

Characterization of the muscular and cardiac diseases of the DMSXL mouse model, a transgenic mouse model for Myotonic Dystrophy type 1

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Characterization of the muscular and cardiac diseases of the DMSXL mouse model, a transgenic mouse model for Myotonic Dystrophy type 1



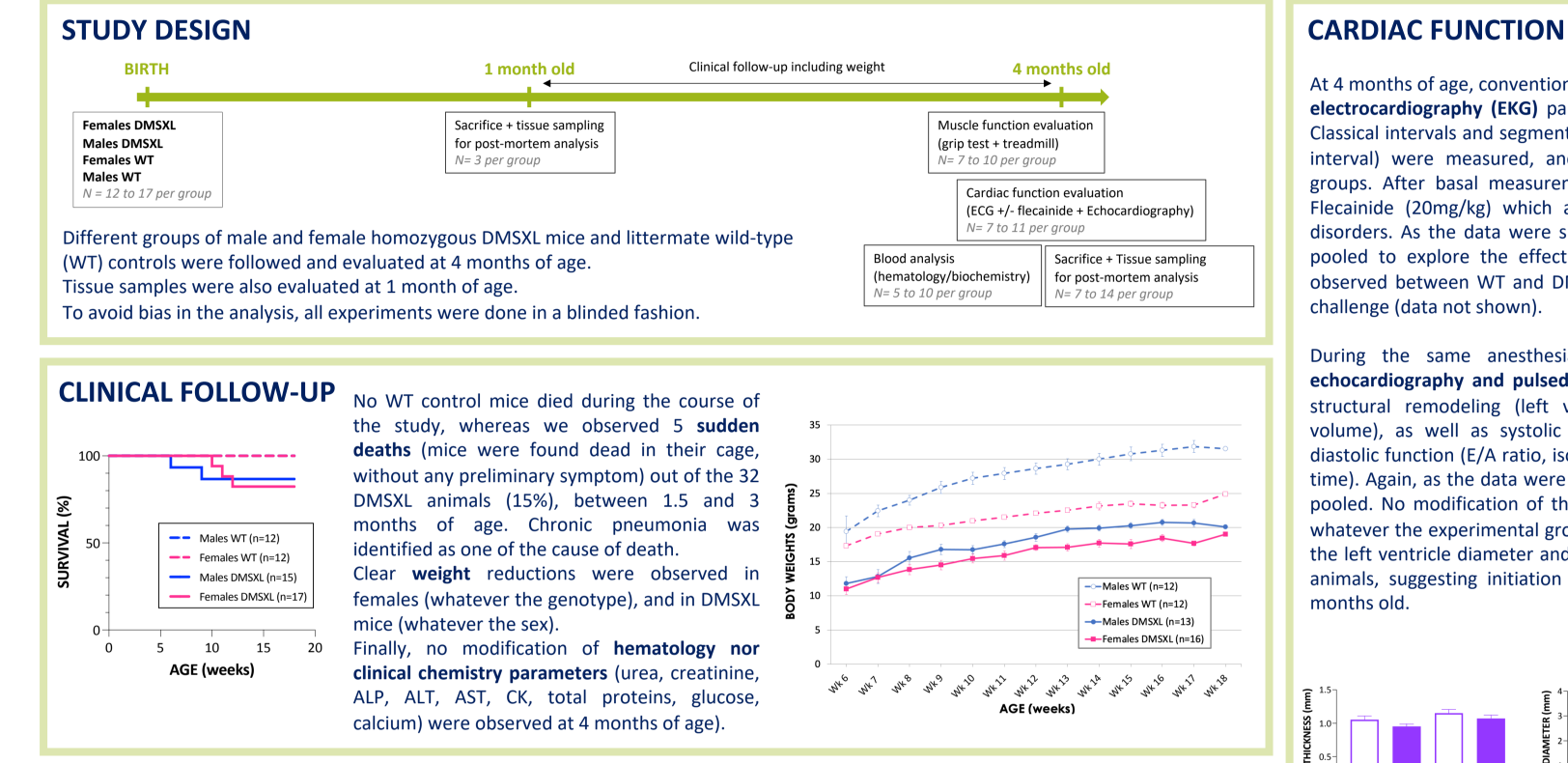
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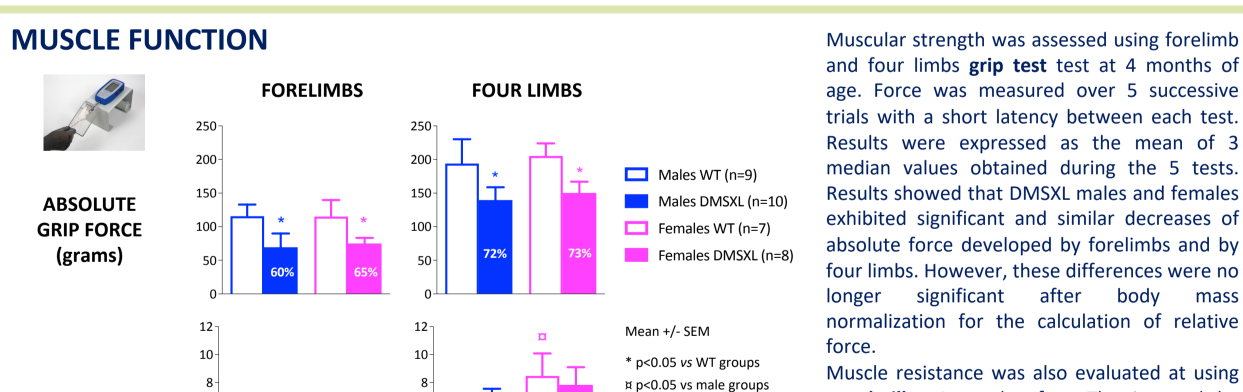
1 Nantes Université, CHU de Nantes, INSERM UMR 1089, Target Lab - Nantes, France / Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Boisbonne, ONIRIS - Nantes, France / Sorbonne Université, INSERM, Centre de Recherche en Myologie - Paris, France / Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Boisbonne, ONIRIS - Nantes, France / Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Boisbonne, ONIRIS - Nantes, France / Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Recherche en Myologie - Paris, France / Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Boisbonne, ONIRIS - Nantes, France / Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Rosklepios BioPharmaceutical, Inc. - Durham, NC, USA / Centre de Roskle



