

Gene therapy for muscular dystrophy: Lessons learned and path forward.

[Mendell JR](#) , [Rodino-Klapac L](#) , [Sahenk Z](#) , [Malik V](#) , [Kaspar BK](#) , [Walker CM](#) , [Clark KR](#) .

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[A phase I trial of adeno-associated virus serotype 1–gammaglobulin gene therapy for limb girdle muscular dystrophy type 2C - Brain 2012; doi: 10.1093/brain/awr342, published on January 11, 2012 and available on this](#)

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Eymard 3, Didier Caizergues 5, Thomas Voit 3, Olivier Benveniste 1

Intervista: □ Il Dr. Thomas Voit discute la terapia genica per le gamma-sarcoglicanopatie

<http://www.nationwidechildrens.org/dr-thomas-voit-discusses-gamma-sarcoglycan-gene-therapy-march-2012>

[Sustained alpha-sarcoglycan gene expression after gene transfer in limb-girdle muscular dystrophy, type 2D.](#)

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