

MYELOPROLIFERATIVE NEOPLASMS

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September 15, 2015

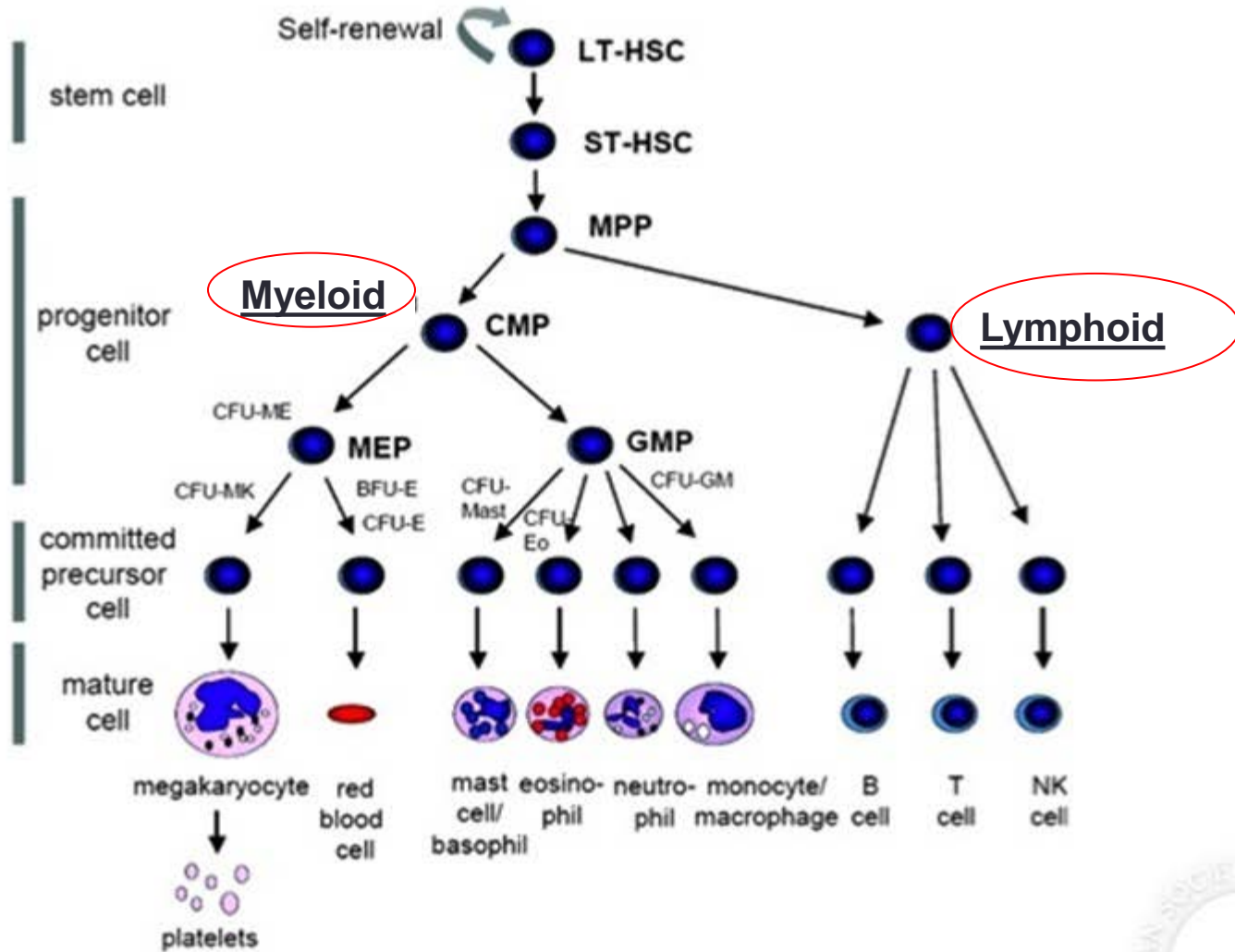
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Objectives

- Discuss an approach to increased cell counts
- Discuss clinical and laboratory diagnosis of myeloproliferative neoplasms
- Overview of treatment options and potential complications of disease and therapy

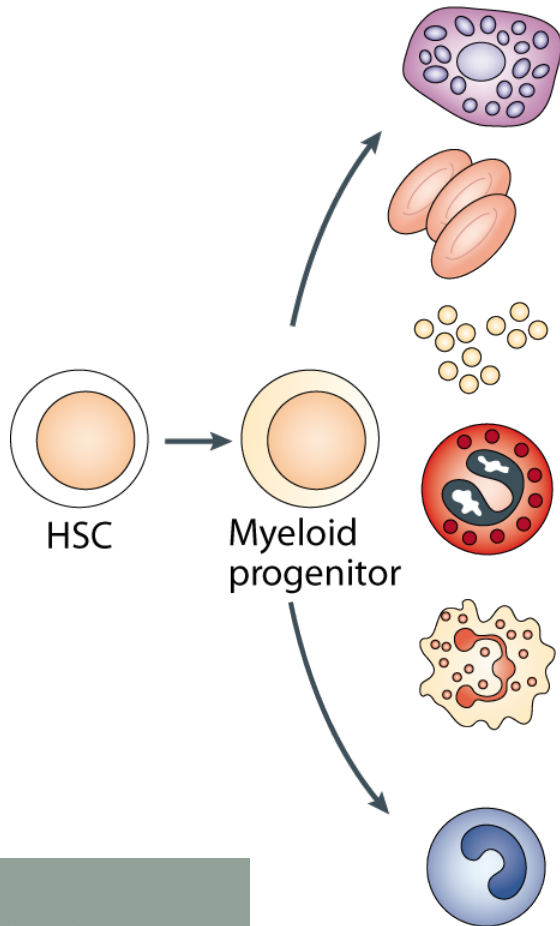
Figure 12-3 Classical hierarchal map of hematopoietic development

ash-sap™



Cantor, A. B. et al. ASH-SAP 2010;2010:331-372

Myeloproliferative Disorders



	MPD	Activating mutation
Mast cell	→ Systemic mastocytosis	KITD816V FIP1L1-PDGFRA
Red blood cells	→ Polycythemia vera	
Platelets	→ Essential thrombocytosis	
Eosinophils	→ Chronic eosinophilic leukemia	FIP1L1-PDGFRA
Neutrophils	{ Chronic myeloid leukemia Chronic myelomonocytic leukemia	BCR-ABL TEL-PDGFRB BCR-PDGFRA TEL-JAK2 other fusion TKs
Monocytes		
		CALR MPL

Marrow Production and Peripheral Blood Half-Life

	<u>Output/day</u>	<u>Blood Count</u>	<u>Lifespan</u>
RBC	200×10^9	$\sim 5 \times 10^6/\mu\text{L}$	120 days
WBC	10×10^9	$\sim 3 \times 10^3/\mu\text{L}$ (neutrophils)	< 1/2 day
Plts	400×10^9	$\sim 200 \times 10^3/\mu\text{L}$	10 days

Leukocytosis

- A word to discourage from clinical use
 - Be more specific!
- For diagnosing MPNs – focus on Absolute counts, not %
- Specific type of cell will help build your differential
 - Neutrophilia: leukemoid reaction/reactive, CML, myelofibrosis
 - Lymphocytosis: CLL, MBL, pertussis,
 - Monocytosis: CMML, TB/fungal,
 - Eosinophilia : allergy/atopy, parasites, adrenal insufficiency, CEL
 - Basophilia: CML
 - Peripheral Blasts: Acute leukemia, high-grade MDS

Case 1 - Presentation

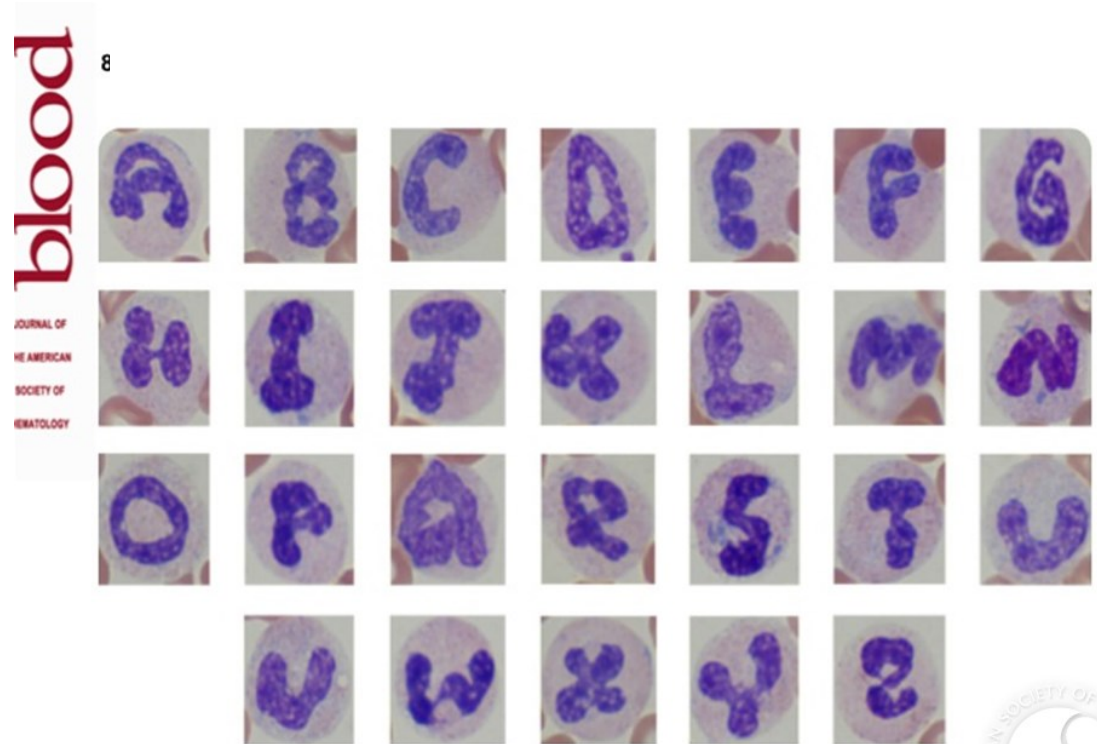
- 32yo resident presents with sore throat and fever
- Cervical adenopathy is present on exam

- CBC: 35>45%<455k

Case 1 - Differential

- 86% Neutrophils
- 12% Immature Granulocytes
- 2% Lymphocytes

- Rapid strep test is positive
- He improves with a course of antibiotics



Morgan A S , and Yang D T Blood 2013;121:3546-3546

©2013 by American Society of Hematology



Origin of MPN

MF: Dr. Gustav Heuck 1879 Two cases of leukemia with peculiar blood and bone marrow findings, respectively

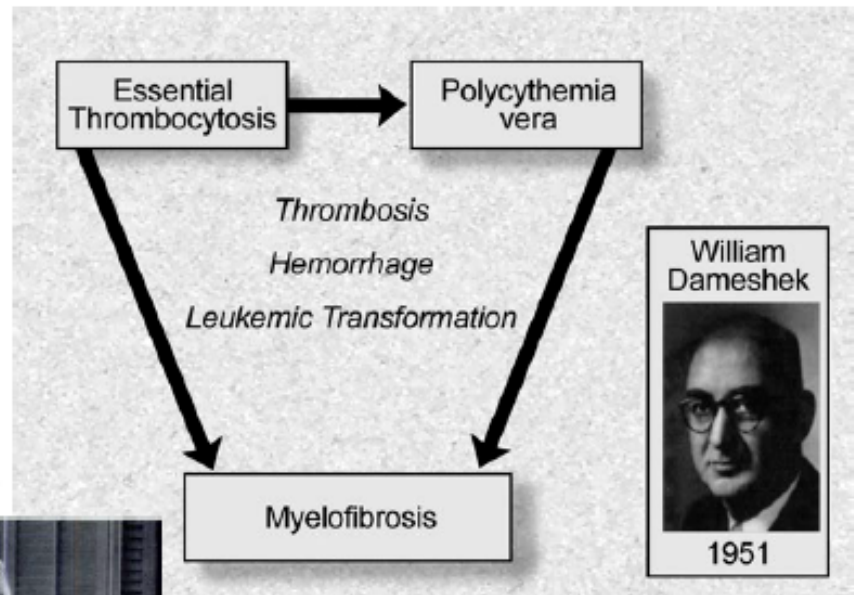
PV: Dr. Louis Henri Vaquez 1892 On a special form of cyanosis accompanied by excessive and persistent erythrocytosis Dr. Osler coins "Vaquez's disease" in 1903 chronic cyanosis with polycythemia and enlarged spleen

