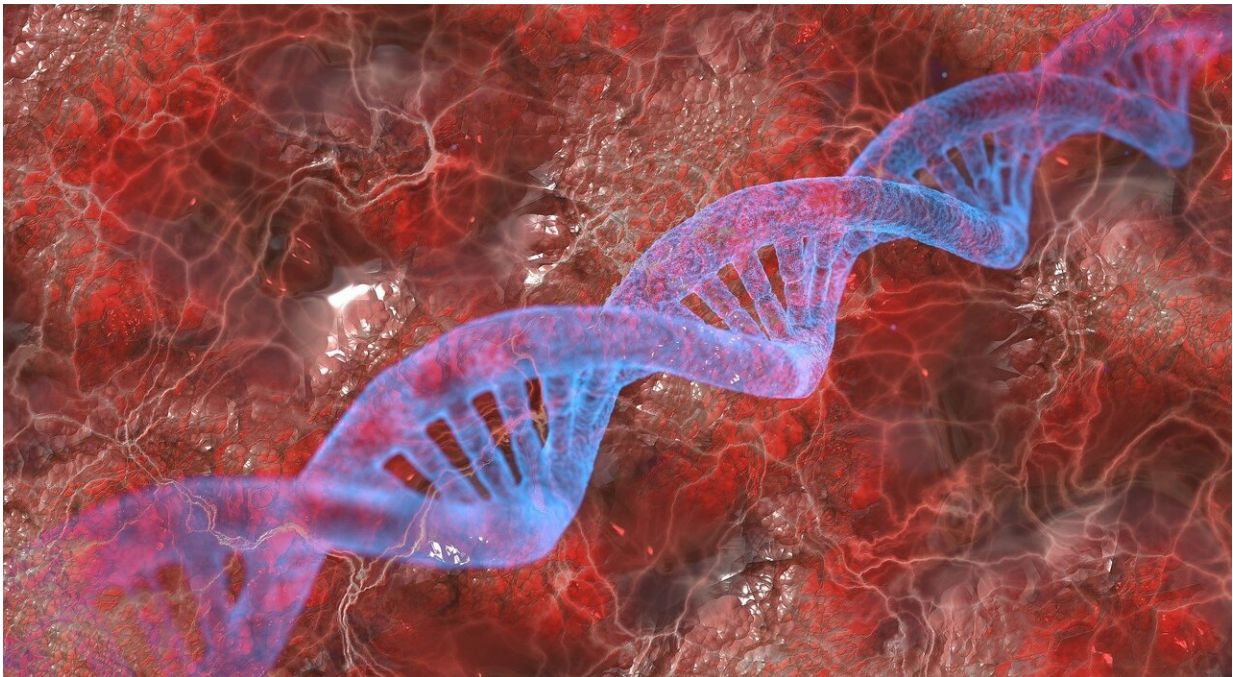


Researchers identify DNA elements that affect MECP2 expression

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Researchers at Baylor College of Medicine and the Jan and Dan Duncan Neurological Research Institute at Texas Children's Hospital (NRI) have identified and characterized two regions of DNA required for the proper expression of *Mecp2*/MECP2 in mice and humans.

These findings, published in *Genes & Development*, are helping to shed

light on the function of these DNA regions and how they could be potential targets for diagnostic and therapeutic interventions for intellectual disabilities such as Rett Syndrome and MECP2 Duplication Syndrome.

Both of these intellectual disabilities are examples of the importance of precise MeCP2 [protein](#) levels for proper brain function. A decrease in this protein leads to Rett Syndrome, while an increase in this protein causes MECP2 Duplication Syndrome. Both are severe neurological disorders characterized by learning disabilities, features of autism and motor difficulties.

Dr. Huda Zoghbi, professor at Baylor, director of the NRI, and a Howard Hughes Medical Institute investigator, underscored the importance of understanding how the levels of RNA encoding this protein are regulated. Researchers in her lab identified two DNA regions that, when mutated, lead to either a decrease or an increase in MECP2 RNA and protein levels, resulting in partial behavioral deficits seen in Rett Syndrome and MECP2 Duplication Syndrome, respectively.

Yingyao Shao, first author on the study and graduate student in the Zoghbi lab, explains that when one of the two DNA regions were altered, results showed a moderate reduction in MeCP2 levels mimicking behavioral and molecular changes observed in Rett mice models. Alteration of the other DNA [region](#) caused a slight increase in MeCP2 protein levels similar to mice models with double expression of MeCP2.

"These findings provide hope that future treatments targeting these DNA regions could have clinically relevant benefits, even when only slightly correcting MeCP2 protein levels," said Shao. "Moreover, mutations in either of these two DNA regions are likely to cause [intellectual disability](#) or autism in humans, thus it is important to sequence these regions when

screening for genetic causes of neurodevelopmental disorders," added Zoghbi.

More information: Yingyao Shao et al, Identification and characterization of conserved noncoding cis-regulatory elements that impact *Mecp2* expression and neurological functions, *Genes & Development* (2021). [DOI: 10.1101/gad.345397.120](https://doi.org/10.1101/gad.345397.120)

Provided by Baylor College of Medicine

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