



Recent Updates in
Pediatric and Adolescent
Young Adult (AYA) Acute
Lymphocytic Leukemia (ALL)

Webinar

Sponsors:

Platinum

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Welcome and Introductions

Rupert Handgretinger, MD
Children's University Hospital and
Department of Hematology/Oncology
Germany



Agenda

Time	Topic	Presenter
2.00 PM – 2.05 PM CDT/GMT-5 3.00 PM – 3.05 PM EDT 4.00 PM – 4.05 PM BRT 21.00 – 21.05 CET	Welcome and Introductions	Rupert Handgretinger, MD
2.05 PM – 2.15 PM CDT/GMT-5 3.05 PM – 3.15 PM EDT 4.05 PM – 4.15 PM BRT 21.05 – 21.15 CET	 Current Paradigm and Long-Term Toxicities for Pediatric ALL Integration of innovative immunotherapies Role of MRD in treatment Long-term toxicities 	Franco Locatelli, MD
2.15 PM – 2.30 PM CDT/GMT-5 3.15 PM – 3.30 PM EDT 4.15 PM – 4.30 PM BRT 2 21.15 – 21.30 CET	 Bispecifics for Pediatric/AYA ALL Review of trial results in pediatric/AYA ALL Role of MRD in research and treatment AYA considerations 	Patrick Brown, MD
2.30 PM – 2.40 PM CDT/GMT-5 3.30 PM – 3.40 PM EDT 4.30 PM – 4.40 PM BRT 21.30 – 21.40 CET	 CAR T Cells for Pediatric/AYA ALL Benefits and risks of CAR Ts and bispecifics Role of MRD in research and treatment AYA considerations 	Franco Locatelli, MD
2.40 PM - 3.00 PM CDT/GMT-5 3.40 PM - 4.00 PM EDT 4.40 PM - 5.00 PM BRT 21.40 - 22.00 CET	Questions to Experts	Rupert Handgretinger, MD







Current Paradigm and Long-Term Toxicities for Pediatric ALL

Franco Locatelli, MD

University of RomelRCCS Ospedale Pediatrico Bambino Gesù of Rome Italy







Current Paradigm and Long-Term Toxicities for Pediatric ALL

Franco Locatelli, MD Università Sapienza, Roma

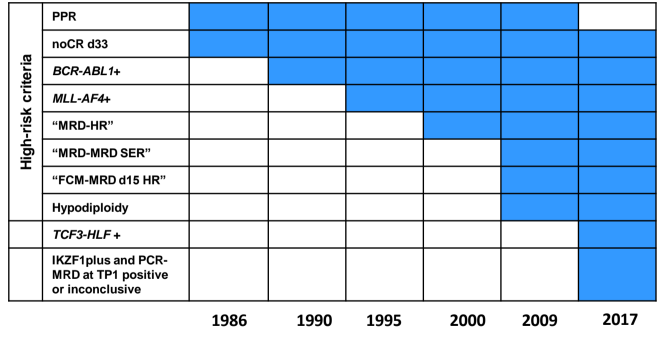
Dept. Pediatric Hematology/Oncology and Cell/Gene Therapy



The essentials in pediatric ALL: Risk stratification and therapy

- Approximately 80% 5-year EFS can be achieved in unselected populations of pediatric patients
- The early treatment response in particular through MRD detection has been established to be the strongest prognostic factor
- New molecular subgroups have been described (eg, Ph-like or BCR/ABL-like pB-ALL;
 MPAL) and their prognostic role defined
- Translation of novel molecular findings into improved treatment outcome is under investigation in various trials
- Reduction of long-term toxicities, especially in adolescents, is a priority
- Novel treatment approaches based on immunotherapy; evidence regarding long-term benefit is yet to be established

Identification of new high-risk groups and reducing relapses in high-risk patients

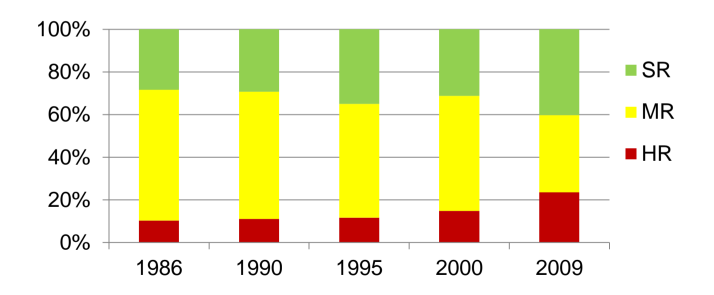


Studies ALL-BFM

More and more patients with "intermediately unfavorable" outcome have been identified and shifted to the high-risk arm

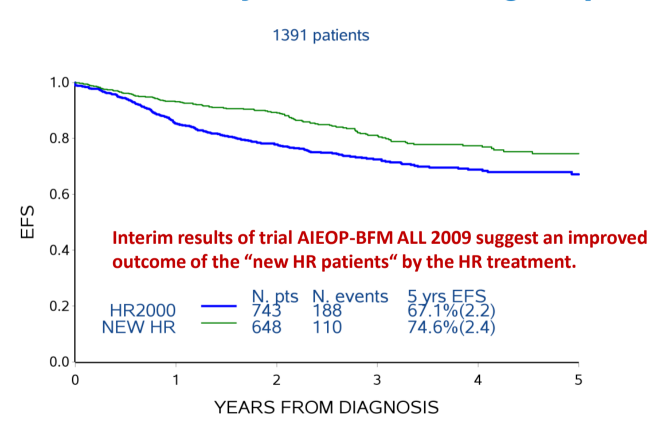
Identification of new high-risk groups and reducing relapses in high-risk patients

→ In AIEOP-BFM ALL 2009, the HR group comprised >20% of the patients



Studies ALL-BFM

AIEOP-BFM ALL 2009 – Interim analysis of the HR group



IKZF1^{plus} Defines a New Minimal Residual Disease–Dependent Very-Poor Prognostic Profile in Pediatric B-Cell Precursor Acute Lymphoblastic Leukemia

Martin Stanulla, Elif Dagdan, Marketa Zaliova, Anja Möricke, Chiara Palmi, Giovanni Cazzaniga, Cornelia Eckert, Geertruy te Kronnie, Jean-Pierre Bourquin, Beat Bornhauser, Rolf Koehler, Claus R. Bartram, Wolf-Dieter Ludwig, Kirsten Bleckmann, Stefanie Groeneveld-Krentz, Denis Schewe, Stefanie V. Junk, Laura Hinze, Norman Klein, Christian P. Kratz, Andrea Biondi, Arndt Borkhardt, Andreas Kulozik, Martina U. Muckenthaler, Giuseppe Basso, Maria Grazia Valsecchi, Shai Izraeli, Britt-Sabina Petersen, Andre Franke, Petra Dörge, Doris Steinemann, Oskar A. Haas, Renate Panzer-Grümayer, Hélène Cavé, Richard S. Houlston, Gunnar Cario, Martin Schrappe, and Martin Zimmermann, for the TRANSCALL Consortium and the International BFM Study Group

DOI: https://doi.org/10.1200/JCO.2017.

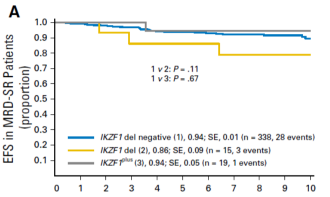
74.3617

New prognostic pattern: Definition of IKZF1^{plus}

- Deletion of IKZF1 and
 - PAX5 and/or
 - CDKN2A and/or
 - CDKN2B and/or
 - CRLF2 (PAR) and

Negativity for *ERG* deletion

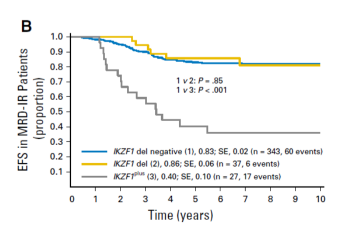
IKZF1^{plus} and MRD: Impact on EFS

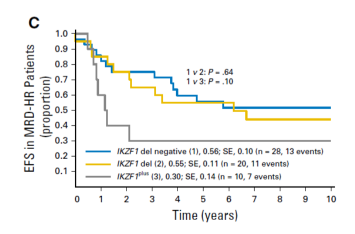


A: MRD – Standard risk (MRD neg at 5w and 12w)

B: MRD – Intermediate risk (MRD non SR/HR)

C: MRD – High risk (MRD pos $\geq 10^{-4}$ at 12w)





Perspectives for new trials in ALL

- Avoid additional toxic agents in most patients
- Utilize novel genetic approaches
- Improve risk stratification by wider combination of genetic factors and response (MRD)
- Introduce novel agents under controlled conditions

AIEOP-BFM ALL 2017

International collaborative treatment protocol for children and adolescents with acute lymphoblastic leukemia

Randomized phase III study conducted by the AIEOP-BFM study group

EudraCT Number: 2016-001935-12

Sponsor: Universitätsklinikum Schleswig-Holstein, Campus Kiel

New in trial AIEOP-BFM ALL 2017

- Modified workflow and timing in genetic diagnostics
- Genetic profiles and early MRD response may be combined to characterize previously not identified pts at high risk to relapse, eg, IKZF1^{plus}
- Randomized evaluation of blinatumomab in de novo ALL in all non-SR patients
- Selective addition of novel agents in HR group
- Limitation of pCRT (only if age ≥4y, only if CNS-3, and/or if T-ALL with WBC ≥100K)
- TDM for ASP activity only in reintensification (P-II, P-III, HR-1/2/3)

AIEOP-BFM ALL 2017 – Risk criteria for pB-ALL MR and HR

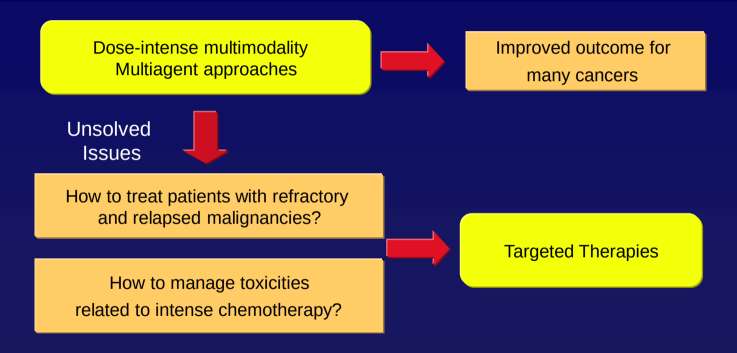
High Risk (HR)

- No complete remission on day 33 *or*
- Positivity for KMT2A-AFF1 or
- Positivity for *TCF3-HLF or*
- Hypodiploidy <45 chromosomes or
- FCM-MRD in BM on day 15 ≥10% and <u>not</u> ETV6-RUNX1 positive or
- IKZF1^{plus} and PCR-MRD at TP1 positive or inconclusive and <u>not</u> positive for ETV6-RUNX1, TCF3-PBX1 or KMT2A rearr. other than KMT2A-AFF1 or
- PCR-MRD at TP1 $\geq 5 \times 10^{-4}$ and positive $< 5 \times 10^{-4}$ at TP2 (PCR-MRD SER)
- PCR-MRD at TP2 \geq 5×10⁻⁴ (PCR-MRD-HR)
- Age <1 year and any KMT2A rearrangement

Medium Risk (MR)

- No HR criteria and
- PCR-MRD either positive at TP1 and/or TP2 or PCR-MRD not evaluable

Why immunotherapy for childhood tumors?