ET: Drs. Emil Epstein and Alfred Goedel 1934 Hemorrhagic thrombocytopenia with a cascular, sclerotic spleen

TABLE 1.—The Myeloproliferative Disorders
(Myelostimulatory Factor's)

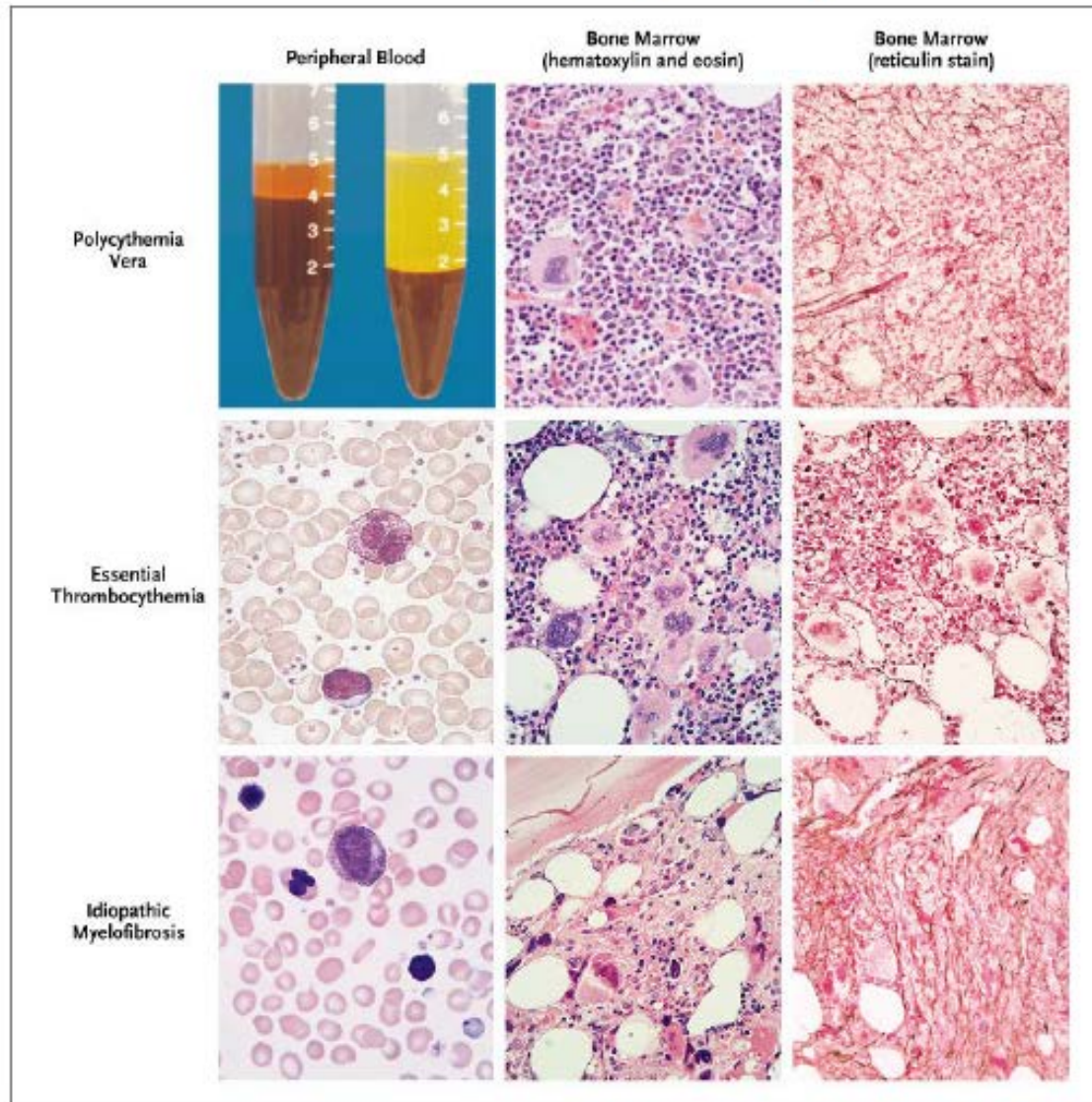
Syndromes	Bone marrow				Potential bone marrow
	Erythroblasts	Granulocytes	Megakaryocytes	Fibroblasts	Myeloid metaplasia of spleen and liver
Chronic Granuloeytic Leukemia	±	+++	+ to +++	+	++
Polycythemia Vera	+++	++	++ to +++	+ to +++	+ to +++
Idiopathic or Agnogenic Myeloid Metaplasia of Spleen	±	±	+++	+ to +++	+++
Megakaryocytic Leukemia	±	±	+++	+	+ to +++
Erythroleukemia (including diGuglielmo syndrome)	+++	+	±	±	+ to +++

Degrees of Proliferation: + slight
++ moderate
+++ marked

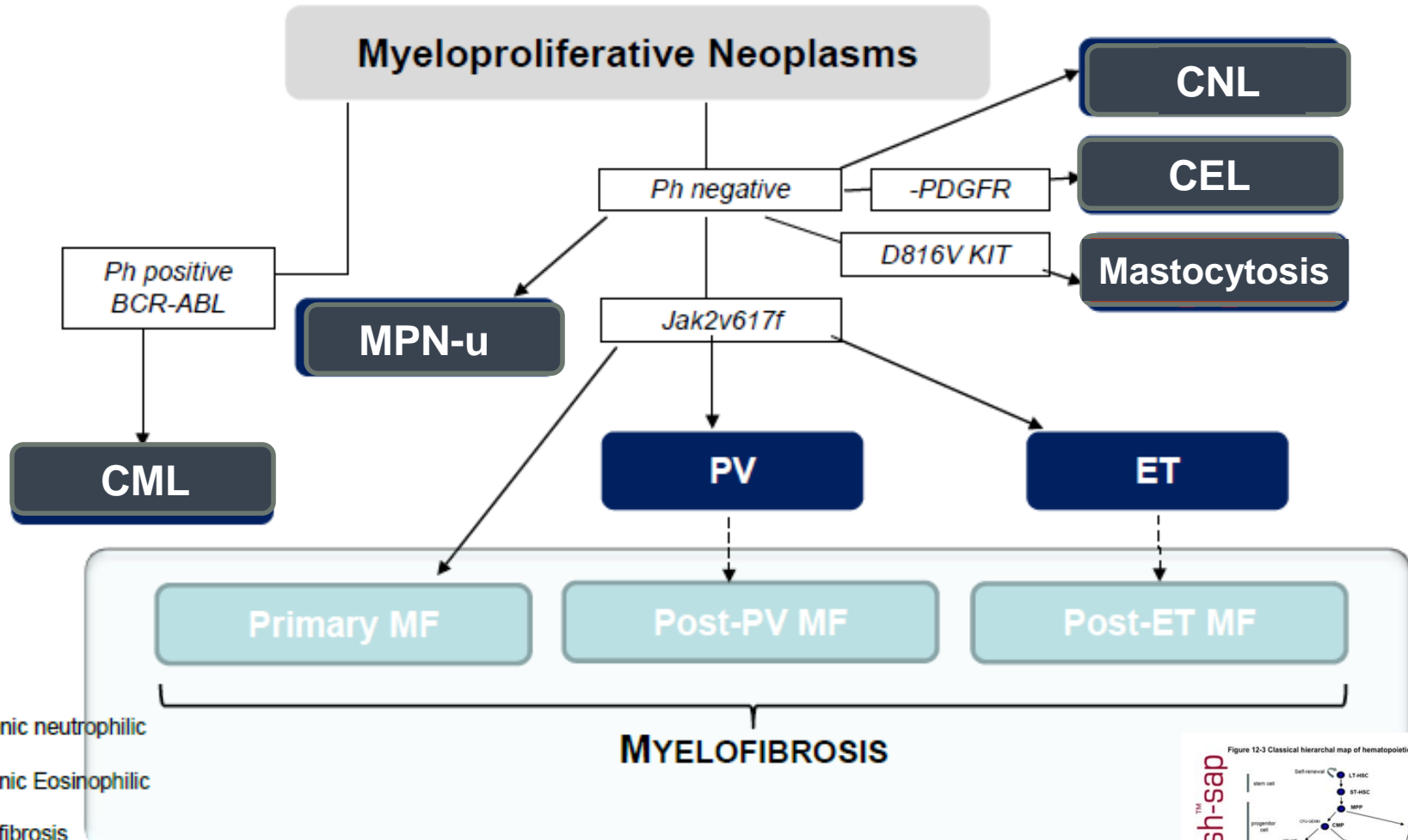


Dameshek et al. *Blood* 1951;6:372-375
Levine and Gilliland *Blood* 2008;112:2190-2198

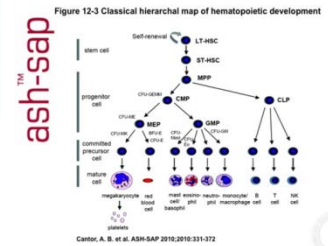
Lab Features of PV, ET, and MF



Making a Molecular Diagnosis

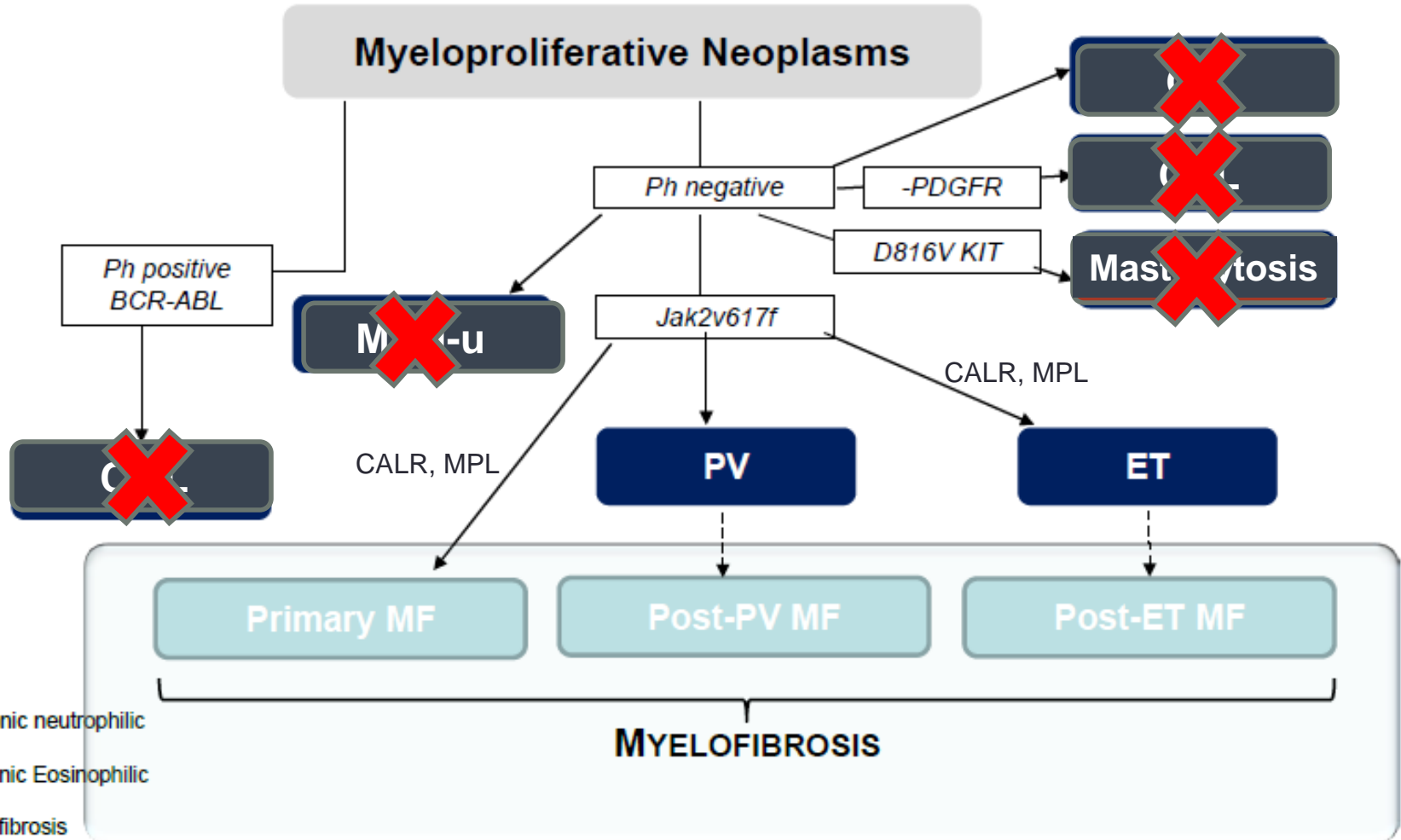


CNL=Chronic neutrophilic leukemia
 CEL=Chronic Eosinophilic Leukemia
 MF=myelofibrosis
 PV=polycythemia vera
 ET=essential thrombocythemia
 CML=chronic myeloid leukemia



Tefferi A, Vardiman JW. *Leukemia*. 2008;22:14-22; Vardiman JW, et al. *Blood*. 2009;114(5):937-951 Mesa RA. *Blood*. 2009;113(22):5394-5400; Tam CS, et al. *J Clin Oncol*. 2009;27:5587-5593.

