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	NCT Number Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
1	NCT0411 5345 A Study of a Renal Autologous Cell Therapy (REACT®) in Patients With Chronic Kidney Disease (CKD) From Congenital Anomalies of the Kidney and Urinary Tract (CAKUT). Study Documents:	Title Acronym: Other Ids: REGEN-004	Recruiting	Chronic Kidney Disease Congenital Anomalies of Kidney and Urinary Tract	Biological: Renal Autologous Cell Therapy (REACT®) Autologous selected renal cells (SRC)	Phase: Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Intervention Model Description: Open-label Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Assess change in eGFR and observe incidence of renal-specific procedure and/or product related adverse events (AEs) through 24 months following two Renal Autologous Cell Therapy (REACT) injections [Safety]. [Time Frame: 12 months following last REACT injection] The primary objective is to assess the safety and optimal delivery of Renal Autologous Cell Therapy (REACT) injected at one site in a recipient kidney as measured by procedure-and/or product related adverse events (AEs) through 12 months post-treatment. Secondary Outcome Measures: Number of subjects with renal-specific adverse events over a 24-month period following injection of Renal Autologous Cell Therapy (REACT). [Time Frame: 24 months following last REACT injection] The number of subjects with renal-specific adverse events over a 24-month period following injection of Renal Autologous Cell Therapy (REACT) will be observed utilizing renal-specific laboratory assessments. The secondary objective will compare the results of laboratory tests from baseline through 12 months following REACT injection, followed by an additional observational period of 18 months for a total of 24 months of observation. Each subject's baseline rate of CKD disease progression serves as his/her own "control" to monitor for changes in renal insufficiency over time.	Actual Enrollment: Estimated Enrollment: 15 Original Estimated Enrollment: Same as current Age: 18 Years to 65 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: CTI Clinical Trial and Consulting Services	Study Start: August 13, 2019 Primary Completion: March 31, 2023 (Final data collection date for primary outcome measure) Study Completion: May 30, 2023 First Posted: October 4, 2019 Results First Posted: Last Update Posted: September 10, 2022
2	NCT0523 7986 Cognitive Aftereffects of Neurotoxicity in Children and Young Adults With Relapsed/Refract ory Hematologic Malignancies Who Receive CAR T-cell Therapy Study Documents:	Title Acronym: Other Ids: 10000631 000631-C	Not yet recruiting	• Lymphom a • Leukemia	Not Provided	Study Type: Observational Phase: Study Design: Observational Model: Cohort Time Perspective: Prospective Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 60 Original Estimated Enrollment: Same as current Age: 5 Years and older (Child, Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: September 19, 2022 Primary Completion: April 30, 2024 (Final data collection date for primary outcome measure) Study Completion: April 30, 2025 First Posted: February 14, 2022 Results First Posted: Last Update Posted: September 14, 2022

	NCT Number Title	e	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
3	0328 Ch An Re T Co on Re of Im Fu Stu	ne Effect of nimeric ntigen ecceptor (CAR)- Cell Therapy n the econstitution HIV-specific numne unction udy ocuments:	Title Acronym: Other Ids: 20170407V3	Recruiting	HIV/AIDS	Biological: CAR-T cells HIV-1 specific chimeric antigen receptor cells	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Intervention Model Description: No control. Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Incidence of Treatment-Emergent Adverse Events of CAR-T cell therapy [Time Frame: 6 Months] The adverse events of VC-CAR-T cell therapy on HIV- infected patients during the clinical trial Secondary Outcome Measures: The HIV reservoir [Time Frame: 6 Months] To assay the HIV loads in the peripheral blood Mono-nuclear cells and plasma	Actual Enrollment: Estimated Enrollment: 40 Original Estimated Enrollment: Same as current Age: 18 Years to 60 Years (Adult) Sex: All	Study Sponsors: Same as current Collaborators: Sun Yat-sen University	Study Start: October 4, 2017 Primary Completion: December 31, 2023 (Final data collection date for primary outcome measure) Study Completion: December 31, 2030 First Posted: August 7, 2017 Results First Posted: Last Update Posted: September 14, 2022
