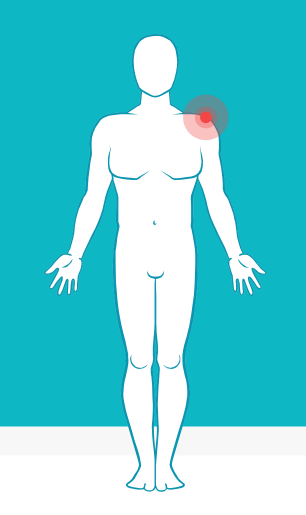
Hematopoietic & Lymphoid System White Cell disorders

Ghadeer Hayel, M.D. Assistant professor of Pathology Mutah University Consultant hematopathologist 4/8/2025



2. Neoplastic Proliferations of White Cells

- Myeloid Neoplasms

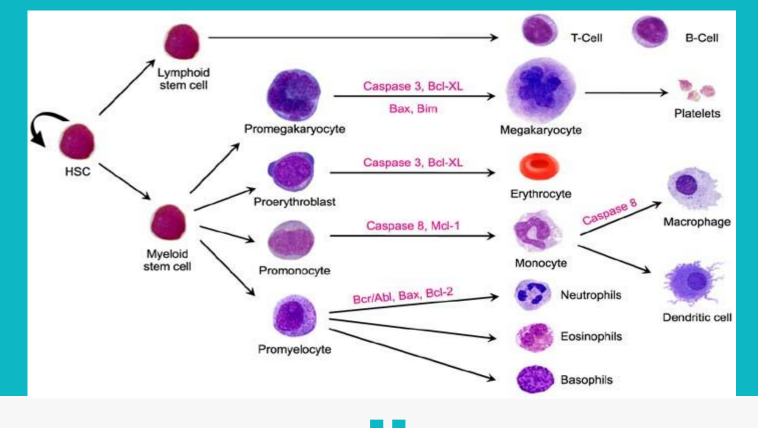
Myeloid Neoplasms

- Neoplasms originated from hematopoietic progenitors.
- Primarily involve the bone marrow & replace normal marrow elements.
- Lesser secondary Hematopoietic organs involvement (LN, spleen & liver).

Myeloid Neoplasms

Three broad categories of myeloid neoplasia:

- Myeloproliferative neoplasms (MPN): neoplastic clone continues to terminal differentiation but with increased or dysregulated growth.



Acute myeloid leukemia (AML)

Acute myeloid leukemia (AML)

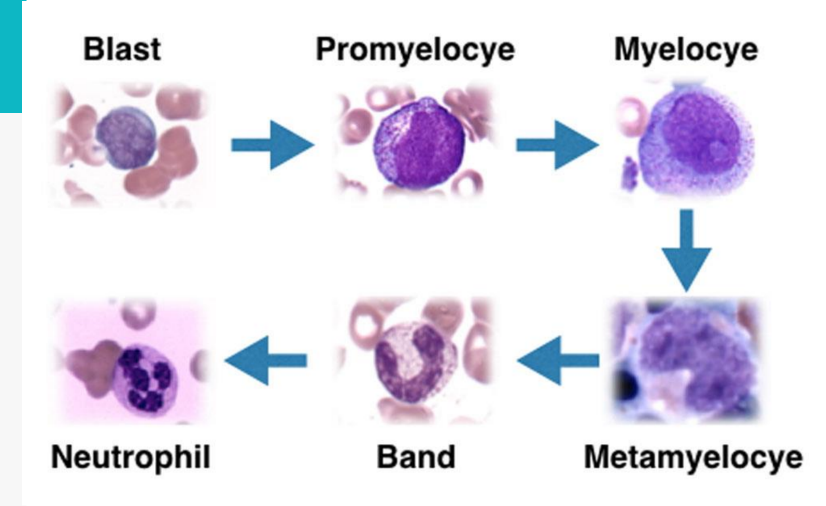
- Affects all age group, peak > 60 years.
- Clinical signs & symptoms; result from the replacement of normal marrow elements by leukemic blasts; symptoms related to anemia, thrombocytopenia, & neutropenia.
- Acute: present within a few weeks of the onset of symptoms.
- Splenomegaly & lymphadenopathy are less prominent than in ALL (Acute Lymphoblastic leukemia)