Making a Molecular Diagnosis



CNL=Chronic neutrophilic
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Klampfl NEJM 2013

Jak 2 Testing in MPN



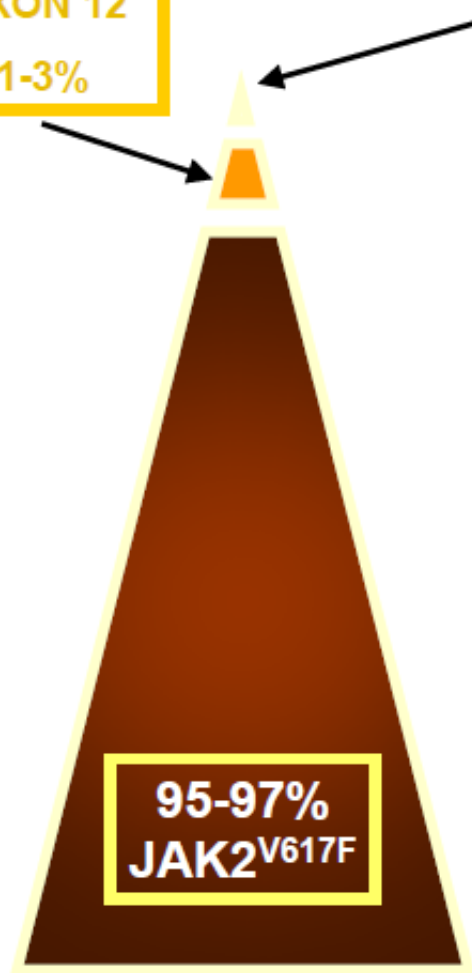
<u>Reference</u>	<u>Assay</u>	<u>Source*</u>	<u>PV % (N)</u>	<u>ET % (N)</u>	<u>MMM % (N)</u>	<u>Controls</u>
Baxter et al. [#]	AS-PCR	PB & BM	97% (73)	57% (51)	50% (16)	0% (90)
Levine et al. ^{#\$}	PCR	PB	74% (164)	32% (115)	35% (46)	0% (270)
James et al. [#]	PCR	PB & BM	89% (45)	43% (21)	43% (7)	0% (45)
Kralovics et al. ^{#\$^}	PCR	PB	65% (128)	23% (93)	57% (23)	0% (82)
Zhao et al.	PCR	PB	83% (24)	N/A	N/A	0% (12)
Tefferi et al.	PCR	PB	95% (38)	55% (22)	30% (10)	0% (30)
Jones et al.	AS-PCR	PB	81% (72)	41% (59)	43% (35)	0% (160)

* purified granulocytes
[#]T-Lymphocytes, ^{\$}Buccal mucosal cells, and [^]hair follicles were negative

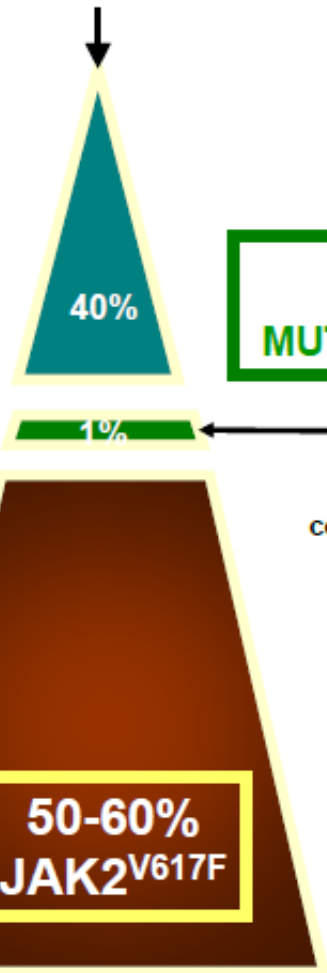
Baxter et al *Lancet* 2005. 365:1054
 Levine et al *Cancer Cell* 2005. 7:387.
 James et al. *Nature* 2005. 434: 1144

**? OTHER MUTATIONS
Genetic/Host Interaction**

JAK2
EXON 12
1-3%

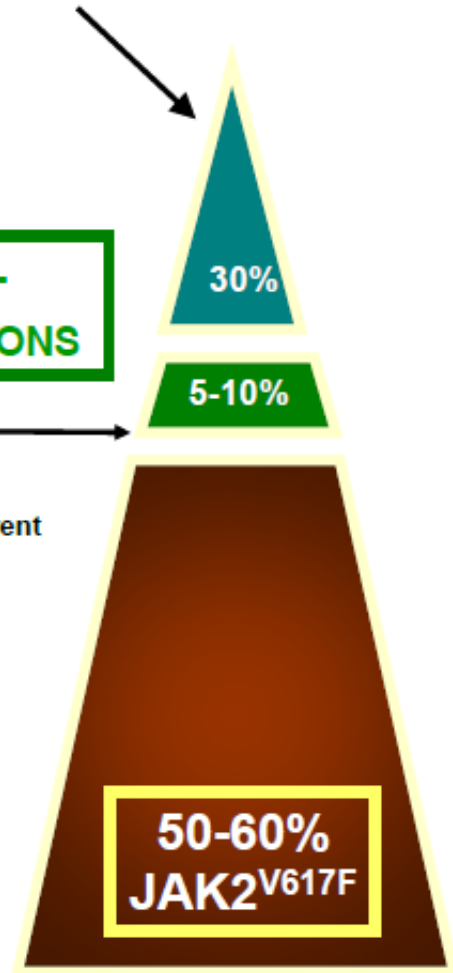


PV



ET
Heterozygous

MPL
MUTATIONS



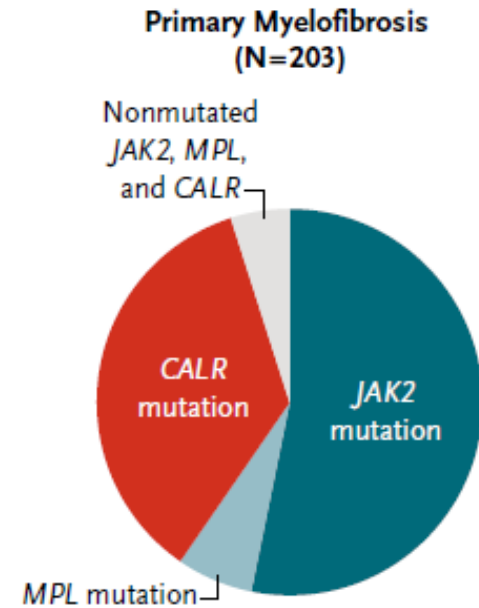
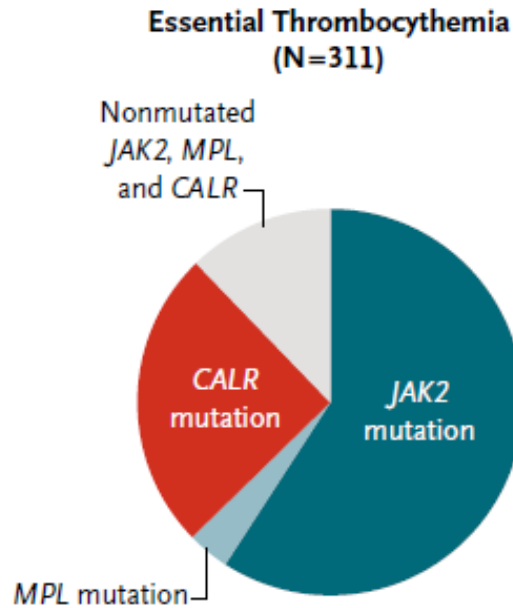
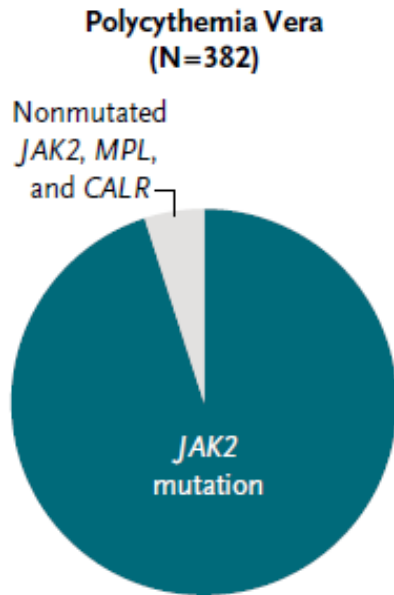
MF

concurrent

Pardanani et al. *Blood* 2006;108:3472-3476
Scott et al. *N Engl J Med* 2007;356:459-468
Kralovics et al. *N Engl J Med* 2005;352:1779-1790

Calreticulin as the 'other mutation'

A Distribution of *JAK2*, *MPL*, and *CALR* Mutations in Philadelphia Chromosome–Negative Myeloproliferative Neoplasms



PV and ET Diagnostic Criteria

WHO Criteria¹: PV

Major Criteria (first major + 2 minor or both + 1 minor)

- Hgb > 18.5 g/dL in men, 16.5 g/dL in women or other evidence of RCV*
- *Jak2V617F* or other mut *Jak2* exon 12

Minor Criteria (first major + 2 minor or both + 1 minor)

- BM Trilineage proliferation
- Low Epo level
- Endogenous ECF in vitro

*Hgb or Hct > 99th% of reference range
or Hgb > 17 g/dL in men, 15 g/dL in women if at least 2 g/dL above baseline not attributed to correction of Fe def.
or elevated RCM > 25% above predicted

WHO Criteria¹: ET

Major Criteria (all required)

- Plt Count $\geq 450 \times 10^9/L$ sustained*
- Megakaryocyte proliferation with increased # of enlarged mature megakaryocytes
- Does not meet criteria for other myeloid d/o (PV[¶], MF[†], CML[‡], MDS[§])
- Clonal marker (*Jak2V617F*) or no evidence of reactive thrombosis[§]

*during the w/u

¶ failure of Fe to increase Hgb in setting of a low ferritin

† absence of relevant reticulin or collagen fibrosis, leukoerythroblastosis, or abnml meg morphology (n/c ratio, hyperchromatic, bulbous, irregularly folded nuclei, and dense clustering)

‡ absence of BCR-ABL1.

§ absence of erythroid and granulocytic dysplasia

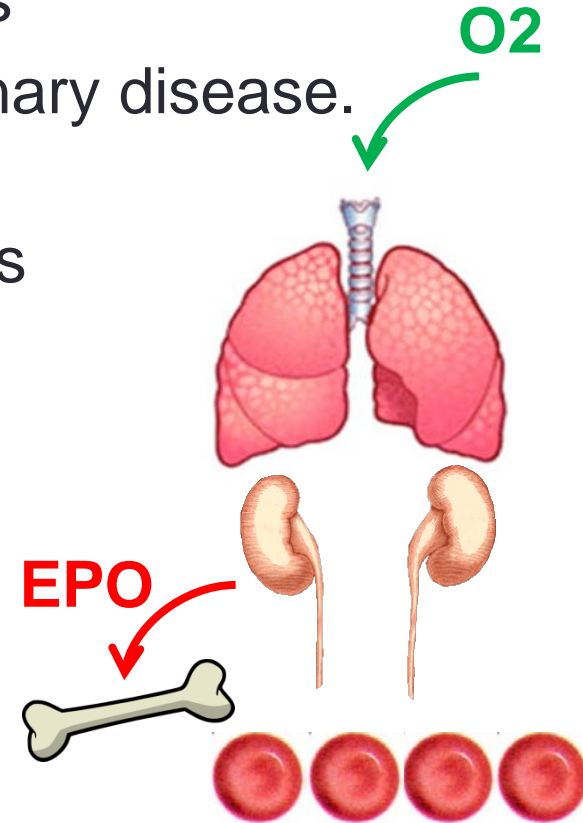
§ the presence of a condition associated with reactive thrombocytosis (Fe def, infection, inflammation, met cancer, connective tissue disease, lymphoproliferative d/o) does not exclude possibility of ET

¹Vardiman JW, et al. *Blood*. 2009;114(5):937-951.

