CURE SICKLE CELL.

CURRENT & ONGOING GENETIC THERAPY CLINICAL TRIALS IN SICKLE CELL DISEASE

It's time to rewrite the story of sickle cell.

Clinical trials are medical studies aimed at finding safe and effective ways to prevent, detect, or treat diseases. Some clinical trials are testing ways of treating disease by changing a patient's genetic material. These are called genetic therapies, and most often they work by adding a healthy copy of a defective gene into the patient's cells. In the future, these techniques may allow doctors to treat a disorder by inserting a gene into a patient's cells instead of using drugs or surgery. Researchers are testing other approaches in addition to replacing the mutated gene with a healthy copy. They are inactivating, or "knocking out," a mutated gene that is functioning improperly. They also are introducing a new gene into the body to help fight a disease. Although genetic therapy is a promising new treatment option for many rare diseases, the technique is still being studied to ensure that it is safe and effective.

How can you help rewrite the story of sickle cell disease? Clinical trial participation is one of the most important ways we can move towards a future without sickle cell disease. We encourage adults, as well as children, healthy volunteers, those living with sickle cell disease, and people from diverse ethnic and racial backgrounds to consider participating in clinical trials.

Researchers are currently studying a number of potential new treatment options and also working towards cures. This document includes genetic therapy trials only. For information on other sickle cell disease trials, please visit: https://clinicaltrials.gov.

Have questions and want to learn more about clinical trials? Go to www.nih.gov/health-information/nih-clinical-research-trials-you.





INTERVENTIONAL

► Gene Transfer for Patients With Sickle Cell Disease: Phase 1/2

ACTIVE / NOT RECRUITING	
Age Range	18-45 Years
Trial Time Frame	7/2014-7/2038
Ref. No.	NCT02186418

This study will assess the safety and efficacy of gene transfer using a gammaglobin lentiviral vector. Gene transfer will occur ex-vivo into human bone marrow or mobilized peripheral blood CD34+ hematopoietic stem cells of subjects with sickle cell disease.

https://ClinicalTrials.gov/show/NCTo2186418

Pilot and Feasibility Study of Hematopoietic Stem Cell Gene Transfer for Sickle Cell Disease: Phase 1 *

Age Range	3-40 Years
Trial Time Frame	2/2018-10/2024
Ref. No.	NCT03282656

ACTIVE / NOT RECRUITING

DECDUITING

This is an open-label, non-randomized, single center, pilot and feasibility, single-arm cohort study involving a single infusion of autologous bone marrow derived CD₃4+ HSC cells transduced with the lentiviral vector containing a short-hairpin RNA targeting BCL₁₁a.

https://ClinicalTrials.gov/show/NCTo3282656

Clinical Research Study of Autologous Stem Cell Transplantation for Sickle Cell Disease: Phase 1/2

RECRUITING	
Age Range	18 Years and Older
Trial Time Frame	12/2014-12/2025
Ref. No.	NCT02247843

This Phase I clinical trial will assess the safety and efficacy of an autologous transplant of lentiviral vector modified peripheral blood for adults with severe sickle cell disease.

https://ClinicalTrials.gov/show/NCTo2247843

▶ Safety of Blood Stem Cell Mobilization With Plerixafor in Patients With Sickle Cell Disease: Phase 1

RECRUITING		This study will investigate whether up to two injections of plerixafor represent
Age Range	18-40 Years	a safe and effective strategy for mobilizing adequate numbers of CD34+
Trial Time Frame	9/2018-12/2023	hematopoietic stem progenitor cells (HSPC) for autologous hematopoietic cell
Ref. No.	NCT03664830	transplantation (HCT).
		https://ClinicalTrials.gov/show/NCTo3664830

▶ Safety Trial of Escalation of Plerixafor for Mobilization of Cells and Evaluation of Gene Transfer: Phase 1

RECRUITING		This study will look at the safety and efficacy of a drug called Plerixafor.
Age Range	18-65 Years	Plerixafor is approved by the U.S. Food and Drug Administration (FDA) for use
Trial Time Frame	9/2014-7/2025	in increasing blood stem cell counts before collection in cancer patients.
Ref. No.	NCT02193191	https://ClinicalTrials.gov/show/NCT02193191

► Evaluation of the Safety and Efficacy of the LentiGlobin BB305 Drug Product in Severe Sickle Cell Disease: Phase 1/2

ACTIVE / NOT RECRUIT	ING	This is a non-randomized, open-label, multi-site, single-dose study in
Age Range Trial Time Frame	12-50 Years 8/2014-7/2024	approximately 50 adults and adolescents with severe SCD. The study will evaluate hematopoietic stem cell (HSC) transplantation (HSCT) using
Ref. No.	NCT02140554	LentiGlobin BB305 Drug Product.
		https://ClinicalTrials.gov/show/NCT02140554

▶ A Safety and Efficacy Study Evaluating CTX001 in Subjects With Severe Sickle Cell Disease: Phase 1/2/3

