A LABORATORY GUIDE TO CLINICAL HEMATOLOGY

Valentin Villatoro and Michelle To Northern Alberta Institute of Technology



Book: A Laboratory Guide to Clinical Hematology (Villatoro and To)

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CHAPTER OVERVIEW

1: Red Blood Cells- Normal Morphology

- 1.1: Red Blood Cell Maturation
- 1.2: Red Blood Cell Indices, Colour, and Size

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1.1: Red Blood Cell Maturation

Pronormoblast (Rubriblast, Proerythroblast)

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1.2: Red Blood Cell Indices, Colour, and Size

RBC Indices

Red blood cell indices are useful parameters when investigating suspected anemia. They help provide a general idea of the clinical picture, predict the red blood cell appearance, and aid in the classification of anemia. These indices may be calculated using the red blood cell count, hematocrit, and hemoglobin values generated by automated hematology analyzers, or directly measured in the case of MCV, depending on the model of instrument being used.^{1,2}

1. Mean Cell Volume (MCV)

MCV (fL, $10x^{-15}L$) = <u>Hct (L/L)</u> x 1000 RBC Count (x10-12/L)

*Reference range: 80-100 fL

MCV is the measurement of the average red blood cell volume and is used to classify red blood cells based on size ^{3,4}

<80 fL	Microcytic
80-100 fL	Normocytic
>100 fL	Macrocytic

Note: If the MCV is measured directly, it may be increased if there are many reticulocytes present.³

2. Mean Cell Hemoglobin (MCH)

MCH (pg, $10x^{-12}g$) = <u>Hb(g/L)</u> RBC Count (x10-12/L)

*Reference range: 28-36 pg

MCH is the measurement of the average hemoglobin weight in a red blood cell.³

3. Mean Cell Hemoglobin Concentration (MCHC)

MCHC (g/L) = Hb(g/L)Hct (L/L)

*Reference range: 310-360 g/L

MCHC is the measurement of the hemoglobin concentration in a population of red blood cells. This is used to denote the colour of the red blood cell population.^{3,5}

<310 g/L	Hypochromic
310-360 g/L	Normochromic
>360 g/L	Check for spherocytes or errors in Hb/Hct measurement (interferences)

4. Red Blood Cell Distribution Width (RDW)

RDW is the coefficient of variation or standard deviation of the MCV. Similar to the RBC indices, it is determined by automated cell counting instruments and is used to predict the degree of red blood cell size variation, known as anisocytosis.²⁻⁴





An increase in the RDW would indicate a higher presence of anisocytosis on the peripheral blood smear.²⁻⁴

A decrease in the RDW is not associated with any known abnormalities.²⁻⁴

*Reference range: 11.5-14.5%

*Please be aware that the reference ranges provided in this book were obtained from multiple sources and may not accurately reflect the values used in your laboratory. References ranges vary depending on institution, patient population, methodology and instrumentation. Laboratories should establish their own ranges based on these factors for their own use.

Size

As previously described, MCV is used to classify red blood cells based on their size.

1. Normocytic RBCs

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Peripheral blood smears showing normochromic, normocytic red blood cells. From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R3RJ4995W

Image 2: 60x oil immersion. https://doi.org/10.7939/R35M62P2N

The MCV of normocytic RBCs fall within the normal reference ranges of 80-100 fL and the size should be around 7-8µm.^{6,7}

Size comparison: Mature red blood cells are about the size of the nucleus of a small lymphocyte. It i also a pproximately three normal red blood cells should fit within a normal neutrophil. ⁶

2. Microcytic RBCs

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Peripheral blood smear images show numerous microcytic red blood cells. A small lymphocyte is present and can be used for a size comparison. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3599ZH07

Image 2: 100x oil immersion. https://doi.org/10.7939/R3WS8J199

A microcytic red blood cell measures less than 7-8µm, and has an MCV that is <80 fL. The hemoglobin concentration (MCHC) can be normal or decreased, and can help differentiate different clinical conditions or severities of anemia.

Microcytes are commonly seen with any abnormalities involving hemoglobin synthesis and thus cells often also appear hypochromic. ^{3,6,7}

Size comparison: Microcytes are smaller than the size of the nucleus of a normal small lymphocyte. If a normal neutrophil is being used for comparison, more than three microcytes can easily fit in a normal neutrophil.

Associated Disease/Clinical States ^{7,8}:

TAILS: Thalassemias Anemia of chronic inflammation Iron Deficiency Anemia Lead poisoning S ideroblastic Anemia

3. Macrocytic RBCs (Round/Oval)





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CHAPTER OVERVIEW

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2.1: Poikilocytosis

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Peripheral blood smears demonstrating marked poikilocytosis. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3KD1R163

Image 2: https://doi.org/10.7939/R3RV0DG2H

General Peripheral Blood Smear Description:

Poikilocytosis is a general term used to describe the collective presence of various abnormal red blood cell shapes on a peripheral blood smear. Normal red blood cell morphology is described in the previous two chapters but under certain clinical conditions, they can take on various shapes or morphologies. When certain red blood cell shapes are predominant, this may be associated with specific disease states.¹⁻³

Associated Disease/Clinical States: ³⁻⁴

Hemolytic anemias

Thalassemia

Myelofibrosis

Hereditary pyropoikilocytosis

Note: See the rest of the chapter for other disease states related to a specific predominant abnormal morphology.

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

3. Turgeon ML. Erythrocyte morphology and inclusions. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 99-111.

4. Coetzer TL, Zail S. Introduction to hemolytic anemias: intracorpusculardefects: I. hereditary defects of the red cell membrane. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 176-95.

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2.2: Acanthocytes (Spur Cells)



An image of a peripheral blood smear containing some acanthocytes. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3MP4W32X







An image of a peripheral blood smear containing some acanthocytes (examples shown with arrows). 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3MP4W32X

Cell Description:

Red blood cells appear small and dense, lacking an area of central pallor with multiple spiky projections (spicules) of varying lengths protruding from the membrane. Projections are irregularly distributed around the cell membrane.¹⁻³

Cell Formation:

Acanthocyte formation occurs as a result of either hereditary or acquired membrane defects. Defects that cause an imbalance between the membrane cholesterol and lipid content affect the RBC's ability to deform resulting in more rigid plasma membrane. Red blood cells are then remodelled in circulation, resulting in an acanthocyte.^{1,3,4}

Associated Disease/Clinical States: 1-5

Abetalipoproteinemia (Inherited) Lecithin-cholesterol acyltransferase (LCAT) Deficiency Liver Disease Post-splenectomy Pyruvate Kinase (PK) Deficiency

References:





1. Cochran-Black D. Hemolytic anemia: membrane defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 317-33.

2. Manchanda N. Anemias: red blood morphology and approach to diagnosis. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 284-96.

3. Turgeon ML. Normal erythrocyte lifecycle and physiology. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 71-98

4. Harmening D. The red blood cell: structure and function. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 759-792.

5. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

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2.3: Agglutination



A peripheral blood smear demonstrating agglutination of red blood cells. 10x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3C824V90





A peripheral blood smear demonstrating severe agglutination of red blood cells. 10x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R34J0BC72





A peripheral blood smear demonstrating autoagglutination of red blood cells. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3599ZH26







An image from a peripheral blood smear showing agglutination of red blood cells. The arrow points to a cluster of red blood cells. 50x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3SB3XD80

Cell Description:

This alteration of RBC distribution presents as irregular and random grape-like clusters or clumps. Agglutination is differentiated from Rouleaux by the lack of linear chains or "coin staking". ¹⁻⁴

The outlines of individual cells may not be evident.

Cell Formation:

Agglutination is caused by the formation of antibody-antigen complexes and occurs at room temperatures. Auto-agglutination is produced as a result of a complex formed between the patient's own RBC antigens and antibodies, mediated by cold-reacting antibodies. Agglutination can be reversed when the blood sample is warmed to 37°C.^{1,2,5}

Associated Disease/Clinical States: ²⁻⁴

Cold Hemagglutinin Disease

Paroxysmal Cold Hemoglobinuria

Cold Autoimmune Hemolytic Anemia

Note: Formation is NOT reversed with the addition of saline.⁵

References:

1. Hemolytic anemia: membrane defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 317-33.





2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

3. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

4. Turgeon ML. Normal erythrocyte lifecycle and physiology. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 71-98.

5. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

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2.4: Bite (Keratocyte) and Blister (Helmet) Cells



An image of a peripheral blood smear with bite cells present (indicated with arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3CC0V83F







An image of a peripheral blood smear with bite cells present (indicated with arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3V698T3M





A peripheral blood smear demonstrating a blister cell (shown with an arrow). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3H12VP5B







A peripheral blood smear with a blister cell present (shown with an arrow). From MLS Collection, University of Alberta, https://doi.org/10.7939/R3HQ3SD75

Cell Description:

Bite cells are red blood cells that contain a semi-circular indent on the edge of their membrane, giving the appearance of a bite being taken out of the cell.¹ Blister cells on the other hand, have cytoplasmic projections that fuse together, creating a vacuole on the edge of the membrane, giving the appearance of a blister.²

Cell Formation:

Bite and Blister cells are often seen together, and may form through various mechanisms. Red blood cells originally containing inclusions are "pitted" or removed by macrophages in the spleen, resulting in bite or blister cells.³ When the red blood cell is impaled by fibrin strands, the membrane can reform and produce a vacuole which results in a blister cell.^{2,3}

Bite cells can also form when a blister cell ruptures.⁴

Associated Disease/Clinical States:^{2,4,5}

Microangiopathic Hemolytic Anemias (MAHAs)

Mechanical Hemolysis (i.e. mechanical heart-valves)

Heinz body hemolytic anemias (G6PD Deficiency, Thalassemia)

Note: Bite and blister cells are mainly seen in clinical states where Heinz bodies are formed.²

References:





1. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

2. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

3. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

4. Turgeon ML. Erythrocyte morphology and inclusions. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 99-111.

5. Julius CJ, Schaub CR. Hypoproliferative anemia: anemia associated with systemic diseases. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 280-304

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2.5: Dimorphic Population

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Images show peripheral blood smears containing a dimorphic population (hypochromic-microcytic, and normochromic-normocytic red blood cells). From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R3T14V447

Image 2: 50x oil immersion. https://doi.org/10.7939/R3V11W18D

Cell Description:

The peripheral blood smear shows that there are two distinct red blood cell populations present. The different red blood cell populations that may be seen are normocytic/normochromic, microcytic/hypochromic, macrocytic/normochromic.^{1,2}

Cell Formation:

The cause for the formation of a dimorphic red blood cell population varies depending on the clinical condition.

Associated Disease/Clinical States: 1-2

Sideroblastic Anemia

Myelodysplastic Syndrome (MDS)

Iron, Vitamin B12, Folate deficiency (and during the early treatment stage)

Post-transfusion

Erythropoietin Therapy

Note: $RDW > 14.5\%^3$

References:

1. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

2. Constantino BT. The red cell histogram and the dimorphic red cell population. Lab Med [Internet]. 2011 May 1 [cited 2018 Jul 23];42(5):300–8. Available from: http://dx.doi.org/10.1309/LMF1UY85HEKBMIWO

3. Rodak BF, Carr JH. Variations in size and color of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 89-92.

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2.6: Echinocytes (Burr Cells)

Cell Description:

The red blood cell has multiple evenly distributed projections that are of equal length that cover the entire surface of the cell.¹ Cells usually have an area of central pallor.²

Cell Formation:

Commonly form due to a "glass effect" during peripheral blood smear preparation with glass slides. Glass slides can release basic substances that can induce echinocyte formation.³

Another cause of echinocyte formation is due to storage conditions. Echinocytes can naturally form in whole blood that has been stored at 4°C after a few days (i.e. Blood to be transfused).³

The formation of echinocytes is a reversible process and can reform a natural discoid shape.³

Echinocytes are often considered artifact from the smear making process (drying or staining) and may not be reported, depending on individual laboratory protocol.

Associated Disease/Clinical States: ¹⁻³

Artifact Post-transfusion Burns Liver Disease Pyruvate Kinase (PK) Deficiency Uremia Microangiopathic Hemolytic Anemias (MAHAs)

References:

1. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

2. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

3. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

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2.7: Elliptocytes and Ovalocytes



A peripheral blood smear with numerous elliptocytes present. 100x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R35H7C887







A peripheral blood smear showing elliptocytes along with various other poikilocytosis. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3H98ZV4C

Cell Description:

Elliptocytes: Red blood cells are cigar or pencil shaped with parallel sides and an area of pallor.^{1,2}

<u>Ovalocytes:</u> Are red blood cells that are oval or egg shaped.^{1,2}

<u>Macro-ovalocytes:</u> Ovalocytes that are larger than a normal red blood cells.³

Southeast Asian Ovalocytes: Ovalocytes show two transverse zones of pallor separated by a transverse zone of cytoplasm.⁴

Cell Formation:

Elliptocytes and ovalocytes are formed only after the red blood cell has reached its normal and mature morphology. Elliptical features develop over time as the cell undergoes stress in the circulation.^{1,2}

Formation occurs due to erythrocyte membrane protein defects resulting in an increase in mechanical weakness and membrane fragility.^{1,3,5}

Hereditary elliptocytosis occurs due to defects in the horizontal protein linkages between the membrane and cytoskeleton. (α -spectrin, *B*-spectrin, protein 4.1, glycophorin C).^{5,6}

Associated Disease/Clinical States: 1,4,7

Elliptocytes:	Ovalocytes:
Hereditary elliptocytosis	Hereditary ovalocytosis (Southeast Asian Ovalocytosis)
Thalassemia	Megaloblastic Anemia (Macro-ovalocytes)
Iron deficiency Anemia	

References:





1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

2. Manchanda N. Anemias: red blood morphology and approach to diagnosis. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 284-96.

3. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

4. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

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2.8: Pyknocytes

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Images of peripheral blood smears with pyknocytes present. Pyknocytes are indicated by the arrows. 100x oil immersion. From MLS Collection, University of Alberta.

Image 1: https://doi.org/10.7939/R33B5WQ09

Image 2: https://doi.org/10.7939/R3VX06J4H

Image 3: https://doi.org/10.7939/R3KS6JM01

Cell Description:

Pyknocytes appear as small, dark, pyknotic RBCs that lack central pallor and have an irregular, non-spherical shape.

Cell Formation:

Pyknocytes are rare, but may form as a result of red blood cell dehydration or oxidative damage.

Associated Disease/Clinical States: (6 Mary Louise ch 6 pg 103)

Pyruvate Kinase Deficiency

Glucose-6-phosphate (G6PD) Deficiency

Acute and severe hemolytic anemias

Infantile pyknocytosis

References:

1. Turgeon ML. Normal erythrocyte lifecycle and physiology. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 103.

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2.9: Rouleaux

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Images of peripheral blood smears with rouleaux present. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3445HT5R

Image 2: 10x magnification. https://doi.org/10.7939/R3HQ3SD56

Image 3: 50x oil immersion. https://doi.org/10.7939/R3HM5313R

Cell Description:

Red blood cells are arranged into rows or linear chains, appearing on top of one another in a "coin stacking" fashion. The outlines of the the individual cells are usually seen.^{1,2}

Cell Formation:

Can form naturally after blood is collected and allowed to sit for a long period of time.¹

Similar morphology can be seen in the thick areas of a blood smear.¹ Pathological rouleaux is only reported when seen in the thin areas of a peripheral blood smear where a differential would usually be performed.³

In pathological states, the increase of plasma proteins (e.g. fibrinogen, globulins) will coat the red blood cells and cause them to become "sticky" and result in rouleaux formation.^{1,4}

Associated Disease/Clinical States:^{2,5}

*Associated with any condition that results in the increase of plasma proteins

Acute and chronic inflammatory disorders

Plasma Cell Myeloma (Multiple Myeloma)

Polyclonal or monoclonal hyperglobulinemia

Note: Unlike with agglutination, the formation of rouleaux can be reversed with the addition of saline.²

References:

1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

2. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

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4. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

5. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

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2.10: Schistocytes

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Images show peripheral blood smears with schistocytes present (indicated by the arrow). From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R34Q7R51T

Image 2: 50x oil immersion. https://doi.org/10.7939/R3N29PN8H

Image 3: https://doi.org/10.7939/R31R6NG3M

Cell Description:

Fragmented red blood cells with varying shapes and sizes. Cells often appear small, with multiple pointed and angular ends and lack an area of central pallor.^{1,2}

Cell Formation:

Formed in circulation when a red blood cell is damaged by mechanical means (e.g. damaged by fibrin strands or mechanical heart valves). The presence of schistocytes suggests an intravascular hemolytic process is occurring^{2,3}

Associated Disease/Clinical States: 1,2,4,5

Microangiopathic Hemolytic Anemia (MAHAs)

Macroangiopathic Hemolytic Anemias

Renal graft rejection

Severe burns

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

2. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

3. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

4. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

5. Manchanda N. Anemias: red blood morphology and approach to diagnosis. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 284-96.

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2.11: Sickle Cells (Drepanocytes)

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A peripheral blood smear demonstrating sickle cells (indicated by arrows). From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R3TQ5RV8M

Image 2: 100x oil immersion. https://doi.org/10.7939/R3X05XT8X

Cell Description:

Red blood cells that lack an area of central pallor, are thin, and appear curved or S-shaped (cells resemble a sickle or crescent). The ends of the cell are pointed.¹

Cell Formation:

A genetic mutation in the β globin chain results in the production of abnormal hemoglobin S. The mutation results in an amino acid substitution in the 6th position from glutamine to valine. Red blood cells have normal morphology under normal conditions but under hypoxic conditions (decreased oxygen tension), hemoglobin S polymerizes and causes the red blood cell to assume the characteristic sickle shape.² Sickle cell formation causes the red blood cell to become rigid and inflexible.³

Factors that contribute to hemoglobin S polymerization and RBC sickling include:

- Low oxygen saturation
- Decreased pH
- Increased 2,3-BPG
- Dehydration

These factors affect either the oxygenation or concentration of hemoglobin S inside the red blood cell, leading to polymerization and sickling.

Note: Formation of sickle cells can be reversible when hypoxic conditions are corrected however not all sickle cells have the ability to revert back to a normal morphology.^{2,3}

Associated Disease/Clinical States:¹

Sickle Cell Disease (Homozygous Hemoglobin S disease)

Hemoglobin SC Disease

Note: Sickle cells not usually seen in heterozygous hemoglobin S (Hemoglobin AS or Sickle Cell Trait).¹

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

2. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

3. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

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2.12: Spherocytes

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Images show peripheral blood smears with numerous spherocytes present (examples are indicated by arrows). From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R39883320

Image 2: 50x oil immersion. https://doi.org/10.7939/R3XG9FS24

Cell Description:

Round red blood cells that lack an area of central pallor. Cells often appear darker and smaller than a normocytic red blood cell.¹

Cell Formation:

Formation of spherocytes in circulation occurs due to a partial loss of the red blood cell membrane. This can occur when RBCs are not fully phagocytosed by macrophages during extravascular hemolysis.² Cellular content remains the same and this leads to a decrease in the surface to volume ratio and spherocyte formation.³

Hereditary Spherocytosis: the formation of spherocytes occurs due to the defects in the vertical protein linkages between the membrane and cytoskeleton, resulting in a loss of unsupported RBC membrane and spherocyte formation.⁴

Associated Disease/Clinical States:^{1,5-7}

Hereditary Spherocytosis

Warm Auto-Immune Hemolytic Anemia (WAIHA)

Drug-Induced Immune Hemolytic Anemia

Allo-Immune-mediated hemolysis (delayed hemolytic transfusion reactions, Hemolytic Disease of the Fetus and Newborn)

Glucose-6-Phosphate (G6PD) Deficiency

Transfused cells (storage lesion)

Severe burns

Note: Spherocytes have an increased MCHC (>360 g/L).⁵

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

2. Doig K. Introduction to increased destruction of erythrocytes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 348-66.