Case 2 - Presentation

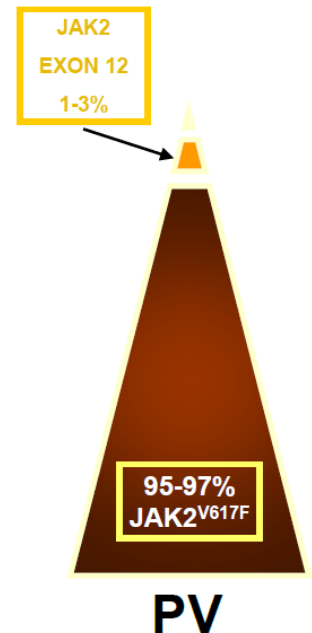
- 65yo woman is referred for 'abnormal labs'
- Nonsmoker, no OSA, no history of pulmonary disease. She does not live at altitude.
- She reports pruritis but no other symptoms
- O2 saturation 98% RA
- Hb = 19
- WBC 9 Plt 400k

Next Tests?



Case 2 – Diagnostics: Polycythemia Vera

- EPO = 5 (2-18)
- JAK2 V617F mutation positive
- There is no need for a bone marrow with positive JAK2 in PV
- (Potential causes of secondary polycythemia include altitude, lung disease/hypoxia, renal cell carcinoma and hepatocellular carcinoma as well as testosterone/anabolic steroid use or exogenous EPO)



Case 2 – Treatment: Back to the Future

- Goal Hct is $<45\%$ (better than $<50\%$ in randomized trial by Marchioli et al. *NEJM* 2013 368:22)
 - Phlebotomy
 - Hydroxyurea
- ASA



Ancient Greek Painting



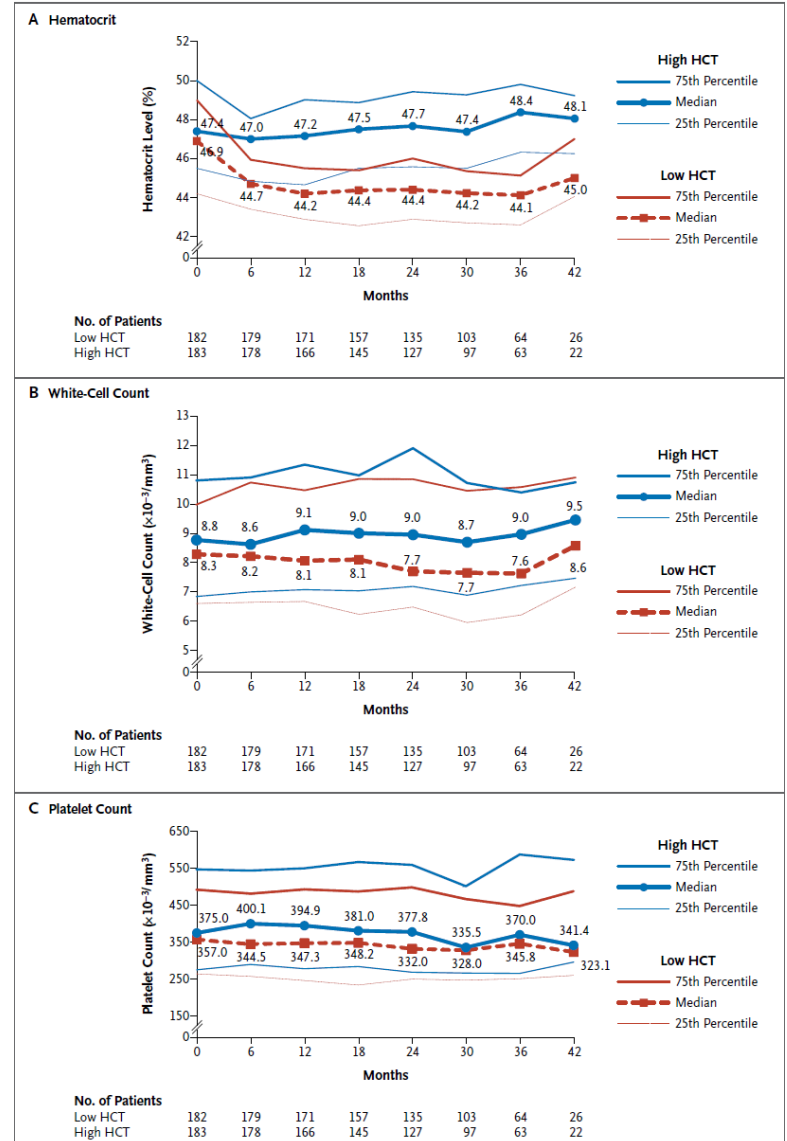
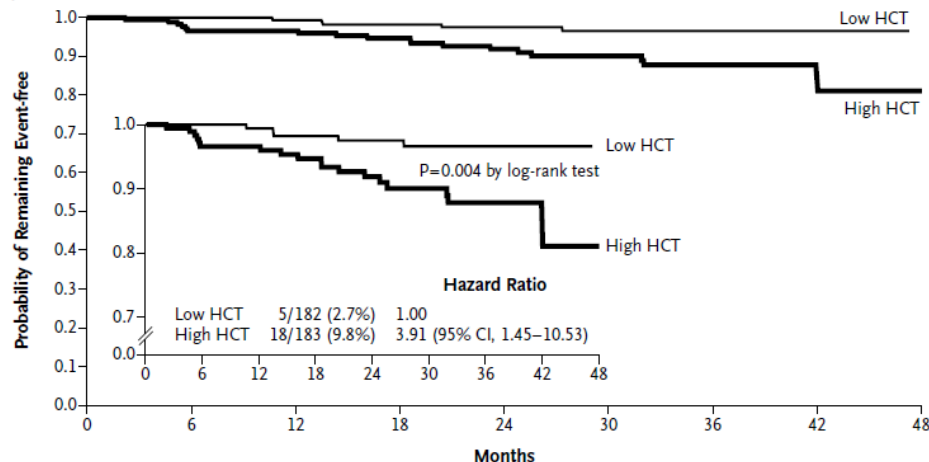
Photograph from the Burns Archive 1860

Patients with PV cannot donate blood,
but patients with hemochromatosis can

CYTO-PV Study: 45% vs 50%

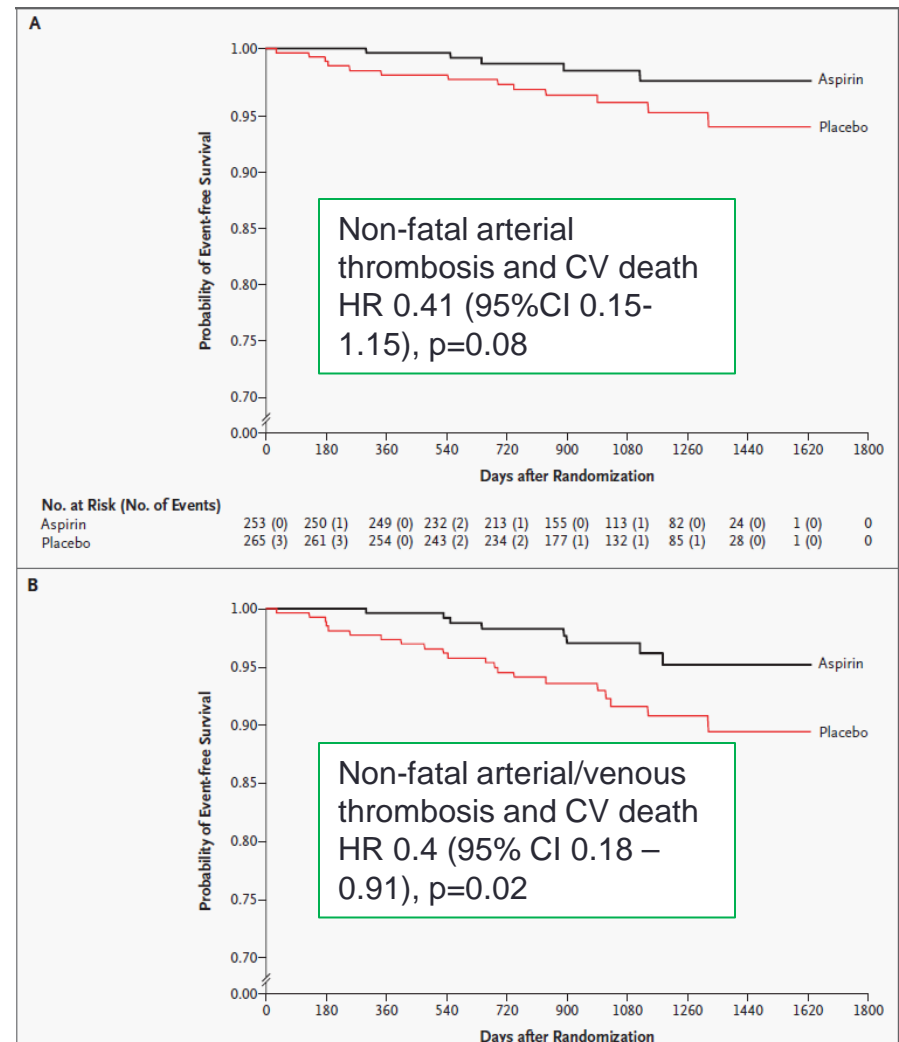
- 365 patients, randomized
- Primary end point
 - death from cardiovascular causes or thrombotic events
- HU or phlebotomy allowed

A Primary End Point



ECLAP: ASA vs Placebo in PV

- Efficacy and Safety of Low Dose Aspirin in PV
 - Multicenter European Study
- 518 patients, randomized
- Mean follow up 3 years
- More smokers in ASA arm
- Other tx as needed
 - Cytoreduction (HU)
 - Phlebotomy
- No difference in overall mortality
- NS reduction in major thrombosis
- Major bleeding not different

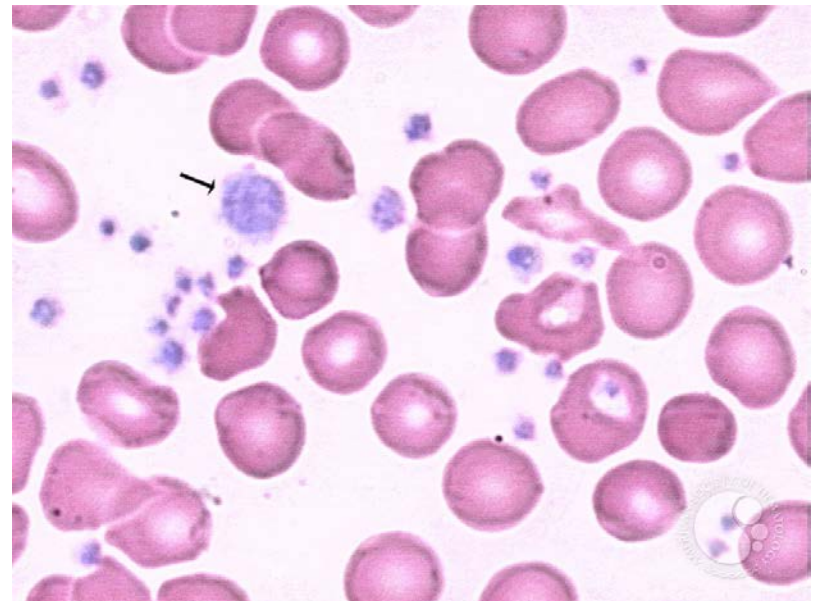


Case 3 - Presentation

- 55yo man presents with fatigue, and abnormal labs prior to upcoming hernia surgery.
- He has no active infections. Exam reveals no major findings and his hernia is easily reducible without associated erythema or tenderness.

