Recessive COL4A2 Mutation Leads to Intellectual Disability, Epilepsy, and Spastic Cerebral Palsy

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Neurol Genet 2021;7:e583. doi:10.1212/NXG.0000000000000583

Dominant negative or haploinsufficient mutations in the collagen genes COL4A1 and COL4A2 are characterized by arterial basement membrane thickening resulting in a multisystem microangiopathy targeting the CNS but also potentially affecting the ocular, renal, cardiac, and muscular systems. Within the brain, such changes predispose affected individuals to recurrent ischemic and/or hemorrhagic strokes beginning during early fetal development but extending into the postnatal period and even into adulthood. Mutations affecting glycine residues of the Gly-Xaa-Yaw (typically representing glycine-proline-4-trans-hydroxyproline in vertebrates) repeat domains that typify collagens usually manifest in an autosomal dominant (AD) fashion. However, recent work suggests that tissue-specific mutation effects may also occur, with mutations leading to gain of function effects in some tissues and loss of function effects in others. Stroke-related complications may be insidious and clinically silent. Neuroimaging phenotypes of COL4A-associated disease include chronic white matter disease, porencephaly/hydranencephaly, encephalomalacia, cerebral calcifications, schizencephaly and hydrocephalus and corresponding clinical diagnoses of cerebral palsy, intellectual disability, cortical visual impairment, and epilepsy.

Type IV collagen α chains form heterotrimers with β chains in a 2:1 ratio, and incompletely penetrant AD inheritance is typical. Although autosomal recessive mutations of COL4A1 have been described, prior reports of COL4A2-associated disease have all featured AD inheritance. We describe 2 children from a consanguineous Iranian family with intellectual disability, spastic cerebral palsy, and epilepsy who each harbored the same homozygous mutation in COL4A2.

Pregnancies for both children were uncomplicated, and both were born at term by vaginal delivery. The couple’s 8 year-old son was bedridden and exhibited cortical visual impairment. He did not fix or track stimuli. He had never been able to sit unassisted or control his head. He did not speak or demonstrate communicative intent. He had focal epilepsy that began at age 6 months, partially controlled with carbamazepine. Motor examination revealed spastic quadriplegia with ophthalmoplegia, nystagmus, and skew deviation. Brain MRI revealed bilateral colpocephaly and irregular ventricular contours. His sister was 20 years old at the time of evaluation. Milestone attainment had been globally delayed. She also had focal epilepsy with onset at 18 months. Seizures were controlled with phenobarbital and carbamazepine. She began walking independently at age 3 years. At the time of evaluation, she did not speak, and her
examination was significant for asymmetric spastic-dystonic quadriparesis with right-sided predominance. Her brain MRI showed irregular lateral ventricle contour with coalescent porencephaly and generalized cortical atrophy (figure, A). Neither patient was known to have ocular, kidney, skeletal, or cardiac muscle disease.

Both affected children and their parents provided informed consent for study participation in accordance with local ethics oversight and underwent whole-exome sequencing with filtering and variant prioritization as described previously. This revealed a homozygous c.3472G>C (p.G1158R) variant in COL4A2 (NM_001846) segregating with disease status in the family (both parents are neurologically healthy, although neuroimaging was not able to be performed) (figure, B). This variant is novel (not found in gnomAD or the Greater Middle Eastern Variome server) and predicted to be deleterious (MetaSVM, CADD, SIFT, and PolyPhen), putatively by disrupting a glycine residue within a highly conserved Gly-Xaa-Yaw collagen triple helix repeat domain. Such glycine residues are known to be critical determinants of collagen stability and we anticipate that the substitution of an arginine residue to have a destabilizing effect.

In comparison to previously described incompletely penetrant dominantly inherited mutations in COL4A2, this homozygous (p.G1158R) variant appears to exhibit autosomal recessive inheritance, although our observations will benefit from subsequent confirmation. Our results indicate that both dominant and recessive forms of COL4A2 disease exist. This finding has potentially important implications for the interpretation of clinical genetic testing in cases of porencephaly, hydrocephalus or idiopathic antenatal or perinatal ischemic/hemorrhagic stroke and for associated clinical phenotypes including epilepsy, intellectual disability, and cerebral palsy.

Data Availability
Full data are available to qualified investigators on reasonable request to the corresponding author.

Acknowledgment
The authors thank the patients and their family members for their support of this work.

Study Funding
NIH; 1R01 NS106298.

Disclosure
MCK serves as a consultant for PTC Therapeutics and the United States National Health Services Administration and has performed grant review for the United States Department of Defense. The remaining authors report no disclosures.
related to the conduct of this manuscript. Bioinformatic and statistical analysis performed by SB and SCJ. Go to Neurology.org/NG for full disclosures.

**Publication History**

Received by Neurology: Genetics September 21, 2020. Accepted in final form February 15, 2021.

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**Appendix**

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**References**


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**Appendix (continued)**

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*Neurol Genet* 2021;7;
DOI 10.1212/NXG.0000000000000583

This information is current as of April 26, 2021
Recessive COL4A2 Mutation Leads to Intellectual Disability, Epilepsy, and Spastic Cerebral Palsy

In the Clinical/Scientific Notes article “Recessive COL4A2 Mutation Leads to Intellectual Disability, Epilepsy, and Spastic Cerebral Palsy” by Bakhtiari et al.,1 the sixth author’s name should be listed as “Saghar Ghasemi Firouzabadi.” The editorial staff regret the error. Additionally, Dr. Hossein Darvish’s affiliation should have been listed as “Neuroscience Research Center, Faculty of Medicine, Golestan University of Medical Sciences, Gorgan, Iran.” The authors regret the error.

Reference