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5. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

6. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

7. Turgeon ML. Erythrocyte morphology and inclusions. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 99-111.





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2.13: Stomatocytes

Cell Description:

Red blood cells that appear to have an area of central pallor that is slit-like (stoma) instead of circular.¹ Cells are normal in size but lack it's normal biconcavity.² By using electron microscopy, cells instead appear "cup" or "bowl" shaped.³

Cell Formation:

Cell formation is due to a membrane defects (acquired or inherited) that results alterations in cell volume. Both an increase (hydrocytosis) and a decrease (xerocytosis) in cell volume can cause the production of stomatocytes.²

Associated Disease/Clinical States:^{1,2,4,5}

Hereditary Stomatocytosis Artifact

Alcoholism

Liver disease

Rh Null Disease

Drugs (effects are often reversible)

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

3. Bain BJ. Morphology of blood cells. In: Blood cells: a practical guide [Internet]. 5th ed. Chichester, UK: John Wiley & Sons, Ltd; 2015 [cited 2018 Jul 10]: 67-185. Available from: http://doi.wiley.com/10.1002/9781118817322

4. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

5. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

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2.14: Target Cells (Codocytes)

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Images show peripheral blood smears with numerous target cells present (examples are indicated by arrows). From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R3R49GR23

Image 2: 40x magnification. https://doi.org/10.7939/R3NG4H71X

Image 3. 60x oil immersion. https://doi.org/10.7939/R3R78644B

Cell Description:

Target cells adopt a "bullseye" morphology where hemoglobin is concentrated in the center and on the periphery with a colourless zone in between the two areas. Other target cells may also look folded or bell shaped.¹⁻³

Note: The target cell membrane is thinner than normal cells.^{1,4}

Cell Formation:

Liver Disease: membrane cholesterol concentration is reduced, decreasing the tensile strength of the membrane, resulting in target cell formation.^{2,3,5}

Artifact: Target cell formation occurs when blood smears are made when humidity is high.¹

Hemoglobinopathies: There is a uneven distribution of hemoglobin within the cell, and an increased surface area to volume ratio.¹

Note: Target cells have an increased surface area to volume ratio and decreased osmotic fragility.^{1,3}

Associated Disease/Clinical States:^{1,2,5}

Hemoglobinopathies (Hemoglobin C Disease, Sickle cell Disease, Thalassemia, etc.)

Iron deficiency anemia

Obstructive Liver disease

Splenectomy

Artifact

References:

1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

2. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

3. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

4. Bain BJ. Morphology of blood cells. In: Blood cells: a practical guide [Internet]. 5th ed. Chichester, UK: John Wiley & Sons, Ltd; 2015 [cited 2018 Jul 10]: 67-185. Available from: http://doi.wiley.com/10.1002/9781118817322

5. Turgeon ML. Erythrocyte morphology and inclusions. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 99-111.

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2.15: Tear Cells (Dacrocytes, Teardrops)

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Images show peripheral blood smears with numerous tear cells (examples indicated by arrows). From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R37S7J768

Image 2: https://doi.org/10.7939/R3D21S07J

Image 3: https://doi.org/10.7939/R38P5VR5R

Cell Description:

Red blood cells that are teardrop or pear shaped with one blunt projection.¹ The size of these cells are variable.²

Cell Formation:

Red blood cells with inclusions: Teardrop cells form from these cells when the cells attempt to pass through the microcirculation resulting in the pinching the cell as the part containing the inclusion is left behind.²

Myelophthisis: displacement of normal hematopoietic tissue in the bone marrow by abnormal cells (malignancies) or fibrosis, leading to bone marrow crowding and pinching of RBCs as they as pushed out of the bone marrow.

Associated Disease/Clinical States:¹⁻³

Primary myelofibrosis

Thalassemia

Megaloblastic Anemia

Sideroblastic Anemia

Myelophthisic Anemia

Drug-induced Heinz body formation

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106, 289.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

3. Bain BJ. Morphology of blood cells. In: Blood cells: a practical guide [Internet]. 5th ed. Chichester, UK: John Wiley & Sons, Ltd; 2015 [cited 2018 Jul 10]: 67-185. Available from: http://doi.wiley.com/10.1002/9781118817322

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CHAPTER OVERVIEW

3: Red Blood Cells- Abnormal RBC Inclusions

- 3.1: Basophilic Stippling
- 3.2: Cabot Rings
- 3.3: Heinz Bodies
- 3.4: Hemoglobin H (Hb H)
- 3.5: Hemoglobin C Crystals
- 3.6: Hemoglobin SC Crystals
- 3.7: Howell-Jolly Bodies
- 3.8: Pappenheimer Bodies (Siderotic Granules)
- 3.9: Bacteria and Fungi
- 3.10: Malaria
- 3.11: Babesia
- 3.12: Trypanosomes

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3.1: Basophilic Stippling

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Images show peripheral blood smears with basophilic stippling in the red blood cells (indicated by arrows). From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R3G15TS2N

Image 2: 100x oil immersion. https://doi.org/10.7939/R3RJ4993X

Image 3: 100x oil immersion. https://doi.org/10.7939/R3J960R3H

Appearance:

Multiple dark blue-purple granules that are distributed throughout the red blood cell. Granules can appear coarse, fine, round, and/or irregularly shaped, and are present in numerous numbers.^{1,2} Typically, only coarse basophilic stippling is reported.

Inclusion composition:¹⁻³

Aggregates of ribonucleic Acid (RNA)

Associated Disease/Clinical States: 1,2

Lead toxicity

Thalassemia

Abnormal heme synthesis

References:

1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

3. Bain BJ. Morphology of blood cells. In: Blood cells: a practical guide [Internet]. 5th ed. Chichester, UK: John Wiley & Sons, Ltd; 2015 [cited 2018 Jul 10]: 67-185. Available from: http://doi.wiley.com/10.1002/9781118817322

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3.2: Cabot Rings



An image from a peripheral blood smear showing a cabot ring. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3B854027

Appearance:

Red-purple inclusions that appear as a loop, ring, or figure-eight shape and span the diameter of the red blood cell. 1-2 cabot rings may be seen in a single cell.¹

Note: Finding is rare, and not to be confused with malaria.

Inclusion composition:¹

Remnant microtubules of mitotic spindle

Associated Disease/Clinical States:¹⁻³

Myelodysplastic Syndrome (MDS; Dyserythropoiesis)

Megaloblastic Anemia

Lead poisoning

References:

1. Rodak BF, Carr JH. Inclusions in erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 107-14.





2. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

3. Turgeon ML. Erythrocyte morphology and inclusions. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 99-111.

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3.3: Heinz Bodies



A peripheral blood smear stained with a supravital stain demonstrating numerous heinz bodies (indicated by arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R35718396







A supravital stained peripheral blood smear showing numerous heinz bodies (indicated by arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3901ZX4N

Appearance:

Inclusions are not visible on Wright or Romanowsky-stained blood smears. Inclusions can only be visualized with supravital stains. After staining, Heinz body inclusions appear dark blue-purple and are located at the periphery of the red blood cell at the membrane. The inclusions are round and look as if they are being ejected out of the cell.^{1,2}

Note: Heinz bodies are usually not seen, as they are normally removed by splenic macrophages.³ Their presence indicates an increase in hemoglobin denaturation and precipitation, seen in numerous conditions that result in hemoglobin instability, oxidative damage, or excess globin chains.

Inclusion composition:^{1,2}

Denatured or precipitated hemoglobin

Associated Disease/Clinical States:^{1,2}

Glucose-6-phosphate dehydrogenase (G6PD) Deficiency

Hemoglobinopathies (may result in the formation of unstable hemoglobins)

Thalassemia

Post-splenectomy

Oxidizing drugs

Note: Appear in conditions where unstable hemoglobin can form.²



References:

1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

2. Rodak BF, Carr JH. Inclusions in erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 107-14.

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3.4: Hemoglobin H (Hb H)

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Images show supravital stained peripheral blood smears with numerous hemoglobin H inclusions (examples indicated by arrows). Note the golf ball-like appearance of the red blood cells. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3GM8243H

Image 2: 100x oil immersion. https://doi.org/10.7939/R33F4M37Z

Appearance:

Hemoglobin H inclusions can only be visualized with supravital stains and not Wright or Romanowsky stains. With supravital stains, such as Brillian Cresyl Blue, the red blood cells are covered with numerous small, dark blue dots that give the cells a "golf ball" or "raspberry" appearance.^{1,2}

Inclusion composition:^{1,2}

Hemoglobin H is made up of 4 $\hat{\boldsymbol{v}}\hat{\boldsymbol{v}}$ globin chains in a tetramer formation. Hemoglobin H is unstable and will precipitate over time in the RBC, leading to Heinz Body formation. Precipitation can be induced using supravital stains, and can be used to visualize Hemoglobin H inclusions.

Associated Disease/Clinical States:1,2

Hemoglobin H Disease

References:

1. Randolph TR. Thalassemia. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 251-276.

2. Rodak BF, Carr JH. Inclusions in erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 107-14.

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3.5: Hemoglobin C Crystals

Appearance:^{1,2}

Dark red hexagonal crystals with blunt ends. The crystal is prominent within the red blood cell, or may be found extra-cellularly. Usually only one crystal is present per single cell. Hemoglobin C crystals are rarely found, as the spleen will remove them from circulation, though patients who have undergone a splenectomy have may numerous hemoglobin C crystals present on their peripheral blood smear.

Inclusion composition:²

Crystalized Hemoglobin C.

Associated Disease/Clinical States:¹

Homozygous Hemoglobin C Disease

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

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3.6: Hemoglobin SC Crystals

Appearance:

Crystals appear as a combination of sickle cells and hemoglobin C crystals. They are dark red inclusions with blunt ended projections.¹ The crystals are longer than Hemoglobin C crystals, but shorter and thicker than Hemoglobin S.

Inclusion composition:¹

Hemoglobin S and hemoglobin C

Associated Disease/Clinical States:¹

Hemoglobin SC Disease (compound heterozygosity)

References:

1. Rodak BF, Carr JH. Variations in shape and distribution of erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 93-106.

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3.7: Howell-Jolly Bodies



A peripheral blood smear with Howell-Jolly bodies. A. shows Howell-Jolly bodies. B. shows platelets on top of a red blood cell. Note the clear space surrounding the platelet. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R30R9MK3C



A peripheral blood smear with a Howell-Jolly body and a platelet on the same red blood cell. A. shows Howell-Jolly body. B. shows platelets on top of a red blood cell. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3B853Z9R



A peripheral blood smear with Howell-Jolly bodies (indicated with arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3028PV21

Appearance:

Under Wright/Romanowksy stains, Howell-Jolly Bodies appear as dark blue/purple round inclusions located at the periphery of the RBC. They usually present as a single inclusion inside the cell. Howell-Jolly Bodies are also visible under supravital stains.¹⁻⁴

Inclusion composition:^{2,3}

Nuclear fragments/remnants made up of DNA ¹⁻⁴

Associated Disease/Clinical States:

Thalassemia

Megaloblastic Anemia

Myelodysplastic Syndrome

Post-splenectomy



References:

1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

3. Fritsma GA. Bone marrow examination. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 253-68.

4. Ford J. Red blood cell morphology. Int J Lab Hematol [Internet]. 2013 Mar 9 [cited 2018 Jul 12];35(3):351–7. Available from: https://doi.org/10.1111/ijlh.12082

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3.8: Pappenheimer Bodies (Siderotic Granules)



An iron stained peripheral blood smear with pappenheimer bodies present (indicated with arrows). Perls Prussian Blue. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3FN1173T





An iron stained peripheral blood smear with pappenheimer bodies present (indicated with arrows). Perls Prussian Blue. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R36689100







A peripheral blood smear with pappenheimer bodies present (indicated with arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R35X25V0R







A peripheral blood smear with pappenheimer bodies present (indicated with arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3251G16Q

Appearance:

Inclusions are visible under both Wright/Romanowsky stains and Perls Prussian Blue stain. Pappenheimer inclusions appear as clusters of fine and irregular granules located at the periphery of the red blood cell.¹⁻³

Inclusion composition:³

Iron Associated Disease/Clinical States:^{1,2} Splenectomy Sideroblastic Anemia Thalassemia Sickle Cell Disease Hemachromatosis

References:

1. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.





3. Rodak BF, Carr JH. Inclusions in erythrocytes. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 107-14.

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3.9: Bacteria and Fungi

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Images show peripheral blood smears with bacteria present. Neutrophils show toxic changes (toxic vacuolation and granulation are most prominent) and contain ingested bacteria. Bacteria is also present extracellularly. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3NV99S2S

Image 2: 100x oil immersion. https://doi.org/10.7939/R3J38KZ9T

Image 3: 100x oil immersion. https://doi.org/10.7939/R38K75C0V

Appearance:

The morphology of a microorganism depends on the type of microorganism (fungi or bacteria) present in the blood. Can be seen extracellularly or intracellularly when patient is septic.¹

Organisms:¹

Fungi:	Bacteria:
Yeast	Clostridium perfringens
Histoplasma capsulatum	Bartonella bacilliformis
Cryptococcus neoformans	Cocci
	Other Bacilli/Rods

References:

1. Smith LA. Hemolytic anemia: nonimmune defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 372-87.

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3.10: Malaria



An image from a peripheral blood smear showing multiple malarial rings (Plasmodium falciparum) inside red blood cells. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3891263S





An image of a peripheral blood smear showing a crescent-shaped gametocyte (characteristic of Plasmodium falciparum). From MLS Collection, University of Alberta, https://doi.org/10.7939/R30R9MK4V







An image of a peripheral blood smear showing malaria at the gametocyte stage in the center. 60x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R37941903







An image of a peripheral blood smear showing a malarial parasite at the trophozoite stage in a red blood cell. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3WS8J22C

Appearance:

The morphology of malarial parasites seen in the red blood cell varies depending on the stage of maturation and species present. Malaria can appear as rings, trophozoites, schizonts, and gametocytes. Ring forms appear as a pale blue ring with a pink/purple chromatin dot, and more than one may be present in a single red blood cell. Malarial parasites are most often seen intracellular to the red blood cell with various forms.¹

Parasites can be visualized using the Giemsa stain during the screening of thin and thick smears.¹

Note 1: Banana shaped gametocytes seen are characteristically in *Plasmodium falciparum* infections.

Note 2: Malarial rings may be confused with platelets when the appear on top of a red blood cell. Platelets may be differentiated by a showing a slight clearing or halo around the platelet.²

Organisms:¹

The malarial parasite is spread to humans by the female Anopheles sp. mosquito.

Malaria parasites:^{1,2} Plasmodium falciparum Plasmodium vivax Plasmodium ovale Plasmodium malariae Plasmodium knowlesi





References:

1. Keohane EM. Extrinsic defects leading to increased erythrocyte destruction – nonimmune causes. In: Rodak's hematology clinical applications and principles. St. Louis, Missouri: Saunders; 2015. p. 394-410.

2. Rodak BF, Carr JH. Microorganisms. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 195-202.

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3.11: Babesia

Appearance:

Like malaria, Babesia species can be seen both intracellularly and extracellularly and visualed with the Giemsa stain. Babesia parasites appear as ring forms in the red blood cell with variable morphology.¹

Note: Tetrad of rings may appear as a "maltese cross".¹

Organisms:

The *Babesia microti* parasite is carried by the *Ixodes scapularis* tick.¹ Humans are incidental dead-end hosts of Babesia.

References:

1. Keohane EM. Extrinsic defects leading to increased erythrocyte destruction – nonimmune causes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 394-410.

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3.12: Trypanosomes

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Images of peripheral blood smears showing C shaped trypanosomes (center). From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3QZ22Z5K

Image 2: 100x oil immersion. https://doi.org/10.7939/R3M61C52N

Appearance:

Hemoflagellates that are often visualized using Giemsa stain during screening of thin and thick smears, though they are visible by regular Romanowsky staining procedures.¹

Trypomastigotes appear C or U shaped in the peripheral blood, usually seen extracellular to the red blood cells.²

Organisms:

The Reduviid bug ("kissing bug") carries *Trypanosoma cruzi* in it's gut where it matures. During a blood meal, the Reduviid bug releases *T.cruzi* via feces onto the feeding sites or mucous membranes where it can cause infection in the bloodstream.²

Trypanosoma cruzi is the causative agent of Chagas Disease.²

References:

1. Rodak BF, Carr JH. Microorganisms. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 195-202.

2. Ahmad N, Drew WL, Lagunoff M, Pottinger P, Reller LB, Sterling CR. Sarcomastigophora-the amebas. In: Ryan KJ, Ray CG, editors. Sherris medical microbiology. 6th ed. McGraw-Hill Education; 2014. p. 823-44.

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CHAPTER OVERVIEW

4: Red Blood Cells- Hypochromic, Microcytic Anemias

- 4.1: Iron Deficiency Anemia (IDA)
- 4.2: Anemia of Chronic Inflammation/Disease (ACI/ACD)
- 4.3: Sideroblastic Anemia
- 4.4: Thalassemia
- 4.5: Iron Studies

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4.1: Iron Deficiency Anemia (IDA)



An image from a peripheral blood smear demonstrating hypochromic and microcytic red blood cells. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3P84499C



An image from a peripheral blood smear showing hypochromic, microcytic red blood cells with occasional targets which can been in IDA. 60x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3988333G



An image from a peripheral blood smear showing hypochromic, microcytic red blood cells. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3KSGJK8F

Cause(s):1,2

Chronic Blood Loss (heavy menstruation, intermittent GI bleeding, etc.)

Increased Need (periods of rapid growth, pregnancy)

Inadequate intake (diet)

Impaired absorption (malabsorption)

Notes: When there is not enough iron to meet the requirements of the body, iron stores begin to deplete, and IDA occurs. Development of IDA occurs over a period of time.¹⁻⁴

IDA development is said to occur in three stages:³

Stage 1	Stage 2	Stage 3
(Storage Iron Depletion):	(Transport Iron Depletion):	(Functional Iron Depletion- IDA):
- Normal RBC maturation	- Decreased serum iron and ferritin	 Anemia is evident
- Decline in storage iron (decreased ferritin, decreased iron stores	- Transferrin and TIBC levels increase	- PBS shows microcytic, hypochromic RBCs
in the bone marrow)	- Absent iron stores in the bone marrow	- RBC development is affected
- No other evidence of anemia development.	 Evidence of anemia is not as apparent. 	- Same iron study results as stage 2
		- Hb is decreased
		 Hepcidin is decreased
		 Erythropoietin is increased

Laboratory Features of Iron Deficiency Anemia (At Stage 3):¹⁻⁴

CBC: RBC Count: Decreased PLT: Variable (increased in chronic bleeding) Hb: Decreased Hct: Decreased MCV, MCH, MCHC: Decreased RDW: Increased	PBS: Microcytic, hypochromic RBCs Target cells Elliptocytes Teardrop cells Normal WBC morphology	BM: M:E Ratio: Decreased Erythroid hyperplasia Iron Stores: Absent or severely decreased (not sustainable)
Iron Studies: Serum Iron: Decreased Ferritin: Decreased Transferrin: Increased Transferrin Saturation: Decreased TIBC: Increased	Other Tests: Prussian Blue stain of the BM shows absent or decreased iron Reticulocyte count decreased	

References:

1. McKenzie SB. Anemias of disordered iron metabolism and heme synthesis. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 198-230.

2. Miller JL. Iron deficiency anemia: A common and curable disease. Cold Spring Harb Perspect Med [Internet]. 2013 Jul 1 [cited 2018 Jun 28];3(7):10.1101/cshperspect.a011866 a011866. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3685880/

3. Doig K. Disorders of iron kinetics and heme metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 297-313.





4. Finnegan K. Iron metabolism and hypochromic anemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 117-37.

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4.2: Anemia of Chronic Inflammation/Disease (ACI/ACD)

Cause(s):

Anemia that occurs in patients with conditions that result in chronic inflammatory states such as rheumatoid arthritis, infections, and malignancies. Anemia is reversed when underlying condition is treated and inflammation subsides.¹ Chronic inflammation leads to an increase in inflammatory cytokines and actue phase reactants that alter iron metabolism and decrease RBC production and lifespan.

Additional notes:

Acute phase reactants are serum proteins whose levels are increased by the liver in response to inflammation. Three related acute phase reactants to ACI are:²

- 1. Hepcidin
- 2. Lactoferrin
- 3. Ferritin

Mechanisms of anemia development in ACI:^{1,3}

1. Decrease in iron available for erythropoiesis (increased hepcidin causes inhibition of iron release from macrophages and decreased iron absorption from the diet, iron bound to lactoferrin and ferritin is not available to developing RBCs)

2. EPO production in the kidneys is inhibited by cytokines

3. Erythropoiesis is inhibited by cytokines (diminishes developing erythroid precursor's response to EPO)

4. Decreased RBC survival (increased activation of macrophages by inflammatory cytokines leads to increased removal of RBCs from circulation and decreased survival)

Laboratory Features: 1,2,4

CBC: RBC: Decreased WBC: Normal or Increased depending on the cause of the inflammation PLT: Normal Hb: Decreased Hct: Normal to Decreased MCV, MCH, MCHC: Normal to Decreased RDW: Increased	PBS: Normochromic, Normocytic OR Microcytic, hypochromic RBCs	BM: M:E Ratio: Increased (Decreased production of erythroids, +/- increased production of myeloids) Iron Stores: Increased
Iron Studies: Serum Iron: Decreased Ferritin: Increased Transferrin: Decreased Transferrin saturation: Normal to Decreased TIBC: Decreased	Other Tests: N/A	

References:

1. McKenzie SB. Anemias of disordered iron metabolism and heme synthesis. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 198-230.