4	6704 for Ad Th Max	eukapheresis r CAR or doptive Cell nerapy anufacturing udy ocuments:	Title Acronym: Other Ids: 170137 17-C-0137	Enrolling by invitation	 Leukemia Lymphom Acute Lymphobl astic Leukemia Diffuse Large B Cell Lymphom Non-Hodgkin's Lymphom 	Not Provided	Study Type: Observational Phase: Study Design: Observational Model: Cohort Time Perspective: Prospective Primary Outcome Measures: Fraction of subjects who can enroll on a CAR-T study within approximately 6 months of undergoing apherisis [Time Frame: 6 months] Secondary Outcome Measures: Fraction of patients who experience a grade 4 toxicity associated with apherisis [Time Frame: completion of apherisis procedure]	Actual Enrollment: Estimated Enrollment: 120 Original Estimated Enrollment: Same as current Age: 3 Years to 65 Years (Child, Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: August 14, 2017 Primary Completion: January 31, 2030 (Final data collection date for primary outcome measure) Study Completion: July 31, 2030 First Posted: July 24, 2017 Results First Posted: Last Update Posted: September 10, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
5	NCT0553 1708	Exploratory Study of Novel MSLN CAR-T Cell Therapy in Patients With MSLN-positive Advanced Refractory Solid Tumors Study Documents:	Title Acronym: Other Ids: 2021-IIT-004-E02	Recruiting	Mesothelin- positive Advanced Refractory Solid Tumors	 Biological: Anti-mesothelin CAR-T cells D0: Anti-mesothelin CAR-T cells are autologous genetically modified T cells. Cells will be infused intravenously. Drug: Fludarabine D-7 to D-3: Fludarabine (25 mg/m^2/day) will be administered intravenously for 5 days. Other Name: Fludara Drug: Cyclophosphamide D-7 and D-6: Cyclophosphamide (60 mg/kg/day) will be administered intravenously for 2 days. Other Name: Cytoxan 	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 20 Original Estimated Enrollment: Same as current Age: 18 Years to 70 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: UTC Therapeutics Inc.	Study Start: April 2, 2021 Primary Completion: April 30, 2025 (Final data collection date for primary outcome measure) Study Completion: April 30, 2026 First Posted: September 8, 2022 Results First Posted: Last Update Posted: September 8, 2022
6	NCT0553 4269	Stress Urinary Incontinence Study to Assess Safety and Efficacy of Muvon's Muscle Precursor Cell Therapy Study Documents:	Title Acronym: Other Ids: SUISSE MPC2	Not yet recruiting	Female Stress Urinary Incontinence	Biological: autologous muscle precursor cells Patients own Muscle Precursor Cells are isolated and injected into the rhabdomyosphincter	Study Type: Interventional Phase: Phase 2 Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description: Low dose and High dose evaluation Masking: Single (Participant) Masking Description: Neither patient nor sponsor will know which patient gets which dose Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 70 Original Estimated Enrollment: Same as current Age: 20 Years to 65 Years (Adult, Older Adult) Sex: Female	Study Sponsors: Same as current Collaborators: GCP-Service International Ltd. & Co. KG	Study Start: September 2022 Primary Completion: November 2024 (Final data collection date for primary outcome measure) Study Completion: November 2025 First Posted: September 9, 2022 Results First Posted: Last Update Posted: September 9, 2022

NCT Nui	mber Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	t Dates
7 NCT02: 5027	Mesenchymal Stem Cell Therapy in Multiple System Atrophy Study Documents:	Title Acronym: Other Ids: 12- 005950 R01FD004789 (U.S. FDA Grant/Contract) R01NS092625 (U.S. NIH Grant/Contract)	Active, not recruiting	MSA	 Biological: Autologous Mesenchymal Stem Cells single dose of 1 × 10(7) cells intrathecally Biological: Autologous Mesenchymal Stem Cells 2 doses of 5 × 10(7) cells intrathecally each 1 month (±4 days) apart Biological: Autologous Mesenchymal Stem Cells 2 doses of 1 × 10(8) cells intrathecally each 1 month apart Biological: Autologous Mesenchymal Stem Cells Ten doses of 5 x 10(7) (±20%) cells intrathecally six months (±1 month) apart Biological: Autologous Mesenchymal Stem Cells Ten doses of 2.5 x 10(7) (±20%) cells intrathecally six months (±1 month) apart 	Study Type: Interventional Phase: Phase 1 Phase 2 Study Design: Allocation: Non-Randomized Intervention Model: Sequential Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: 24 Estimated Enrollment: Original Estimated Enrollment: Age: 30 Years to 80 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Food and Drug Administr ation (FDA) National Institute of Neurologi cal Disorders and Stroke (NINDS)	Study Start: October 2012 Primary Completion: March 2024 (Final data collection date for primary outcome measure) Study Completion: March 2024 First Posted: December 11, 2014 Results First Posted: Last Update Posted: September 9, 2022

NCT Number Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