Acute myeloid leukemia (AML) – Risk factors

- Increase age.
- Male sex
- Previous cancer treatment.
- Exposure to radiation. (e.g., survivors of a nuclear reactor accident).
- Dangerous chemical exposure. (e.g., benzene)
- Smoking; AML is linked to cigarette smoke (contains benzene & other chemicals)
- Other blood disorders (MDS, MPN)
- Genetic disorders. (e.g., Down syndrome)

Acute myeloid leukemia (AML) - Pathogenesis

- Most AMLs harbor mutations in genes encoding transcription factors that are required for normal myeloid cell differentiation → interfere with the differentiation of early myeloid cells → accumulation of myeloid precursors (blasts) in BM.
- Examples: t(15;17) in acute promyelocytic Leukemia (APL) → fusion of retinoic acid receptor α (RARA) gene on chr. 17 & PML gene on chr. 15 → PML/RARA fusion protein → blocks myeloid differentiation at promyelocytic stage.



Acute myeloid leukemia (AML) - Pathogenesis

- ► Treatment with all-trans retinoic acid (ATRA), an analogue of vitamin A, overcomes this block → induce the neoplastic promyelocytes to differentiate into neutrophils rapidly → clears the tumor.
- The effect is very specific; AMLs without t(15;17) don't respond to ATRA.
- This is an important example of a highly effective therapy targeted at a tumor-specific molecular defect.
- ► t(15;17) AML have the best prognosis of any type → curable in > 90%

Acute myeloid leukemia (AML) – Classification

- AMLs are very diverse in terms of genetics, cellular lineage, and degree of maturation.
- WHO classification relies on all of these features to divide AML into four categories:
- (1) AMLs ass with specific genetic aberrations: important coz they predict outcome & they guide therapy.
- (2) AMLs with dysplasia: arise from MDSs.

- (3) AMLs occurring after genotoxic chemotherapy.
- (4) AMLs, Not otherwise specified: subclassified based on the predominant line of differentiation

9, 2012.

TABLE 1. WHO classifications for AML subtypes

Туре	Name
MO	Minimally differentiated acute myeloblastic leukernia
M1	Acute myeloblastic leukemia (t(8;21)(q22,q22))
M2	Acute myeloblastic leukemia (t(6,9))
МЗ	Acute promyelocytic leukemia (APL)
M4	Acute myelomonocytic leukemia
M4eo	Myelomonocytic leukemia with bone marrow eosinophilia
M5	 Acute monoblastic leukemia (M5a) Acute monocytic leukemia (M5b)
Mб	Acute erythroid leukemias, including —Erythroleukemia (M6a) —Very rare pure erythroid leukemia (M6b)
M7	Acute megakaryoblastic leukemia
M8	Acute basophilic leukemia
Source: Acu	cute myeloid leukemia; t, translocation; WHO, World Health Organization. te myeloid leukemia classification. News-Medical.net Web site. http://www. cal.net/health/Acute-Myeloid-Leukemia-Classification.aspx. Accessed March

Table 12.11 WHO Classification of AML

Class	Prognosis		
I. AML With Recurrent Chromosomal Translocations			
AML with t(8;21)(q22;q22); RUNXT1/RUNX1 fusion gene	Favorable ⁻		
AML with inv(16)(p13;q22); CBFB/MYH11 fusion gene	Favorable		
AML with t(15;17)(q22;q21.1); PML/RARA fusion gene	Favorable		
AML with t(11q23;variant); <i>MLL</i> fusion genes	Poor		
AML with mutated NPM1	Variable		
II. AML With Multilineage Dysplasia			
With previous MDS	Very poor		
Without previous MDS	Poor		
III. AML, Therapy-Related			
Alkylating agent–related	Very poor		
Epipodophyllotoxin-related	Very poor		
IV. AML, Not Otherwise Classified			
Subclasses defined by extent and type of differentiation (e.g., myelocytic, monocytic)	Intermediate .		

