

# Immunotherapy for the Treatment of Hematologic Malignancies

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- No relevant financial relationships to disclose
- I will be discussing non-FDA approved indications during my presentation.





# Outline: Major immunotherapies under development

- Immune checkpoint inhibitors
- Antibody-drug conjugates
- Bispecifics
- Cellular therapies



### Society for Immunotherapy of Cancer Immune checkpoint inhibitors **ADVANCES IN** IMMUNOTHERAPY Anti-PD-1 **TCR/MHC** interaction CD80/CD28 interaction **Tumor** T cell cell



Anti-PD-L1



### FDA-approved checkpoint inhibitors: lymphoma

Drug	Indication	Dose
Nivolumab	Classical <b>Hodgkin lymphoma</b> , relapsed after HSCT and brentuximab vedotin or ≥3 previous therapies	240 mg Q2W or 480 mg Q4W
Pembrolizumab	Adult relapsed/refractory classical <b>Hodgkin</b> <b>lymphoma</b> Pediatric refractory <b>cHL</b> or <b>cHL</b> relapsed	200 mg Q3W or 400 mg Q6W adults 2 mg/kg (up to 200 mg) Q3W
	after <a>2 lines of therapy</a>	(pediatric)
Pembrolizumab	Adult/pediatric refractory <b>primary</b> <b>mediastinal large B-cell lymphoma</b> or relapsed after 2 previous therapies**	200 mg Q3W or 400 mg Q6W adults 2 mg/kg (up to 200 mg) Q3W (pediatric)

\*\*Not recommended for patients with PBMCL that require urgent cytoreductive therapy.





# Efficacy of approved checkpoint inhibitors: lymphoma

Study	Treatment	Patient population	Overall response rate	Complete response rate	Landmark OS
CheckMate 205	Nivolumab	Brentuximab vedotin-naïve <b>cHL</b>	65%	29%	1-year: 92%
		Bretuximab vedotin after auto-HCT cHL	68%	13%	1-year: 93%
		Bretuximab vedotin before/after auto-HCT cHL	73%	12%	1-year: 90%
KEYNOTE-087	Pembrolizumab	cHL progressed after ASCT and BV	78.3%	26%	3-year: 86.3%
		<b>cHL</b> after salvage chemo and BV, ineligible for ASCT	64.2%	26%	3-year: 85.7%
		<b>cHL</b> progressed after ASCT without BV treatment	71.7%	31.7%	3-year: 87.6%
KEYNOTE-013	Pembrolizumab	<b>PMBCL</b> with relapse/ineligible for ASCT	48%	33%	1-year: 65%
KEYNOTE-170	Pembrolizumab	<b>PMBCL</b> ineligible for ASCT with progression on <u>&gt;</u> 2 previous therapies	45%	13%	1-year: 58%

cHL: Classical Hodgkin lymphoma; PMBCL: primary mediastinal B cell lymphoma

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# In development: Immune checkpoint inhibitors in AML

Study	Population	Treatment(s)	ORR	Median OS (months)	Status
NCT02775903	Untreated AML	Azacitidine + durvalumab	20%	13.0	Active, not recruiting
		Azacitidine	23%	14.4	recruiting
NCT02397720	Relapsed/refractory AML	Azacitidine + nivolumab	33%	6.4	Recruiting
		Azacitidine + nivolumab + ipilimumab	44%	10.5	
NCT02768792	Relapsed/refractory AML	HiDAC followed by pembrolizumab	46%	8.9	Active, not recruiting
	Relapsed/refractory AML	Azacitidine + pembrolizumab	31%	10.8	Recruiting
	Newly diagnosed AML, <u>&gt;65</u> years of age		70.5%	13.1	

