



*CGPH / Custom Gene Panel, Hereditary, Next-Generation Sequencing, Varies can be used to modify any epilepsy panel or test any single gene included on any epilepsy panel.

**Segregation studies can be performed to determine if a variant segregates with the condition in a family and/or occurred *de novo*, which may clarify the significance of a variant. For more information, contact the Laboratory Genetic Counselors at 800-533-1710.

Metabolic Tests to Consider
AACSF / Amino Acids, Quantitative, Spinal Fluid
AAQP / Amino Acids, Quantitative, Plasma
CDG / Carbohydrate Deficient Transferrin for Congenital Disorders of Glycosylation, Serum
CRDPU / Creatine Disorders Panel, Urine
LACS1 / Lactate, Plasma
OAU / Organic Acids Screen, Urine
OLIGU / Oligosaccharide Screen, Urine
PIPA / Picoic Acid, Serum
PIPU / Picoic Acid, Urine, if newborn
PLSD / Lysosomal and Peroxisomal Storage Disorders Screen, Blood Spot, if <18 years of age
POXP / Fatty Acid Profile, Peroxisomal (C22-C26), Plasma
PUPYP / Purines and Pyrimidines Panel, Plasma
PYP / Pyruvic Acid, Blood
PYRC / Pyruvate, Spinal Fluid
Autoimmune Evaluations to Consider
EPC2 / Epilepsy, Autoimmune/Paraneoplastic Evaluation, Spinal Fluid
EPS2 / Epilepsy, Autoimmune/Paraneoplastic Evaluation, Serum