Oncology Clinical Pathways Acute Myeloid Leukemia (AML)

April 2025 - V1.2025







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Acute Myeloid Leukemia – Presumptive Conditions

VA automatically presumes that certain disabilities were caused by military service. This is because of the unique circumstances of a specific Veteran's military service. If a presumed condition is diagnosed in a Veteran within a certain group, they can be awarded disability compensation.

Atomic Veterans – Exposure to Ionizing Radiation

• All forms of leukemia

Gulf War and Post 9/11 Veterans

If the patient served on or after Sept. 11, 2001, in Afghanistan, Djibouti, Egypt, Jordan, Lebanon, Syria, Uzbekistan, or Yemen or if the patient served in the *Southwest Asia theater of operations, or Somalia, on or after Aug. 2, 1990, specific conditions include:

• Acute leukemias

*The Southwest Asia theater of operations refers to Iraq, Kuwait, Saudi Arabia, the neutral zone between Iraq and Saudi Arabia, Bahrain, Qatar, the United Arab Emirates, Oman, the Gulf of Aden, the Gulf of Oman, the Persian Gulf, the Arabian Sea, the Red Sea, and the airspace above these locations.

For more information, please visit <u>U.S. Department of Veterans Affairs - Presumptive Disability Benefits (va.gov); VA makes</u> several cancers presumptive for service connection Jan 08 2025; <u>eCFR :: 38 CFR 3.320b -- Presumptive service connection for</u> leukemias, multiple myelomas, myelodysplastic syndromes, and myelofibrosis.







Acute Myeloid Leukemia – First Line



* Diagnosis must include flow cytometry, karyotype, rapid order (<72 hours) molecular tests (to include: FLT3, NPM1, IDH1, and IDH2), and myeloid NGS test (at minimum must include: ASXL1, BCOR, CEBPA, EZH2, FLT3, IDH1, IDH2, NPM1, RUNX1, SF3B1, SRSF2, STAG2, TP53, U2AF1, and ZRSR2); additional optional genes include: CBL, DDX41, KIT, KR5S, NRAS, and other genes associated with myeloid neoplasms; AML FISH testing can also be performed, either up front, or at the discretion of the pathologist (can include: -5/59, -7/70, KMT2A, t(8,21) RUNX1:-RUNX111, t(15, 17) PML::RARA, t(16:16) or inv(16) CBFB::MYH11, t(9;22) BCR::ABL1, and TP53)

Supportive Care includes transfusions with leukocyte depleted/irradiated units for patients who are transplant candidates; platelet transfusion for platelet <u><</u> 10 per 10,000/mm³, pRBC transfusion for Hgb < 7 g/dL, cryoprecipitate for fibrinogen < 150 mg/dL; tumor lysis syndrome monitoring, allopurinol and IV fluid prophylaxis, and rasburicase treatment if needed for patients with high WBC, hyperuricemia, and/or renal dysfunction; infection prophylaxis is recommended e.g., fungal, HSV/VZV, and bacterial

^c Candidate for Intensive Therapy assess by age, performance status, comorbidities, and social factors; useful tool is the Fred Hutch Treatment Related Mortality Calculator; echocardiogram is required if considering intensive therapy; candidates for intensive therapy assumes that the patient is a transplant candidate early HLA typing recommended

^d Risk Group Classification determined via guidelines such as European LeukemiaNet (ELN) or National Comprehensive Care Network (NCCN)

Anthracycline either daunorubicin or idarubicin is appropriate

Persistent Disease second induction may be appropriate based on depth of response and regimen used

PHDAC "High Dose" Cytarabine Consolidation; dosing schedule may be on days 1-3 or days 1, 3, and 5; monitoring for neurologic (cerebellar) toxicity required; supportive care with steroid eye drops required

^b FLT3 Status mutation defined as point mutation in the TKD or ITD mutation.

Quizartinib has a boxed warning for risk of QT prolongation, Torsades de pointes, and cardiac arrest; perform EKG at baseline, weekly during induction and consolidation phases and weekly for at least the first month of maintenance, then periodically; refer to prescribing information for EKG monitoring nmendations; monitor serum electrolytes (magnesium, potassium) at baseline and as clinically indicated

Other Groups includes AML secondary to myeloproliferative neoplasms as well as unfavorable molecular features

Venetodiax has mary drug-drug interactions: consultation with nonclogy pharmacits is recommended, anti-infection prophylasis recommended particularly when the patient has neutropenia, e.g., fungal, HSV/VZV, and bacteriak dose modifications (duration, dose, frequency) of venetodax are frequently needed bases and follow response are needed with the continuous therapy in block del country, regular bote many broines to assess and follow response are needed with the continuous therapy.

Bone Marrow Biopsy after decitabine and venetoclax variability in response times

ⁿ Adequate Response defined as at minimum a partial remission

AML Acute Myeloid Leukemia

ITD Internal Tandem Duplication TKD Tyrosine Kinase Domain







Acute Myeloid Leukemia – Relapsed



Every Step of the Way

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Acute Myeloid Leukemia – Molecular Testing Table

Eligibility	Test Category	Test Type	Recommended Vendors	NPOP Coverage	Specimen Type		
Acute Myeloid Leukemia (AML)	Flow cytometry	Leukemia/lymphoma panel	Local VA or locally contracted vendor	No	Bone Marrow Biopsy, Blood		
	FISH	-5/-5q, -7/-7q, KMT2A, t(8;21) RUNX1::RUNX1T1, t(15;17) PML::RARA, t(16;16) or inv(16) CBFB::MYH11; t(9;22) BCR::ABL1; TP53	Local VA or locally contracted vendor	No	Bone Marrow Biopsy, Blood		
	Karyotyping	Karyotype	Local VA or locally contracted vendor	No	Bone Marrow Biopsy, Blood		
	Rapid Molecular Tests (<1 week TAT)	FLT3 ITD and TKD, IDH1/2, NPM1 (quantitative preferred), CEBPA (optional)	Local VA or locally contracted vendor	No	Bone Marrow Biopsy, Blood		
	Somatic NGS	NGS panel ASXL1, BCOR, CEBPA, EZH2, FLT3, IDH1, IDH2, NPM1, RUNX1, SF3B1, SRSF2, STAG2, TP53, U2AF1, and ZRSR2. Optional genes include: CBL, DDX41, KIT, KRAS, NRAS, and other genes associated with myeloid neoplasms.	GLA	GLA Grant*	Bone Marrow Biopsy, Blood		
	Somatic NGS	Consider CGP if no driver mutation detected	Foundation Medicine	Yes	Bone Marrow Biopsy, Blood		
	Germline NGS	Consider germline testing and genetic counseling if VAF >40% in AML predisposition genes (CEBPA, DDX41, RUNX1, ANKRD26, ETV6, GATA2, SAMD9, SAMD9L, BLM, NF1, CBL)	Local VA or locally contracted vendor	No	Blood		
* Reach out to GLA for information on use of NGS testing under a VA sponsored grant, with no cost to your local facility							