2. Doig K. Disorders of iron kinetics and heme metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 297-313.

3. Turgeon ML. Hypochromic anemias and disorders of iron metabolism. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 131-144.

4. Finnegan K. Iron metabolism and hypochromic anemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 117-37.

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4.3: Sideroblastic Anemia



A peripheral blood smear picture showing a dimorphic population of red blood cells: hypochromic, microcytic and normochromic, normocytic red cells. Dimorphissm is commonly seen in Sideroblastic Anemia cases. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3P844B3X



A peripheral blood smear picture showing a dimorphic RBC population and tear cells in sideroblastic anemia. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3CF9JN85

Cause(s): Development of sideroblastic anemia can be due to hereditary or acquired causes that lead to abnormal heme synthesis.^{1,2}

Hereditary: Sex-linked or autosomal recessive mutations

Acquired: Idiopathic, MDS and other malignancies, drugs, lead toxicity

Laboratory Features of Sideroblastic Anemia:¹⁻³

CBC:	PBS:	BM:
RBC: Decreased	Dimorphic population (Normochromic/Normocytic alongside	M:E Ratio: Decreased
WBC: Variable	Hypochromic/Microcytic)	Erythroid hyperplasia (ineffective erythropoiesis)
PLT: Variable	Tears	Ringed sideroblasts
Hb: Decreased	Schistocytes	Macrophages have increased iron (Increased iron stores)
MCV, MCH, MCHC: Normal to Decreased (as they are averages of	Pappenheimer bodies	
the RBC appearance)	Basophilic stippling	
RDW: Increased		
RETIC: Decreased		







Iron Studies: Serum Iron: Increased Ferritin: Increased Transferrin: Normal to Decreased TIBC: Normal to Decreased Other Tests: Bilirubin: Increased Haptoglobin: Decreased LD: Increased Prussian blue stain of BM shows increased iron levels and ringed sideroblasts

References:

1. McKenzie SB. Anemias of disordered iron metabolism and heme synthesis. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 198-230.

2. Doig K. Disorders of iron kinetics and heme metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 297-313.

3. Finnegan K. Iron metabolism and hypochromic anemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 117-37.

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4.4: Thalassemia

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Images show thalassemia peripheral blood smears with hypochromic, microcytic red blood cells and poikilocytosis. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3DR2PQ4J Image 2: 50x oil immersion. https://doi.org/10.7939/R3V698T05 Image 3: 50x oil immersion. https://doi.org/10.7939/R3HD7P773

Thalassemias are classified as a group of genetic hemoglobin disorders where the production of α and β globin chains is affected. This is considered to be a quantitative hemoglobin disorder and is categorized by the affected globin chain (alpha or beta), and as major or minor depending on the severity of the disease.^{1,2}

Alpha-Thalassemia:

Cause(s):

 α globin chain genes are located on chromosome 16 and there are normally four genes in total ($\alpha\alpha/\alpha\alpha$), two inherited from each parent. α -thalassemia results when there is a deletion in any number of the α globin gene. The severity of anemia and amount of α globin chain production is dependent the number of genes that are deleted.³

α-Thalassemia Silent Carrier (αα/α-):^{1,2}

Occurs when one α gene is deleted. There is still adequate production of α to ensure normal hemoglobin synthesis. Patient is asymptomatic and the mutation is benign.

In newborns, there is an excess production of γ globin chains. These γ globin chains tend to also form tetramers and result in Hemoglobin Barts (Hb Barts). Hb Barts has a high oxygen affinity and is inefficient for oxygen delivery to the tissues of the developing fetus. In the silent carrier state, there is only a small amount of Hb Barts produced.

α-Thalassemia Minor (αα/–) or (α-/α-): 1,2

Occurs when two α genes are deleted. There is now a 50% reduction in normal α globin chain production.

In adults, increased production of red blood cells is able to compensate for the decrease in α chain production, and α and β globin chain production is balanced. Patients are asymptomatic and any anemia present is mild.

There is between 5-15% hemoglobin Barts present at birth, but this decreases once β globin chain production takes over and γ globin chain production decreases. In adults, globin chain production is balances, so no Hemoglobin H is formed.

Hemoglobin H Disease (α-/–):^{1,2}

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4.5: Iron Studies

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Images of iron stained bone marrow particle smears showing various amounts of iron stores (indicated by the amount of blue present). Perls Prussian Blue. From MLS Collection, University of Alberta.

Image 1: 10x magnification. Low iron stores. https://doi.org/10.7939/R3NC5ST92

Image 2: 10x magnification. Normal iron stores. https://doi.org/10.7939/R3XP6VK0C

Image 3: 10x magnification. High iron stores. https://doi.org/10.7939/R3ZP3WG13

Table 1. Iron studies of hypochromic and microcytic anemias.¹⁻³

Апетіа	Serum Iron	Ferritin	Transferrin	Transferrin Saturation	TIBC	BM Iron stores
IDA	D	D	Ι	D	Ι	Absent/D
Thalassemia Minor	N/I	N/I	N/I	N/I	Ν	N/I
Anemia of Chronic Inflammation	D	Ι	Ν	N/D	D	Ι
Sideroblastic Anemia	I	Ι	N/D	I	N/D	Ι
N = Normal I = Increased D = Decreased						

References:

1. McKenzie SB. Anemias of disordered iron metabolism and heme synthesis. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 198-230.

2. Doig K. Disorders of iron kinetics and heme metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 297-313.

3. Finnegan K. Iron metabolism and hypochromic anemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 117-37.

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CHAPTER OVERVIEW

5: Red Blood Cells- DNA Metabolism Abnormalities and Bone Marrow Failure

- 5.1: Megaloblastic Anemia
- 5.2: Non-Megaloblastic Macrocytic Anemia
- 5.3: Aplastic Anemia

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5.1: Megaloblastic Anemia



An image of a megaloblastic bone marrow showing nuclear-cytoplasmic asynchrony in a polychromatic normoblast. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3Q81576M



An image of a megaloblastic bone marrow showing nuclear-cytoplasmic asynchrony in erythroid precursors. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3KK94T57



An image of a megaloblastic bone marrow demonstrating erythroid hyperplasia and nuclear-cytoplasmic asynchrony. 40x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3C24R36T



An image of a megaloblastic bone marrow demonstrating a giant metamyelocyte. From MLS Collection, University of Alberta, https://doi.org/10.7939/R33J39G7T



An image of a megaloblastic bone marrow showing a hypersegmented neutorphil and a giant band. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3794191K



An image of a megaloblastic bone marrow demonstrating a hypersegmented neutrophil. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3V11W200







An image of a megaloblastic peripheral blood smear showing a hypersegmented neutrophil and oval macrocytes. From MLS Collection, University of Alberta, https://doi.org/10.7939/R36970D1B



An image of a megaloblastic peripheral blood smear showing a hypersegmented neutrophil and oval macrocytes. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3B27Q653



A megaloblastic peripheral blood smear demonstrating oval macrocytes. Small lymphocytes are present for size comparison. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3XS5JZ3P

Cause(s):

Megaloblastic anemia occurs when there are defects in DNA synthesis that cause problems with blood cell production and maturation (all cells are affected, not just red blood cells). Megaloblastic anemia is most commonly caused by deficiencies in Vitamin B_{12} (cobalamin) and folate (folic acid). Both Vitamin B_{12} and folate are important factors used in the process of DNA synthesis.¹

Cellular characteristics of Megaloblastic Anemia:

1. A characteristic finding in bone marrow smears for megaloblastic anemia would the appearance of nuclear-cytoplasmic (N:C) asynchrony in all cell lines.¹ N:C asynchrony describes the inability of the cell's chromatin to mature normally giving the nucleus a more immature, more fine, looser, and larger appearance than expected compared to that of the cytoplasm. Cytoplasm maturation is not affected and matures normally. Due to these characteristics, the cells are described as megaloblastic.^{1,2}

2. Another characteristic finding on the peripheral blood smear would be the appearance of hypersegmented neutrophils. Hypersegmentation is described when either observation is present:^{1,2}

- 5% or more neutrophils have 5 lobes
- One neutrophil with ≥ 6 lobes
- 3. Oval macrocytes are also indicative of megaloblastic anemia.²

Function of Folate and Vitamin $B_{12}\ in$ DNA synthesis:

Folate is ingested as folic acid which is inactive. The process of converting folic acid to its active form (Tetrahydrofolate, THF) requires the help of Vitamin B_{12} .¹ Vitamin B_{12} is used as a cofactor in a reaction that converts inactive folate (N5-methylTHF) into the active Tetrahydrofolate (THF) form which is then used to continue DNA synthesis.^{1,3} Without Vitamin B_{12} or folate, the nucleotide thymidine cannot be produced and DNA synthesis is impaired.²

Vitamin B12 (Cobalamin)

Absorption:

Available in eggs, milk, and meat. The low pH in the stomach causes Vitamin B_{12} to be released from ingested proteins. Vitamin B_{12} then binds to haptocorrin to be transported into the duodenum. In the duodenum, proteases release the Vitamin B_{12} and then it is picked up by intrinsic factor where it transports it to enterocytes of the ileum to be absorbed.¹

Transport in circulation:

Once absorbed by the gastrointestinal tract, the transport protein called transcobalamin binds the Vitamin B₁₂ to be transported to the rest of the body in circulation.^{1,2}

Vitamin B₁₂ deficiency:

Can occur to due to a variety of causes such as: malabsorption, bacterial and parasitic infection, inadequate intake in the diet, or impaired utilization by the body.^{1,2}

Vitamin B₁₂ deficiency can develop secondary to the absence of intrinsic factor (IF) which is used to help absorb Vitamin B₁₂ into the body. IF deficiency can be caused by autoantibodies against IF and gastric cells resulting in a type of anemia called Pernicious anemia.^{1,3}

Folate (Folic acid)

Absorption:



Folate can be found in yeast, milk, eggs, mushrooms, and leafy greens and is easily destroyed by heat. Folate is absorbed throughout the gastrointestinal tract as folic acid and converted into N5-methylTHF in the cells.¹

Folate deficiency:

Causes of folate deficiency can be due to inadequate intake in the diet, malabsorption, drugs that interfere with use, and an increased need (such as during pregnancy or rapid growth).^{1,3}

Laboratory Features of Megaloblastic Anemia: ^{1,3}				
CBC: RBC, WBC, PLT, Hb, Hct: Decreased *MCV: Usually > 110 fL MCH: Increased MCHC: Normal RETIC: Normal to decreased	PBS: *Ovalmacrocytes Howell-Jolly Bodies *Hypersegmented neutrophils Schistocytes Teardrop Cells	BM: M:E Ratio: decreased (Ineffective erythropoiesis) Hypercellular *N:C asynchrony Enlarged precursors Giant metamyelocytes and bands		
Other Tests: Folate deficiency – Serum Folate: Decreased Vitamin B ₁₂ deficiency – Serum Vitamin B ₁₂ : Decreased – IF blocking antibodies – Antibody assays (Pernicious anemia)				
* Indicates the characteristic morphological findings in m	negaloblastic anemia blood smears			

References:

1. Hubbard J, Robinson S. Megaloblastic and nonmegaloblastic macrocytic anemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 277-301.

2. Goossen LH. Anemias caused by defects of DNA metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 314-30.

3. Nagao T, Hirokawa M. Diagnosis and treatment of macrocytic anemias in adults. J Gen Fam Med [Internet]. 2017 Oct 13 [cited 2018 Jun 25];18(5):200–4. Available from: http://doi.wiley.com/10.1002/jgf2.31

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5.2: Non-Megaloblastic Macrocytic Anemia



An image from a peripheral blood smear demonstrating round macrocytes and poikilocytosis in liver disease. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3MP4W32X

Cause(s):

A group of anemias that present with macrocytes without megaloblastic features. Most often, non-megaloblastic macrocytic anemias are caused by: alcoholism, liver disease, bone marrow failure, and myelodysplastic syndromes (MDS).^{1,2}

Chronic and heavy consumption of alcohol can lead to macrocytosis due to a variety of effects it has in erythrocyte development. Alcohol can not only interfere with folate metabolism but also is also directly toxic on bone marrow precursors. ^{1,3}

Liver disease is commonly associated with alcoholism and it is thought that macrocytosis is caused by an increase in cholesterol and lipids in the red blood cell membrane.³

Note: Additional information about bone marrow failure and MDS will be discussed in later chapters.

Laboratory Features: ¹⁻⁴

CBC: PLT: Decreased *MCV: usually 100-110 fL (MCV is rarely >110 fL) *RETIC: Increased (if hemolytic anemia is present)	PBS: *Round Macrocytes Target cells *NO hypersegmented neutrophils	BM: Nomorcellular or hypercellular Erythroid hyperplasia *Megaloblastic features are absent in precursors
Iron Studies (Liver Disease): Serum Iron: Decreased to Normal Ferritin: Increased Transferrin: Normal Transferrin Saturation: Normal to Increased	Other Tests: Liver enzyme tests	
*Features that differentiate megaloblastic from non-megaloblastic anemias		





References:

1. Hubbard J, Robinson S. Megaloblastic and nonmegaloblastic macrocytic anemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 277-301.

2. Nagao T, Hirokawa M. Diagnosis and treatment of macrocytic anemias in adults. J Gen Fam Med [Internet]. 2017 Oct 13 [cited 2018 Jun 25];18(5):200-4. Available from: http://doi.wiley.com/10.1002/jgf2.31

3. Taghizadeh M. Megaloblastic anemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 138-55.

4. Goossen LH. Anemias caused by defects of DNA metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 314-30.

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5.3: Aplastic Anemia

Bone Marrow Failure:

Bone marrow failure is characterized by reduced hematopoiesis in the bone marrow resulting in cytopenias in one or more cell lines. Decreased hematopoiesis can be attributed to:¹

- 1. Destruction of hematopoietic stem cells due to acquired causes
- 2. Destruction of hematopoietic stems cells due to inherited causes
- 3. Ineffective erythropoiesis
- 4. Disruption of bone marrow microenvironment
- 5. Reduced production of growth factors and hormones related to hematopoiesis
- 6. Infiltration of the bone marrow

Aplastic anemia is a bone marrow failure syndrome that is characterized by a decreased cell count in all cell lines (pancytopenia) and a hypocellular (aplastic) bone marrow. (McKenzie ch 16 pg 303, Rodak ch 22 pg 332) Unlike other anemias, hepatosplenomegaly and lymphadenopathy are absent. (McKenzie ch 16 pg 307)

Cause(s):

There is no known, single cause of aplastic anemia but it's development can be associated with a variety of clinical states and agents which can be either acquired or inherited.^{1,2} It is thought that acquired causes of aplastic anemia can lead to an immunologic response against one's own hematopoietic stem cells.¹

Table 1. List of acquired and inherited causes of aplastic anemia.¹⁻³

Acquired:	Inherited:
Idiopathic	Fanconi Anemia (autosomal recessive, rare X-linked recessive)
Drugs and Chemicals	Dyskeratosis congenita
Radiation	Shwachman-Diamond Syndrome
Infectious agents	
Clonal Disorders (e.g. MDS, PNH)	

Laboratory Features of Aplastic Anemia:1,3

CBC: RBC: Decreased WBC: Decreased PLT: Decreased Hb: Decreased RETIC: Decreased MCV: Normal to increased	PBS: Pancytopenia (Thrombocytopenia, neutropenia, anemia) Normochromic Normocytic or macrocytic	BM: Hypocellular or dry tap Fatty infiltration
Iron Studies: Serum Iron: Increased Ferritin: Increased Transferrin: Normal Transferrin Saturation: Normal to increased	Other Tests: Molecular testing Flow cytometry	

References:

1. Lo C, Glader B, Sakamoto KM. Bone marrow failure. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 331-47.

2. Laudicina R. Hypoproliferative anemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 302-16.

3. Perkins SL. Aplastic anemia including pure red cell aplasia, congenital dyserythropoietic anemia, and paroxysmal nocturnal hemoglobinuria. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 156-75.

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CHAPTER OVERVIEW

6: Red Blood Cells- Introduction to Hemolytic Anemias

6.1: Introduction to Hemolytic Anemias

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6.1: Introduction to Hemolytic Anemias

Hemolytic anemia refers to a process where there is increased red blood cell destruction or decreased red blood cell survival (hemolysis) leading to a drop in the measured hemoglobin (anemia).¹

The type of hemolysis can be categorized into different categories based on the location of the hemolysis (intravascular or extravascular) or the cause (intrinsic or extrinsic).

Intravascular and extravascular refers to the location of the hemolytic process, whether the process is taking place within the blood vessels (intra) or outside the blood vessels (extra).²

Intrinsic and extrinsic refers to the cause of red blood cell destruction relative to the red blood cell itself. If the cause is due to an issue with the red blood cell (e.g. inherited defects of the RBC), it is referred to as being intrinsic. If the cause is due to factors from outside the red blood cell (e.g. environment), it is referred to as being extrinsic.²

Compensated hemolysis refers to the ability of the bone marrow to increase red blood cell production in order to compensate for the rate of hemolysis. As a result, anemia does not develop.²

Extravascular (Macrophage-mediated) Hemolysis (EVH)

Process:1, 3-5

- 1. RBCs are phagocytized in the spleen, bone marrow, or liver by macrophages.
- 2. Hemoglobin is broken down into iron, globin, and the protoporphyrin ring.
- 3. Iron is carried by transferrin to the bone marrow be reused or stored as ferritin or hemosiderin.
- 4. Globin is broken down into amino acids to be recycled.
- 5. The protoporphyrin ring is further is broken down to biliverdin and then to unconjugated bilirubin in the macrophage.
- 6. Unconjugated bilirubin is then released and carried by albumin to the liver.
- 7. In the liver, unconjugated bilirubin is converted to conjugated bilirubin (bilirubin diglucuronide).
- 8. Conjugated bilirubin is excreted with bile into the intestines where it is converted into urobilinogen by bacteria.

9. A majority of the urobilinogen is then excreted in feces, a small amount is reabsorbed by the kidney, and another portion is excreted into the urine.

Intravascular (Fragmentation) Hemolysis (IVH)

Process:³⁻⁵

- 1. RBC hemolysis occurs in the blood vessels and hemoglobin is released into circulation.
- 2. Hemoglobin dissociates into $\alpha\beta$ dimers and is picked up by Haptoglobin where it is carried to the liver.
- 3. Subsequent catabolic steps are the same as extravascular hemolysis from the liver onwards.
- 4. If haptoglobin is not available, the $\alpha\beta$ dimers become oxidized into methemoglobin where it is broken down into metheme and globin.
- 5. Metheme is carried by hemopexin to the liver.
- 6. If hemopexin is not available, metheme binds albumin instead to form methalbumin,.
- 7. Methemalbumin continues to circulate the body until hemopexin becomes available.

Table 1. Comparison of IVH and EVH Laboratory Findings¹⁻³

Test	IVH	EVH
RBC, Hct, Hb	Decreased	Decreased
Total Bilirubin	Increased	Increased
LDH	Increased	Slightly Increased
Haptoglobin	Decreased	Slightly Decreased
Hemopexin	Decreased	Slightly Decreased
Hemosiderinuria	Present	Absent



Hemoglobinuria	Present	Absent
RBC Morphology	Schistocytes	Spherocytes
Examples	PNH, PCH, MAHAs, Mechanical trauma, Bacterial Infections, Thermal Injury	Thalassemia (Other hemoglobinopathies), Enzymopathies, Membranopathies, Megaloblastic anemia, Autoimmune hemolytic anemia, Drug-induced hemolytic anemia

References:

1. Barcellini W, Fattizzo B. Clinical Applications of Hemolytic Markers in the Differential Diagnosis and Management of Hemolytic Anemia. Dis Markers [Internet]. 2015 Dec 27 [cited 2018 Jun 26];2015:635670. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4706896/

2. McKenzie SB, Otto CN. Introduction to anemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.178–97.

3. Doig K. Introduction to increased destruction of erythrocytes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 348-66.

