diathesis of polycythemia is obscure. Bleedings do not ordinarily occur spontaneously but only in response to trauma as with a blow or following operative procedures. Extreme degrees of postoperative hematomas are common, and excessive bleeding occurs after dental extractions, tonsillectomy, polypectomy and similar procedures. Based on the analysis of 50 cases from the literature and 5 of his own, Gunz [42] defined hemorrhagic thrombocythemia (HT) as clinical syndrome of recurrent spontaneous hemorrhages often preceded by thromboses, extremely high platelet count in excess of 1000 × 10^9/L, frequently splenomegaly, and hypochromic anemia with a tendency towards polycythemia between hemorrhages. Inclusion and exclusion

Bleeds from nose gums and gastrointestinal tract were most frequent followed by bruises and bleedings after trauma or surgery, whereas “thrombocytopenic purpura” were never seen. HT is obviously related to cases of primary or essential thrombocythemia, polycythemia vera, and agnogenic myeloid metaplasia (AMM). About half of the patients diagnosed as HT first come to medical attention via an emergency episode of acute bleeding or impending vascular occlusion. Spontaneous hemorrhages in HT typically develop at superficial cutaneous and mucosal surfaces, manifesting primarily as epistaxis, bruising, painful hematomas, ecchymoses, and bleeding from the gastrointestinal tract. Life threatening hemorrhage is infrequent and usually follows accidental or surgical trauma. When death occur, it was usually due to major thrombosis and much less frequent to hemorrhages. Deep tissue bleeding like joints bleeds seen in hemophilia, muscles hematomas, and intracerebral hemorrhage is very rare. Petechiae specific for thrombocytopenia of Glanzmann or Bernard Soulier or severe thrombocytopenia are never seen.

Figure 1: Spectrum of aspirin sensitive erythromelalgia in myeloproliferative thrombocythemia.

Figure 2: Discovery of platelet-mediated arteriolar inflammation and thrombosis (erythromelalgia) in thrombocythemia of patients with essential thrombocythemia and polycythemia vera (Michiels et al. 1984 and 1985).
Delineation of Erythromelalgic Thrombotic Thrombocythemia (ETT) in ET and PV

Vreeken and van Aken [11] described a patient with aspirin-responsive recurrent painful toes and fingers (erythromelalgia) in ET. Peripheral gangrene [21], microvascular ischemic or occlusive disease [18,65], and peripheral arterial thrombosis has been recognized as presenting symptoms of ET. Bernstein et al. [66] found ischemic signs on the electrocardiograms (ECG) in two ET patients with angina pectoris and "Prinz Metal" angina pectoris with transient increased distress that persisted for a few days. Bab et al. [64] observed attack of amaurosis vasospasm followed by transient dilopia. Sometime there had been luminous visual phenomena, and at times he had transient attacks of giddiness. Neurologic manifestations in ET patients varied from recurrent attacks of transient ischemic attacks (TIA) to persistence of neurological deficits. Unilateral migraine-like headache, amaurosis fugax preceded by scotomas trigeminus neuralgia, aphasia, dysarthria, central facialis paresis, transient paresis of one arm, leg or hemiparesis, persistent hemiparesis and organic vascular dementia have been frequently noted in ET patients [69-75]. Two ET patient became blind caused by thrombotic occlusion of the arteria centralis retinae. Singer [73] and Mundall et al. [75] observed attack of amaurosis vasospasm followed by white-yellow particle flowing from central to peripheral retina on fundoscopy, which disappeared together with complete recovery of the visus.