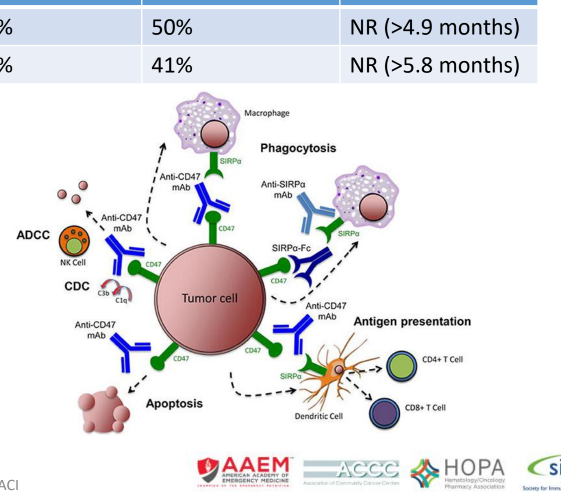




# In development: Macrophage checkpoint: CD47

Treatment	Populations	ORR	CRR	Median DOR
Azacitidine +	Untreated MDS	91.7%	50%	NR (>4.9 months)
magroliumab	Untreated AML	63.6%	41%	NR (>5.8 months)

- CD47 is expressed on some cancer cells
- CD47 signaling through SIRPα prohibits macrophage phagocytosis of cancer cells – "don't eat me"
- Blocking interaction of CD47 and SIRPα promotes adaptive immune responses and boosts tumor cell phagocytosis







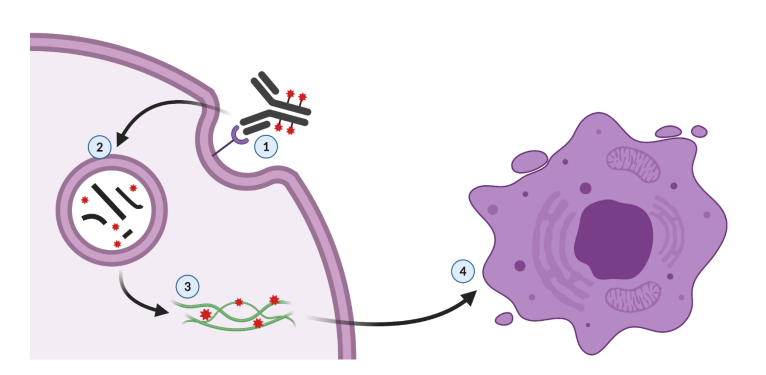
- Immune checkpoint inhibitors
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## Antibody-drug conjugates

- 1. Antibody binds to receptor on tumor cell
- 2. ADC is internalized and broken down
- 3. Drug payload performs its MOA (here, microtubule disruption)
- 4. Apoptosis is induced in target cell







# FDA-approved antibody-drug conjugates

Drug	Target antigen	Indication		
		<b>Classical Hodgkin lymphoma,</b> relapsed after HSCT or ≥2 previous therapies		
Brentuximab vedotin	CD30	Cutaneous anaplastic large cell lymphoma or CD30+ mycosis fungoides ≥ 1 previous therapies		
		Classical Hodgkin lymphoma - first line with combination chemo		
		Classical Hodgkin lymphoma consolidation after auto-HSCT		
Inotuzumab ozogamicin	CD22	Relapsed/refractory/MRD+ B-cell ALL		
Polatuzumab vedotin (w/ bendamustine & rituximab)	CD79b	<b>DLBCL</b> ≥ 2 previous therapies		
Gemtuzumab ozogamicin	CD33	R/R or newly-diagnosed CD33+ AML in adults or pediatric patients		
Belantamab mafodotin	BCMA	<b>R/R multiple myeloma</b> after <u>&gt;</u> 4 prior therapies		

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# Efficacy of approved ADCs – brentuximab vedotin

33%	5-year: 41%	
	- ,	
56%	5-year: 60%	
2-year modified PFS rate: 82.1%		
d PFS rate: 7	7.2%	
Median PFS: 42.9 months		
Median PFS: 24.1 months		
d P d P 2.9	PFS rate: 8 PFS rate: 7 months	

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## Efficacy of approved ADCs