Acute myeloid Leukemia

History				
Chemotherapy ± → Radiotherapy	Myeloid neoplasm post cytotoxic therapy (e.g. AML with <i>KMT2A::MLLT3</i> fusion post cytotoxic therapy)			
	AML with defining genetic abnormalities			
	Acute promyelocytic leukemia with PML::RARA fusion			
	AML with RUNX1::RUNX1T1 fusion			
	AML with CBFB::MYH11 fusion			
	AML with DEK::NUP214 fusion			
	AML with RBM15::MRTFA fusion			
	AML with BCR::ABL1 fusion			
	AML with KMT2A rearrangement			
	AML with MECOM rearrangement			
	AML with NUP98 rearrangement			
	AML with NPM1 mutation	AML with RUNX1T3::GLIS2 fusion		
	AML with CEBPA mutation	AML with KAT6A::CREBBP fusion		
		AML with FUS::ERG fusion		
MDS or MDS/MPN —	AML, myelodysplasia-related	AML with MNX1::ETV6 fusion		
	ANN with allowed fined and the ellowed and	AML with NPM1::MLF1 fusion		
	AML with other defined genetic alterations			
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	AML defined by differentiation			
	AML with minimal differentiation			
	AML without maturation			
	AML with maturation			
	Acute basophilic leukemia			

Acute myelomonocytic leukemia

Acute monocytic leukemia

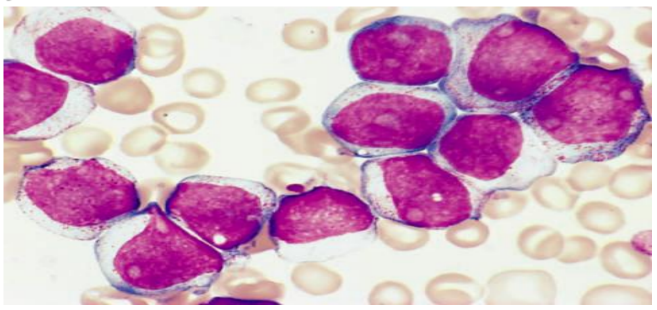
Acute erythroid leukemia*

Acute megakaryoblastic leukemia

*the only type in this family that supersedes AML-MR

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▶ By definition → AML: the presence of at least 20% myeloid blasts or promyelocytes of BM cellularity.



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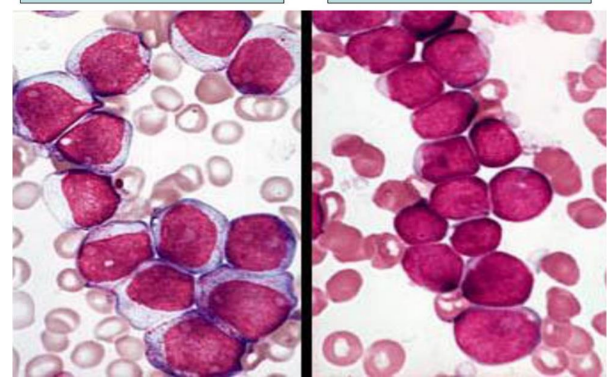


Myeloblasts: have delicate nuclear chromatin, 2-4 nucleoli, larger cytoplasm than lymphoblasts & fine azurophilic cytoplasmic granules.

