



# TREATING ADOLESCENTS AND YOUNG ADULTS (AYA) WITH BLOOD CANCER

October 11, 2023



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## WELCOME AND INTRODUCTIONS

**Lesley Hoerst, BSN, RN**  
*Senior Manager, Professional Education Programs*  
The Leukemia & Lymphoma Society  
Rye Brook, NY



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## SPEAKERS

### **Sharon M. Castellino MD, MSc**

Professor, Department of Pediatrics  
Emory University School of Medicine  
Director, Leukemia and Lymphoma Program  
Aflac Cancer and Blood Disorders Center  
Children's Healthcare of Atlanta  
Atlanta, GA

### **Julie Anna Wolfson, MD, MSHS**

Associate Professor, Pediatric Hematology-Oncology  
Institute for Cancer Outcomes and Survivorship  
Director, AYA Oncology and Oncofertility Program  
University of Alabama at Birmingham  
Birmingham, AL



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## DISCLOSURES

Sharon M. Castellino, MD, MSc, has a financial interest/relationship or affiliation in the form of:  
Advisory Board/Consultant: Bristol Myers Squibb

The following relationships have ended within the last 24 months:  
Advisory Board/Consultant: Seagen Inc.

#### Unlabeled Uses in Pediatrics

- Nivolumab
- Brentuximab vedotin (BV) (approved in high-risk; front line)
- Pembrolizumab (approved in rr/HL)

Julie Anna Wolfson, MD, MSHS, has no relevant financial relationships with ineligible companies to disclose for this educational activity.



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## DISCLOSURE

### Disclosure & Conflict of Interest Policy

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### Planning Committee and Content/Peer Reviewers

The planners and content/peer reviewers from Medical Learning Institute, Inc. and The Leukemia & Lymphoma Society do not have any relevant financial relationships to disclose with ineligible companies unless listed below.

#### Planner

Lauren Berger, MPH  
The Leukemia & Lymphoma Society

Lauren Berger, MPH, has a financial interest/relationship or affiliation in the form of:  
Stock Ownership with Bristol Myers Squibb, Gilead Sciences, Inc., Merck & Co., Inc., Organon & Co., Pfizer Inc., and Viatrix Inc.

All of the relevant financial relationships of individuals for this activity have been mitigated.

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### Disclaimer

Participants have an implied responsibility to use the newly acquired information to enhance patient outcomes and their own professional development. The information presented in this CE activity is not meant to serve as a guideline for patient management. Any procedures, medications, or other courses of diagnosis or treatment discussed or suggested in this CE activity should not be used by clinicians without evaluation of their patient's conditions and possible contraindications and/or dangers in use, review of any applicable manufacturer's product information, and comparison with recommendations of other authorities.

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## TARGET AUDIENCE

This activity is intended for hematologist/oncologists, nurses, social workers, and other healthcare professionals involved in the care of patients with blood cancer.

## EDUCATIONAL OBJECTIVES

*After completing this activity, the participant should be better able to:*

- Describe blood cancers common in adolescent and young adults (AYAs)
- Identify signs and symptoms of common blood cancers in AYAs and diagnostic tests used
- Explain treatment options, including new and emerging data and the role of clinical trials
- Discuss the management of short and long-term effects, as well as unique considerations for AYAs
- List resources to support patients and their caregivers



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## CE DESIGNATION

### Accreditation, Credit and Support



In support of improving patient care, this activity has been planned and implemented by Medical Learning Institute, Inc., and The Leukemia & Lymphoma Society. Medical Learning Institute, Inc. is jointly accredited by the Accreditation Council for Continuing Medical Education (ACCME), the Accreditation Council for Pharmacy Education (ACPE), and the American Nurses Credentialing Center (ANCC), to provide continuing education for the healthcare team.

### Physician Continuing Medical Education

Medical Learning Institute, Inc. (MLI) designates this live activity for a maximum of 1.0 *AMA PRA Category 1 Credits™*. Physicians should claim only the credit commensurate with the extent of their participation in the activity.

### Nursing Continuing Professional Development



Approval for nurses has been obtained by the National Office of The Leukemia & Lymphoma Society under Provider Number CEP 5832 to award 1.0 continuing education contact hour through the California Board of Registered Nursing.

### Social Worker Continuing Education

The Leukemia & Lymphoma Society (LLS) Provider Number 1105, is approved as an ACE provider to offer social work continuing education by the Association of Social Work Boards (ASWB) Approved Continuing Education (ACE) program. Regulatory boards are the final authority on courses accepted for continuing education credit. ACE provider approval period: 12/10/2020-12/10/2023. Social workers completing this course receive 1.0 clinical continuing education credit.

The Leukemia & Lymphoma Society (LLS) is recognized by the New York State Education Departments State Board for Social Work as an approved provider of continuing education for licensed social workers #0117. LLS maintains responsibility for the program. Social workers will receive 1.0 clinical CE contact hour for this activity.

### Interprofessional Continuing Education Credit



This activity was planned by and for the healthcare team, and learners will receive 1.0 Interprofessional Continuing Education (IPCE) credit for learning and change.

### Support Statement

There is no commercial support associated with this CE activity.

### Providers

This activity is provided by The Leukemia & Lymphoma Society and Medical Learning Institute, Inc.



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## METHOD OF PARTICIPATION

There are no fees for participating in or receiving credits for this accredited activity. For information on applicability and acceptance of continuing education credit for this activity, please consult your professional licensing board.

Learners must participate in the entire CE activity and submit the online evaluation form to earn credit, by clicking the link at the end of the presentation. Once submitted, the certificate will be generated. If you have questions regarding your certificate, please contact via email at [ndane@mlieducation.org](mailto:ndane@mlieducation.org).



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Institute for Cancer Outcomes and Survivorship



## Adolescents and Young Adults (AYA) with Blood Cancer

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### The Leukemia & Lymphoma Society

Julie Anna Wolfson, MD, MSHS  
Associate Professor, Pediatric Hematology-Oncology  
Member, Institute for Cancer Outcomes and Survivorship  
Director, AYA Oncology & Oncofertility Program  
October 11, 2023

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## Polling Question #1

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What is the NCI definition of an adolescent or young adult (AYA)?

- a) 15 years – 39 years
- b) 15 years – 24 years
- c) 12 years – 21 years
- d) 12 years – 24 years
- e) They act like a teenager

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# Adolescents and Young Adults (AYA) with Cancer

- Share diagnoses with older and/or younger patients

**Have not seen same improvement in survival over time**

**AYA GAP**

Keegan, Cancer 2016.

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## What Blood Cancers are AYAs Diagnosed with?

### Rates of New Cases by Cancer Type and Sex

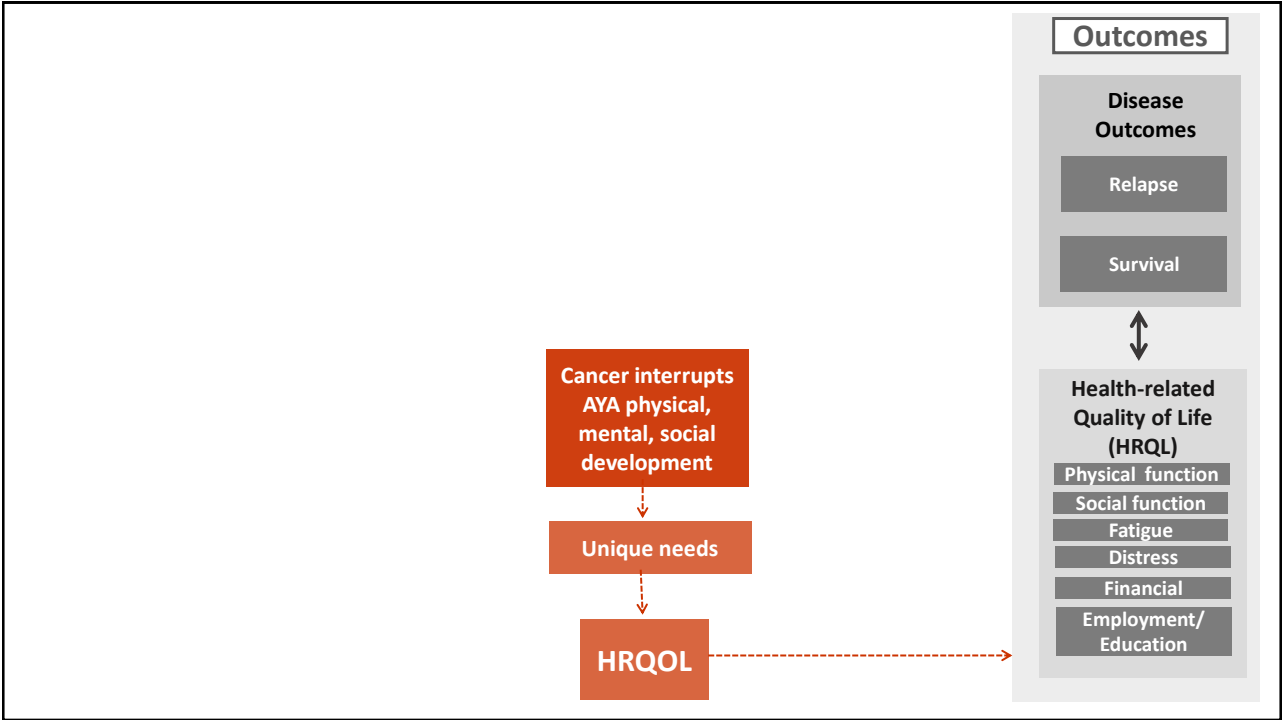
Cancer Type	Male	Female
Breast	0.1	23.0
Thyroid	4.4	18.5
Testis	11.6	N/A
Melanoma of the skin	3.9	7.4
Colon and rectum	5.1	5.2
<b>Non-Hodkin Lymphoma</b>	<b>4.6</b>	<b>3.4</b>
Leukemia	4.1	3.0
Hodgkin Lymphoma	3.5	3.4
Cervix	N/A	6.4
Brain & ONS	3.2	2.4

**New Cancer Cases, 2023**

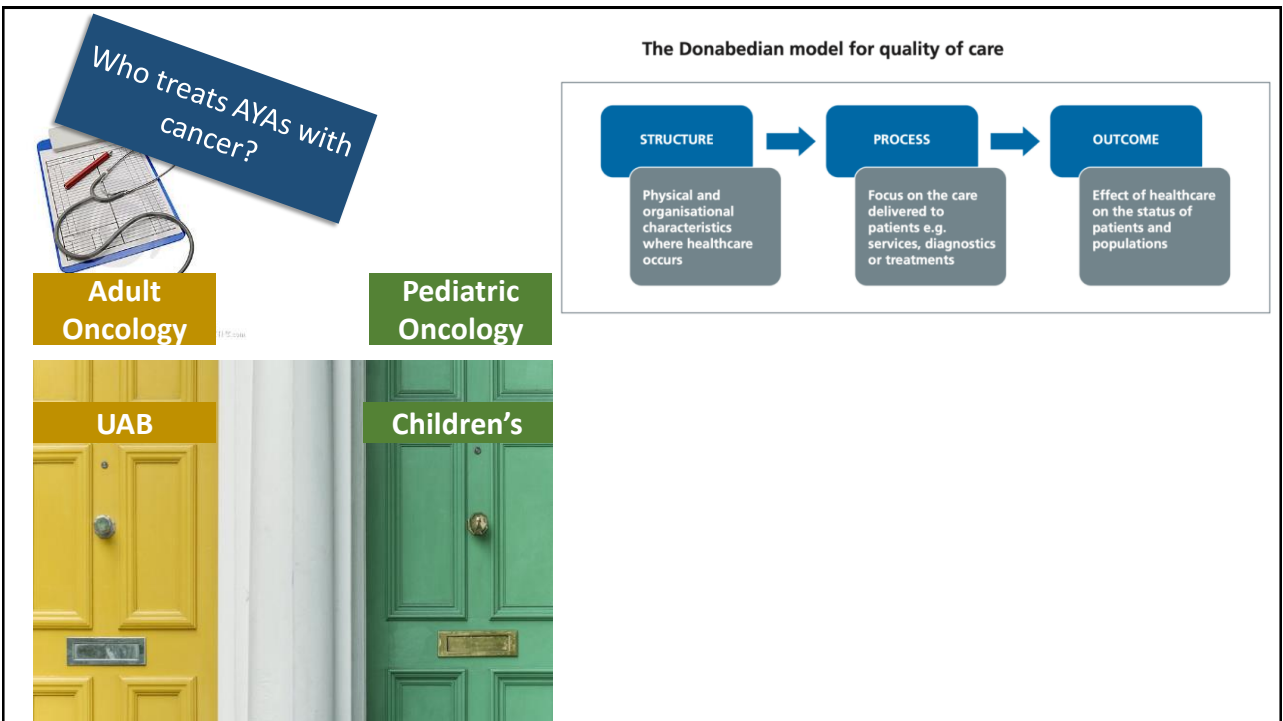
Estimated New Cancers Among AYAs in the U.S. in 2023	85,980
% of All New Cancer Cases at Any Age	4.4%

Age-adjusted rates of new cases per 100,000. SEER 22, 2016-2020. Cancer Stat Facts: Cancer Among AYAs (Ages 15-39); <https://seer.cancer.gov/statfacts/html/aya.html>

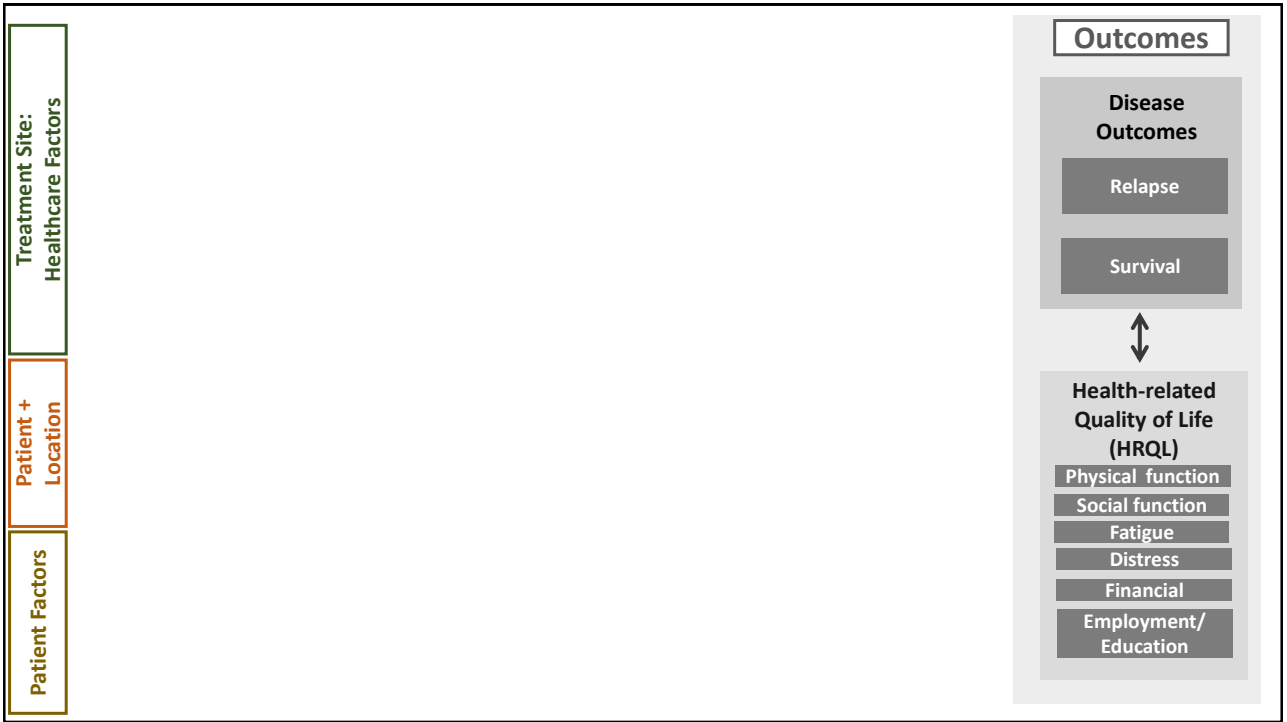
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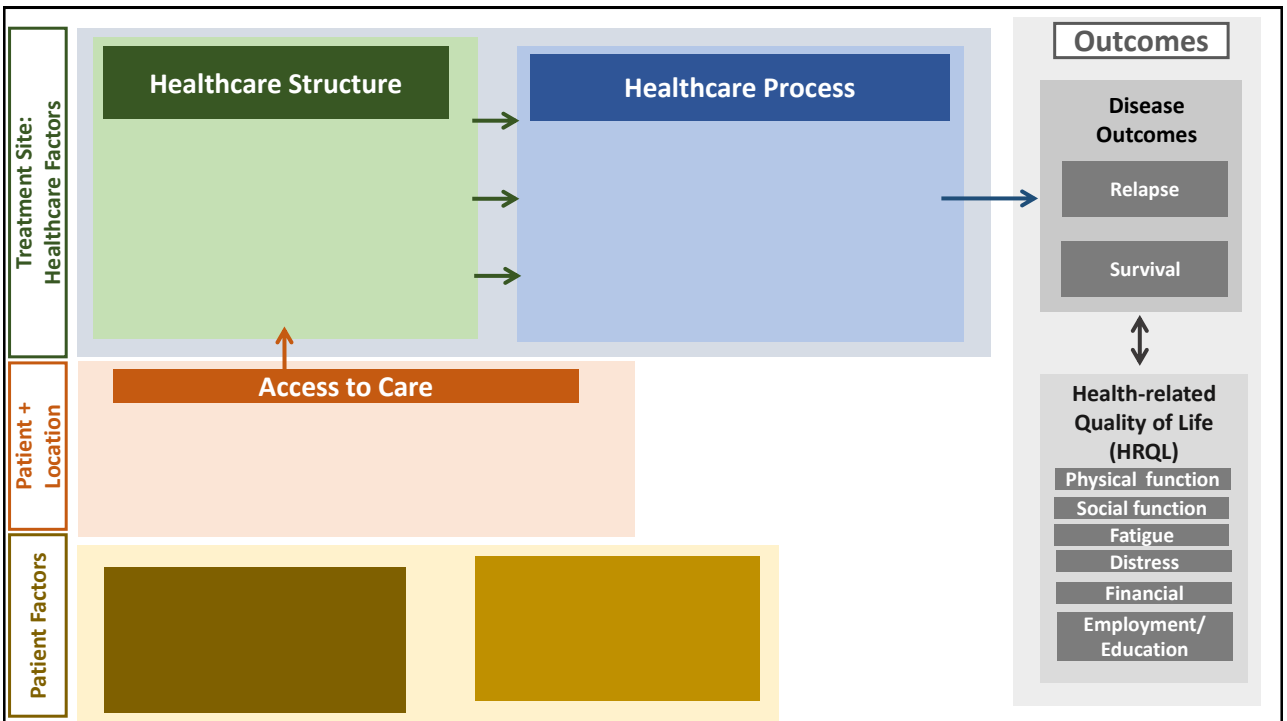
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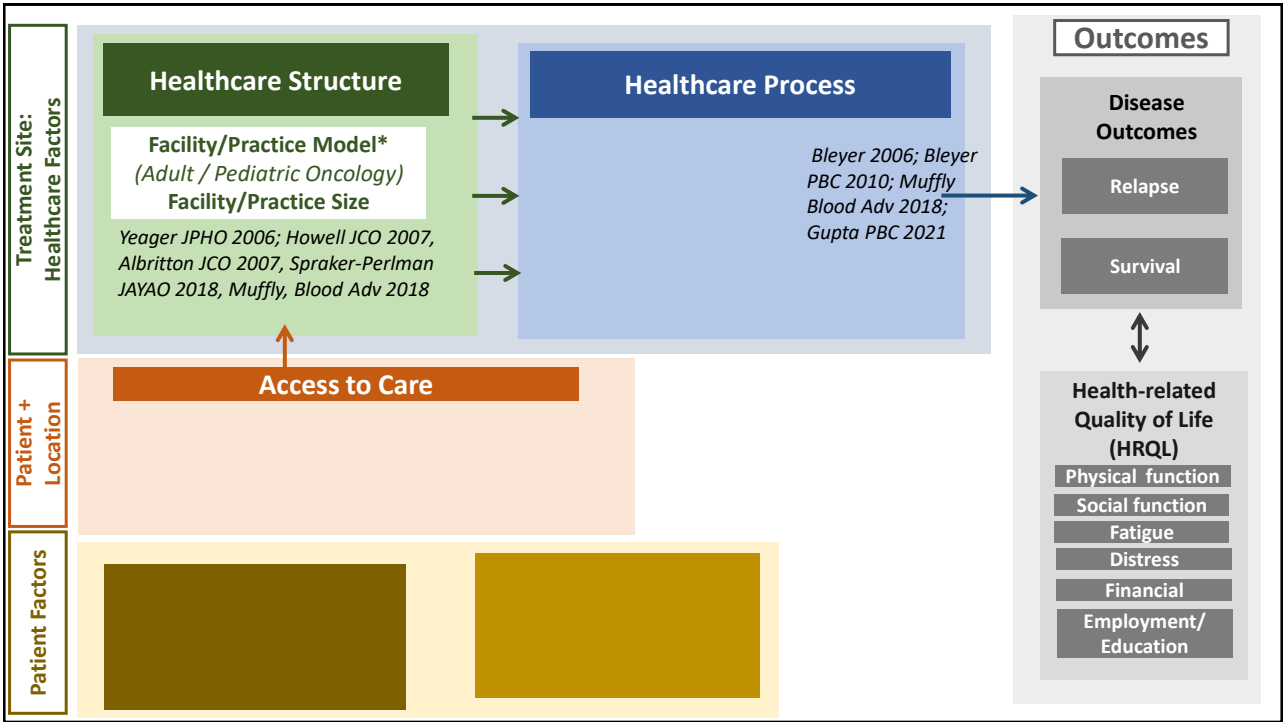