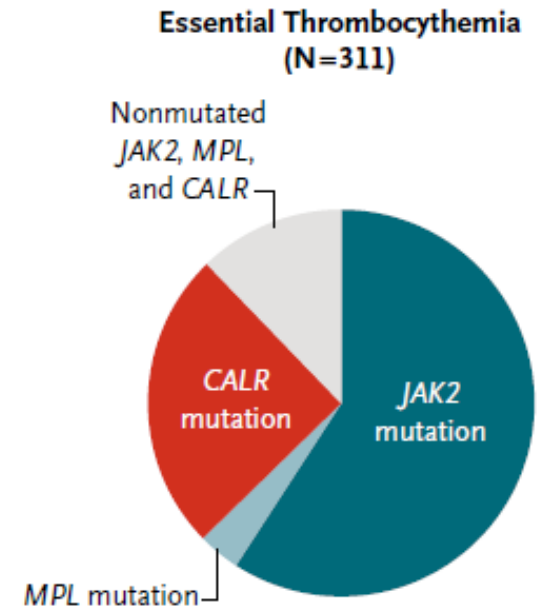
- CBC: $27 > 45\% < 750$
- N65%, L25%, M8%, E2%

Next Tests?

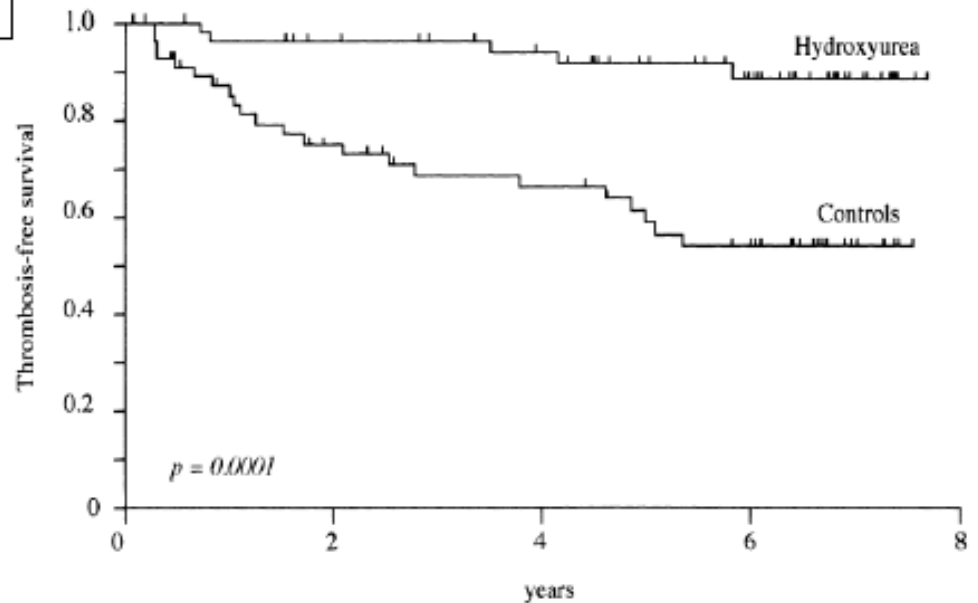
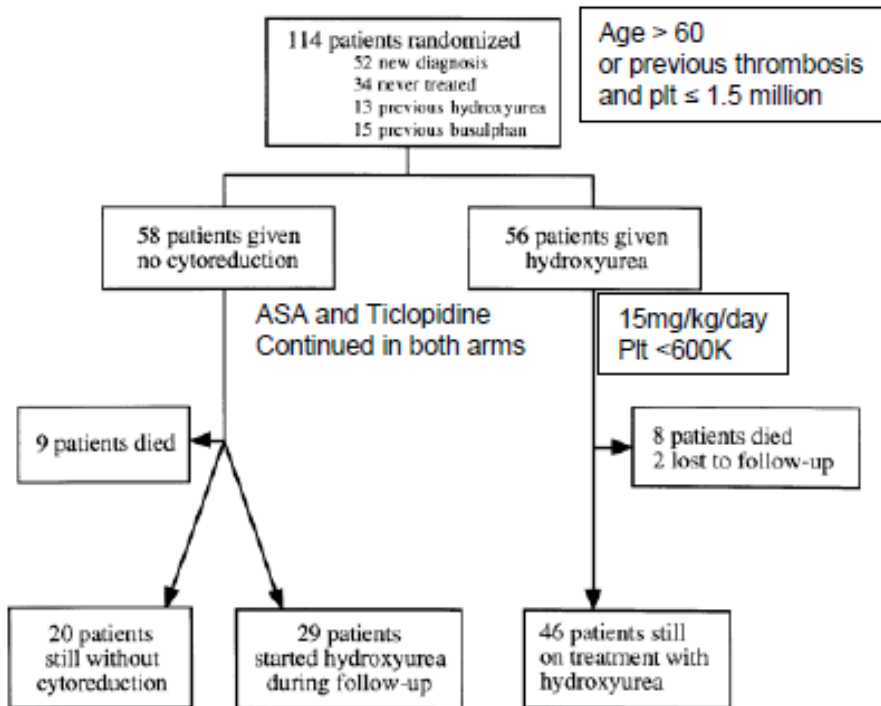


Case 3 - Diagnostics

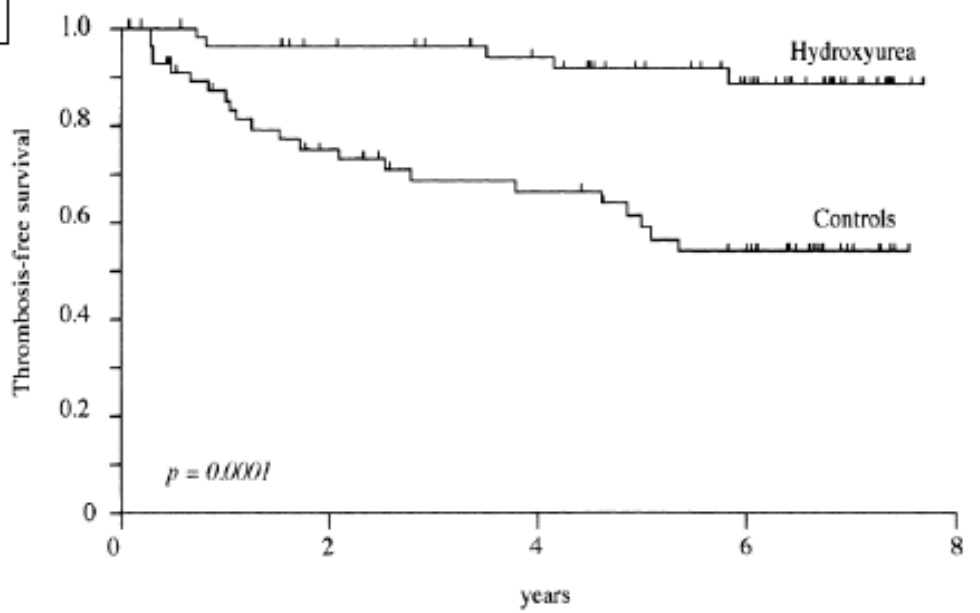
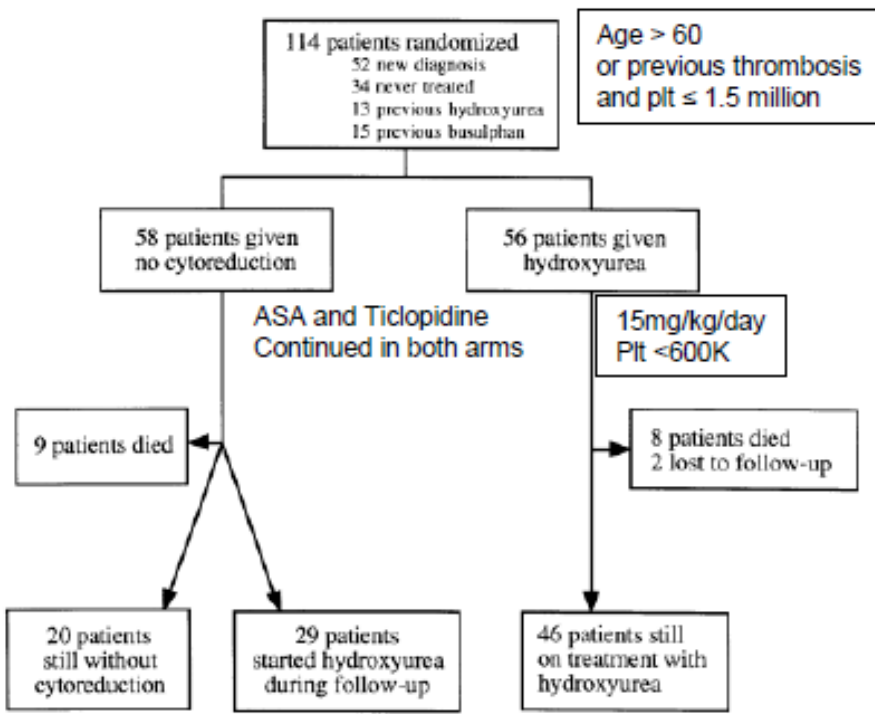
- JAK2 V617F mutation negative
- BCR/ABL negative
- CALR positive
- Bone Marrow - increased megakaryocytes, some are increased in size but not abnormal. No increase in fibrosis.
- Diagnosis of Essential Thrombocythemia



HU in High-Risk ET



HU in High-Risk ET



Who gets treated with ET (and who just phones home)?

	Age <60yo	Age >60yo
No prior Thrombosis	NO CYTOREDUCTION	Cytoreduce
Prior Thrombosis*	Cytoreduce	Cytoreduce

Barbui, JCO. 2011;29: 761.

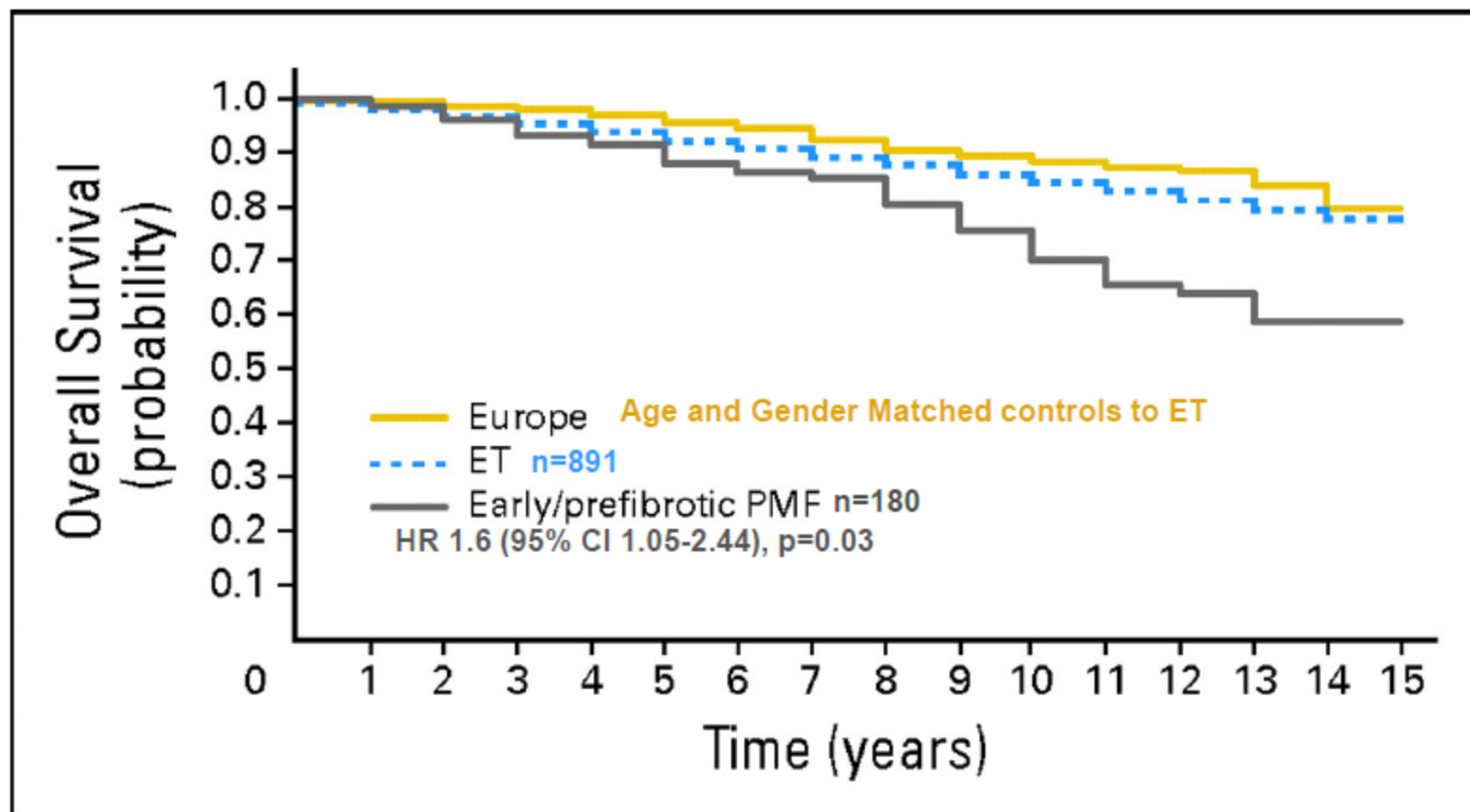
* Includes CVA, TIA, AMI, Arterial thrombus, or VTE