Context of the study

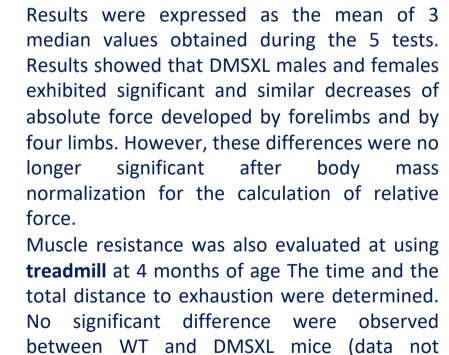
Myotonic Dystrophy type 1 (DM1) is an autosomal dominant, progressive, and muscle atrophy. Other clinical manifestations include cardiac conduction defects, cardio-respiratory problems, cataracts, endocrine dysfunction and frequent of the clinical manifestations include cardiac conduction defects, cardio-respiratory problems, cataracts, endocrine dysfunction and frequent of the clinical manifestations include cardiac conduction defects, cardio-respiratory problems, cataracts, endocrine dysfunction and frequent of the clinical manifestations include cardiac conduction defects, cardio-respiratory problems, cataracts, endocrine dysfunction and frequent of the clinical manifestations include cardiac conduction defects, cardio-respiratory problems, cataracts, endocrine dysfunction and frequent of the cardiac conduction defects at the cardiac conduction and frequent of the cardiac neurological manifestations. The disease is caused by unnaturally expanded repeats of CTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients, the number of cTG trinucleotide in the 3'-untranslated region of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the sum of the DMPK (dystrophia myotonica protein kinase) gene. In DM1 patients are the sum of the sum repeats correlated with earlier onset, more severe symptoms and shorter lifespan). CTG-containing mutant DMPK transcripts are toxic. They aggregate as nuclear foci and impact the expression and function of RNA-binding proteins (such as MBNL1 and CELF1), resulting in spliceopathy of downstream effector genes, which accounts for much of the disease phenotype. Several therapeutics approaches either pharmacological or gene-therapy based, are under investigation to address this unmet medical need. One current limitation for the efficient evaluation and development of the most investigation to address this unmet medical need. One current limitation for the efficient evaluation and development of the most investigation to address this unmet medical need. One current limitation for the efficient evaluation and development of the most investigation to address this unmet medical need. One current limitation for the efficient evaluation and development of the most investigation to address this unmet medical need. relevant remains the DMSXL mouse model, which carries a 45-kb human genomic fragment including the DMPK transgene is under the control of its own promoter and has been shown to have an almost ubiquitous expression. Initial characterization studies demonstrated that homozygous DMSXL mice display severally severally severally to the control of its own promoter and has been shown to have an almost ubiquitous expression. Initial characterization studies demonstrated that homozygous DMSXL mice display severally severally to the control of its own promoter and has been shown to have an almost ubiquitous expression. Initial characterization studies demonstrated that homozygous DMSXL mice display severally severally to the control of its own promoter and has been shown to have an almost ubiquitous expression. Initial characterization studies demonstrated that homozygous DMSXL mice display severally se manifestations of the human DM1 pathology, including growth retardation, muscle defects, cognitive impairments, nuclear foci, and splicing abnormalities. After establishing a colony in our hands the most relevant and cardiac diseases, in both genders.

