## ClinicalTrials.gov: cell therapy | Last update posted in the last 7 days

|   | 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) | Study Type: Interventional  Phase: Phase 1  Study Design: Allocation: N/A Intervention Model: Single Group Assignment Intervention Model Description:  | 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 16, 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 27, 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 22, 2022 |

|   | NCT Number      | Title  | Other Names  | Status                 | Conditions   | Interventions   | Characteristics   | Population   | Sponsor/Collaborators  | Dates   |
|---|-----------------|--|--|------------------------|--|---|---|--|--|---|
| 3 | NCT0332<br>5101 | Dendritic Cell Therapy After Cryosurgery in Combination With Pembrolizumab in Treating Patients With Stage III-IV Melanoma That Cannot Be Remove by Surgery Study Documents: | Title Acronym:  Other Ids: MC1771 NCI-2017- 01967 (Registry Identifier: CTRP (Clinical Trial Reporting Program)) MC1771 (Other Identifier: Mayo Clinic) P30CA015083 ( U.S. NIH Grant/Contract) | Active, not recruiting | Stage III Cutaneous Melanoma AJCC v7  Stage IIIA Cutaneous Melanoma AJCC v7  Stage IIIB Cutaneous Melanoma AJCC v7  Stage IIIC Cutaneous Melanoma AJCC v7  Stage IV Cutaneous Melanoma AJCC v7  and v7 | <ul> <li>Procedure: Cryosurgery Undergo cryosurgery Other Names: <ul> <li>cryoablation</li> <li>cryosurgical ablation</li> </ul> </li> <li>Biological: Pembrolizumab Given IV Other Names: <ul> <li>Keytruda</li> <li>Lambrolizumab</li> <li>MK-3475</li> <li>SCH 900475</li> </ul> </li> <li>Procedure: Pheresis Undergo apheresis Other Names: <ul> <li>Apheresis</li> <li>Blood Component Removal</li> <li>Collection, Apheresis/Leukapheresis</li> <li>Hemapheresis</li> </ul> </li> <li>Biological: Therapeutic Autologous Dendritic Cells Given IT</li> </ul> | 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: 7  Estimated Enrollment:  Original Estimated Enrollment: 39  Age: 18 Years and older (Adult, Older Adult)  Sex: All                 | Study Sponsors:  Same as current  Collaborators: National Cancer Institute (NCI)                           | Study Start: November 15, 2017  Primary Completion: October 31, 2022 (Final data collection date for primary outcome measure)  Study Completion: October 31, 2022  First Posted: October 30, 2017  Results First Posted: Last Update Posted: September 22, 2022 |
| 4 | NCT0554<br>9921 | Phase II Study of TAEST16001 in Soft Tissue Sarcoma Study Documents:   | Title Acronym: Other Ids: TAEST16001 Phase II  | Recruiting             | Soft Tissue<br>Sarcoma   | Biological: NY-ESO-1(TCR Affinity Enhancing Specific T cell Therapy) NY-ESO-1(TCR Affinity Enhancing Specific T cell Therapy)   | Study Type: Interventional  Phase: 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: 70  Original Estimated Enrollment: Same as current  Age: 18 Years to 70 Years (Adult, Older Adult)  Sex: All | Study Sponsors:  Same as current  Collaborators: Guangdong Xiangxue Precision Medical Technology Co., Ltd. | Study Start: July 8, 2022  Primary Completion: September 1, 2024 (Final data collection date for primary outcome measure)  Study Completion: September 1, 2024  First Posted: September 22, 2022  Results First Posted: Last Update Posted: September 22, 2022  |

| NCT Number Ti   | tle Other Names  | Status     | Conditions  | Interventions  | Characteristics  | Population  | Sponsor/Collaborat<br>ors  | Dates  |
|-----------------|--|------------|---|--|--|---|--|--|
| NCT0535<br>9211 | NKTR-255 in Combination With CAR-T Cell Therapy for the Treatment of Relapsed or Refractory Large B-cell Lymphoma Study Documents:  Study Consortium  Study Documents:  Title Acronym: Other Ids: RG1122036 NCI-2022- 02316 ( Registry Identifier: CTRP (Clinical Trial Reporting Program)) 10802 ( Other Identifier: Fred Hutch/University of Washington Cancer Consortium) | Recruiting | Recurrent Diffuse Large B- Cell Lymphom a Recurrent Diffuse Large B- Cell Lymphom a, Not Otherwise Specified Recurrent Grade 3b Follicular Lymphom a Recurrent Primary Mediastina I (Thymic) Large B- Cell Lymphom a Refractory Diffuse Large B- Cell Lymphom a Transform ed Indolent B-Cell Non- Hodgkin Lymphom a Transform ed Indolent B-Cell Non- Hodgkin Lymphom a to Diffuse | Prug: Cyclophosphamide Given IV Other Names: ○ (-)-Cyclophosphamide ○ 2H-1,3,2-Oxazaphosphorine, 2-[bis(2-chloroethyl)amino]tetrahydro-, 2-oxide, monohydrate ○ Carloxan ○ Ciclofosfamida ○ Ciclofosfamide ○ Ciclofosfamide ○ Clafen ○ Clafen ○ Claphene ○ CP monohydrate ○ CTX ○ CYCLO-cell ○ Cycloblastin ○ Cyclophosphamide Monohydrate ○ Cyclophosphamide Monohydrate ○ Cyclophosphamidum ○ Cyclophosphamidum ○ Cyclophosphane ○ Cyclostine ○ Cytophosphane | Characteristics  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: 24  Original Estimated Enrollment:  Same as current  Age: 18 Years and older (Adult, Older Adult)  Sex: All | Study Sponsors: Same as current Collaborators: Nektar Therapeutics | Study Start: October 20, 2022  Primary Completion: January 31, 2024 (Final data collection date for primary outcome measure)  Study Completion: December 31, 2024  First Posted: May 3, 2022  Results First Posted: Last Update Posted: September 22, 2022 |