Study	Treatment(s)	Patient population	Key outcomes
INO-VATE	Inotuzumab ozogamicin Standard-of-care chemo	Relapsed/refractory <b>B cell precursor</b> ALL	CR/CRi rate: 73.8% vs 30.9% Median OS: 7.7 vs 6.2 months 2-year OS: 22.8% vs 10%
GO29365	Polatuzumab vedotin + bendamustine & rituximab Bendamustine & rituximab	Relapsed/refractory <b>DLBCL</b>	CRR: 40.0% vs 17.5% Median PFS: 9.5 vs 3.7 months Median OS: 12.4 vs 4.7 months
ALFA-0701	Gemtuzumab ozogamicin + daunorubicin + cytarabine Daunorubicin + cytarabine	De novo acute <b>myeloid leukemia</b>	CR/CRp rate: 81.5% vs 73.6% Median OS: 27.5 vs 21.8 months Median EFS: 17.3 vs 9.5 months
DREAMM-2	Belantamab mafodotin	R/R <b>multiple myeloma</b> after IMiD, PI, and anti-CD38	ORR: 31% Median PFS: 2.9 months



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# **In development:** Novel ADCs in clinical trials

Trial	Indication	Treatment(s)	ADC target antigen	Phase
NCT03544281	R/R multiple myeloma	GSK2857916 + lenalidomide + dexamethasone	BCMA	2
		GSK2857916 + bortezomib + dexamethasone		
NCT03386513	CD123+ AML, BPDCN or ALL	IMGN632	CD123	1/2
NCT03424603	R/R B cell malignancies	STRO-001	CD74	1
NCT03682796	R/R B cell lymphoma	TRPH-222	CD22	1
NCT04240704	CLL or NHL	JBH492	CCR7	1
NCT03833180	Pre-treated hematologic malignancies	VLS-101	ROR1	1





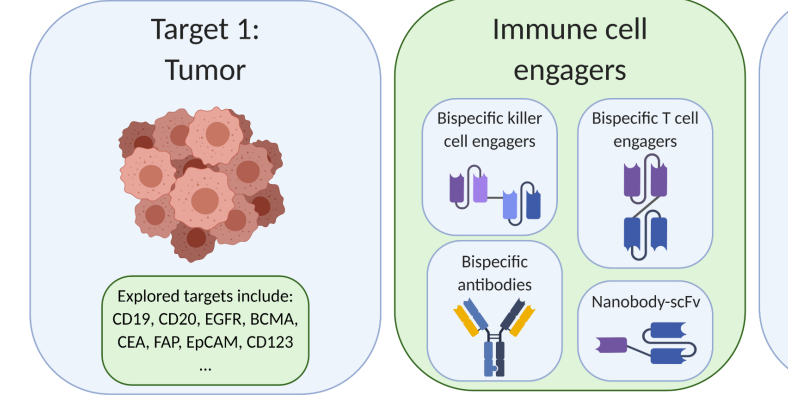


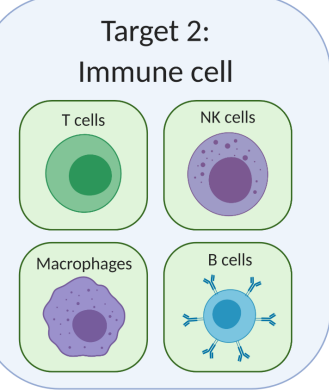
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## **Bispecifics in immunotherapy**





*Commonly CD3 on T cells, CD16 for NK and macrophages, etc* 





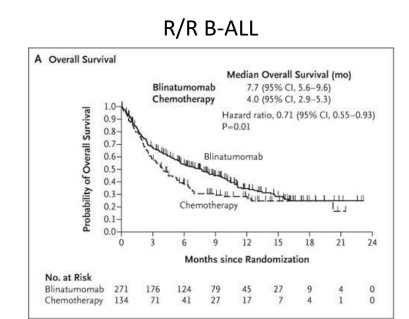
## Clinical use of immune cell engagers

Drug	Indications	CD19
	Relapsed/refractory B-ALL	
Blinatumomab	B-ALL in 1 <sup>st</sup> or 2 <sup>nd</sup> complete response with MRD ≥ 0.1%	CD3



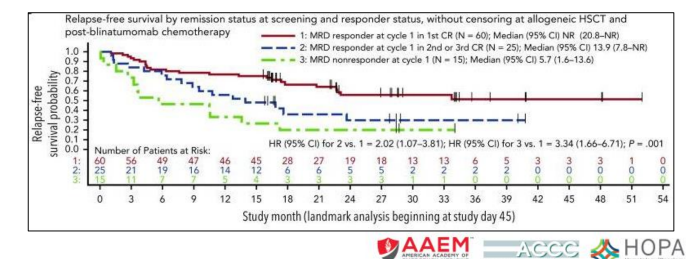


## Blinatumomab in R/R B-ALL



Trial	Patient population	Treatment	Key outcomes
NCT02013167	Adults with R/R B-ALL	Blinatumomab	Median OS: 7.7 vs 4.0 months
		Chemotherapy	Median DOR: 7.3 vs 4.6 months
NCT01207388	Adults with MRD+ B-ALL	Blinatumomab	Complete MRD response rate: 78% Median OS: 36.5 months

