# 2. Neoplastic Proliferations of White Cells

# ~ Myeloid Neoplasms II

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#### The 2016 WHO Classification of MPN

Chronic myeloid leukemia, BCR-ABL1-positive

Chronic neutrophilic leukemia

Polycythemia vera

Primary myelofibrosis (PMF) Primary myelofibrosis, prefibrotic/early stage Primary myelofibrosis, overt fibrotic stage

Essential thrombocythemia

Chronic eosinophilic leukemia, not otherwise specified (NOS)

Myeloproliferative neoplasm, unclassifiable

# **Myeloproliferative** Neoplasms (MPN)

- A group of disorders characterized by the presence of mutated, constitutively activated tyrosine kinases or other related molecules in signaling pathways 
  ightarrow lead to growth factor independence.
- **Tyrosine kinase** Mutations do not impair differentiation.
- So the most common consequence is increase in production of one or more mature blood elements

- ► The neoplastic progenitors tend to seed secondary hematopoietic organs (spleen, liver, & LNs) → hepatosplenomegaly (neoplastic extramedullary hematopoiesis).
- MPNs often transform to AML

- Four major diagnostic entities are recognized:
- 1) Chronic myeloid leukemia (CML).
- 2) Polycythemia vera (PCV).
- 3) Primary myelofibrosis (PM).
- 4) Essential thrombocythemia (ET).

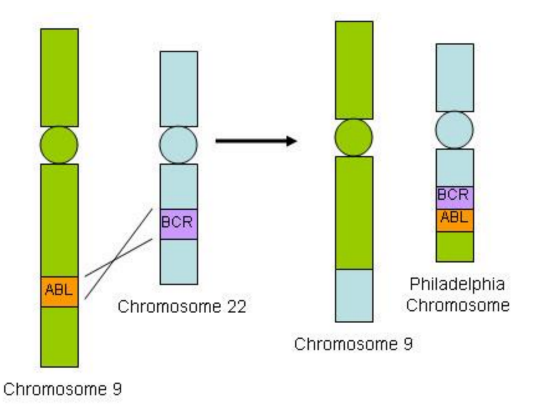
- The most common genetic abnormalities in "BCRABL-negative" MPNs are activating mutations in the tyrosine kinase JAK2.
- all MPNs have variable propensities to transform to:
- 1) a "spent phase": resembling primary myelofibrosis
- 2) a "blast crisis" identical to AML
- Both triggered by the acquisition of other somatic mutations

# Chronic Myeloid Leukemia (CML)

### Pathogenesis

- CML is distinguished from other MPN by the presence of a chimeric BCR-ABL gene, derived from portions of the BCR gene on chr. 22 & the ABL gene on chr. 9
- 95% of cases, the BCR-ABL gene is the product of a balanced t(9;22) translocation that moves ABL from chr. 9 to a position on chr. 22 adjacent to BCR.
- Translocation identified in some B-ALL.

### **CML - Pathogenesis**



# Chronic Myeloid Leukemia (CML)

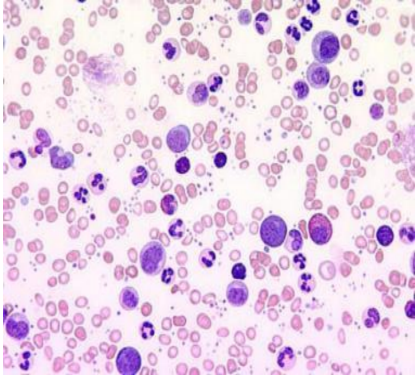
#### Pathogenesis

- Because BCR-ABL does not inhibit differentiation, the early disease course is marked by <u>excessive</u> production of relatively normal blood cells, particularly granulocytes & platelets.

# CML - Morphology

### **Peripheral blood**

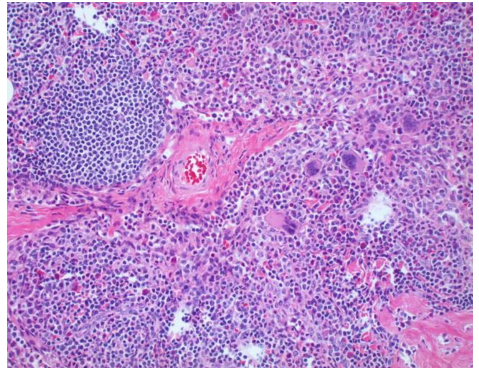
- Leukocyte count is ↑↑ (often >100,000 cells/µL).
- Circulating cells are predominantly neutrophils, metamyelocytes & myelocytes.
- Basophils, eosinophils & platelets are increased



# CML - Morphology

### **BM & spleen**

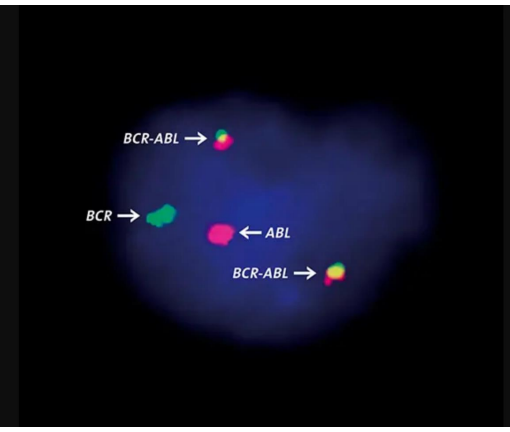
- The bone marrow is hypercellular, ^ numbers of maturing granulocytic & megakaryocytic precursors.
- ▷ Spleen resembles BM → extensive extramedullary hematopoiesis.