Smith and Allen first discovered that a single dose of acetyl salicylic acid (aspirin) produced marked relief of the burning erythromelalgic distress that persisted for a few days. Bab et al. [76] distinguished primary from secondary erythromelalgia associated with PV, systemic lupus erythematoses (SLE) or rheumatoid arthritis (RA) and stated that the response to aspirin is characteristic enough to constitute a valuable diagnostic clue for the diagnosis of erythromelalgia. Alarcon-Segovia et al. documented that aspirin-responsive erythromelalgia was a clue to early diagnosis in 8 PV patients associated thrombocythemia in 4 (Table 2). The spectrum of microvascular manifestations, cerebrovascular accidents, acute coronary artery syndromes, superficial thrombophlebitis, deep vein thrombosis and splanchic vein thrombosis at time of diagnosis in the 1970s of 226 PV patients are shown in Table 3. Post-splenectomy thrombocytosis in MPD patients is typically persistent and complicated by bleeding and thrombotic complications [77-79]. Reactive thrombocytosis is not associated with thrombotic or bleeding complications (McClure et al., Zucker and Mielke, Ginsburg, Coons et al.) [80]. The incidence of venous thrombosis is low in transient reactive post-splenectomy thrombocytosis [78,81,82]. Bleeding and thrombosis are rare in Ph+CML associated with persistent thrombocytosis [83].

Objective of the ETT vs. HT Literature Study Anno 1980 (Michiels Thesis 1981)

Between 1975 and 1980, the author of this monography discovered that erythromelalgic thrombotic complications in thrombocythemia (ETT) occurred at slightly increased platelet counts (>400 x 10^9/L) in ET and PV patients, whereas hemorrhagic complications in thrombocythemia (HT) in myeloproliferative disorders tend to occur.
at platelet counts in excess of 1000 × 10^9/L. In order to document a relationship between platelet count and clinical complication of thrombosis or hemorrhages I reviewed in 1980 the spectrum erythromelalgic, thrombotic and hemorrhagic manifestations and hematological findings in 200 consecutive thrombocytopenia cases published in the literature between 1929 and 1980. The 99 ETT cases histories from the literature could diagnosed as ET (N=67) and PV (N=32) and the 100 HT cases in thrombocytopenia of various MPDs were reported in the literature between 1908 1980 and published in my thesis [13].

Results in 99 ETT and 100 HT patients

The platelet number in ETT patients subdivided in ET and PV and the age distribution of 85 case histories of ETT is shown in Figure 3. Burning painful red or blue extremities, in retrospect very suggestive of erythromelalgia, were reported in 80 ETT cases. Peripheral (digital) gangrene developed in 31 cases. Eleven patients presented with peripheral gangrene without a record on burning. In 51 of the 62 evaluable cases, the peripheral arterial pulses were reported to be normal indicating the absence of significant atherosclerotic disease. Sympathectomy in 10 cases with digital gangrene (thromboangiitis obliterans) never induced improvement of the peripheral vascular insufficiency. Transient ischemic or thrombotic complications of the cerebral or coronary circulation were recorded in 12% and 19% of the cases, respectively. Deep vein thrombosis, pulmonary embolism and superficial thrombophlebitis of one or both leg rarely occurred (3% each).

The hemorrhagic symptoms of 100 case histories from the literature between 1933 to 1978 are shown in Table 4. The age distribution of 98 HT patients is shown in Figure 3. Gastrointestinal blood loss usually occurred as melena and/or hematemesis. A bleeding source could not be detected, neither roentgenologically in 19 out of 24, nor gastroscopically in 8 out of 9 patients investigated. In a few cases, the bleeding source could not be localize even at laparotomy. Nose bleeds in 33 patients were generally profuse and relapsed frequently. The typical gingival bleeding in 15 patients usually occurred after tooth extraction. Skin bleedings were either recorded as bruises in 18 as subcutaneous hematomas in 15, and as ecchymoses or sugillations in 10 out of 34 patients. In some cases painful subcutaneous hematomas started with or had a central nodule (very likely a thrombotic clot), which extended to large hemorrhagic effusions is. Petechiae characteristic for severe thrombocytopenia or congenital thrombosthenia of Glanzmann were never seen.