Table 2. Significant risk factors for thrombosis in 891 patients with WHO-defined ET and associated prognostic scores

Risk factor	HR	Score
Age > 60 y	1.50	1
Cardiovascular risk factors	1.56	1
Previous thrombosis	1.93	2
JAK2V617F	2.04	2

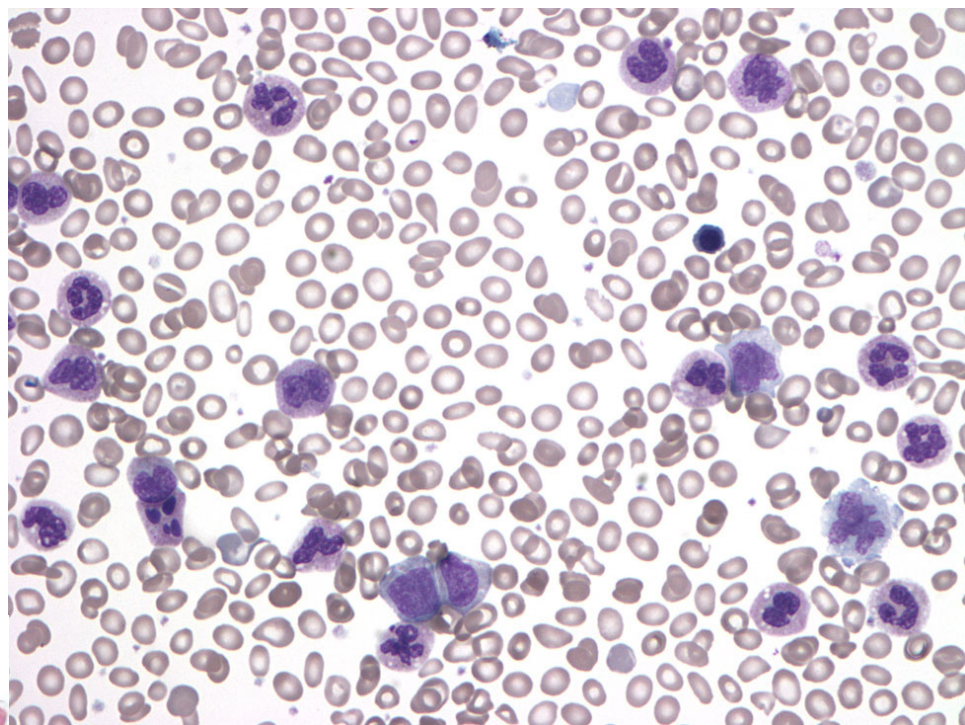
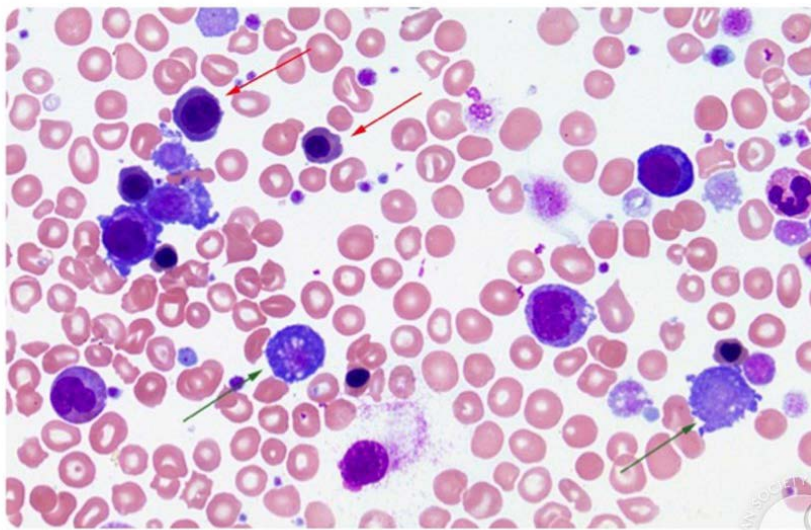
Low risk implies a score = 0-1; intermediate risk, score = 2; and high risk, score \geq 3.

ET vs. MF vs. Control



Case 4 - Presentation

- 62yo woman presents with LUQ abdominal pain, early satiety and weight loss x 3 months
- Examination reveals splenomegaly 8cm below the costal margin
- Next Test?



Sanford D , and Hsia C Blood 2013;122:4163

MF Diagnostic Criteria

WHO Criteria¹: Primary MF

Major criteria (all required)

- Megakaryocyte proliferation and atypia
 - Reticulin or collagen fibrosis
- Does not meet criteria for other myeloid disorders (e.g., PV,[¥] CML,[‡] MDS[§])
- Clonal marker (e.g., *MPLW515K/L*, *JAK2V617F*) or no evidence for secondary marrow fibrosis[§]

Minor criteria (must meet 2)

- Increase in serum LDH
- Palpable splenomegaly
- Leukoerythroblastosis
- Anemia

[¥] failure of Fe to to increase Hgb in setting of a low ferritin

[‡] absence of BCR-ABL1.

[§] absence of erythroid and granulocytic dysplasia

[§] infection, autoimmune, chronic inflammatory, hairy cell leukemia or other lymphoid neoplasm, met malignancy, or toxic chronic myelopathies

IWG Criteria²: Post-ET MF & Post-PV MF

Major criteria (all required)

- Previous diagnosis of ET or PV
- Grade 2-3 bone marrow fibrosis (on 0-3 scale) or Grade 3-4 bone marrow fibrosis (on 0-4 scale)

Minor criteria (must meet 2)

- ≥ 5 cm increase in palpable splenomegaly or new splenomegaly
- Leukoerythroblastosis
- One or more constitutional symptoms
- Increase in serum LDH (Post-ET MF only)
- Anemia with a Hgb ≥ 2 mg/mL decrease from baseline (Post-ET MF only)
- Anemia or sustained loss of requirement for either cytoreductive treatment or phlebotomy (Post-PV MF only)

¹Vardiman JW, et al. *Blood*. 2009;114(5):937-951.

²Barosi G, et al. *Leukemia*. 2008;22(2):437-438.

MF Disease Features

- 85% or more of MF patients present with palpable splenomegaly at the time of diagnosis¹
- 60% to 80% of MF patients report spleen-related symptoms²
 - e.g., abdominal pain / discomfort, early satiety
- Other MF symptoms that can be present include³
 - Pruritus
 - Night sweats
 - Bone pain



Splenomegaly in MF Patient

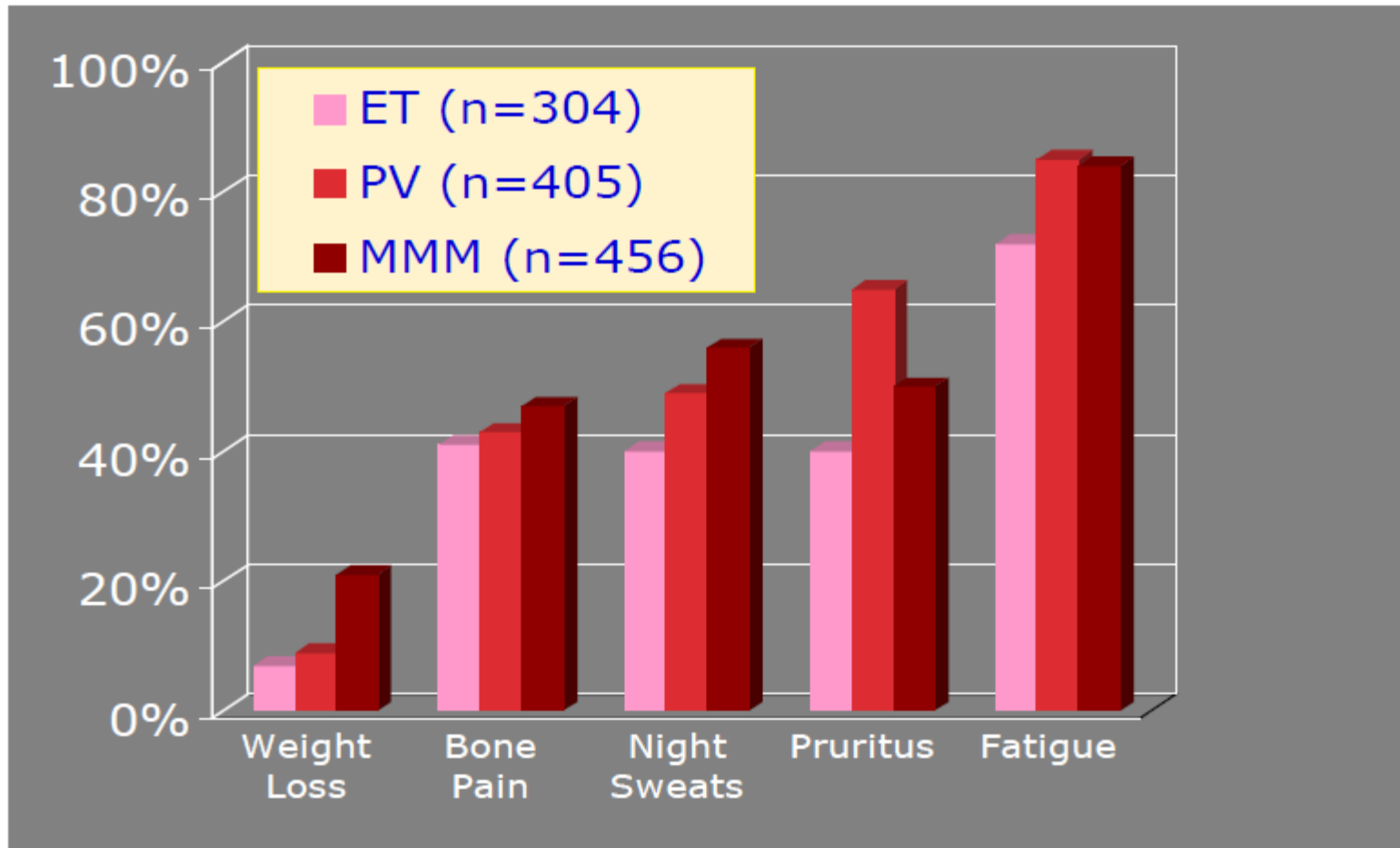
*Image courtesy of
MD Anderson Cancer Center*

¹Barosi G. *J Clin Oncol.* 1999;17:2954-2970.

²Scherber RM, et al. *Blood.* 2011;118(2):401-408.

