

بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ



# Pathology

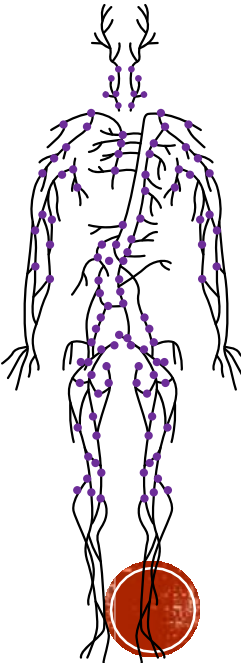
﴿وَقُلْ رَبِّ أَدْخِلْنِي مُدْخَلَ صِدْقٍ وَأَخْرِجْنِي مُخْرَجَ صِدْقٍ وَاجْعَلْ لِي مِنْ لَدُنْكَ سُلْطَانًا نَصِيرًا﴾  
ربنا آتانا من لَدُنْكَ رَحْمَةً وَهَيِّئْ لَنَا مِنْ أَمْرِنَا رَشَدًا

MID | Lecture #1

# Polycythemia

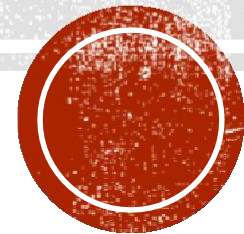
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# POLYCYTHEMIA

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# DEFINITION

- Polycythemia: Increase in total RBC **mass (not number)** above normal range, **and is defined by a numerical value, not by symptoms.**
- Erythrocytosis: increased RBC s number **(above the normal limit).**
- **In most cases, erythrocytosis will lead to polycythemia.**
- **Polycythemia can be either absolute or relative:**
  - 1) Relative polycythemia: secondary to decreased plasma volume **(seen in cases of water deprivation, severe diarrhea, vomiting, and excessive use of diuretics); a decrease in plasma volume (fluid part of the blood) leads to a high concentration of RBCs and hemoglobin, and hence polycythemia.**

Normal RBC number but low plasma volume = high RBC concentration



# ABSOLUTE POLYCYTHEMIA

Erythropoietin: a hormone produced by the kidney that stimulates bone marrow to produce RBCs.

- 2) Absolute (**true**) polycythemia: true increase in RBC mass (**and concentration**), secondary to increased RBC production by bone marrow **rather than a decrease in plasma volume**.
- Normal plasma volume but increase in RBC production = high RBC concentration
  - Can be primary (**no obvious cause for polycythemia, and the problem is within the bone marrow itself**) or secondary (**another obvious factor is causing polycythemia**):
    - Primary: autonomous, high RBC production by a bone marrow neoplasm that arose from a mutation. Polycythemia resulting from a bone marrow neoplasm is termed **polycythemia vera**, and is characterized by low erythropoietin as a result of the negative feedback response created by high numbers of RBCs.
    - Secondary: **caused by** systemic hypoxia, **which is sensed by the kidneys** → kidneys **secrete** high **amounts of** erythropoietin → increased erythropoiesis. **Therefore, unlike primary absolute polycythemia (polycythemia vera), secondary polycythemia is characterized by high erythropoietin. More on secondary polycythemia is on the next slide.**



# SECONDARY POLYCYTHEMIA

Causes of secondary polycythemia can be:

- Adaptive (a physiological response to hypoxia) : living in high altitude (where atmospheric oxygen levels tend to be lower), cyanotic heart disease (congenital heart defects that prevent the heart from pumping enough blood, leading to systemic hypoxia), and chronic pulmonary diseases including sleep apnea.
- Paraneoplastic: renal cancer, liver cancer which may secrete erythropoietin and lead to polycythemia. Recall that paraneoplastic syndrome is when a tumor, whether benign or malignant, secretes a hormone. This hormone could be endogenous to the organ from which the tumor originated, or belonging to a completely different gland (ectopic secretion).



