

Targeted gene therapy enhances treatment for Pompe disease

June 25 2012

Gene therapy to replace the protein missing in Pompe disease can be effective if the patient's immune system does not react against the therapy. Targeted delivery of the gene to the liver, instead of throughout the body, suppresses the immune response, improving the therapeutic effect, according to an article published in Human Gene Therapy, a peerreviewed journal from Mary Ann Liebert, Inc. The article is available free online at the Human Gene Therapy website.

"The current unmet medical need in Pompe disease is for prevention of immune responses against standard-of-care <u>enzyme replacement therapy</u> ," says coauthor Dwight Koeberl, MD, PhD. "However, we foresee a future application of the dual vector strategy described in this paper, including a liver-expressing vector along with a ubiquitously expressing vector, which might achieve much higher efficacy than either vector alone."

In the article "Immunodominant Liver-Specific Expression Suppresses Transgene-Directed Immune Responses in Murine Pompe Disease," Ping Zhang and coauthors from Duke University Medical Center (Durham, NC), targeted a gene delivery vector carrying the therapeutic gene to the livers of mice with Pompe disease. Not only did the liverspecific expression of the protein induce <u>immune tolerance</u>, but when combined with non-targeted delivery of the therapeutic gene it also boosted the overall effectiveness of the treatment.

More information: DOI: 10.1089/hum.2011.063



Provided by Mary Ann Liebert, Inc

Citation: Targeted gene therapy enhances treatment for Pompe disease (2012, June 25) retrieved 6 May 2024 from https://medicalxpress.com/news/2012-06-gene-therapy-treatment-pompe-disease.html

This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.