|   | NCT Number      | Title   | Other Names                        | Status             | Conditions  | Interventions  | Characteristics  | Population   | Sponsor/Collaborators   | Dates  |
|---|-----------------|---|------------------------------------|--------------------|---|--|--|--|---|--|
| 6 | NCT0463<br>7763 | CRISPR-Edited Allogeneic Anti- CD19 CAR-T Cell Therapy for Relapsed/Refract ory B Cell Non- Hodgkin Lymphoma Study Documents: | Title Acronym: Other Ids: CB10A    | Recruiting         | <ul> <li>Lymphom <ul> <li>a, Non-Hodgkin</li> </ul> </li> <li>Relapsed <ul> <li>Non</li> <li>Hodgkin</li> <li>Lymphom</li> <li>a</li> </ul> </li> <li>Refractory <ul> <li>B-Cell</li> <li>Non-Hodgkin</li> <li>Lymphom</li> <li>a</li> <li>Lymphom</li> <li>a</li> </ul> </li> <li>Lymphom <ul> <li>a</li> <li>Edl</li> <li>Lymphom</li> <li>a</li> </ul> </li> <li>B Cell</li> <li>Non-Hodgkin's</li> <li>Lymphom</li> </ul> <li>B Cell <ul> <li>Non-Hodgkin's</li> <li>Lymphom</li> </ul> </li> | Genetic: CB-010 CB-010 is a CRISPR-edited allogeneic CAR-T cell therapy targeting CD19.  Drug: Cyclophosphamide Chemotherapy for lymphodepletion  Drug: Fludarabine Chemotherapy for lymphodepletion | Study Type: Interventional  Phase: Phase 1  Study Design: Allocation: Non-Randomized Intervention Model: Sequential Assignment Intervention Model Description:  The CB10A clinical study consists of 3 + 3 design with three dose levels.  Masking: None (Open Label) Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Not Provided  | Actual Enrollment:  Estimated Enrollment: 50  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: May 26, 2021  Primary Completion: August 2025 (Final data collection date for primary outcome measure)  Study Completion: September 2025  First Posted: November 20, 2020  Results First Posted: Last Update Posted: September 19, 2022 |
| 7 | NCT0554<br>1549 | A Phase 2 Study Evaluating JCPyV-specific T Cell Therapy for PML Study Documents:   | Title Acronym: Other Ids: 20210001 | Not yet recruiting | Progressive<br>Multifocal<br>Leukoencephalo<br>pathy  | Biological: CE-VST01-JC CE-VST01-JC at a dose of 1 × 10^8 cells administered as an intravenous (IV) infusion every 28 days for 4 total infusions   | Study Type: Interventional  Phase: Phase 2  Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description:     randomized, double- blinded, Phase 2 trial in patients with PML due to JCPyV.  Masking: Triple (Participant, Care Provider, Investigator) Primary Purpose: Treatment  Primary Outcome Measures: To evaluate the effect of CE-VST01-JC on time to disease progression, as measured by mRS (modified Rankin Score) [ Time Frame: 1 year ] Time to progression as measured by mRS. A progression event is defined as an increase of 2 points on mRS attributable to disease progression* that is durable (not reversed over two consecutive measurements, at least 14 days apart), or an increase to mRS of 5 or 6 (severe disability or death, respectively).  Secondary Outcome Measures: Not Provided | Actual Enrollment: Estimated Enrollment: 60 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: February 2023  Primary Completion: March 2024 (Final data collection date for primary outcome measure)  Study Completion: April 2025  First Posted: September 15, 2022  Results First Posted: Last Update Posted: September 16, 2022    |

|   | NCT Number      | Title  | Other Names                       | Status     | Conditions                             | Interventions   | Characteristics   | Population  | Sponsor/Collaborat  | Dates  |
|---|-----------------|--|-----------------------------------|------------|--|---|---|---|---|--|
| 8 | NCT0444<br>5454 | Mesenchymal Stromal Cell Therapy for Severe Covid- 19 Infection Study Documents:   | Title Acronym: Other Ids: TJT2012 | Recruiting | Coronavirus Infection                  | Biological: Mesenchymal stromal cells  Bone marrow collection and MSC expansion cultures will be carried out at the Laboratory of Cell and Gene Therapy (LTCG) at the University of Liège as described in IMPD and its SOPs.  Other Name: MSC | Study Type: Interventional  Phase: Phase 1 Phase 2  Study Design: Allocation: N/A Intervention Model: Single Group Assignment Intervention Model Description:  This study is a monocentric prospective phase I/II clinical trial, aiming at evaluating the safety and efficacy of 3 intravenous administrations of BM-MSC in 20 patients with severe to critical COVID-19 pneumonia.  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: Not Provided           | Study Start: June 12, 2020  Primary Completion: September 30, 2024 (Final data collection date for primary outcome measure)  Study Completion: September 30, 2024  First Posted: June 24, 2020  Results First Posted: Last Update Posted: September 21, 2022 |
| 9 | NCT0459<br>9556 | Clinical Trial for the Safety and Efficacy of Anti-CD7 CAR- T Cell Therapy for Patients With Relapsed or Refractory CD7 Positive Hematological Malignancy Study Documents: | Title Acronym: Other Ids: CD7-001 | Recruiting | • CD7+ Acute Leukemia • CD7+ Lymphom a | Biological: anti-CD7 CAR-T Lymphodepleting chemotherapy followed by anti-CD7 CAR-T infusion   | Study Type: Interventional  Phase: Phase 1 Phase 2  Study Design: Allocation: Non-Randomized Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Not Provided   | Actual Enrollment: Estimated Enrollment: 108 Original Estimated Enrollment: Same as current Age: 3 Years to 80 Years (Child, Adult, Older Adult) Sex: All | Study Sponsors: Same as current  Collaborators: Yake Biotechnology Ltd. | Study Start: November 20, 2021  Primary Completion: December 2022 (Final data collection date for primary outcome measure)  Study Completion: December 2023  First Posted: October 22, 2020  Results First Posted: Last Update Posted: September 21, 2022    |

