

Leukemia for the Primary Care Physician (Non-CLL)

Karilyn Larkin, MD Assistant Professor of Medicine Division of Hematology The Ohio State University Wexner Medical Center

MedNet21

THE OHIO STATE UNIVERS

OBJECTIVES:

- Presenting symptoms
- · Context and basic biology
- Diagnostics
- Prognostics
- Therapeutics



CASE 1

- Your longtime patient Mr. Smith, a 57-year-old man who works on his farm, presents with progressive fatigue and dyspnea on exertion over the last two weeks.
- He has shortness of breath with minimal activity and chest pain with climbing stairs, one month ago he was carrying 50lbs without any difficulty. He notes a headache that has been constant for the past day.
- · He is able to sleep while laying flat on one pillow



CASE 1, CONTINUED

PMH

HTN, HLD, Rheumatoid Arthritis

SH

20pk/year smoking history, 1-2 drinks a few times a week, he lives on his farm with his wife and they have a large dog and 2 cats, 3 children whom are grown

FH:

CAD, prostate CA

Meds: lisinopril, simvastatin, methotrexate



CASE 1, CONTINUED

- Physical examination
- Pale but not ill-appearing, with rapid heart rate; not short of breath at rest
- Hypertrophied gums with areas of bleeding
- No pitting edema, lungs are clear
- No lymphadenopathy
- +Ecchymoses on arms and legs



CASE 1, CONTINUED

- Complete blood count
- WBC count: 55,000 cells/µL
- Hemoglobin: 6 g/dL
- Platelet count: 15,000 cells/µL
- CMP
 - Creatinine 1.5 (baseline 1.1), otherwise WNL
 - AST/ALT minimally elevated



WHAT DOES THE HEMATOLOGIST WANT TO KNOW?

- What are his coags? PT/PTT/INR AND Fibrinogen
- Uric Acid
- · Has he had any fevers?
- Any headaches? Vision changes? Difficulty breathing or hypoxemia? Any chest pain?

WHAT ARE THE IMMEDIATE NEXT STEPS?

Report to the closest ER

(acute leukemia treating center if possible) • These are the patients that keep me up all night

ACUTE LEUKEMIA PRESENTING SYMPTOMS

- Cytopenias
- Hyperleukocytosis → leukostasis
- Extramedullary disease
- Tumor lysis syndrome
- Disseminated Intravascular Coagulation



CASE 2

- Your longtime patient Mr. Habib, a 57-year-old man who works on his farm, presents with progressive fatigue and early satiety over the past several months.
- He denies any shortness of breath with minimal activity but notes some discomfort with deep inspiration and frequent sharp pains on his left side.
- He has been sleeping well and doesn't understand why he's feeling so fatigued



CASE 2, CONTINUED

- PMH: HTN, HLD
- SH: 20pk/year smoking history, 1-2 drinks a few times a week, he lives on his farm with his wife and they have a small dog and 2 hamsters, 2 children whom are grown
- FH:CAD, prostate CA
- · Meds: lisinopril, simvastatin



CASE 2, CONTINUED

- Physical examination
- Appears well, non-toxic, normal vitals
- Cardiac exam unremarkable
- No pitting edema, lungs are clear
- No lymphadenopathy, but spleen is palpable 4 cm below left costal margin
- No rashes or bruises



CASE 2, CONTINUED

- Complete blood count
- WBC count: 55,000 cells/µL
- Hemoglobin: 10 g/dL
- Platelet count: 325,000 cells/µL
- CMP
 - Creatinine 1.2 (baseline 1.1), otherwise WNL
 - AST/ALT normal



WHAT DOES THE HEMATOLOGIST WANT TO KNOW?

- What does his PB smear look like?
- Uric acid
- When was his last CBC and what did it look like?