#### MRD+ B-ALL



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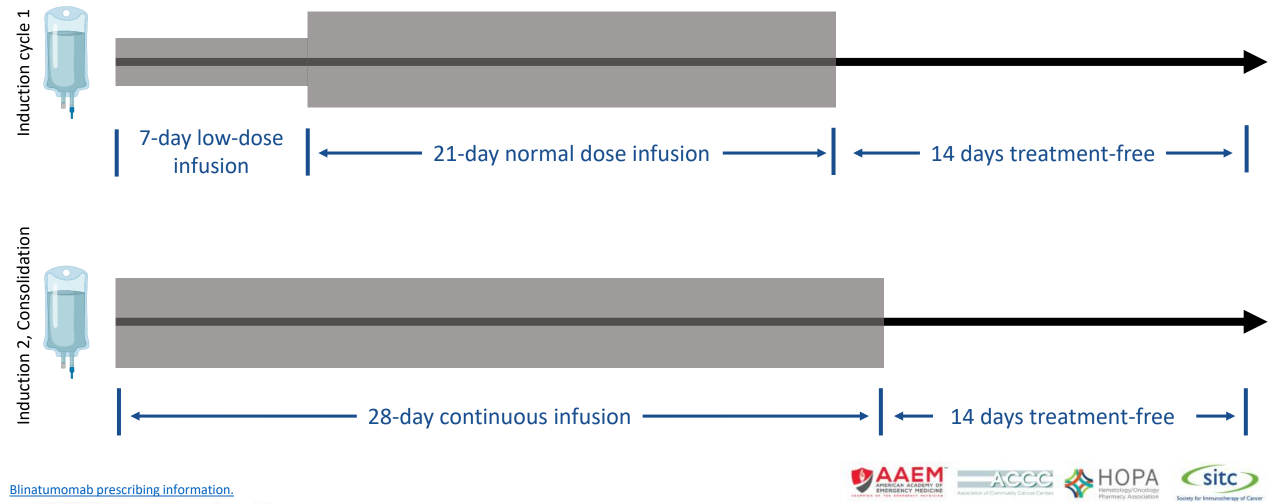
## Dosing regimens for blinatumomab

	Cycle		Patients weighing 45 kg or more (Fixed-dose)	Patients weighing less than 45 kg (BSA-based dose)
MRD-	Induction cycle 1	Days 1-28	28 mcg/day	15 mcg/m <sup>2</sup> /day (not to exceed 28 mcg/day)
positive B		Days 29-42	14-day treatment-free interval	14-day treatment-free interval
ALL	Consolidation cycles 2-4	Days 1-28	28 mcg/day	15 mcg/m <sup>2</sup> /day (not to exceed 28 mcg/day)
		Days 29-42	14-day treatment-free interval	14-day treatment-free interval
	Cycle		Patients weighing 45 kg or more (Fixed-dose)	Patients weighing less than 45 kg (BSA-based dose)
	Induction cycle 1	Days 1-7	9 mcg/day	5 mcg/m <sup>2</sup> /day (not to exceed 9 mcg/day)
		Days 8-28	28 mcg/day	15 mcg/m <sup>2</sup> /day (not to exceed 28 mcg/day)
		Days 29-42	14-day treatment-free interval	14-day treatment-free interval
R/R B-	Induction cycle 2	Days 1-28	28 mcg/day	15 mcg/m <sup>2</sup> /day (not to exceed 28 mcg/day)
ALL		Days 29-42	14-day treatment-free interval	14-day treatment-free interval
	Consolidation cycles 3-5	Days 1-28	28 mcg/day	15 mcg/m <sup>2</sup> /day (not to exceed 28 mcg/day)
		Days 29-42	14-day treatment-free interval	14-day treatment-free interval
	Continued therapy cycles	Days 1-28	28 mcg/day	15 mcg/m <sup>2</sup> /day (not to exceed 28 mcg/day)
	6-9	Days 29-42	56-day treatment-free interval	56-day treatment-free interval
natumomab prescr	ibing information.			AAAEM AMERICAN ACADEMY OF EMERGENCY MEDICINE Auscator of Community Carters Certers