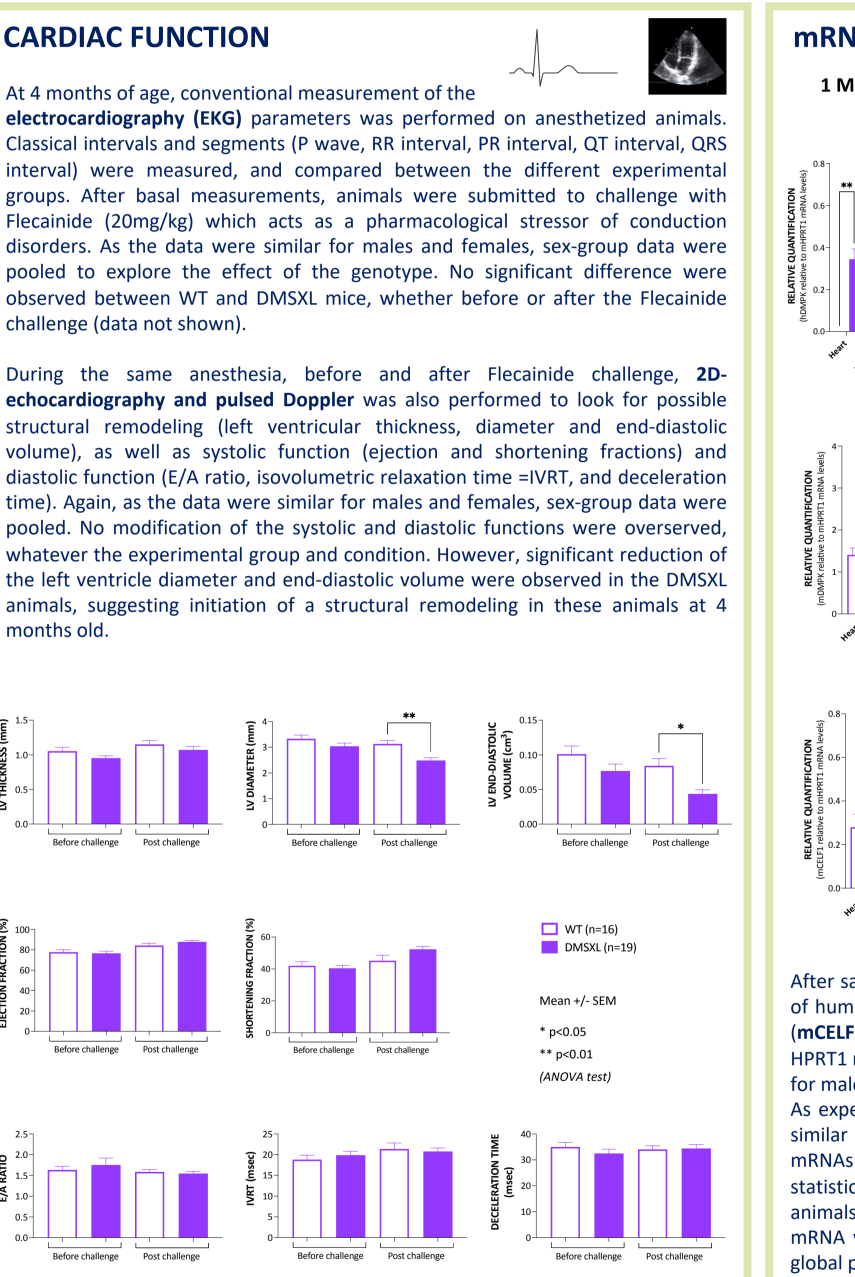


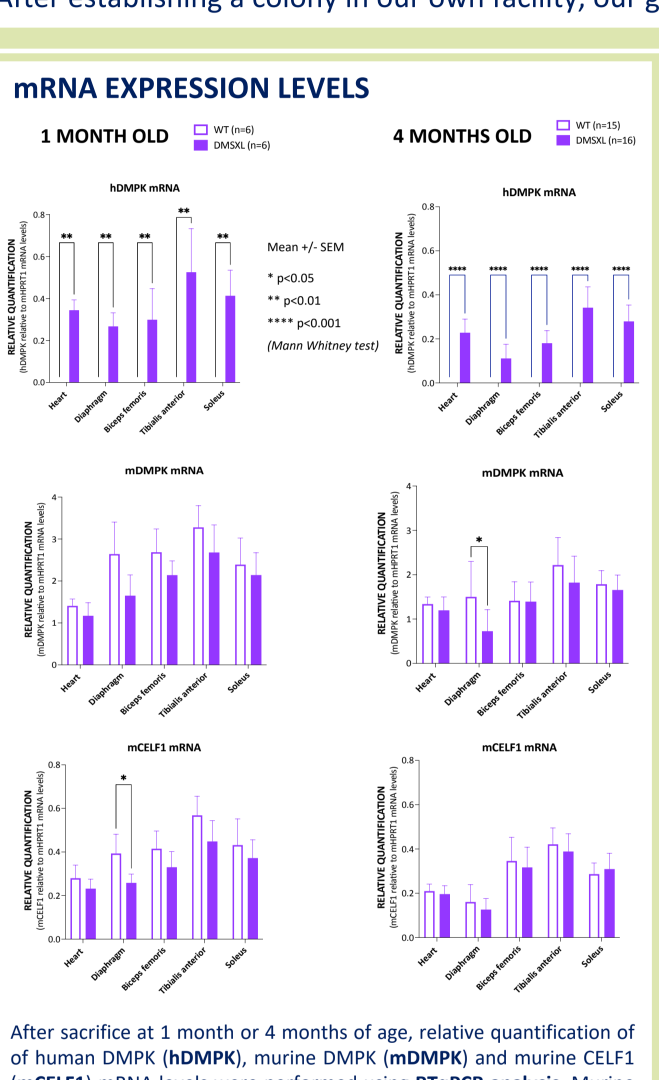


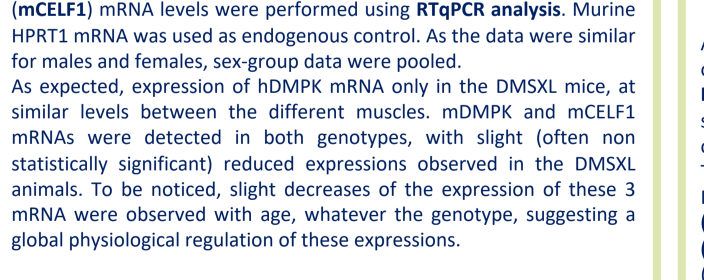
(Mann Whitney test)