8 NCT0342 5526 Donor T Cell Therapy in Treating Immunocompro mised Patients With Adenovirus- Related Disease Study Documents:	Title Acronym: Other Ids: 2017- 0350 NCI-2018- 00929 (Registry Identifier: CTRP (Clinical Trial Reporting Program)) 2017-0350 (Other Identifier: M D Anderson Cancer Center)	Recruiting	Hematopoi etic and Lymphoid Cell Neoplasm Immunoco mpromise d	Biological: Allogeneic Adenovirus-specific Cytotoxic T Lymphocytes Given IV Other Name: Allogeneic Adenovirus-specific CTLs	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: • Toxicity of T Cells for Therapy of Adenovirus Related Disease in Immunocompromised Patients defined by the NCI Common Terminology Criteria for Adverse Events (CTCAE), Version 4.0). [Time Frame: 45 days after last CTL dose] • T Cells for Therapy of Adenovirus Related Disease in Immunocompromised Patients Determined Feasible if at Least 50% of the Enrolled Eligible Patients Receive One CTLs Infusion [Time Frame: 1 year] Secondary Outcome Measures: • Overall Survival (OS) of T Cells for Therapy of Adenovirus Related Disease in Immunocompromised Patients [Time Frame: 2 years] Overall survival (OS) defined from treatment start date to date of death. OS estimated using the Kaplan-Meier method. • Relapse-Free Survival (RFS) of T Cells for Therapy of Adenovirus Related Disease in Immunocompromised Patients [Time Frame: 2 years] Relapse-free survival (original malignancy) (RFS) defined from treatment start date to the date of documented disease recurrence or death. RFS estimated using the Kaplan-Meier method. • Cumulative Incidence of Adenovirus Reactivation After Infusion of T Cells for Therapy of Adenovirus Related Disease in Immunocompromised Patients [Time Frame: 2 years] Cumulative incidence of adenovirus reactivation after therapy assessed using the competing risks method. The competing risks include relapse and death and patients who are still alive without disease progression at end of study will be censored. • Cumulative Incidence of Grade 2-4 Graft Versus Host Disease (GVHD), Grade 3-4 GVHD, and Chronic GVHD [Time Frame: 2 years] Cumulative incidence of grade 2-4 Graft Versus Host Disease (GVHD), Grade 3-4 GVHD, and Chronic GVHD [Time Frame: 2 years] Cumulative incidence of grade 2-4 Graft Versus Host Disease (GVHD), Grade 3-4 GVHD, and Chronic GVHD [Time Frame: 2 years] Cumulative in	Actual Enrollment: Estimated Enrollment: 16 Original Estimated Enrollment: Same as current Age: Child, Adult, Older Adult Sex: All	Study Sponsors: Same as current Collaborators: National Cancer Institute (NCI)	Study Start: March 15, 2018 Primary Completion: January 1, 2024 (Final data collection date for primary outcome measure) Study Completion: January 1, 2024 First Posted: February 7, 2018 Results First Posted: Last Update Posted: September 10, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collabora	t Dates
9	NCT0425 7578	Acalabrutinib and Anti-CD19 CAR T-cell Therapy for the Treatment of B- cell Lymphoma Study Documents:	Title Acronym: Other Ids: RG1006269 NCI-2020- 00238 (Registry Identifier: NCI / CTRP) 10418 (Other Identifier: Fred Hutch/University of Washington Cancer Consortium)	Recruiting	B-Cell Non-Hodgkin Lymphom a Diffuse Large B-Cell Lymphom a, Not Otherwise Specified High Grade B-Cell Lymphom a Primary Mediastina l (Thymic) Large B-Cell Lymphom a Transform ed Follicular Lymphom a to Diffuse Large B-Cell Lymphom a Grade 1 Follicular Lymphom a Grade 2 Follicular Lymphom a Grade 3 Follicular Lymphom a Grade 3 Follicular Lymphom a	Drug: Acalabrutinib Given PO Other Names: 1420477-60-6 ACP-196 Bruton Tyrosine Kinase Inhibitor ACP-196 Calquence Biological: Axicabtagene Ciloleucel Given IV Other Names: KTE C19 KTE-C19 KTE-C19 Yescarta Testandary	Study Type: Interventional Phase: Phase 1 Phase 2 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 50 Original Estimated Enrollment: 20 Age: 18 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: AstraZeneca	Study Start: December 2, 2020 Primary Completion: March 1, 2024 (Final data collection date for primary outcome measure) Study Completion: March 1, 2029 First Posted: February 6, 2020 Results First Posted: Last Update Posted: September 9, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
								-	ors	
10	NCT0400 7029	Modified Immune Cells	Title Acronym:	Recruiting	• CD19	Biological: Chimeric Antigen Receptor T-Cell	Study Type: Interventional	Actual Enrollment:	Study Sponsors: Same as current	Study Start: October 4, 2019
	702)	(CD19/CD20	Other Ids: 18-		Positive	Therapy	Phase: Phase 1			
		CAR-T Cells) in	001989 NCI-2019-		• CD20 Positive	Given Autologous anti-CD19/anti-CD20 CAR- expressing naive/memory T cells IV	Study Design: Allocation: N/A	Estimated Enrollment: 24	Collaborators: Parker Institute	Primary Completion:
		Treating Patients With	03190 (Registry		Recurrent	Other Names:	Intervention Model: Single Group Assignment		for Cancer	August 1, 2023