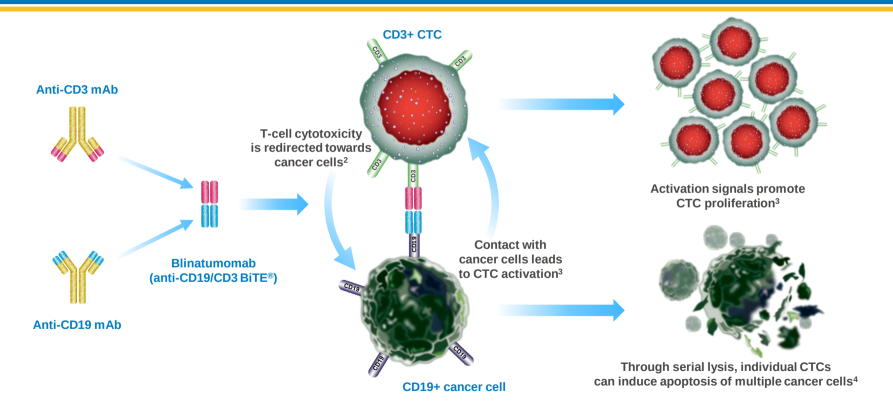


Immunotherapy for de novo pediatric ALL in trial AIEOP-BFM ALL 2017

• Anti-CD19/anti-CD3 (Blinatumomab/BiTE®): prospective evaluation in MR and HR patients

Allogeneic hSCT in predefined subgroups

The BiTE® blinatumomab: Designed to bridge cytotoxic T cells (CTCs) to CD19-expressing cancer cells, resulting in cancer cell death¹



BiTE®, bispecific T-cell engager; mAb, monoclonal antibody.

1. Baeuerle PA, Reinhardt C. Cancer Res. 2009;69:4941-4944; 2. Bargou R, et al. Science. 2008;321:974-977; 3. Klinger M, et al. Blood. 2012;119:6226-6233; 4. Hoffmann P, et al. Int J Cancer. 2005;115:98-104.

Blinatumomab activity and toxicity in ALL

- Antileukemic activity demonstrated in both adults and children
- 80% MRD negativity in ALL patients treated in hematological CR but with molecularly-resistant disease
- Toxicity in patients without overt disease limited
- Toxicity profile different from chemotherapy

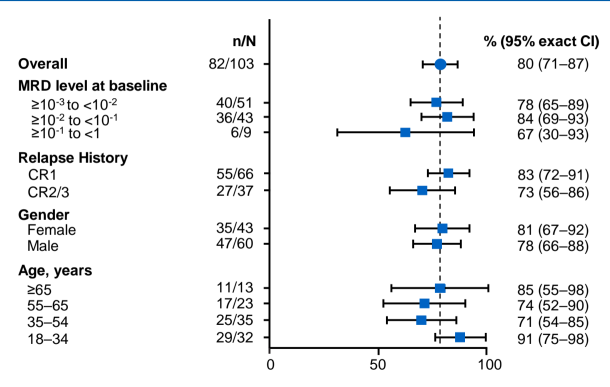
Phase I/Phase II Study of Blinatumomab in Pediatric Patients With Relapsed/Refractory Acute Lymphoblastic Leukemia

Arend von Stackelberg, Franco Locatelli, Gerhard Zugmaier, Rupert Handgretinger, Tanya M. Trippett, Carmelo Rizzari, Peter Bader, Maureen M. O'Brien, Benoît Brethon, Deepa Bhojwani, Paul Gerhardt Schlegel, Arndt Borkhardt, Susan R. Rheingold, Todd Michael Cooper, Christian M. Zwaan, Phillip Barnette, Chiara Messina, Gérard Michel, Steven G. DuBois, Kuolung Hu, Min Zhu, James A. Whitlock, and Lia Gore

VOLUME 34 · NUMBER 36 · DECEMBER 20, 2016

Complete MRD response after cycle 1 by clinical characteristics

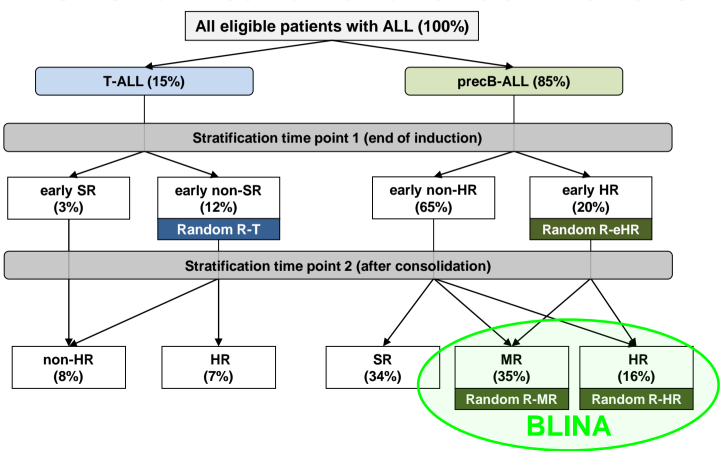
Phase 2 study MT103-203 (BLAST) in adults with MRD-positive B-precursor ALL



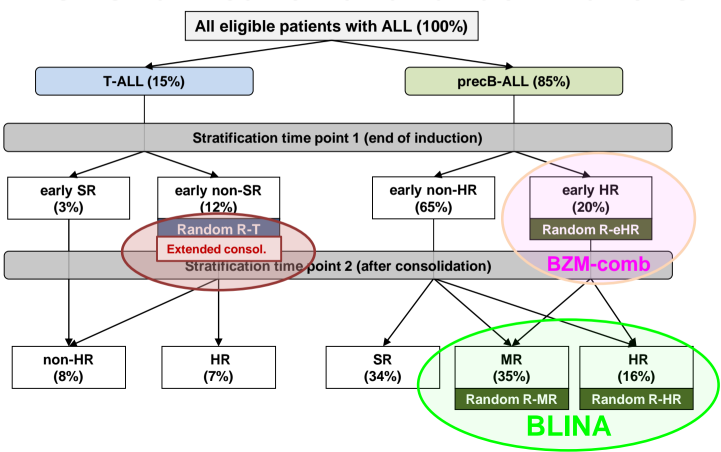
- Complete MRD response (primary endpoint, FAS[†]): 78% (88/113)
- Complete MRD response (efficacy set*): 80% (82/103)

Complete MRD response rate, % (95% CI)

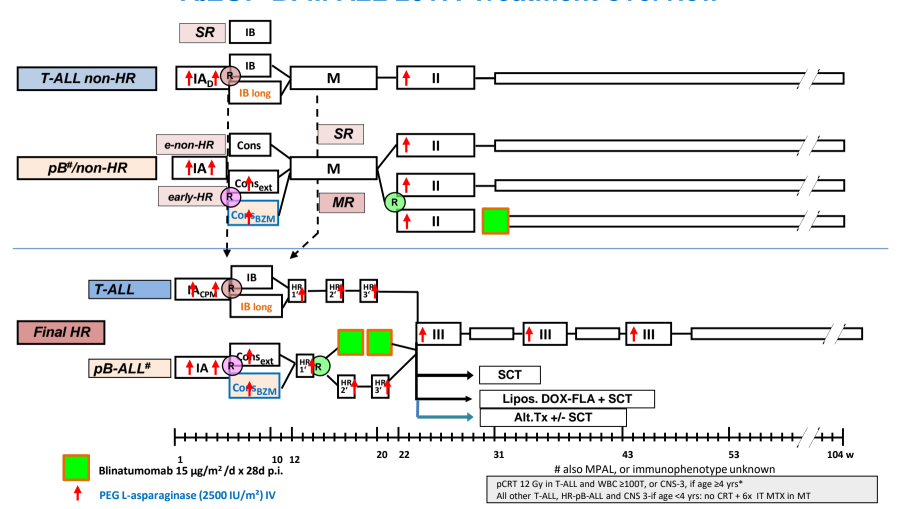
AIEOP-BFM ALL 2017 – Risk stratification and randomizations