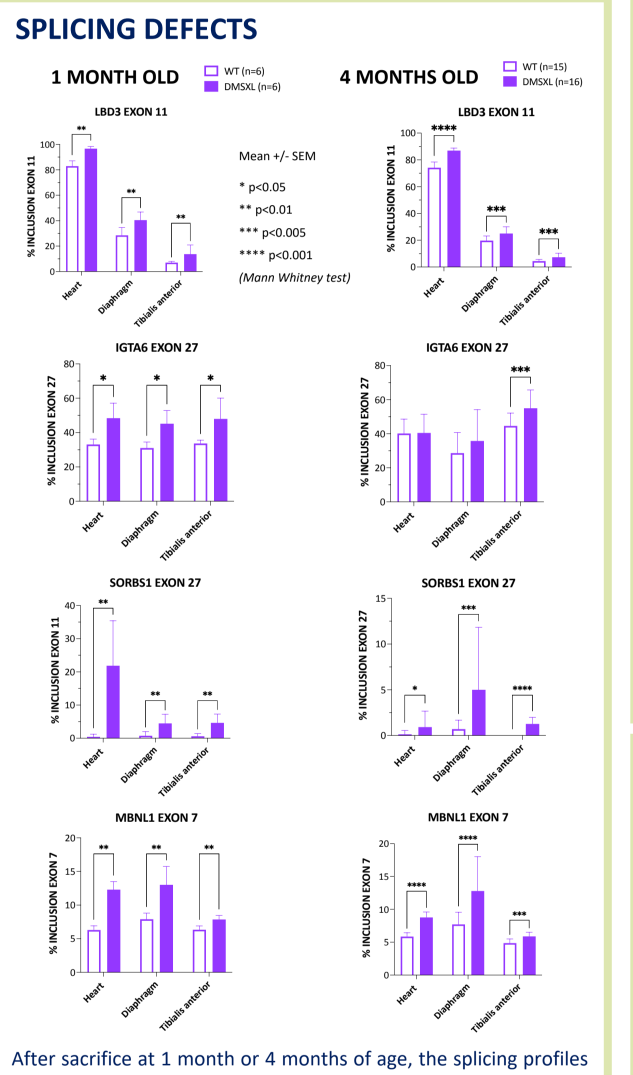
% = values of DMSXL group

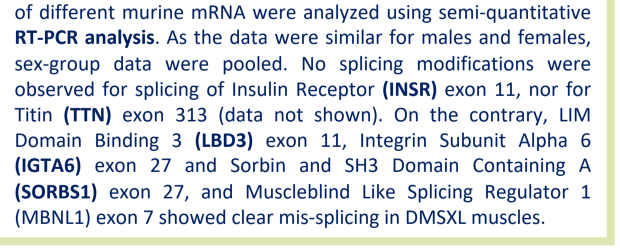


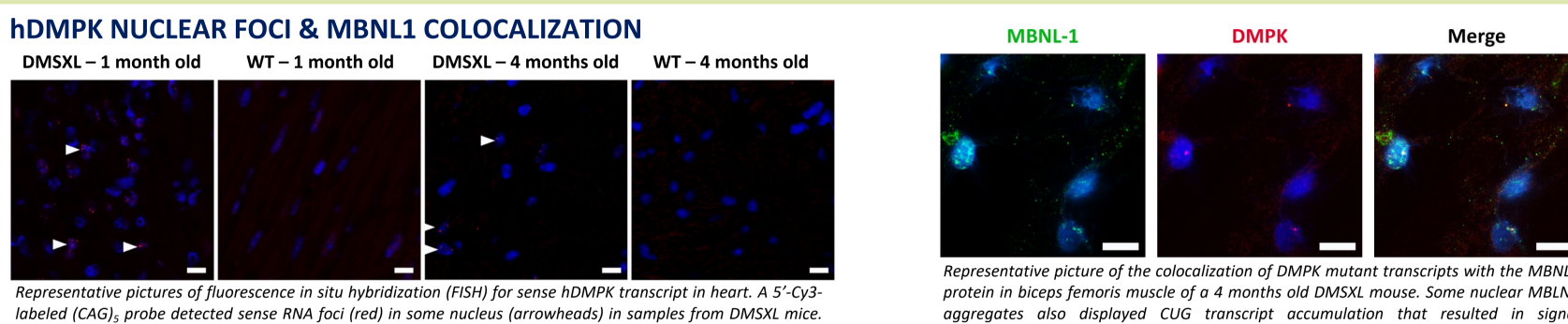


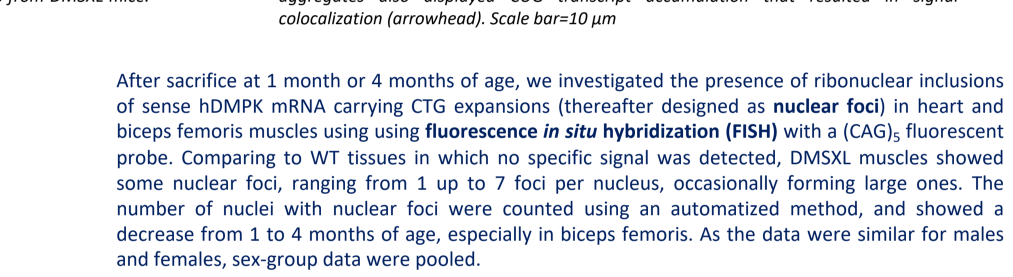












In DMSXL skeletal muscles and heart, IHC using an antibody specific for MBNL1 followed by the FISH technique showed sequestration of the MBLN1 protein next to the hDMPK nuclear foci.