| NCT Number      | Title                                       | Other Names  | Status     | Conditions             | Interventions   | Characteristics   | Population                           | Sponsor/Collabora<br>ors   | t Dates                                     |  |  |
|-----------------|---|--|------------|------------------------|---|---|--------------------------------------|--|---|--|--|
| NCT0435<br>9784 | Anakinra for the Prevention of              | Title Acronym:   | Recruiting | B-Cell Non-<br>Hodgkin | Biological: Anakinra<br>Given SC  | Study Type: Interventional  | Actual Enrollment:                   | Study Sponsors: Same as current  | Study Start:<br>December 27                 |  |  |
|                 | Cytokine Release Syndrome and               | Other Ids:<br>RG1006866<br>NCI-2020-                             |            | Lymphoma               | Other Names: • Kinaret  | Phase: Phase 2 Study Design: Allocation: N/A  | Estimated Enrollment: 25             | Collaborators:<br>Swedish Orphan   | 2021<br>Primary                             |  |  |
|                 | Neurotoxicity in Patients With B- Cell Non- | 01861 ( Registry<br>Identifier:<br>CTRP (Clinical                |            |                        | <ul> <li>Kineret</li> <li>rIL-1ra</li> <li>rIL1RN</li> <li>143090-92-0</li> </ul> | Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Prevention  | Original Estimated Enrollment:       | Biovitrum  | Completion:<br>March 3, 202<br>(Final data  |  |  |
|                 | Hodgkin Lymphoma Receiving                  | Trial Reporting<br>Program))<br>10373 (Other<br>Identifier: Fred |            |                        | 143070-72-0   | Primary Outcome Measures: Absence of any grade cytokine release syndrome (CRS) [ Time Frame: Up to 90 days after axicabtagene ciloleucel (Axi-cell) infusion ]                              | Age: 18 Years                        |  | for primary outcome measure)                |  |  |
|                 | CD19-Targeted CAR-T Cell Therapy            | Hutch/University of Washington Cancer                            |            |                        |   | Will assess the efficacy of anakinra in preventing the occurrence of any grade CRS using the Bayesian optimal phas 2 design. Assessed based on the ASTCT Consensus Grading                  | and older<br>(Adult, Older<br>Adult) |  | Study<br>Completion:                        |  |  |
|                 | Study<br>Documents:                         | Consortium )   |            |                        |   | for CRS and Neurotoxicity Associated with Immune Effector Cell.   | Sex: All                             |  | March 3, 20 First Posted                    |  |  |
|                 |   |  |            |                        |   | <ul><li>Secondary Outcome Measures:</li><li>CRS grade [ Time Frame: Up to 90 days after Axi-cell infusion ]</li></ul>   |                                      |  | April 24, 20 Results Firs Posted:           |  |  |
|                 |   |  |            |                        |   | Graded according to the American Society for Transplantation and Cellular Therapy (ASTCT) Consensus Grading for CRS and Neurotoxicity Associated with Immune Effector Cell.                 |                                      |  | Last Update<br>Posted:<br>September<br>2022 |  |  |
|                 |   |  |            |                        |   | <ul> <li>Neurotoxicity grade [ Time Frame: Up to 90 days after<br/>Axi-cell infusion ]</li> </ul>   |                                      |  | 2022  |  |  |
|                 |   |  |            |                        |   |   |                                      | Graded according to the American Society for<br>Transplantation and Cellular Therapy (ASTCT)<br>Consensus Grading for CRS and Neurotoxicity<br>Associated with Immune Effector Cell. |   |  |  |
|                 |   |  |            |                        |   | <ul> <li>Disease response to Axi-cel [ Time Frame: At 28 and 90 days after Axi-cell infusion ]</li> </ul>   |                                      |  |   |  |  |
|                 |   |  |            |                        |   | Objective responses to the therapeutic regimen will be assessed based on institutional standard using physical examination, imaging (CT or PET-CT), and if necessary, bone marrow biopsies. |                                      |  |   |  |  |
|                 |   |  |            |                        |   | <ul> <li>Adverse events (AEs) [ Time Frame: Within 28 days<br/>after Axi-cell infusion ]</li> </ul>   |                                      |  |   |  |  |
|                 |   |  |            |                        |   | Graded according to the National Cancer Institute<br>Common Terminology Criteria for Adverse Events<br>version 5.0.   |                                      |  |   |  |  |

|    |                 |                              |                               | _          |                        |   |   |                              | Sponsor/Collaborat              | _                                   |
|----|-----------------|------------------------------|-------------------------------|------------|------------------------|---|---|------------------------------|---------------------------------|-------------------------------------|
|    | NCT Number      | Title                        | Other Names                   | Status     | Conditions             | Interventions   | Characteristics   | Population                   | ors                             | Dates                               |
| 11 | NCT0369<br>6030 | HER2-CAR T Cells in Treating | Title Acronym:                | Recruiting | Malignant              | Biological: Chimeric Antigen Receptor T-Cell Therapy                    | Study Type: Interventional  | Actual Enrollment:           | Study Sponsors: Same as current | Study Start:<br>August 31, 2018     |
|    | 0030            | Patients With                | Other Ids: 17237<br>NCI-2018- |            | Neoplasm  • Metastatic | Given HER2-CAR T cells via intraventricular administration Other Names: | Phase: Phase 1  | Estimated                    | Collaborators:                  | Primary                             |
|    |                 | Recurrent Brain or           | 01270 (Registry               |            | Malignant              | <ul><li>CAR T Infusion</li><li>CAR T Therapy</li></ul>                  | Study Design: Allocation: Non-Randomized<br>Intervention Model: Single Group Assignment                                       | Enrollment: 39               | National                        | Completion:                         |
|    |                 | Leptomeningeal<br>Metastases | Identifier:<br>CTRP (Clinical |            | Neoplasm in the        | CAR T-cell therapy  | Masking: None (Open Label)  | Original                     | Cancer<br>Institute             | August 31,<br>2023 (Final           |
|    |                 | Study                        | Trial Reporting<br>Program))  |            | Brain  • Metastatic    | Chimeric Antigen Receptor T-cell Infusion                               | Primary Purpose: Treatment  | Estimated<br>Enrollment: 21  | (NCI)                           | data collection<br>date for primary |
|    |                 | Documents:                   | 17237 (Other Identifier: City |            | Malignant              |   | Primary Outcome Measures:  • Incidence of dose limiting toxicities (DLTs) [ Time  | Age: 18 Years                | • California Institute          | outcome<br>measure)                 |
|    |                 |                              | of Hope<br>Medical Center )   |            | Neoplasm in the        |   | Frame: 21 days post T cell infusion ]   | to 75 Years<br>(Adult, Older | for<br>Regenerati               | Study                               |
|    |                 |                              | Wedical Center)               |            | Leptomeni<br>nges      |   | Rate and associated 90% Clopper and Pearson binomial confidence limits (90% CI) will be estimated for                         | Adult)                       | ve<br>Medicine                  | Completion:<br>August 31, 2023      |
|    |                 |                              |                               |            | • Breast               |   | participants experiencing DLTs at the recommended phase 2 dose schedule.  | Sex: All                     | (CIRM)                          | First Posted:                       |
|    |                 |                              |                               |            | Cancer • HER2-         |   | Number of participants with treatment related adverse   |                              |                                 | October 4, 2018                     |
|    |                 |                              |                               |            | positive<br>Breast     |   | events as assessed by CTCAE v5.0. [ Time Frame: Up to 15 years ]  |                              |                                 | Results First<br>Posted:            |
|    |                 |                              |                               |            | Cancer                 |   | Tables will be created to summarize all toxicities and side effects by dose, time post treatment, organ, severity             |                              |                                 | Last Update                         |
|    |                 |                              |                               |            |                        |   | and arm.  |                              |                                 | Posted:<br>September 16,            |
|    |                 |                              |                               |            |                        |   | Secondary Outcome Measures:   |                              |                                 | 2022                                |
|    |                 |                              |                               |            |                        |   | <ul> <li>HER2-CAR T cells in cerebrospinal fluid (CSF) and<br/>peripheral blood [ Time Frame: Measured over time</li> </ul>   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | from baseline through 1 year, the number of measurements is determined by whether or not the                                  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | participant has progressed (progressed: baseline, 1, 3,   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | 6,and 12 months, not progressed: baseline, 1, 3,6,8,10 and 12 months) ]   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | Statistical and graphical methods will be used to describe the data.  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | <ul> <li>Endogenous B cells in cerebrospinal fluid (CSF) and<br/>peripheral blood [ Time Frame: Measured over time</li> </ul> |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | from baseline through 1 year, the number of   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | measurements is determined by whether or not the participant has progressed (progressed: baseline, 1, 3,                      |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | 6,and 12 months, not progressed: baseline, 1, 3,6,8,10 and 12 months) ]   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | Statistical and graphical methods will be used to describe the data.  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | • T cells in cerebrospinal fluid (CSF) and peripheral blood [Time Frame: progressed: baseline, 1, 3, 6, and 12                |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | months, not progressed: baseline, 1, 3, 6, 8,10 and 12  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | months) ] Statistical and graphical methods will be used to   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | describe the data.  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | <ul> <li>Myeloid cells in cerebrospinal fluid (CSF) and<br/>peripheral blood [ Time Frame: Measured over time</li> </ul>      |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | from baseline through 1 year, the number of measurements is determined by whether or not the                                  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | participant has progressed (progressed: baseline, 1, 3, 6, and 12 months, not progressed: baseline, 1, 3, 6, 8, 10            |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | and 12 months) ]  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | Statistical and graphical methods will be used to describe the data.  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | Host immune subsets (e.g. T cell inhibitory/exhaustion<br>markers, activation markers, and effector memory T                  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | cells) in cerebrospinal fluid (CSF) and peripheral blood.   |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | [ Time Frame: Measured over time from baseline through 1 year, the number of measurements is                                  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | determined by whether or not the participant has progressed (progressed: baseline, 1, 3, 6, and 12 months,                    |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | not progressed: baseline, 1, 3,6,8,10 and 12 months) ]  |                              |                                 |                                     |
|    |                 |                              |                               |            |                        |   | Statistical and graphical methods will be used to describe the data.  |                              |                                 |                                     |

