



The 65th ASH Annual Meeting Abstracts

ORAL ABSTRACTS

801.GENE THERAPIES

Exagamglogene Autotemcel for Severe Sickle Cell Disease

Haydar Frangoul, MD¹, Franco Locatelli, MD PhD², Akshay Sharma, MBBS³, Monica Bhatia, MD⁴, Markus Mapara, MD⁵, Lyndsay Molinari, MD⁶, Donna Wall, MD⁷, Robert I. Liem, MD⁸, Paul Telfer, MD⁹, Ami J. Shah, MD¹⁰, Marina Cavazzana, MD PhD¹¹, Selim Corbacioglu, MD PhD¹², Damiano Rondelli, MD¹³, Roland Meisel, MD¹⁴, Laurence Dedeken, MD¹⁵, Stephan Lobitz, MSc¹⁶, Mariane de Montalembert, MD PhD¹⁷, Martin H. Steinberg, MD¹⁸, Mark C. Walters, MD¹⁹, Laura Bower, MD²⁰, Suzan Imren, MD²⁰, Christopher Simard, MD²⁰, Fengjuan Xuan, PhD²⁰, Weiyu Zhou, PhD²⁰, Phuong Khanh Morrow, MD²¹, William Hobbs, MD PhD²⁰, Stephan Grupp, MD PhD²²

¹ Sarah Cannon Research Institute at The Children's Hospital at TriStar Centennial, Nashville, TN

² IRCCS, Ospedale Pediatrico Bambino Gesù Rome, Catholic University of the Sacred Heart, Rome, Italy

³ St. Jude Children's Research Hospital, Memphis, TN

⁴ Department of Pediatrics, Columbia University Irving Medical Center, New York - Presbyterian-Morgan Stanley Children's Hospital, New York, NY

⁵ Division of Hematology and Oncology, Columbia University, New York, NY

⁶ Sarah Cannon Pediatric Transplant and Cellular Therapy Program at Methodist Children's Hospital, San Antonio, TX

⁷ SickKids, Toronto, Canada

⁸ Ann & Robert H. Lurie Children's Hospital of Chicago, Chicago, IL

⁹ Royal London Hospital, Barts Health NHS Trust, London, United Kingdom

¹⁰ Center for Definitive and Curative Medicine, Stanford University, Palo Alto, CA

¹¹ Necker-Enfants Malades Hospital, Assistance Publique-Hôpitaux de Paris (AP-HP), University of Paris, Paris, France

¹² University of Regensburg, Regensburg, Germany

¹³ Division of Hematology/Oncology, University of Illinois at Chicago, Chicago, IL

¹⁴ Division of Pediatric Stem Cell Therapy, Department of Pediatric Oncology, Hematology and Clinical Immunology, Medical Faculty, Heinrich-Heine-University, Duesseldorf, Germany

¹⁵ Hôpital Universitaire des Enfants Reine Fabiola, Brussels, Belgium

¹⁶ Gemeinschaftsklinikum Mittelrhein, Koblenz, Germany

¹⁷ Necker-Enfants Malades Hospital, Assistance Publique-Hôpitaux de Paris (AP-HP), University of Paris-Cité, Paris, France

¹⁸ Boston University Chobanian & Avedisian School of Medicine, Boston, MA

¹⁹ UCSF Benioff Children's Hospital, Oakland, CA

²⁰ Vertex Pharmaceuticals, Boston, MA

²¹ CRISPR Therapeutics, Cambridge, MA

²² Children's Hospital of Philadelphia, Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA

Background: Exagamglogene autotemcel (exa-cel) is a non-viral cell therapy designed to reactivate fetal hemoglobin via ex vivo CRISPR-Cas9 gene-editing of autologous CD34+ hematopoietic stem and progenitor cells (HSPCs) at the erythroid-specific enhancer region of the BCL11A gene in patients (pts) with severe sickle cell disease (SCD). We report that in a pre-specified interim analysis, the pivotal CLIMB SCD-121 trial of exa-cel met primary and key secondary endpoints.