AIEOP-BFM ALL 2017 – Risk stratification and randomizations

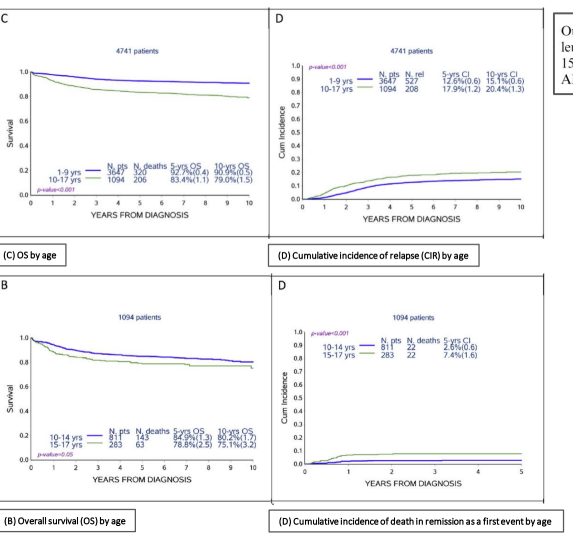


AIEOP-BFM ALL 2017: Treatment overview



A brief focus on adolescents

Acute and late toxicities



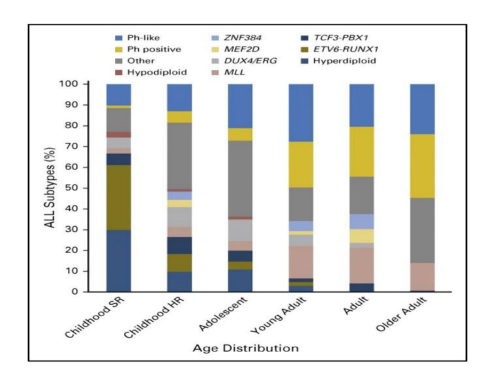
Outcome of adolescent patients with acute lymphoblastic leukaemia aged 10-14 years as compared with those aged 15-17 years: Long-term results of 1094 patients of the AIEOP-BFM ALL 2000 study

European Journal of Cancer 122 (2019) 61-71

ADOLESCENT AND YOUNG ADULT MALIGNANT HEMATOLOGY

Acute lymphoblastic leukemia in adolescent and young adults: treat as adults or as children?

Nicolas Boissel^{1,2} and André Baruchel^{2,3}



		Trial	Median Age	No.	% CR	% Survival		
R		FRALLE93	1993-1999	16	77	94	DFS	72
		INALLESS	1333-1333	10	,,	34	EFS	67
E		LALA94	1994-2000	18	100	83	DFS	49
T		LALAS4	1334-2000	10	100	83	EFS	41
D	6	CCG	1988-2001	16	196	96	EFS	63
R	S	CALGB	1988-2001	19	103	93	EFS	34
O	T	AIEOP	1996-2003	15	150	94	os	80
S	u	GIMEMA	1996-2003	16	95	89	os	71
	U	DOCG	1985-1999	12	47	98	DFS	71
P	D	Doca	1383-1333	12	47	36	EFS	69
E		HOVON	1985-1999	20	73	91	DFS	37
		HOVOIV	1383-1333	20	73	31	EFS	34
C	E	MRC97/99	1997-2002	15-17	61	98	os	71
T	S	WINC57/33	1997-2002	15-17	01	36	EFS	65
		HEALIVII	1997-2002	15-17	67	94	os	56
		UKALLXII	1997-2002	15-17	07	34	EFS	49
V		NOPHO	1990-2004	13	128	96	EFS	67
E		FINNISH LEUKEMIA	1990-2004	19	97	97	EFS	60

	Trial	Trial		No.	% CR	% Survival	
	FRALLE93	1993-1999	16	77	94	DFS	72
FRANCE	THALLESS	1333 1333	10	,,	34	EFS	67
FRANCE	LALA94	1994-2000	18	100	83	DFS	49
	EREAJT	1334-2000			03	EFS	41
UNITED STATES	ccg	1988-2001	16	196	96	EFS	63
	CALGB	1988-2001	19	103	93	EFS	34
	AIEOP	1996-2003	15	150	94	os	80
ITALY	GIMEMA	1996-2003	16	95	89	os	71
	DOCC	1005 1000	42	47	00	DFS	71
	DOCG	1985-1999	12	4/	98	EFS	69
HOLLAND	HOVON	VON 1985-1999	20	73	91	DFS	37
	HOVON	1985-1999	20	/3		EFS	34
	MRC97/99	1997-2002	15-17	61	98	os	71
UNITED						EFS	65
KINGDOM	HIVALLYII	1997-2002	15-17	67	94	os	56
	UKALLXII					EFS	49
	NOPHO	1990-2004	13	128	96	EFS	67
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UNITED STATES	CCG	1988-2001	16	196	96	EFS EFS	41 63	
ONITED STATES								
	AIEOP	1996- 2003	15	15	94		OS	80
ITALY								
	GIMEMA	1996-	16	9	5 89		OS	71
		2003				DEC	37	
HOLLAND	HOVON	1985-1999	20	73	91	DFS EFS	34	
						os	71	
UNITED KINGDOM	MRC97/99	1997-2002	15-17	61	98	EFS	65	
		4007.0000	45.45	67	0.4	os	56	
	UKALLXII	1997-2002	15-17	67	94	EFS	49	
FINLAND	NOPHO	1990-2004	13	128	96	EFS	67	
	FINNISH LEUKEMIA	1990-2004	19	97	97	EFS	60	

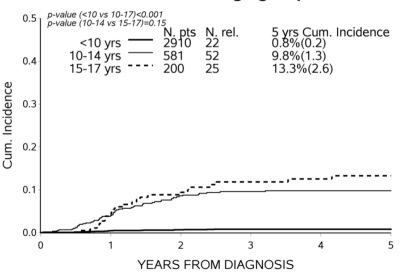
Support as adults, as children, or as AYAs?

- Adolescents, considered high-risk patients per se by pediatricians, for a long time have been considered good-risk patients when seen by adult hematologists
- Most of the studies have shown that the induction death rate, death in remission, delays in chemotherapy administration, and occurrence of severe adverse events are higher in the AYA population compared with children, thus affecting survival as well as quality of life
- Increasing toxicities with age were reported in almost all cohorts of patients treated with fully pediatric or pediatric-like approaches, including in younger age ranges
- The use of intensified regimens and the following improvements in survival in AYAs with ALL raise the need for monitoring and preventing acute and late effects in this population of patients

Correspondence: Osteonecrosis in childhood acute lymphoblastic leukemia: a retrospective cohort study of the Italian Association of Pediatric Haemato-Oncology (AIEOP)

Parasole et al. *Blood Cancer Journal* (2018)8:115

a. Overall incidence in the age groups



 $\label{lem:condition} \textbf{Five-year cumulative incidence of ON according to patient's age at ALL diagnosis}$

AVN - CCG 1961

❖7/769 patients <10 years developed AVN – 1%

❖126/1287 patients ≥10 years developed AVN – 9.8%

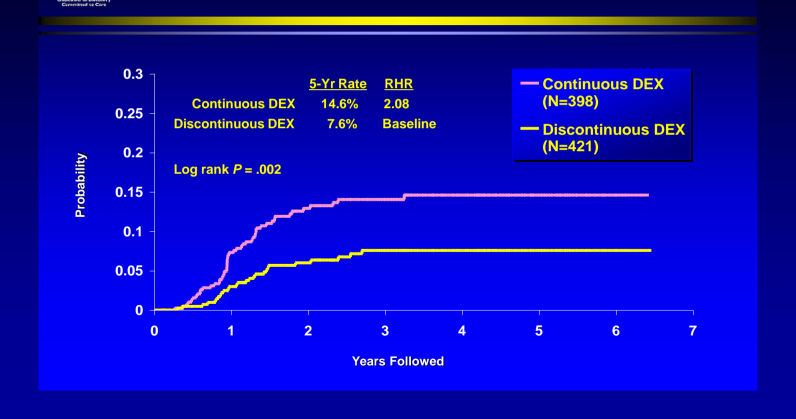
❖10-12 years 32/505 7%

♦13-15 years 53/520 12.6%

❖16+ years 41/262 18.5%

❖Incidence of AVN twice as high in females

CCG-1961 AVN by RER groups (Age 10+ Yrs)







Bispecifics for Pediatric/AYA ALL

Patrick A. Brown, MD

Johns Hopkins University School of Medicine

USA











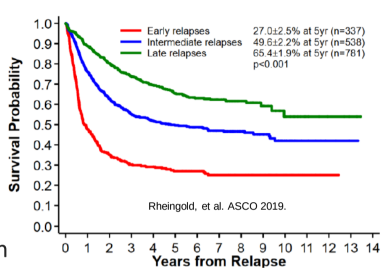
Bispecifics for Pediatric/AYA ALL

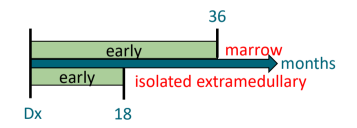
Patrick Brown, MD

Associate Professor of Oncology, Johns Hopkins University
Director, Pediatric Leukemia Program, Sidney Kimmel Comprehensive Cancer Center
Vice Chair for Relapse, COG ALL Committee
Chair, NCCN ALL Guidelines Panel

Background

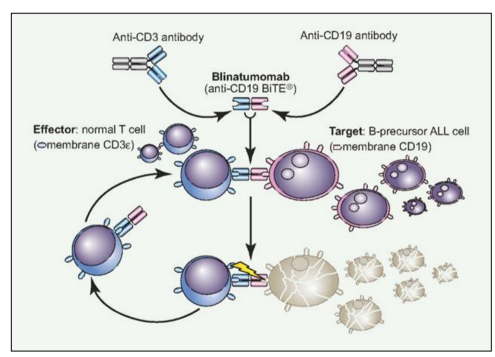
- Poor survival for 1st relapse B-ALL in children, adolescents and young adults (AYA), especially early relapses
- Standard treatment approach
 - Reinduction chemotherapy → 2nd remission
 - Consolidation
 - <u>Early relapse</u>: Intensive chemo → HSCT
 - Goal: MRD-negativity prior to HSCT
 - Late relapse
 - "MRD high": same as early
 - "MRD low": Intensive chemo → maintenance therapy







Blinatumomab (CD19 BiTE)



Adapted from Brown P. Blood. 2018;131:1497-1498

CHILDREN'S ONCOLOGY GROUP

- In multiple relapsed/refractory setting (pediatrics)
 - CR 35%-40%
 - MRD-negative CR 20%–25%