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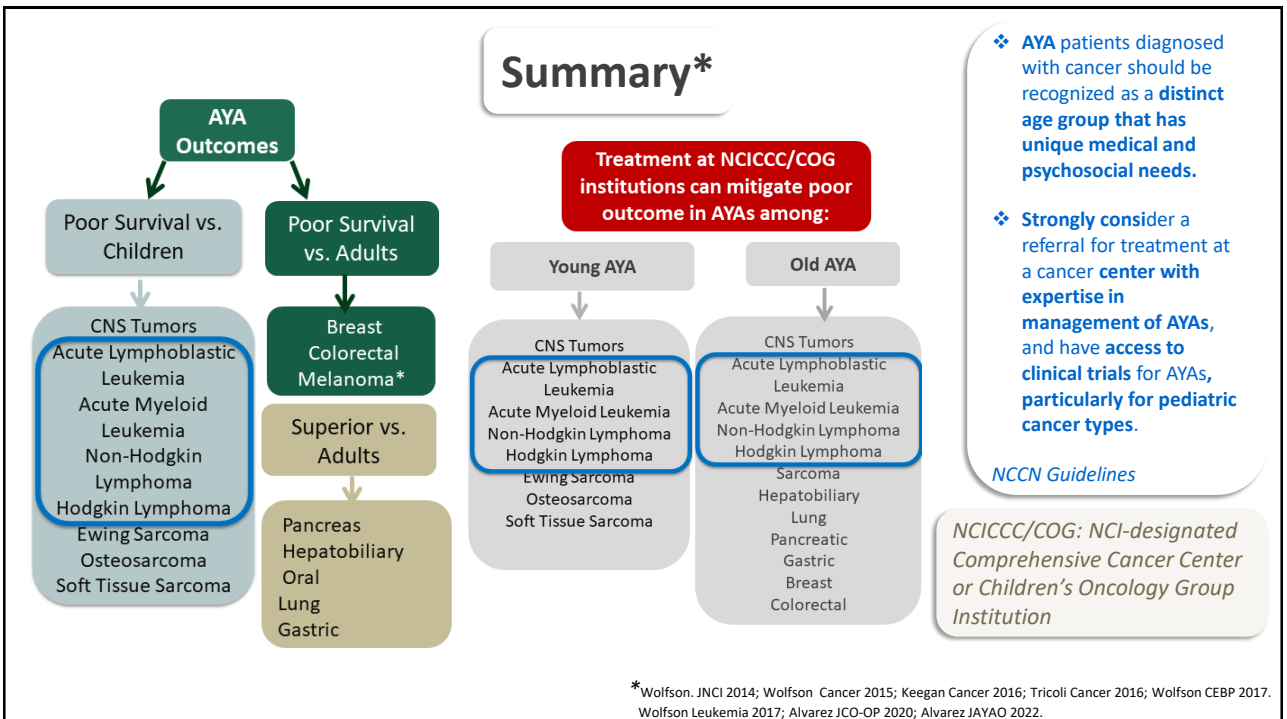


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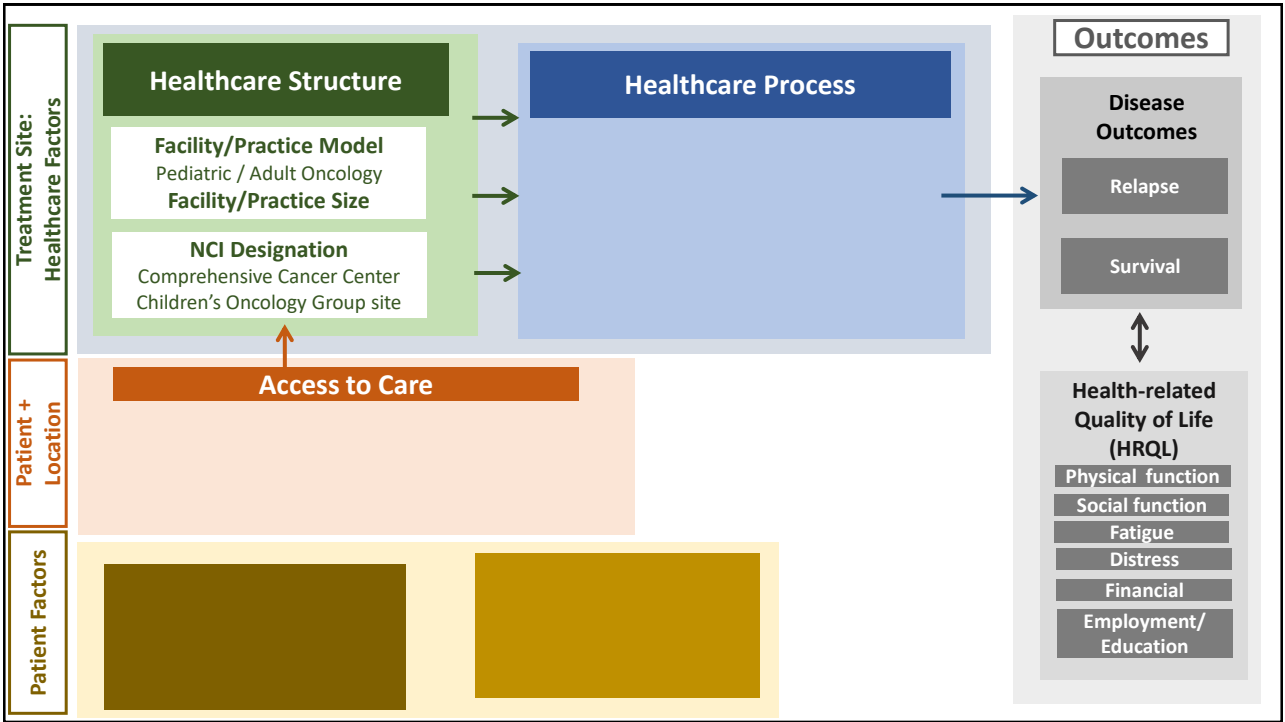




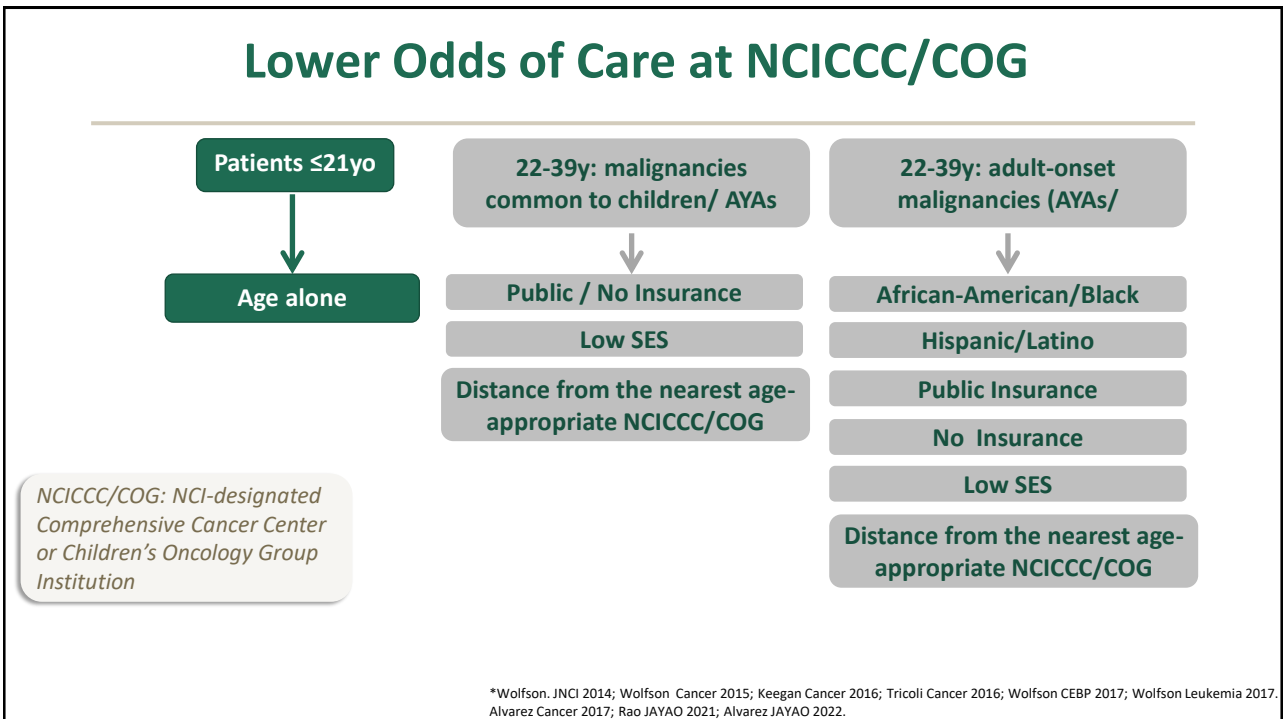
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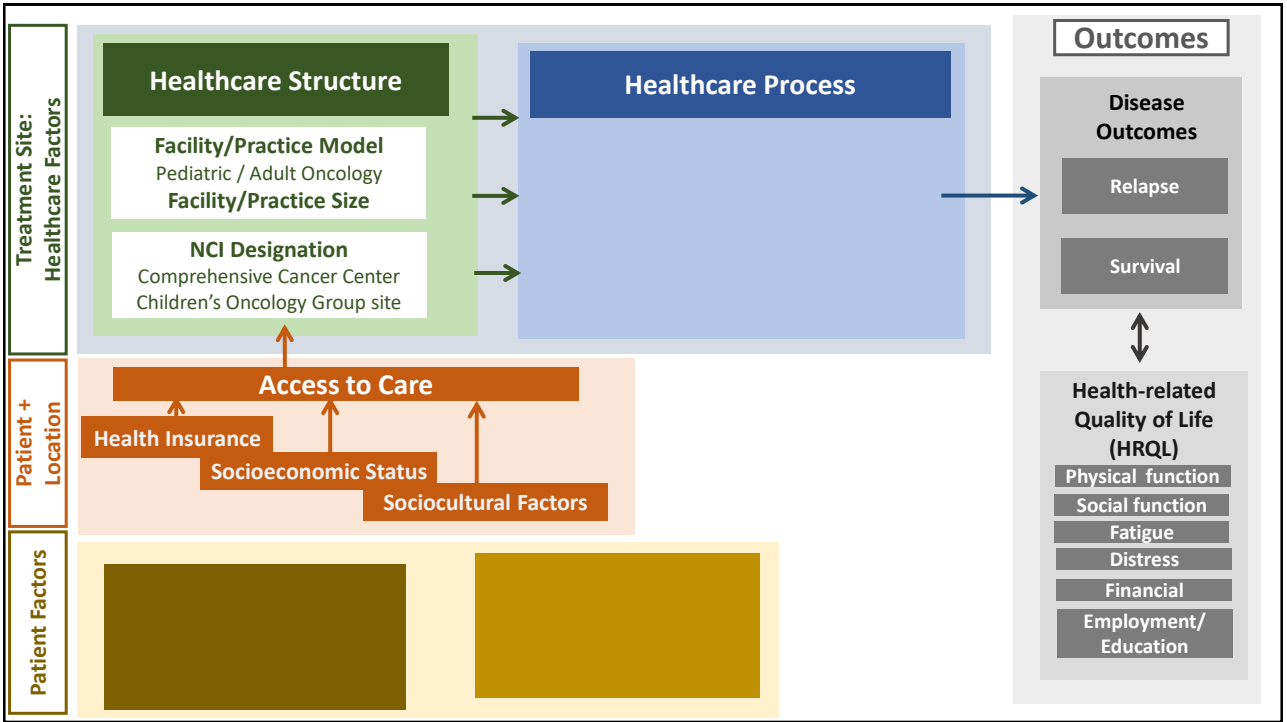
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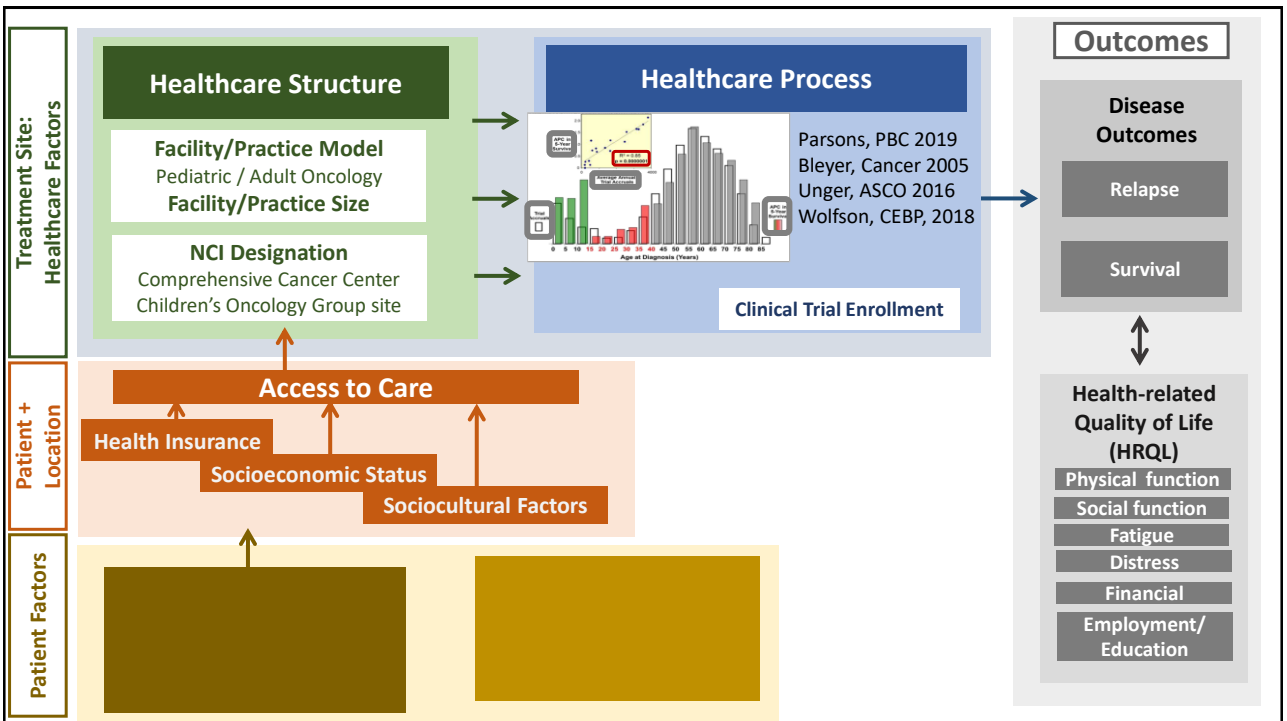
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**2005-2006**  
**AYA Progress Review Group**  
**NCI + Lance Armstrong**  
**Foundation**  
 Closing the Gap: Research & Care Imperatives for AYAs with Cancer



**Launched 2014 (est. 2010)**  
**NCI National Clinical Trials Network**

- Focus on increasing awareness of trials for AYAs
- 4 medical oncology, 1 pediatric oncology, 1 international group
- Proportion of CTEP treatment trials represented by AYAs pre-/post-creation of NCTN: 9.5% (95% CI, 7.6-11.8) vs, 14.0% (95% CI, 9.9-18.3). Mean difference in proportions: 4.4% (0.7%-8.3%).