Methods: CLIMB SCD-121 is an ongoing, 24-mo, phase 3 trial of exa-cel in pts age 12-35y with SCD and a history of ≥ 2 VOCs/y in 2y prior to screening. Primary efficacy endpoint is proportion of pts free of severe VOCs for ≥ 12 consecutive months (mos) (VF12); key secondary efficacy endpoints are proportion of pts free from inpatient hospitalization for severe VOCs for ≥ 12 consecutive mos (HF12) and proportion of pts free from severe VOCs for ≥ 9 consecutive mos (VF9). Evaluable pts for VF12 and HF12 had ≥ 16 mos follow-up after exa-cel infusion; pts evaluable for VF9 had ≥ 12 mos follow-up after infusion.

Evaluation of primary and key secondary endpoints began 60 days after last RBC transfusion for post-transplant support or SCD management. Pts completing trial enrolled in long-term follow-up Study 131. Mean (SD) are shown except where noted. **Results:** As of 10 Feb 2023, 42 pts with SCD (age 21.2[range 12-34]y; 12[28.6%] age ≥ 12 to < 18 y; 4.2 VOCs/y at baseline) received exa-cel. Following infusion, all pts engrafted neutrophils and platelets (median 27 and 34.5 days, respectively). 19/20

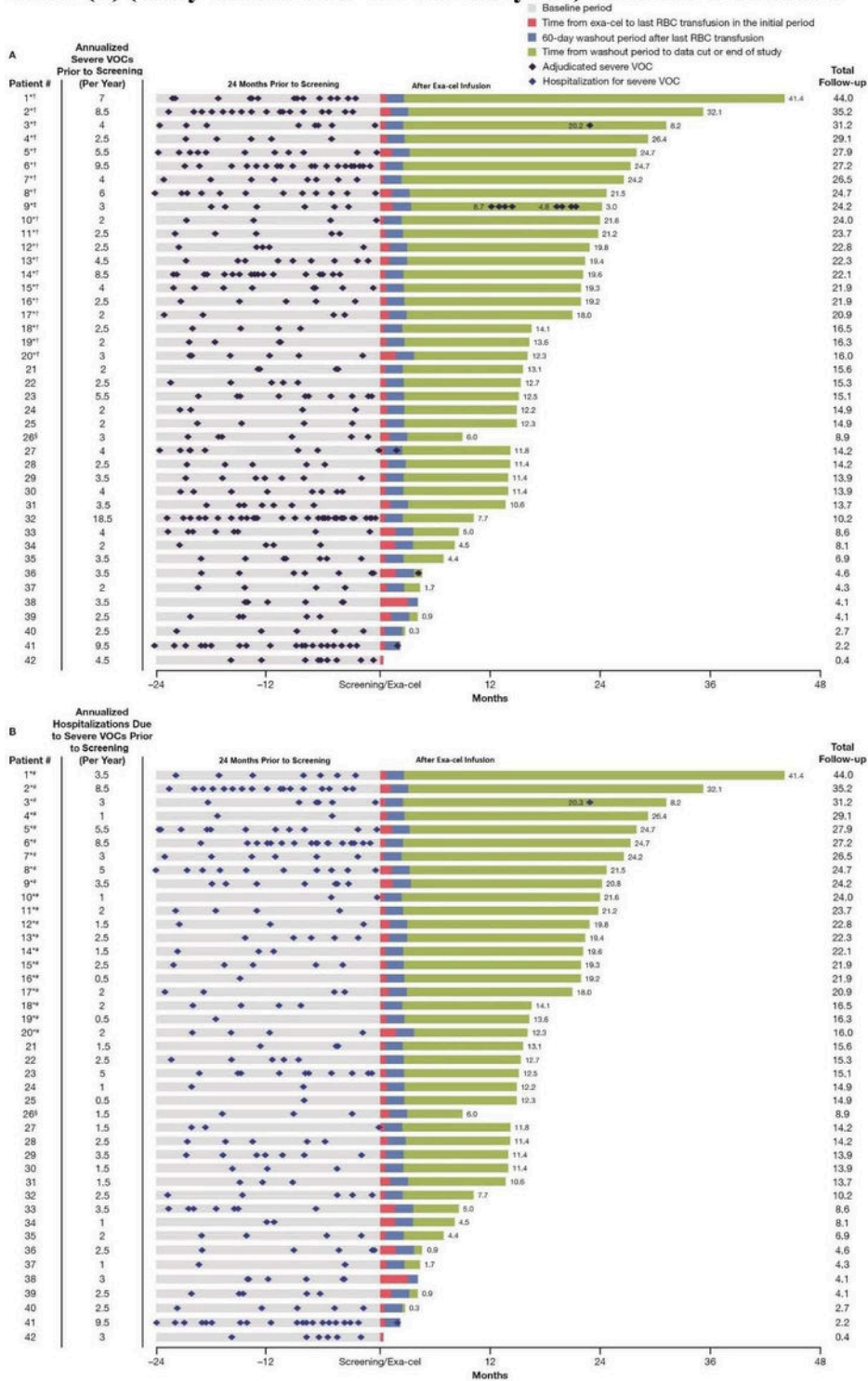