WHAT ARE THE IMMEDIATE NEXT STEPS? Look at PB smear



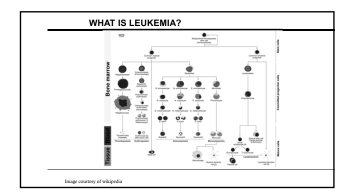
Send BCR/ABL

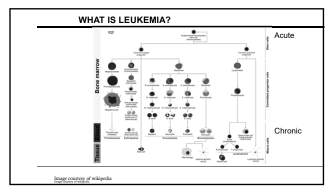
Can follow up in clinic

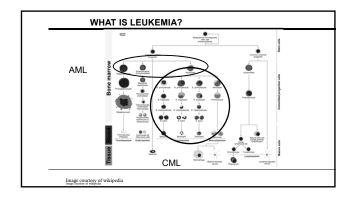
CML PRESENTING SYMPTOMS

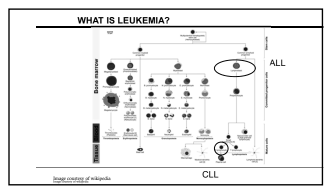
- Mild Anemia, thrombocytosis
- Hyperleukocytosis
- Splenomegaly, +/- infarcts
- Hyperuricemia

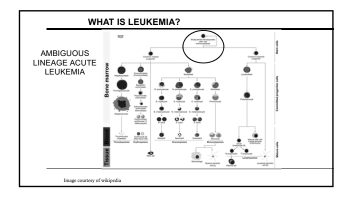


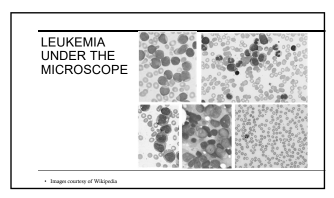












TESTING: BONE MARROW BIOPSIES

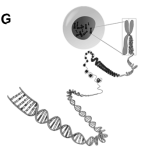
- Morphology
- Flow cytometry (immunophenotype)
- Cytogenetics
- Molecular genetics



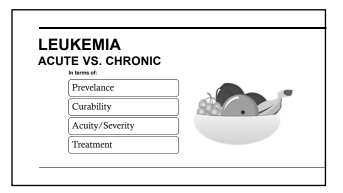
By Photographer's Mate 2nd Class Chad McNeeley - Navy News Service, 021204-N-0696M-180, Public Domain, https://commons.wikimedia.org/w/index.php?curid=1337397

GENETIC TESTING

- Karyotyping
- •Fluorescence in situ hybridization (FISH)
- Molecular genetics (next generation sequencing, NGS)



OTHER USEFUL TESTS/PROCEDURES Testing AML ALL **CML** Lumbar If high Multiple puncture suspicion PET or CT Myeloid Presenting with LAD or sarcoma scans masses Tuneled line Yes Yes No



EPIDEMIOLOGY

AML

CML

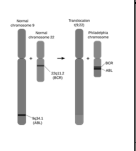
- ~20,000 new cases yearly in US
- yearly in US
- ~ 6,600 new cases ~8,800 new cases yearly in US
- > 11,000 deaths/year
- > 1,500 deaths/year
 > 1,200 deaths/year
- 5yr survival 30.5%
- Median age: 68 years Median age: 17 years Median age: 65 years ■ 5 yr survival 70.8% ■ 5yr survival 70.4%

SEER data. Cancer.gov

CHRONIC MYELOGENOUS LEUKEMIA

DIAGNOSING CML

- · Clinical History
- Physical Exam
- Labs
- BCR-ABL
- BM Biopsy



SYMPTOMS

Up to 50% of patients asymptomatic

46-76% p/w splenomegaly

Fatigue, night sweats

Symptoms of anemia, bleeding d/t platelet dysfunction

<5% p/w hyperviscosity symptoms (usually WBC >250,000)

CBC AND PERIPHERAL SMEAR

Absolute leukocytosis (median 100,000) Left shift

Myelocytes outnumber mature metamyelocytes on PB smear
Blasts usually <2%

Absolute basophilia (100%)

Absolute eosinophilia (90%)

