X-linked myotubular myopathy: A prospective international natural history study.

Anoussamy M¹, Lilien C¹, Gidaro T¹, Gargaun E¹, Chê V¹, Schara U¹, Gangfuß A¹, D’Amico A¹, Dowling JJ¹, Darras BT¹, Daron A¹, Hernandez A¹, de Lattre C¹, Arnal JM¹, Mayer M¹, Cuisset JM¹, Vuillerot C¹, Fontaine S¹, Bellance R¹, Biancalana V¹, Buj-Bello A¹, Hogrel JY¹, Landy H¹, Servais L².

Abstract

OBJECTIVES: Because X-linked myotubular myopathy (XLMTM) is a rare neuromuscular disease caused by mutations in the MTM1 gene with a large phenotypic heterogeneity, to ensure clinical trial readiness, it was mandatory to better quantify disease burden and determine best outcome measures.

METHODS: We designed an international prospective and longitudinal natural history study in patients with XLMTM and assessed muscle strength and motor and respiratory functions over the first year of follow-up. The humoral immunity against adeno-associated virus serotype 8 was also monitored.

RESULTS: Forty-five male patients aged 3.5 months to 56.8 years were enrolled between May 2014 and May 2017. Thirteen patients had a mild phenotype (no ventilation support), 7 had an intermediate phenotype (ventilation support less than 12 hours a day), and 25 had a severe phenotype (ventilation support 12 or more hours a day). Most strength and motor function assessments could be performed even in very weak patients. Motor Function Measure 32 total score, grip and pinch strengths, and forced vital capacity, forced expiratory volume in the first second of exhalation, and peak cough flow measures discriminated the 3 groups of patients. Disease history revealed motor milestone loss in several patients. Longitudinal data on 37 patients showed that the Motor Function Measure 32 total score significantly decreased by 2%. Of the 38 patients evaluated, anti-adeno-associated virus type 8 neutralizing activity was detected in 26% with 2 patients having an inhibitory titer >1:10.

CONCLUSIONS: Our data confirm that XLMTM is slowly progressive for male survivors regardless
of their phenotype and provide outcome validation and natural history data that can support clinical
development in this population.

**CLINICALTRIALSGOV IDENTIFIER:** NCT02057705.


PMID: 30902907    DOI: 10.1212/WNL.0000000000007319