(95.0%) pts evaluable for primary endpoint were free of VOCs for ≥ 12 consecutive mos (VF12; 95% CI, 75.1% to 99.9%; $P < 0.0001$), 20/20 (100%) were free from hospitalizations for VOCs for ≥ 12 consecutive mos (HF12; 95% CI, 83.2 to 100.0; $P < 0.0001$), and 29/30 (96.7%) were free of VOCs for ≥ 9 consecutive mos (VF9; 95% CI, 82.8 to 99.9; $P < 0.0001$). In pts achieving VF12, VOC free duration was 21.8 (range 12.3-41.4) mos; 18 pts remained VOC free through follow-up and 1 pt had an adjudicated VOC in the setting of parvovirus infection ~22.8 mos after exa-cel; pt recovered fully and has since been VOC free (Fig). For all pts, total Hb was 12.1 g/dL at Month 3 and was maintained at ≥ 11.0 g/dL from Month 6 onward; HbF was 36.0% at Month 3 and was generally maintained at $\geq 40.0\%$ from Month 6 onward with pancellular distribution ($\geq 95\%$ RBCs expressing HbF). Proportion of edited *BCL11A* alleles was stable over time in bone marrow CD34⁺ and peripheral blood nucleated cells. 36/39 pts with ≥ 60 days follow-up after last RBC transfusion (including those not yet evaluable) remained VOC free (up to 41.4 mos; Fig). Quality-of-life (QOL) measures showed clinically significant improvements from baseline.