4. McKenzie SB. Hemoglobin. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 77-96.

5. Harmening DM. The red blood cell: structure and function. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 64-81.

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CHAPTER OVERVIEW

7: Red Blood Cells- Hemoglobinopathies

- 7.1: Normal Hemoglobin Structure
- 7.2: Sickle Cell (Hemoglobin SS) Disease
- 7.3: Sickle Cell Trait (Hemoglobin AS)
- 7.4: Hemoglobin C (Hb CC) Disease
- 7.5: Hemoglobin SC Disease

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7.1: Normal Hemoglobin Structure

Location of globin genes:¹

Chromosome 16: α globin genes

Chromosome 11: β globin genes

One α globin gene and one β globin gene are inherited from each parent.

Normal hemoglobin A is made up of two α globin chains, two β globin chains, and four heme molecules.

Heme is formed from protoporphyrin ring precurosors and ferrous iron (Fe²⁺).

Table #1 Normal hemoglobin content at various stages of life.¹

Embryonic:	Fetal:	Adult:
Hb Gower I ($\zeta_2 \varepsilon_2$)	HbF ($\alpha_2\gamma_2$)	HbA $(\alpha_2\beta_2) > 95\%$
Hb Gower II ($\alpha_2 \epsilon_2$)		HbA ₂ ($\alpha_2 \delta_2$) <3.5%
Hb Portland ($\zeta_2 \gamma_2$)		HbF $(\alpha_2\gamma_2)$ 1-2%

References:

1. Keohane EM. Hemoglobin metabolism. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 124-36.

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7.2: Sickle Cell (Hemoglobin SS) Disease



An image of a peripheral blood smear of a patient with sickle cell disease. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3G737K46



©••\$



An image of a peripheral blood smear containing sickle cells, target cells, and increased polychromasia. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3XP6VJ98

Cause(s):

 β globin chain amino acid substitution in the 6th position from glutamic acid (Glu) to valine (Val). In the homozygous form of the disease, both β globin genes are affected.¹

Inheritance:

Autosomal dominant²

Demographics:³

Tropical Africa

Mediterranean Areas

Sickle cell disease is common in areas where malaria is prominent and it is suggested that the disease acts as a protective factor for malaria. This protection is only seen in heterozygotes, as homozygotes often lose splenic function, which is essential for combating the parasite.

Cellular Features:¹⁻⁴

See sickle cell (drepanocytes) under RBC morphology for more information about cell formation.

The formation of sickle cells becomes irreversible over time leading to the formation of rigid and "sticky" sickle cell aggregates resulting in many complications.

Complications:¹⁻⁴

Chronic hemolytic anemia

Vaso-occlusion (can lead to ischemic tissue injury, splenic sequestration of RBCs, autosplenectomy)

Prone to infections

Nephropathies

Stroke

Laboratory Features of Sickle Cell Disease:²⁻⁴

CBC: RBC: Decreased WBC: Increased PLT: Increased Hb: Decreased RETIC: Increased RDW: Increased	PBS: Sickle cells Normochromic, normocytic RBCs Target cells Polychromasia nRBCs Howell-Jolly bodies Pappenheimer bodies Basophilic Stippling	BM: Erythroid Hyperplasia Iron stores: often increased
Hemoglobin Electrophoresis: Hb S: 80-95% Hb A: None Hb A ₂ : 2-% Hb F: 5-20%	Other Tests: Solubility Screen: Positive Metasulfite Sickling Test: Positive HPLC Hemoglobin Electrophoresis	

References:



1. Chonat S, Quinn CT. Current standards of care and long term outcomes for thalassemia and sickle cell disease. Adv Exp Med Biol [Internet]. 2017 [cited 2018 Jun 5];1013:59–87. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC5720159/

1. Randolph TR. Hemoglobinopathies (structural defects in hemoglobin). In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 426-453.

3. Laudicina RJ. Hemoglobinopathies: qualitative defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.231-50.

4. Harmening DM, Yang D, Zeringer H. Hemolytic anemias: extracorpuscular defects. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 250-79).

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7.3: Sickle Cell Trait (Hemoglobin AS)

Cause(s):

 β globin chain amino acid substitution in the 6th position from glutamic acid (Glu) to valine (Val). Only one β globin genes is affected.^{1,2}

Inheritance:

Heterozygous state where one normal β globin gene and one affected β globin gene are inherited.³

Clinical Findings:¹⁻³

Due to the presence of Hb A and reduced concentration of HB S, polymerization of Hb S and sickling of red blood cells does not normally occur. As a result, condition is mostly benign and asymptomatic.

Sickling can still occur under extremely low hypoxic conditions.

Laboratory Features:¹⁻³

CBC: All parameters (Even Hb) are normal	PBS: Absence of sickle cells PBS appears normal (may see a slight increase in target cells)	BM: N/A
Hemoglobin Electrophoresis: Hb S: 35-45% Hb A: 50-65% Hb A ₂ : Normal Hb F: Normal Hb A:Hb S is ~60:40	Other Tests: Solubility Screen: Positive Metasulfite Sickling Test: Positive HPLC Hemoglobin Electrophoresis	

References:

1. Laudicina RJ. Hemoglobinopathies: qualitative defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.231-50.

2. Harmening DM, Yang D, Zeringer H. Hemolytic anemias: extracorpuscular defects. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 250-79).

3. Randolph TR. Hemoglobinopathies (structural defects in hemoglobin). In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 426-453.

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7.4: Hemoglobin C (Hb CC) Disease

Cause(s):

 β globin chain amino acid substitution in the 6th position from glutamic acid (Glu) to lysine (Lys).^{1,2}

Inheritance:

Autosomal dominant¹

Demographics:

West Africa¹

Clinical Features:¹⁻³

See Hemoglobin C under RBC inclusions for additional information.

Less splenic sequestration and milder chronic hemolysis compared to sickle cell disease. Patients are usually asymptomatic.

Laboratory Features:^{1,2}

CBC: Hb: Decreased Hct: Decreased	PBS: Normochromic, normocytic RBCs *Hb C crystals Target cells nRBCs	BM: N/A
Hemoglobin Electrophoresis: Hb S: >90 Hb A: None Hb A ₂ : Normal Hb F: <7% *Not always seen, more likely in patients who have had a splenectomy.	Other Tests: Solubility Screen: Negative Metasulfite Sickling Test: Negative HPLC Hemoglobin Electrophoresis	

References:

1. Laudicina RJ. Hemoglobinopathies: qualitative defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.231–50.

2. Randolph TR. Hemoglobinopathies (structural defects in hemoglobin). In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 426-453.

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7.5: Hemoglobin SC Disease

Cause(s):

Both β globin chains are affected as both genes for hemoglobin S and hemoglobin C are both inherited. 1,2

Clinical Features:¹

Complication is less severe than sickle cell disease but more severe than hemoglobin C disease. Cells are still prone to sickling under decreased oxygen tension.

Complications are similar to those seen in sickle cell anemia and vaso-occlusion can occur.

Laboratory Features:^{1,2}

CBC: Hb: Decreased Hct: Decreased MCHC: Increased	PBS: Normochromic Normocytic Target Cells HbSC crystals	BM: N/A
Hemoglobin Electrophoresis: Hb S: 45% Hb C: 45% Hb A: None Hb A ₂ : 2-4% Hb F: 1%	Other Tests: Solubility Tests: Positive HPLC Hemoglobin Electrophoresis	

References:

1. Laudicina RJ. Hemoglobinopathies: qualitative defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.231–50.

2. Randolph TR. Hemoglobinopathies (structural defects in hemoglobin). In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 426-453.

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CHAPTER OVERVIEW

8: Red Blood Cells- Extrinsic Defects Causing Hemolytic Anemias

- 8.1: Microangiopathic Hemolytic Anemias (MAHAs)
- 8.2: Macroangiopathic Hemolytic Anemias
- 8.3: Immune-Mediated Hemolytic Anemias
- **8.4: Infectious Agents**

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8.1: Microangiopathic Hemolytic Anemias (MAHAs)

An interactive or media element has been excluded from this version of the text. You can view it online here: https://pressbooks.library.ualberta.ca/mlsci/?p=464

Images of peripheral blood smears demonstrating features of microangiopathic hemolytic anemia. Note the presence of schistocytes, increased polychromasia, and lack of platelets. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3FX74D2V

Image 2: 100 oil immersion. https://doi.org/10.7939/R3B56DK8D

Image 3: 50x oil immersion. https://doi.org/10.7939/R3639KM46

Introduction:^{1,2}

Microangiopathic hemolytic anemias are a group of disorders that involve the fragmentation of red blood cells in the circulation due to the formation of microthrombi in the microvasculature. This results in intravascular hemolysis and thrombocytopenia.

Red blood cells are physically damaged as they pass through blood vessels resulting in the formation of schistocytes (intravascular hemolysis). The damaged red blood cells are then often removed from circulation by the spleen resulting in extravascular hemolysis.

These features can be found in certain clinical states:

1. Disseminated Intravascular Coagulation (DIC)

- 2. Thrombotic thrombocytopenic purpura (TTP)
- 3. Hemolytic-uremic syndrome (HUS)

4. HELLP Syndrome (Hemolysis, Elevated liver enzymes and Low platelets)

General Laboratory Findings of MAHAs:³

CBC:	PBS:	Other Tests:
PLT: Decreased	Schistocytes	Unconjugated Bilirubin: Increased
Hb: Decreased	Polychromasia	LDH: Increased
RETIC: Increased	nRBCs	Haptoglobin: Decreased
		Urine urobilinogen: Increased
		Variable hemoglobinuria and hemoglobinemia

References:

1. Smith LA. Hemolytic anemia: nonimmune defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.372-87.

2. Harmening DM, Yang D, Zeringer H. Hemolytic anemias: extracorpuscular defects. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 250-79).

2. Keohane EM. Extrinsic defects leading to increased erythrocyte destruction – nonimmune causes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 394-410.

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8.2: Macroangiopathic Hemolytic Anemias

Traumatic Cardiac Hemolytic Anemia

In this condition, hemolysis is due to mechanical trauma caused by prosthetic cardiac valves. High blood flow around the prosthetic causes red blood cells to fragment leading to intravascular hemolysis. Any damaged cells that do not hemolyze in circulation are removed by the spleen via extravascular hemolysis.^{1,2}

Hemolytic anemia due to traumatic cardiac causes is uncommon and platelet count is not usually decreased drastically. Any hemolysis that occurs is often compensated by the bone marrow,^{1,2}

Laboratory Findings for Traumatic Cardiac Hemolytic Anemia:²

CBC:	PBS:	Other Tests:
PLT: Normal	Schistocytes	Unconjugated Bilirubin: Increased
Hb: Decreased		LD: Increased
RETIC: Increased		Haptoglobin: Decreased

Exercise-induced Hemoglobinuria

Transient hemolysis that occurs due to stress caused by exercise. Most often due to activities involving contact with hard surfaces such as running. Red blood cells become damaged as they pass through small vessels. Anemia usually does not develop unless hemolysis is severe.¹

Laboratory Findings for Exercise-induced Hemoglobinuria:²

CBC:	PBS:	Other Tests:
Hb: Increased	Schistocytes are NOT present	Unconjugated Bilirubin: Increased
RETIC: Increased		LDH: Increased
MCV: Slight increase		Haptoglobin: Decreased
-		Hemoglobinuria

Thermal Injury

Hemolytic anemia can develop after thermal burns to the body. Degree of hemolysis is dependent on the amount of surface area affected. Hemolysis is due to direct thermal damage to the red blood cells.¹

Laboratory Findings for Thermal Injury:¹

CBC:	PBS:	Other Tests:
Hb: Decreased	Schistocytes Micro-Spherocytes	Hemoglobinuria

References:

1. Smith LA. Hemolytic anemia: nonimmune defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p.372–87.

2. Keohane EM. Extrinsic defects leading to increased erythrocyte destruction – nonimmune causes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 394-410.

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8.3: Immune-Mediated Hemolytic Anemias

Hemolytic anemias can be caused by antibodies that can be directed against self (auto-antibodies) or foreign (allo-antibodies) antigens. Antibodies implicated vary in immunoglobulin class and optimal temperature of reactivity.

Table 1. Comparison of Warm and Cold Reacting Antibodies.¹

	Warm	Cold
Immunoglobulin Class	IgG	IgM (Exception: autoanti- P is IgG)
Optimal Temperature	37°℃	<30°C (often at 4°C) Pathological cold agglutinins react closer to body temperature
Mechanism of Hemolysis	Extravascular IgG or C3b attachment to macrophages	Extravascular or Intravascular Complement-mediated
Specificity	anti-Rh (Broad specificity)	Autoanti-I Autoanti-i Autoanti-P

Warm Auto-Immune Hemolytic Anemia (WAIHA)

Associated Conditions:^{2,3}

Lymphoproliferative diseases (e.g. Chronic lymphocytic leukemia)

Other autoimmune diseases: systemic lupus erythematosus, rheumatoid arthritis

Ingestion of certain drugs

Some non lymphoid neoplasms

Some inflammatory diseases

Affected age: Usually old age (>40 years old).¹

Antibody Specificity: Panreactive, polyclonal anti-Rh (IgG).³

Pathophysiology:

Often due to a Pan-reacting antibody against the Rh blood group system causing extravascular hemolysis. The antibodies bind to the red blood cells, resulting in their removal by macrophages in the spleen. Incomplete phagocytosis results in the removal of only some of the red blood cell membrane allowing the rest to reform. This reformation changes the red blood cell shape, and it becomes as spherocyte. Red blood cells can also be coated with complement along with IgG antibodies as another mechanism of opsonization and removal from circulation.³

Laboratory Findings for WAIHA:¹⁻³

CBC: WBC: Normal to increased PLT: Normal to increased Hb: Decreased RETIC: Increased	PBS: Normochromic Normocytic Spherocytes Polychromasia +/- nRBCs	BM: Erythroid hyperplasia Erythrophagocytosis by macrophages
Other Tests: DAT: Positive (IgG & C3b) Osmotic Fragility: Increased Antibody Screen: Positive with all cells Autocontrol: Positive Bilirubin: Increased Haptoglobin: Decreased LD: Increased Hemoglobinuria: Positive Hemosiderinuria: Positive		

()



Cold Agglutinin Disease (CAD)

Associated Conditions:²

Primarily idiopathic

B-cell lymphoproliferative neoplasms

Mycoplasma pneumoniae (anti-I) Infectious mononucleosis (anti-i)

Affected age: > 50 years old.²

Antibody Specificity: Autoanti-I and autoanti-i.²

Autoantibody is an IgM antibody that reacts optimally below body temperature, usually around 4°C². Pathological cold agglutinins will react closer to body temperature (around 30°C).

Can be polyclonal (i.e. infections) or monoclonal (Monoclonal is more pathogenic).¹

Pathophysiology:

Under cold temperatures (circulation in the extremities), the autoantibodies bind to the red blood cells causing them to agglutinate. As the autoantibodies are strong complement activators, complement (C3b) also binds the red blood cells.^{1,2,4}

When the cells return to body temperature (central circulation), the autoantibody unbinds allowing cells to separate and leaves C3b behind remaining on the red blood cell. This leads to complement-mediated hemolysis by macrophages in the liver (extravascular hemolysis).^{1,2,4}

Can cause acrocyanosis and hemolysis is self-limiting.⁵

Laboratory Findings for CAD:^{1,2,4}

CBC:	PBS:	BM:
RBC: Decreased	Agglutination at room temperature (Not present if sample is heated	Erythroid hyperplasia
WBC: Normal	to 37°C)	
PLT: Normal	Spherocytes	
RETIC: Increased	Normohromic	
MCV, MCH, MCHC: Falsely increased	Normocytic	
(due to cold-agglutination)	+/- nRBCs	
Other Tests:		
DAT: Positive for C3b, but negative for IgG or IgM		
IAT: Reactive at < 25°C		
Screen Cells: Positive		
Autocontrol: Positive		
Cord cells: Negative (If autoanti-I, otherwise positive if		
autoanti-i)		
Bilirubin: Increased		
Haptoglobin: Decreased		
Hemoglobinemia		
Hemoglobinuria (Acute)		
Hemosiderinuria (Chronic)		

Paroxysmal Cold Hemoglobinuria (PCH)

Associated Conditions:^{2,3}

Can develop following viral infections or upper respiratory infections

Affected age: Primarily in children.²

Antibody Specificity: autoanti-P (IgG, polyclonal, binds optimally at 4-20°C, reactive at 37°C).^{1,4,5}

Pathophysiology: 1,4,5

Attachment of autoanti-P to cells do not cause the cells to agglutinate but does result in an intravascular, complement-mediated hemolysis.

Autoanti-P is a biphasic antibody meaning that it activates only partial complement at cold temperatures (<37 °C) and full complement at warmer temperatures (37 °C) leading to hemolysis.

Laboratory Findings for PCH:²⁻⁴

CBC:	PBS:	Other Tests:
WBC: Normal	Spherocytes	DAT: Positive for C3d only
Hb: Decreased	Polychromasia	LD: Increased
RETIC: Increased	+/- nRBCs	Haptoglobin: Decreased
	Some Schistocytes	Hemoglobinemia
		Hemoglobinuria
		Donath-Landsteiner Test is Positive:
		Control incubated at 37°C: Hemolysis absent
		Patient sample incubated at 37°C only: Hemolysis absent
		Patient sample incubated at 4°C and 37°C: Hemolysis present

Table 2. Comparative Table of Warm and Cold Immune-Related Hemolytic Anemias





	WAIHA	CAD	РСН
Age (years old)	>40	>50	Children (After viral infection)
Antibody Class	IgG	IgM	IgG
Antibody Specificity	Anti-Rh (most-often)	Anti-I, Anti-i	Anti-P
Optimal Binding Temperature	37°C	4°C	<20°C
DAT Reactivity	IgG	IgG, C3d	C3d
Donath-Landsteiner Test	N/A	Negative	Positive
Type of Hemolysis	Extravascular	Extravascular	Intravascular

Drug-Induced Immune Hemolytic Anemia

Immune hemolytic anemias can also be induced when certain drugs are administered into the body. There are four mechanisms in which they are able to do this:

1. Autoantibody Induction^{1,4,6}

Most commonly caused by methyldopa.

Mechanism mimics that found in warm autoimmune hemolytic anemia. The drug induces the production of warm-reactive antibodies against the red blood cell membrane (self-antigens). Antibodies bind at 37°C and affected red cells are removed by the spleen through extravascular hemolysis.

2. Drug Adsorption (Hapten)^{1,4,6}

Most commonly caused by penicillins.

The drug is non-specifically adsorbed onto the red blood cells and antibodies are produced against the drug itself. As red blood cell pass through the spleen, they are removed by macrophages.

3. Immune Complex Formation (Innocent Bystander)^{1,4,6}

Most commonly caused by quinidine.

An IgG or IgM antibody is produced against the drug when it loosely binds to the red blood cells (antibody-drug immune complex). The immune complex induces the activation of complement, leading to the formation of membrane attack complexes and intravascular hemolysis.

4. Membrane Modification^{1,6}

Most commonly caused by cephalosporins.

Drug modifies the red blood cell membrane causing it to become "sticky". This results in red blood cells becoming coated with many plasma proteins. No hemolysis occurs, but DAT testing will be positive.

Table 3. Comparison of Mechanisms Leading to Drug-Related Immune Hemolytic Anemia

	Autoantibody Induction	Drug Adsorption	Immune Complex	Membrane Modification
Drug Example	Methyldopa	Penicillins	Quinidine	Cephalosporins
Antibody Class	IgG	IgG	IgG or IgM	N/A, due to plasma proteins
DAT	IgG Positive C3 Negative	IgG Positive C3 Negative	C3 Positive IgG Variable	Positive, due to plasma proteins
Eluate	Positive	Usually Negative	Usually Negative	Usually Negative
Type of Hemolysis	Extravascular	Extravascular	Intravascular	No hemlysis

Alloimmune Hemolytic Anemias

Hemolytic anemias can also occur with there is sensitization of red blood cells due previous exposure to another individual's red blood cells.

1. Hemolytic Transfusion Reactions⁷

Hemolytic transfusion reactions occur when there is an incompatibility between the patient's blood (contain alloantibodies) and the transfused cells. Alloantibodies present in the patient's blood binds the antigens on the transfused cells and this results in hemolysis. Transfusion reactions are classified as being acute or delayed.





Table 4. Comparison of Acute and Delayed Transfusion Reactions.