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# Dosing regimens for blinatumomab – R/R B-ALL



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## Common side effects of T cell engagers

#### Cytokine release syndrome

- Characterized by initial flu-like symptoms, which progress into a shock-like syndrome with elevation in cytokine levels
- Patients display fever, vascular leakage, and organ dysfunction
- Variable onset and course
- Pre-treatment with dexamethasone required
- Management:
  - IL-6 and IL-6R antagonism
  - Corticosteroids
  - Other cytokine receptor antagonists

#### **B** cell aplasia

- Due to current clinical agents targeting CD19, which is expressed by both normal and neoplastic B cells
- May result in hypogammaglobulinemia
- Increased risk of infection
- Managed through administration of intravenous immunoglobulin

Stay tuned: more information on toxicity management later in this program

#### Neurotoxicity

- Also known as "immune effector cell-associated neurotoxicity syndrome" (ICANS)
- Manifests as confusion, delirium, seizures, cerebral edema
- Largely unknown mechanisms
- Incidence increases with more doses, increased age, more prior therapies
- Management:
  - Supportive care for low-grade
  - Corticosteroids for highergrade









# **In development:** Novel immune cell engagers in clinical trials

Trial	Indication	Treatment	Target antigens	Phase
NCT03214666	HR myelodysplastic syndromes, R/R AML, systemic mastocytosis	GTB-3550 (TriKE)	CD16, IL-15, CD33	1/2
NCT03516591	High-risk myelodysplastic syndromes	AMV564	CD33, CD3	1
NCT03739606	CD123+ R/R blood cancers	Flotetuzumab	CD123, CD3	2
NCT02730312	CD123+ R/R blood cancers	XmAb14045	CD123, CD3	1
NCT03888105	R/R B cell NHL	Odronextamab	CD20, CD3	2
NCT03309111	Previously treated multiple myeloma	GBR 1342	CD38, CD3	1/2
NCT03761108	R/R multiple myeloma	REGN5458	BCMA, CD3	1/2







- Immune checkpoint inhibitors
- Antibody-drug conjugates
- Bispecifics
- Cellular therapies





### Comparing T cell engagers and CAR T therapy

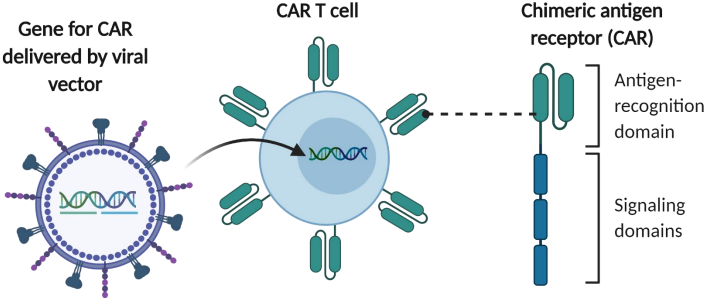
	CAR T cells	T cell engagers (BiTEs)
Structure	Synthetic gene construct encoding an scFv against tumor antigen linked to activation/costimulatory motifs	Recombinant protein with two specificities: one for tumor antigen and one for T cell antigen (usually CD3)
Effector cell types	Engineered CD8+ and CD4+ T cells	Endogenous CD8+ and CD4+ T cells
Immune synapse	Atypical	Typical
Serial killing	Yes	Yes
Killing mechanisms	Perforin and granzyme B, Fas-Fas-L, or TNF/TNF-R	Perforin and granzyme B
Trafficking	Active	Passive
<b>Clinical applications</b>	Pre-treatment lymphodepletion followed by a single infusion	No lymphodepletion; repeat administration and continuous infusions.
Specificity	Manufactured for each patient	"Off-the-shelf"