|    | NCT Number      | Title  | Other Names                                   | Status                 | Conditions   | Interventions  | Characteristics   | Population  | Sponsor/Collaborators  | Dates  |
|----|-----------------|--|---|------------------------|--|--|---|---|--|--|
| 12 | NCT0513<br>5091 | FIH Study of NRTX-1001 Neural Cell Therapy in Drug-Resistant Unilateral Mesial Temporal Lobe Epilepsy Study Documents: | Title Acronym: Other Ids: NTE001              | Recruiting             | Mesial<br>Temporal Lobe<br>Epilepsy With<br>Hippocampal<br>Sclerosis                 | Biological: NRTX-1001 is an experimental neural cell therapy product candidate derived from an allogeneic human embryonic stem cell line. The stem cells were converted into inhibitory nerve cells that produce GABA.  Other Name: GABA-secreting interneurons  Procedure: Sham Comparator Sham Comparator. | Phase: Phase 1 Phase 2  Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description:  This is a two-stage study. Stage 1 is an open-label, single arm, sequential dose escalation. Stage 2 is a parallel, randomized, 2-arm, sham controlled study.  Masking: Triple (Participant, Investigator, Outcomes Assessor) Masking Description:  This is a two-stage study. Stage 1 is open-label and unmasked. Stage 2 is masked with participant, part of investigator team, and outcomes assessor masked to treatment assignment.  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 to 55 Years (Adult)  Sex: All              | Study Sponsors:  Same as current  Collaborators: California Institute for Regenerative Medicine (CIRM) | Study Start: June 16, 2022  Primary Completion: May 2025 (Final data collection date for primary outcome measure)  Study Completion: May 2026  First Posted: November 26, 2021  Results First Posted: Last Update Posted: September 21, 2022                     |
| 13 | NCT0382<br>7343 | Retrospective Study of Immunotherapy Related Toxicities in Children and Adults With Cancer Study Documents:            | Title Acronym: Other Ids: 999919044 19-C-N044 | Active, not recruiting | Macropha ge     Activation Syndrome     Primary Hemophag ocytic Lymphohi stiocytosis | Not Provided   | Phase:  Study Design: Observational Model: Cohort Time Perspective: Retrospective  Primary Outcome Measures: To develop a retrospective study to allow for comparison of immunotherapy related toxicity profiles and risk factors across a set of protocols in the NCI. [ Time Frame: 2 years ]  To develop a retrospective study to allow for comparison of immunotherapy related toxicity profiles and risk factors across a set of protocols in the NCI  Secondary Outcome Measures:  • Evaluate the incidence, risk factors for, and treatment of HLH/MAS in patients who receive CAR-T cell therapy [ Time Frame: 2 years ]  Evaluate the incidence, risk factors for, and treatment of HLH/MAS in patients who receive CAR-T cell therapy  • Evaluate infectious complications and their risk factors in patients who receive CAR-T cell therapy for cancer [ Time Frame: 2 years ]  Evaluate infectious complications and their risk factors in patients who receive CAR-T cell therapy for cancer [ | Actual Enrollment: 500 Estimated Enrollment: Original Estimated Enrollment: Same as current Age: 1 Month and older (Child, Adult, Older Adult) Sex: All | Study Sponsors: Same as current Collaborators: Not Provided  | Study Start: January 23, 2019  Primary Completion: December 31, 2025 (Final data collection date for primary outcome measure)  Study Completion: December 31, 2025  First Posted: February 1, 2019  Results First Posted: Last Update Posted: September 21, 2022 |