Arterial and/or venous thrombosis preceded or followed the hemorrhagic periods in 15 out of 100 HT cases (15%) (Table 5). One HT case developed recurrent gangrene of the toes 4 years following the hemorrhagic manifestations [29,30]. The thrombotic manifestations preceding the hemorrhagic diathesis were described as painfull acrocyanosis in 3, gangrene of toes in 5, transient neurologic ischemic attacks in 3 and hemiparesis in 2, superficial thrombophlebitis in 3, priapism in 3 and deep vein leg thrombosis in one case only (Table 5). In 7 of 99 ETT cases (7%) hemorrhagic complications are recorded; subcutaneous hematomas in 2, ecchymoses in 10 out of 34 patients. In some cases painful subcutaneous hematomas started with or had a central nodule (very likely a thrombotic clot), which extended to large hemorrhagic effusions is. Petechiae characteristic for severe thrombocytopenia or congenital thrombosthenia of Glanzmann were never seen.

The spleen in 64 evaluable ETT patients was not palpable in 27, slightly to moderately enlarged in 36 and greatly enlarged to below the costal margin in only 3. The spleen in 67 evaluable HT patients was not palpable in 17, slightly to moderately enlarged in 26 and greatly enlarged to below the umbilical level in 21 cases. In 2 ETT patients a splenectomy had been performed. In 20 HT patients, splenectomy was performed (spleen weight in excess of 1 kilogram) had induced a post-splenectomy thrombocytopenia complicated by bleedings (see section post-splenectomy HT) (Table 6). At time of splenectomy portal vein thrombosis was diagnosed in 2 and thrombosis of vena lienalis in 4 patients. Platelet counts in these 6 patients with splanchic vein thrombosis was normal in 4 and strongly increased in 2. The majority of post-splenectomy HT patients (90%) had no bleeding manifestations at time of splenectomy. Platelet count at time of splenectomy was normal in 8, increased between 400-1000 × 10^9/L in 3, increased above 1000 × 10^9/L in 3, and not stated in 6 cases (Table 6). In the 3 cases with platelet counts of >1000 × 10^9/L had bleeding manifestation during or
Platelet and leukocyte counts, and bone marrow findings in ETT and HT patients

Platelet counts ranged from 735 to 5000 × 10^9/L in HT and from 352 to 3500 × 10^9/L in ETT (Figure 3). The frequency distribution of the platelet and leukocyte counts in ETT and HT is shown in Table 7. The platelet count of 1059±507 × 10^9/L (mean±SD) in 96 ETT patients was significantly lower (P<0.001, Mann-Whitney-U test) than the platelet count of 2050±1107 × 10^9/L in 100 PHT patients. The mean leukocyte count of 25±17 × 10^9/L in 86 PHT patients was significantly higher immediately after splenectomy (Table 6). In 17 post-splenectomy cases hemorrhagic manifestation became evident after 1 month in 4, after 3 months in 4, after 6 months in 1 and after one or a few years in 6 cases. At time of HT the platelet counts were far above 1000 × 10^9/L (range 1289-4353, Table 6). The weight of splenectomized spleen was significantly increased except in 1 and not stated in 4 (Table 6). One case of post-splenectomy HT in a 43-years old case was complicated by angina pectoris, myocardial infarction, and transient hemiparesis at platelet count of 1700 × 10^9/L [84], and by erythromelalgia (painful red swollen toes) at platelet count of 900 × 10^9/L in another case [85].
This pseudohyperkaliemia in thrombocythemia of various normal renal function and normal plasma potassium levels [86-89]. In 4 PHT patients. Between 1960 and 1970, Silverstein observed 47 HT patients with peripheral blood and bone marrow findings were consistent with excluding erythroleukemia, preleukemia and MDS. In addition, clumps, normoblastic erythropoiesis and normal myelopoiesis thereby patients showed major increase of mature megakaryocytes and platelet counts between 375 and 1000 × 10^9/L. There was no correlation between increased number of platelets and leukocytes in each of the cases had leukocytosis between 20 and 90 × 10^9/L. In some cases of HT the white blood cell differential count was left-shifted indicative for more advanced MPD disease [19,31,40,54,59,61,63].