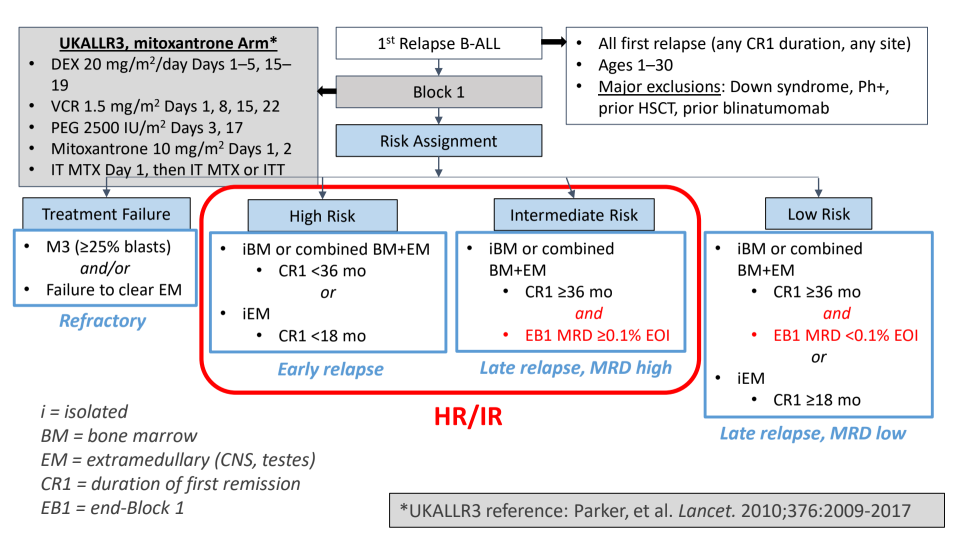
von Stackelberg et al. J Clin Oncol. 2016;34:4381-4389

- In MRD+ setting (adults)
 - 80% MRD clearance
 - 60% subsequent DFS (bridge to HSCT)

Gokbuget et al. Blood. 2018;131:1522-1531

Objective of COG AALL1331:

To determine if substituting blinatumomab for intensive consolidation chemotherapy improves survival in 1st relapse of childhood/AYA B-ALL



Stratifications

- Risk group (HR vs IR)
- For HR
 - Site (BM vs iEM)
 - For BM: CR1 duration (<18 vs 18– 36 mo)

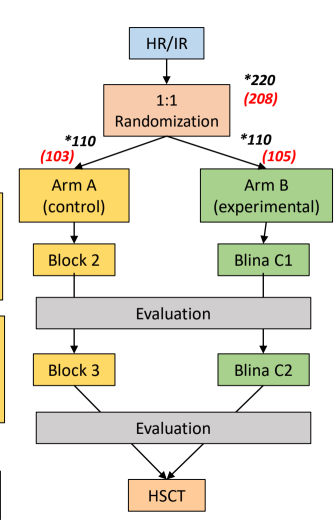
UKALLR3, Block 2*

- VCR, DEX week 1
- ID MTX, PEG week 2
- CPM/ETOP week 3
- IT MTX or ITT

UKALLR3, Block 3*

- VCR, DEX week 1
- HD ARAC, Erwinia Weeks 1–2
- ID MTX, Erwinia Week 4
- IT MTX or ITT

*UKALLR3 reference: Parker, et al. Lancet. 2010;376:2009-2017



Endpoints

- Primary: DFS
- Other: OS, MRD response, ability to proceed to HSCT

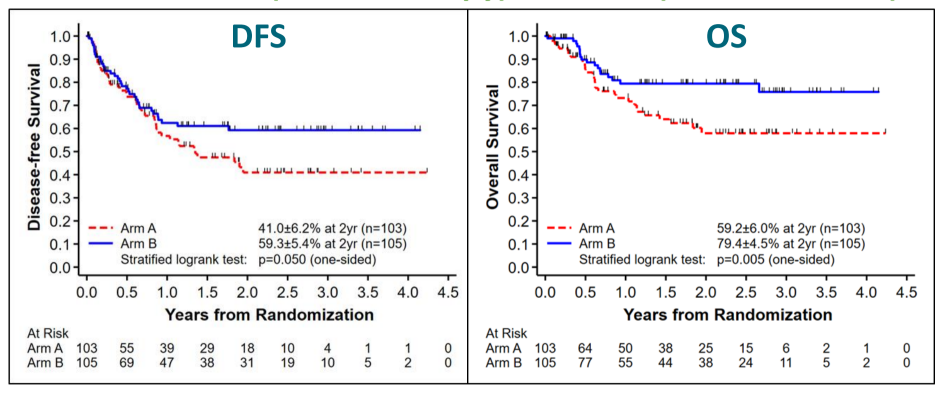
Sample size n = 220 (110 per arm)

- Power 85% to detect HR 0.58 with 1-sided α = 0.025
- Increase 2-yr DFS from 45% to 63%

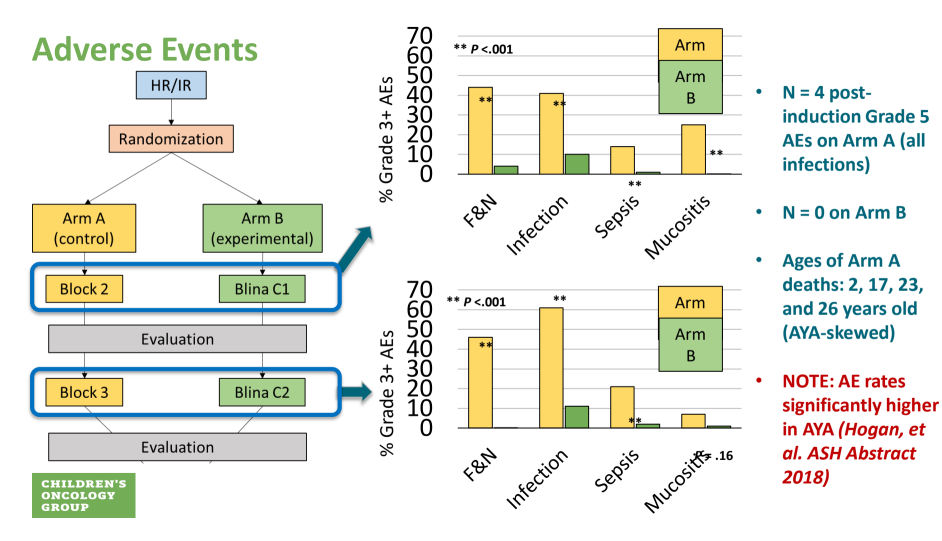
Blina C1 and Blina C2

- Blinatumomab 15 ug/m²/day × 28 days, then 7 days off
- Dex 5 mg/m²/dose × 1 premed (C1 only)
- First patient randomized
 Jan 2015
- Randomization halted Sep 2019 (95% projected accrual)

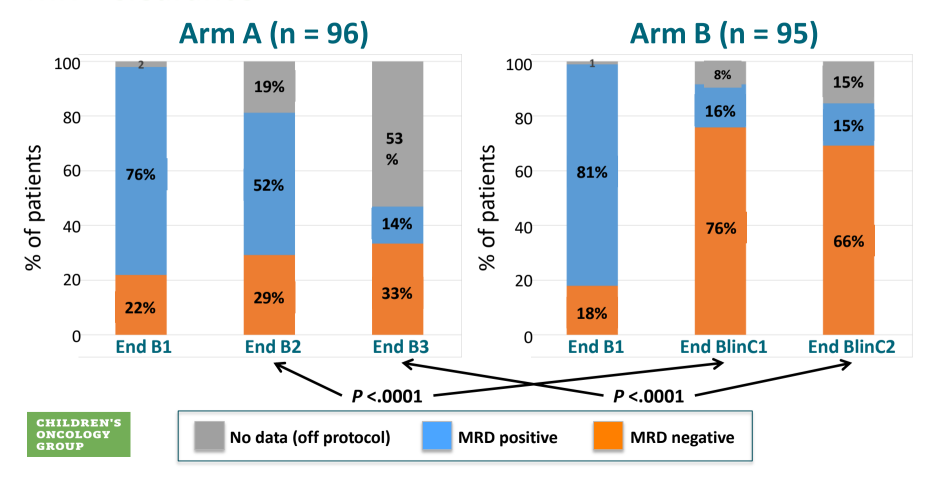
Survival: Arm A (chemotherapy) vs Arm B (blinatumomab)



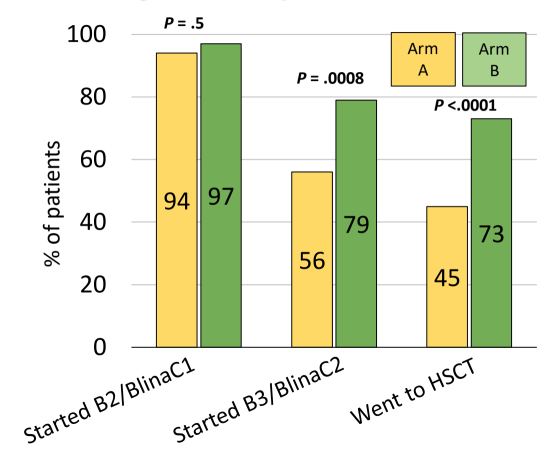




MRD Clearance

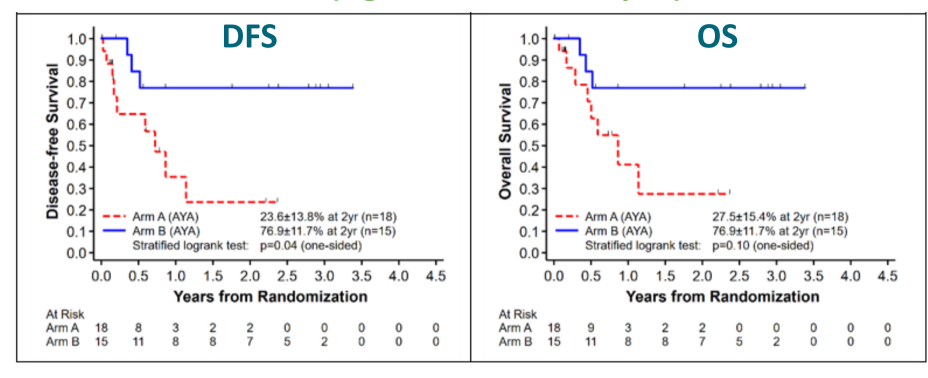


Proceeding to Transplant: Arm A vs Arm B



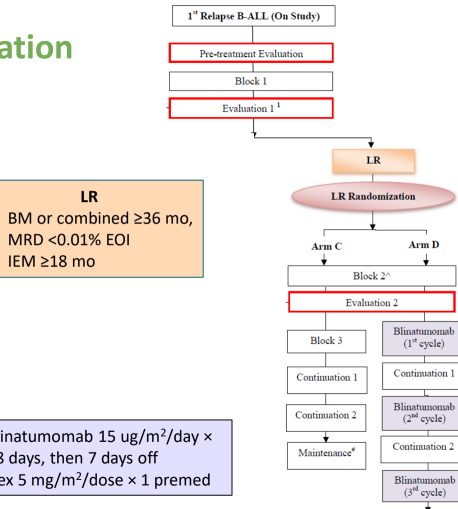
A significant contributor to the improved survival for Arm B (blina) vs Arm A (chemo) in HR/IR relapses may be the ability of blinatumomab to successfully bridge to HSCT