		Recurrent or	Identifier:		Chronic	o CAR T Infusion	Masking: None (Open Label) Primary Purpose: Treatment	Original Estimated	Immunotherapy	(Final data
		Refractory B- Cell Lymphoma	CTRP (Clinical Trial Reporting		Lymphocy	CAR T TherapyCAR T-cell therapy		Enrollment:		collection date for primary
		or Chronic	Program))		tic Leukemia	CAR 1-cell therapy Chimeric Antigen Receptor T-cell Infusion	Primary Outcome Measures: Same as current	Same as current		outcome
		Lymphocytic	18-001989 (Other Identifier:		Recurrent	Drug: Cyclophosphamide	Secondary Outcome Measures:	Age: 18 Years		measure)
		Leukemia	UCLA / Jonsson		Diffuse	Given IV	• Clinical response [Time Frame: Up to 15 years]	to 70 Years (Adult, Older		Study
		Study Documents:	Comprehensive Cancer Center)		Large B- Cell	Other Names:	Descriptive statistics including simple summary	Adult)		Completion: August 1, 2024
		Documents.	Cancer Center)		Lymphom	(-)-Cyclophosphamide2H-1,3,2-Oxazaphosphorine, 2-[bis(2-	measures and plots appropriate for longitudinal data will be used.	Sex: All		
					a	chloroethyl)amino]tetrahydro-, 2-oxide,	Duration of remission [Time Frame: Time from	SCA. All		First Posted: July 5, 2019
					Recurrent	monohydrate	complete remission (CR)/partial remission (PR)			
					Follicular Lymphom	CarloxanCiclofosfamida	measurement criteria are first met until the first date that recurrent or progressive disease is objectively			Results First Posted:
					a	 Ciclofosfamide 	documented, or until death, assessed up to 15 years]			
					Recurrent	CicloxalClafen	Descriptive statistics including simple summary			Last Update Posted:
				Mantle Cell	o Claphene	measures and plots appropriate for longitudinal data will			September 13,	
				Lymphom	CP (cyclophosphamide) monohydrate	be used. Will also be summarized descriptively (mean, standard deviation, median, first and third quartiles,			2022	
				a	CTX (cytoxan)CYCLO-cell	minimum, maximum). Figures showing the Kaplan-				
				Recurrent	o Cycloblastin	Meier estimates will also be presented.				
					Primary Mediastina	 Cycloblastine Cyclophospham	Objective response rate (ORR) [Time Frame: Up to 15 years]			
					1	 Cyclophosphamid monohydrate 	Descriptive statistics including simple summary			
				(Thymic) Large B-	CyclophosphamidumCyclophosphan	measures and plots appropriate for longitudinal data will				
					Cell Cell	Cyclophosphane	be used. ORR and the individual rate for CR and PR will			
					Lymphom a	o Cyclophosphanum	be summarized with the frequency count and the percentage of subjects in each category, along with a 2-			
					-	CyclostinCyclostine	sided 95% exact confidence interval.			
					Recurrent Small	 Cytophosphan 	Progression-free survival [Time Frame: From time of			
					Lymphocy	CytophosphaneCytoxan	study entry to documentation of objective disease progression or death due to any cause assessed up to 15			
					tic Lymphom	• Fosfaseron	years]			
					a	o Genoxal	Descriptive statistics including simple summary			
					• Refractory	Genuxal Ledoxina	measures and plots appropriate for longitudinal data will			
					Chronic	o Mitoxan	be used. Will also be summarized descriptively (mean, standard deviation, median, first and third quartiles,			
					Lymphocy tic	NeosarRevimmune	minimum, maximum). Figures showing the Kaplan-			
					Leukemia	 Syklofosfamid 	Meier estimates will also be presented.			
					Refractory	o WR- 138719	 Overall survival [Time Frame: From date of enrollment until death, assessed up to 15 years] 			
					Diffuse Large B-	Drug: Fludarabine Phosphate	Descriptive statistics including simple summary			
					Cell	Given IV	measures and plots appropriate for longitudinal data will			
					Lymphom	Other Names: o 2-F-ara-AMP fludarabine: 2-Fluoroadenine 9-beta-	be used. Will be summarized with figures using the			
					Refractory	D-Arabinofuranoside 5'-Monophosphate	Kaplan-Meier method. The Kaplan-Meier estimates for the 1-year OS rates and the 2-sided 95% confidence			
					• Refractory Follicular	• 9H-Purin-6-amine, 2-fluoro-9-(5-O-phosphono-	interval of the rates using the Greenwood?s formula will			
					Lymphom	.betaD-arabinofuranosyl)- o Beneflur	be reported. Will also be summarized descriptively (mean, standard deviation, median, first and third			
					a D. C.	∘ Fludara	quartiles, minimum, maximum).			
					Refractory Mantle	o SH T 586	Chimeric antigen receptor (CAR) T-cell (T) 19/20			
					Cell	Biological: Tocilizumab Civen IV	bispecific transgenic T-cell persistence [Time Frame:			
					Lymphom a	Given IV	Up to 5 years post-infusion]			
					Refractory	Other Names: o Actemra	Descriptive statistics of T-cell counts over time, including simple summary measures and plots			
					Refractory Primary	o Immunoglobulin G1, Anti-(Human Interleukin 6	appropriate for longitudinal data will be used.			