Platelet count usually normal or elevated

Thrombocytopenia= alternative dx OR advanced stage CML

CML PHASES

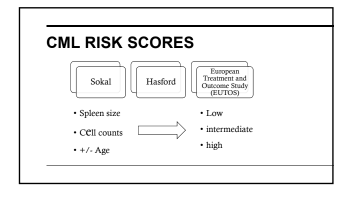
Chronic

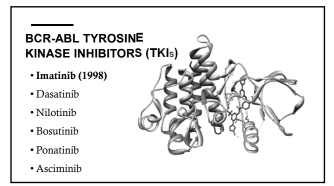
• most patients present in early phase Accelerated

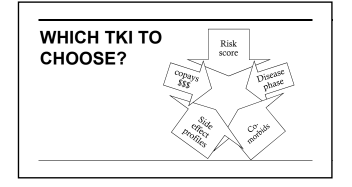
 more aggressive disease, less likely to respond as well to therapy
 most commonly

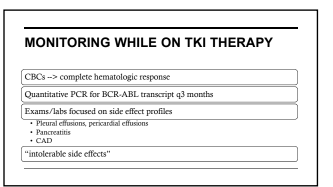
• most commonly seen after treatment failure Blast

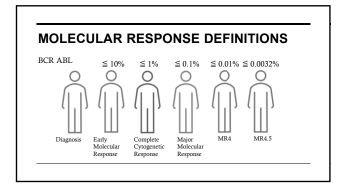
AML or ALL

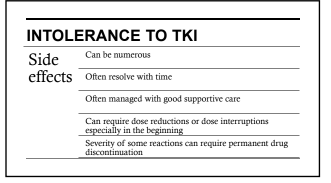








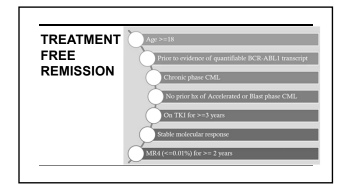




LOSS OF RESPONSE TO TKI

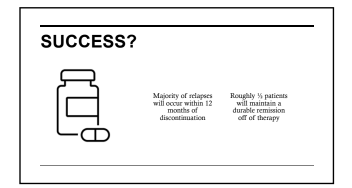
- $\bullet \ Adherence$
- Adherence
- Adherence
- Taking correctly (PPIs, food)
- Check TKI resistance panel
 - BCR-ABL kinase domain mutational analysis





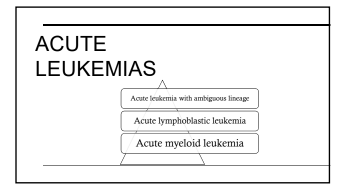
MONITORING AFTER DISCONTINUATION

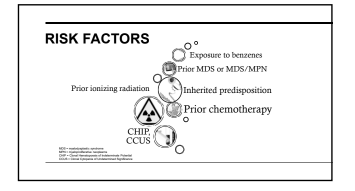
- •Enhanced Monitoring off drug Can continue to hold TKI as long as maintain
 - MMR (<=0.1%)
 - First 6 months monthly
 - Second 6 months decrease to Q2 months
 - Forevermore Q3 months

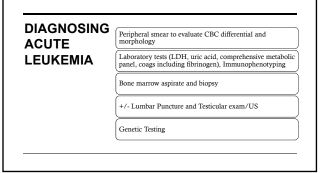


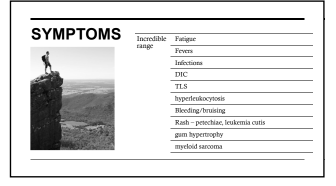
CML SUMMARY

- · CBC and peripheral smear are very helpful in distinguishing
 - · *Peripheral basophilia
- · PB looks like BM
- \bullet Diagnosis from PB t(9;22) and BM Biopsy establishes stage
- Multiple TKI treatment options- depends on disease factors/risk score, patient factors
- Can now consider discontinuing TKI with very close monitoring and follow up

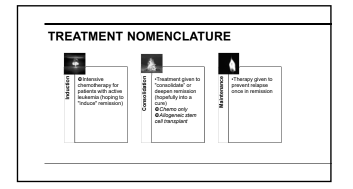








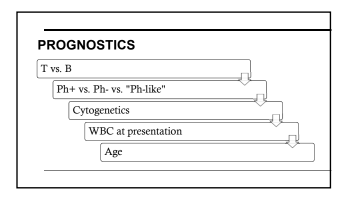
Profound cytopenias	Neutropenia
	Leukocytosis (predominantly blasts)
	possibly dysplastic neutrophils
	Anemia without schistocytes or other abnormal indices
	thrombocytopenia, no clumping