ACTIVE / NOT RECRUIT	TING	This is a single-arm, open-label, multi-site, single-dose study to evaluate
Age Range	12-35 Years	the safety and efficacy of autologous CRISPR-Cas9 Modified CD34+ Human
Trial Time Frame	11/2018-10/2024	Hematopoietic Stem and Progenitor Cells (hHSPCs) using CTX001.
Ref. No.	NCT03745287	https://ClinicalTrials.gov/show/NCTo3745287

^{*} Funded by the Cure Sickle Cell Initiative

INTERVENTIONAL CONTINUED

▶ A Study Evaluating Gene Therapy With BB305 Lentiviral Vector in Sickle Cell Disease: Phase 3

RECRUITING Age Range Trial Time Frame Ref. No.	2–50 Years 2/2020–4/2027 NCT04293185	This is a non-randomized, open-label, multi-site, single-dose study in approximately 35 adults and pediatric subjects with sickle cell disease. The study will evaluate hematopoietic stem cell (HSC) transplantation (HSCT) with LentiGlobin BB305 Drug Product for SCD.
		https://clinicaltrials.gov/ct2/show/NCTo4293185

▶ Study of Safety and Efficacy of Genome-edited Hematopoietic Stem and Progenitor Cells in Sickle Cell Disease: Phase 1/2

RECRUITING Age Range Trial Time Frame Ref. No.	2–40 Years 8/2020–9/2026 NCT04443907	This study will evaluate two genome-edited, autologous, hematopoietic stem and progenitor cell (HSPC) products - OTQ923 and HIX763 - each reducing the biologic activity of BCL11A, increasing fetal hemoglobin (HbF) and reducing complications of sickle cell disease.
		https://clinicaltrials.gov/ct2/show/NCT04443907

► Study to Evaluate the Safety and Efficacy of EDIT-301 for Autologous HSCT in Subjects With Severe Sickle Cell Disease: *Phase* 1/2

RECRUITING		The purpose of this study is to evaluate the efficacy, safety and tolerability of
Age Range	18-50 Years	treatment with EDIT-301 in adult subjects with severe sickle cell disease (SCD).
Trial Time Frame	5/2021-8/2025	
Ref. No.	NCT04853576	https://clinicaltrials.gov/ct2/show/NCTo4853576

► Transplantation of CRISPRCas9 Corrected Hematopoietic Stem Cells (CRISPR_SCD001) in Patients With Severe Sickle Cell Disease: *Phase 1/2**

NOT YET RECRUITING		This is an open label, non-randomized, 2-center, phase 1/2 trial of a
Age Range Trial Time Frame Ref. No.	12–35 Years 6/2024–6/2028 NCT04774536	single infusion of sickle allele modified cluster of differentiation (CD34+) hematopoietic stem progenitor cells (HSPCs) in subjects with in subjects ≥12 years old to 35 years old severe Sickle Cell Disease (SCD). The study will evaluate the hematopoietic stem cell transplantation (HSCT) using CRISPR/ Cas9 edited red blood cells (known as CRISPR_SCD001 Drug Product).
		https://clinicaltrials.gov/ct2/show/NCT04774536

▶ Evaluation of Safety and Efficacy of CTX001 in Pediatric Participants With Severe Sickle Cell Disease: Phase 3

RECRUITING Age Range Trial Time Frame Ref. No.	2-11 Years 5/2022–5/2026 NCT05329649	This is a single-dose, open-label study in pediatric participants with severe SCD and hydroxyurea (HU) failure or intolerance. The study will evaluate the safety and efficacy of autologous CRISPR-Cas9 modified CD34+ human hematopoietic stem and progenitor cells (hHSPCs) (CTX001).
		https://clinicaltrials.gov/ct2/show/NCTo5329649

▶ A Gene Transfer Study Inducing Fetal Hemoglobin in Sickle Cell Disease (GRASP, BMT CTN 2001): Phase 2 *

RECRUITING Age Range Trial Time Frame Ref. No.	13-40 Years 7/2022-5/2026 NCT05353647	This is an open-label, non-randomized, multi-center, phase 2 study involving a single infusion of autologous bone marrow derived CD34+ HSC cells transduced with the lentiviral vector containing a short-hairpin RNA targeting BCL11a. 25 patients ages 13 to 40 will be enrolled at sites across the US. The main goal of this study is to determine whether the treatment will lead to a complete absence of severe vaso-occlusive events (VOEs) in patients with severe SCD.
		https://clinicaltrials.gov/ct2/show/NCTo5353647

INTERVENTIONAL CONTINUED

▶ Haploidentical Hematopoietic Stem Cell Transplantation (HSCT) for Patients With Severe Sickle Cell Disease

RECRUITING		The purpose of this study is to develop a safe and curative stem cell transplant
Age Range	2-25 Years	approach to treating sickle cell disease by assessing the safety of haploidentical
Trial Time Frame	04/2020-11/2027	hematopoietic stem cell transplantation using a $oldsymbol{eta}_+$ T-cell depletion for children
Ref. No.	NCT04207320	and adolescents with severe sickle cell disease (SCD).