					Mediastina	Receptor) (Human-Mouse Monoclonal MRA Heavy Chain), Disulfide with Human-Mouse	• Frequency of T cell phenotypic markers on CART19/20			
					l (Thymic)	Monoclonal MRA Kappa-Chain, Dimer	cells using flow cytometry [Time Frame: Up to 5 years post-infusion]			
					Large B-	 MRA (myeloma receptor antibody) 	The frequency of CART19/20 cell properties will be			
					Cell Cell	o R-1569o RoActemra	assessed using flow cytometry to indicate the % and/or			
					Lymphom	1101.101111111	total number of CART10/20 cells expressing critical			

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
11	NCT0468 4459	Dual-targeting HER2 and PD-	Title Acronym:	Recruiting	Peritoneal	Biological: Dual-targeting HER2 and PD-L1 CAR-T cells	Study Type: Interventional	Actual Enrollment:	Study Sponsors: Same as current	Study Start: March 12, 2021
	4137	L1 CAR-T for Cancers With	Other Ids: MCART-002		Carcinoma Metastatic	serosal cavity infusion	Phase: Early Phase 1	Estimated	Collaborators:	Primary
		Pleural or Peritoneal			 Pleural Effusion, 		Study Design: Allocation: N/A Intervention Model: Single Group Assignment	Enrollment: 18	Not Provided	Completion: January 1,
		Metastasis			Malignant		Masking: None (Open Label) Primary Purpose: Treatment	Original Estimated		2023 (Final data collection
		Study Documents:					Primary Outcome Measures: Same as current	Enrollment: Same as current		date for primary outcome
		2 ocuments.					Secondary Outcome Measures: Same as current	Age: 18 Years		measure)
								to 70 Years (Adult, Older		Study Completion:
								Adult)		January 1, 2024
								Sex: All		First Posted: December 24,
										2020
										Results First Posted:
										Last Update Posted:
										September 13, 2022
12	NCT0516	CD19- and	Title Acronym:	Not yet	Acute	Drug: IMJ995 single agent	Study Type: Interventional	Actual	Study Sponsors:	Study Start:
	8748	CD22-directed CAR-T Cell	Other Ids: CIMJ995A1210	recruiting	Lymphoblastic Leukemia	Single intravenous administration of IMJ995	Phase: Phase 1	Enrollment: Estimated	Same as current	December 14, 2022
		Therapy in Patients With	1				Study Design: Allocation: N/A Intervention Model: Single Group Assignment	Enrollment: 35	Collaborators: Not Provided	Primary
		Acute Lymphoblastic	2021-000677-89 (EudraCT				Masking: None (Open Label) Primary Purpose: Treatment	Original Estimated		Completion: July 3, 2026
		Leukemia	Number)				Primary Outcome Measures: Same as current	Enrollment: Same as current		(Final data collection date
		Study Documents:					Secondary Outcome Measures: Same as current	Age: 1 Year and		for primary outcome
								older (Child, Adult, Older		measure) Study
								Adult)		Completion:
								Sex: All		July 3, 2026 First Posted:
										December 23, 2021
										Results First
										Posted:
										Last Update Posted:
										September 9, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	t Dates
13	NCT0553 2761	Multidimensiona 1 Assessment of Quality of Life, Social and Professional Life and Care Utilization in Patients With Diffuse Large Cell B-cell Lymphoma Treated With CAR-T Cells Study Documents:	Title Acronym: Other Ids: 69HCL22_0430	Not yet recruiting	Diffuse Large B- cell Lymphom a (DLBCL) CAR-T Cells Treatment	Other: self-administered questionnaires In order to describe the experience of CAR-T cell therapy of DLBCL patients, a pharmaceutical follow-up is carried out the day before the injection (baseline) and at 1, 3, 6, 9, 12 and 18 months. These follow-ups consist of interviews with the patient and the delivery of self-administered questionnaires. The interviews will investigate drug consumption, the use of self-medication and complementary alternative therapies and the adverse effects of interest. The self-questionnaires will focus on exploring multidimensional quality of life, social and professional life, anxiety-depression or uncertainty tolerance through internationally validated questionnaires. No supplementary visits will be needed: interviews with the research team will occur at the end of hematologic consultations.	Study Type: Observational Phase: Study Design: Observational Model: Cohort Time Perspective: Prospective Primary Outcome Measures: Same as current Secondary Outcome Measures: Not Provided	Actual Enrollment: Estimated Enrollment: 30 Original Estimated Enrollment: Same as current Age: 18 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: September 2022 Primary Completion: March 2025 (Final data collection date for primary outcome measure) Study Completion: March 2025 First Posted: September 8, 2022 Results First Posted: Last Update Posted: September 8, 2022