Results AYA Patients (Ages 18–30 at Relapse)





LR Randomization



Maintenance#

Blinatumomab 15 ug/m²/day × 28 days, then 7 days off

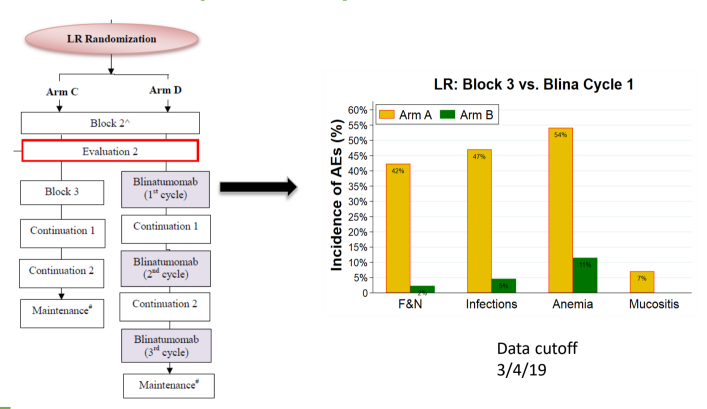
LR

MRD < 0.01% EOI

IEM ≥18 mo

Dex 5 mg/m 2 /dose × 1 premed

Adverse Events: LR (Grade 3+)





Superior Event-Free Survival With Blinatumomab Versus Chemotherapy in Children With High-risk First Relapse of B-cell Precursor Acute Lymphoblastic Leukemia: A Randomized, Controlled Phase 3 Trial

Franco Locatelli, Gerhard Zugmaier, Carmelo Rizzari, Joan Morris, Bernd Gruhn, Thomas Klingebiel, Rosanna Parasole, Christin Linderkamp, Christian Flotho, Arnaud Petit, Concetta Micalizzi, Noemi Mergen, Abeera Mohammad, Cornelia Eckert, Anja Moericke, Mary Sartor, Ondrej Hrusak, Christina Peters, Vaskar Saha, and Arend von Stackelberg

Open-Label, Randomized, Phase III Trial: 47 Centers, 13 Countries

Key eligibility criteria

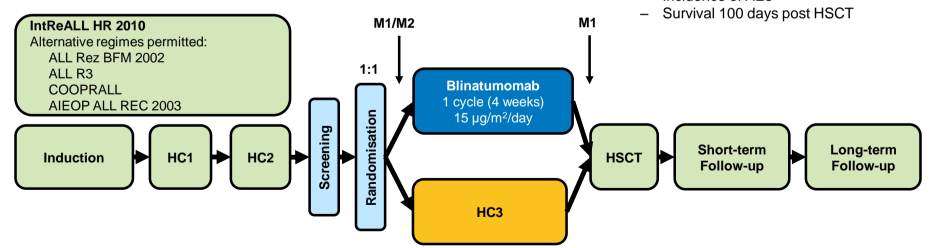
- Age >28 days <18 years
- HR 1st relapse Ph- BCP-ALL
- M1 or M2 marrow at randomization
- No CNS disease, unless treated before enrollment
- No clinically relevant CNS pathology

Stratification

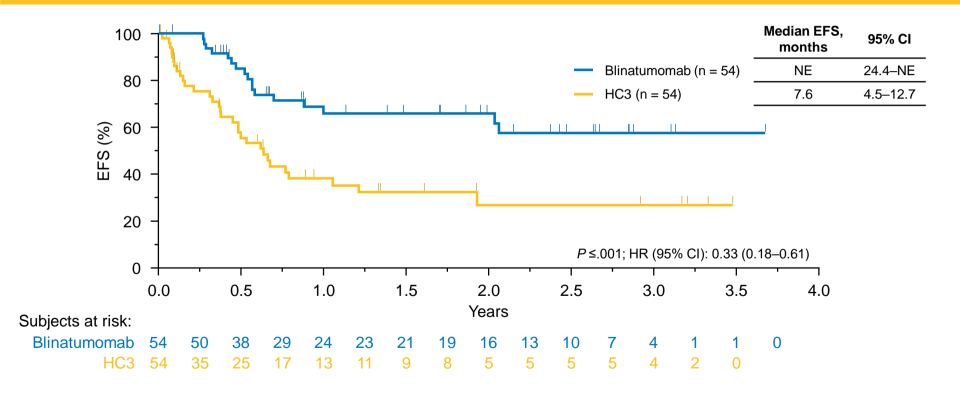
- Age: <1 year, 1 to 9 years, >9 years
- BM status at end of HC2
 - M1 with MRD >10⁻³
 - M1 with MRD <10⁻³
 - M2

Endpoints

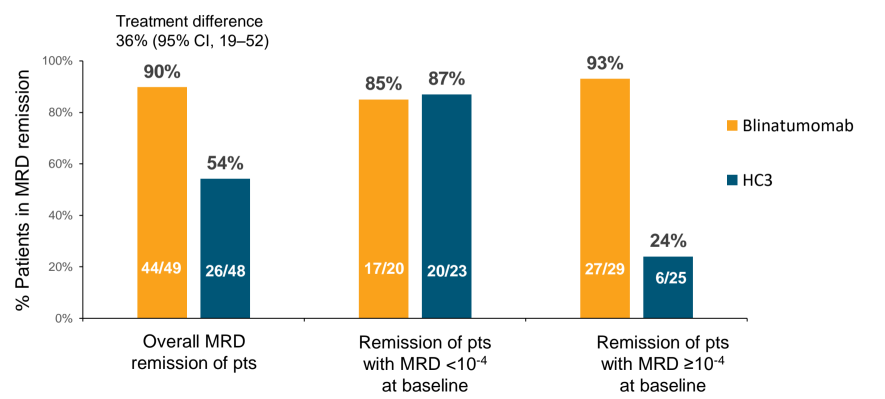
- Primary: EFS
- Secondary
 - OS
 - MRD response (end of blinatumomab or HC3)
 - Cumulative incidence of relapse
 - Incidence of AEs



Superior EFS in the Blinatumomab Arm



Superior MRD Remission by PCR in the Blinatumomab Arm (overall and by baseline* MRD status)



^{*}Baseline: end of HC2 (screening sample before enrollment). PCR, polymerase chain reaction.

Conclusions for Trials

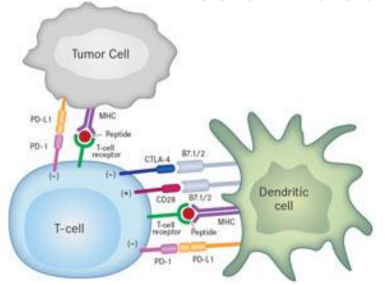
- For children and AYA patients with HR/IR first relapse of B-ALL, blinatumomab is superior to standard chemotherapy as post-reinduction consolidation prior to HSCT, resulting in
 - Fewer and less-severe toxicities
 - Higher rates of MRD response
 - Greater likelihood of proceeding to HSCT
 - Improved disease-free and overall survival
- Blinatumomab constitutes a new standard of care in this setting
- Future: Optimizing immunotherapy in relapsed ALL
 - Combination of blinatumomab and checkpoint inhibitors
 - Immunotherapy to replace or augment reinduction chemotherapy
 - CAR T cells to replace or augment HSCT



What Happens When Blinatumomab Doesn't Work?

Endogenous T-cell "exhaustion"

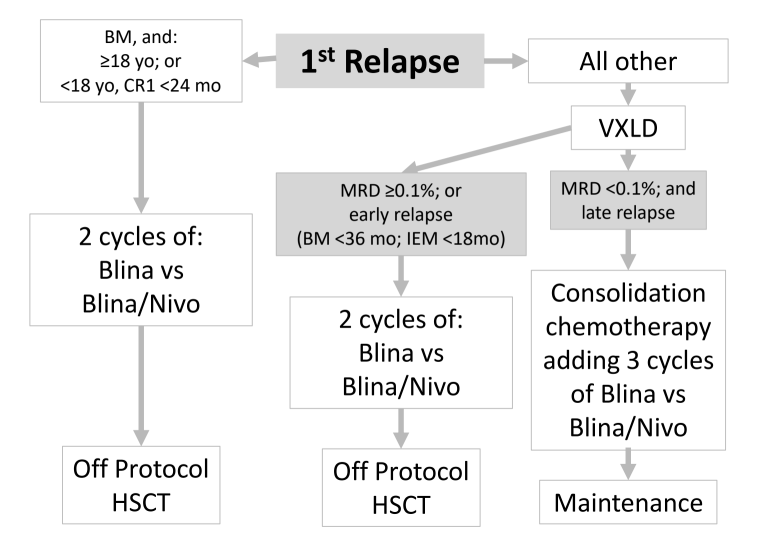




PD-1	PD-L1	CTLA-4
Nivolumab	Atezolizumab	Ipilimumab
Pembrolizumab	Avelumab	
	Durvalumab	

Reports of efficacy in patients relapsing after blina/CAR T cells

• Feucht, et al. *Oncotarget*. 2016;7(47):76902-76919



Can We Predict When Blinatumomab Won't Work?



