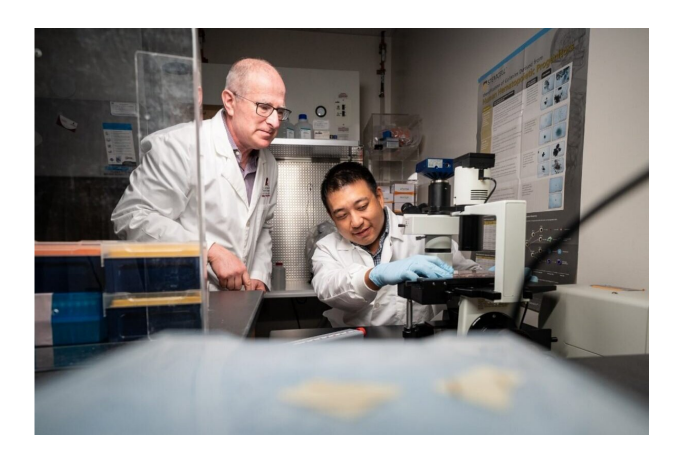


New insight into red blood cell machinery offers clues to treating sickle cell disease

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Corresponding author Mitchell Weiss, M.D., Ph.D., chair of the St. Jude Department of Hematology (left), helped uncover how the body generates protective fetal hemoglobin against sickle cell disease and beta thalassemia. Credit: St. Jude Children's Research Hospital

Hematologists at St. Jude Children's Research Hospital have discovered



key molecular details of how genetic variants in blood-forming machinery enable some people to retain the ability to generate red blood cells with a form of hemoglobin that is normally expressed only before birth. Persistence of fetal hemoglobin expression after birth can protect patients from the deleterious effects of mutations that cause beta thalassemia and sickle cell disease. The findings are already aiding development of gene therapies. The work was published online today in *Nature Genetics*.

Fetuses have a different form of hemoglobin that binds oxygen more strongly, enabling them to thrive in the low-oxygen environment of the womb. Researchers are developing a gene therapy whereby a patient's own blood-forming cells, called hematopoietic stem cells, or HSCs, are genetically reprogrammed to favor production of protective fetal globin after birth, rather than transitioning to the aberrant adult form.

"More than 50 years ago, researchers had observed that people with beta thalassemia or sickle cell <u>disease</u> mutations who also inherited a genetic variant that gave them persistently high levels of fetal hemoglobin didn't get as sick," said corresponding author Mitchell Weiss, M.D., Ph.D., chair of the St. Jude Department of Hematology. "So, it has been a holy grail to figure out how to use drugs or genetic manipulation to enhance fetal hemoglobin therapeutically in these common, serious diseases.

"Our current study builds upon an extraordinary amount of prior research to offer important, therapeutically relevant insights into the mechanisms by which these variants repress the transition from fetal to adult hemoglobin," Weiss added.

Mapping the genetic machinery of fetal hemoglobin

The researchers mapped the genetic machinery controlling how six fetal hemoglobin-persistent variants inhibit the fetal-to-adult hemoglobin



switch. These variants either destroy regulatory DNA that represses fetal hemoglobin or create new regulatory sequences that activate it.

"Genes are controlled by a balance between activation and repression," said first author Phillip Doerfler, Ph.D., of St. Jude Hematology. "Our work shows how the unique interplay between activators and repressors ultimately controls the expression of fetal hemoglobin."

The work identified two distinct <u>activator</u> sequences, each paired with a nearby <u>repressor</u> sequence. Fetal hemoglobin expression was turned on by using genome editing to disrupt a repressor sequence and turned off by disruption of an activator.

A gene therapy treatment to turn on fetal hemoglobin

The close proximity of activator and repressor sequences has important therapeutic implications for treating beta thalassemia and sickle cell disease. The researchers are already developing clinical trials of a geneediting treatment that disrupts the repressor binding, yet still enables the fetal hemoglobin gene to be turned on. This genome editing approach must be precise, so as to not disrupt the adjacent activator sequence.

"Now we understand that because these activator and repressor sites are so close to each other, when we perform gene editing to manipulate them, we have to be careful that by trying to interfere with the repressor we don't block the activator," Weiss said.

The researchers' objective is to take hematopoietic stem cells from sickle cell disease or beta thalassemia patients, edit the cells to disrupt the repressor binding site and return them to the patients. The red blood cells then made by those stem <u>cells</u> will be tipped toward expressing more fetal hemoglobin, which will alleviate the disease.



The same genetic strategy of enhancing fetal hemoglobin can treat both sickle cell disease and beta thalassemia. Although the two diseases are different genetically, both involve mutations of the gene for adult hemoglobin that disrupt its function or expression.

More information: Phillip A. Doerfler et al, Activation of γ-globin gene expression by GATA1 and NF-Y in hereditary persistence of fetal hemoglobin, *Nature Genetics* (2021). DOI: 10.1038/s41588-021-00904-0

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