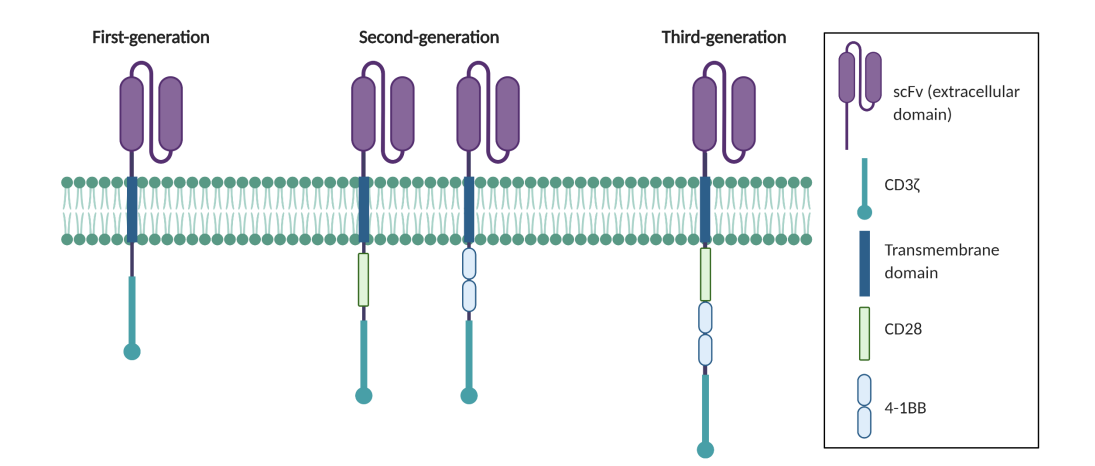
## **Chimeric antigen receptors**

- Overcome immune tolerance
- Targets surface molecules in native conformation
- Independent of antigen presenting cell and MHC complex





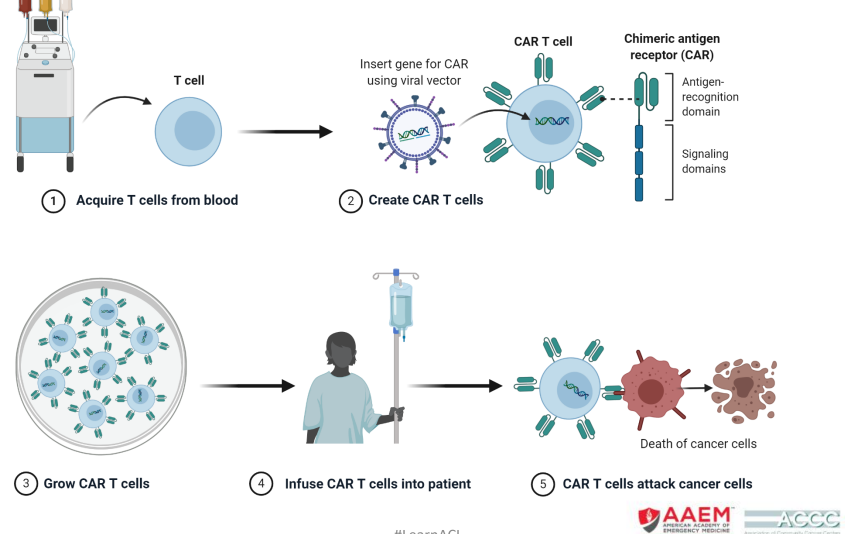
### **Evolution of CAR constructs**







# CAR T manufacturing and administration



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## FDA-approved CAR T cell therapies