14	NCT0478 9408	Study Evaluating the Safety of KITE- 222 in Participants With Relapsed/Refract ory Acute Myeloid Leukemia Study Documents:	Title Acronym: Other Ids: KT- US-486-0201 2020-000962-40 (EudraCT Number)	Recruiting	Acute Myeloid Leukemia	 Drug: Cyclophosphamide Administered intravenously Drug: Fludarabine Administered intravenously Biological: KITE-222 A single infusion of chimeric antigen receptor (CAR)-transduced autologous T cells administered intravenously 	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 40 Original Estimated Enrollment: Same as current Age: 18 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Gilead Sciences Collaborators: Not Provided	Study Start: July 19, 2021 Primary Completion: January 2024 (Final data collection date for primary outcome measure) Study Completion: January 2039 First Posted: March 9, 2021 Results First Posted: Last Update Posted: September 9, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
15	NCT Number NCT0434 8643	Safety and Efficacy of CEA-Targeted CAR-T Therapy for Relapsed/Refract ory CEA+ Cancer Study Documents:	Other Names Title Acronym: Other Ids: PBC017	Status Recruiting	Solid Tumor Lung Cancer Colorectal Cancer Liver Cancer Pancreatic Cancer Gastric Cancer Breast Cancer	Interventions Biological: CEA CAR-T cells CEA-CAR-T cells will be administered intravenously.	Characteristics Study Type: Interventional Phase: Phase 1 Phase 2 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: • The response rate of CEA CAR-T treatment in patients with relapse/refractory CEA+ Cancer that treatment by CEA CAR-T cells therapy [Time Frame: 6 months] The response rate of CEA CAR-T treatment will be recorded and assessed according to the irRECIST Version 1.1 • Duration of Response (DOR) of CEA CAR-T treatment in patients with refractory/relapsed CEA+ Cancer [Time Frame: 2 years] DOR will be assessed from the first assessment of CR/PR/SD to the first assessment of recurrence or progression of the disease or death from any cause • Progress-free survival(PFS) of CEA CAR-T treatment in patients with refractory/relapsed CEA+ Cancer [Time Frame: 2 years] PFS will be assessed from the first CAR-T cell infusion to death from any cause or the first assessment of progression • Overall survival(OS) of CEA CAR-T treatment in patients with refractory/relapsed CEA+ Cancer [Time Frame: 2 years] OS will be assessed from the first CAR-T cell infusion to death from any cause • Serum tumor marker change level [Time Frame: 2 years] In vivo quantity of CEA, AFP, etc. • Rate of CEA CAR-T cells in peripheral blood [Time Frame: 2 years]	Actual Enrollment: Estimated Enrollment: 40 Original Estimated Enrollment: Same as current Age: 18 Years to 75 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: February 20, 2020 Primary Completion: January 31, 2023 (Final data collection date for primary outcome measure) Study Completion: April 30, 2023 First Posted: April 16, 2020 Results First Posted: Last Update Posted: September 10, 2022
							 Progress-free survival(PFS) of CEA CAR-T treatment in patients with refractory/relapsed CEA+ Cancer [Time Frame: 2 years] PFS will be assessed from the first CAR-T cell infusion to death from any cause or the first assessment of progression Overall survival(OS) of CEA CAR-T treatment in patients with refractory/relapsed CEA+ Cancer [Time Frame: 2 years] OS will be assessed from the first CAR-T cell infusion to death from any cause Serum tumor marker change level [Time Frame: 2 years] In vivo quantity of CEA, AFP, etc. Rate of CEA CAR-T cells in peripheral blood [Time 			

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	t Dates
16	NCT0231 5599	Follow-Up Evaluation for Gene-Therapy- Related Delayed Adverse Events After Participation in Pediatric Oncology Branch Clinical Trials Study Documents:	Title Acronym: Other Ids: 150028 15-C-0028	Enrolling by invitation	Pediatric Cancers Hematolog ic Malignanc ies Solid Tumors	Not Provided	Study Design: Observational Model: Cohort Time Perspective: Prospective Primary Outcome Measures: Conduct long term safety evaluations after gene therapy [Time Frame: Every 3 months X 1 year then annually X 15 years] Secondary Outcome Measures: Not Provided	Actual Enrollment: Estimated Enrollment: 500 Original Estimated Enrollment: Same as current Age: 1 Year to 99 Years (Child, Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: December 23, 2014 Primary Completion: April 1, 2035 (Final data collection date for primary outcome measure) Study Completion: August 1, 2050 First Posted: December 12, 2014 Results First Posted: Last Update Posted: September 8, 2022