|    | NCT Number      | Title   | Other Names  | Status             | Conditions   | Interventions   | Characteristics  | Population  | Sponsor/Collaborat  | Dates   |
|----|-----------------|---|--|--------------------|--|---|--|---|---|---|
| 14 | NCT0505<br>0006 | ITIL-168 in Advanced Melanoma  Study Documents:                           | Title Acronym: Other Ids: ITIL- 168-101 2020-003862-37 ( EudraCT Number )  | Recruiting         | Advanced<br>Cutaneous<br>Melanoma  | Biological: ITIL-168 ITIL-168 is a cell therapy product derived from a patient's own TILs. A tumor sample is removed from each patient to make a personalized ITIL-168 product. Once ITIL-168 has been made, the patient is treated with 5 days of lymphodepleting chemotherapy including cyclophosphamide and fludarabine, followed by a single infusion of ITIL-168, and up to 8 doses of IL-2. | Study Type: Interventional  Phase: Phase 2  Study Design: Allocation: Non-Randomized Intervention Model: Parallel Assignment Intervention Model Description:  All enrolled participants are assigned to be treated with a single dose of ITIL-168 Masking: None (Open Label) Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Same as current  | Actual Enrollment:  Estimated Enrollment: 130  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: October 7, 2021  Primary Completion: March 2024 (Final data collection date for primary outcome measure)  Study Completion: August 2028  First Posted: September 20, 2021  Results First Posted: Last Update Posted: September 16, 2022              |
| 15 | NCT0554<br>6723 | LMY-920 for Treatment of Relapsed or Refractory Myeloma  Study Documents: | Title Acronym:  Other Ids: LMY- 920-002 LUMT1A22 ( Other Identifier: Cleveland Clinic Taussig Cancer Institute ) | Not yet recruiting | Multiple     Myeloma,     Refractory     Multiple     Myeloma     in Relapse | Biological: Autologous CAR-T cell therapy expressing the BAFF-ligand. LMY-920   | Study Type: Interventional  Phase: Phase 1  Study Design: Allocation: N/A Intervention Model: Sequential Assignment Intervention Model Description:  Open label, dose escalation study with up to four dose levels of LMY-920. The maximum tolerated dose (MTD) of LMY-920 will be determined using dose-escalation 3+3 design.  Masking: None (Open Label) Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Same as current | 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:  The Cleveland Clinic  Case Western Reserve University | Study Start: November 1, 2022  Primary Completion: October 31, 2024 (Final data collection date for primary outcome measure)  Study Completion: December 31, 2024  First Posted: September 21, 2022  Results First Posted: Last Update Posted: September 21, 2022 |

|    | NCT Number      | Title   | Other Names          | Status                  | Conditions   | Interventions   | Characteristics  | Population                                     | Sponsor/Collaborators          | Dates  |
|----|-----------------|---|----------------------|-------------------------|--|---|--|--|--------------------------------|--|
| 16 | NCT0231<br>5599 | Follow-Up<br>Evaluation for                         | Title Acronym:       | Enrolling by invitation | Pediatric  | Not Provided  | Study Type: Observational  | Actual Enrollment:                             | Study Sponsors:                | Study Start:<br>December 23,                                   |
|    | 3399            | Gene-Therapy-                                       | Other Ids:<br>150028 | invitation              | Cancers  |   | Phase:   | Estimated                                      | Same as current Collaborators: | 2014   |
|    |                 | Adverse Events After                                | 15-C-0028            |                         | Hematolog     ic     Malignanc                         |   | Study Design: Observational Model: Cohort<br>Time Perspective: Prospective   | Enrollment: 500                                | Not Provided                   | Primary<br>Completion:   |
|    |                 | Participation in Pediatric Oncology Branch Clinical |                      |                         | ies • Solid Tumors                                     |   | Primary Outcome Measures: Conduct long term safety evaluations after gene therapy [ Time Frame: Every 3 months X 1 year then annually X 15 years ] | Original Estimated Enrollment: Same as current |                                | April 1, 2035<br>(Final data<br>collection date<br>for primary |
|    |                 | Trials Study  |                      |                         |  |   | Secondary Outcome Measures: Not Provided   | Age: 1 Year to<br>99 Years<br>(Child, Adult,   |                                | outcome<br>measure)  |
|    |                 | Documents:  |                      |                         |  |   |  | Older Adult)  Sex: All                         |                                | Study<br>Completion:<br>August 1, 2050                         |
|    |                 |   |                      |                         |  |   |  | Jon III  |                                | First Posted:<br>December 12,<br>2014                          |
|    |                 |   |                      |                         |  |   |  |  |                                | Results First Posted:  |
|    |                 |   |                      |                         |  |   |  |  |                                | Last Update<br>Posted:<br>September 21,<br>2022                |
| 17 | NCT0547         | Dual-targeting                                      | Title Acronym:       | Recruiting              | Malignant  | Biological: Dual-targeting VEGFR1 and PD-L1 CAR-T   | Study Type: Interventional   | Actual   | Study Sponsors:                | Study Start:   |
|    | 7927            | VEGFR1 and<br>PD-L1 CAR-T                           | Other Ids:           |                         | Peritoneal<br>Effusion                                 | cells  In the dose escalation part, the dose levels will be escalated   | Phase: Phase 1   | Enrollment:                                    | Same as current                | October 30,<br>2022  |
|    |                 | Patients With Pleural or Peritoneal Metastases      | MCART-006            |                         | <ul><li>Malignant<br/>Ascites</li><li>Serous</li></ul> | following a traditional escalation scheme for 3+3 design.  In the dose expansion part, patients will be assigned to different groups based on pleural or peritoneal metastases condition. | Study Design: Allocation: N/A Intervention Model: Single Group Assignment Masking: None (Open Label) Primary Purpose: Treatment                    | Estimated Enrollment: 58  Original Estimated   | Collaborators:<br>Not Provided | Primary Completion: December 31, 2024 (Final                   |
|    |                 | Study   |                      |                         | Cavity<br>Metastatis                                   |   | Primary Outcome Measures: Same as current  | Enrollment: Same as current                    |                                | data collection<br>date for primary                            |
|    |                 | Documents:  |                      |                         | es   |   | Secondary Outcome Measures: Same as current  | Age: 18 Years to 65 Years                      |                                | outcome<br>measure)  |
|    |                 |   |                      |                         |  |   |  | (Adult, Older<br>Adult)                        |                                | Study<br>Completion:   |
|    |                 |   |                      |                         |  |   |  | Sex: All                                       |                                | December 31,<br>2024   |
|    |                 |   |                      |                         |  |   |  |  |                                | First Posted:<br>July 28, 2022                                 |
|    |                 |   |                      |                         |  |   |  |  |                                | Results First<br>Posted:                                       |
|    |                 |   |                      |                         |  |   |  |  |                                | Last Update<br>Posted:<br>September 19,<br>2022                |