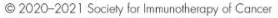
Drug	Target/co-stimulatory domain	Indication	Dose	
Axicabtagene ciloleucel	CD19/CD28	Adults with r/r large <b>B-cell lymphoma</b> , Including diffuse large B-cell lymphoma, primary mediastinal large B-cell lymphoma, high-grade B- cell lymphoma, DLBCL arising from follicular lymphoma, and R/R follicular lymphoma	2 x 10 <sup>6</sup> CAR-positive, viable T cells per kg bodyweight (up to 2x10 <sup>8</sup> )	
Tisagenlecleucel	CD19/4-1BB	Patients ≤25 yr with refractory <b>B-cell acute lymphoblastic leukemia</b> or in 2+ relapse	0.2-0.5x10 <sup>6</sup> CAR-positive, viable T cells per kg if under 50 kg 0.1-2.5x10 <sup>8</sup> CAR-positive, viable T-cells if over 50 kg	
Tisagenlecleucel	CD19/4-1BB	Adults with r/r large <b>B-cell lymphoma</b> after 2+ therapies Including DLBCL, high-grade B-cell lymphoma, DLBCL arising from follicular lymphoma	0.6-6.0 x $10^8$ CAR-positive, viable T cells	
Brexucabtagene autoleucel	CD19/CD28	Adults with <b>mantle cell lymphoma</b> (MCL) who have not responded to or who have relapsed following other treatments	2 x 10 <sup>6</sup> CAR-positive, viable T cells per kg bodyweight (up to 2x10 <sup>8</sup> )	
Lisocabtagene maraleucel	CD19/4-1BB	Adults with R/R large <b>B-cell lymphoma</b> after at least 2 prior therapies	50-110 x 10 <sup>6</sup> CAR-positive viable T cells (1:1 CD4:CD8)	
Idecabtagene vicleucel	BCMA/4-1BB	Adults with R/R <b>multiple myeloma</b> after 4+ prior therapies	300-460x10 <sup>6</sup> CAR-positive T cells	
		#LearnACI	ACCCC Association al Community Carses Centers	



### Comparing clinical trials of CD19 CAR T therapies

Trial	Indication	Treatment(s)	ORR	Landmark OS	Grade 3+ toxicity rates
ZUMA-2	R/R mantle cell lymphoma	Brexucabtagene autoleucel (KTE- X19)	86% CRR: 57%	1-year: 86%	CRS: 18% NE: 46%
ZUMA-1	Refractory large B cell lymphoma	Axicabtagene ciloleucel	83% CRR: 58%	2-year: 50%	CRS: 11% NE: 32%
JULIET	R/R diffuse large B cell lymphoma	Tisagenlecleucel	52% CRR: 40%	1-year: 49%	CRS: 22% NE: 12%
ELIANA	R/R B cell acute lymphoblastic leukemia	Tisagenlecleucel	82% CRR: 62%	18-month: 70%	CRS: 48% NE: 13%

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### CAR T side effects

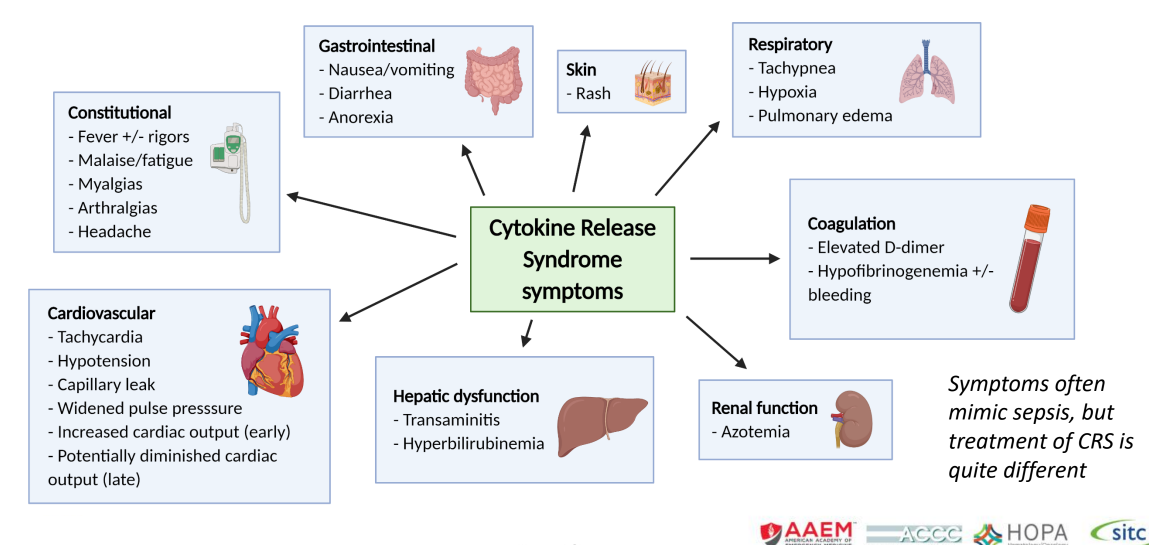
- Cytokine Release Syndrome (CRS)
- Neurotoxicity
  - ICANS: Immune effector cell-associated neurotoxicity syndrome
  - NE: Neurologic events
- B cell aplasia
- Macrophage Activation Syndrome (MAS)/HLH