https://clinicaltrials.gov/ct2/show/NCT04207320

► A Study Evaluating the Safety and Efficacy of BEAM-101 in Patients With Severe Sickle Cell Disease (BEACON): Phase 1/2

RECRUITING		This is an open-label, single-arm, multicenter, Phase 1/2 study evaluating the
Age Range	18-35 Years	safety and efficacy of the administration of autologous base edited CD34+
Trial Time Frame	8/2022-2/2027	HSPCs (BEAM-101) in patients with severe SCD.
Ref. No.	NCT05456880	https://clinicaltrials.gov/ct2/show/NCTo5456880

► Evaluation of Efficacy and Safety of a Single Dose of CTX001 in Participants With Transfusion-Dependent β-Thalassemia and Severe Sickle Cell Disease: *Phase* 3

RECRUITING		This is a single-dose, open-label study in participants with transfusion-
Age Range Trial Time Frame	12–25 Years 8/2022–2/2025	dependent β-thalassemia (TDT) or severe SCD. The study will evaluate the safety and efficacy of autologous CRISPR-Cas9 modified CD34+ human
Ref. No.	NCT05477563	hematopoietic stem and progenitor cells (hHSPCs) using CTX001.
		https://clinicaltrials.gov/ct2/show/NCTo5477563

▶ Non-myeloablative Haploidentical HCT Study for Patients With Sickle Cell Disease, Including Compromised Organ Function: *Phase* 1/2

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NOT YET RECRUITING		Haploidentical hematopoietic cell transplantation offers a widely available
Age Range	4-100 Years	curative option for individuals with sickle cell disease. The goal is to reverse
Trial Time Frame	, ,	SCD while avoiding unacceptable graft rejection, graft-versus-host disease,
Ref. No.		infectious complications, and hyperinflammatory responses. We hypothesize
		that a moderate amount of immunosuppression will maximize efficacy while
		avoiding unacceptable toxicity.
		https://clinicaltrials.gov/ct2/show/NCTo6145282
		nctps://chincarchais.gov/ct2/show/ivc100145282

Long-term Follow-up of Subjects With Transfusion-Dependent β-Thalassemia (TDT) Treated With Ex Vivo Gene Therapy

RECRUITING BY INVITATION ONLY

Age Range Trial Time Frame Ref. No.

Up to 50 Years 1/2014-11/2035 NCT02633943 This is a multi-center, long-term safety and efficacy follow-up study for subjects with transfusion-dependent β-thalassemia (TDT) who have been treated with ex vivo gene therapy drug product in bluebird bio-sponsored clinical studies.

https://ClinicalTrials.gov/show/NCTo2633943

▶ Long-term Follow-up of Subjects With Sickle Cell Disease Treated With Ex Vivo Gene Therapy

RECRUITING BY INVITATION ONLY

Age Range Trial Time Frame Ref. No.

2-53 Years 10/2020-5/2038 NCT04628585

This is a multi-center, long-term safety and efficacy follow-up study for subjects who have been treated with ex vivo gene therapy drug product in bluebird bio-sponsored clinical studies. After completing the parent clinical study (approximately 2 years), eligible subjects will be followed for an additional 13 years for a total of 15 years post-drug product infusion.

https://ClinicalTrials.gov/show/NCTo4628585

▶ Long-term Follow-up Study in Subjects Who Received CTX001

RECRUITING BY INVITATION ONLY

Age Range 18 Years and Older **Trial Time Frame** 1/2021-9/2039

Ref. No. NCT04208529 This is a multi-site, observational study to evaluate the long-term safety and efficacy of CTX001 in subjects who received CTX001 in Study CTX001-111 (NCT03655678) or Study CTX001-121 (NCT03745287).

https://ClinicalTrials.gov/show/NCTo4208529

► Cooperative Assessment of Late Effects for SCD Curative Therapies (COALESCE)

RECRUITING

Age Range 4-65 Years **Trial Time Frame** 7/2022-2/2026 Ref. No. NCT05153967 The primary goal of this study is to determine whether curative therapies for individuals with SCD will result in improved or worsening heart, lung, and kidney damage when compared to individuals with SCD receiving standard therapy. The investigators will also explore whether certain genes are associated with a good or bad outcome after curative therapy for SCD.

https://ClinicalTrials.gov/show/NCTo5153967

▶ Discarded Bone Marrow for Hematology Research

RECRUITING

Age Range Child, Adult, Older Adult **Trial Time Frame** 7/2022-1/2035 Ref. No. NCT04671212 The primary objective of this study is to establish a mechanism to obtain discarded bone marrow-containing bone samples from hemoglobinopathy, as well as non-hemoglobinopathy individuals. The processing of samples will help to understand how best to manipulate HSPC's from hemoglobinopathy patients with gene therapy and gene technologies in the laboratory environment. It will also allow us to establish a reservoir of samples that can be studied in the future to assess cellular function and fitness for transplant.

https://ClinicalTrials.gov/show/NCTo4671212