# SECONDARY POLYCYTHEMIA

Causes of secondary polycythemia can be:

- Surreptitious (meaning hidden, also known as blood doping): endurance athletes who take erythropoietin or even blood units to improve their athletic performance. This enhances oxygen delivery to skeletal muscles, which is particularly useful for long distance runners. These athletes exhibit high hemoglobin levels (polycythemia).
- Alcohol which causes both relative and secondary absolute polycythemia by different mechanisms: frequent urination (causing relative polycythemia because of fluid loss and leading to a high hemoglobin concentration), and depressed respiration (especially during sleep, leading to hypoxia).
- Smoking which causes hypoxia.
- In secondary polycythemia, regardless of the cause, you will see: high erythropoietin, no splenomegaly



# POLYCYTHEMIA VERA

Explanation on next page

- The most important cause of primary polycythemia
- Myeloproliferative neoplasm. **Myelo : bone marrow, proliferative = production of a large number of cells (neoplastic).**
- Mutation in tyrosine kinase JAK2 in bone marrow stem cells
- Normally acts in the signaling pathway of erythropoietin receptor and other growth factor receptors
- Hematopoietic cells become less dependent on growth factors
- Excessive proliferation of erythroid, myeloid cells and megakaryocytes (panmyelosis)
- Not only increase in RBCs but WBCs and other blood cells, although Erythrocytosis is most prominent, results in polycythemia
- Splenomegaly is common **with all myeloproliferative diseases, including polycythemia vera (a primary polycythemia), but is not seen in secondary polycythemia.**



# POLYCYTHEMIA VERA

As mentioned on slide 4, polycythemia vera is a type of primary polycythemia caused by a neoplasm. It is caused by a mutation in tyrosine kinase JAK2 in bone marrow stem cells, which is important in the signaling pathway of the erythropoietin receptor. This mutation results in the permanent activation of the erythropoietin receptor (the signaling pathway proceeds even in the absence of erythropoietin). When hematopoietic stem cells are affected by this mutation, all types of blood cells are produced in large numbers (panmyelosis), with RBCs being produced the most. As previously mentioned, low erythropoietin is a characteristic of polycythemia vera.

Key words to look for with polycythemia vera: neoplasm, panmyelosis, low erythropoietin, splenomegaly, primary polycythemia, JAK2.



# SYMPTOMS OF POLYCYTHEMIA

Whether primary  
or secondary

- Plethora (red skin) / cyanosis (blue skin because of hypoxia that cannot be resolved, e.g. in patients with heart disease)
- Headache and dizziness (from hypertension resulting from the high RBC mass)
- Slow circulation (especially in small circulation and delicate organs like the retina because of the crowdedness of RBCs) and hyperviscosity cause cyanosis, blurred vision, tissue ischemia and necrosis
- Thrombosis, or bleeding (disturbed function of vWF; high numbers of RBCs may also interfere with the function of platelets and the coagulation system)

## In polycythemia vera: similar symptoms plus:

- Pruritus (aquagenic, meaning itching when exposed to hot water). Recall that polycythemia vera causes panmyelosis, so WBCs are also produced in large amounts. Basophils in particular cause a variety of symptoms as they secrete histamine, one of which is pruritus.



# SYMPTOMS OF POLYCYTHEMIA

## In polycythemia vera: additional findings:

- Peptic ulcer as histamines produced by basophils also increase stomach acidity.
- Secondary gout, which is associated with neoplasms of the bone marrow as the high turnover rate of RBCs produces high levels of uric acid (recall nucleotide metabolism) which precipitate in joints and soft tissue (arthritis, kidney stones, tophi)
- Occurs in old adults as do most tumors
- Chronic disease (cancer is always chronic)
- Treated using phlebotomy (drawing blood from a vein) to decrease the patient's hemoglobin concentration
- Spent phase: a phase that occurs after an interval of 10 years of symptoms, hematopoietic cells die and bone marrow becomes fibrotic, hematopoiesis shifts to spleen and splenomegaly exacerbates.
- Rarely and after a long time some patients enter the blast phase/crisis, where the accumulation of more mutations over time leads to the transformation from polycythemia vera into acute myeloid leukemia, which is a much more aggressive disease. It is classified as high grade with visible myeloblasts (very immature).