17	NCT0483 8171	Prospective Study of White Blood Cells Study Documents:	Title Acronym: Other Ids: QEL-RP-001	Recruiting	Autoimmu ne Diseases Inflammati on Rejection; Transplant , Liver	Other: White blood cell and blood collection Mononucleocytes will be collected via apheresis	Study Type: Observational Phase: Study Design: Observational Model: Case-Only Time Perspective: Prospective Primary Outcome Measures: Same as current Secondary Outcome Measures: Not Provided	Actual Enrollment: Estimated Enrollment: 10 Original Estimated Enrollment: Same as current Age: Child, Adult, Older Adult Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: March 24, 2021 Primary Completion: December 2024 (Final data collection date for primary outcome measure) Study Completion: December 2024 First Posted: April 8, 2021 Results First Posted: Last Update Posted: September 8, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
18	NCT0431 0592	Natural Killer Cell (CYNK-	Title Acronym:	Recruiting	• Leukemia	Biological: CYNK-001 CYNK-001 is an allogeneic off the shelf cell therapy	Study Type: Interventional	Actual Enrollment:	Study Sponsors: Same as current	Study Start: March 12, 2020
		001) Infusions	Other Ids: CYNK-001-		 Leukemia, Myeloid 	enriched for CD56+/CD3- NK cells expanded from human	Phase: Phase 1	Estimated	Collaborators:	Primary
		in Adults With AML Study Documents:	AML-001		Leukemia, Myeloid, Acute Neoplasms by Histologic Type Neoplasms Immunosu	placental CD34+ cells.	Study Design: Allocation: Non-Randomized Intervention Model: Parallel Assignment Intervention Model Description: Experimental: Minimal Residual Disease (MRD) positive AML patients; Cyclophosphamide + Fludarabine + CYNK-001. On Days 0, 7, and 14, (and 21 in certain arms) CYNK-001 at 3 varying dose levels. Experimental: Relapsed/Refractory AML patients; Cyclophosphamide + Fludarabine + CYNK-001. On Days 0, 7, and 14, (and 21 at certain dose levels) CYNK-001 at	Enrollment: 94 Original Estimated Enrollment: 22 Age: 18 Years to 80 Years (Adult, Older Adult)	Not Provided	Completion: June 3, 2024 (Final data collection date for primary outcome measure) Study Completion: December 3,
					ppressive Agents		3 varying dose levels. Masking: None (Open Label)	Sex: All		2024
					Immunolo gic Factors		Primary Purpose: Treatment Primary Outcome Measures:			First Posted: March 17, 2020
					Physiologi		Number of Participants who experience a Dose-limiting			Results First
					cal Effects of		Toxicity (DLT) [Time Frame: Day +28]			Posted:
					Drugs • Alkylating		The number of participants who experience a DLT will be measured.			Last Update Posted:
					Agents • Antimetab		 Determine the Maximum Tolerated Dose (MTD) or Maximum Planned Dose (MPD) of CYNK-001 [Time Frame: up to 28 days] 			September 14, 2022
					olites, Antineopla stic		The maximum dose safely administered for the treatment of patients with AML.			
				Antiviral Agents		Frequency and Severity of Adverse Events (AEs) [Time Frame: up to 12 months]				
				• Analgesics , Non-		Frequency and severity of Adverse Events will be evaluated.				
					narcotic • Anti-		Secondary Outcome Measures:			
					infective Agents • Analgesics		 Number of Participants who experience Minimal Residual Disease (MRD) Response [Time Frame: up to 12 months] 			
					Peripheral Nervous		The number of participants who convert from MRD positive to MRD negative.			
					System		• Time to MRD Response [Time Frame: up to 12 months]			
					Agents • Hematolog		The time it takes to convert from MRD positive to MRD negative.			
					ic Diseases • Hematolog		Duration of MRD Response [Time Frame: up to 12 months]			
					ic Neoplasms		The measure of how long participants remain MRD negative.			
					• Leukemia in		 Progression-free Survival (PFS) [Time Frame: up to 12 months] 			
					Remission • Relapsed		Date of first CYNK-001 infusion to date of disease progression.			
					Adult AML		Time to Progression (TTP) [Time Frame: up to 12 months]			
					Refractory AML		Date of first CYNK-001 infusion to date of disease progression.			
							• Duration of Morphologic Complete Remission (CR) [Time Frame: up to 12 months]			
							Duration from first Morphologic CR observation to time of disease progression.			
							• Overall Survival (OS) [Time Frame: up to 12 months]			
							Date of first CYNK-001 infusion to date of death.			