Correspondence 🙃 Free Access

Day 15 bone marrow minimal residual disease predicts response to blinatumomab in relapsed/refractory paediatric B-ALL

Patrick Brown 🗷, Gerhard Zugmaier, Lia Gore, Catherine A. Tuglus, Arend von Stackelberg

First published: 03 December 2019 | https://doi.org/10.1111/bjh.16306

Efficacy Outcomes in Patients Enrolled in Phase I/II Study

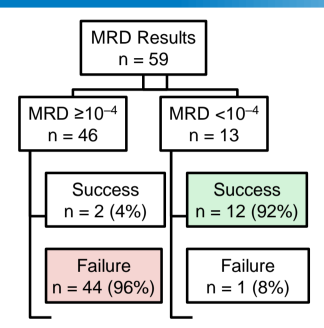
Response	Patients at Recommended Dose That Had Response Assessment (N = 64) ^a			
	n/N (%)	95% CI		
CR within the first 2 cycles	27/64 (42)	30, 55		
Non-responders (did not achieve CR)	→ 37/64 (58)	45, 70		
Partial remission	4			
Blast-free or aplastic bone marrow	2			
Progressive disease	10			
No response	21			
MRD response in patients who achieved CR within the first 2 cycles				
Complete MRD response	14/27 (52)	32, 71		
No MRD response	12/27 (44)	26, 64		
No data available	1/27 (4)			

- Study definitions
 - "Success" was defined as complete MRD response in CR (n = 14)
 - "Failure" was defined as anything other than success (n = 50)

^a70 patients treated at 5/15 μg/m²/d in phase I or II; Six patients died (n = 5) or withdrew consent (n = 1) before the first response assessment. CR, complete remissions; HSCT, hematopoietic stem cell transplantation; MRD, minimal residual disease. Adapted from von Stackelberg, et al. *J Clin Oncol.* 2016;34:4381-4389.

Biomarkers to Predict Blinatumomab Success/Failure

Overall, Day 15
 MRD results
 predicted best
 response after 2
 cycles with 95%
 accuracy (correctly
 in 56 of 59 patients)



•	Study	defin	itions
	– 10.0.,		

- "Success" was defined as complete MRD response in CR (n = 14)
- "Failure" was defined as anything other than success (n = 50)

Parameter	Accuracy (n/N)	Accuracy (%)
Day 8 PB morphology (clearance of blasts)	19/40	49
Day 15 BM morphology (M1)	54/60	90
Day 29 BM morphology (M1)	42/51	84
Day 15 BM MRD (< 10-4)	56/59	95
Day 29 BM MRD (< 10 ⁻⁴)	42/49	86

NOTE: Day 8 PB is an especially poor predictor of subsequent response

As patients with MRD ≥10⁻⁴ at Day 15 could potentially pursue alternative therapies, such as dose escalation or combination therapies, Day 15 MRD results may allow personalized treatment and improve outcomes in pediatric patients with relapsed/refractory B-ALL

Blinatumomab: Questions and Discussion

- HSCT after MRD clearance with blinatumomab?
- Role of HTS (ClonoSEQ) MRD?
- Ability of checkpoint inhibition to safely enhance blinatumomab response?
- Earlier (pre-treatment) predictive biomarkers of blinatumomab response?
- Risk of prior blinatumomab exposure and CD19 escape after subsequent CD19 CAR T therapy?





CAR T Cells for Pediatric/AYA ALL

Franco Locatelli, MD

University of RomelRCCS Ospedale Pediatrico Bambino Gesù of Rome Italy







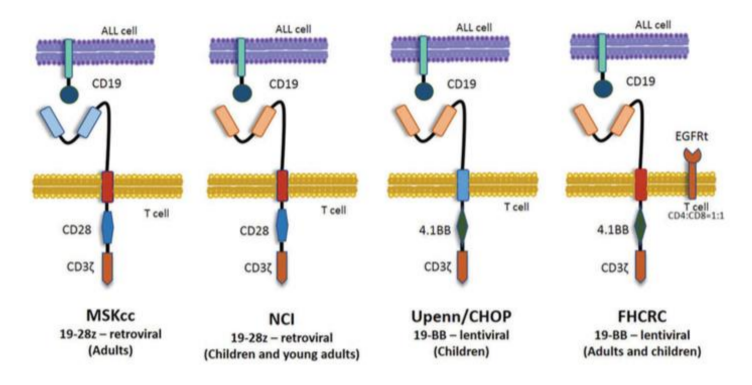
CAR T Cells for Pediatric/AYA ALL

Franco Locatelli, MD
Università Sapienza, Roma
Dept. Pediatric Hematology/Oncology and Cell/Gene Therapy
IRCCS Ospedale Bambino Gesù, Roma, Italy



Published constructs of second-generation CD19 CARs for ALL

CAR design important for persistence and sustained efficacy



Published studies of second-generation CD19 CAR-T cells for R/R ALL

Reference	Treated Patients (n)	CAR Vector	Response + Consolidation
Maude SL, et al. <i>N Engl J Med.</i> 2014;371:1507-1517	30	FMC63-41BB-ζ	27 CR; 22 MRD-negative
	(18 post-HSCT)	lentivirus	3 → allogeneic HSCT
Lee DW, et al.	20	FMC63-CD28-ζ	13 CR + 1 CRi; 12 MRD-negative
Lancet. 2015;385:517-528	(7 post-HSCT)	retrovirus	10 → allogeneic HSCT
Gardner RA, et al. <i>Blood.</i> 2017;129:3322-3331	43	FMC63-41BB-ζ	41 CR; 41 MRD-negative
	(28 post-HSCT)	lentivirus	11 → allogeneic HSCT
Maude SL, et al. <i>N Engl J Med.</i> 2018;378:439-448	75	FMC63-41BB-ζ	61 CR/CRi; 61 MRD-negative
	(46 post-HSCT)	lentivirus	8 → allogeneic HSCT
Turtle CJ, et al.	30	FMC63-41BB-ζ	29 CR; 25 MRD-negative
J Clin Invest. 2016;126:2123-2138	(11 post-HSCT)	lentivirus	13 → allogeneic HSCT
Park JH, et al.	53	SJC25C1-CD28-ζ	44 CR; 32 MRD-negative
N Engl J Med. 2018;378:449-459	(19 post-HSCT)	retrovirus	17 → allogeneic HSCT

251 patients treated: 85% CR, 76% MRD-negative



ELIANA study design

Key Eligibility Criteria

Inclusion

- R/R B-cell ALL, aged 3-21 years^a
- Bone marrow with ≥5% lymphoblasts

Exclusion

- Isolated extra-medullary disease relapse
- Prior CD19-directed or gene therapy

Endpoints

- Primary endpoint: Overall remission rate (CR + CRi) within 3 months
- 4-week maintenance of remission
- IRC assessment
- Secondary endpoints
 - MRD status, DOR, OS, EFS, cellular kinetics, safety

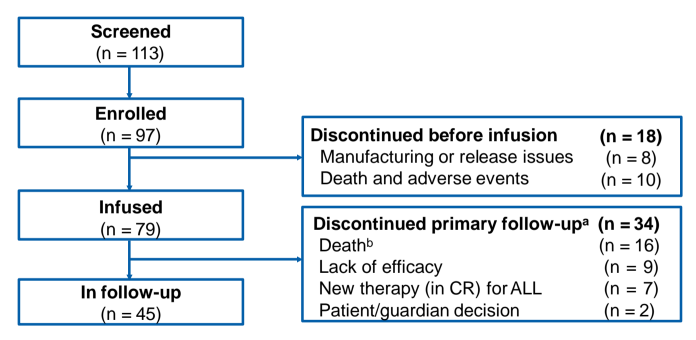
Study Treatment

- Lymphodepleting chemotherapy prior to infusion
 - Fludarabine 30 mg/m² IV daily for 4 doses
 - Cyclophosphamide 500 mg/m² IV daily for 2 doses
- Tisagenlecleucel dose range (single infusion)
 - 0.2 to 5.0 × 10⁶ cells/kg for patients ≤50 kg
 - 0.1 to 2.5 × 108 cells for patients >50 kg

^a Age of 3 years at the time of screening to age of 21 years at time of initial diagnosis.

CR, complete response; CRi, CR with incomplete blood count recovery; DOR, duration of response; IRC, Independent Review Committee; MRD, minimal residual disease; OS, overall survival; R/R B-ALL, relapsed or refractory B-cell acute lymphoblastic leukemia.

ELIANA Patient disposition



Median time from infusion to data cut-off (13 April 2018) was 24.2 months (range, 4.5-35.1 months)

a Patients followed for survival.

^b One death occurred while the patient was in remission; other deaths occurred after treatment failure or relapse.