The individual platelet in ETT and HT patients is shown in Figure 3. About half of 67 ETT patients had thrombotic complications at platelet counts above 400 × 10^9/L (Michiels et al. [15]). Focusing on the causal relation between erythromelalgia and thrombocythemia in ET and PV patients, we were able to document since 1975 the very early stage of ET by the use of the Rotterdam Clinical and Pathological (RCP) criteria for ET and PV [92,93]. The 1980 RCP criteria of ET and PV were determined by careful prospective documentation of peripheral blood and bone marrow smears and bone marrow biopsy material [93]. Platelets in excess of 400 × 10^9/L, and an increase of clustered enlarged megakaryocytes in a bone marrow biopsy material was found to be diagnostic for ET and excluded reactive thrombocytosis. On top of the clinical PVSG criteria for PV [46] we introduced in 1980 bone marrow histopathology and erythocyte count above 6 × 10^12/L; Dameshek 1940 [2].

Bone marrow findings, mainly aspirates, in 36 HT and 31 ETT patients showed major increase of mature megakaryocytes and platelet clumps, normoblastic erythropoiesis and normal myelopoiesis thereby excluding erythroleukemia, preleukemia and MDS. In addition peripheral blood and bone marrow findings were consistent with chronic myeloid leukemia in 5 PHT and agnogenic myeloid metaplasia in 4 PHT patients. Between 1960 and 1970, Silverstein observed 47 HT patients in whom the thrombocythemia was associated with chronic granulocytic leukaemia, myeloid metaplasia, or polycythaemia vera in eight patients each.

At platelet counts in excess of 800 × 10^9/L, serum potassium concentrations are always significantly elevated in the presence of normal renal function and normal plasma potassium levels [86-89]. This pseudohyperkaliemia in thrombocythemia of various myeloproliferative disorders is explained by release of potassium from the platelets during coagulation and clot retraction [23,90].

