

With gene editing, researchers cure blood disorder in fetal mice

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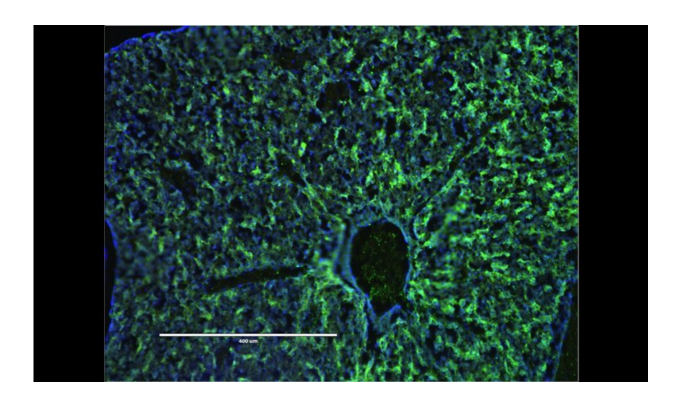


Image of nanoparticles accumulating in the liver of a fetal mouse. Credit: Yale University

With the combined efforts of three Yale laboratories, researchers conducted the first demonstration of site-specific gene editing in a fetus, correcting a mutation that causes a severe form of anemia.



The technique, described in a paper published June 26 in *Nature Communications*, involves an <u>intravenous injection</u> of <u>nanoparticles</u> carrying a combination of donor DNA and synthetic molecules known as <u>peptide nucleic acids</u> (PNAs). The PNAs, which mimic DNA, bind to the target gene and form a triple helix—an aberration that triggers the cells' repair mechanisms. As part of this process, the healthy donor DNA—paired with the PNA in a nanoparticle—is used to fix the mutation.

For the study, this gene-editing package was injected into the fetuses of mice. At four months after birth, the mice had been cured of thalassemia, an inherited defect in oxygen-carrying <u>red blood cells</u>.

"The treated mice had normal blood counts, their spleens returned to normal size, and they lived a normal life span—whereas, the untreated ones died much earlier," said Peter M. Glazer, M.D., professor of therapeutic radiology and of genetics, who developed the technique of combining the PNA and DNA to repair gene mutations. "So we have a long-term survival benefit, which is pretty dramatic."

Using nanoparticles for delivery is crucial, said the researchers. When injected intravenously without nanoparticles, PNAs are cleared from circulation within about 30 minutes. When delivered by a nanoparticle and taken up by a cell, though, they remain much longer. Mark Saltzman, Yale's Goizueta Foundation Professor of Biomedical Engineering, Chemical and Environmental Engineering, and professor of physiology, made the nanoparticles with a degradable polymer and designed them small enough—200 to 300 nanometers—to readily accumulate in the liver of the fetus, where the stem cells are located before migrating to the bone marrow.

In a previous study, the researchers yielded promising results when they conducted a similar procedure on <u>adult mice</u>. By editing the gene in the



fetus for the new study, though, the researchers greatly minimized the amount of damage caused by the disease.

More information: Adele S. Ricciardi et al. In utero nanoparticle delivery for site-specific genome editing, *Nature Communications* (2018). DOI: 10.1038/s41467-018-04894-2

Provided by Yale University

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