*Sankaran...Seibel (Cancer, 2022) Cancer.gov*

**AYA Discipline Committees**

**CHILDREN'S ONCOLOGY GROUP** | **Southwest Oncology Group** A National Clinical Research Group | **Alliance for Clinical Trials in Oncology** | **ECOG-ACRIN** cancer research group Reshaping the future of patient care

**2022**  
**Joint Adult + Pediatric NCTN/SARC AYA Clinical Trials Sarcoma Working Group**  
*Whiteway et al, JAYAO 2023*

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**CTEP Cancer Therapy Evaluation Program**

**NCTN Adolescent to Young Adult (AYA) Cancer Trials Portfolio (Open as of 9/15/2023)**

Click on trial number to go to the associated ClinicalTrials.gov webpage, to view the protocol title and the study information.

**NCTN Trial Portfolios by Disease**

*Note: For full functionality, it is recommended that users download the PDF file, and open with a PDF reader.*

**NCTN Trial Portfolio (including all disease areas) (pdf)**

Cancer trials by disease area:

- Brain (pdf)
- Breast (pdf)
- Gastrointestinal (pdf)
- Genitourinary (pdf)
- Gynecological (pdf)
- Head and Neck (pdf)
- Leukemia (pdf)
- Lymphoma (pdf)
- Myeloma (pdf)
- Sarcoma (pdf)
- Skin (Mainly Melanoma) (pdf)
- Thoracic (pdf)
- Adolescent and Young Adult (AYA) (pdf)

**Newly Diagnosed Disease**

- Rhabdomyosarcoma (RMS)**: ARST2031 (High risk)
- Germ Cell Tumors**: AGCT1531, AGCT1532
- Osteosarcoma (Pulmonary Mets)**: AOST2031
- Osteosarcoma (Metastatic)**: AOST2032
- Classical Hodgkin Lymphoma**: AHOD2131
- Acute Lymphoblastic Leukemia (ALL)**: A041501
- Nongerminomatous germ cell tumor of the CNS**: ACNS2021
- Mediastinal (thymic) large B-cell lymphoma**: ANHL1931

**Recurrent Disease**

- Acute Lymphoblastic Leukemia (ALL)**: AALL1821
- Classical Hodgkin Lymphoma**: E4412
- Osteosarcoma (Pulmonary Mets)**: AOST2031

**Legend by Disease Types**

- Blue = ALL
- Light Green = Osteosarcoma
- Light Purple = CNS NGGCT
- Gray = PMBCL

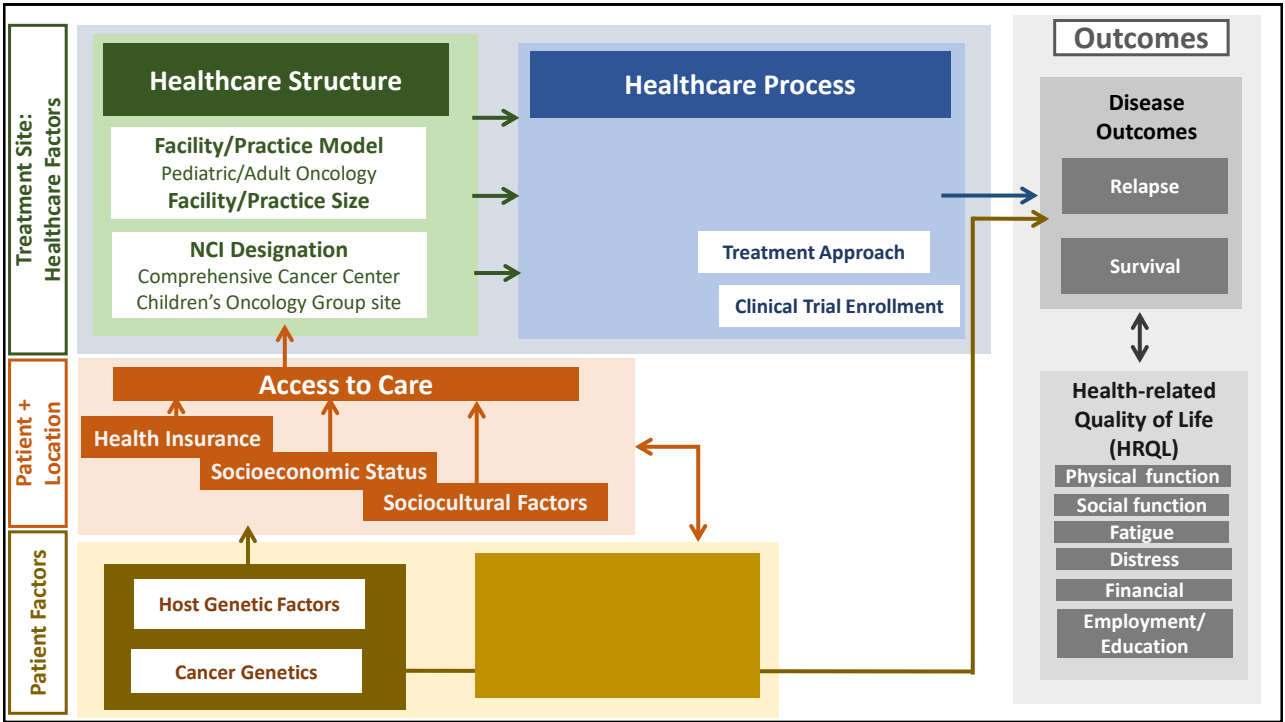
**CTSU (NCI Clinical Trials Support Unit)**

- AYA (NCTN)
- ALLIANCE
- A151804
- COG
- ACNS2021
- AGCT1531
- AGCT1532
- ANHL1931
- AOST2031
- ARST2031
- ECOG-ACRIN
- EAY191
- SWOG
- S1823

Protocol Number	Phase	Protocol Title
A041501	III	A Phase II Trial to Evaluate the Efficacy of the Addition of Intratumoral Ozogamicin (a Conjugated Anti-CD22 Monoclonal Antibody) to Routine Therapy in Young Adults (Ages 18-39 Years) with Newly Diagnosed Precursor B-Cell ALL.
AALL1821	II	A Phase 2 Study of Brentuximab (NCT0155866) in Combination with Vinorelbine (NCT01482726), a Checkpoint Inhibitor of PD-L1 in B-ALL Patients Aged 1 to 31 Years Old with First Relapse
ACNS2021	II	A Phase 2 Trial of Chemotherapy Followed by Response-Based Whole Ventricular & Spinal Canal Irradiation (WVSCI) for Patients with Localized Non-Germinalomatous Central Nervous System Germ Cell Tumor.
ARST2031	II	A Phase 3 Study of Active Surveillance for Low Risk and a Randomized Trial of Carboplatin vs. Cisplatin for Standard Risk Pediatric and Adult Patients with Germ Cell Tumors
AGCT1531	III	A Randomized Phase 3 Trial of Accelerated Versus Standard BEP Chemotherapy for Patients with Intermediate and Poor-Risk Germ Cell Tumors
AGCT1532	III	A Randomized Phase 3 Trial of Neovulvum (NCT01748326) in Combination with Chemo-Immunotherapy for the Treatment of Newly Diagnosed Primary Mediastinal B-Cell Lymphoma
ANHL1931	III	A Randomized Phase 3 Interim Response Adapted Trial Comparing Standard Therapy with Immuno-oncology Therapy for Children and Adults with Newly Diagnosed Stage I and II Classic Hodgkin Lymphoma
AHOD2131	III	A Phase 3 Randomized Controlled Trial Comparing Open vs Thoracoscopic Management of Pulmonary Metastases in Patients with Osteosarcoma
AOST2031	III	A Feasibility and Randomized Phase 2/3 Study of the VEGFR2/MET Inhibitor Cabozantinib in Combination with Cytotoxic Chemotherapy for Newly Diagnosed Osteosarcoma
AOST2032	II/III	A Randomized Phase 3 Trial of Vinorelbine, Doxorubicin, and Cyclophosphamide (VINO AC) Plus Maintenance Chemotherapy with Vinorelbine and Oral Cyclophosphamide (VINO-CPO) vs Vinorelbine, Doxorubicin, and Cyclophosphamide (VAC) Plus VINO-CPO Maintenance in Patients with High Risk Rhabdomyosarcoma (HR-RMS)
ARST2031	III	A Phase 1 Study with an Expansion Cohort/Randomized Phase II Study of the Combinations of topotecan, Neovulvum and Brentuximab in Patients with Relapsed/Refractory Hodgkin Lymphoma
E4412	III	

This portfolio shows NCTN trials. For information about NCORP trials, please see: <https://ncorp.cancer.gov/find-a-study/>

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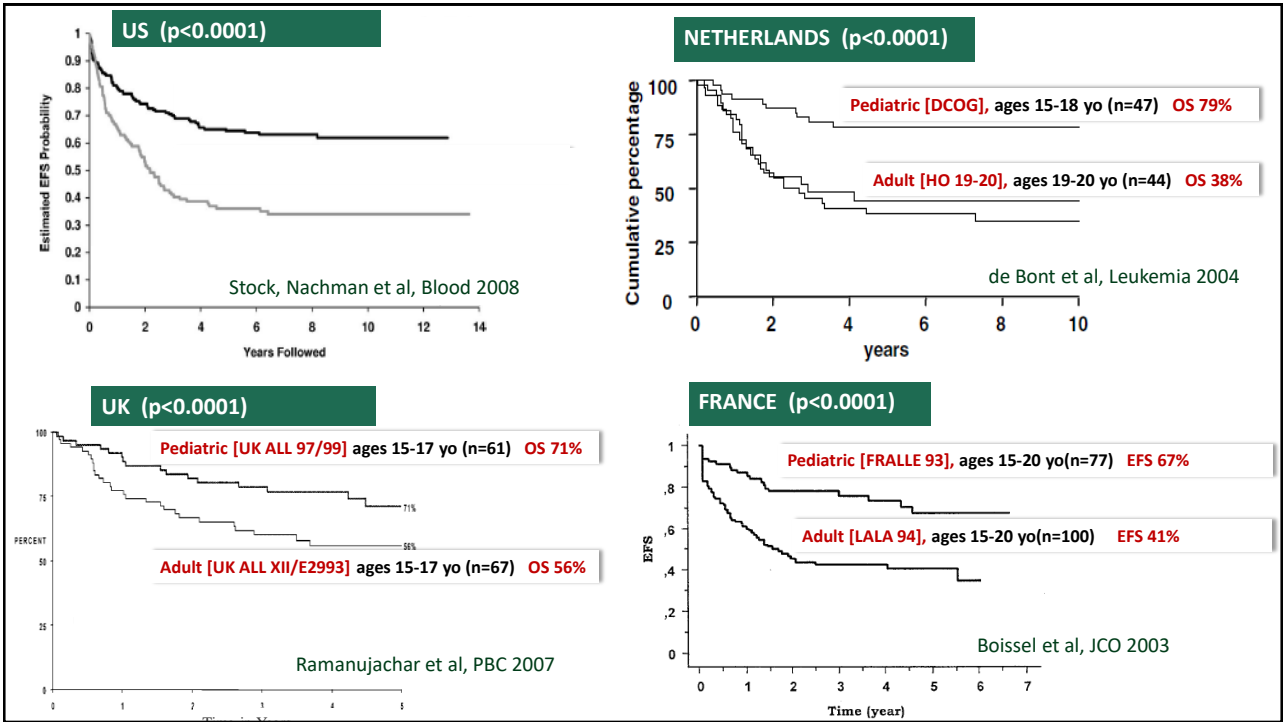


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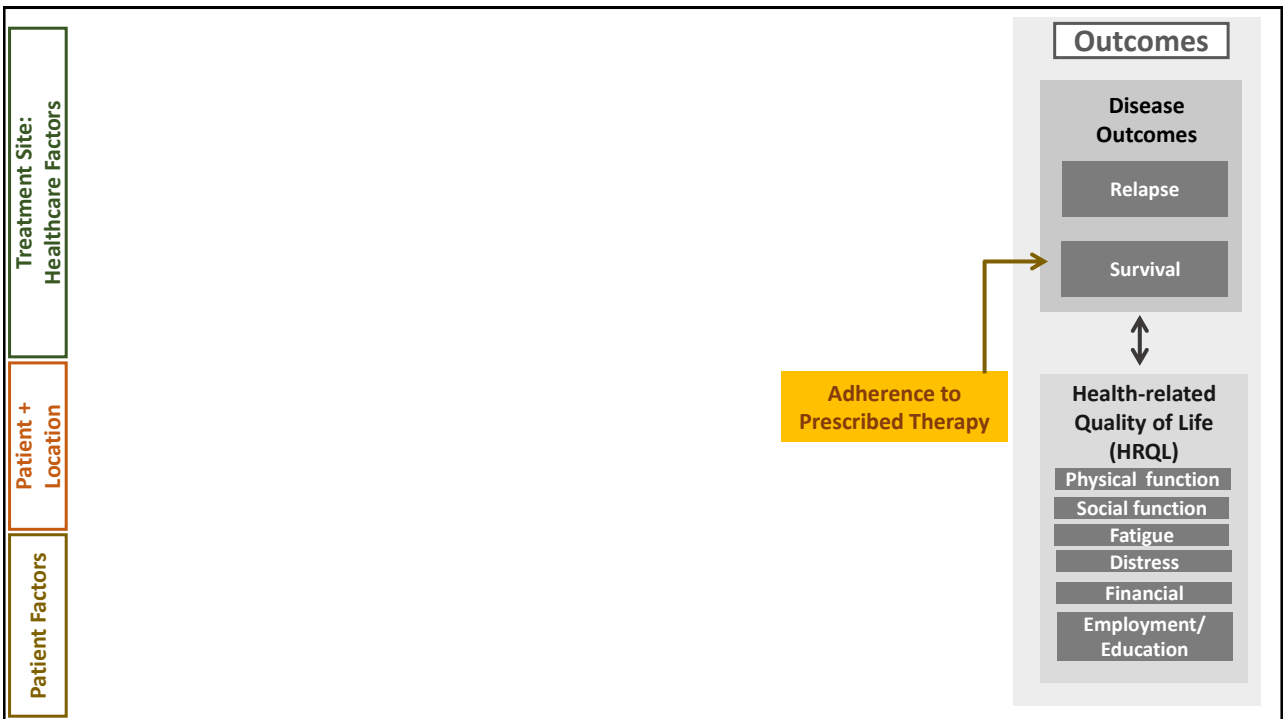
**Retrospective review of clinical trial data among AYAs (of the same age) with ALL treated on pediatric and adult trials**

Superior survival on pediatric trials

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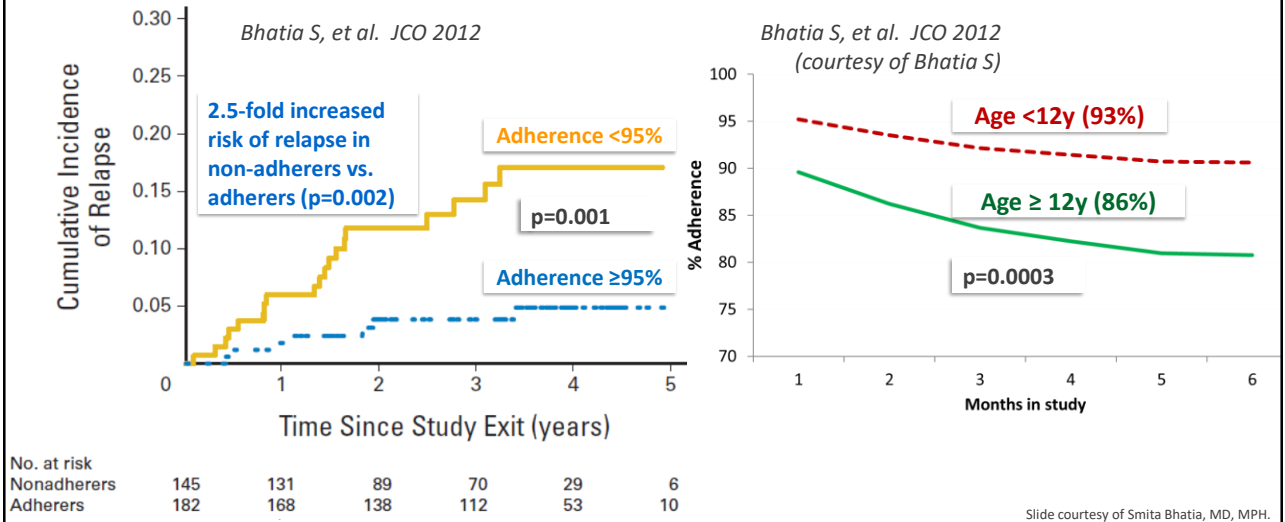
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## Childhood, Adolescent and Young Adult ALL: Relapse and Adherence

- Multisite Children's Oncology Group study (>94 sites) [Bhatia, Landier et al]
- Electronic monitoring of medication adherence (MEMS)
- Median age 6y, range 1-21y



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**Adherence is the critical factor for achieving molecular responses in patients with chronic myeloid leukemia who achieve complete cytogenetic responses on imatinib**