	Acute	Delayed
Time	Immediate; minutes to hours	Days to weeks
Related Blood Groups	ABO	Other non-ABO blood groups
Symptoms	Fever, chills, back pain, pain at infusion site, difficulty breathing, hypotension, urticaria, tachycardia	Usually show no clinical symptoms but may develop a fever
Type of Hemolysis	Intravascular	Extravascular
DAT	Negative (if all transfused cells have all been hemolyzed)	Positive for IgG and/or C3d (Can be negative depending on time of sample collection)
Other Laboratory Findings	Hemoglobinemia Hb: Decreased Bilirubin: Increased after a few days Haptoglobin: Decreased	Hemoglobinuria Hb: Variable Bilirubin: Increased Eluate is positive for offending antibody.

2. Hemolytic Disease of the Fetus and Newborn (HDFN)

Hemolysis that occurs in the fetus or newborn due to incompatibility between the mother's alloantibodies and the fetus's/newborn's blood groups.

Mother's immune system can become sensitized and produce alloantibodies against the blood group antigens that she lacks during a previous pregnancy or transfusion. If the fetus/newborn contains the blood group antigens that the mother has alloantibodies against, HDFN can develop. During pregnancy, alloantibodies are able to pass through the placenta and bind to the red blood cells in the fetus/newborn resulting in hemolysis of the fetal red blood cells.^{4,6}

Newborns appear jaundiced and have high levels of bilirubin at birth.^{4,7} The peripheral blood smear will show increased spherocytes, polychromasia, and increased nucleated red blood cells (normoblastemia).

Alloantibodies can be produced against Rh, ABO, and other blood groups.⁷

References:

1. Smith LA. Hemolytic anemia: immune anemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 348-71.

2. Barcellini W, Fattizzo B. Clinical Applications of Hemolytic Markers in the Differential Diagnosis and Management of Hemolytic Anemia. Dis Markers [Internet]. 2015 Dec 27 [cited 2018 Jun 26];2015:635670. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4706896/

3. Berentsen S. Role of complement in autoimmune hemolytic anemia. Transfus Med Hemotherapy [Internet]. 2015 Sep 7 [cited 2018 Jun 27];42(5):303–10. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4678321/

4. Przekop KA. Extrinsic defects leading to increased erythrocyte destruction – immune causes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 411-25.

5. Packman CH. The clinical pictures of autoimmune hemolytic anemia. Transfus Med Hemotherapy [Internet]. 2015 Sep 11 [cited 2018 Jun 26];42(5):317–24. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4678314/

6. Harmening DM, Yang D, Zeringer H. Hemolytic anemias: extracorpuscular defects. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 250-79).

7. Landis-Piwowar K, Landis J, Keila P. The complete blood count and peripheral blood smear evaluation. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 154-77.

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8.4: Infectious Agents

Malarial Infection

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CHAPTER OVERVIEW

9: Red Blood Cells- Intrinsic Defects of the RBC Membrane Causing Hemolytic Anemia

- 9.1: Hereditary Spherocytosis
- 9.2: Hereditary Elliptocytosis and Related Variants
- 9.3: Hereditary Stomatocytosis Syndromes
- 9.4: Hereditary Acanthocytosis (Abetalipoproteinemia)
- 9.5: Paroxysmal Nocturnal Hemoglobinuria (PNH)
- 9.6: Glucose-6-phosphate Dehydrogenase (G6PD) Deficiency
- 9.7: Pyruvate Kinase (PK) Deficiency

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9.1: Hereditary Spherocytosis

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Images of hereditary spherocytosis peripheral blood smears demonstrating marked spherocytosis and polychromasia. From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R34B2XK9F

Image 2: 50 x oil immersion. https://doi.org/10.7939/R3HM53096

Note:

The hereditary condition results in the formation of spherocytes with a decreased life span, decreased deformability, and a reduced surface-to-volume ratio causing increased osmotic fragility.¹

Mutation:

Genetic mutations in the vertical protein linkages between the membrane and cytoskeleton: α -spectrin, β -spectrin, band 3, ankyrin, and protein 4.2.¹⁻⁴ Results in loss of unsupported membrane overtime, and spherocyte formation.

Inheritance:^{1,3}

Autosomal dominant or recessive depending on which mutations are inherited.

Clinical Features:^{1,3,4}

Jaundice

Fatigue

Pallor

Splenomegaly

Iron Overload

Extramedullary erythropoiesis

Laboratory Results:¹⁻³

CBC:		
Hb: Decreased	PBS:	
MCV: Decreased to Normal	Spherocytes (Variable amounts)	BM.
MCH: Normal to Increased	Polychromasia	M:E Patio: Docroasod
MCHC: Increased	Increased inclusions (HJ bodies,	Frythroid Hyporplacia
(>360 g/L)	pappenheimer bodies)	Erythold Hyperplasia
RETIC: Increased	+/- NRBCs	
RDW: Increased		





Other Tests: Osmotic Fragility: Increased Eosin -5'-maleimide Binding Test: Decreased fluorescence DAT: Negative (AIHA with spherocytes are DAT positive) Markers of EVH: Bilirubin: Increased LD: Increased Urobilinogen: Increased

References:

1. Gallagher PG. Abnormalities of the erythrocyte membrane. Pediatr Clin North Am [Internet]. 2013 Dec 15 [cited 2018 Jun 26];60(6):1349–62. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4155395/

2. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

3. Andolfo I, Russo R, Gambale A, Iolascon A. New insights on hereditary erythrocyte membrane defects. Haematologica [Internet]. 2016 Nov 22 [cited 2018 Jun 26];101(11):1284–94. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC5394881/

4. Da Costa L, Galimand J, Fenneteau O, Mohandas N. Hereditary spherocytosis, elliptocytosis, and other red cell membrane disorders. Blood Rev [Internet]. 2013[cited 2018 Jul 24];27(4):167–78. Available from: http://www.sciencedirect.com/science/article/pii/S0268960X13000192

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9.2: Hereditary Elliptocytosis and Related Variants

Hereditary Elliptocytosis

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Images of hereditary elliptocytosis peripheral blood smears showing numerous elliptocytes. From MLS Collection, University of Alberta.

Image 1: 100x oil immersion. https://doi.org/10.7939/R3F18SW6B

Image 2: 100x oil immersion. https://doi.org/10.7939/R3JS9HQ0S

Hereditary elliptocytosis encompasses group of hereditary conditions that result in the formation of elliptocytes with a decreased erythrocyte lifespan.¹ Variants of hereditary elliptocytosis include Hereditary pyropoikilocytosis and Southeast Asian Ovalocytosis.

Mutation:

Genetic mutations involving the horizontal protein linkages between the membrane and cytoskeleton: α -spectrin, β -spectrin, protein 4.1, glycophorin C).²⁻⁴ These mutations result in a decreased red blood cell lifespan and increased susceptibility to hemolysis (primarily extravascular). Hemolysis is often mild.

Inheritance:1,3,4

Autosomal dominant

Clinical Features:

Patients are usually asymptomatic and discovery of hereditary elliptocytosis is often incidental.^{1,3}

Laboratory Results for Hereditary Elliptocytosis:^{1,5}

CBC:	PBS:	Other Tests:
MCV: Normal to Increased	Elliptocytes (Variable amounts)	Osmotic Fragility: Normal
MCH, MCHC: Normal	During hemolytic episodes may see:	Thermal Stability: Decreased
	Normocytic, normochromic anemia	PCR
	Increased Polychromasia	Hyperbilirubinemia
		LDH: Increased

Hereditary Pyropoikilocytosis (HPP)

A rare variant of hereditary elliptocytosis that presents with severe hemolytic anemia.^{4,5}

Inheritance:^{2,4}

Autosomal recessive

Mutation:^{2,4}

Defects in spectrin that results in red blood cell fragmentation.

Laboratory Findings for HPP:^{2,4,5}

CBC:	PBS:	Other Tests:	
RBC: Decreased	Microspherocytes	Osmotic Fragility: Increased	
Hb: Decreased	Schistocytes	Thermal Sensitivity: Increased	
MCV: Always decreased	Elliptocytes	Eosin -5'-maleimide Binding Test: Decreased	
MCHC: Increased		fluorescence	

Southeast Asian Ovalocytosis (SAO)

A variant of hereditary elliptocytosis that and clinical symptoms are mainly asymptomatic. Ovalocytes are large and may show one or more transverse bars in the cytoplasm of the cell. These ovalycotes are much more rigid than normal red blood cells.⁵ Patients are usually asymptomatic.²

Inheritance:²⁻⁴

Autosomal dominant

Mutation:^{2,3}

Mutation in the Band 3 protein.

Laboratory Findings for SAO:⁵

PBS: Elliptocytes (May show one or more transverse bars)

References:

1. Gallagher PG. Abnormalities of the erythrocyte membrane. Pediatr Clin North Am [Internet]. 2013 Dec 15 [cited 2018 Jun 26];60(6):1349–62. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4155395/





2. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

3. Da Costa L, Galimand J, Fenneteau O, Mohandas N. Hereditary spherocytosis, elliptocytosis, and other red cell membrane disorders. Blood Rev [Internet]. 2013[cited 2018 Jul 24];27(4):167–78. Available from: http://www.sciencedirect.com/science/article/pii/S0268960X13000192

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5. Coetzer T, Zail S. Hereditary defects of the red cell membrane. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 176–95).

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9.3: Hereditary Stomatocytosis Syndromes

Overhydrated Hereditary Stomatocytosis

Condition results in cells with altered intracellular concentrations of sodium (Na+) and potassium (K+) ions. There is an increased permeability of K+ into the cell and increased permeability of Na+ out of the cell. This results in cells with increased volume (cells are overhydrated), a decreased surface-to-volume ratio, and decreased cytoplasm viscosity.¹⁻³

Mutation:^{1,3}

Hereditary defect leading to alterations in the permeability of the red blood cell membrane.

Inheritance: 1-3

Autosomal dominant (Severe hemolysis may indicate a autosomal recessive inheritance)

Laboratory Results:¹⁻³

CBC:	PBS:	Other Tests:
MCV: Increased	Macorcytes	Osmotic Fragility: Increased
MCHC: Decreased	Stomatocytes	

Dehydrated Hereditary Stomatocytosis (Hereditary Xerocytosis)

Defects lead to an increased movement of K+ out of the cell and results in the dehydration of cell. Unlike Stomatocytes, cells have an increased surface-to-volume ratio.²

Inheritance:²⁻⁴

Autosomal dominant

Laboratory Results:¹⁻⁴

CBC:	PBS:	Other Tests:
MCV: Increased	Stomatocytes	Osmotic Fragility: Increased
MCHC: Increased	Target cells	
	Echinocytes	
	Macrocytes	
	RBCs with Hb concentrated at the periphery of the cell	

References:

1. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

2. Coetzer T, Zail S. Hereditary defects of the red cell membrane. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 176–95).

3. Da Costa L, Galimand J, Fenneteau O, Mohandas N. Hereditary spherocytosis, elliptocytosis, and other red cell membrane disorders. Blood Rev [Internet]. 2013[cited 2018 Jul 24];27(4):167–78. Available from: http://www.sciencedirect.com/science/article/pii/S0268960X13000192

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9.4: Hereditary Acanthocytosis (Abetalipoproteinemia)



An image from a peripheral blood smear showing acanthocytes and other poikilocytosis. Hereditary acanthocytosis would typically show acanthocytes as the main red blood cell abnormality. From MLS Collection, University of Alberta, https://doi.org/10.7939/R31J97P8T

Mutation:^{1,2}

Microsomal triglyceride transfer protein (MTP) gene mutation that results in a lack of apolipoprotein B. An increase in sphingomyelin concentration in the RBC membrane leads to increased membrane rigidity and acanthocyte formation.

Inheritance:,1,2

Autosomal recessive

Laboratory Results:^{,1,2}

CBC:	PBS:
MCV: Normal	Acanthocytes
MCH: Normal	
MCHC: Normal	
RETIC: Normal to increased	

Hereditary Acanthocytosis References:

1. Cochran-Black D. Hemolytic anemia: membrane defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 317-33.





2. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

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9.5: Paroxysmal Nocturnal Hemoglobinuria (PNH)

Paroxysmal Nocturnal Hemoglobinuria is an acquired clonal disorder that starts at the stem cell level. Cells produced become susceptible and are destroyed by chronic complement-mediated hemolysis.^{1,2}

Cause(s):^{1,2}

Deficiency in glycosylphosphatidylinositol anchor proteins (GPIs). Normally, CD55 and CD59 act as complement regulators to prevent autologous complement-mediated hemolysis. Without GPIs, cells lack CD55 and CD59 and undergo spontaneous intravascular hemolysis.

Hemolytic episodes (Paroxysms) can be exacerbated by stressors such as inflammation or infections.

Complications:²

Hemolytic Anemia

Bone Marrow Failure

Thrombophilia

Laboratory Results for PNH:^{1,3,4}

CBC: RBC: Decreased WBC: Decreased PLT: Decreased Hb: Decreased MCV: Increased RETIC: Increased	PBS: May see: -nRBCs -Polychromasia If Iron Deficiency Present: -Hypochromic -Microcytic If Folate Deficiency Present: -Oval Macrocytes If BM Failure Present: -Pancytopenia	BM: If BM Failure: -Impaired hematopoiesis -Hypocellular If not BM Failure, may be: -Normo to hypercellular -Erythroid hyperplasia Iron stores: Decreased, or absent Note: Any dysplastic findings may be indicative of MDS.
Iron Studies: Same as iron deficiency anemia if patient becomes iron deficient.	Other Tests: Sucrose Hemolysis Test: Positive for hemolysis Ham's (Acidified Serum Lysis) Test: Positive Flow cytometry (for CD55 and CD59) DAT: Negative Osmotic Fragility: Normal	Tests for IVH: Indirect bilirubin: Increased Haptoglobin: Decreased LD: Increased Hemoglobinemia Hemoglobinuria Hemosiderinuria

References:

1. DeZern AE, Brodsky RA. Paroxysmal Nocturnal Hemoglobinuria. A Complement-Mediated Hemolytic Anemia. Hematol Oncol Clin North Am [Internet]. 2015 Jun 7 [cited 2018 Jun 26];29(3):479–94. Available from: http://www.ncbi.nlm.nih.gov/pmc/articles/PMC4695989/

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3. Cochran-Black D. Hemolytic anemia: membrane defects. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 317-33.

4. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

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9.6: Glucose-6-phosphate Dehydrogenase (G6PD) Deficiency



A peripheral blood smear stained with a supravital stain demonstrating numerous heinz bodies (indicated by arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R35718396

Normal G6PD Function: Hexose Monophosphate (HMP) Shunt and Oxidative Damage^{1,2}

The HMP Shunt describes the glycolytic pathway where glucose is transferred to and catabolized in the cell to produce ATP and maintain sustainable amounts of reduced glutathione (GSH). GSH acts to protect the red blood cell from oxidative damage by reducing oxidative molecules (Peroxides and free oxygen radicals) and becoming oxidized itself.

In the HMP shunt, NADPH is generated by G6PD from NADP which than acts to regenerate reduced glutathione from its oxidized state to allow continued protection from oxidative damage. A lack of G6PD results in increased red blood cell destruction.

G6PD activity is highest in younger cells (i.e. reticulocytes) compared to that of older and more mature red blood cells.¹

Mutation:²

G6PD gene mutation results in decreased G6PD levels.

Inheritance:¹

X-linked

Complications:^{1,2}

Acute hemolytic Anemia

Neonatal Hyperbilirubinemia

Chronic Nonspherocytic Hemolytic Anemia

Causes of Hemolytic Episodes:^{1,2}

Exposure to oxidative agents:

-Drugs (e.g. antimalarials, sulfonamides)

-Infections

-Fava Beans (Favism)





Notes:

G6PD deficiency is commonly associated with bite and blister cells. Oxidized hemoglobins become denatured and form heinz bodies within the red blood cells. These heinz bodies can be removed by splenic macrophages through a "pitting" mechanism, which results in the formation of bite and blister cells.¹

It has been suggested that G6PD deficiency acts as a protective mechanism from malaria as the prevalence of G6PD deficiency coincides with that of malaria.¹

Laboratory Results for G6PD Deficiency:¹

CBC:	PBS:	Other Tests:
WBC: Increased	Bite and Blister Cells	Dye Reduction Test: No colour change
PLT: Normal	Polychromasia	G6PD Assay: Decreased activity
Hb: Decreased	Occasional spherocytes	DAT: Negative
MCHC: Increased	Heinz bodies (Supravital staining)	Hemolysis Markers:
RETIC: Increased		Indirect Bilirubin: Increased
		LDH: Increased
		Haptoglobin: Decreased
		Hemoglobinemia
		Hemoglobinuria

References:

1. Lake M, Bessmer D. Hemolytic anemia: enzyme deficiencies. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 334-47.

2. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

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9.7: Pyruvate Kinase (PK) Deficiency



A peripheral blood smear with a pyknocyte* shown by the arrow (A). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3VX06J4H

Normal PK Function:

Pyruvate Kinase catalyzes the conversion of phosphoenolpyruvate to pyruvate which results in the production of ATP from ADP. A lack of pyruvate kinase results in the ability of cells to maintain proper cell shape, normal lifespan, and low levels of 2,3-PBG.^{1,2}

Mutation:^{1,2}

Mutations in the PKLR gene leading to decreased levels of pyruvate kinase.

Inheritance:^{1,2}

Autosomal recessive

Complications:^{1,2}

Chronic hemolytic anemia

- Splenomegaly
- Jaundice

Gallstones

Notes:1

WBCs contain more pyruvate kinase than RBCs.

Laboratory Results for PK Deficiency:¹

CBC:	PBS:	Other Tests:
Hb: Decreased	Normocytic	Osmotic Fragility: Normal
RETIC: Increased	Normochromic	Pyruvate Kinase Assay: Decreased
*Note: The term "pyknocyte" is not universally	Echinocytes	Indirect bilirubin: Increased
used. It refers to a small, dehydrated, dark-	Pyknocytes*	LDH: Increased
colored RBC.	Post-splenectomy shows varying degrees of anisocytosis and	Haptoglobin: Decreased or absent
	poikilocytosis	

References:

1. Lake M, Bessmer D. Hemolytic anemia: enzyme deficiencies. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 334-47.

2. Keohane EM. Intrinsic defects leading to increased erythrocyte destruction. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 367–93.