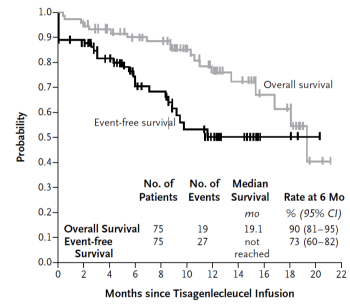
Summary of ELIANA study

ORIGINAL ARTICLE

Tisagenlecleucel in Children and Young Adults with B-Cell Lymphoblastic Leukemia

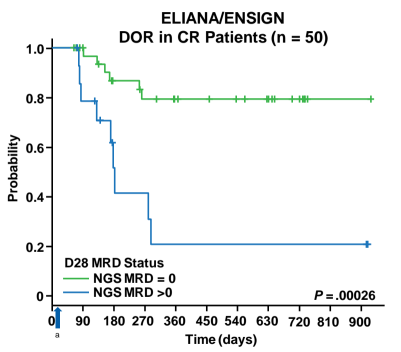
- 92 patients enrolled, 75 treated
- 73% Grade 3–4 AEs related to CAR T
- 81% → CR/CRi, all MRD negative; 66% in intention-to-treat analysis
- 1-year EFS at 50%, no relapses after this
- Demonstrates feasibility of delivery in multiple centers
- FDA approval for R/R pediatric ALL: August 2017
- Also approved in the EU, Canada, and Switzerland

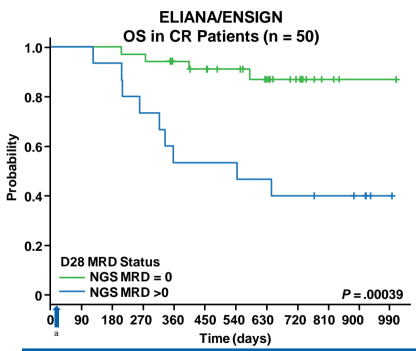
B Event-free and Overall Survival



No. at Risk												
Overall survival	75	72	64	58	55	40	30	20	12	8	2	0
Event-free survival	75	64	51	37	33	19	13	8	3	3	1	0

Patients with no MRD detected in D28 bone marrow by NGS had superior outcomes





Pulsipher MA, et al. Molecular Detection of Minimal Residual Disease Precedes Morphological Relapse and Could be Used to Identify Relapse in Pediatric and Young Adult B-Cell Acute Lymphoblastic Leukemia Patients Treated with Tisagenlecleucel. ASH 2018 Abstract 1551

^a Tisagenlecleucel infusion at Day = 1.

Frequency of high-risk cytogenetic abnormalities in ELIANA and ENSIGN

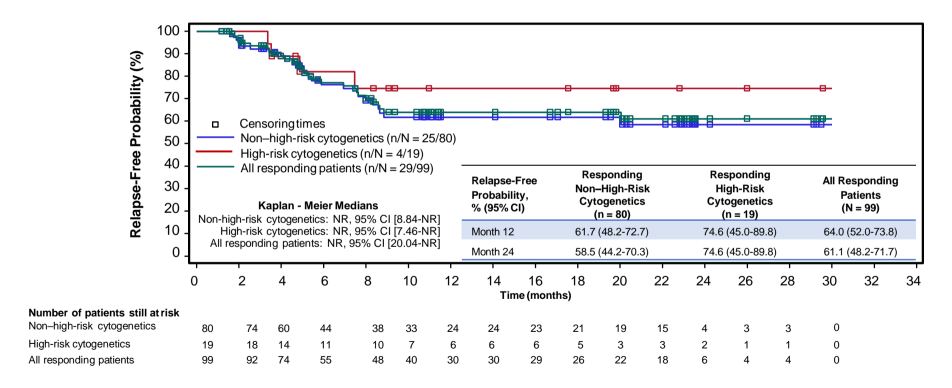
29 of 137 infused patients had high-risk cytogenetic abnormalities

High-Risk Cytogenetic Abnormality	n
Hypodiploidy ^a	3
t(9;22)(q34;q11.2)/BCR-ABL1	5
KMT2A (MLL) rearrangement	4
Intrachromosomal amplification of chromosome 21 (iAMP21)	7
t(17;19)(q23;p13), encoding <i>TCF3-HLF</i> fusion	1
BCR-ABL1-like	6
CRLF2 rearrangement	2
TP53 mutation/deletion	1

High response rate in evaluable patients

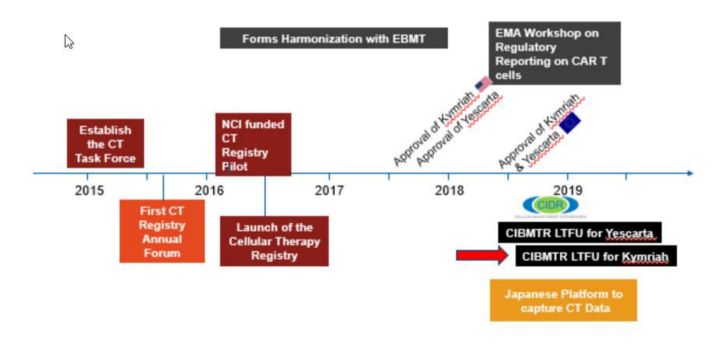
	High-Risk Cytogenetics (n = 25)	Non–High-Risk Cytogenetics (n = 104)	All Patients (N = 129)
ORR, n (%)	19 (76.0)	80 (76.9)	99 (76.7)
CR	17 (68.0)	72 (69.2)	89 (69.0)
CRi	2 (8.0)	8 (7.7)	10 (7.8)
Responding patients with MRD-negative disease, an/N (%)	18/19 (94.7)b	78/80 (97.5) ^c	96/99 (97.0)
HSCT post-infusion while in remission, n (%)	1 (4.0)	9 (8.6)	10 (7.8)

Median RFS was not reached in responding high-risk and non-high-risk cytogenetics subgroups



Development of the Cellular Therapy Registry



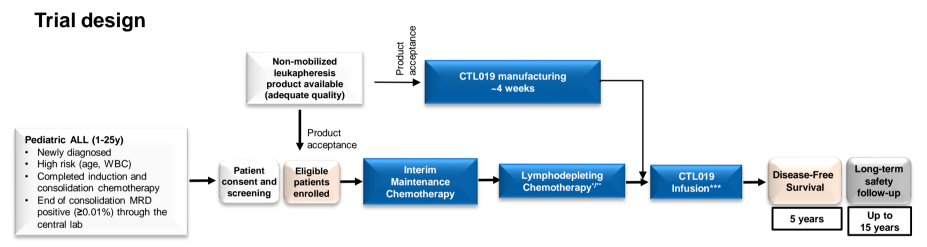


Acute lymphoblastic leukemia

ALL Characteristic	N (%)
No. of patients	144
Disease status at CT	
Primary induction failure	18 (12)
Morphologic CR	50 (35)
Disease relapse	76 (53)
≥5% blast in marrow prior to CT	45 (31)
Extramedullary disease prior to CT	20 (14)
Ph+ ALL	11 (8)
≥3 of lines of prior therapies	70 (49)
Prior allogeneic HCT	48 (33)

- Median time from diagnosis to CT –
 33 months
- CR rate was 87% and among patients with MRD assessment (N = 58), 98% were negative
- DOR at 6m 71%
- EFS at 6m 66%
- OS at 6m 91%

Study of efficacy and safety of tisagenlecleucel in HR B-ALL EOC MRD-positive patients (CASSIOPEIA)



*Lymphodepleting chemotherapy

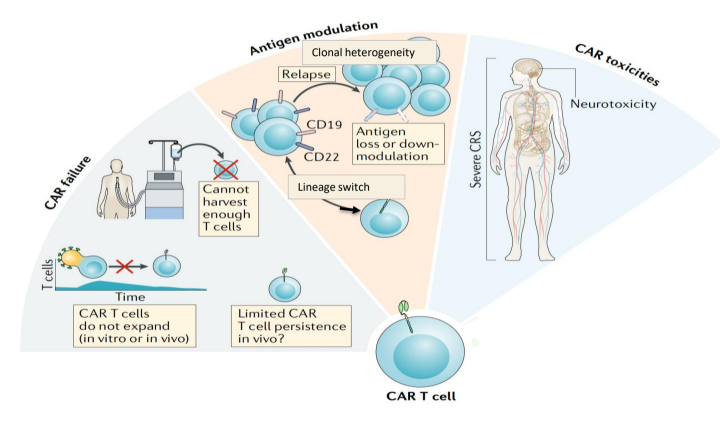
- Fludarabine (25 mg/m² IV daily for 4 days)
- Cyclophosphamide (500 mg/m² IV daily for 2 days starting with the first dose of fludarabine)

**Second Infusion: If the patient satisfies certain criteria, a second infusion may be possible. The patient would then restart all visits starting from the LD chemo visit

***Single IV infusion

- ≤50 kg body weight: 0.2 to 5 × 10⁶ tisagenlecleucel transduced cells/kg
- >50 kg body weight: 0.1 to 2.5 × 108 tisagenlecleucel transduced cells

Current limitations of CAR T cells



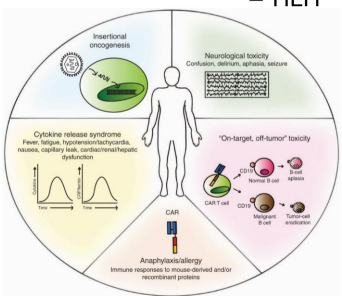
Peculiar toxicities associated with CD19 CAR T cells

"On-target, off-tumor" toxicities

B cell-aplasia

Non-antigen-specific toxicities

- Cytokine release syndrome (CRS)
- Neurotoxicity
- HLH



Overall safety and AEs of special interest within 8 weeks after infusion

AESI ^a	Patients (N = 79)		
AESI	All Grades, %	Grade 3, %	Grade 4, %
Cytokine release syndrome	77	22	27
Infections	43	20	4
Cytopenias not resolved by day 28	42	18	18
Neurological events	39	13	0
Tumor lysis syndrome	5	5	0

- Majority of AEs occurred in the first 8 weeks after tisagenlecleucel infusion
- No cases of cerebral edema reported

^a Occurring within 8 weeks of tisagenlecleucel infusion.

^b Cytokine release syndrome was graded using the Penn scale. AESI, adverse events of special interest.

