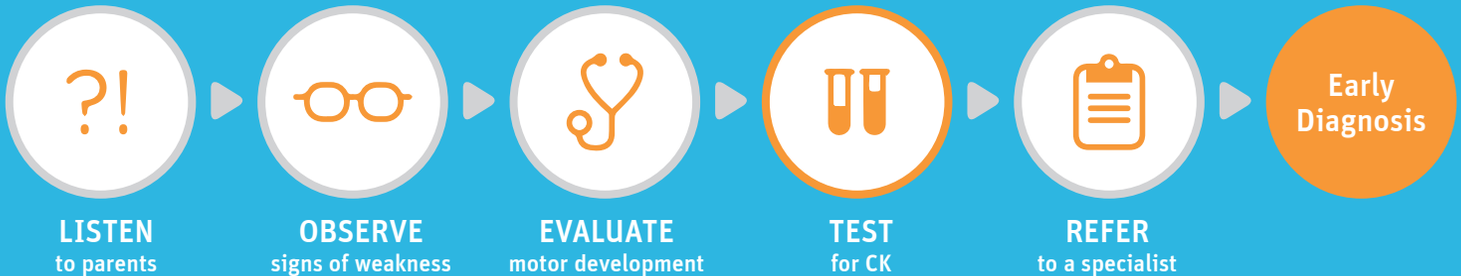


ChildMuscleWeakness.org

Early diagnosis makes a difference.

Learn the steps to identify pediatric muscle weakness and signs of peripheral neuromuscular diseases.



Developmental Delay > Do a CK

In partnership with Parent Project Muscular Dystrophy



Developmental Delay > Do a CK

The National Task Force for the Early Identification of Childhood Neuromuscular Disorders presents ChildMuscleWeakness.org

Early diagnosis of neuromuscular disorders can help target interventions, improve outcomes, reduce family stress related to diagnostic delay, and allow for accurate genetic counseling.

Visit ChildMuscleWeakness.org to access a web-based tool that provides:

- ▶ motor surveillance aids
 - ▶ video clips to illustrate early signs of motor weakness
 - ▶ recommendations on follow up and referral when motor delays are suspected, and more
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- ▶ If you suspect Duchenne muscular dystrophy, FREE genetic testing and interpretation is available through the Decode Duchenne program at ParentProjectMD.org/Decode.