Stay tuned: more information on toxicity management later in this program





### CAR T side effects - CRS



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## **Eligibility considerations for CAR**

- Disease
  - Relative stability during CAR T manufacturing (~2-6 weeks)
  - Bridging therapy (polatuzumab, chemo, RT, steroids, lenalidomide, ibrutinib)
  - CNS control
- Patient
  - Adequate cell counts
  - DVT, bleeding, infection, neuro disorders
  - Functional status: at screen vs. day of CAR T infusion
- Other
  - Social support, reimbursement
  - Availability of tocilizumab for CRS management





# **In development:** Novel CAR T therapies in clinical trials

Trial	Indication	Treatment	Target antigen	Phase
NCT03971799	R/R pediatric AML	CD33CART	CD33	1/2
NCT04186520	R/R B cell malignancies	CAR-20/19 T cells	CD19, CD20	1/2
NCT04109482	R/R BPDCN, AML, HR MDS	MB-102	CD123	1/2
NCT03287817	Diffuse large B cell lymphoma	AUTO3	CD19, CD22	1/2
NCT02690545	R/R HL and NHL	ATLCAR.CD30	CD30	1/2





### Conclusions

- Many immunotherapy options for hematological malignancies
- Checkpoint inhibitors for Hodgkin lymphoma and PMBCL high response rate, excellent tolerance, durable responses if CR
- Blinatumomab and inotuzumab for ALL effective salvage, deeper remissions
- Polatuzumab vedotin for DLBCL effective salvage, potential to become frontline
- CAR T therapy ever-increasing indications; patient selection and toxicity management still concerns





## **Additional Resources**









### Acknowledgements

- University of Virginia, Division of Hematology/Oncology
- SITC leadership
- Some figures created using Biorender.com





### **Case Studies**





### Case Study #1

- 60 year old woman with PMHx of splenic marginal zone lymphoma transformed to stage IVB DLBCL s/p R-CHOP x 6 with scattered FDG-avid uptake of the bilateral forearms on post-treatment PET-CT. Patient feels well but complains of moderate bilateral arm pain.
- What would you do next?
  - A) Start therapy directed at DLBCL
  - B) Start therapy directed at MZL
  - C) Watch and wait with short follow-up PET-CT
  - D) Obtain biopsy of FDG-avid lesion





- A) Start therapy directed at DLBCL
- B) Start therapy directed at MZL

Unclear which histology to target with therapy

- C) Watch and wait with short follow-up PET-CT
  - Reasonable although risk of rapidly progressive disease
- D) Obtain biopsy of FDG-avid lesion
  - Allows us to choose appropriate next therapy





- Biopsy of FDG-avid bone lesion shows recurrent DLBCL and short follow up PET-CT shows rapidly progressive disease involving axial and appendicular skeleton as well as diffuse FDG-avid lymphadenopathy.
- Patient initiated on salvage immunochemotherapy for DLBCL
  - R-ICE complicated by ifosfamide neurotoxicity and renal electrolyte wasting
  - R-GDP without complication
- Repeat PET with partial response to therapy





- What would you do next?
  - A) Initiate workup for high dose therapy followed by autoSCT
  - B) Initiate workup for CAR-T therapy
  - C) Administer another cycle of salvage immunochemotherapy
  - D) Transition to hospice





- A) Initiate workup for high dose therapy followed by autoSCT
  - Primary refractory disease with lack of CR to salvage therapy, low likelihood of benefit with autoSCT
- B) Initiate workup for CAR-T therapy
  - Next appropriate step as patient is young and desires curative therapy
- C) Administer another cycle of salvage immunochemotherapy
  - Reasonable to consider as bridging therapy although would probably benefit from novel agent (polatuzumab) as bridging instead
- D) Transition to hospice
  - Appropriate to consider but patient young and many treatment options available