|    | NCT Number      | Title  | Other Names                                | Status     | Conditions   | Interventions   | Characteristics   | Population  | Sponsor/Collabora<br>ors   | t Dates   |
|----|-----------------|--|--|------------|--|---|---|---|--|---|
| 18 | NCT0000<br>1405 | Recruitment and Apheresis Collection of Peripheral Blood Hematopoietic Stem Cells, Mononuclear Cells and Granulocytes Study Documents: | Title Acronym: Other Ids: 940073 94-I-0073 | Recruiting | <ul> <li>Granulom a</li> <li>Granulom atous Disease, Chronic</li> <li>Leukocyte Disease</li> <li>Genetic Disease, X-Linked</li> <li>Genetic Disease, Inborn</li> </ul> | Not Provided  | Study Design: Observational Model: Cohort Time Perspective: Other  Primary Outcome Measures: Not Provided  Secondary Outcome Measures: Not Provided   | 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, 1994  Primary Completion: Not Provided  Study Completion: Not Provided  First Posted: November 4, 1999  Results First Posted: Last Update Posted: September 19, 2022  |
| 19 | NCT0506<br>8674 | Human Embryonic Stem Cell- Derived Cardiomyocyte Therapy for Chronic Ischemic Left Ventricular Dysfunction Study Documents:            | Title Acronym: Other Ids: 60978            | Recruiting | Chronic Ischemic Left Ventricular Dysfunction  | <ul> <li>Drug: Human Embryonic Stem Cell-Derived Cardiomyocyte 50M cells 50 million (M) cells delivered in a dose of 5M cells per injection over 10 injections. Other Name: Human ESC-CMs</li> <li>Drug: Human Embryonic Stem Cell-Derived Cardiomyocyte 150 cells 150M cells delivered in a dose of 15M cells per injection over 10 injections Other Name: Human ESC-CMs</li> <li>Drug: Human Embryonic Stem Cell-Derived Cardiomyocyte 300M cells 300M cells delivered in a dose of 30M per injection over 10 injections Other Name: Human ESC-CMs</li> </ul> | Study Type: Interventional  Phase: Phase 1  Study Design: Allocation: Randomized Intervention Model: Sequential Assignment Intervention Model Description: Phase I will be a standard 3+3 dose-escalation study to evaluate 3 doses of allogeneic hESC-CMs Masking: None (Open Label) Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Not Provided | Actual Enrollment:  Estimated Enrollment: 18  Original Estimated Enrollment:  Same as current  Age: 21 Years to 80 Years (Adult, Older Adult)  Sex: All | Study Sponsors:  Same as current  Collaborators: California Institute for Regenerative Medicine (CIRM) | Study Start: March 22, 2022  Primary Completion: October 2025 (Final data collection date for primary outcome measure)  Study Completion: October 2025  First Posted: October 6, 2021  Results First Posted: Last Update Posted: September 19, 2022 |

|    | NCT Number      | Title  | Other Names                           | Status             | Conditions  | Interventions  | Characteristics   | Population   | Sponsor/Collaborat  | Dates  |
|----|-----------------|--|---------------------------------------|--------------------|---|--|---|--|---|--|
| 20 | NCT0400<br>5989 | Adipose Stromal Cells Injection in the Myocardium for Induction of Revascularizatio n Study Documents: | Title Acronym: Other Ids: ADMIRE      | Withdrawn          | Ischemic Heart Disease     Myocardia I Ischemia     Coronary Artery Disease | Biological: Stromal Cells Injection Adipose stromal cell   | Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description:  It is a randomized, prospective, double-blind, placebocontrolled clinical study.  Eligible patients will be randomly divided into 4 groups (one control group and 3 active treatment groups): 10 PATIENTS EACH ARM Control Group - saline injection Low dose group (BD) - hASC injection (1x106 / kg body weight) Intermediate dose group (DI) - injection of hASC (2x106 / kg body weight) High dose group (AD) - hASC injection (4x106 / kg body weight)  Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor) Masking Description: DOUBLE BLIND Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Same as current | Actual Enrollment: 0  Estimated Enrollment:  Original Estimated Enrollment: 40  Age: 18 Years to 80 Years (Adult, Older Adult)  Sex: All | Study Sponsors:  Same as current  Collaborators: Not Provided | Study Start: December 1, 2021  Primary Completion: November 30, 2022 (Final data collection date for primary outcome measure)  Study Completion: December 20, 2022  First Posted: July 2, 2019  Results First Posted: Last Update Posted: September 22, 2022 |
| 21 | NCT0554<br>4526 | CAR T Cells to Target GD2 for DMG Study Documents:   | Title Acronym: Other Ids: UCL/ 150853 | Not yet recruiting | Diffuse Midline<br>Glioma, H3<br>K27M-Mutant                                | Biological: GD2 CAR T cells Infusion with: GD2 CAR T-cells | 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: 12  Original Estimated Enrollment:  Same as current  Age: up to 16 Years (Child)  Sex: All     | Study Sponsors:  Same as current  Collaborators: Not Provided | Study Start: December 2022  Primary Completion: December 2025 (Final data collection date for primary outcome measure)  Study Completion: December 2039  First Posted: September 16, 2022  Results First Posted: Last Update Posted: September 16, 2022      |

| NCT Number Title   | Other Names                       | Status       | Conditions   | Interventions   | Characteristics  | Population   | Sponsor/Collabora<br>ors                                     | t Dates   |
|--|-----------------------------------|--------------|--|---|--|--|--|---|
| NCT0545 1849  A Phase Trial of In Paties Advance Mesothe Express Cancer Study Docume | Other Ids: TCR2-21-01  TCR2-21-01 | : Recruiting | Mesotheli oma     Mesotheli omas Pleural     Mesotheli oma, Malignant     Mesotheli oma Peritoneu m     Ovarian Cancer     Ovarian Serous Adenocarc inoma     Pancreatic Cancer     Pancreatic Adenocarc inoma     Colorectal Cancer     Triple Negative Breast Cancer     Triple-Negative Breast Cancer     Triple-Negative Breast Cancer     Triple-Negative Breast Cancer     Ovarian Adenocarc inoma     Pancreatic Negative Breast Cancer     Ovarian Adenocarc inoma     Pancreatic Neoplasms     Colorectal Neoplasms     Ovarian Neoplasms | Biological: TC-510 TC-510  Drug: Fludarabine Fludarabine Drug: Cyclophosphamide Cyclophosphamide  Order  Biological: TC-510 TC-510  Drug: Fludarabine Fludarabine  Drug: Cyclophosphamide  Order  Biological: TC-510  Drug: Fludarabine Fludarabine Fludarabine  Drug: Cyclophosphamide  Order  Biological: TC-510  Drug: Fludarabine Fludarabine Fludarabine Fludarabine  Drug: Cyclophosphamide  Order  Biological: TC-510  Drug: Fludarabine Fludarabine Fludarabine Fludarabine  Drug: Cyclophosphamide  Order  Drug: Cyclophosphamide  Order  Drug: Cyclophosphamide  Order  Drug: Cyclophosphamide  Order  Drug: Order  Drug: Cyclophosphamide  Order  Drug: Order  Drug | 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: Not Provided | Actual Enrollment:  Estimated Enrollment: 115  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: June 21, 2022  Primary Completion: June 2025 (Final data collection date for primary outcome measure)  Study Completion: December 2027  First Posted: July 11, 2022  Results First Posted:  Last Update Posted: September 22, 2022 |