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CHAPTER OVERVIEW

10: White Blood Cells and Platelets- Normal Morphology

10.1: Granulocytes and Granulocyte Maturation
10.2: Lymphocytes
10.3: Plasma Cells
10.4: Monocytes
10.5: Macrophages
10.6: Megakaryocytes
10.7: Platelets

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10.1: Granulocytes and Granulocyte Maturation

Myeloblast/Blast

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10.2: Lymphocytes



An image from a peripheral blood smear showing a small lymphocyte (A) and a large lymphocyte (B). 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3JH3DH6Q





An image from a normal peripheral blood smear showing two large lymphocytes. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R30Z71C10





An image from a peripheral blood smear showing a small lymphocyte. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3H12VP79







An image from a peripheral blood smear with a neutrophil (left) and a small lymphocyte (right). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3930P92Z

Notes: Can be characterized as being small or large depending on the amount of cytoplasm. Small lymphocytes are more uniform in appearance whereas large lymphocytes have a variable appearance.¹

Nucleus-to-Cytoplasm Ratio: 5:1 to 2:1 ^{1,2} Nucleoli: May be visible ^{1,2} Nucleus:^{1,2} Round, oval, or indented Dark purple, dense chromatin (heterochromatin) Cytoplasm:^{1,2} Pale blue Scant to moderate Vacuoles may be present Granules:^{1,2} Large: Azurophilic granules may be present Small: typically lack granules (agranular) Normal % in Bone Marrow: 5-15% ² Normal % in Peripheral Blood: 20-40% ²





Lymphocyte Lineage

Lymphocytes can be characterized into two cell types depending on the site of cell maturation:

1. B Cells

Lymphocytes that mature in the bone marrow. These cells are lymphocytes that are able to mature into plasma cells and take part in antibody production.¹

Specific surface markers:^{1,3}

CD10, CD19, CD20, D21, CD22, D24, CD38

2. T Cells

Lymphocytes that mature in the thymus and lymphoid tissues. When these cells become activated, they are able to take part in cell-mediated immunity.¹

Specific surface markers:^{1,3}

CD2, CD3, CD4, CD5, CD7, CD8, CD25

References:

1. Williams L, Finnegan K. Lymphocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 122-43.

2. Rodak BF, Carr JH. Lymphocyte maturation. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 79-88.

3. Czader M. Flow cytometric analysis in hematologic disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 543-60.

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10.3: Plasma Cells



An image from a bone marrow smear showing a plasma cell. Note the perinuclear clearing surrounding the nucleus. 100z oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R30C4T111





An image from a bone marrow smear with a plasma cell (indicated by an arrow). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3JM23X5J

Notes: The maturation of a lymphocyte to a plasma cell marks the production of immunoglobulins. Lymphocytes that mature into plasma cells are of B lineage.¹

Nucleus-to-Cytoplasm Ratio: 2:1 to 1:1²
Nucleoli: None ^{1,2}
Nucleus:^{1,2}
Round or oval
Eccentrically located
Coarse, clumpy, dark purple staining chromatin
Cytoplasm:^{1,2}
Abundant
Darkly basophilic
Perinuclear (clear) zone may be seen around the nucleus (Representing the golgi body)
Vacuoles may be present
Normal % in Bone Marrow: 0-1% ^{1,2}

References:



1. Williams L, Finnegan K. Lymphocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 122-43.

2. Rodak BF, Carr JH. Lymphocyte maturation. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 79-88.

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10.4: Monocytes



An image from a peripheral blood smear showing a monocyte in the center. 50x oil immersion.From MLS Collection, University of Alberta, https://doi.org/10.7939/R3Q52FV0F



An image from a peripheral blood smear showing a monocyte in the top left corner (indicated with an arrow). 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3FN1172B



An image from a peripheral blood smear showing two monocytes (indicated by arrows) with prominent vacuoles. 60x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3MG7GB06

Notes: Monocyte nuclear and cytoplasmic morphology can be highly variable.¹

Nucleus-to-Cytoplasm Ratio: Variable²

Nucleoli: Not visible ²

Nucleus:^{1,2}

Variable shapes (Folds, kidney shaped)

Loose, lacy, violet chromatin

Cytoplasm:^{1,2}

Blue-gray cytoplasm (Ground glass appearance due to fine, diffuse granules)

May have pseudopods

May have vacuoles

Normal % in Bone Marrow: 2%²

Normal % in Peripheral Blood: 3-11%²





References:

1. Landis-Piwowar K. Granulocytes and Monocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 97-121.

2. Rodak BF, Carr JH. Monocyte maturation. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 55-64.

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10.5: Macrophages



An image from a bone marrow smear showing a macrophage with a valuolated and granular cytoplasm. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3DV1D40B





An image from a Cerebrospinal Fluid (CSF) cytospin slide showing erythrophagocytosis in a macrophage. Ingested red blood cells, vacuolation, and hemosiderin granules can be seen within the cell. 60x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R36H4D570

Notes: Macrophages represent the mature form of monocytes when they leave the circulation and enter the tissues.¹

Nucleus-to-Cytoplasm Ratio: N/A ²

Nucleoli: 1-2²

Nucleus:²

Variable shapes (kidney, rounded, indented, oval)

Eccentrically located

Dark purple, coarse, clumped chromatin

Cytoplasm:^{1,2}

Abundant

Irregular shaped

Many azurophilic granules

May contain ingested material and/or storage granules (hemosiderin, red blood cells, lipids, microorganisms, debris)

May contain vacuoles

References:





1. Landis-Piwowar K. Granulocytes and Monocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 97-121.

2. Rodak BF, Carr JH. Monocyte maturation. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 55-64.

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10.6: Megakaryocytes



An image from a bone marrow smear showing a normal megakaryocyte with multiple nuclear lobes. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3FF3MF8N



An image from a bone marrow smear showing a megakaryocyte (indicated by an arrow) in the tails of the smear. 10x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3TQ5RV93



An image from a bone marrow smear showing three megakaryocytes in the tail of the smear. 10x magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3K64B82D

Notes: Develop and are mainly found in the bone marrow. Maturation usually involves the division of nucleus but not the division of the cytoplasm, this gives rise to a polyploid cell.¹

Nucleus-to-Cytoplasm Ratio: Variable²

Nucleoli: N/A²

Nucleus:

Variable number of lobes $(2-32)^2$

Cytoplasm:²

Abundant

Blue to purple cytoplasm

Reddish blue granules may be visible

% in Bone Marrow: 5-10 (per field at 100x magnification)²

% in Peripheral Blood: None





References:

1. Lynne Williams J. The Platelet. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 144–53.

2. Rodak BF, Carr JH. Megakaryocyte maturation. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 31-40.

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10.7: Platelets



An image from the thicker section of a peripheral blood smear showing platelet satellitism around three neutrophils. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R31Z4280R



An image from a peripheral blood smear showing platelet satellitism around a neutrophil. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3F766P1H



An image from a peripheral blood smear demonstrating platelet clumping. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3XD0RD0V

Notes: Platelets are cytoplasmic fragmentations from a megakaryocyte. Fragmentation occurs by the megakaryocyte demarcation membrane system.¹

Nucleus-to-Cytoplasm Ratio: N/A²

Nucleoli: N/A²

Nucleus: N/A²

Cytoplasm:^{1,2}

Light blue to colourless

Azurophilic granules present

% in Bone Marrow: N/A

% in Peripheral Blood: 7-25 per field (100x oil immersion field)

Reticulated Platelets

Immature platelets that contain an abundant amount of RNA.¹





Platelet Clumps and Satellitism

Platelet satellitism is a phenomenon that can occur in vitro when a blood sample is collected in an EDTA anticoagulant tube. Platelets *adhere* to neutrophils by an antibody mediated process and this results in falsely decreased platelet counts.^{3,4}

Platelet clumping can also occur when blood is collected in an EDTA tube. Platelets become activated and *aggregate*. EDTA causes some cell antigens to be unmasked and react with antibodies in the serum.^{3,4}

In both cases, the issue may be corrected when blood samples are collected in sodium citrate anti-coagulated tubes.^{3,4}

References:

1. Lynne Williams J. The Platelet. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 144–53.

2. Rodak BF, Carr JH. Megakaryocyte maturation. In: Clinical hematology atlas. 5th ed. St. Louis, Missouri: Elsevier Inc.; 2017. p. 31-40.

3. Burns C, Dotson M. Hematology Procedures. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 782-814.

4. Clark KS, Hippe TGl. Manual, semiautomated, and point-of-care testing in hematology. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p.187-207.

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CHAPTER OVERVIEW

11: White Blood Cells- Non-Malignant Leukocyte Disorders

- 11.1: Neutrophil Hyposegmentation
- 11.2: Neutrophil Hypersegmentation
- 11.3: Toxic Changes
- 11.4: Pelger-Huet Anomaly
- 11.5: Chediak-Higashi Syndrome
- 11.6: Alder-Reilly Anomaly
- 11.7: May-Hegglin Anomaly
- 11.8: Chronic Granulomatous Disease
- 11.9: Infectious Mononucleosis/Reactive Lymphocytes

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11.1: Neutrophil Hyposegmentation

Hyposegmentation

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11.2: Neutrophil Hypersegmentation

Neutrophil Hypersegmentation

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11.3: Toxic Changes



An image from a peripheral blood smear showing a band with a blue dohle body inclusion found in the center of the cell and toxic granulation. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3930P95D



An image from a peripheral blood smear showing toxic changes (Granulation, vacuolation) in two neutrophils. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3891261T



A peripheral blood smear picture showing granulocytes with toxic changes. The Band (center) shows toxic granulation and a dohle body. The neutrophil (bottom) shows toxic vacuolation. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3HT2GT1K

Cell Features:^{1,2}

Toxic morphological changes are seen in neutrophils. A left shift with an increase in immature granulocytes typically accompanies toxic changes. In order to report toxic changes, typically two out of the three features should be seen in the majority of neutrophils:

1. Toxic Granulation:¹⁻³

Dark blue-black peroxidase positive granules that appear in the cytoplasm of the neutrophil. Appear very similar to Alder-Reilly bodies found in Alder-Reilly anomaly but is commonly found with other features of toxicity. Can be found in mature neutrophils, bands, and metamyelocytes.

2. Toxic Vacuolation:¹⁻³

Clear, circular, and unstained cytoplasmic areas that represent phagocytosis or autophagocytosis. Vacuoles may contain bacteria or yeast if the patient is septic.

3. Dohle Bodies:¹⁻³

Pale blue, round or elongated cytoplasmic inclusions containing remnant ribosomal ribonucleic acid (RNA) in parallel rows (rough endoplasmic reticulum). Often present in mature neutrophils and bands near the periphery of the cell. Bodies are non-specific and





can appear in several conditions such as pregnancy, cancer, burns, and infections.

Note: A left shift is usually seen on the peripheral blood smear when toxicity is present. A Left shift refers to the increase presence of immature bands and myeloid precusors.

Cause:¹

Reaction to infection, inflammation, stress, and granulocyte colony-stimulating factor therapy

Laboratory Features:^{1,2}

CBCD:	Peripheral Blood Smear:
Moderate leukocytosis	At least 2 of 3 toxic changes in the majority of neutrophils
Neutrophilia	Left shift (often)

References:

1. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

2. Jones KW. Evaluation of cell morphology and introduction to platelet and white blood cell morphology. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 93-116.

3. Landis-Piwowar K. Nonmalignant disorders of leukocytes: granulocytes and monocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 388-407.

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11.4: Pelger-Huet Anomaly

An interactive or media element has been excluded from this version of the text. You can view it online here: https://pressbooks.library.ualberta.ca/mlsci/?p=643

Images of Pelger-Huet Anomaly in various peripheral blood smears showing numerous hyposegmented neutrophils with mature clumped chromatin. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3DB7W572

Image 2: 50x oil immersion. https://doi.org/10.7939/R32Z13500

Image 3: 50x oil immersion. https://doi.org/10.7939/R3Z60CH59

PBS Key Features:¹⁻⁴

Neutrophil nuclei appear hyposegmented – can appear as a single round nucleus (unilobed, homozygous Pelger-Huet Anomaly) or dumbbell shaped (bilobed, heterozygous Pelger-Huet Anomaly). Anomaly is differentiated from a left shift by displaying mature chromatin pattern, abundant cytoplasm (low nuclear:cytoplasmic ratio), mature granulation, and an absence of toxic changes.

<u>Congenital Pelger-Huet:</u> granulocytes show normal granulation, 50-90% of neutrophils are affected.

<u>Pseudo Pelger-Huet:</u> seen in leukocyte malignancies and Myelodysplastic Syndrome, hypogranulation and other Dy's plastic features may be present, 10-30% of neutrophils are affected.

Clinical Significance and Cause:^{1,3,5}

Pelger-Huet Anomaly is benign and cell function is normal. Psuedo Pelger-Huet may indicate leukocyte malignancies and myelodysplasia.

<u>Congenital</u>: Lamin β -receptor gene mutation.

<u>Acquired (Pseudo-Pelger-Huet)</u>: Hematologic malignancies such as myelodysplastic syndrome (MDS), acute myeloid leukemia (AML), myeloproliferative neoplasms (MPNs). Pseudo-Pelger-Huet may also be seen during infections, and drug interactions.

Inheritance Pattern:^{1-3,5}

Autosomal dominant

CBC:²

Congenital Pelger-Huet: Cytopenias often absent

<u>Pseudo-Pelger-Huet:</u> Cytopenias often present

References:

1. Turgeon ML. Nonmalignant Disorders of Granulocytes and monocytes. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 206-16.

2. Cunningham JM, Patnaik MM, Hammerschmidt DE, Vercellotti GM. Historical perspective and clinical implications of the Pelger-Huet cell. Am J Hematol [Internet]. 2009 Oct 20 [cited 2018 Jul 10];84(2):116–9. Available from: https://doi.org/10.1002/ajh.21320

3. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

4. Harmening DM, Marty J, Strauss RG. Cell biology, disorders of neutrophils, infectious mononucleosis, and reactive lymphocytosis. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 305-30.

5. Landis-Piwowar K. Nonmalignant disorders of leukocytes: granulocytes and monocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 388-407.

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11.5: Chediak-Higashi Syndrome



An image of a peripheral blood smear demonstrating neutrophils with abnormally large fused granules seen in Chediak-Higashi Syndrome. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R39S1M158



An image of a peripheral blood smear demonstrating a neutrophil with abnormally large fused granules (top) and a lymphocyte containing a single large granule in the cytoplasm (bottom) seen in Chediak-Higashi syndrome. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3707X414



An image of a peripheral blood smear demonstrating a lymphocyte containing a single large granule in the cytoplasm seen in Chediak-Higashi syndrome. 100X oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3377697R

PBS Key Features:^{1,2}

Leukocytes contain abnormally large lysosomal granules in the cytoplasm. Granules represent the aggregation of primary granules combined with the fusion of secondary granules.

Cause:³

Mutation in the CHS1/LYST gene which encodes for a vesicle transport protein.

Inheritance Pattern:³⁻⁵

Autosomal recessive

Clinical Significance:^{1,3-5}

Development of lysosomes are abnormal resulting in the fusion of granules. The syndrome results in impaired chemotaxis, defective degranulation, and defective killing of bacteria. Granulocytes, Platelets, Monocytes, and lymphocytes are dysfunctional.





Patients often present with oculocutaneous albinism, recurrent bacterial infections and bleeding tendencies. Complications develop during early childhood.

CBC:^{1,2,5}

Anemia

Neutropenia

Thrombocytopenia

References:

1. Harmening DM, Marty J, Strauss RG. Cell biology, disorders of neutrophils, infectious mononucleosis, and reactive lymphocytosis. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 305-30.

2. Bain BJ. Morphology of blood cells. In: Blood cells: a practical guide [Internet]. 5th ed. Chichester, UK: John Wiley & Sons, Ltd; 2015 [cited 2018 Jul 10]: 67-185. Available from: http://doi.wiley.com/10.1002/9781118817322

3. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

4. Turgeon ML. Nonmalignant Disorders of Granulocytes and monocytes. In: Clinical hematology: theory and procedures. 4th ed. Philadelphia, PA: Lippincott Williams & Wilkins; 1999. p. 206-16.

5. Landis-Piwowar K. Nonmalignant disorders of leukocytes: granulocytes and monocytes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 388-407.

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11.6: Alder-Reilly Anomaly

PBS Key Features:^{1,2}

Granulocytes show metachromatic and darkly staining inclusions (Alder-Reilly bodies) containing partially digested mucopolysaccharides that resemble toxic granulation but are permanent (non-transient). Anomaly is differentiated from toxicity by a lack of Dohle bodies, left shift, and neutrophilia. Abnormal granules may also be seen in lymphocytes and monocytes.

Cause:1,2

Incomplete degradation of mucopolysaccharides (Mucoplysaccharidosis disorder)

Inheritance Pattern:¹⁻³

Autosomal recessive

Clinical Significance:¹⁻⁵

Leukocyte function is not impaired. Associated syndromes include Tay-Sachs disease, Hunter syndrome, Hurler syndrome, and Maroteaux-Lamy polydystrophic dwarfism which all result in different clinical symptoms.

CBC:

N/A

References:

1. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

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11.7: May-Hegglin Anomaly



An image from a peripheral blood smear demonstrating a blue dohle body-like inclusion in a neutrophil and a giant platelet seen in May-Hegglin anomaly. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3W669Q4B



An image from a peripheral blood smear demonstrating blue dohle body-like inclusions in two neutrophils seen in May-Hegglin anomaly. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R34Q7R53S



An image from a peripheral blood smear demonstrating blue dohle body-like inclusions in two neutrophils along with enlarged platelets seen in May-Hegglin anomaly. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R30Z71B9D

PBS Key Features:¹⁻⁴

Graunulocyte and Monoyte cytoplasms contain large basophilic inclusions that resemble Dohle bodies but are much larger and elongated. Inclusions are composed of precipitated myosin heavy chains.

Giant platelets and thrombocytopenia are also associated with this anomaly.

Cause:1

MYH9 gene mutation

Inheritance Pattern:^{1,5}

Autosomal dominant

Clinical Significance:^{1-3,5}

May-Hegglin anomaly is a platelet disorder that can cause mild bleeding tendencies but majority of patients are asymptomatic. Degree of bleeding is correlated to the degree of thrombocytopenia. Leukocyte function is unaffected.





CBC:^{1,5}

Variable thrombocytopenia

References:

1. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

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11.8: Chronic Granulomatous Disease

PBS:1

Leukoctye morphology is normal.

Cause:²

Mutations in the NADPH oxidase subunit genes.

Inheritance Pattern:²⁻⁴

Autosomal recessive, X-linked recessive

Clinical Significance:²⁻⁴

Antmicrobial activity defect where neutrophils and monocytes are unable to kill catalase positive organisms after ingestion. The respiratory burst is not activated and cells are unable to produce reactive oxygen species and superoxide. Disease results in recurrent and life-threatening bacterial and fungal infections in the first year of life.

Infections occur often in the lung, skin, lymph nodes, and liver. Granuloma formation can be found in various organs and cause obstruction.

Additional Tests:¹

Nitroblue Tetrazolium Test (NBT)

Flow Cytometry

References:

1. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

2. Harmening DM, Marty J, Strauss RG. Cell biology, disorders of neutrophils, infectious mononucleosis, and reactive lymphocytosis. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 305-30.

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11.9: Infectious Mononucleosis/Reactive Lymphocytes

An interactive or media element has been excluded from this version of the text. You can view it online here: https://pressbooks.library.ualberta.ca/mlsci/?p=667

Images of peripheral blood smears with heterogeneous reactive lymphocytes with prominent basophilic skirting of the cytoplasm. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3G44J57B

Image 2: 50x oil immersion. https://doi.org/10.7939/R34M91R9B

Image 3: 60 x oil immersion. https://doi.org/10.7939/R3GQ6RH6F

Cell Features:¹⁻³

Large reactive (atypical) lymphocytes that represent activated T cells. The cytoplasm shows characteristic basophilic skirting in areas where there is contact with red blood cells. Red blood cells look as if they are creating indents in the cytoplasm. The population of reactive lymphocytes is heterogeneous with diverse shapes and sizes in cytoplasm and nuclear shapes.

Cause:1,3-5

Epstein-Barr Virus (EBV) infection that is usually acute, benign, and self-limiting.

Age group affected:^{1,3,5}

Young adults (approx. 15-25 years of age)

Common Clinical Symptoms:^{1,4}

Classic Triad: Pharyngitis, fever, lymphadenopathy.

Laboratory Features:^{1,3,5}

CBC:	PBS
Leukocytosis	Reactive lymphocytes
Absolute lymphocytosis	

Other Tests:^{1,3,5}

Positive heterophile antibody*

Positive EBV specific antigen and antibody (ELISA)*

Elevated C-reactive protein (CRP)

Viral cultures

Flow cytometry (to rule out malignancies with similar cell morphologies)

*Positivity for antigen or antibody varies depending on the date of testing. Some antigen or antibodies may appear only after a few weeks of infection.

Immunologic Markers:²

CD3, CD4 or CD8

References:

1. Manonneaux S. Nonmalignant leukocyte disorders. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 475-97.

2. Holmer LD, Bueso-Ramos CE. Chronic lymphocytic leukemia and related lymphoproliferative disorders. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 440-65.

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CHAPTER OVERVIEW

12: White Blood Cells- Acute Leukemia

- 12.1: Introduction to Leukemias
- 12.2: Acute Lymphoblastic Leukemia (ALL)
- 12.3: Acute Myelogenous Leukemia (AML)
- 12.4: Acute Promyelocytic Leukemia (APL)
- 12.5: Cytochemical Testing
- 12.6: Flow Cytometry, Cytogenetics and Molecular Genetics

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12.1: Introduction to Leukemias



An image from a peripheral blood smear demonstrating blasts with a loose, immature chromatin pattern, a high nuclearcytoplasmic ratio, and basophilic cytoplasm seen in an Acute Leukemia. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3TB0Z99Q





An image from a bone marrow smear demonstrating an abundance of blasts and a reduction of normal hematopoietic cells seen in an Acute Leukemia. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3Z31P43G

Leukemia: Describes tumors that originate from the bone marrow.¹

Lymphoma: Describes tumors that originate from the lymphatic tissues.¹

The causes of acute leukemia are vast. There are a number of factors that can lead to the development of leukemia, such as: genetic mutations, environmental factors (e.g. exposure to drugs, chemicals, radiation), inherited syndromes (e.g. Down syndrome, fanconi anemia), viral infections (e.g. HIV), immunologic dysfunction (e.g. immunosuppressants), or idiopathic factors.¹

There are different classification systems that exist to categorize acute and chronic leukemias. Two examples are the French-American-British (FAB) system and the World Health Organization (WHO) system.¹⁻⁴

Note: Please be aware that these schemes are updated periodically and the sources used in this ebook may not reflect the most current classification systems used.