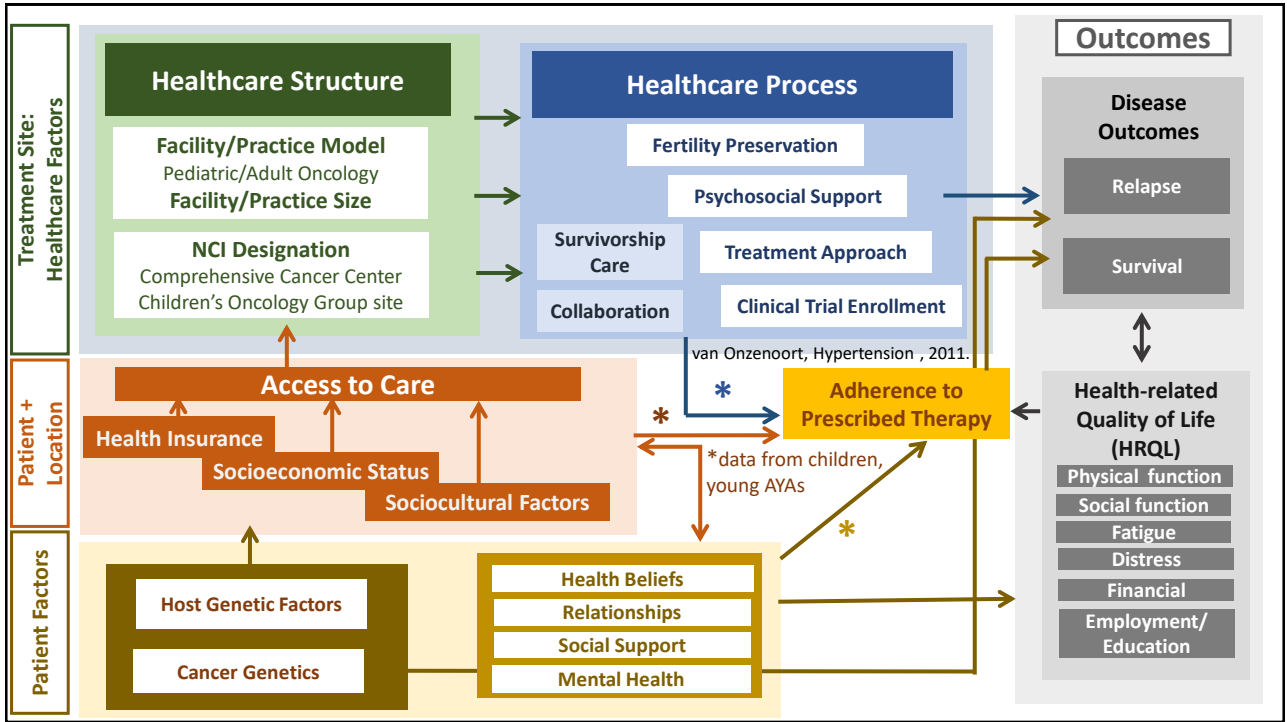
*Marin D et al. J Clin Oncol. 2010;28:2381-8*

- Median adherence rate 98% (24% to 104%).
- 26.4% had adherence  $\leq 90\%$ ; 14% had adherence  $\leq 80\%$
- Strong correlation between adherence ( $\leq 90\%$  or  $> 90\%$ ) and 6-year probability of MMR (28.4% v 94.5%;  $P < .001$ )
- Multivariate analysis: adherence was independent predictor for response
- No molecular responses observed when adherence was  $\leq 80\%$  ( $P < .001$ )

“Imatinib works better if you take it!”

Slide courtesy of Ravi Bhatia, MD.

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## Cancer Therapy Confers a Risk for Infertility for AYAs

- National guidelines recommend discussing fertility risks and offering referral for preservation *before* starting gonadotoxic therapy
- However, fertility preservation referrals are very inconsistent

Fertility Preservation in Patients With Cancer: ASCO Clinical Practice Guideline Update  
Kaufak Olan, Zimmmer E, Harris, Ann H, Partridge, Gonsky P, Quinn, Joyce R, Kotecki, Hugh S, Stefan, W, Harshbarger, Erica T, Wang, and Adam W Laven

**Unique AYA unmet needs:** Financial, Mental Health, Support Group Services

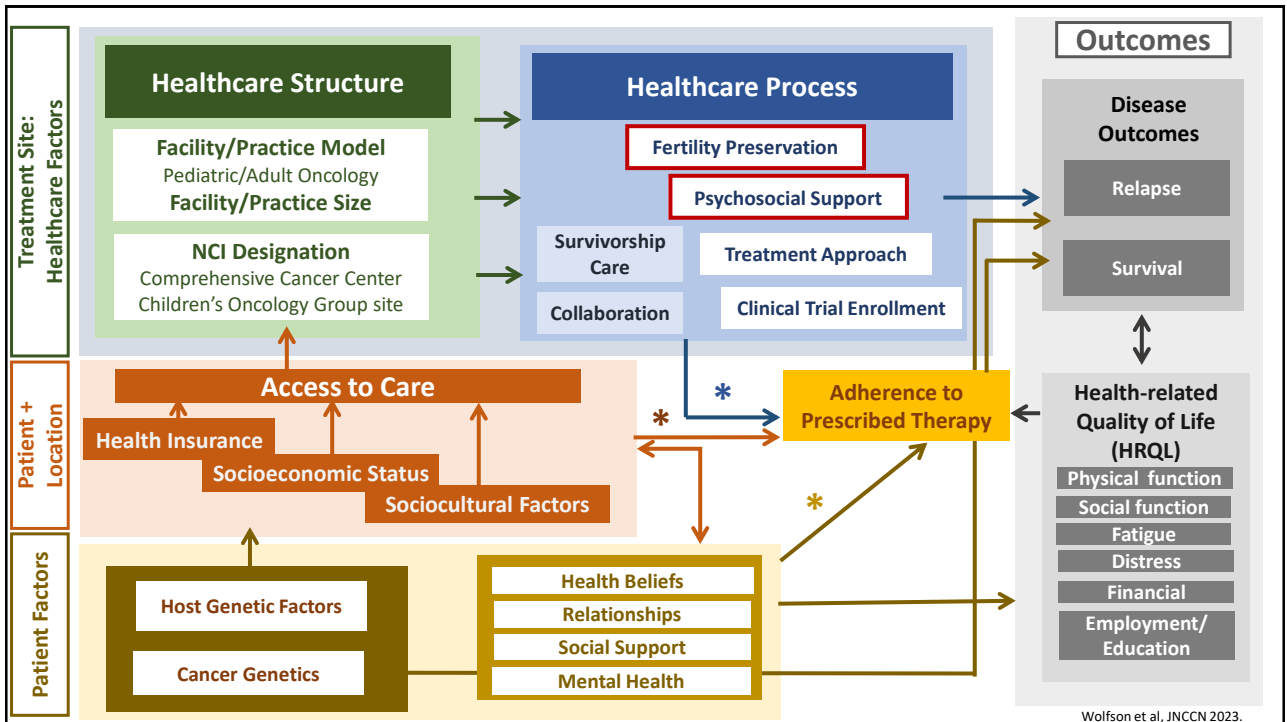
**Fertility Preservation Discussion** leads to:

- Health-related Quality of Life (HRQL)
- Functioning (physical, emotional, social, school/work)
- Fatigue
- Mental Health

Forman EJ, Fertil Steril 2010; Neal MS, Cancer 2007  
Ginsberg JP, PBC 2008; Schover LR, JCO 2002  
Partridge AH, JCO 2004; Schover LR, Med Ped Onc 1999  
Schover LR, JCO 2002; Schover LR, PBC 2009  
Zelbrack BJ, Psychooncology 2004; Smith AW, PBC 2019  
Burns KC, J Ped Hem Onc 2006; Keegan T, JCS 2012  
Smith AW, Frontiers Onc 2013;  
Wettergren L, Psychooncology 2017

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## Polling Question #2

**When you refer an AYA for a fertility preservation consultation before they start chemotherapy, how much is able to be done at your institution vs. outside your institution?**

- In-house: Oocyte cryopreservation, embryo cryopreservation, ovarian tissue cryopreservation, sperm banking; Outside referrals: none
- In-house: Oocyte cryopreservation, embryo cryopreservation; Outside referrals: ovarian tissue cryopreservation, sperm banking
- In-house: sperm banking; Outside referrals: Oocyte cryopreservation, embryo cryopreservation, ovarian tissue cryopreservation,
- In-house: none; Outside referrals: Oocyte cryopreservation, embryo cryopreservation, ovarian tissue cryopreservation, sperm banking
- Other

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# Treatment Options and New Emerging Data: Leukemias

## Chronic Myeloid Leukemia: CML

Acute Lymphoblastic Leukemia: ALL

Acute Myeloid Leukemia: AML

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## CML: Staging and Disease Response

### Staging of CML (MD Anderson criteria)

<b>Chronic phase</b>
None of the criteria for accelerated or blastic phase
<b>Accelerated phase</b>
Blasts $\geq$ 15% in blood or BM
Blasts plus progranulocytes $\geq$ 30% in blood or BM
Basophilia $\geq$ 20% in blood or BM
Platelets $<$ $100 \times 10^9/L$ unrelated to therapy
Cytogenetic clonal evolution
<b>Blast phase</b>
$\geq$ 30% blasts in blood or BM
Extramedullary disease with localized immature blasts

### Response to TKI is the most important prognostic factor

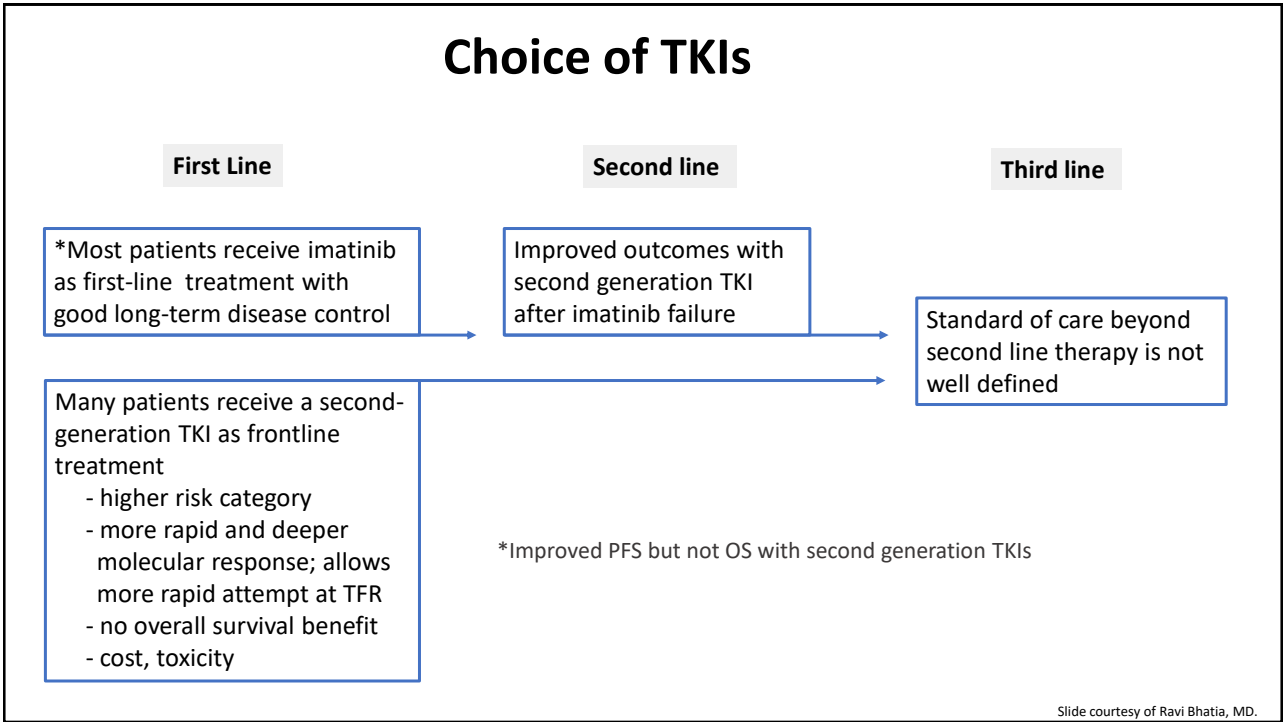
- Initial response to therapy provides a sensitive measure of future clinical outcome
- Measurement of BCR-ABL1 transcript levels using RT-Q-PCR standardized to the international reporting scale (IS)
- Based on achievement of CCyR or MMR at key time points
- Treatment failure defined as *BCR-ABL1*  $>$ 10% at 6 months and  $>$ 1% at 12 months

<i>BCR-ABL1</i> (IS)	3 months	6 months	12 months <sup>1</sup>
$>$ 10% <sup>m</sup>	YELLOW	RED	RED
$>$ 1%–10%	GREEN	GREEN	YELLOW
$>$ 0.1%–1% (CCyR)	GREEN	GREEN	LIGHT GREEN
$\leq$ 0.1% (MMR)	GREEN	GREEN	GREEN

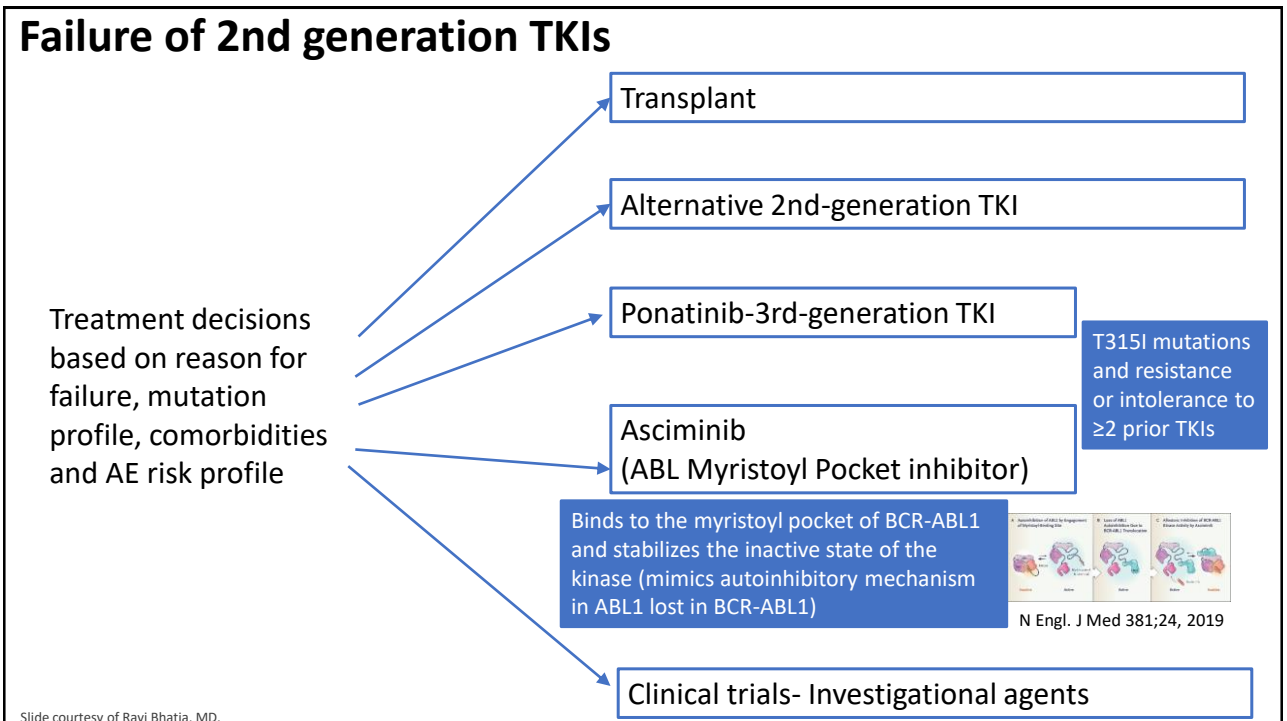
NCCN Guidelines Version 1.2022:  
Chronic Myeloid Leukemia

Slide courtesy of Ravi Bhatia, MD

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## Clinical Trials

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- **HQP1351 (Olverembatinib)**: a **3G TKI** with in vitro activity against **T315I and other mutants**
- **PF-114**: **3G TKI** with efficacy at nanomolar concentrations against **mutated BCR-ABL1, including the T315I mutation**; similar to Ponatinib but designed to **minimize interaction with VEGFR**
- **K0706 (Vodobatinib)**: a **2G TKI** effective against **wild-type and mutated BCR-ABL1** isoforms with **reduced off-target activity** compared to existing TKIs
- **Non BCR-ABL targets**

Slide courtesy of Ravi Bhatia, MD.

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## Important Considerations for AYAs

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- **Women** who take **TKIs** are at **risk of miscarriage and birth defects**, and are strongly advised to use birth control
- Women on TKI who **become pregnant** must **choose between ending the pregnancy or stopping the TKI temporarily**
- For women who choose to stop TKI treatment and continue with the pregnancy, and require **treatment, options include apheresis, and treatment with interferon alfa**
- **Breastfeeding** women are **advised to avoid TKIs** because these **medications are passed into breast milk**

Slide courtesy of Ravi Bhatia, MD.

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# Treatment Options and New Emerging Data: Leukemias

Chronic Myeloid Leukemia: CML

**Acute Lymphoblastic Leukemia: ALL**

Acute Myeloid Leukemia: AML

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## Polling Question #3

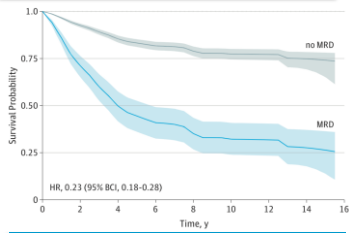
**What therapy is the best practice based on guidelines to treat AYAs with Ph-negative Acute Lymphoblastic Leukemia (ALL)?**

- a) CALGB 10403 or AALL1732
- b) DFCI ALL (001, etc)
- c) GRALLE-2005
- d) PETHEMA ALL-96
- e) Hyper-CVAD (without addition of other agents)
- f) Hyper-CVAD + Rituximab
- g) Hyper-CVAD + other targeted agent(s)
- h) Linker 4-drug regimen
- i) USC-MSKCC regimen (based on CCG1882)

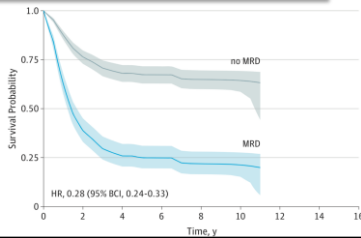
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Presence of MRD is BAD in Adult and Pediatric ALL...