| NCT Number | Title                                   | Other Names           | Status     | Conditions               | Interventions   | Characteristics   | Population  | Sponsor/Collabora<br>ors        | Dates Dates   |
|------------|---|-----------------------|------------|--------------------------|---|---|---|---------------------------------|---|
| NCT0462    | NEO-PTC-01 in Patients With Advanced or | Other Ids: NTC-       | Recruiting | • Unresecta ble Melanoma | Biological: NEO-PTC-01<br>Administered via intravenous (IV) infusion. | Study Type: Interventional  | Actual  | Study Sponsors: Same as current | December 1, 2020  Primary Completion: November 2023 (Final data collectio date for prima outcome measure)  Study Completion: November 20  First Posted: November 12 |
| 5205       |   |                       |            |                          |   | Phase: Phase 1  | Enrollment:  Estimated Enrollment: 52  Original Estimated |                                 |   |
|            | Metastatic<br>Melanoma                  | 001<br>2019-003908-13 |            | Metastatic               |   | Study Design: Allocation: Non-Randomized<br>Intervention Model: Sequential Assignment<br>Masking: None (Open Label)<br>Primary Purpose: Treatment   |   | Collaborators:<br>Not Provided  |   |
|            | Study<br>Documents:                     | ( EudraCT<br>Number ) |            | Melanoma                 |   |   |   |                                 |   |
|            |   |                       |            |                          |   | Primary Outcome Measures: Rate of adverse events (AEs), including serious adverse events (SAEs) and AEs leading to treatment discontinuation [Time Frame: Day 1 to week 52]   | Age: 18 Years to 75 Years                                 |                                 |   |
|            |   |                       |            |                          |   | Rate of AEs, including SAEs and AEs leading to treatment discontinuation and those adverse events and serious adverse   | (Adult, Older Adult) Sex: All                             |                                 |   |
|            |   |                       |            |                          |   | events detected during symptom-directed physical<br>examinations (changes in safety laboratory evaluations,<br>physical examination findings, and vital signs.  |   |                                 |   |
|            |   |                       |            |                          |   | Secondary Outcome Measures:   |   |                                 |   |
|            |   |                       |            |                          |   | Progression-free survival (PFS), defined as the time from the date of first dosing of NEO-PTC-01 to the date      (C) (1) (1) (1) (2) (1) (2) (2) (3) (4) (4) (4) (4) (4) (4) (4) (4) (4) (4  |   |                                 | 2020<br>Results Fire  |
|            |   |                       |            |                          |   | of first documented progressive disease (PD) or death, whichever comes first [ Time Frame: Day 1 to week 52 ]   |   |                                 | Posted:   |
|            |   |                       |            |                          |   | Clinical activity endpoints, based on Investigator assessment of serial radiographic evaluations [Computed Tomography (CT) or Magnetic Resonance Imaging (MRI)] to determine response to treatment and progression of disease based on response criteria in solid tumors (RECIST) v1.1. |   |                                 | Last Upda<br>Posted:<br>Septembe:<br>2022   |
|            |   |                       |            |                          |   | <ul> <li>Overall response rate (ORR), defined as the proportion<br/>of patients who achieve complete response (CR) or<br/>partial response (PR) based on RECIST v1.1 [Time<br/>Frame: Day 1 to week 52]</li> </ul>  |   |                                 |   |
|            |   |                       |            |                          |   | <ul> <li>Duration of response (DOR), defined as the date of the<br/>first documentation of a confirmed response to the date<br/>of the first documented PD [ Time Frame: Day 1 to week<br/>52 ]</li> </ul>  |   |                                 |   |
|            |   |                       |            |                          |   | <ul> <li>Clinical benefit rate (CBR), defined as the proportion of<br/>patients who achieve CR, PR, or stable disease (SD)<br/>based on RECIST [ Time Frame: Day 1 to week 52 ]</li> </ul>  |   |                                 |   |
|            |   |                       |            |                          |   | • Time to first subsequent therapy, defined as the time from the date of first dosing to the start date of first subsequent therapy [ Time Frame: Day 1 to week 52 ]  |   |                                 |   |