### Discussion

The PVSG criteria in 1975 for the diagnosis of hemorrhagic thrombocythemia were and crude and required a minimal platelet count in excess of 1000 × 10^9/L [91] thereby overlooking symptomatic ET patients who presented with ET at platelet counts above 400 × 10^9/L (Michiels et al. [15]). Focusing on the causal relation between erythromelalgia and thrombocythemia in ET and PV patients, we were able to document since 1975 the very early stage of ET by the use of the Rotterdam Clinical and Pathological (RCP) criteria for ET and PV [92,93]. The 1980 RCP criteria of ET and PV were determined by careful prospective documentation of peripheral blood and bone marrow smears and bone marrow biopsy material [93]. Platelets in excess of 400 × 10^9/L, and an increase of clustered enlarged megakaryocytes in a bone marrow biopsy material was found to be diagnostic for ET and excluded reactive thrombocytosis. On top of the clinical PVSG criteria for PV [46] we introduced in 1980 bone marrow histopathology and erythrocyte count above 6 × 10^12/L proposed by Dameshek in 1940 [94] as specific clues to the diagnosis of PV to clearly differentiate PV from all variant of primary and secondary erythrocytosis. The 1980 RCP modifications of the 1975 PVSG criteria for PV include 4 main changes [13]. First, the major criterion O_2-saturation of >92% is replaced by absence of primary or secondary erythrocytosis by clinical and laboratory tests. Second; splenomegaly is replaced by bone marrow histology as a major criterion (A3). Third, the 1980 RCP diagnostic set used splenomegaly as a minor criterion . Fourth, we skipped raised B12 (>900 ng/L) or raised B12 binding capacity (>2200 ng/L) as completely irrelevant for the diagnosis of early and overt stage PV.
The follow-up of the Rotterdam MPD Working Group (1975-1995) on the efficacy and safety for the prevention and treatment of thrombotic complications in 68 ET patients seen in the Academic Hospital Rotterdam has been critically evaluated by Perry van Genderen et al. [95,96]. The therapeutic implications from the prospective Rotterdam ET studies are completely in line with the concept that microcirculatory thrombotic complications in thrombocythemia already occur at platelet counts in excess of 400 × 10^9/L, which are relieved by reduction of platelet counts to normal (<400 × 10^9/L) or by control of platelet function with low dose aspirin. At time of presentation 54 patients had ET related microvascular thrombotic complications at platelet counts between 575 and 1031, mean 750 × 10^9/L and treated with low-dose aspirin. In patients receiving platelet lowering agents either busulfan or hydroxyurea and no aspirin, thrombotic complications occurred frequently during follow-up at platelet counts above 400 × 10^9/L, indicating that inadequate control of platelet number in thrombocythemia already occur at platelet counts in excess of 400 × 10^9/L, which are relieved by reduction of platelet counts to normal (<400 × 10^9/L) or by control of platelet function with low dose aspirin. At time of presentation 54 patients had ET related microvascular thrombotic complications at platelet counts between 575 and 1031, mean 750 × 10^9/L and treated with low-dose aspirin. In patients receiving platelet lowering agents either busulfan or hydroxyurea and no aspirin, thrombotic complications occurred frequently during follow-up at platelet counts above 400 × 10^9/L (624 ± 255 × 10^9/L), indicating that inadequate control of platelet number in thrombocythemia of ET and PV patients is associated with a persistent high incidence of microvascular disturbances including MIAs when not on aspirin at platelet counts above 400 × 10^9/L. One low dose aspirin reduces the risk of microvascular ischemic or thrombotic complications from above 50% to less than 5% during 6.2 years of follow-up, and thereby prevented the progression from low thrombotic risk to high risk thrombotic in ET and PV (Van Genderen et al. [95,96]). Aspirin in ET at platelet counts above 1000 × 10^9/L in general daily practices is not safe enough in terms of bleedings elicited by aspirin indicating the need to add non-leukemogenic platelet lowering agents. Thrombocythemia with platelet counts in excess of 1000+250 × 10^9/L is frequently associated with the paradoxical occurrence of thrombosis and bleeding. Mucocutaneous bleedings spontaneously occur at platelet count in excess of 1000+250 × 10^9/L due to an acquired type II-like von Willebrand syndrome (AVWS, absence of high and intermediate von Willebrand factor (VWF) multimers resulting in an acquired von Willebrand disease (AVWD) type 2A. (Right) Loss of large VWF multimers at increasing platelet count is associated with low VWF:RCO/VWF:Ag(open circles) and VWF:CB/VWF:Ag (black dots) ratios indicative for a typical acquired von Willebrand disease (AVWD) type 2 A.

References

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Figure 4: (Left) The Cigar of ETT and Wedge concept of erythromelalgic thrombotic thrombocythemia (ETT) and hemorrhagic thrombocythemia (HT). The transition of ETT into HT is related to the increase of platelet counts from below to far above 1000×10^9/L caused by a platelet mediated proteolysis of large von Willebrand factor (VWF) multimers resulting in an acquired von Willebrand disease (AVWD) type 2A. (Right) Loss of large VWF multimers at increasing platelet count is associated with low VWF:RCO/VWF:Ag(open circles) and VWF:CB/VWF:Ag (black dots) ratios indicative for a typical acquired von Willebrand disease (AVWD) type 2 A. ETT=erythromelalgic thrombotic thrombocythemia. HT=hemorrhagic thrombocythemia. ET essential thrombocythemia. PV=polycythemia vera. VWF:RCO=von Willebrand ristocetin cofactor activity. VWF:CB= von Willebrand factor collagen binding.
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