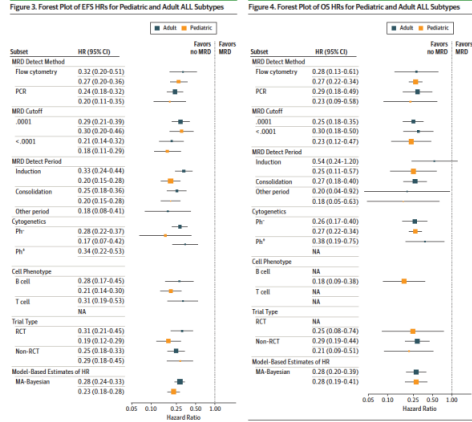
EFS for Pediatric ALL  
20 studies with 11,249 patients



EFS for Adult ALL  
16 studies with 2,065 patients



... Regardless of MRD Method, Detection Period, or ALL Subgroup

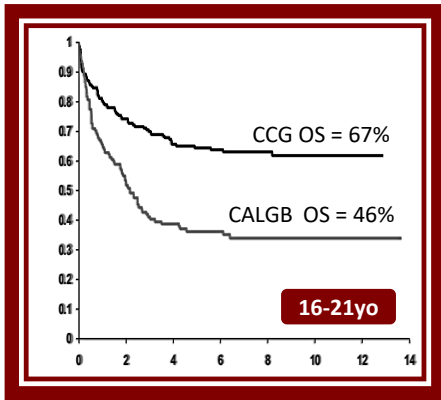


Berry et al, JAMA Onc, 2017  
Slide courtesy of Wendy Stock, MD.

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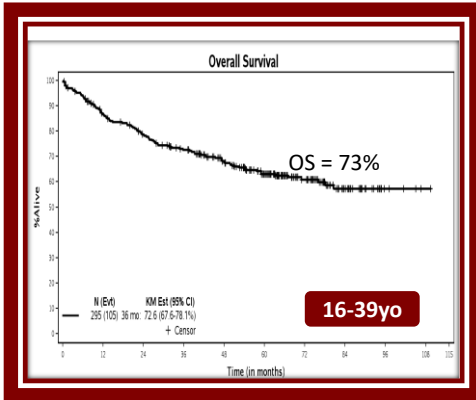
Young Adults: Treatment Standard has Changed

2000: Historical CALGB vs. CCG



Stock et al, Blood, 2008

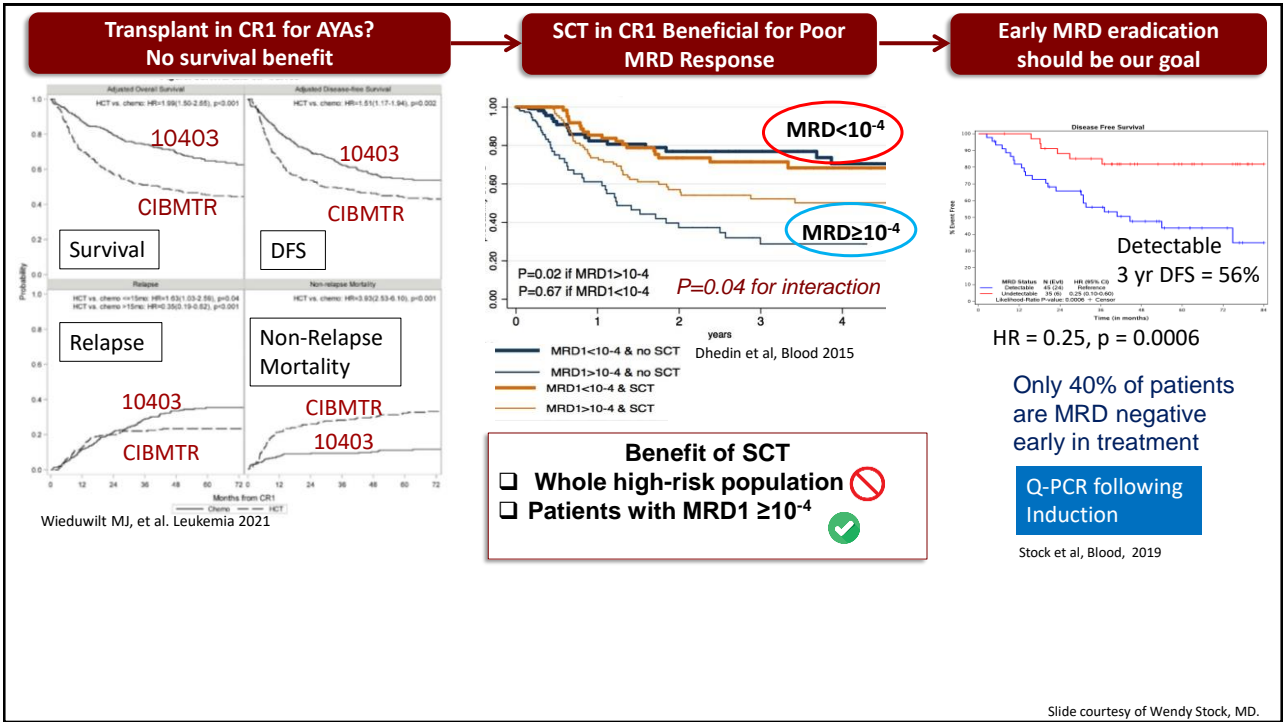
2019: CALGB 10403



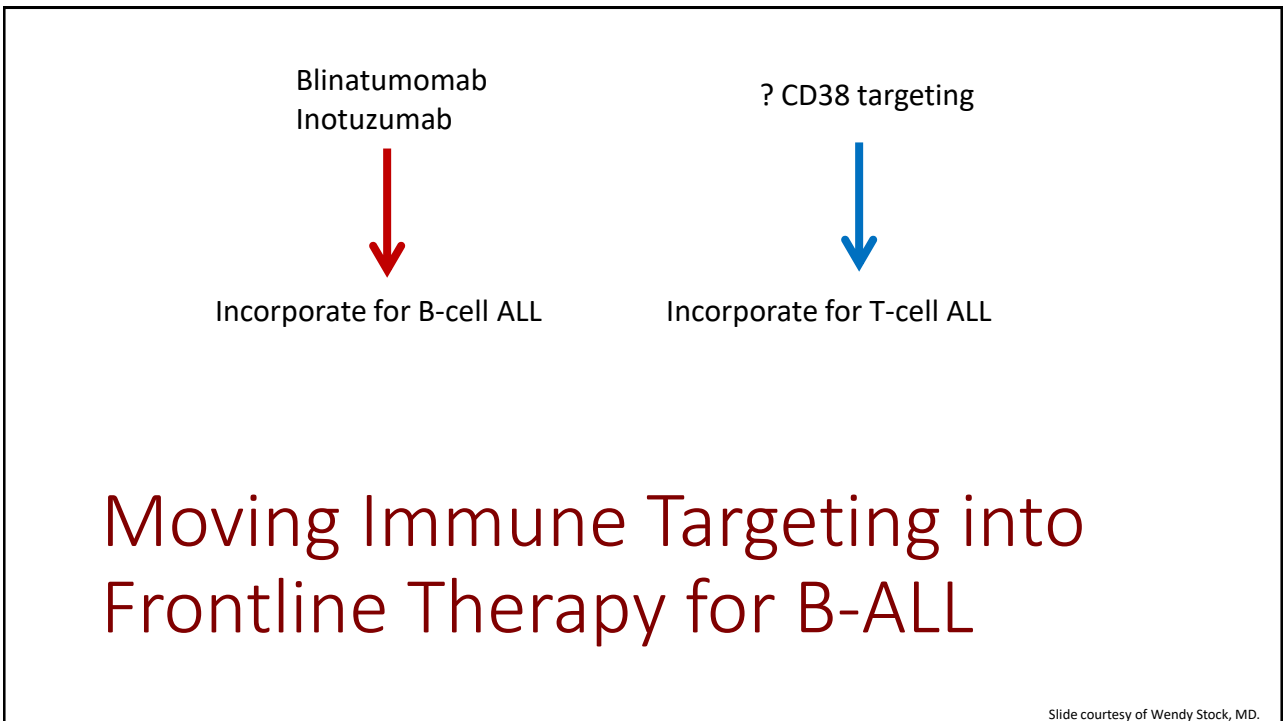
Stock et al, Blood, 2019

Slide courtesy of Wendy Stock, MD.

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# Blina in Frontline Phase III: E1910 for untreated B-ALL 30-60 years old

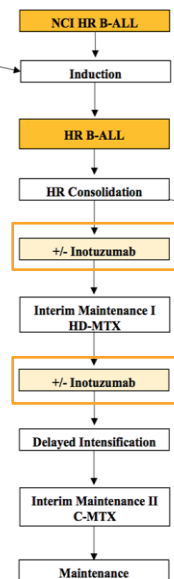
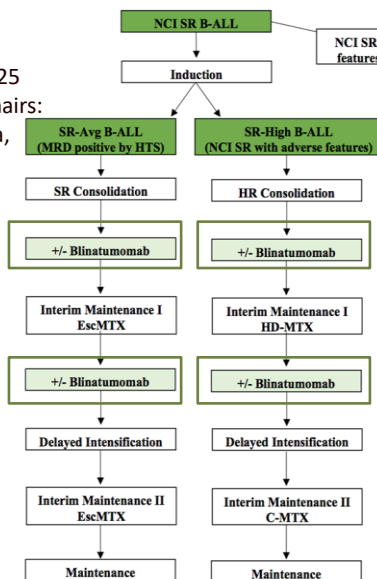
- Phase III randomized trial adding blina treatment modules at several treatment timepoints in a modified BFM backbone
  - 4 cycles of Blina are given; 2 cycles after intensification; 2 during late consolidation
- Initial goal was to evaluate efficacy of blinatumomab in frontline as treatment for both MRD- and MRD+ disease
- With approval of blinatumomab for MRD+ in 2018, only MRD- were subsequently randomized
- Completed enrollment fall 2019
- ASH 2022: Median follow-up 43 months, survival advantage of Blina (manuscript pending)

Slide courtesy of Wendy Stock, MD.

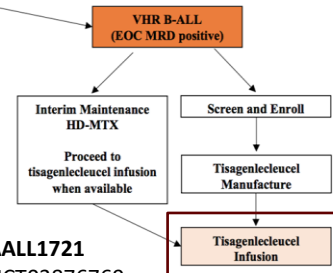
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## B-ALL: The Future - Incorporation of Immunotherapy into Frontline Therapy

**AALL1731**  
NCT03914625  
Study Co-Chairs:  
Sumit Gupta,  
Rachel Rau



**AALL1732**  
NCT03959085  
Study Co-Chairs:  
Jennifer McNeer,  
Maureen O'Brien



**AALL1721**  
NCT03876769  
Study Chair:  
Shannon Maude

**Pediatric (AYA) Trials**

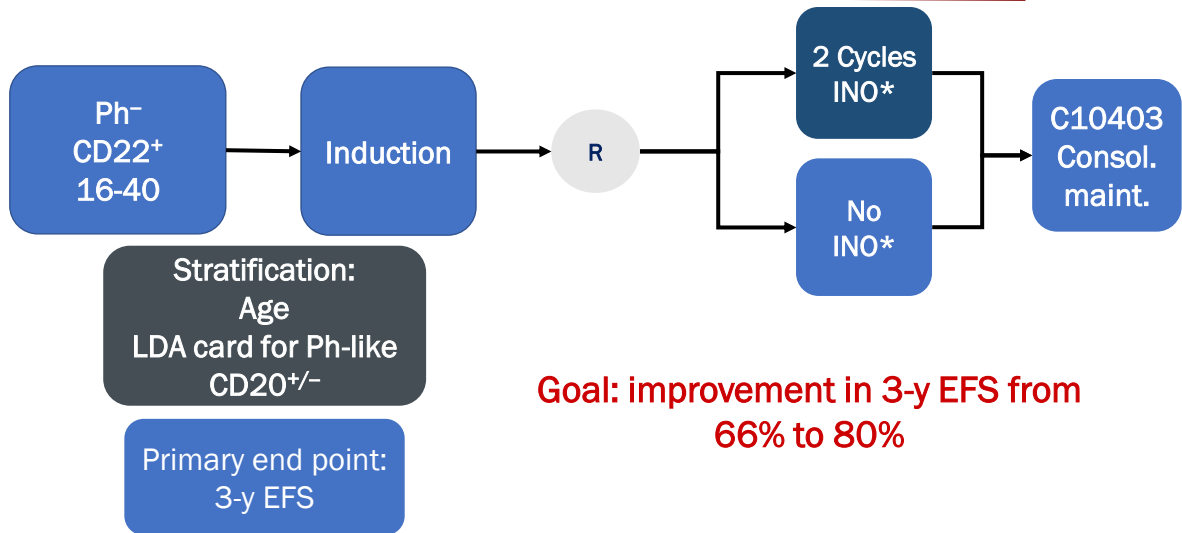
Slide courtesy of Jennifer McNeer, MD.

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# A041501 for AYAs 18-39 years: Can We Improve EFS to 80%?

Adult AYA Trial



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### Disease status predicts survival after CAR-T

**A** Event-free Survival, According to Disease Burden

Disease burden pre CAR-T

Low disease burden  
High disease burden

No. at Risk

Months since T-Cell Infusion	0	10	20	30	40	50	60
Low burden	20	10	7	5	4	2	1
High burden	31	8	0	0	0	0	0

P=0.01

### Allogeneic Transplant post CAR-T therapy improves EFS

**A** EFS probability vs Time after allogeneic HCT (months)

**B** OS probability vs Time after allogeneic HCT (months)

**C** Cumulative incidence vs Time after allogeneic HCT (months)

Non-relapse mortality  
Relapse

### Pediatric ELIANA trial: HCT necessary?

**A** Duration of Remission

Probability of Continued Remission vs Months since Onset of Remission

**B** Event-free and Overall Survival

Event-free survival  
Overall survival

Months since Tisagenlecleucel Infusion	0	2	4	6	8	10	12	14	16	18	20	22
No. of Patients	61	54	43	33	23	18	8	7	3	3	1	0

12 mo OS 76%, EFS 50%;  
Median duration not reached

Maude et al, NEJM, 2018.

**Overall response rate**

- Within 3 mos (all MRD neg): 81%
- Intent to treat: 66%
- SCT rate: 13% (all alive)

Association with EFS: HR 0.39, p=.088

Slide courtesy of Wendy Stock, MD.

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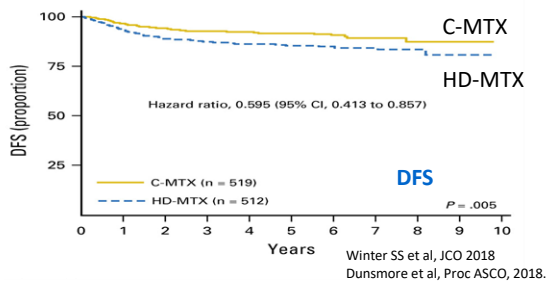
# Cellular Therapy: New Directions

- **"Off the shelf" CART**
  - Faster, doesn't require patient's cells to manufacture
  - Efficacy proven in early trials
- **Dual Targeted CART**
  - CD19, CD22 targeted CART cells have high response rates
  - May minimize emergence of resistant CD19 negative clones
- **Early phase CD5 targeted CAR-T**
  - Ongoing work;
  - Being viewed as bridge to transplant
- **Natural Killer (NK)-CAR**
  - No need for HLA full matching; NK cells may be derived from cord blood
  - Activity has been demonstrated using CAR-NK cells in CD19+ Lymphoma, CLL,

Lu et al, Abstract 284, ASH 2019; Huang et al, Cells 2022.  
Schultz et al, Abstract 744, ASH 2019; Spiegel et al, Nat Med. 2021.  
Liu et al, NEJM, 2020.  
Slide courtesy of Wendy Stock, MD

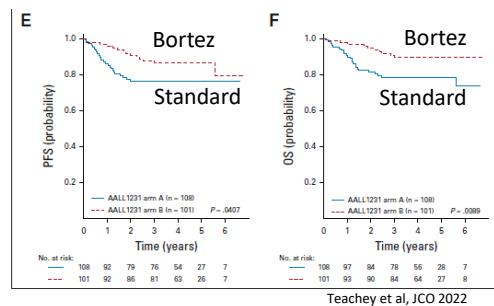
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## T-ALL: Nelarabine Improves Survival in COG AALL0434



- Nelarabine incorporated into ABFM; six 5-day courses
- **AYAs 20-30yo: 3% of the 1895 patients**
- **4yr DFS was 88.9% with nelarabine vs. 83% DFS without**

## TLLy: Improved EFS and OS with Bortezomib (AALL1231)



- Bortezomib incorporated into frontline therapy
- **4-year EFS**
  - **76.5% ± 5.1% vs. 86.4% ± 4.0% (p = 0.041)**
- **4-year OS**
  - **78.3% ± 4.9% vs. 89.5% ± 3.6% (p = 0.009)**

Slide courtesy of Wendy Stock, MD.

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# T-ALL: Immunotherapy

## Target

- **CD38**
  - Daratumumab
- **Phase 2 DELPHINUS study**
  - Daratumumab + chemo
  - 24 pediatric, 5 young adult pts
  - ORR
    - Pediatric ALL: 83.3%
    - Young Adult ALL: 60%
    - Lly: 40%

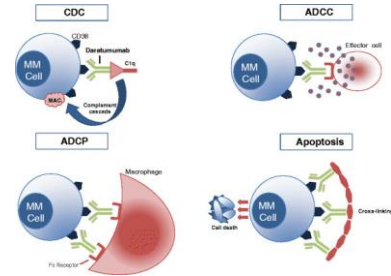


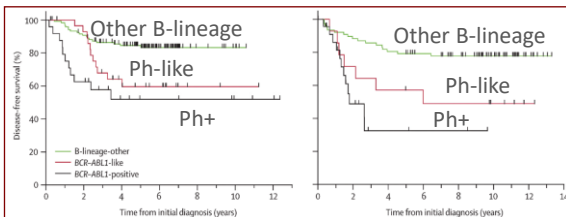
Figure from: Sanchez, et al. J Hematol Oncol. 2016  
Hogan, et al. JCO 2022, ASCO Annual Meeting.