# LABORATORY FINDINGS OF POLYCYTHEMIA

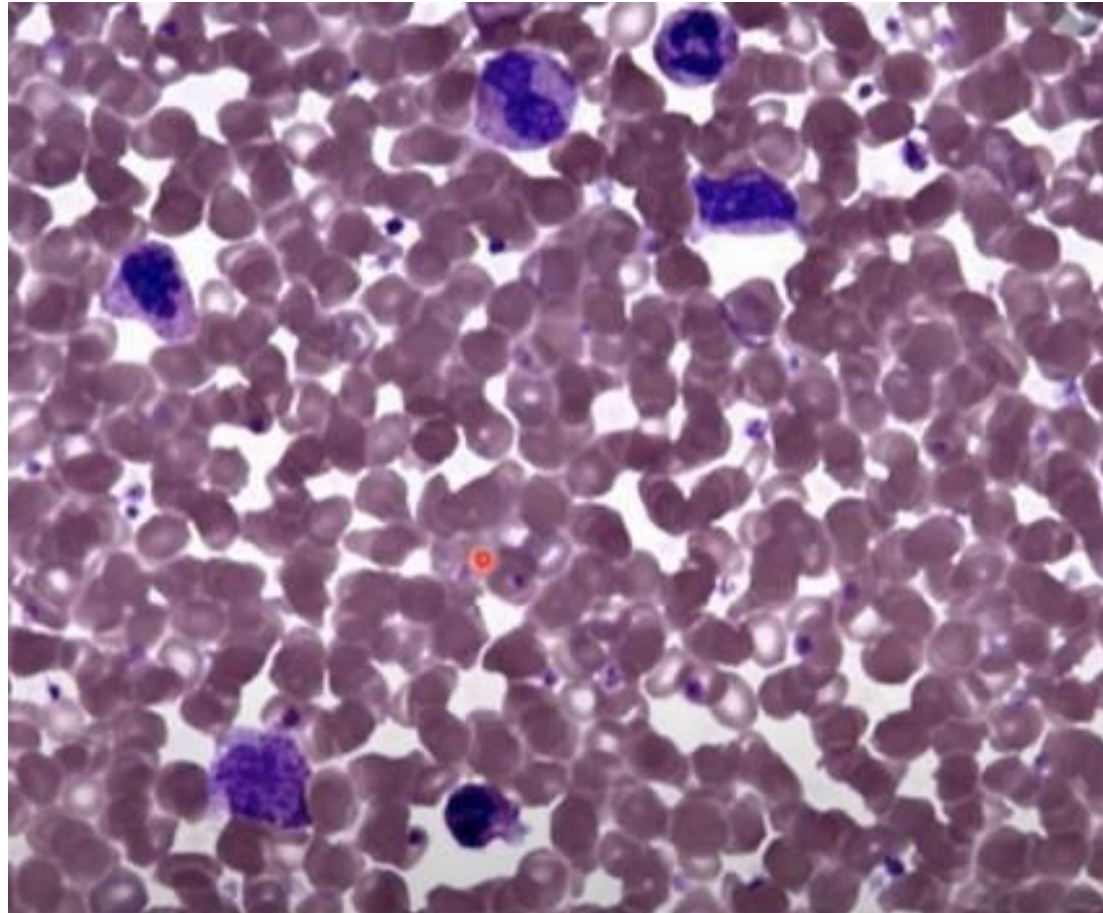
- High hemoglobin concentration (>16.5 g/dL in men, 16 in women) and high hematocrit (>49% in men, 48% in women). These numbers must be persistent in order to diagnose the patient with polycythemia vera. However, slightly lower values may be accepted if the patient has iron deficiency.
- High RBCs count (erythrocytosis!)
- These tests might be masked if iron deficiency develops, high RBC number but low iron thus low Hb content in each RBC, masking laboratory tests NOT symptoms

## In polycythemia vera: additional findings:

- Leukocytosis (including basophils, as previously mentioned) and thrombocytosis are common
- JAK2 gene mutation: positive. This mutation occurs in almost all patients with polycythemia vera. Although it is also found in other diseases, it is most common in polycythemia vera.
- Low erythropoietin level
- Hypercellular bone marrow with panmyelosis

Pan: all





Compact, crowded,  
dense RBCs. A single  
RBC's contouring  
and biconcave shape  
cannot be seen.

- Peripheral blood smear in polycythemia: packed RBCs, boundaries can't be seen



# Quiz



# For any feedback, scan the code or click on it.



Corrections from previous versions:

| Versions | Slide # and Place of Error | Before Correction | After Correction |
|----------|----------------------------|-------------------|------------------|
| V0 → V1  |                            |                   |                  |
| V1 → V2  |                            |                   |                  |