|    | NCT Number Title |  | Other Names                                | Status     | Conditions   | Interventions  | Characteristics   | Population   | Sponsor/Collaborators   | Dates  |
|----|------------------|--|--|------------|--|--|---|--|---|--|
| 24 | NCT0554<br>8712  | The Effect of PEMF for Patients With Knee OA  Study Documents: | Title Acronym: Other Ids: 2021.491         | Recruiting | Knee Osteoarthr itis     Knee Pain Chronic   | <ul> <li>Device: Pulse Electromagnetic Field         Subjects will receive PEMF treatment with the         duration of 8 weeks, twice a week with total 16         treatment sessions.</li> <li>Device: Sham Pulse Electromagnetic Field         Subjects will receive sham PEMF treatment with the         duration of 8 weeks, twice a week with total 16         treatment sessions.</li> </ul>   | Phase: Not Applicable  Study Design: Allocation: Randomized Intervention Model: Parallel Assignment Intervention Model Description:  Participants will be assigned into either treatment group or sham group.  Masking: Triple (Participant, Care Provider, Investigator) Masking Description:  It is doubled blinded, a subject 's specific ID card will be provided for each of the enrolled subject and the investigators don't know whether it is PEMF or Sham treatment for that subject's specific ID card. And the investigator will ask the manufactory about the number of grouping at the end of the study.  Primary Purpose: Treatment  Primary Outcome Measures: Same as current  Secondary Outcome Measures: Same as current | Actual Enrollment:  Estimated Enrollment: 80  Original Estimated Enrollment: Same as current  Age: 50 Years and older (Adult, Older Adult)  Sex: All   | Study Sponsors:  Same as current  Collaborators: Not Provided | Study Start: May 5, 2022  Primary Completion: September 15, 2024 (Final data collection date for primary outcome measure)  Study Completion: September 15, 2024  First Posted: September 21, 2022  Results First Posted: Last Update Posted: September 21, 2022      |
| 25 | NCT0410<br>2436  | Non-Viral TCR Gene Therapy  Study Documents:                   | Title Acronym: Other Ids: 190143 19-C-0143 | Recruiting | Endocrine/<br>Neuroendo<br>crine     Non-<br>Small Cell<br>Lung<br>Cancer     Breast<br>Cancer     Gastrointe<br>stinal/Geni<br>tourinary<br>Cancers     Ovarian<br>Cancer | <ul> <li>Drug: Fludarabine Days -7 to -3: Fludarabine 25 mg/m2/day IVPB daily over 30 minutes for 5 days.</li> <li>Drug: Cyclophosphamide Days -7 and -6: Cyclophosphamide 60 mg/kg/day x 2 days IV in 250 mL D5W infused simultaneously with mesna 15 mg/kg/day over 1 hour x 2 days.</li> <li>Drug: Aldesleukin Aldesleukin 720,000 IU/kg or 72,000 IU/kg (based on total body weight) IV over 15 minutes approximately every 8 hours beginning within 24 hours of cell infusion and continuing for up to 4 days (maximum 10 doses).</li> <li>Biological: Sleeping Beauty Transposed PBL Day 0: Cells are to be infused at a dose not to exceed 1.5e11 in 400 mL intravenously on the Patient Care Unit over 20-30 minutes or as clinically determined by an investigator for patient safety (between 2-4 days after the last dose of fludarabine).</li> </ul> | 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:  • Phenotypic and functional characteristics of PBL [ Time Frame: 2-4 years post cell infusion ]  Patient PBL will be obtained from whole blood and then evaluated for function and phenotype  • Safety and tolerance [ Time Frame: 6 weeks (+/- 2 weeks) following administration of the cell product ]  Using standard CTCAE 5.0  | Actual Enrollment: Estimated Enrollment: 210  Original Estimated Enrollment: Same as current  Age: 18 Years to 70 Years (Adult, Older Adult)  Sex: All | Study Sponsors:  Same as current  Collaborators: Not Provided | Study Start: September 27, 2022  Primary Completion: December 31, 2028 (Final data collection date for primary outcome measure)  Study Completion: December 31, 2029  First Posted: September 25, 2019  Results First Posted: Last Update Posted: September 22, 2022 |

|    | NCT Number      | Title   | Other Names                                | Status                 | Conditions  | Interventions  | Characteristics   | Population  | Sponsor/Collaborators   | Dates   |
|----|-----------------|---|--|------------------------|---|--|---|---|---|---|
| 26 | NCT0409<br>9797 | C7R-GD2.CAR T Cells for Patients With GD2-expressing Brain Tumors (GAIL-B) Study Documents:           | Title Acronym: Other Ids: H- 45668 GAIL-B  | Recruiting             | Diffuse Intrinsic Pontine Glioma     High Grade Glioma     Embryonal Tumor     Ependyma 1 Tumor | <ul> <li>Genetic: (C7R)-GD2.CART cells <ul> <li>single transduced without C7R with lymphodepletion chemotherapy</li> <li>Dose Level 1: 1 x 10^7 C7R-GD2.CART cells with lymphodepletion chemotherapy</li> <li>Dose Level 2: 3 x 10^7 C7R-GD2.CART cells with lymphodepletion chemotherapy</li> <li>Dose Level 2: 3 x 10^7 C7R-GD2.CART cells with lymphodepletion chemotherapy</li> </ul> </li> <li>Drug: Cyclophosphamide <ul> <li>Patients at all dose levels will receive lymphodepletion chemotherapy. They will receive 2 daily doses of cyclophosphamide (500mg/m2/day) finishing at least 24 hours before T-cell infusion. The drug will be given intravenously (through an IV needle).</li> <li>Other Name: Cytoxan</li> </ul> </li> <li>Drug: Fludarabine <ul> <li>Patients at all dose levels will receive lymphodepletion chemotherapy. They will receive 3 daily doses of fludarabine (30mg/m2/day) finishing at least 24 hours before T-cell infusion. The drug will be given intravenously (through an IV needle).</li> <li>Other Name: Fludara</li> </ul> </li> </ul> | 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: 34  Original Estimated Enrollment: Same as current  Age: 12 Months to 21 Years (Child, Adult)  Sex: All | Study Sponsors:  Same as current  Collaborators: Center for Cell and Gene Therapy, Baylor College of Medicine | Study Start: February 3, 2020  Primary Completion: February 2023 (Final data collection date for primary outcome measure)  Study Completion: February 2038  First Posted: September 23, 2019  Results First Posted: Last Update Posted: September 22, 2022  |
| 27 | NCT0360<br>2612 | T Cells Expressing a Novel Fully- Human Anti- BCMA CAR for Treating Multiple Myeloma Study Documents: | Title Acronym: Other Ids: 180125 18-C-0125 | Active, not recruiting | Myeloma-<br>Multiple     Myeloma,<br>Plasma-<br>Cell  | <ul> <li>Drug: Cyclophosphamide 300 mg/m^2 IV over 30 minutes on days -5, -4, and -3</li> <li>Drug: Fludarabine 30 mg/m^2 IV infusion over 30 minutes administered immediately following the cyclophosphamide on day -5, -4, -3</li> <li>Biological: Anti-BCMA CAR T cells 0.75x10^6 - 12.0X10^6 CAR+ T cells per kg of recipient bodyweight one time dose on day 0</li> </ul>   | Study Type: Interventional  Phase: Phase 1  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: Not Provided | Actual Enrollment: 35  Estimated Enrollment:  Original Estimated Enrollment: 42  Age: 18 Years to 73 Years (Adult, Older Adult)  Sex: All         | Study Sponsors:  Same as current  Collaborators: Not Provided   | Study Start: September 14, 2018  Primary Completion: January 1, 2023 (Final data collection date for primary outcome measure)  Study Completion: January 1, 2024  First Posted: July 27, 2018  Results First Posted: Last Update Posted: September 21, 2022 |