Move to frontline therapy?

Slide courtesy of Jennifer McNeer, MD.

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## B-ALL: Molecular Diagnostics - Ph-like B-ALL



### 5y DFS (+ validation cohort)

- Ph-Like: 59.5%
- Ph+: 51.9%
- Other B-lineage: 84%

## Current Trials for Philadelphia chromosome-like ALL

Table 1. Kinase Fusions Identified in Ph-like Acute Lymphoblastic Leukemia.

Kinase Gene	Tyrosine Kinase Inhibitor	Fusion Partners number	Patients number	5' Genes
ABL1	Dasatinib	6	14	ETV6 <sup>11</sup> NUP214 <sup>11</sup> RCSD1 <sup>12</sup> RANBP2 <sup>13</sup> SNK2 <sup>14</sup> ZMI21 <sup>15</sup>
ABL2	Dasatinib	3	7	PAG1 <sup>16</sup> RCSD1 <sup>12</sup> ZC3H4V1 <sup>17</sup>
CSF3R	Dasatinib	1	4	SSBP2 <sup>18</sup>
PDGFRB	Dasatinib	4	11	EBF1 <sup>19,20</sup> SSBP2 <sup>18</sup> TNIP1 <sup>21</sup> ZEB2 <sup>22</sup>
CRLF2	JAK2 inhibitor	2	30	IGH <sup>23</sup> P381B <sup>24</sup>
JAK2	JAK2 inhibitor	10	19	ATF7IP <sup>25</sup> BCR <sup>26</sup> EBF1 <sup>27</sup> ETV6 <sup>28</sup> PAK5 <sup>29</sup> PPF1BP1 <sup>30</sup> SSBP2 <sup>31</sup> STNNA1 <sup>32</sup> TERF2 <sup>33</sup> TPR <sup>34</sup>
EPOR	JAK2 inhibitor	2	9	IGH <sup>35</sup> IGH <sup>36</sup>
DGKH	Unknown	1	1	ZFAND3 <sup>37</sup>
IL2RB	JAK1 inhibitor, JAK3 inhibitor, or both	1	1	MYH9 <sup>38</sup>
NTRK3	Crizotinib	1	1	ETV6 <sup>39,40</sup>
PTK2B	FAK inhibitor	2	1	KDM6A <sup>41</sup> STAG2 <sup>42</sup>
TSLP	JAK2 inhibitor	1	1	IQGAP2 <sup>43</sup>
TYK2	TYK2 inhibitor	1	1	MYB <sup>44</sup>

- Driven by a variety of signaling pathways
- Potential for targeted therapy in Ph-like ALL
  - **JAK/STAT pathway**
    - Ruxolitinib (AALL1521, recently closed to accrual)
  - **ABL-class fusions**
    - Dasatinib, Imatinib (AALL1631)

Den Boer, et al. Lancet Oncol 2009  
Roberts et al, NEJM, 2014  
Slide courtesy of Jennifer McNeer, MD.

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# Treatment Options and New Emerging Data: Leukemias

Chronic Myeloid Leukemia: CML

Acute Lymphoblastic Leukemia: ALL

Acute Myeloid Leukemia: AML

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## 2022 ELN AML Risk-Stratification (Adult)

Risk Category	Cytogenetic and Molecular Classification	Transplant Recommendation
<b>Favorable</b>	<ul style="list-style-type: none"> <li>t(15;17)</li> <li>inv(16)(p13.1q22) or t(16;16)(p13.1;q22)/CBFB::MYH11</li> <li>t(8;21)(q22;q22.1)/RUNX1::RXN1T1</li> <li>Mutated <i>NPM1</i> without <i>FLT3-ITD</i></li> <li>bZIP in-frame mutated <i>CEPBA</i></li> </ul>	CR2
<b>Intermediate</b>	<ul style="list-style-type: none"> <li>Mutated <i>NPM1</i> with <i>FLT3-ITD</i> mutation</li> <li>Wild-type <i>NPM1</i> with <i>FLT3-ITD</i> (w/o adverse genetic lesions)</li> <li>t(9;11)(p21.3;q23.3)/MLLT3::KMT2A</li> <li>Cytogenetic and/or abnormalities not classified as favorable or adverse</li> </ul>	CR1 for the majority of patients
<b>Adverse</b>	<ul style="list-style-type: none"> <li>t(6;9)(p23.3;q34.1)/DEK::NUP214</li> <li>t(v;11q23.3)/KMT2A-rearranged</li> <li>t(9;22)(q34.1;q11.2)/BCR::ABL1</li> <li>t(8;16)(p11.2;p13.3)/KAT6A::CREBBP</li> <li>Inv(3)(q21.3q26.2) or t(3;3)(q21.3;q26.2)/GATA2, MECOM(EV11)</li> <li>T(3q26.2;v)/MECOM(EV11) –rearranged</li> <li>-5 or del(5q); -7; -17/abn(17p)</li> <li>Complex karyotype, monosomal karyotype</li> <li>Mutated ASXL1, BCOR, EZH2, RUNX1, SF3B1, SRSF2, STAG2, U2AF1, and/or ZRSR2</li> <li>t(9;11)(p21.3;q23.3)/MLLT3::KMT2A</li> <li>Mutated TP53</li> </ul>	CR1

Slide courtesy of Kristen O'Dwyer, MD.  
Dohner H et al. Blood 2022; 140(12):1345-1377.

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# Risk Stratification – Pediatrics (AAML1831)

Table 3.1 Risk Stratification

	Low Risk							High Risk						
	LR1 4 chemo courses				LR2 5 chemo courses			HR 3 chemo courses and HSCT						
FLT3 ITD allelic ratio > 0.1	-	-	-	-	-	-	+	-	-	-	+	+/-	+	
FLT3 activating mutations (not ITD)	-	-	-	+	-	-	+/-	+/-	+/-	+	+	+/-	+/-	+/-
t(8;21) or inv(16)/t(16;16)	+1	-	-	+/-	+	-	-	+2	-	+	-	+/-	+/-	-
NPM1 or CEBPA	-	+	-	+/-	-	+	+	-	-	-	+	-	+/-	+
RAM phenotype or any unfavorable cytogenetic and/or NGS marker EXCEPT FLT3/ITD allelic ratio > 0.1*	-	-	-	-	-	-	-	-	-	-	-	+/-	EXCEPT	-
Measurable residual disease after Induction 1	-	-	-	-	+	+	-	-	+	+	+	+/-	+/-	+

FAVORABLE PROGNOSTIC MARKERS	
Cytogenetics	Genes
t(8;21)(q21.3;q22)	RUNX1-RUNX1T1
inv(16)(t(16;16)(p13.1q22.1))	CBFB-MYH11
No associated cytogenetic abnormality	NPM1 mutation positive
No associated cytogenetic abnormality	CEBPA bZIP mutation positive

UNFAVORABLE PROGNOSTIC MARKERS	
Cytogenetics	Genes
inv(3)(q21.3q26.2) / t(3;3)(q21.3q26.2)	RPNI-MECOM
t(3;21)(26.2;q22)	RUNX1-MECOM
t(3;5)(q25;q34)	NPM1-MLF1
t(6;9)(p22.3;q34.1)	DEK-NUP214
t(8;16)(p11.2;p13.3) (if 90 days or older at diagnosis)	KAT6A-CREBBP (if 90 days or older at diagnosis)
t(16;21)(p11.2;q22.2)	FUS-ERG
inv(16)(p13.3q24.3)	CBFA2T3-GLIS2
t(4;11)(q21;q23.3)	KMT2A-AFF1 (MLL-MLLT2)
t(6;11)(q27;q23.3)	KMT2A-AFDN (MLL-MLLT4)
t(10;11)(p12.3;q23.3)	KMT2A-MLLT10
t(10;11)(p12.1;q23.3)	KMT2A-ABI1
t(11;19)(q23.3;p13.3)	KMT2A-MLLT1(MLL-ENL)
11p15 rearrangement	NUP98- any partner gene
12p13.2 rearrangement	ETV6- any partner gene
Deletion 12p to include 12p13.2	Loss of ETV6

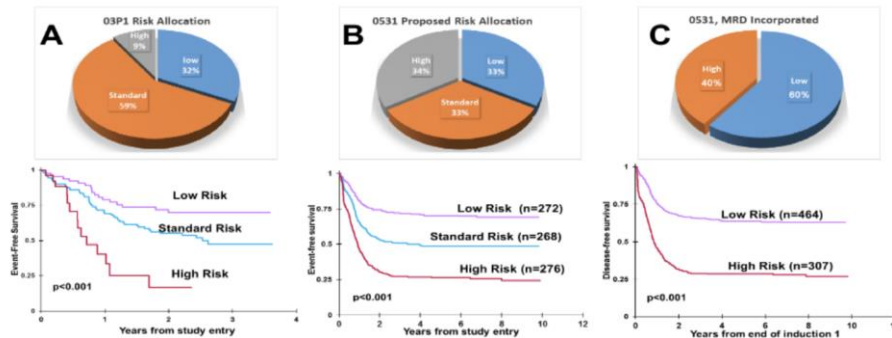
UNFAVORABLE PROGNOSTIC MARKERS	
Cytogenetics	Genes
Monosomy 5/De1(5q) to include 5q31	Loss of EGRI
Monosomy 7	No associated gene
10p12.3 rearrangement	MLLT10- any partner gene
No associated cytogenetic abnormality	FLT3/ITD + with allelic ratio > 0.1%
RAM phenotype as evidenced by flow cytometry	

Slide courtesy of Jennifer McNeer, MD.

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## Updated karyotype/FISH, new immunophenotypic, and NGS data applied retrospectively to prior COG AML patients

Figure 2.1 Risk Allocation



Thus, AAML1831 risk stratification more nuanced than prior studies.

Risk groups from AAML03P1  
 SR 60%  
 LR 30%  
 HR 10%

Initially proposed risk groups applied to AAML0531 data (~30% in each group)

Applying EO1 MRD 0.1% to SR group then further stratifies them into HR vs LR groups

Slide courtesy of Jennifer McNeer, MD.

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## New/Targeted Therapies in AML

- **Gemtuzumab**
  - Anti-CD33 conjugated to calicheamicin
  - AAML0531: outcome benefit (Gamis, JCO 2014)
    - CD33 expression (Pollard, JCO 2016)
    - FLT3/ITD (Tarlock, Clin Cancer Res 2016)
    - KMT2A (Pollard, JCO 2021)
    - Thus – added for all patients (**AAML1831**)
  - May increase risk of SOS with HSCT
  - FDA-approved 2017: adult CD-33+ AML, and peds ≥ 2 yrs with R/R CD33+ AML
- **Sorafenib, Gilteritinib**
  - Sorafenib: Multi-target TKI that targets *FLT3*, *c-KIT*, *PDGF*, *VEGF*, *RAF/MED/ERK* (**AAML0531**)
  - Gilteritinib: Multi-target TKI that targets FLT3 (ITD and TKD), with weak activity against c-Kit, and inhibits AXL (implicated in FLT3 inhibitor resistance) – **AAML1831**
- **CPX-351**
  - Liposomal 5:1 preparation of cytarabine:daunorubicin
  - Less cardiotoxicity
  - FDA-approved 2017 for adults with t-AML, or AML with MDS-related changes
    - COG AAML1421 (r/r), **COG AAML1831** (de novo)
- **Venetoclax**
  - BCL2 inhibitor (BCL2 is anti-apoptotic)
  - 2016/2017 – Breakthrough designation for AML
- **Azacitidine, Decitabine**
  - Epigenetic Modifiers
  - AML16 (St. Jude trial)

Slide courtesy of Jennifer McNeer, MD.

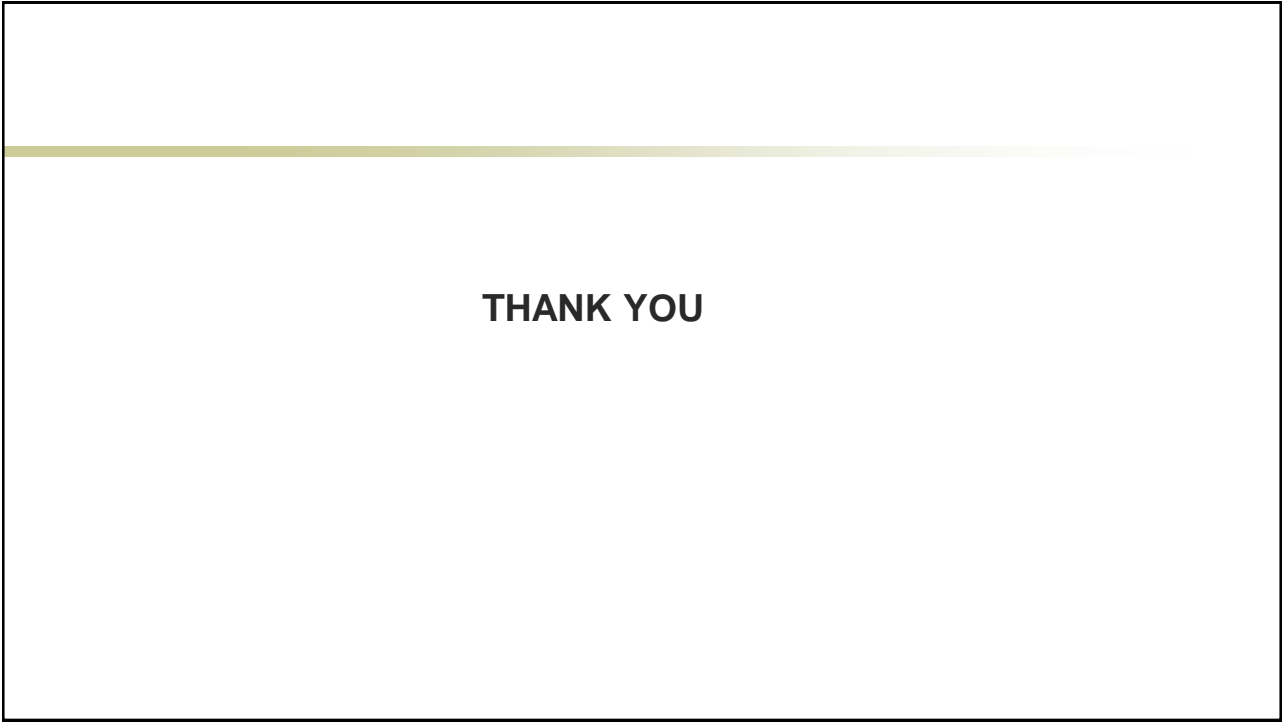
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## Recent AML Updates

- **AAML1031: Up front study**
  - Randomization ± Bortezomib
    - No benefit with bortezomib
  - 4 cycles of chemo
    - 4 cycles inferior to 5 when compared to historical data
- **AAML1331 (recently closed)**
  - Phase 3 study of arsenic and ATRA for APML
  - Omit anthracycline for standard-risk patients
  - Minimize anthracycline for high-risk patients
- **AAML1531**
  - Disease-response based treatment for DS-AML
    - >90 days and <4 years
  - Original study – omit HD-AraC for standard-risk patients.
    - Worse outcomes than historical control
      - 2-year EFS 85.6% vs 93.5%, p=0.0002
      - HD-AraC re-introduced

Slide courtesy of Jennifer McNeer, MD.

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**Diagnostic and Management Considerations for AYA with Lymphoma**

LLS Webinar  
Sharon M Castellino, MD, MSc  
Professor of Pediatrics, Emory School of Medicine  
Program Leader: Pediatric Leukemia and Lymphoma  
October 11, 2023

-

A decorative horizontal line consisting of 20 small, light grey dots arranged in two rows of ten.