After sacrifice at 1 month or 4

months of age, muscle fibers

size was measured in biceps

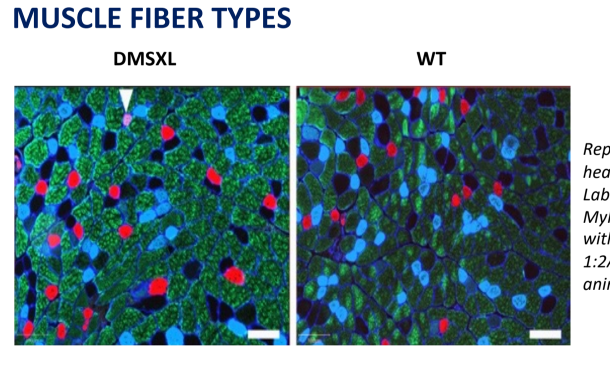
femoris muscle using minimal

Ferret diameter and a laminin

As the data were similar for

males and females, sex-group

MUSCLE FIBER SIZE



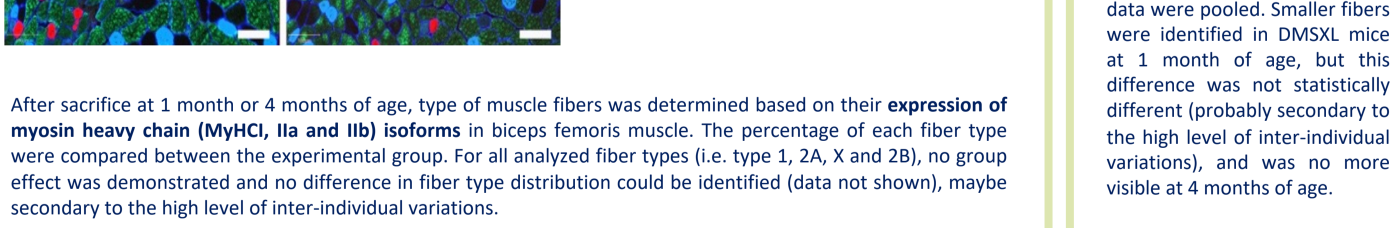
secondary to the high level of inter-individual variations.

Mean +/- SEM

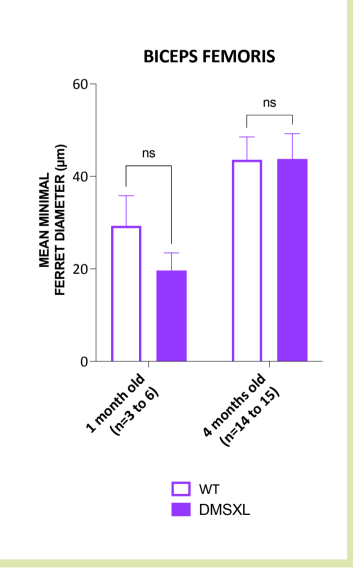
** p<0.01

**** p<0.001

vith no labelling are type X (black). Some few hybrid type :2A fibers (arrowhead) were only observed in DMSXL nimals. Scale=100 μm



immunolabelling





Conclusion

The data obtained in the frame of this study confirm that the DMSXL mouse model exhibits several features of the muscles and heart at both 1 and 4 months of age. This mutant transcript aggregates as foci in the nuclei of the muscles and heart at both 1 and 4 months of age. This mutant transcript aggregates as foci in the nuclei of the muscles and heart at both 1 and 4 months of age. This mutant transcript aggregates as foci in the nuclei of the n muscle cells, where sequestration of the MBNL1 protein was also observed. As a consequence, and even if expression of the murine CELF1 mRNA was not modified, splicing defects of several downstream messengers was found reduced in 4 ariations, no clear skeletal muscle histological abnormalities were observed. However, grip force was found reduced in 4 ariations, no clear skeletal muscle histological abnormalities were observed. However, grip force was found reduced in 4 ariations, no clear skeletal muscle histological abnormalities were observed. months old DMSXL animals. At the same age, the heart exhibited no EKG abnormalities, but an initiation of structural remodeling. Except for body weights, no gender effect was raised during this study.

All these data will be of importance to design future preclinical studies for the evaluation of the efficacy and safety of different therapeutic products designed to treat DM1 at the muscular and cardiac levels, and using this unique DMSXL mouse model.