Cytokine release syndrome

	Patients Infused (N = 79)
Patients developed CRS, n (%)	61 (77)
Time to onset, median (range), days	3.0 (1-22)
Duration of CRS, median (range), days	8.0 (1-36)
ICU admission, n (%)	38 (48)
Anticytokine therapy, n (%)	31 (39)
Tocilizumab, n (%)	31 (39)
1 dose	18 (23)
2 doses	10 (13)
3 doses	3 (4)
Corticosteroids, n (%)	16 (20)
Hypotension that required intervention, n (%)	42 (53)
High-dose vasopressors, n (%)	19 (24)
Intubation, n (%)	12 (15)
Dialysis, n (%)	8 (10)

CRS was graded using the Penn scale and managed by a protocol-specific algorithm¹

Positive association of CRS grade and neurological event grade

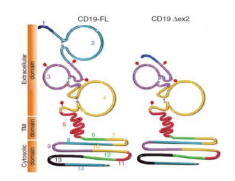
CRS	N	Any-Grade Neurological Events, n (%)	Grade 3 Neurological Events, n (%)
None	18	4 (22)	1 (6)
Grade 1/2	23	7 (30)	1 (4)
Grade 3	17	7 (41)	2 (12)
Grade 4	21	13 (62)	6 (29)

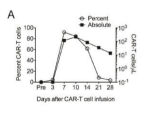
- Grade 3 neurological events were more frequent with grade 4 CRS compared with grade 0-3 CRS (95% CI, –2% to 45%)
- Median onset of any-grade CRS (day 3) preceded median onset of neurological events (day 7)
- Grade 3 or 4 CRS and neurological events occur earlier than grade 1 or 2

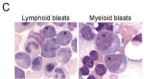
Mechanisms of leukemia escape after CAR T-cell therapy

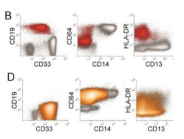
Tumor evasion systems in BCP-ALL: CD19-negative relapses

- ➤ Loss of CAR-recognized epitope as a result of alternative exon splicing forms of the CD19 gene where exon 2 was lost (Sotillo et al. Cancer Discov. 2015)
- ➤ Altered trafficking of CD19 protein to the cell membrane of blast cells (Braig et al. *Blood.* 2016)
- ➤ Myeloid switch and loss of CD19 in patients with mixed-phenotype leukemia and MLL rearrangement (Gardner et al. *Blood.* 2016)
- ➤ Induction of resistance to CAR T-cell therapy by transduction of a single leukemic B cell (Ruella et al. Nat Med. 2018)









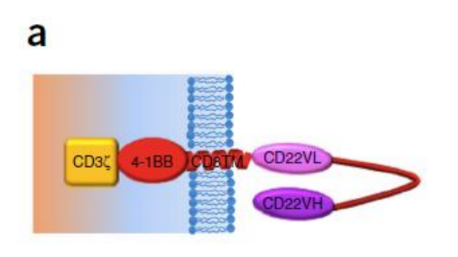
Characteristics of remission and relapse

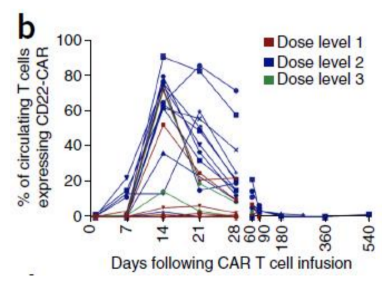
- Relapses tended to occur early (within the first year)
- Most relapses were CD19-negative:

CD19-Negative	CD19-Positive	Unknown CD19 Status
14/19 (73.7% of relapses)	3/19 (15.8% of relapses)	2/19 (10.5% of relapses)

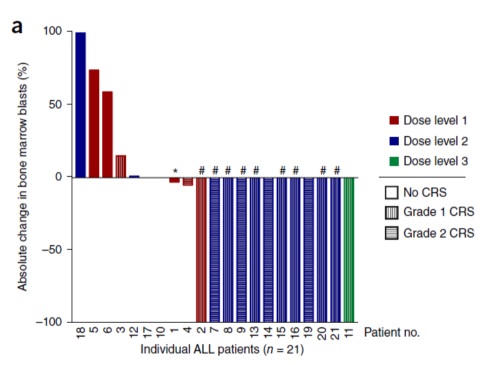
- All CD19-negative relapses occurred in the context of persistent B-cell aplasia
- One CR patient with B-cell recovery at 12 months is still in ongoing CR for 27 months at the time of data cut-off

CD22.CAR





CD22.CAR

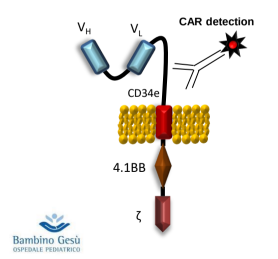


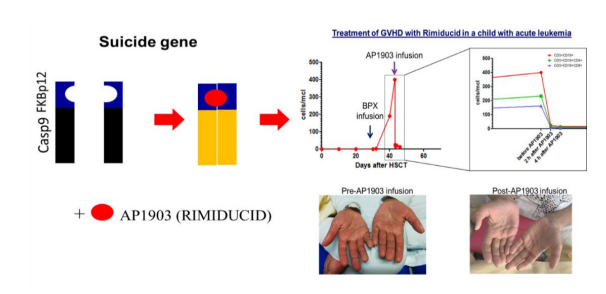
- Twenty-one children/young adults treated with CD22.4-1BB CAR T cells
- Antileukemia activity is dose-dependent
- CR obtained in 73% (11/15) of patients receiving ≥1×10⁶ CD22-CAR T cells/kg, including 5 of 5 patients with CD19dim or CD19− B-ALL
- Eight patients relapsed (reduced CD22 surface site density in 7 of them)

CAR.CD19 strategy at OPBG: Second-generation CAR targeting CD19



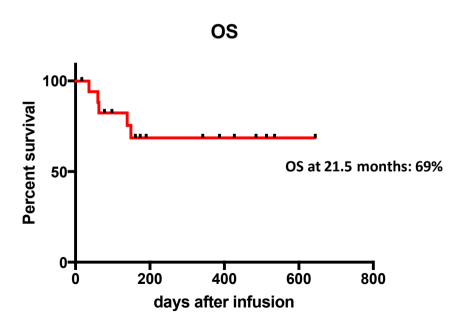
CAR.CD19-4.1BB-ζ





Outcome of ALL patients treated with CD19-CAR T cells at OPBG

Fourteen out of the 17 (82%) patients with Bcp-ALL infused obtained CR with MRD negativity after DP infusion

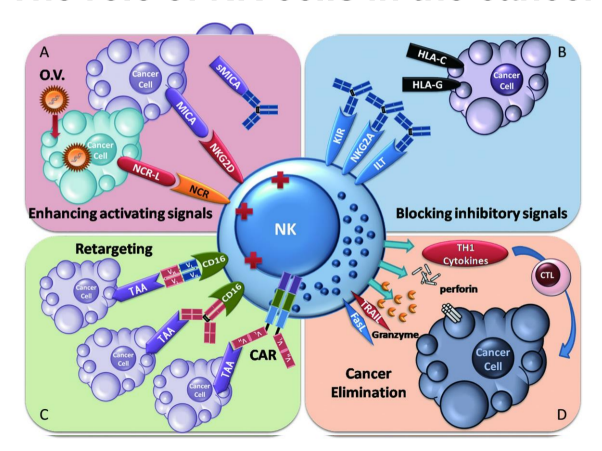




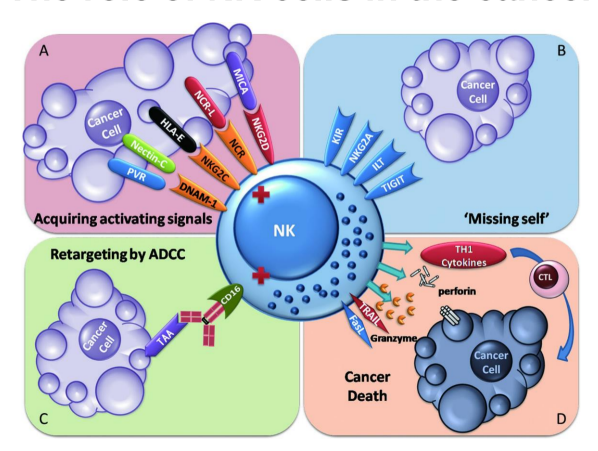
Role of CAR T cells in AYA

- Commercially available CAR T-cell products are approved for patients until 25 years of age
- No data are available on safety and efficacy outcome of CAR T cells in AYA as compared with children below the age of 13 years
- Considering the relevant toxicities and the risk of treatment-related fatality observed in AYA with intensive chemotherapy protocols and HSCT, CAR T cells could represent an attractive option to be considered for relapsed/refractory patients

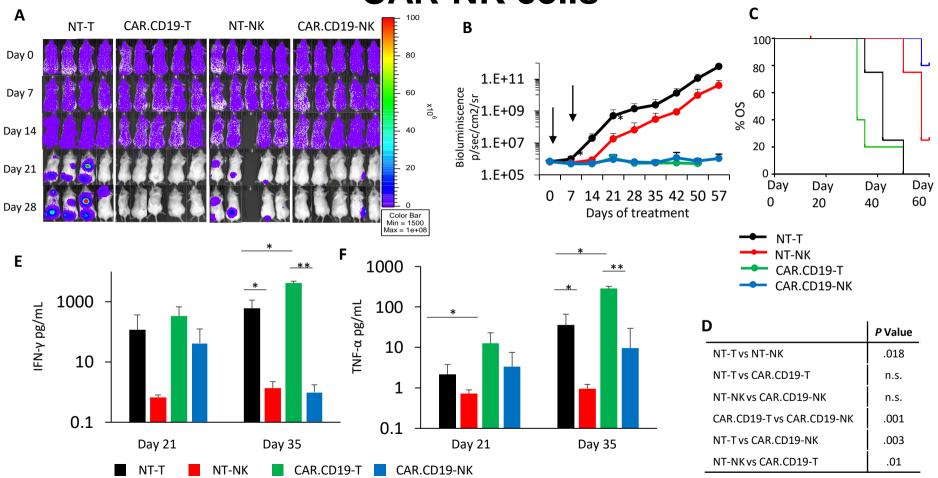
The role of NK cells in the cancer



The role of NK cells in the cancer



CAR NK cells





Questions to Experts

Rupert Handgretinger, MD
Children's University Hospital and
Department of Hematology/Oncology
Germany



Thank you to all participants!

And thank you to Amgen and Adaptive Biotechnologies for their sponsorship

- > Please complete the evaluation form using the provided link
- > To obtain a copy of the meeting slides and access other educational materials, please visit the GLA website at www.globalleukemiaacademy.com
 - Meeting materials will be available in approximately 1 week
- If you have a question for any of our experts that was not answered today, you can submit it through the GLA website at https://globalleukemiaacademy.com/ask-the-expert/