Leukemias are described as being "acute" or "chronic" and specified as to which cell lineage and maturation stage is affected.⁵

Table 1. Comparison of Acute and Chronic Leukemias.^{5,6}

Acute	Chronic
All ages affected	Mainly adults affected
Rapid onset	Insidious onset
Involve immature cells	Involve mature cells
≥20%* blasts in PBS or BM	≤20% blasts in PBS or BM
*WHO Classification Criteria	





References:

1. McKenzie SB. Introduction to hematopoietic neoplasms. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 424-45.

2. Roquiz W, Gandhi P, Kini AR. Acute leukemias. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 543-60.

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12.2: Acute Lymphoblastic Leukemia (ALL)

WHO Classification (2008):1

As of 2008, the WHO has classified ALL into categories based on the the lymphoblast origin and genetic abnormalities:

1. B Lymphoblastic Leukemia/Lymphoma

With recurrent genetic abnormalities

• Not otherwise specified (NOS)

2. T Lymphoblastic Leukemia/Lymphoma

Affected Age: Primarily children 2-5 years old.²

Affected Cell:1

B and T lymphoblasts

ALL Blasts Cell Features: 1,3,4

Size: Blasts are variable in size, usually smaller than myeloblasts

Nucleoli: Often indistinct, 0-2

Chromatin: Fine to coarse, dispersed chromatin

Cytoplasm: Usually scant and basophilic, vacuoles may also be present

Auer Rods: Not present

Auer Rods are thought to be fused primary granules and are only found in myeloblasts. Presence of auer rods is distinctive of AML and can be used to differentiate the condition from ALL if they are present.²

Laboratory Results for ALL:^{1,3,5}

CBC: RBC: Decreased WBC: Variable PLT: Decreased	PBS: Lymphoblasts Normocytic Normochromic Anisocytosis, poikilocytosis, and nRBCs are usually not present.	BM: ≥20% Lymphoblasts Hypercellular with replacement hematopoietic tissue	of	normal
Immunologic Markers: <u>B cell:</u> TdT, CD10, CD19, CD22, CD24, CD34 <u>T cell:</u> TdT, CD1, CD2, CD3, CD4, CD5, CD7, CD8, CD10	Other Tests: LD: Increased Hyperuricemia Hypercalcemia Cytogenetics Cytochemical Stains			

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

2. McKenzie SB. Introduction to hematopoietic neoplasms. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 424-45.

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12.3: Acute Myelogenous Leukemia (AML)



An image from a bone marrow smear showing an abundant amount of myeloid precursors and a few myeloblasts seen in a patient with acute myeloid leukemia. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R32V2CR7N



An image from a peripheral blood smear showing myeloblasts. One myeloblast contains a faint, pink staining, auer rod. 60x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3C24R34V



An image from a peripheral blood smear showing a myeloblast with multiple auer rods (stained pink) seen in AML. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3DV1D397

WHO Classification (2008):

As of 2008, acute myeloid leukemias have been classified into different subcategories based on the type of genetic abnormalities, type of myeloid cell type affected, and by cell characteristics:¹

- 1. Acute myeloid leukemia with recurrent genetic abnormalities
- 2. Acute myeloid leukemia with myelodysplastic changes
- 3. Therapy related myeloid neoplasms
- 4. Acute myeloid leukemia, not otherwise specified (NOS)
- 5. Myeloid Sarcoma
- 6. Myeloid Proliferations related to Down syndrome
- 7. Blastic plasmacytoid dendritic cell neoplasm

Affected Age: Adults.²

Cells of myeloid lineage are affected:¹

- Myeloblast
- Monocytes
- Erythrocytes
- Megakaryocytes
- Dendritic cells

AML Blasts Cell Features:³

Size: Myeloblasts are usually larger compared to lymphoblasts and have a consistent appearance.

Nucleoli: Prominent, 1-4 present

Chromatin: Loose open chromatin

Cytoplasm: Often abundant and granules may be visible

Auer Rods: may be present (stained faint pink with Wright's stain)

Auer Rods are thought to be fused primary granules and are only found in myeloblasts. Presence of auer rods is distinctive of AML and can be used to differentiate the condition from ALL if it is present.¹

Laboratory Results for AML:^{1,4,5}





CBC: RBC: Decreased WBC: Variable PLT: Decreased Hb: Decreased RDW: Increased	PBS: Myeloblasts ± Auer rods Macrocytic RBCs May see hypogranular PLT, Giant PLT Neutropenia (Can appear dysplastic) May see Basophilia, Eosinophilia, Monocytosis	BM: ≥20% Myeloblasts ± Auer rods Hypercellular Decreased fat
Immunologic Markers: Depending on the subgroup, cells may be positive for: CD11b, CD13, CD14, CD33, CD34, CD117	Other Tests: LD: Increased Hyperuricemia Hyperphosphatemia Hypocalcemia Hypokalemia Cytogenetics Cytochemistry	

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

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3. Gatter K, Cruz F, Braziel R. Introduction to leukemia and the acute leukemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 331-370.

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5. Bentley G, Leclair SJ. Acute Myeloid Leukemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 500-21.

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12.4: Acute Promyelocytic Leukemia (APL)



An image from a bone marrow smear demonstrating numerous myeloid precursors arrested at the promyelocyte stage, and cells with bundles of auer rods known as faggot cells in acute promyelocytic leukemia. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3WH2DW4D



An image from a bone marrow smear showing faggot cells with bundles of auer rods seen in acute promyelocytic leukemia. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3N010854



An image from a bone marrow smear showing faggot cells with bundles of auer rods seen in acute promyelocytic leukemia. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3H708G10

Note: APL is a subtype of AML where the promyelocytes specifically are affected. It is classified under the "AML with recurrent genetic abnormalities" and is associated with a specific genetic abnormality: t(15;17)(q22;q12); PML-RARA.

Affected Age:

Most often middle age adults, but APL can develop at any age,^{1,2}

Affected Cell: 1

Promyelocyte

Cell Description:^{1,2}

<u>Nucleus:</u> The shape of the nucleus is variable and can be bilobed, multilobed, indented, or folded. May demonstrate a typical "butterfly shape" appearance.

<u>Cytoplasm</u>: The hypergranular variant of APL is characterized by numerous promyelocytes with abundant abnormal, coarse, and dense granulation. The granules stain light pink to reddish-purple and heavily cover and obscure the nucleus of the cell. In the microgranular variant, the abnormal promyelocyte cells demonstrate a lack of granulation but often have abnormal bi-lobed nuclei.





Cells may contain characteristic multiple or bundles of auer rods (light pink) which the cells are then termed "Faggot cells."

The granules in the neoplastic promyelocytes have procoagulant activity. Because of this, Disseminated Intravascular Coagulation (DIC) is associated as a complication of APL. Coagulation studies including fibrinogen and DDimer measurement can aid in the diagnosis of DIC in these patients.

References:

1. Bentley G, Leclair SJ. Acute Myeloid Leukemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 500-21.

2. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

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12.5: Cytochemical Testing

Cytochemistry involves staining cells in vitro to visualize certain cellular components that will help determine the lineage of the cell. After staining, cells are examined microscopically.¹

Myeloperoxidase (MPO)

MPO is an enzyme that is found in the primary granules of all granulocytes and monocytes and not present in lymphocytes. MPO is useful for differentiating between ALL and AML blasts.^{2,3}

Results:^{2,3}

Granulocytes, Myeloblasts, Auer rods: Positive

Monocytes: Negative to weak positive

Lymphocytes: Negative

Sudan Black B

Stains lipids present in the primary and secondary granules of granulocytes and monocyte lysosomes. Similar to MPO, it is useful for differentiating between AML and ALL but it is less specific.¹⁻³

Results:¹⁻³

Granulocytes, Myeloblasts: Positive

Monocytes: Negative to weakly positive

Lymphocytes: Negative

Nonspecific Esterases (Alpha-napthyl acetate esterase)

An enzymatic stain that is used to differentiate granulocytes from monocytes. The stain is considered nonspecific because other cells may also be stained.¹⁻³ **Results:**¹⁻³

Monocytes: Diffusely positive (Positivity can be inhibited by sodium fluoride)

Granulocytes: Negative

Lymphocytes: Negative (Except T lymphocytes which show focal positivity)

Specific Esterase (Chloroacetate esterase)

An enzymatic stain that is specific for granulocytes.¹⁻³ **Results:¹⁻³** Granulocytes (Neutrophils), Myeloblasts, Auer rods: Positive Monocytes: Negative to weak positive

Periodic Acid Schiff (PAS)

PAS stains glycogen related compounds.³ PAS is useful for the identification of lymphoid cells.

Results:¹

Leukemic erythroblasts: Positive (normal erythroblasts are not positive)

Lymphoblasts: Block positivity

Leukocyte Alkaline Phosphatase (LAP)

LAP is an enzyme present in the secondary granules of neutrophils and not present in eosinophils or basophils. LAP is useful for distinguishing between chronic myelogenous leukemia (CML) from other conditions that show increased leukocyte counts.¹

Results:¹

CML: Low LAP score Leukemoid Reactions: High LAP score

Acid Phosphatase





An enzyme that is present in the lysosomes of normal leukocytes.¹

Results:¹

T cell ALL: Positive

Hairy cells: Positive

Hairy cells also show positivity for tartrate resistant acid phosphatase (TRAP) whereas other cells would be inhibited by TRAP.¹

Terminal Deoxynucleotidyl transferase (TdT)

TdT is a DNA polymerase found in immature cells.¹ Results are useful in identifying lymphoblastic leukemias.

Results:¹

Immature lymphocytes (ALL): Positive

Table 1. Cytochemistry Staining for ALL and AML Subgroups.¹⁻⁵

Acute Leukemia Subgroup	МРО	Sudan Black B	Nonspecific Esterase	Specific Esterase	PAS
ALL	-	-	±	-	Block +
AML	+	+	-	+	±
AMML	+	+	+	+	-
AMoL	-	±	+	-	_
AEL	+	+	±	+	+
AMkL	_	-	±	+	+
APL	+	+	±	+	-
ALL = Acute lymphoblastic let AML = Acute myeloid leukem AMML = Acute myelomonocy AMoL = Acute monoblastic an AEL = Acute Erythroid Leuker AMkL = Acute megakaryoblas APL = Acute promyelocytic let	ukemia ia rtic leukemia nd monocytic leukemia mia stic leukemia ukemia		+ = Positive - = Negative ± = Can be positive or ne	gative	

References:

1. McKenzie SB. Introduction to hematopoietic neoplasms. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 424-45.

2. Roquiz W, Gandhi P, Kini AR. Acute leukemias. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 543-60.

3. Gatter K, Cruz F, Braziel R. Introduction to leukemia and the acute leukemias. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 331-370.

4. Bentley G, Leclair SJ. Acute Myeloid Leukemias. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 500-21.

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12.6: Flow Cytometry, Cytogenetics and Molecular Genetics

Classification of Leukemia

The classification of Acute Leukemia relies on the use of a variety of laboratory results, including morphology, immunophenotyping, genetic features, and clinical features. Classification allows for appropriate disease management, treatment, prognosis, and monitoring to occur. The laboratory is crucial in this aspect. The following is a brief summary of the type of laboratory testing involved in the classification of Acute Leukemia in addition to what has already been discussed.

Flow Cytometry

Flow Cytometry, also known as immunophenotyping, is a technique that can be used to help determine a cell's lineage based on cell markers (e.g Cluster of Differentiation/CD Markers) present and the stage of maturation of a cell.¹

Principle:

Monoclonal antibodies with fluorescent labels that are specific for the surface antigen of interest are incubated with the sample. Samples are taken up by the flow cytometer and injected into a stream of sheath fluid to allow cells to be positioned centrally, this process is called hydrodynamic focusing. A laser is directed at the cells and the bound antibodies fluoresce. Fluorescence detectors are used to detect the fluorescence and a scatter graph is produced based on the antibodies bound.² Other properties such as light scatter (in the forward and side direction) are combined with fluorescence intensity measurements to distinguish cell populations.

Flow cytometry can be used to help determine what cells are present to help diagnose acute leukemias and other hematological disorders.

Cell Lineage	Surface Markers
Immature Cells	CD34, CD117
Granulocytes, Monocytes	CD13, CD14 ,CD15, CD33
Erythrocytes	CD71, Glycophorin A
Megakaryocytes	CD41, CD42, CD61
T Lymphocytes	CD2,CD3, CD4, CD5, CD7, CD8
B Lymphocytes	CD19, CD20, CD22

Table 1. Common Surface Markers for Blood Cells.²

Cytogenetics

Cytogenetics involve the identification of abnormal karyotypes which may be characteristic to a related disorder.¹

Fluorescence In Situ Hybridization (FISH)

FISH is a molecular method that is a cytogenetic tool that is used to detect chromosomal abnormalities such as translocations, deletions, inversion, and duplications.3

The method involves using a fluorescently labelled DNA or RNA probe that is complementary to a specific target sequence. After denaturing double stranded DNA to single stranded DNA, the labelled probe is allowed to incubate and hybridize with the DNA. After incubation, the sample is washed to remove any unbound probes and then a counterstain is added to assist examination. Samples are examined with a fluorescent microscope to look for any chromosomal abnormalities in the cells.³

Molecular Genetics

Molecular genetics involve the use of molecular techniques to identify specific genetic sequences and mutations that can be characteristic for a diagnosis.¹

Polymerase Chain Reaction (PCR)





PCR is commonly used to amplify a specific target sequence such a mutation.⁴

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1. McKenzie SB. Introduction to hematopoietic neoplasms. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 424-45.

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CHAPTER OVERVIEW

13: White Blood Cells- Mature Lymphoid Neoplasms

- 13.1: Introduction to Mature Lymphoid Neoplasms
- 13.2: Chronic Lymphocytic Leukemia (CLL)
- 13.3: Hairy Cell Leukemia (HCL)
- 13.4: Waldenstrom Macroglobulinemia
- 13.5: Monoclonal Gammopathy of Undetermined Significance (MGUS)
- 13.6: Plasma Cell Myeloma (Multiple Myeloma)

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13.1: Introduction to Mature Lymphoid Neoplasms

These group of disorders are also known as lymphoproliferative disorders and involve the clonal proliferation of mature lymphocytes.¹ The proliferation of these cells causes the formation of lymphomas and leukemias.²

As with acute leukemia, there are also a variety of factors that can lead to the development of these disorders:³

- 1. Acquired mutations leading to altered oncogene and tumor suppressor gene functions.
- 2. Inherited immunodeficiency syndromes that are associated with these neoplasms
- 3. Environmental factors that can lead to the development of neoplasms (e.g. viral and bacterial infections)

The 2008 WHO Classification categorizes the related disorders based on the type of cell that is involved (B or T cell).⁴

List of Mature B cell Neoplasms as per WHO 2008:⁴

Chronic lymphocytic leukemia

Hairy Cell Leukemia

Waldenstrom macroglobulinemia (lymphoplasmacytic lymphoma)

Plasma cell neoplasms

Monoclonal gammopathy of undetermined significance (MGUS)

Plasma cell myeloma/multiple myeloma

Note: WHO 2008 lists additional disorders under mature B-cell neoplasms and as well has mature T-cell neoplasms. Only those listed above will be discussed in this eBook.

References:

1. Holmer LD, Bueso-Ramos CE. Chronic lymphocytic leukemia and related lymphoproliferative disorders. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 440-65.

2. Czader M. Mature lymphoid neoplasms. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p. 619-41.

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13.2: Chronic Lymphocytic Leukemia (CLL)



An image from a peripheral blood smear showing numerous small mature lymphocytes and smudge cells seen in a patient with CLL. 50X oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3SQ8QZ8C



An image from a CLL peripheral blood smear showing a smudge cell (A) and mature small lymphocytes (B). 100x oil magnification. From MLS Collection, University of Alberta, https://doi.org/10.7939/https://doi....7939/R3DJ58X9R

Cell Features:¹⁻⁴

Abundant small mature small lymphocytes

Scant Cytoplasm

Condensed/clumped chromatin (often described as "soccer ball" or "parched earth" appearance)

Cause:¹⁻⁴





A mature B cell neoplasm with no specific agent or cause. Cytogenetic findings show relation to trisomy 12 and other chromosomal deletions.

Age group affected: Mainly older adults >50 years old.^{1,5,6}

Laboratory Features for CLL:^{1,2,6}

CBC: Normochromic, Normocytic Anemia Normal to Decreased reticulocyte count Neutropenia Thrombocytopenia	PBS: Small mature lymphocytes Presence of smudge cells Platelet and granulocyte morphology is normal	BM: Lymphocytic infiltration, reduced numbers of normal hematopietic cells
Immunologic markers: CD5, CD19, CD20, CD23	Other useful tests: FISH PCR Associated with hypogammaglobulinemia	

References:

1. Holmer LD, Bueso-Ramos CE. Chronic lymphocytic leukemia and related lymphoproliferative disorders. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 440-65.

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13.3: Hairy Cell Leukemia (HCL)



An image from a peripheral blood smear demonstrating two hairy cells with abundant pale blue cytoplasm, small cytoplasmic projections, and mature-looking nucleus giving the cell a "fried egg" appearance. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3ZG6GP2C



An image from a peripheral blood smear demonstrating the different forms of hairy cells. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3610W71F



An image from a peripheral blood smear showing a form of hairy cell with hairy cytoplasmic projections (left) and a different form of hairy cell with abundant pale blue cytoplasm (right). 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R32805D72

Cell Features:¹⁻⁴

Two types of characteristic B lymphocyte morphologies can be seen in Hairy Cell Leukemia. Hairy cells are small to medium sized cells with either serrated cytoplasmic projections giving it a "hairy" appearance or more abundant pale blue cytoplasm giving it a "fried egg" appearance. The nucleus can be oval or indented, lacks nucleoli, and has an evenly stained mature chromatin patttern.

Cause:1,5,6

An indolent clonal mature B cell disorder has been associated with the BRAF -V600E mutation.

Age Group Affected:^{2,6}

Middle-aged adults (median age of 55 years).

Laboratory Features for HCL:^{1,2,4,6}

CBCD: Pancytopenia Relative lymphocytosis	Peripheral Blood Smear: Hairy Cells (± Fried egg appearance)	Bone Marrow: Dry Tap due to fibrosis (Aspirate) Hypocellular
Immunologic markers: CD19, CD20, CD22, CD25, CD123, Annexin A1	Other useful tests: Tartrate-resistant acid phosphatase (TRAP) stain positive	

References:

1. Grever MR, Abdel-Wahab O, Andritsos LA, Banerji V, Barrientos J, Blachly JS, et al. Consensus guidelines for the diagnosis and management of patients with classic hairy cell leukemia. Blood [Internet]. 2016 Feb 2 [cited 2018 Jun 22];129(1):553–61. Available from: http://www.bloodjournal.org/cgi/doi/10.1182/blood-2016-01-689422

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13.4: Waldenstrom Macroglobulinemia

Features:

Commonly characterized by an IgM monoclonal gammopathy. Increased igM may also result in cryoglobulinemia. Deposits of IgM into tissues and organs can result in intestinal dysfunction, clotting, and neuropathic complications.¹

Cause:^{1,2}

Genetic mutations

Inherited associations

Following viral infections (e.g. Hepatitis C).