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collabora ors	t Dates
19	NCT0000 1405	Recruitment and Apheresis Collection of Peripheral Blood Hematopoietic Stem Cells, Mononuclear Cells and Granulocytes Study Documents:	Title Acronym:	Recruiting	 Granulom Granulom atous Disease, Chronic 	Not Provided	Study Type: Observational	Actual Enrollment: Estimated Enrollment: 850 Original Estimated Enrollment: Age: 18 Years to 70 Years (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: February 27,
			Other Ids: 940073 94-I-0073				Phase: Study Design: Observational Model: Cohort			1994 Primary
							Time Perspective: Other Primary Outcome Measures: Not Provided			Completion: Not Provided
					Leukocyte Disease		Secondary Outcome Measures: Not Provided			Study Completion:
					 Genetic Disease, X-Linked Genetic Disease, Inborn 					Not Provided First Posted:
										November 4, 1999
										Results First Posted:
										Last Update Posted: September 10, 2022
20	NCT0553	Study on the Safety and Efficacy of Autogenous Tumor Infiltrates Lymphocytes for the Treatment of Advanced Solid Tumor Study Documents:	Title Acronym: Other Ids: HS-IT101-ST001	Not yet	-	Biological: HS-IT101 Adoptive transfer of 1x10^9-6x10^10 autologous TIL to patients i.v. in 30-60 minutes.	Study Type: Interventional	Actual Enrollment: Estimated	Study Sponsors: Same as current Collaborators:	Study Start:
	9768			recruiting			Phase: Early Phase 1 Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label)			October 8, 2022 Primary
								Enrollment: 8 Original	Qingdao Sino- Cell Biomedicine	Completion: December 31, 2023 (Final
							Primary Purpose: Treatment	Estimated Enrollment:	Co.,Ltd.	data collection date for primary
							Secondary Outcome Measures: Same as current Age: 18 Years to 75 Years	Same as current		outcome measure)
								to 75 Years (Adult, Older Adult)		Study Completion: March 31, 2027
										First Posted: September 14, 2022
										Results First Posted:
										Last Update Posted: September 14, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborat	Dates
21	NCT0408 8864	CD22-CAR T Cells in Children and Young Adults With B Cell Malignancies Study Documents:	Other Ids: IRB-50878 CCT6003 (Other Identifier: OnCore) IRB-50878 (Other Identifier: Stanford IRB) NCI-2019- 07285 (Other Identifier: NCI Trial Identifier)	Suspended	B Cell Lymphom a Acute Lymphobl astic Leukemia, Pediatric Lymphom a	 Drug: Fludarabine Fludarabine is a purine antagonist antimetabolite Drug: Cyclophosphamide Cyclophosphamide is a nitrogen mustard derivative alkylating agent Drug: Autologous CD22 CAR T Autologous T cells transduced with lentiviral vector (m971BBZ) Chimeric Antigen Receptor (CD22 CAR) 	Study Type: Interventional Phase: Phase 1 Study Design: Allocation: Non-Randomized Intervention Model: Parallel Assignment Masking: None (Open Label) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 52 Original Estimated Enrollment: Same as current Age: 1 Year to 30 Years (Child, Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: January 10, 2020 Primary Completion: August 2025 (Final data collection date for primary outcome measure) Study Completion: August 2035 First Posted: September 13, 2019 Results First Posted: Last Update Posted: September 13, 2022
22	NCT0518 1540	A Study of the Effects of AB- 205 in Patients With Lymphoma Undergoing Autologous Hematopoietic Cell Transplantation Study Documents:	Title Acronym: Other Ids: AB-205-301	Recruiting	 Hodgkin Lymphom a Non Hodgkin Lymphom a 	Biological: AB-205 Allogeneic genetically engineered human umbilical vein endothelial cells Other Name: E-CEL cells Other: Placebo Placebo	Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Primary Purpose: Treatment Primary Outcome Measures: Same as current Secondary Outcome Measures: Same as current	Actual Enrollment: Estimated Enrollment: 148 Original Estimated Enrollment: Same as current Age: 40 Years and older (Adult, Older Adult) Sex: All	Study Sponsors: Same as current Collaborators: Not Provided	Study Start: February 21, 2022 Primary Completion: June 2025 (Final data collection date for primary outcome measure) Study Completion: December 2025 First Posted: January 6, 2022 Results First Posted: Last Update Posted: September 10, 2022

	NCT Number	Title	Other Names	Status	Conditions	Interventions	Characteristics	Population	Sponsor/Collaborators	Dates
23	NCT0545 8297		Title Acronym: Other Ids: 2140-	Recruiting	Chronic Lymphocy tic Leukemia Mantle Cell Lymphom a Follicular Lymphom a Richter Transform ation Lymphom a	Biological: Zilovertamab vedotin IV infusion	Study Type: Interventional Phase: Phase 2	Enrollment: S	Study Sponsors: Same as current	Study Start: July 21, 2022
			006 MK-2140-006 (Other Identifier: Merck) 2021-004450-36 (EudraCT Number)			Other Name: MK-2140 • Drug: Nemtabrutinib	Study Design: Allocation: Randomized Intervention Model: Parallel Assignment		Collaborators: Not Provided	Primary Completion: March 13, 2027 (Final data collection date for primary outcome measure)
						65 to 80 mg once daily (QD) orally Other Name: MK-1026	Masking: None (Open Label) Primary Purpose: Treatment			
							Primary Outcome Measures: Same as current			
							Secondary Outcome Measures: Same as current			Study Completion: April 26, 2027
										First Posted: July 14, 2022
										Results First Posted:
										Last Update Posted: September 10, 2022
24	NCT0001 2545		Title Acronym: Other Ids: 010122 01-H-0122	Recruiting	Sickle Cell Disease Sickle Cell Trait	Not Provided	Study Type: Observational	Actual Enrollment: Estimated Enrollment: 352 Original Estimated Enrollment:	Study Sponsors: National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) Collaborators:	Study Start: November 1,
							Phase:			2001
							Study Design: Observational Model: Case-Only Time Perspective: Cross-Sectional			Primary Completion: Not Provided
							Primary Outcome Measures: Not Provided Secondary Outcome Measures: Not Provided			Study
		Documents:						Age: 18 Years	Not Provided	Completion: Not Provided
								to 45 Years (Adult)		First Posted: March 12, 2001
								Sex: All		Results First Posted:
										Last Update Posted: September 13, 2022