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## Poll Question #4

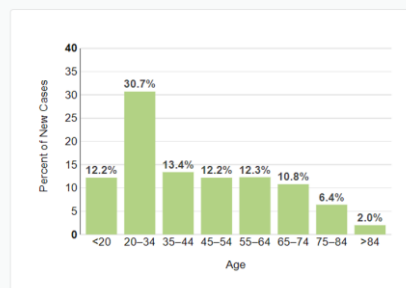
- What is the most common lymphoma in patients age 15-39 in the U.S.
  - A. Nodular Sclerosing Hodgkin Lymphoma (HL)
  - B. DLBCL
  - C. Primary CNS lymphoma
  - D. Nodular lymphocyte predominant HL

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## Epidemiology

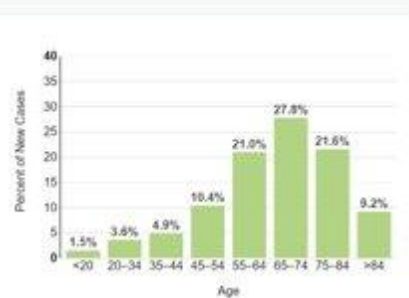
Lymphoma accounts for 20% of cancers in AYA

Percent of New Cases by Age Group: Hodgkin Lymphoma



SEER 21 2014-2018, All Races, Both Sexes

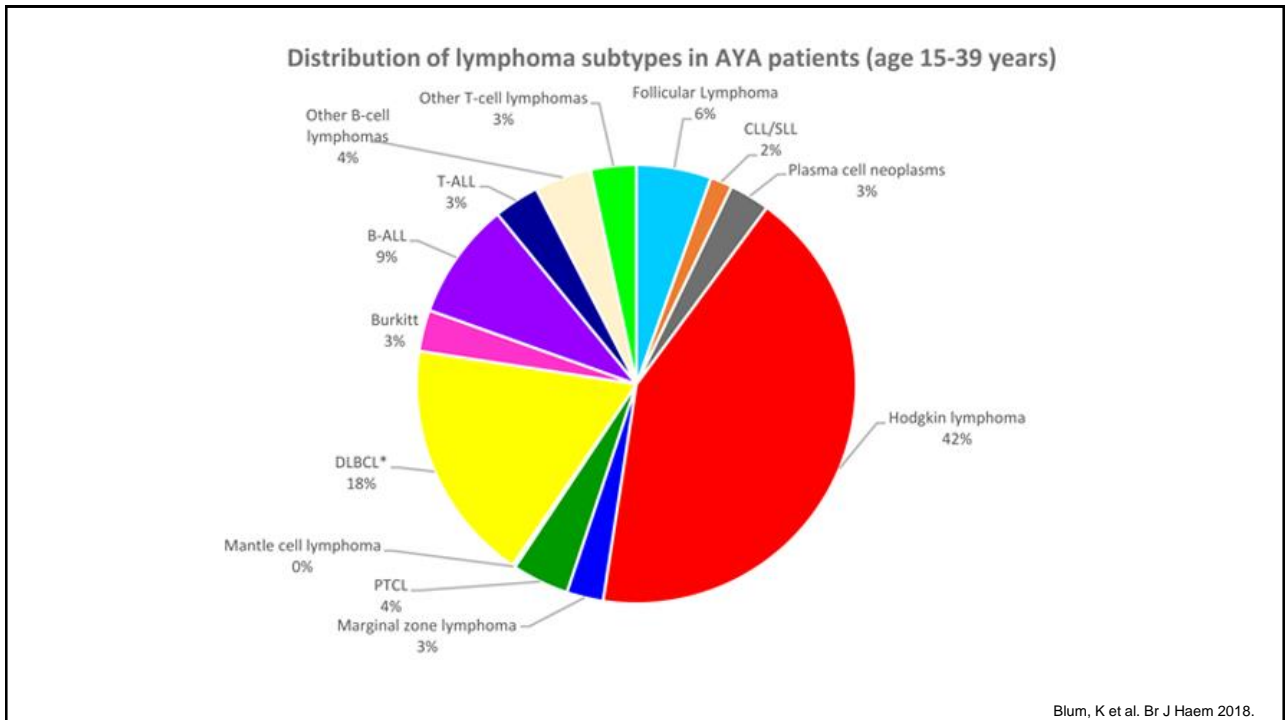
Percent of New Cases by Age Group: Non-Hodgkin Lymphoma



[https://www.researchgate.net/figure/Age-specific-rates-of-incidence-based-on-data-from-SEER-Cancer-Statistics-Review-16\\_fig1\\_49833144](https://www.researchgate.net/figure/Age-specific-rates-of-incidence-based-on-data-from-SEER-Cancer-Statistics-Review-16_fig1_49833144)

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## Disparities in Outcomes for AYA with Lymphoma

- Compared to pediatric patients:
  - AYA patients more likely to present with:
    - Advanced stage disease
    - B symptoms
- Clinical presentation:
  - Indolent: HL
  - Acutely ill with rapid progression in many NHL
- Diagnosis
  - Challenges associated with evolving molecular features in NHL subtypes

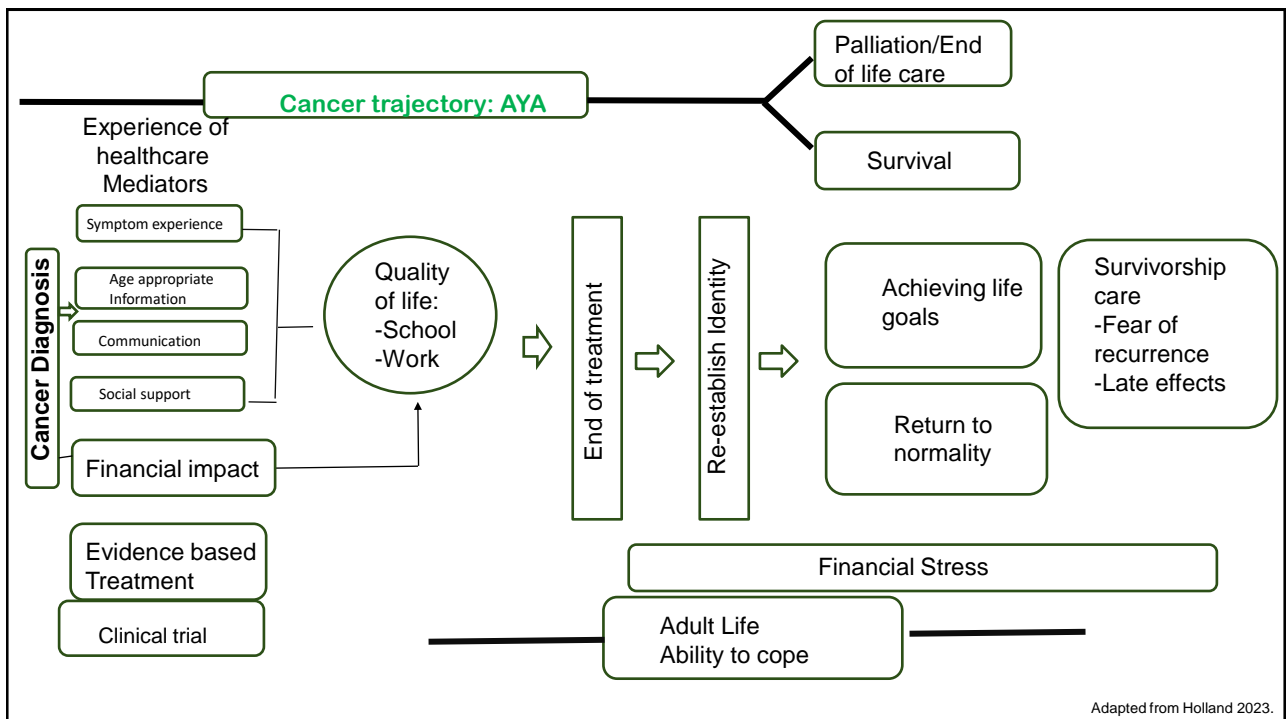
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## Drivers of Disparities in AYA Lymphoma

- Timely Access to care
  - Lack of insurance; underinsurance; non-continuous coverage
  - Distance to care
  - Delays in diagnosis
  - Lack of access to AYA resourced care (i.e. cancer centers)
  - Lack of enrollment to clinical trials
- Non-White Race
- Social determinants of health
- Lack of guideline concordant care through the continuum of post treatment/survivorship
- Unmet psychosocial needs → impact adherence to therapy

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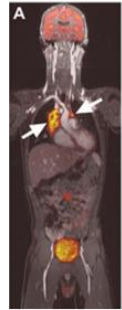
## Diagnostic Workup: Lymphoma in AYA

### Essential for Lymphoma:

- History and physical
  - B symptoms, lymph node, splenomegaly
- Excisional Node Biopsy
- CBC with differential
- ESR and/or CRP
- Complete metabolic panel
- Echocardiogram
- Chest X-ray: PA and lateral views
- FDG-PET/CT or FDG-PET/MRI

### Additional for NHL:

- LDH; uric acid
- Hepatitis B/C testing
- Bilateral bone marrow
- Lumbar puncture
- Immunodeficiency
- Tissue Diagnostics (not comprehensive)
  - IHC
    - CD20, CD30, Ki-67, Tdt
    - ALK
  - Flow cytometry: surface kappa/lambda
  - FISH
    - MYC, BCL2, BCL6; t(8;14)
  - Microarray: 11 q aberrations



[https://www.nccn.org/professionals/physician\\_gls/pdf/b-cell.pdf](https://www.nccn.org/professionals/physician_gls/pdf/b-cell.pdf)

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## AYA Specific Considerations at Diagnosis of Lymphoma

- Pulmonary function test
- HIV
- Health Insurance
- Sexual health assessment
- Pregnancy test
- Fertility preservation-discussion and services
- Psychosocial assessment
- Counseling on substance use and smoking cessation
- Work/school issues
- Social support/network
- Financial toxicity

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## AYA Lymphoma: Goals

- Balancing risk of relapse against:
  - Acute toxicity
  - Late toxicity
  - Quality of life
- During Therapy:
  - Understand tolerability through Patient Reported Outcomes (PRO)
  - Understand how symptomatic and non symptomatic AEs contribute to adherence to dose intensity of therapy



Popali et al. sJHaem 2023.

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## Hodgkin Lymphoma: Peds vs. Adult Oncology Approaches

- Histology distribution varies: younger patients; race/ethnicity
- Risk classification
  - Bulk – definitions differ between adults and peds
  - Prognostic scores- created in older adult cohorts treated with conventional therapy
- Treatment approaches: Risk based, response adapted
  - Chemotherapy back-bone ( ABVE-PC vs. ABVD/BEACOPP)
  - Combined modality
    - Tailored radiation use and dose in older adolescents and YAs
- Trial Endpoints (EFS) – events include subsequent malignant neoplasms (SMN)
  - Goals of care: Person-years of life considered, HRQL
  - Late effects: Cardiac; fertility

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### Survival by age in paediatric and adolescent patients with Hodgkin lymphoma: a retrospective pooled analysis of children's oncology group trials

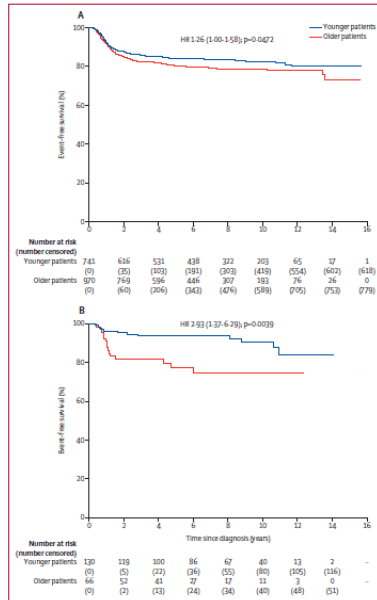
Justine M Kahn, Qinglin Pei, Debra L Friedman, Joel Kaplan, Frank G Keller, David Hodgson, Yue Wu, Burton E Appel, Smrita Bhatia, Tara O Henderson, Cindy L Schwartz, Kara M Kelly, Sharon M Castellino

Supplementary Table 3: Age ≥12 years. Multivariable model of event-free survival (EFS) and overall survival (OS) in N=1,711 patients with non-MC histology treated for HL on Children's Oncology Group (COG) trials (2002 – 2012).

Variables	EFS			OS		
	HR	95% CI	p-value	HR	95% CI	p-value
Age (R: <12 years)						
≥12	1.48	1.03, 2.13	0.03	1.99	0.69, 5.70	0.20
Sex (R: Female)						
Male	1.14	0.90, 1.44	0.28	1.37	0.73, 2.59	0.33
Race (R: Non-Hispanic White)						
Non-Hispanic Black	0.88	0.58, 1.33	0.65	2.19	0.93, 5.17	0.25
Hispanic	1.20	0.86, 1.68		2.08	0.93, 4.67	
Asian/Pacific Island	1.31	0.69, 2.48		0.98	0.13, 7.35	
Other	0.98	0.56, 1.75		0.76	0.10, 5.74	
Study (R: AHOD0031)						
AHOD0431	3.47	2.10, 5.75	<0.0001	*		0.94
AHOD0831	0.48	0.27, 0.85		0.78	0.19, 3.22	
Stage (R: I and II)						
III	1.39	0.93, 2.08	0.0001	0.89	0.32, 2.43	0.94
IV	2.30	1.56, 3.40		1.07	0.37, 3.12	
B symptoms (R: No)						
Yes	2.04	1.44, 2.90	<0.0001	1.93	0.86, 4.33	0.11
Bulky Disease (R: No)						
Yes	2.06	1.44, 2.95	<0.0001	1.59	0.66, 3.85	0.30
Radiation (R: Yes)						
No	1.53	1.18, 1.98	0.001	1.10	0.53, 2.28	0.79
Payment (R: Private)						
Government	0.85	0.65, 1.11	0.22	0.84	0.42, 1.70	0.94
Other + Unknown	1.17	0.71, 1.93		0.80	0.18, 3.48	
Self-Pay or None	0.49	0.20, 1.22		0.67	0.09, 5.03	

Abbreviations: MC: Mixed cellularity; HL: Hodgkin lymphoma; HR: Hazard ratio; CI: confidence interval; cHL, NOS: Classical Hodgkin lymphoma, not-otherwise-specified

Kahn et al. Lancet Haematol 2022.



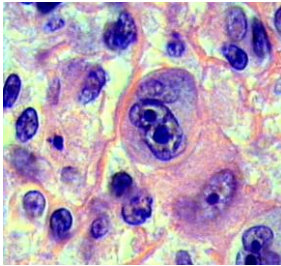
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## Collaboration → Accelerate Novel Approaches ... and AYA enrollment

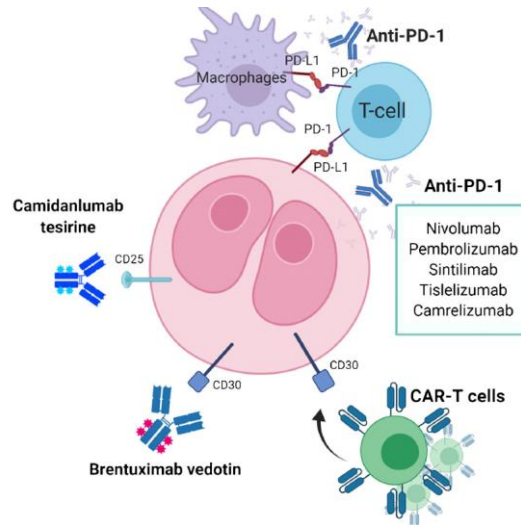
- NCTN (National Clinical Trials Network) - launched 2014
  - Goal: Increase trial participation in rare cancers and in AYA
    - Central support: CTSU; NCI CIRB
  - Increase in phase 3 trials
  - Increase in AYA enrollment ; 9.5% → 14.0%
- Pharma
  - > 10 years (avg.) between regulatory approval and labeling of innovative therapy for adults and children
  - Prolonged off label use in pediatric patients
- International and other consortium partnerships

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## HL: Exploiting biology of HRS cell and the Tumor Microenvironment



HRS = Reed-Sternberg cell



Andrade-Gonzalez, Ansell. Curr Treat Options Oncol. 2021.

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“A Phase III, Randomized Study of Nivolumab (Opdivo) Plus AVD or Brentuximab Vedotin (Adcetris) Plus AVD in Patients (**Age  $\geq$  12 years**) with Newly Diagnosed Advanced Stage Classical Hodgkin Lymphoma.”

**S1826 (NCT03907488)– Activated 7/19/2019**

SWOG Chairs: Alex Herrera, MD; Jonathan Friedberg MD, MMSc  
 Pediatrics/COG Chair: Sharon Castellino, MD, MSc COG Champion: Angela Punnett MD  
 QOL Chair: Susan Parsons, MD, MRP

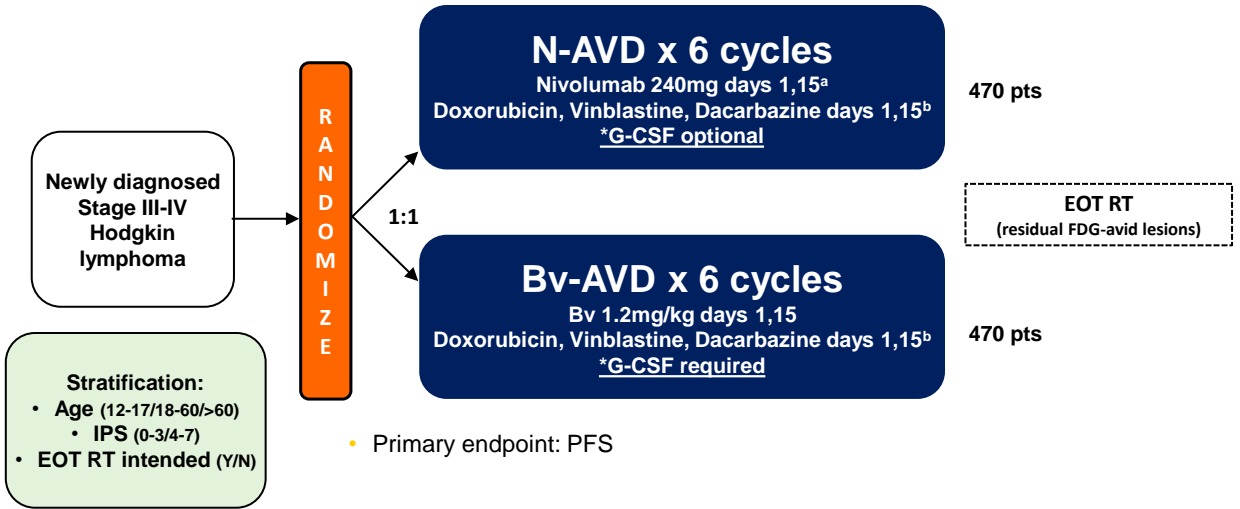


Herrera A. JCO 41, no. 17\_suppl (June 10, 2023)  
 LBA4-LBA4.



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## S1826 Study Design



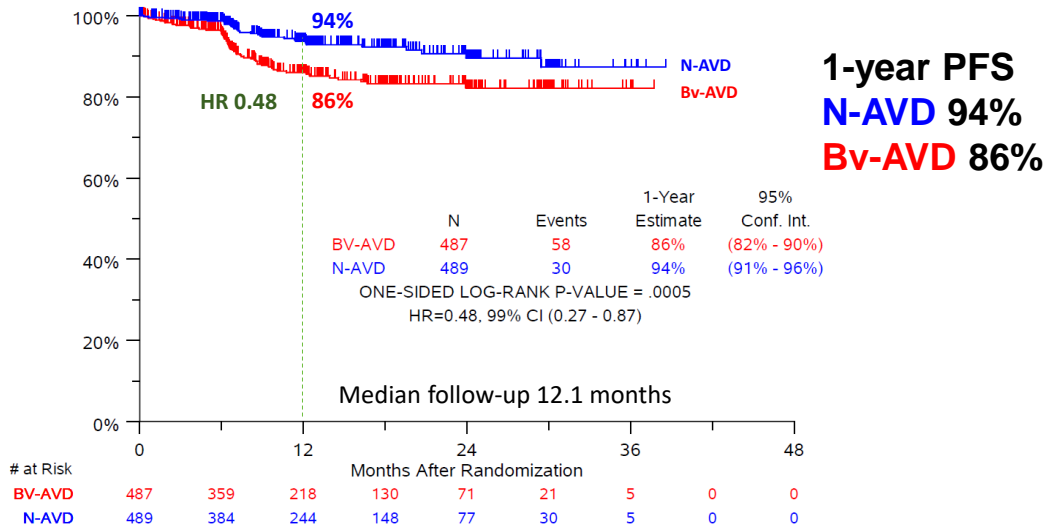
<sup>a</sup> Nivolumab 3mg/kg for ages ≤ 17, max 240mg  
<sup>b</sup> Conventional doses of AVD: Stephens DM et al Blood 2019, Ansell SM et al NEJM 2022

Alex F. Herrera, MD ASCO 2023.



