

BRAIN



JOURNAL ARTICLE ACCEPTED MANUSCRIPT

The new missense G376V-TDP-43 variant induces late-onset distal myopathy but not amyotrophic lateral sclerosis

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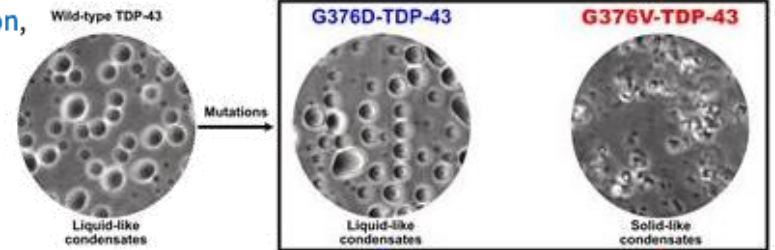
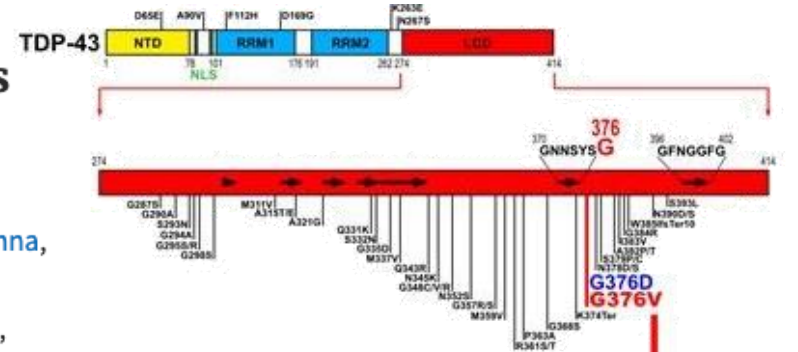
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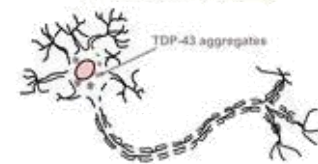
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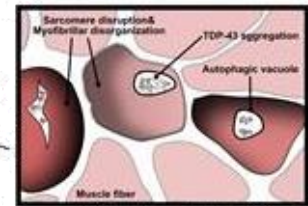


ALS&ALS/FTD
Motor neuron death

Slowly, progressive distal myopathy



Death of patients
6-60 months after the onset



Patients still alive
>20 years after the onset