All pts had ≥ 1 adverse event (AE), most were Grade 1 or 2; 40 (95.2%) pts had AEs of Grade 3 or 4 severity. Most common AEs were nausea (66.7%), stomatitis (61.9%), febrile neutropenia (52.4%), headache (52.4%), and vomiting (52.4%). Most AEs and serious AEs (SAEs) occurred within first 6 mos after infusion. No pts had SAEs considered related to exa-cel. As previously reported, 1 pt died from respiratory failure due to COVID-19 unrelated to exa-cel. There were no study discontinuations or malignancies.

Conclusions: The CLIMB SCD-121 trial met primary and key secondary endpoints, with exa-cel treatment resulting in early and sustained increases in Hb and HbF leading to elimination of VOCs in 95% of pts, elimination of inpatient hospitalization for VOCs in 100% of pts and improved QOL. Safety profile of exa-cel was generally consistent with myeloablative busulfan conditioning and autologous transplantation. These results show exa-cel has the potential to deliver a one-time functional cure to pts with severe SCD.

Disclosures Frangoul: Editas Medicine: Consultancy; Rocket Pharmaceuticals: Consultancy, Other: Member of DSMB for a study; Jazz Pharmaceuticals: Speakers Bureau; Vertex Pharmaceuticals: Consultancy, Membership on an entity's Board of Directors or advisory committees. **Sharma:** Editas Medicine: Consultancy; RCI BMT/NMDP: Honoraria, Other: Clinical Trial Medical Monitor; Medexus Inc: Consultancy; Vertex Pharmaceuticals: Consultancy, Other: Clinical Trial Site PI; Sangamo Therapeutics: Consultancy; CRISPR Therapeutics: Other: Clinical Trial Site PI, Research Funding. **Mapara:** Crispr/vertex: Consultancy; Incyte: Consultancy; Bluebird bio: Consultancy. **Liem:** Bluebird Bio: Research Funding; NIH/NHLBI: Research Funding; NIH/NCATS: Research Funding; Vertex: Research Funding; Editas: Research Funding; Global Blood Therapeutics: Research Funding. **Telfer:** Apopharma: Other: Clinical trial activity, Speakers Bureau; Terumo: Speakers Bureau; Pfizer: Other: Advisory board; Clinical trial activity; Data monitoring committee; ; Napp Pharmaceuticals: Other: Clinical trial activity; Celgene: Other: Clinical trial activity; Kyowa Kirin Limited: Other: Investigator-led funding; Novartis: Other: Advisory board; Clinical trial activity; ; bluebird bio: Other: Advisory board; Investigator-led funding; **Shah:** Vertex: Membership on an entity's Board of Directors or advisory committees. **Rondelli:** Vertex: Other: Steering Committee. **Meisel:** Miltenyi Biotech: Research Funding; medac: Consultancy, Research Funding, Speakers Bureau; Gilead/KITE: Research Funding; Novartis: Consultancy, Research Funding, Speakers Bureau; CELGENE BMS: Consultancy, Research Funding, Speakers Bureau; Bluebird Bio: Consultancy, Speakers Bureau; CRISPR Therapeutics: Consultancy, Research Funding, Speakers Bureau; Vertex: Consultancy, Research Funding, Speakers Bureau. **Lobitz:** AddMedica: Membership on an entity's Board of Directors or advisory committees; Novartis: Membership on an entity's Board of Directors or advisory committees, Other: Patient Booklet and educational papers; Global Blood Therapeutics: Membership on an entity's Board of Directors or advisory committees; Agios: Membership on an entity's Board of Directors or advisory committees; BlueBird Bio: Membership on an entity's Board of Directors or advisory committees, Other: Lecture; Vertex Pharmaceuticals: Membership on an entity's Board of Directors or advisory committees, Other: Steering Committee; German National Disease Management Program: Other: Spokesperson for children and adolescents with sickle cell disease; German National Treatment Guideline: Other: Corresponding author of the treatment guideline for children and adolescents with sickle cell disease. **de Montalembert:** Addmedica: Consultancy, Membership on an entity's Board of Directors or advisory committees, Other: ASH meeting 2022 support; Novartis: Membership on an entity's Board of Directors or advisory committees. **Steinberg:** NIH: Research Funding; Fulcrum Therapeutics: Consultancy; ASH: Other: Lecture at HEM ASH Whiteboard Symposium, Speakers Bureau; Imara Therapeutics: Membership on an entity's Board of Directors or advisory committees. **Walters:** Ensoma, Inc: Consultancy; Vertex Pharmaceuticals: Consultancy; BioChip Labs: Consultancy, Other: Medical Director; AllCells, Inc: Consultancy, Other: Medical Director. **Bower:** Vertex Pharmaceuticals: Current Employment. **Imren:** Vertex Pharmaceuticals: Current Employment, Current holder of stock options in a privately-held company. **Simard:** Vertex Pharmaceuticals: Current Employment. **Xuan:** Vertex Pharmaceuticals: Current Employment. **Zhou:** Vertex Pharmaceuticals: Current Employment. **Morrow:** Vertex Pharmaceuticals: Membership on an entity's Board of Directors or advisory committees; CRISPR Therapeutics: Current Employment, Membership on an entity's Board of Directors or advisory committees. **Hobbs:** Vertex Pharmaceuticals: Current Employment. **Grupp:** Cabaletta: Consultancy, Membership on an entity's Board of Directors or advisory committees; Allogene: Consultancy, Membership on an entity's Board of Directors or advisory committees; Juno: Consultancy, Membership on an entity's Board of Directors or advisory committees; Collectis: Consultancy, Membership on an entity's Board of Directors or advisory committees; Adaptimmune: Consultancy, Membership on an entity's Board of Directors or advisory committees; CBMG: Consultancy, Membership on an entity's Board of Directors or advisory committees; Servier: Research Funding; Kite: Research Funding; Jazz: Consultancy, Membership on an entity's Board of Directors or advisory committees, Research Funding; Vertex: Consultancy, Research Funding; Novartis: Consultancy, Research Funding.

Figure. Duration of Period Free From Vaso-Occlusive Crises (A) and Hospitalizations for Vaso-Occlusive Crises (B) (Study CLIMB SCD-121 and Study 131) After Exa-cel Infusion



All VOCs were adjudicated by the Independent Endpoint Adjudication Committee. *Participant was evaluable for the primary endpoint (VF12) and first key secondary endpoint (HF12); †Participant achieved VF12; ‡Participant did not achieve VF12; §Death from respiratory failure due to COVID-19 infection; #Participant achieved HF12

Figure 1

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