### **CML - Clinical features**

- Peaks in 4<sup>th</sup> & 5<sup>th</sup> decades.
- Initial symptoms usually are nonspecific (e.g., easy fatigability, weakness, weight loss).
- Necessary to distinguish CML from a leukemoid reaction (infection, stress, chronic inflammation..)

#### Fluorescence in situ hybridization (FISH) for the BCR-ABL translocation



https://www.ganeshdiagnostic.com/

### **CML – Clinical features**

Slowly progressive disease

Median survival is 3 years without treatment

- Can progress to accelerated phase
   Anemia, new thrombocytopenia (additional genetic mutations).
- Progress to blast phase:
- 1) 70% AML
- 2) 30% ALL
- Rarely progresses to spent phase with fibrosis.

### **CML – Treatment**

- Tyrosine kinase inhibitors, like Imatinib, induces sustained remissions with manageable toxicity and prevents progression to blast crisis, particularly in patients with early disease. (an example of targeted therapy)
- It suppress the proliferative drive that leads to the acquisition of additional mutations

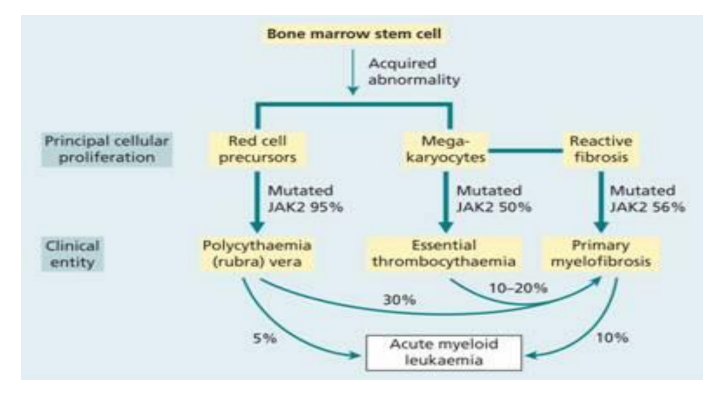
# Polycythemia Vera (PCV)

- Most clinical signs & symptoms are related to an absolute increase in red cell mass.
- Must be distinguished from <u>relative</u> polycythemia results from hemoconcentration.
- Unlike reactive <u>absolute</u> polycythemia, PCV is associated with low serum erythropoietin factor-independent growth of the neoplastic clone.

# **PCV - Pathogenesis**

- Strongly associated (>97%) with activating point mutations in the tyrosine kinase JAK2.
- JAK2 normally acts in the signaling pathways downstream of the erythropoietin receptor.

# **PCV - Pathogenesis**



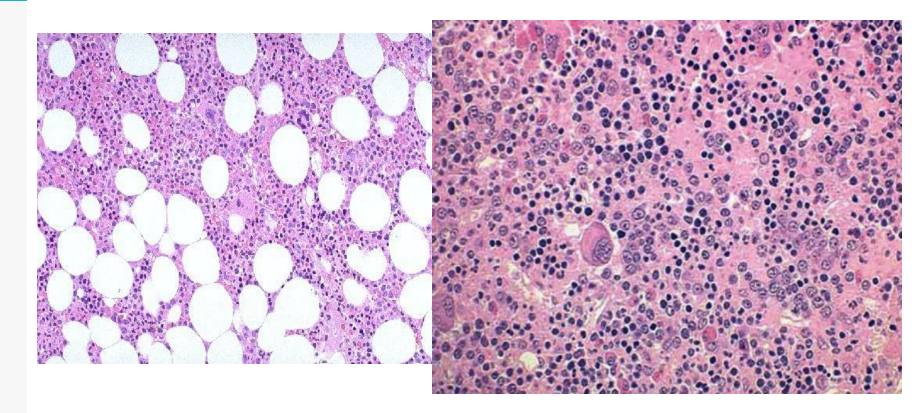
- The major anatomic changes in PCV stem from increases in blood volume and viscosity.
- ▶ Hemoglobin levels (Hb > 18,5 g/dl (♂), > 16.5 g/dl (♀))
- Congestion of many tissues is characteristic.
- Hepatomegaly & small foci of extramedullary hematopoiesis.
- ▷ Spleen usually is slightly enlarged → vascular congestion.

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- ▹ Thromboses & infarctions are common → the increased viscosity and vascular stasis.
- Platelets produced from the neoplastic clone often are dysfunctional 
   → elevated risk of thrombosis and bleeding Hemorrhages; often in GIT, oropharynx or brain.
- ▶ The peripheral blood often shows **basophilia**.

- The bone marrow is hypercellular owing to increased numbers of erythroid, myeloid, and megakaryocytic forms.
- PCV often progresses to a spent phase where the marrow is largely replaced by fibroblasts & collagen -> increase extramedullary hematopoiesis.





## **PCV – Clinical features**

Insidious, usually in late middle age.

- Patients are plethoric & often cyanotic.
- Thrombotic and hemorrhagic tendencies & hypertension.
   Headache, dizziness, GIT (hematemesis & melena) common.

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# **PCV – Prognosis**

- Without treatment, death occurs from vascular complications within months.
- The median survival is increased to about 10 years by lowering the red cell count to near normal repeated phlebotomy.
- Prolonged survival appropensity to evolve to a "spent phase" (resembling PM) ~10 years.
- Extensive marrow fibrosis, hematopoiesis shifts to the spleen, which enlarges markedly.

