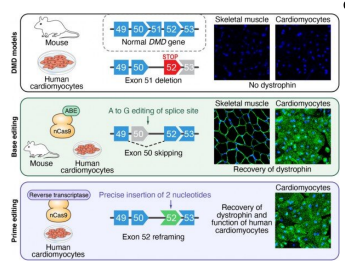


New gene editing strategies developed for Duchenne muscular dystrophy

3 May 2021



In a mouse model and in human heart muscle cells, researchers used gene editing to modify specific DNA sequences and restore dystrophin production in mutant dystrophin genes. Credit: UT Southwestern Medical Center

UT Southwestern scientists successfully employed a new type of gene therapy to treat mice with Duchenne muscular dystrophy (DMD), uniquely utilizing CRISPR-Cas9-based tools to restore a large section of the dystrophin protein that is missing in many DMD patients. The approach, described online today in the journal *Science Advances*, could lead to a treatment for DMD and inform the treatment of other inherited diseases.

"Thousands of different mutations causing Duchenne have been identified, but they tend to cluster into certain parts of the dystrophin gene," says study leader Eric Olson, Ph.D., professor and founding chair of molecular biology at UTSW. Some of these mutations cause <u>muscle cells</u> to produce short, less functional versions of the dystrophin protein. "The power of our method is that you don't need a <u>new gene</u> editing strategy for

every patient with a new mutation; you can correct multiple different mutations with a consolidated approach."

Olson and his colleagues took advantage of the fact that the enormous dystrophin gene is composed of many different segments, called exons, some of which are dispensable. In about 8 percent of boys with DMD, nearly half of the dystrophin protein is missing due to mutations within exon 51 that cause the body to stop protein production. The researchers developed multiple successful CRISPR-Cas9 nucleotide gene editing strategies to skip the errant "stop" signal, restoring production of 97 percent of the protein. Some strategies worked by removing neighboring exons, while others utilized tiny genetic additions or subtractions to get protein production back on track.

When the researchers used the new approach in mice with dystrophin mutations, functioning copies of dystrophin returned to more than half of all leg muscle fibers within three weeks. Moreover, the group showed that they could use isolated <u>cells</u> from mice or humans with DMD to test whether the approach would be successful for a particular patient ahead of treatment. The isolated cells are coaxed to develop into induced <u>pluripotent stem</u> <u>cells</u> (iPS cells) and then heart cells. In a dish, researchers can observe whether the gene editing program helps the heart cells work better.

"Using iPS cell-derived cardiomyocytes from DMD patients, we rapidly tested our nucleotide gene editing approaches, demonstrating the recovery of the dystrophin protein," says Francesco Chemello, Ph.D., a postdoctoral researcher in the Olson lab and first author of the paper.

DMD affects about one in five thousand males at birth and leads to progressively worsening muscle weakness in early childhood. The disease is caused by one of more than 7,000 different



<u>mutations</u> in the gene for dystrophin—a protein that normally acts as a scaffold to support muscle fibers. Without fully functional <u>dystrophin</u>, the skeletal and heart muscles of people with DMD degenerate over time, eventually leading to death.

The gene therapy reported in the study is not ready for humans with DMD yet. Additional safety studies in animals are needed first, as well as more work to optimize the virus that carries the gene editing machinery to muscle and heart cells. But by showing that multiple CRISPR-Cas9 strategies can correct a mutation, the researchers have expanded the toolbox of potential <u>gene therapy</u> options for DMD.

"Every cell in the human body has 3 billion letters of DNA sequence in its genome, and this method makes it possible to correct large deletions in the DMD gene by specifically swapping one of these letters," says Olson. "That level of specificity and efficiency is remarkable."

More information: F. Chemello et al. Precise correction of Duchenne muscular dystrophy exon deletion mutations by base and prime editing, *Science Advances* (2021). <u>DOI:</u> <u>10.1126/sciadv.abg4910</u>

Provided by UT Southwestern Medical Center

APA citation: New gene editing strategies developed for Duchenne muscular dystrophy (2021, May 3) retrieved 2 July 2022 from <u>https://medicalxpress.com/news/2021-05-gene-strategies-duchenne-muscular-dystrophy.html</u>

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