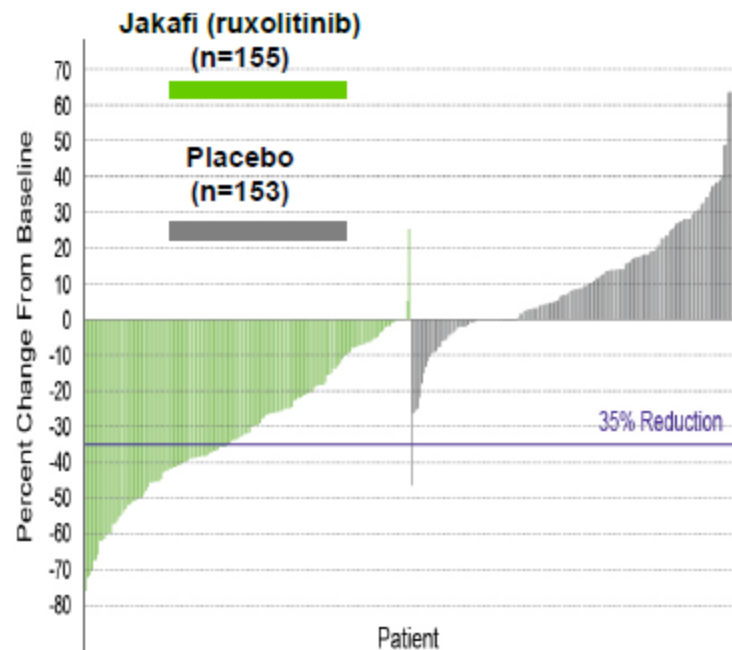
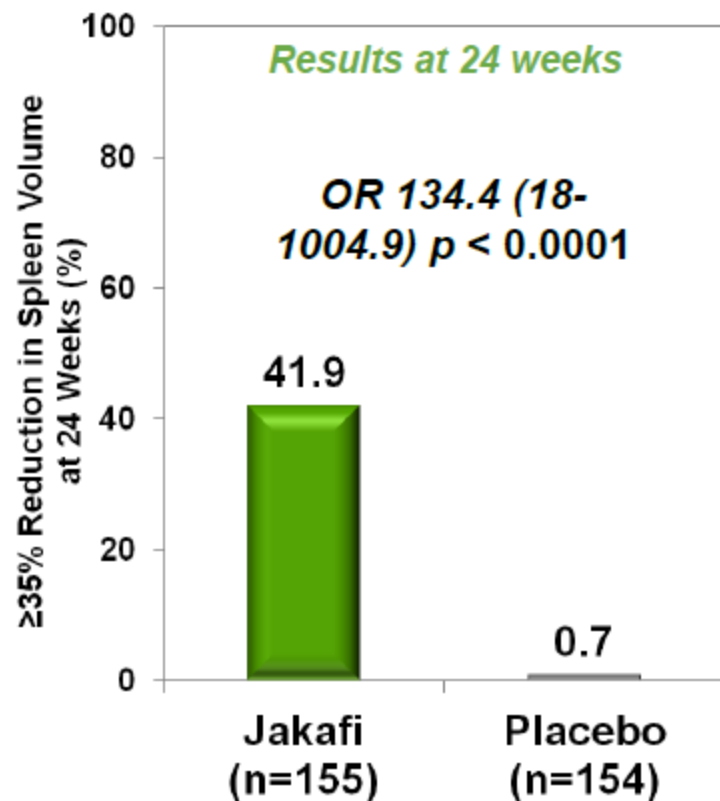
³Mesa RA, et al. *Leuk Res.* 2009;33:1199-1203.

Symptoms in 1179 MPN Patients



COMFORT-I: Spleen Volume Reduction

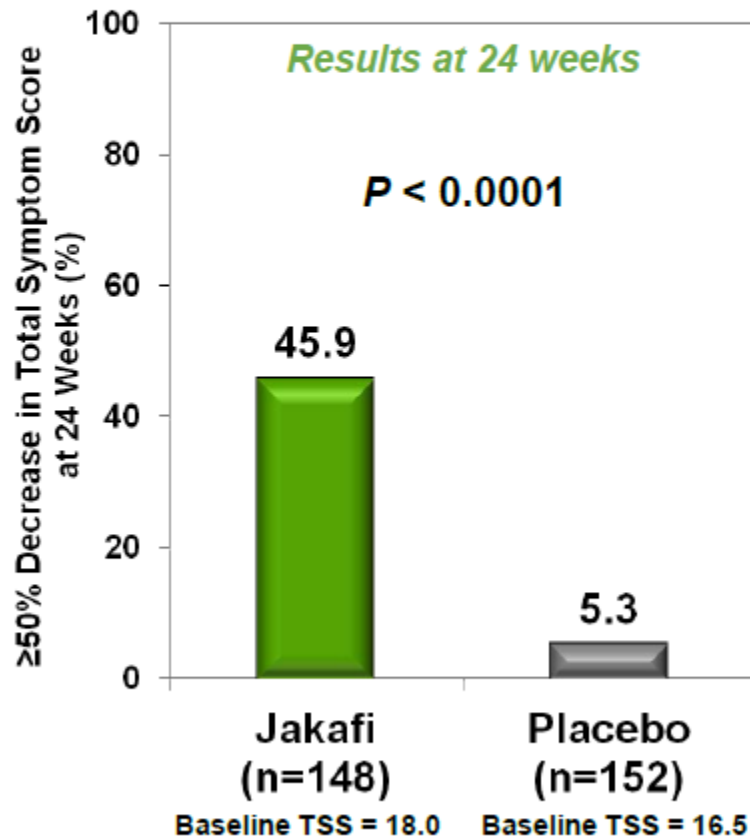
Jakafi (ruxolitinib) provided significant improvement in spleen volume



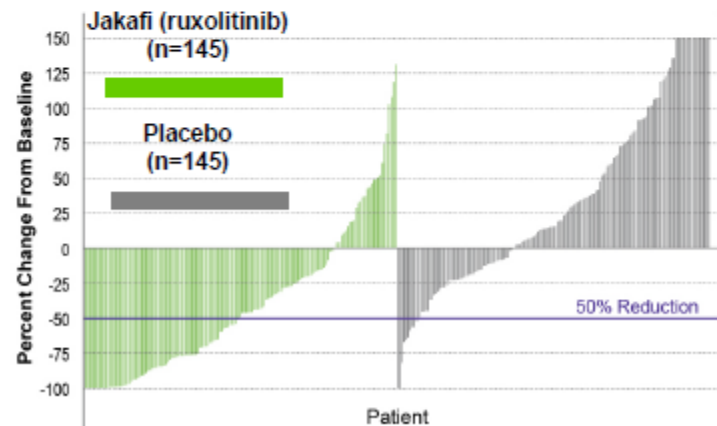
COMFORT-I: Symptom Improvement

Significant improvement in MF symptoms

(MFSAF v2.0)

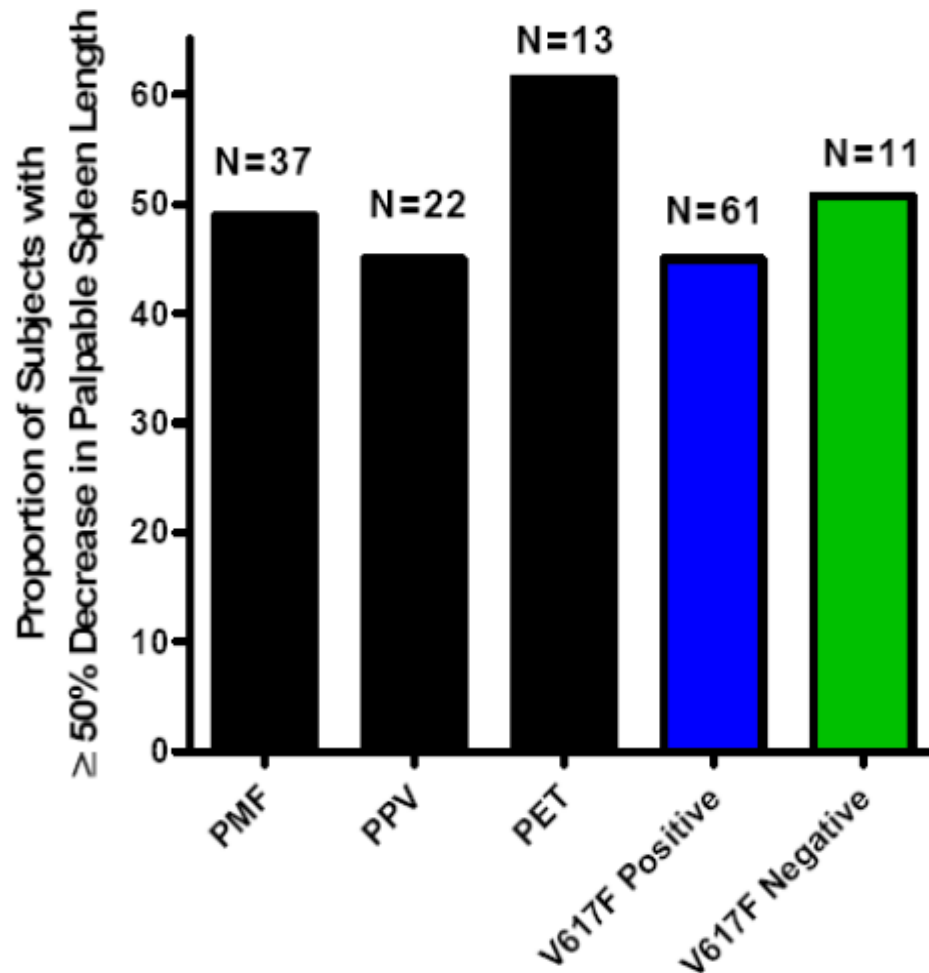


- Jakafi (ruxolitinib) provided significant improvement in total MF symptoms
 - Median time to response: <4 weeks
- Total symptom score (TSS) includes
 - Abdominal discomfort
 - Pain under the left ribs
 - Night sweats
 - Itching
 - Bone/muscle pain
 - Early satiety
- Symptom scores ranged from 0 (absent) to 10 (worst imaginable) and were added to create the daily TSS (maximum of 60)

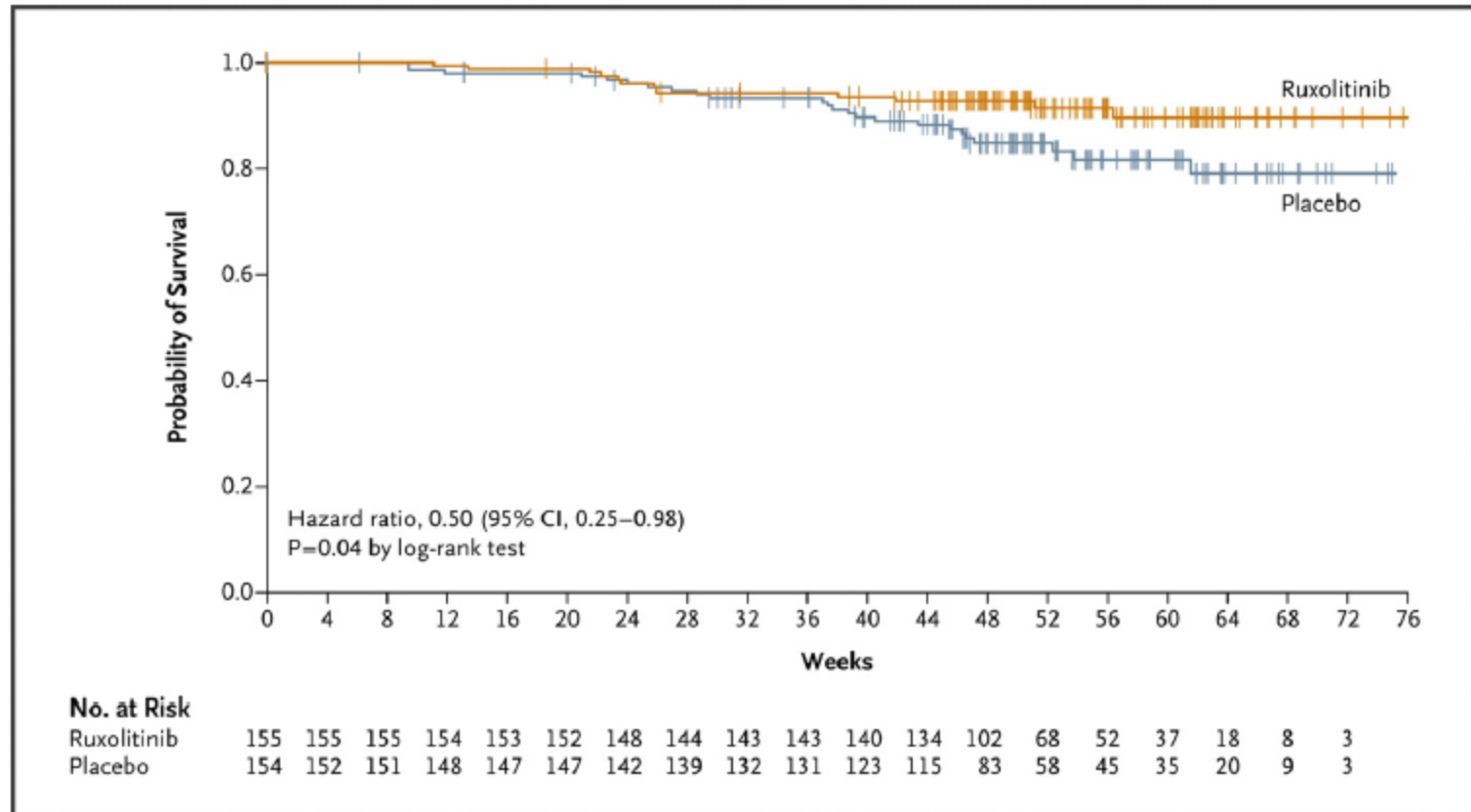


Verstovsek et al. *N Engl J Med* 2012;366:799-807
 Scherber et al. *Blood* 2011;118:401-408

Spleen Size Reduction Is Independent Of JAK Mutation Status Or Disease Subtype



Overall Survival in COMFORT I



Case 5 - Presentation

- 35yo female presents with abdominal pain and jaundice
- She has no history of liver disease, heavy EtOH intake, or thrombosis.
- Exam reveals ascites and RUQ pain, icteric sclerae

Case 5 - Presentation

- 35yo female presents with abdominal pain and jaundice
- She has no history of liver disease, heavy EtOH intake, or thrombosis. No recent surgery, immobility, trauma, or plane flights.
- Exam reveals ascites and RUQ pain, icteric sclerae
- T Bili = 12
- RUQ ultrasound with doppler reveals portal vein thrombosis.