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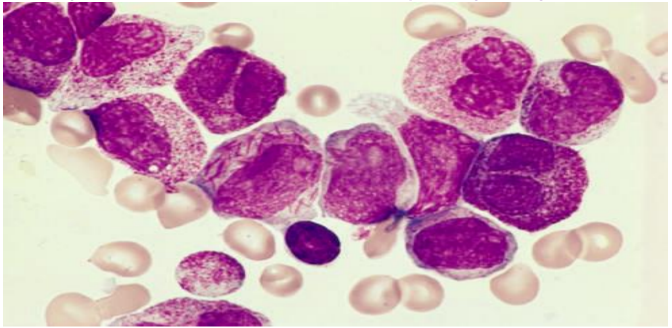
MYELOBLASTS

LYMPHOBLASTS



Auer rods: distinctive red-staining needle-like azurophilic granules, present in many cases. Numerous in acute promyelocytic leukemia

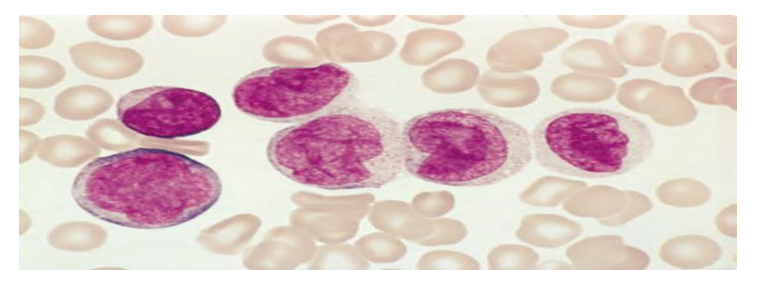
(APL).



- In other subtypes of AML, monoblasts, erythroblasts, or megakaryoblasts predominate.
- Occasionally, blasts are entirely absent from PB (aleukemic leukemia).
- For this reason, BM examination is essential to exclude acute leukemia in pancytopenic patients.

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Monoblasts: have folded or lobulated nuclei, lack Auer rods.



Acute myeloid leukemia (AML) – Immunophenotype

- Immunologic markers are heterogeneous in AML.
- Most tumors express some combination of myeloidassociated antigens; CD13, CD14, CD15, or CD117 (KIT).
- CD34: a marker of hematopoietic stem cells & often present on myeloblasts.
- Myeloperoxidase (MPO), most specific.

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Such markers are helpful in distinguishing AML from ALL and in identifying AMLs with only minimal differentiation.

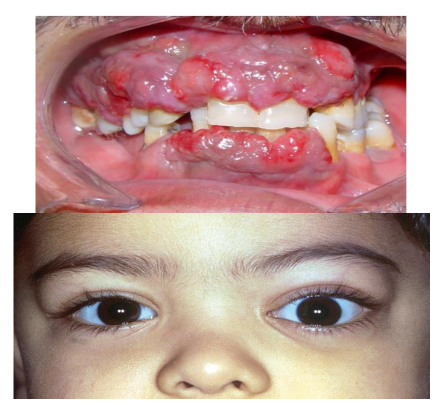
Acute myeloid leukemia (AML) - Clinical features

- Patients present within weeks or a few months of the onset of symptoms.
- Symptoms of anemia, neutropenia, & thrombocytopenia, (fatigue, fever, and spontaneous mucosal & cutaneous bleeding).
- CNS manifestations are less frequent than ALL.
- Procoagulants and fibrinolytic factors released by leukemic cells, especially in AML with the t(15;17) -> high DIC incidence.

Acute myeloid leukemia (AML) – Clinical features

Tumors with monocytic differentiation often infiltrate the skin (leukemia cutis) & the gingiva.

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Acute myeloid leukemia (AML) - Prognosis

AML remains a devastating disease.

- Tumors with "good-risk" karyotypic abnormalities (t[8;21], inv[16]) are associated with a 50% chance of longterm disease-free survival.
- Overall survival in all patients is only 15-30% with conventional chemotherapy.

24 Acute vs Chronic leukemia

Acute leukemia

- Blasts
- Rapid proliferation of cells.
- Rapidly Fatal (<6 months without Tx)
- Lymphoid..ALL
- Myeloid ... AML

Chronic leukemia

- Mature cells
- Gradual proliferation.
- More indolent disease. (2-6 years without Tx)
- ▶ Lymphoid ... CLL
- ▶ MPN...CML