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## N-AVD improves PFS compared to Bv-AVD



Alex F. Herrera, ASCO 2023.



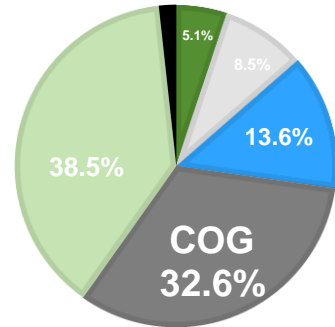
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## Successful Collaboration with the Adult NCTN

- ❑ Earlier access to novel agents for adolescents
- ❑ Harmonize approaches across pediatric and adult providers for AYAs with advanced stage HL
- ❑ Parallel design: Compare Bv-AVD against Bv-AVEPC (AHOD1331)
- ❑ Evaluation of the role of RT in the setting of new agents
- ❑ PROs will facilitate measurement of tolerability of new agents across the age spectrum

**S1826 Accrual (n=994)  
(enrollment closed Oct 5, 2022)**

ACCRUAL BY GROUP



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## Cumulative Chemotherapy Dosing

	AHOD1331* (BV-AVE-PC x 5)	S1826 (N-AVD x 6)
Brentuximab Vedotin	9 mg/kg	
Nivolumab		36 mg/kg
Adriamycin	250 mg/m <sup>2</sup>	300 mg/m <sup>2</sup>
Vincristine	7 mg/m <sup>2</sup>	
Vinblastine		72 mg/m <sup>2</sup>
Etoposide	1875 mg/m <sup>2</sup>	
Prednisone	1400 mg/m <sup>2</sup>	
Cyclophosphamide	6000 mg/m <sup>2</sup>	
Dacarbazine		4500 mg/m <sup>2</sup>
Radiation dose	21 Gy 9 Gy boost to sites of residual avidity on EOT PET	30-36 Gy



\*S. Castellino et al. NEJM 2022

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## Chemotherapy Administration

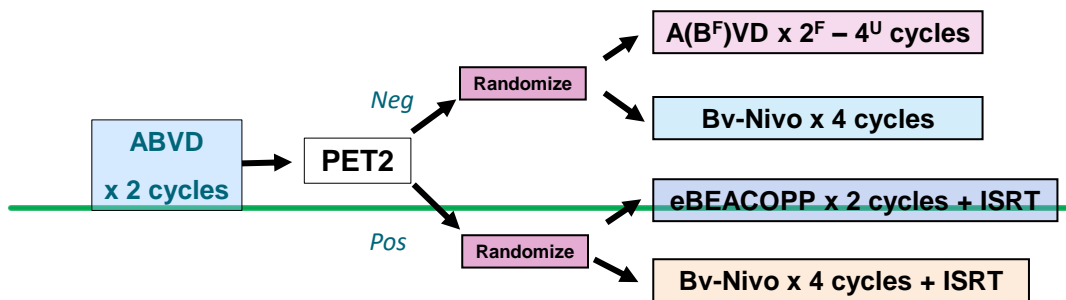
	AHOD1331 (BV-AVE-PC x 5)	S1826 (N-AVD x 6)
Cycle Length	21 days	28 days
Total Duration	105 days	168 days
Days of IV chemo	Day 1, 2, 3, and 8	Day 1 and 15
Total days of IV chemo	20 days	12 days
Growth Factor	Required	Optional
Dexrazoxane	Permitted not required	Permitted not required

CHILDREN'S  
ONCOLOGY  
GROUP

Courtesy: M. Heneghan

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## AHOD2131 (NCT05675410): Stage I/II – Activated April 2023



<sup>a</sup> 1 cycle = 28 days  
<sup>b</sup> PET2 positive defined as Deauville 4 or 5  
 F favorable; U unfavorable

Study Chairs: T. Henderson; K. Kelly; B. Hu

CHILDREN'S  
ONCOLOGY  
GROUP

NCI National Clinical  
Trials Network  
A National Cancer Institute program

NCI Community Oncology  
Research Program  
A program of the National Cancer Institute

Alliance  
for Clinical Trials  
in Oncology

ECOG-ACRIN  
Cancer Research Group  
Reshaping the future of solid tumor care

SWOG  
Leading cancer research. Together.

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## Non-Hodgkin Lymphoma in AYA

- **More Common Pediatric/Adolescent NHL**
  - Mature B-cell lymphomas
    - Diffuse Large B-cell Lymphoma
    - Burkitt Lymphoma
    - Primary Mediastinal B-cell Lymphoma
  - Anaplastic Large Cell Lymphoma
  - Lymphoblastic Lymphoma/Leukemia
    - T differentiation
    - B differentiation
  - Post-transplant lymphoproliferative disease (PTLD)
- **Less Common Pediatric/Adolescent NHL**
  - Pediatric follicular lymphoma
  - Marginal zone & MALT lymphoma
  - Primary CNS lymphoma
  - Peripheral T-cell lymphoma NOS
- **Lack of harmonization in staging systems in NHL**
  - Ann Arbor Staging (adults)
  - International Pediatric NHL Staging System
  - Lack of Prognostic scores relevant to younger patients

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## Novel Agents in combination with chemotherapy in Frontline Regimens for NHL

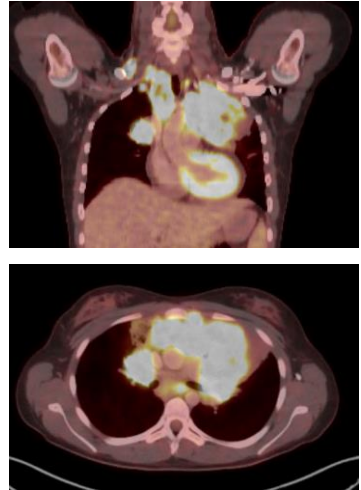
- Anti-CD 30: Brentuximab vedotin
- Anti-CD20: Rituximab
- ALK inhibitor: crizotinib
- Amplified PD1- Checkpoint inhibitors
- Anti-CD 79b –Polatuzumab vedotin
- Bruton tyrosine kinase inhibitor : ibrutinib
- ☐ AYA with a mature B cell lymphoma could receive vastly different therapy depending on point of presentation (adult vs. pediatric provider)
  - ☐ Providers are encouraged to check the NCCN guidelines and to consider offering a clinical trial
- ☐ Many emerging novel agents for NHL are in trial in relapsed setting
- ☐ Most have undergone relatively little study in AYA

*EI-Mallawany et al. eJHaem 2023*

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## Primary Mediastinal B-cell Lymphoma

- Rare subtype of NHL
- Peak incidence in AYA, F>M
- Presents as large mediastinal mass
  - Pleural, pericardial effusions common
- Biology overlaps with classic HL
  - CD30+
  - Overexpression PD-1
  - Sensitive to immune checkpoint blockade



CHILDREN'S  
ONCOLOGY  
GROUP

Courtesy : L Giulino-Roth

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## ANHL1931 (NCT04759586) : Randomized phase III trial of nivolumab in PMBCL

Physician declares  
chemotherapy backbone:  
R-CHOP or DA-EPOCH-R

R-CHOP or  
DA-EPOCH-R  
x 6 cycles

Nivo + R-CHOP or  
Nivo + DA-EPOCH-R  
x 6 cycles

Primary Endpoint: PFS as determined by  
independent review committee

Consolidative RT permitted only  
in the following circumstances:  
1) Physician declares R-CHOP +  
RT regardless of EOT imaging  
2) + biopsy at EOT

**Open NCTN wide  
across all age groups**

Opened to accrual June 2021  
Anticipated to enroll 186 patients  
over 3.8 years

Courtesy: L G Roth.

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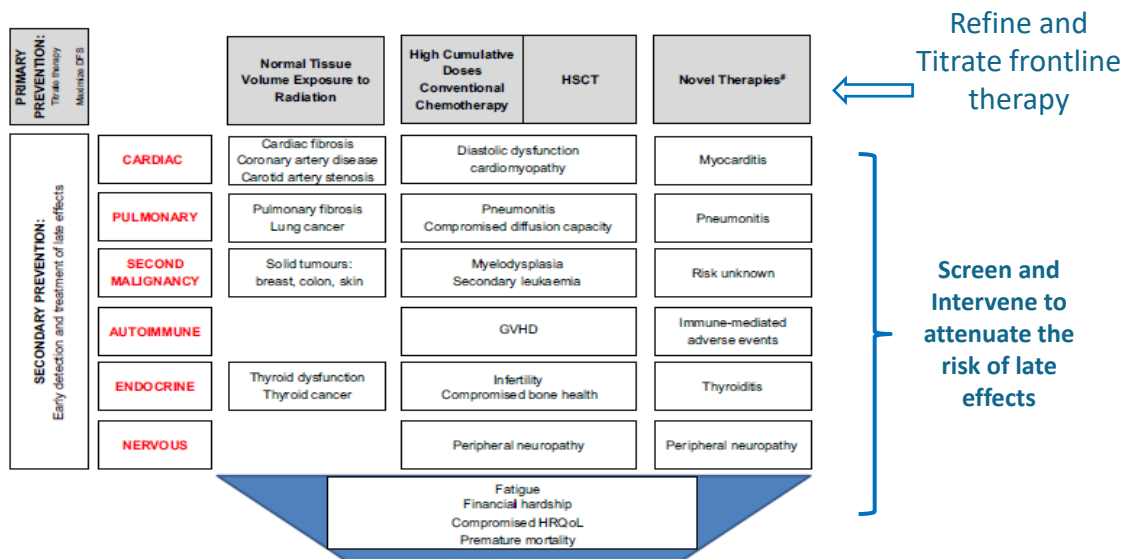
## POLLING QUESTION #5

- What percent of AYA patients with a blood cancer should receive a survivorship care plan?
  - A) 11-25%
  - B) 26-50%
  - C) 50%
  - D) 100%

Aflac Cancer and Blood Disorders Center | Emory University.

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## Addressing the Survivorship Gap... Begin With the End in Mind



Castellino SM, et al. *Br J Haematol.* 2019;187:573.

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## Secondary Prevention

### Children's Oncology Group Guidelines: Exposure based

**CHILDREN'S ONCOLOGY GROUP**  
The world's childhood cancer experts



**OUR MISSION**  
To cure and prevent childhood and adolescent cancer through scientific discovery and compassionate care.

**IMPORTANT: COVID-19 INFORMATION FOR SURVIVORS OF CHILDHOOD, ADOLESCENT, AND YOUNG ADULT CANCERS**  
Download here: [COVID-19 \(English\)](#) [COVID-19 \(Spanish\)](#) [COVID-19 \(French\)](#)

Long-Term Follow-Up Guidelines for Survivors of Childhood, Adolescent, and Young Adult Cancers  
Version 5.0 (October 2018)  
New: Chinese Health Links (in Traditional [TC] and Simplified [SC] Chinese)

The Children's Oncology Group Long-Term Follow-Up Guidelines for Survivors of Childhood, Adolescent, and Young Adult Cancers (COG LTU Guidelines) are a resource for healthcare professionals who provide ongoing care to survivors of pediatric malignancies. The screening recommendations in these guidelines are appropriate for asymptomatic survivors of childhood, adolescent, or young adult cancer presenting

- Need for new models for AYAs who often do not return to treating institution
- Opportunities for mHealth
- Evidence will be a long time in coming for novel agents
- Need biomarkers for late toxicity

### NCCN National Comprehensive Cancer Network® NCCN Guidelines Version 1.202 Survivorship

NCCN Survivorship Panel Members  
NCCN Survivorship Sub-Committee Members  
[Summary of the Guidelines Updates](#)

#### General Survivorship Principles

- [Definition of Survivorship \(SURV-1\)](#)
- [Standards for Survivorship Care \(SURV-2\)](#)
- [General Principles of the Survivorship Guidelines \(SURV-3\)](#)
- [Screening for Subsequent New Primary Cancers \(SURV-4\)](#)
- [Principles of Cancer Risk Assessment and Counseling \(SURV-5\)](#)
- [Assessment by Health Care Provider at Regular Intervals \(SURV-6\)](#)
- [Survivorship Assessment \(SURV-4\)](#)
- [Survivorship Resources for Health Care Professionals and Survivors \(SURV-B\)](#)
- [Principles of Screening for Treatment-Related Subsequent Primary Cancers \(SURV-C\)](#)

#### Preventive Health

- [Healthy Lifestyles \(HL-1\)](#)
- [Physical Activity \(SPA-1\)](#)
- [Nutrition and Weight Management \(SNWM-1\)](#)
- [Supplement Use \(SUSE-1\)](#)
- [Immunizations and Infections \(SIMIN-1\)](#)

#### Late Effects/Long-Term Psychosocial and Physical Problems

- [Cardiovascular Disease Risk Assessment \(SCVD-1\)](#)
- [Antitumor-Induced Cardiac Toxicity \(SCABDIO-1\)](#)
- [Anxiety, Depression, Trauma, and Distress \(SANXDE-1\)](#)
- [Cognitive Function \(SCF-1\)](#)
- [Fatigue \(SFAT-1\)](#)
- [Lymphedema \(SLYMPH-1\)](#)
- [Pain \(SPAIN-1\)](#)
- [Hormone-Related Symptoms \(SHRS-1\)](#)
- [Sexual Health \(SSH-1\)](#)
- [Fertility \(SF-1\)](#)
- [Sleep Disorders \(SSD-1\)](#)
- [Employment and Return to Work \(SWORK-1\)](#)

<http://www.survivorshipguidelines.org/>

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## Survivorship Care Plans

- Document that summarizes an individual patient's treatment-cumulative doses and modalities of therapy received
- Summary of :
  - Therapy associated late effects
  - Recommendations for follow-up care
  - Health promotion for screening and health behaviors

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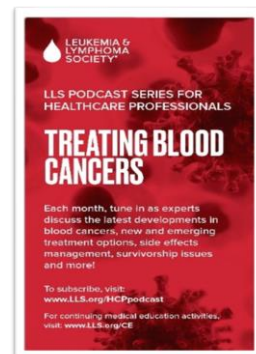
## Thank You



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### FREE LLS RESOURCES FOR HEALTHCARE PROVIDERS

- ❑ CME and CE courses: [www.LLS.org/CE](http://www.LLS.org/CE)
- ❑ Fact Sheets for HCPs: [www.LLS.org/HCPbooklets](http://www.LLS.org/HCPbooklets)
- ❑ Videos for HCPs: [www.LLS.org/HCPvideos](http://www.LLS.org/HCPvideos)
- ❑ Podcast series for HCPs: [www.LLS.org/HCPpodcast](http://www.LLS.org/HCPpodcast)



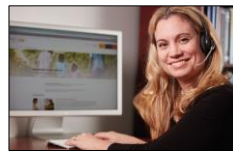
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## FREE LLS RESOURCES FOR PATIENTS

- ❑ **Information Specialists** – Personalized assistance for managing treatment decisions, side effects, and dealing with financial and psychosocial challenges (IRC).
- ❑ **Clinical Trial Nurse Navigators** – RNs and NPs provide a personalized service for patients seeking treatment in a clinical trial, sift through the information and provide information to bring back to their HC team (CTSC).
  - [www.LLS.org/CTSC](http://www.LLS.org/CTSC)
- ❑ **Nutrition Education Services Center (NESC)** – LLS provides **Nutrition Education Services** to patients and caregivers of all cancer types. *Our registered dietitians have expertise in oncology nutrition.* To schedule a free consultation:
  - visit [www.LLSnutrition.org](http://www.LLSnutrition.org)
  - call 800-955-4572

### ❑ Reach out Monday–Friday, 9 am to 9 pm ET

- Phone: (800) 955-4572
- Live chat: [www.LLS.org/IRC](http://www.LLS.org/IRC)
- Email: [infocenter@LLS.org](mailto:infocenter@LLS.org)
- HCP Patient Referral Form: [www.LLS.org/HCPreferral](http://www.LLS.org/HCPreferral)



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## FREE LLS RESOURCES FOR PATIENTS AND CAREGIVERS

### ❑ Webcasts, Videos, Podcasts:

- [www.LLS.org/Webcasts](http://www.LLS.org/Webcasts)
- [www.LLS.org/EducationVideos](http://www.LLS.org/EducationVideos)
- [www.LLS.org/Podcast](http://www.LLS.org/Podcast)

### ❑ [www.LLS.org/youngadults](http://www.LLS.org/youngadults)

### ❑ Support Resources

- ❑ Financial Assistance: [www.LLS.org/Finances](http://www.LLS.org/Finances)
- ❑ Other Support: [www.LLS.org/Support](http://www.LLS.org/Support)
  - LLS Regions
  - Live Online Weekly Chats: "Living with NHL"
    - Facilitated by Oncology SW
  - LLS Community Social Media Platform
  - First Connection Peer to Peer Program



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## FREE LLS RESOURCES FOR YOUR PATIENTS



### BOOKLETS AND FACT SHEETS

English – [www.LLS.org/Booklets](http://www.LLS.org/Booklets)

Spanish – [www.LLS.org/Materiales](http://www.LLS.org/Materiales)



## Questions?



Ask a question by **web**:

- Click “Ask a question”
- Type your question
- Click “Submit”





## INSTRUCTIONS FOR CREDIT

Participants must complete the evaluation to receive credit.  
After completing this process, your certificate will automatically generate.

Link to complete evaluation: <https://mli.link/lisaya>

For questions or concerns, please contact [Profeducation@lls.org](mailto:Profeducation@lls.org)



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# THANK YOU!

We have one goal: A world without blood cancers



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