Additional tests to consider

- Mesenteric/portal vein thrombosis without risk factor (cirrhosis):
 - JAK2 V617F mutation (~32% of all splanchnic vein thromboses associated with this mutation) (Dentali, Blood 2009, 113:5617)
 - ***about half of these patients will have abnormal blood counts at time of clot
 - Flow cytometry to evaluate for PNH (paroxysmal nocturnal hemoglobinuria via CD59, GPI deficient clone) (*rare*)
 - Most of these patients will have intermittent 'hematuria'/hemolysis
 - May also present with cerebral thromboses
 - May also have cytopenias (aplastic anemia, MDS assoc)

Questions?

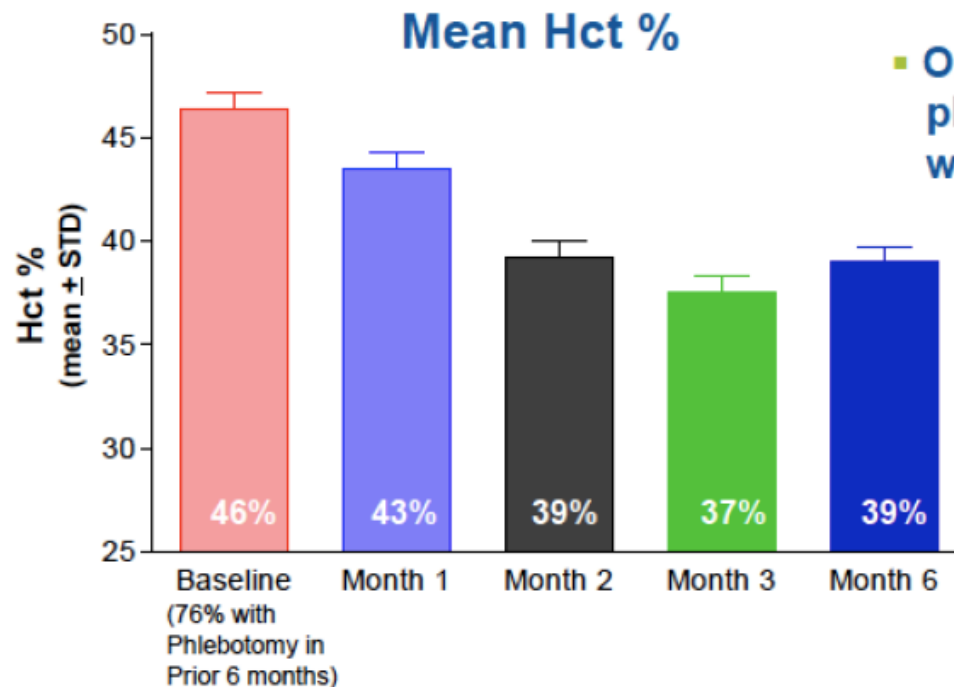


"...AND YOU CANNOT CHANGE A THING, AS YOU ARE COMPLETELY CONTROLLED BY YOUR GENES."

Treating a Molecular Disease

PV Results: Hct % (n=34)

- Normalization of Hct % Achieved in the Absence of Phlebotomy



- Only 2 subjects required phlebotomy in the first 2 weeks, none since

Ruxolitinib (Jakafi) is a JAK2 inhibitor

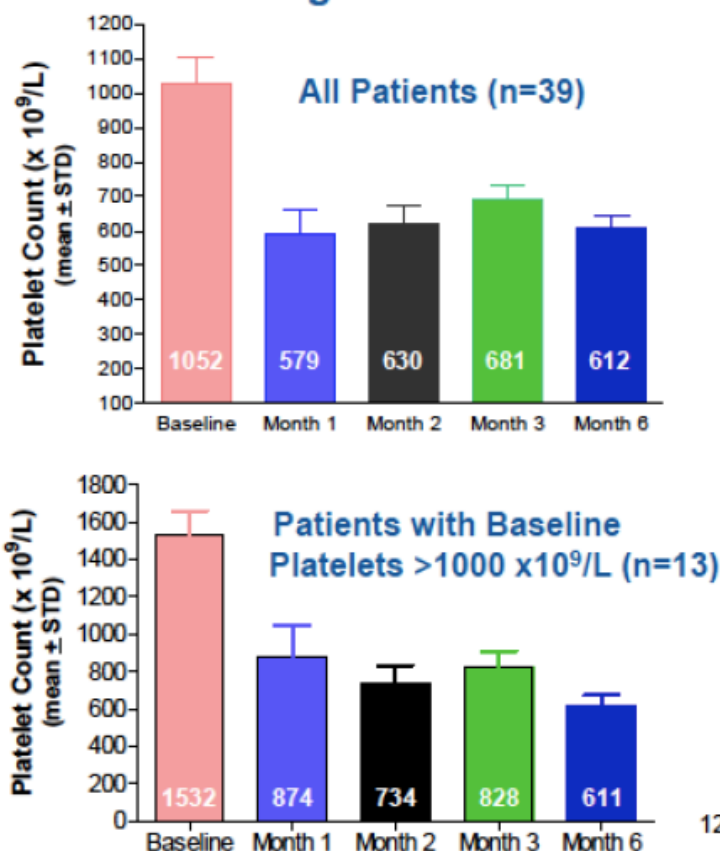


Ruxolitinib (JAK2 inhibitor)

ET Results: Platelets

- **Rapid and Sustained Reduction in Platelets**
- Baseline median platelets of 884 decreased to 558 after 6 months
- At baseline, 13 patients (33%) had platelets $> 1000 \times 10^9/L$
 - Baseline median platelets of 1443 decreased to 553 after 6 months

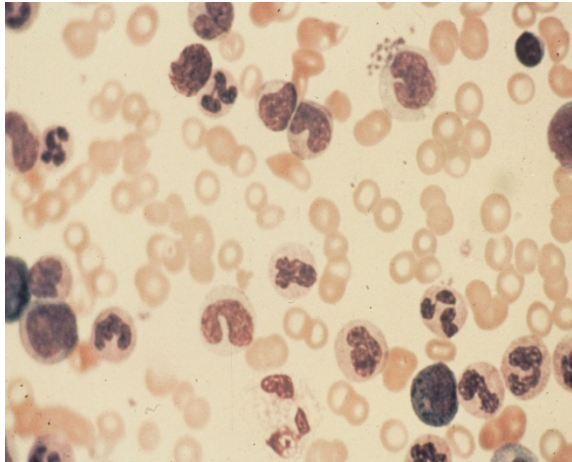
Mean Changes In Platelet Counts



Myeloid Malignancies

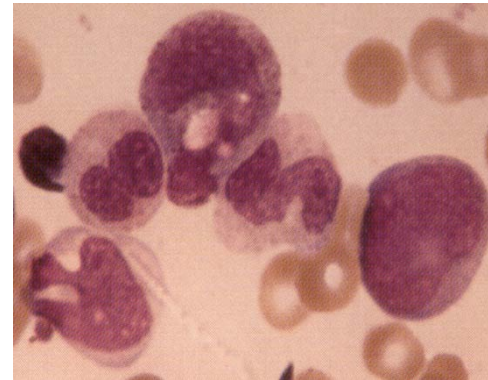
Myeloproliferative neoplasms

- enhanced proliferation/survival
- normal differentiation
- high white blood cell count
- may progress to AML



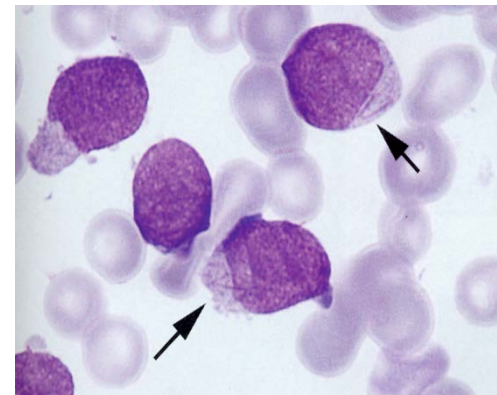
Myelodysplastic syndrome

- impaired differentiation
- low blood cell counts
- may progress to AML

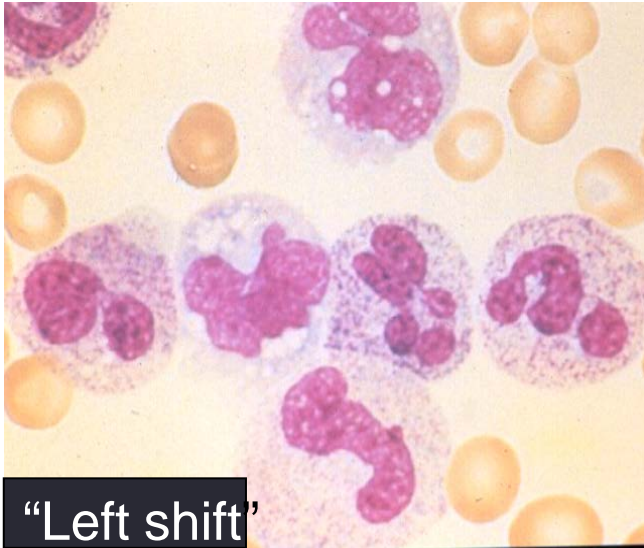


Acute myeloid leukemia (AML)

- enhanced proliferation and survival
- impaired differentiation
- limitless self-renewal

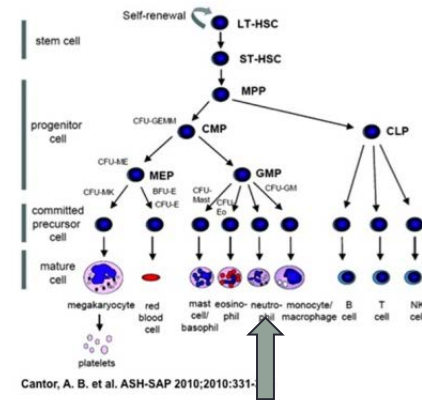


Myeloid Precursors



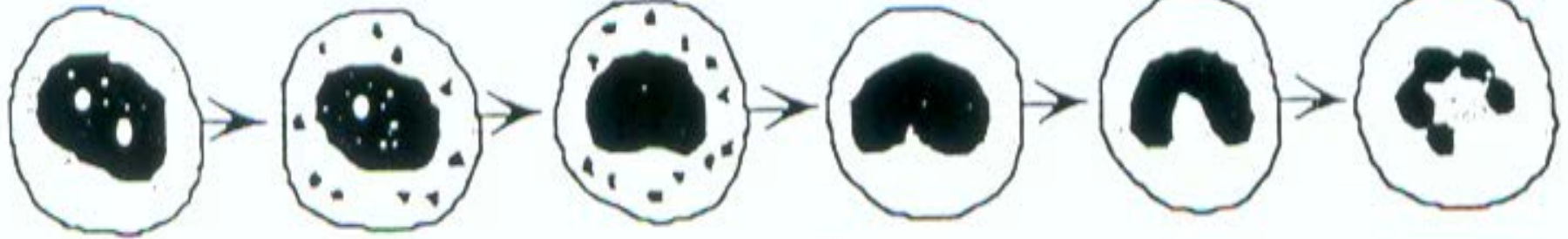
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Figure 12-3 Classical hierarchal map of hematopoietic development



Cantor, A. B. et al. ASH-SAP 2010;2010:331-

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Myeloblast Promyelocyte Myelocyte Metamyelocyte Band Neutrophil

← **“Left Shift”**