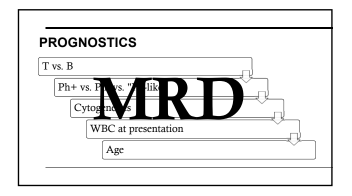


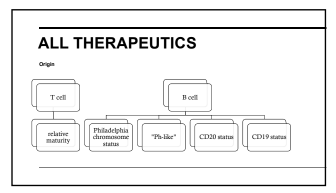
RESPONSE NOMENCLATURE

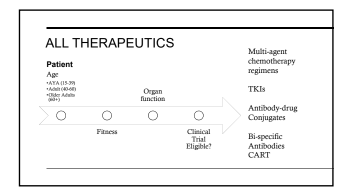
- Complete Response (CR)
- Complete Response with incomplete count recovery (CRi)
- Morphologic leukemia free state (MLFS)
- Remission ≉ Cure
- Minimal (Measurable) Residual Disease (MRD)

ACUTE LYMPHOBLASTIC LEUKEMIA

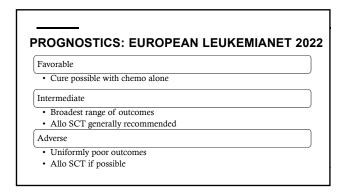


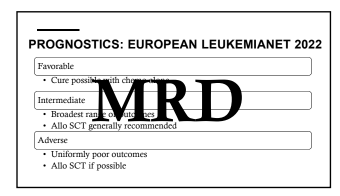


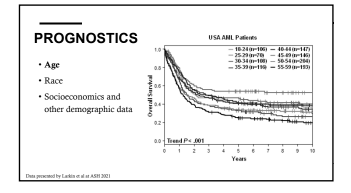


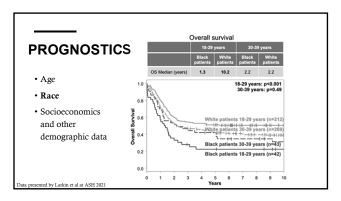


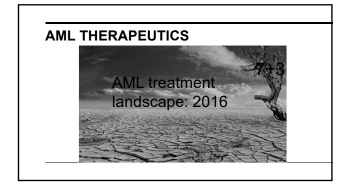
ACUTE MYELOID LEUKEMIA

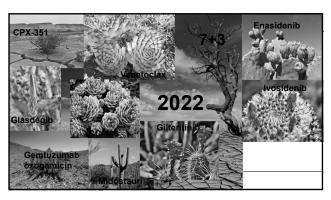


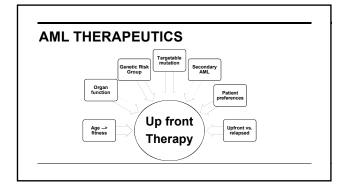


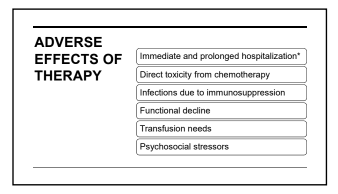












AL SUMMARY

- Onset is typically rapid
- Key historical items can help raise your suspicion in some cases
- \bullet CBC and peripheral smear are very helpful in identifying this urgent/emergent disease
- Diagnosis requires multiple specialized tests
- Prognosis depend on multiple factors
- Treatment options are personalized

HIGH YIELD POINTS

How do you recognize leukemia?

- Patient presentations vary and sometimes require high degree of clinical suspicion
 CBC is very often enough obvious to direct further work-up

CML on TKIs

- Characteristic and non-characteristic side effects
 Adherence is key
 There is hope for treatment free remissions albeit in a minority of patients

Acute Leukemia is a rapidly changing field

- Diagnostics have become more complicated but improved
 Many more tolerable treatment options

