



| *Individual Gene Suspects | | |
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| Type | Gene | OMIM |
| Ceroid lipofuscinosis | CLN2 | 204500 |
| Ceroid lipofuscinosis | CLN3 | 204200 |
| Dravet syndrome, severe myoclonic epilepsy of infancy | SCN1A | 607208 |
| Encephalopathy epileptic, early infantile | PCDH19 | 300088 |
| Encephalopathy epileptic, early infantile; seizures, benign neonatal | KCNQ2 | 613720; 121200 |
| GLUT1-deficiency syndrome | SLC2A1 | 606777 |
| Infantile spasms | ARX | 308350 |
| Myoclonic epilepsy progressive | CSTB | 254800 |
| Myoclonus-nephropathy | SCARB2 | 254900 |
| Pyridoxine-dependent epilepsy | ALDH7A1 | 266100 |
| Rett syndrome (order as MECP2 / MECP2 Gene, Full Gene Analysis) | MECP2 | 312750 |
| Seizures, benign neonatal | KCNQ3 | 121201 |
| Temporal epilepsy familial | LG11 | 600512 |

| Metabolic Tests to Consider |
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| AACSF / Amino Acids, Quantitative, Spinal Fluid |
| AAQP / Amino Acids, Quantitative, Plasma |
| CDG / Carbohydrate Deficient Transferin for Congenital Disorders of Glycosylation, Serum |
| CRDPU / Creatine Disorders Panel, Urine |
| LAA / Lactate, Plasma |
| LABF / Lactate, Body Fluid |
| OUU / Organic Acids Screen, Urine |
| OLIGU / Oligosaccharide Screen, Urine |
| PIPA / PIPeocolic Acid, Serum |
| PIPU / PIPeocolic 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 |
| PYR / Pyruvic Acid, Blood |
| PYRC / Pyruvate, Spinal Fluid |

| Autoimmune Evaluations to Consider |
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| EPC1 / Epilepsy, Autoimmune Evaluation, Spinal Fluid |
| EPS1 / Epilepsy, Autoimmune Evaluation, Serum |

