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Personalized Gene and Cell Therapy for Duchenne Muscular Dystrophy

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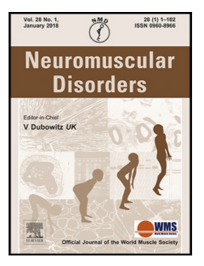
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Highlights

- There is promising therapy being evaluated for Duchenne Musuclar Dystrophy
- Some of them are being evaluated in Human clinical trial
- There is a need of better natural history for assessing efficacy of clinical trial
- There will be a necessity for individualized medecine in the future

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