Age Group Affected: 60-70 years old, slightly more predominant in males.^{1,2}

Laboratory Results for Waldenstrom Macroglobulinemia:^{1,2}

CBC: WBC: Increased (But lower than CLL)	PBS: Increased number of small lymphocytes and maybe plasmacytoid lymphocytes.	BM: Infiltration of small lymphocytes Variable number of plasma cells and plasmacytoid lymphocytes.
Immunologic Markers: CD19, CD20, CD22, CD25, CD27, CD38, CD79a	Other Tests: Monoclonal Paraprotein IgM: Positive	

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

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13.5: Monoclonal Gammopathy of Undetermined Significance (MGUS)

Features:

MGUS is characterized has having an increase in serum M protein, clonal plasma cells, a lack of CRAB symptoms (hyperCalcemia, Renal failure, Anemia, or lytic **B**one lesions), and no diagnosis of any other B cell lymphoproliferative disorder.^{1,2}

Patients are asymptomatic and present with no other physical abnormalities.^{1,3}

Cause:

No specific cause has been linked to the development of MGUS.¹

Age Group Affected: >50 years old.^{1,3}

Laboratory Results for MGUS: ^{1,2}

BM:	Immunologic Markers:	Other Tests:
Aspirates:	CD19, CD38, CD138	Protein electrophoresis
Increased Plasma cells (<10%)		Immunofixation
Biopsies:		Flow cytometry
Slightly increased numbers of plasma cells with minimal morphological abnormalities		FISH

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

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13.6: Plasma Cell Myeloma (Multiple Myeloma)



An image from a peripheral blood smear demonstrating rouleaux, which is a characteristic finding of plasma cell myeloma. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3833ND55



An image from a peripheral blood smear demonstrating rouleaux, which is a characteristic finding of plasma cell myeloma. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3CZ32M1F



An image from a bone marrow direct smear in a patient with plasma cell myeloma demonstrating a plasma cell. Note the perinuclear clearing surrounding the nucleus, and basophilic cytoplasm. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3VM43C6D



An image from a bone marrow direct smear from a patient with plasma cell myeloma demonstrating two plasma cells (indicated by arrows). 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3348GX60



An image from a fluid cytospin slide showing a mott cell (indicated with an arrow) with round globules (Russell bodies). Mott cells may also be seen in plasma cell myeloma bone marrow smears. 60X oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3RV0DG30







An image from a fluid cytospin slide demonstrating a mott cell (indicated with an arrow) with round globules (Russell bodies). Mott cells may be seen in plasma cell myeloma bone marrow smears. 60X oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R31C1TX1V



An image from a bone marrow direct smear from a patient with plasma cell myeloma demonstrating a large plasma cell variant known as a flame cell (indicated by an arrow). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3CF9JN7P



An image from a bone marrow direct smear from a patient with plasma cell myeloma demonstrating a large plasma cell variant known as a flame cell (indicated by arrows). 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3416TF3X

Features:

This disorder is characterized by an increase in M protein (monoclonal gammopathy) in the serum and/or urine with the presence of clonal plasma cells in the bone marrow.¹

Unlike MGUS, patients with plasma cell myeloma often present with CRAB symptoms (hyperCalcemia, Renal failure, Anemia, and lytic Bone lesions). Additional features include osteolytic bone lesions without new bone formation.^{1,2} Some patients may be asymptomatic (smoldering multiple myeloma) but other findings of plasma cell myeloma may still be found.^{1,3}

Due to bone damage, extramedullary hematopoiesis is a common finding.¹

Cause:

Most cases of plasma cell myeloma have developed from MGUS.²

Other causes include infections, exposure to toxic substances, and other chronic diseases which may result in a long term antigenic stimulation.¹

Cytogenetic abnormalities have been found to be associated with the development of multiple myeloma.³

Age Group Affected: >50 years old, more common in males.¹

Plasma Cell Morphology:

Aside from the characteristic appearance of plasma cells, morphologic variants of plasma cells are often seen in Plasma Cell Myeloma. These include bi-lobed plasma cells, flame cells, and mott cells.

Flame Cells:

A reactive plasma cell that has reddish-purple cytoplasms. Colour of the cytoplasm is caused by glycoprotein and ribosomes. The presence of flame cells has been associated with IgA multiple myeloma.⁴

Mott Cells:

Are plasma cells with multiple round inclusions in the cytoplasm. The inclusions are termed "Russell bodies" and are composed of immunoglobulins.³

Laboratory Results for Plasma Cell Myeloma:^{1,2}

CBC:	PBS:	BM:
Cytopenias due to decreased bone marrow	Rouleaux formation (Due to M protein)	Lytic regions
hematopoiesis	Rare circulating plasma cells	Plasmacytosis (>10%)
RBC: Decreased	Normocytic	Bi-lobed plasma cells, Flame cells, and/or Mott cells may
HB: Decreased	Normochromic	be seen.
		Biopsy: interstitial clusters of plasma cells
		Aspirates: shows various heterogeneous forms of plasma
		cells in aggregates or sheets.



Immunologic Markers:	Other Tests:
CD38, CD79a, CD138	Serum FLC: Increased
	Bence-Jones proteinuria
	FISH
	Protein electrophoresis
	Immunofixation
	FLC assays
	Cytogenetics

Reactive Plasmacytosis:

An increase in the number plasma cells and immunoglobulins can also be the result of a non-malignant condition. Bacterial and viral infections (e.g infections mononucleosis, tuberculosis) can evoke a strong antigenic response and lead to an increase in plasma cells in the peripheral blood.⁴ The reactive process should NOT be confused with plasma cell myeloma.

Unlike plasma cell myeloma, there are no findings of CRAB symptoms or clonal plasma cells in the bone marrow. Plasma cells may be increased in the bone marrow, but not above 10%. An increase in M proteins and plasma cells (outside the bone marrow) may be found.³

References:

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CHAPTER OVERVIEW

14: White Blood Cells- Myeloproliferative Neoplasms (MPN)

14.1: Introduction to Myeloproliferative Neoplasms (MPNs)

- 14.2: Chronic Myelogenous Leukemia (CML)
- 14.3: Polycythemia Vera (PV)
- 14.4: Essential Thrombocythemia (ET)
- 14.5: Primary Myelofibrosis (PMF)

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14.1: Introduction to Myeloproliferative Neoplasms (MPNs)

MPNs are a group of clonal disorders that involve the proliferation and accumulation of one or more myeloid cell lines (erythrocytes, granulocytes, or platelets). These disorders are caused by genetic mutations in hematopoietic stem cells.^{1,2}

MPNs are commonly seen in middle age adults but some may occur during childhood.¹

The 2008 WHO classification system lists the following disorders under this category:¹

Chronic Myelogenous Leukemia (CML)

Polycythemia Vera (PV)

Essential Thrombocytopenia (ET)

Primary Myelofibrosis (PMF)

Note: WHO 2008 lists additional disorders under MPNs but only the ones listed above will be discussed in this eBook.

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

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14.2: Chronic Myelogenous Leukemia (CML)



An image from a peripheral blood smear demonstrating myeloid precursors and a few blasts seen in chronic myelogenous leukemia. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3NZ8155H



An image from a peripheral blood smear demonstrating a neutrophilia and myeloid precursors seen in CML. 500x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3SQ8QZ9V





An image from a bone marrow smear demonstrating myeloid hyperplasia and a few blasts seen in chronic myelogenous leukemia. Very few erythroid precursors are present. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R38P5VR48



An image from a bone marrow smear demonstrating myeloid hyperplasia in CML. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R34X54Z0H

Affected Cell Line: Myeloid cell line (platelets and granulocytes are increased, however granulocyte production is most prominent).^{1,2}

Mutation: Philadelphia Chromosome, t(9;22), resulting in the *BCR-ABL1* fusion gene.³ The gene is characterized by a translocation between chromosomes 9 and 22 which is expressed as t(9;22).¹

Age Group Affected: Seen commonly in middle aged adults, 46 to 53 years old.¹

Clinical Features:

The onset of CML is insidious and in some patients may be asymptomatic.³ Other patients may have complications associated with frequent infections, infiltration of leukocytes, bleeding, weight loss, fever, fatigue, and anemia.^{1,4}

Extramedullary hematopoiesis may occur, resulting in organomegaly.¹

The course of CML occurs in three phases:

1. Chronic Phase





Peripheral blood: leukocytosis (usually >100 x10⁹/L), thrombocytosis (Up to >1000 x10⁹/L).³

Bone Marrow: hypercellularity due to increased granulopoiesis. Megakaryocytes are increased and may appear small and hypolobulated.³

2. Accelerated Phase

The bone marrow is hypercellular and often myelodysplastic features are seen. Additionally, there is an increased number of myeloblasts, and a dropping platelet count resulting in thrombocytopenia. Total white blood cell count continues to increase.^{3,4}

3. Blast Phase (Acute Leukemia)

CML has transformed into an acute leukemia, either ALL or AML, and prognosis becomes poor even with treatment.⁴

Blast Phase is diagnosed when either: Bone marrow shows \geq 20% blasts or when extramedullary blast proliferation is present.^{3,4}

Laboratory Findings for CML:^{1,3,4}

CBC:	PBS:	BM:
RBC: Decreased	+/- NRBCs	M:E ratio: Increased
WBC: Increased (Average: >100 x10 ⁹ /L)	Neutrophilia, Eosinophilia, Basophilia	Hypercellular
PLT: Increased to Normal to Decreased (depending	+/- Micromegakaryocytes	Myeloid Hyperplasia
on phase)	Left shift	Megakaryocytes: Increased, may be dysplastic
Hb: Decreased	Dysplastic features may be present in granulocytes and platelets	Fibrosis in later stages
RETIC: Normal to Decreased		
Other Tests:		
LAP: Decreased		
Cytogenetics		
Hyperuricemia		
Uricosuria (May lead to gout)		
PLT Function: Abnormal		

Leukemoid Reaction

A leukemoid reaction is a response to different stress events such as infections, inflammation, hemorrhage, or other malignant disorders that has a similar presentation to CML where there is an increase in the number of white blood cells and a left shift.^{4,5}

A leukemoid reaction should NOT be confused with CML.

Table 1. Comparison of CML and Leukemoid Reaction.^{4,5}

Laboratory Finding	CML	Leukemoid Reaction
PBS	Left shift (Very Immature) Neutrophilia Eosinophilia Basophilia Dysplastic Features	Left shift (mild) Neutrophilia Absent eosinophilia Absent Basophilia Toxic changes are often present
WBC	20-500 x10 ⁹ /L	Rarely $>60 \text{ x}10^9/\text{L}$
PLT	Increased	Normal
Anemia	Present	Absent
LAP Score	Low	High
Chromosomal Abnormality	Philadelphia Chromosome/BCR-ABL1	None

References:

1. Randolph TR. Myeloproliferative neoplasms. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p.561-90.

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14.3: Polycythemia Vera (PV)



An image from a peripheral blood smear showing a thick smear with an abundant red blood cells and platelets often seen in polycythemia vera, 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3WH2DW3X

Affected Cell Line: Mainly erythrocytes, though tri-lineage growth ("panmyelosis") is seen in the bone marrow.^{1,2}

Mutation:

JAK 2 exon 12 and JAK 2 V617F gene mutations have been associated with PV.^{1,3}

Age Group Affected: Average age of diagnosis is 60 years old.³

Features:

PV involves the proliferation of erythrocytes independent of normal erythropoiesis regulating mechanisms (e.g. erythropoietin). There is also an a proliferation of granulocytes and megakaryocytes (resulting in "panmyelosis") but the proliferation of erythrocytes is most prominent.³⁻⁵

Unlike CML, PV does not readily transform into acute leukemia but may result in fibrosis over time.³ Splenomegaly is commonly seen.

Laboratory Findings for PV:^{2,4,5}

CBC: RBC: Increased	PBS: RBCs are normochromic/normocytic, though iron	BM: M:E ratio: Normal to decreased
WBC: Increased	stores may be exhausted, leading to hypo/micro RBCs	Hypercellular due to increased proliferation in all cell lines
Hb: Increased	PBS appears crowded with RBCs (thick smears are due to the elevated Hct)	exhibit lobulated nuclei
Hct: Increased MCV: Increased	A left shift and basophilia may be seen	Iron stores: decreased to absent
RETIC: Normal to increased		
Other Tests: Erythropoietin: Low or normal		

Secondary Polycythemia



Secondary polycythemia is a condition that occurs when there is an increased production of erythrocytes due an increased level of erythropoietin. Bone marrow shows an erythroid hyperplasia.⁵

Causes include: Hypoxia, Inappropriate use of erythropoietin, Familial polycythemia, neonatal polycythemia.⁵

Relative Polycythemia

Polycythemia that occurs due to a decrease in plasma volume, resulting in an elevated hematocrit, RBC count, and hemoglobin. There is no actual increased production of erythrocytes.⁵ The decrease in plasma volume is often the result of dehydration.

Hemoglobin and hematocrit appear increased but other CBC parameters such as white blood cell and platelet counts are normal. The bone marrow is also normal in terms of iron stores, cellularity and number of megakaryocytes.⁵

Table 1. Comparison between the different Polycythemias.⁵

-			
Laboratory Finding	PV	Secondary	Relative
Hct	Increased	Increased	Increased
НЬ	Increased	Increased	Increased
WBC	Increased	Normal	Normal
PLT	Increased	Normal	Normal
Erythropoietin	Decreased	Increased	Normal
Bone Marrow	Panmyelosis	Erythroid hyperplasia	Normal
Chromosomal Abnormalities	JAK 2 exon 12 and JAK 2 V617F	None	None

References:

1. Choi CW, Bang S-M, Jang S, Jung CW, Kim H-J, Kim HY, et al. Guidelines for the management of myeloproliferative neoplasms. Korean J Intern Med [Internet]. 2015 Nov 30 [cited 2018 Jul 9];30(6):771–88. Available from: http://kjim.org/journal/view.php?doi=10.3904/kjim.2015.30.6.771

2. Schaub CR. Chronic Myeloproliferative disorders I: chronic myelogenous leukemia. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 371-84.

3. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

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14.4: Essential Thrombocythemia (ET)

An interactive or media element has been excluded from this version of the text. You can view it online here:

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Image of peripheral blood smears showing a giant platelet (center) and an increase in the number of platelets seen in Essential Thrombocythemia. From MLS Collection, University of Alberta.

Image 1: 50x oil immersion. https://doi.org/10.7939/R3J09WK51

Image 2: 60x oil immersion. https://doi.org/10.7939/R3X63BM52

Affected Cell Line: Megakaryocytes, Platelets.^{1,2}

Mutation: JAK 2 V617F, CALR, and MPL gene mutations. 1,3

Age Group Affected: Diagnosed most commonly at 50-60 years old.¹

Clinical Features:

Most patients are asymptomatic and present with a platelet count of \geq 450 x10⁹/L.¹ Thrombosis, vascular occlusion, and bleeding problems are the most commonly associated complications. Despite having abundant platelets, they are often dysfunctional.

Laboratory Findings for ET:^{1,4,5}

CBC: WBC: Normal to Slightly Increased PLT: Increased (often 1000-5000 x10 ⁹ /L) Hb: Slightly Decreased Hct: Slightly Decreased MCV: Normal	PBS: RBCs, are normocytic and normochromic Marked thrombocytosis Abnormal platelet morphologies: Giant, agranular, clumping, and irregularly shaped	BM: Hypercellular due to increased megakaryopoiesis Abnormal megakaryocyte morphology: Clusters, Enlarged, Hyperlobulated
Other Tests: Platelet function tests: abnormal		

Reactive Thrombocytosis

A non-malignant condition that involves an increased platelet count secondary to other conditions that result in an increase in platelet production. It is associated with infections and inflammatory processes.⁵

Reactive thrombocytosis can be differentiated from essential thrombocythemia by looking at the platelet count. Platelet count rarely reaches >1000 x10⁹/L and platelet function tests are normal.⁵

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

2. Schaub CR. Chronic Myeloproliferative disorders I: chronic myelogenous leukemia. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 371-84.

3. Choi CW, Bang S-M, Jang S, Jung CW, Kim H-J, Kim HY, et al. Guidelines for the management of myeloproliferative neoplasms. Korean J Intern Med [Internet]. 2015 Nov 30 [cited 2018 Jul 9];30(6):771–88. Available from: http://kjim.org/journal/view.php?doi=10.3904/kjim.2015.30.6.771

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14.5: Primary Myelofibrosis (PMF)



An image from a peripheral blood smear showing tears, elliptocytes, schistocytes, and a giant platelet seen in primary myelofibrosis. 50x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3CJ88201



An image from a peripheral blood smear showing tears, elliptocytes, and schistocytes, and a nucleated red blood cell seen in primary myelofibrosis. 100x oil immersion. From MLS Collection, University of Alberta, https://doi.org/10.7939/R3416TF2F

Affected Cell Line: Granulocytes and Megakaryocytes in the bone marrow resulting in secondary fibroblast stimulation and fibrotic desposition in the bone marrow.^{1,2} Mutation: JAK 2 V617F, CALR, and *MPL* gene mutations.^{1,3}





Age Group Affected: >50 years old, occurs equally between males and females.⁴

Features:

Primary myelofibrosis is characterized by two stages:

1. Prefibrotic Stage

The bone marrow is hypercellular and shows minimal reticulin and fibrosis initially, with an increase in megakaryocytes and granulocytes.¹

2. Fibrotic Stage

Peripheral blood shows a characteristic **leukoerythroblastic picture** (immature granulocyte and erythrocyte precursors) with poikilocytosis, especially teardrop cells and elliptocytes.¹

Bone marrows shows marked fibrosis.¹ Extramedullary hematopoiesis is often seen, with cells accumulating in the spleen, liver, and other organs.

Laboratory Findings for Primary Myelofibrosis:^{2,4,5}

CBC:	PBS:	BM:
Early Stage:	Platelets have a dysplastic morphology (Giant,	Often results in a dry tap
RBC: Normal	agranular)	Hypercellular
WBC: Increased	May see micromegakaryocytes	Fibrosis of varying degrees (Marked fibrosis in later stages)
PLT: Increased	Variable poikilocytosis	Megakaryocyte aggregates
Hb: Normal	Fibrotic Stage:	Dysgranulopoiesis
Fibrotic Stage:	Pancytopenia	Dysmegakaryopoiesis
RBC: Decreased	Leukoerythroblastic picture	
WBC:Decreased	Teardrop cells	
PLT: Decreased	Elliptocytes	
Hb: Decreased	nRBCs	
Other Tests:		
PLT Function: Abnormal		

References:

1. Swerdlow SH, Campo E, Harris NL, Jaffe ES, Pileri SA, Stein H, et al. editors. WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues Volume 2. 4th ed. International Agency for Research on Cancer (IARC); 2008.

2. Schaub CR. Chronic Myeloproliferative disorders I: chronic myelogenous leukemia. In: Clinical hematology and fundamentals of hemostasis. 5th ed. Philadelphia: F.A. Davis Company; 2009. p. 371-84.

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4. Randolph TR. Myeloproliferative neoplasms. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 450-78.

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CHAPTER OVERVIEW

15: White Blood Cells- Myelodysplastic Syndromes (MDS)

15.1: Introduction to Myelodysplastic Syndromes (MDS)

15.2: MDS- Dyserythropoiesis, Dysmyelopoiesis and Dysmegakaryopoiesis

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15.1: Introduction to Myelodysplastic Syndromes (MDS)

Myelodysplastic syndromes are a group of clonal disorders that result in cytopenias and defective cell maturation.¹ Morphology of cells during maturation show abnormalities, referred to as dysplasia.² Dysplastic features seen varies in terms of the types of dysplasia seen, and the cell lines affected. Progression toward acute leukemia is often seen later in the disease, leading to an increase in blasts seen in the bone marrow and peripheral blood.

Age Group Affected: Commonly age of diagnosis is 70 years old.¹

Affected Cell Line(s): Can affect one, two, or all three hematopoietic cell lines (erythroid, myeloid, megakaryocyte).¹

Cause(s):

Chromosomal abnormalities

Mutations in oncogenes and tumor suppressor genes

General Laboratory Findings for MDS:²

PBS:	BM:	Other Tests:
Anemia	Dysplastic hematopoietic precursors	Cytogenetic testing
Anisocytosis (Dimorphic)	Usually hypercellular (though hematopoiesis is	FISH
Poikilocytosis	ineffective)	Iron Studies:
Sideroblasts	+/- Increased blasts	Serum Iron: Normal to Increased
Dysplastic granulocytes		Serum Ferritin: Normal to Increased
Dysplastic thrombocytes		TIBC: Decreased to Normal
+/- Increased blasts		

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1. Rodak BF. Myelodysplastic syndromes. In: Rodak's hematology clinical applications and principles. 5th ed. St. Louis, Missouri: Saunders; 2015. p.591-603.

2. Lawrence LW, Taylor SA. Myelodysplastic syndromes. In: Clinical laboratory hematology. 3rd ed. New Jersey: Pearson; 2015. p. 479-99.

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15.2: MDS- Dyserythropoiesis, Dysmyelopoiesis and Dysmegakaryopoiesis

As previously discussed, MDS is a clonal disorder that results in defective cell maturation and results in dysplastic changes. The dysplasia can be seen in both the peripheral blood and in the bone marrow. Dysplasia may be seen in one or more cell lines, and the types of dysplasia seen vary. Below are descriptions that may be seen, organized by cell lineage.

Dyserythropoiesis

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Glossary

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Sample Word 1 | Sample Definition 1



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