

CGT Essential

| chrom | OMIM (gene) | Gene | Previous symbol | OMIM (phen) | DISEASE | MOI |
|-------|-------------|---------|-----------------|-------------------|---|---|
| 1 | 607008 | ACADM | | 201450 | Medium-chain acyl-CoA dehydrogenase deficiency | Autosomal recessive |
| 2 | 604285 | AGXT | | 259900 | Hyperoxaluria, primary, type 1 | Autosomal recessive |
| 22 | 607574 | ARSA | | 250100 | Metachromatic leukodystrophy | Autosomal recessive |
| 3 | 609019 | BTD | | 253260 | Biotinidase deficiency | Autosomal recessive |
| 21 | 613381 | CBS | | 236200 | Homocystinuria due to cystathionine beta-synthase | Autosomal recessive |
| 7 | 602421 | CFTR | | 219700 | Cystic fibrosis | Autosomal recessive |
| 11 | 602858 | DHCR7 | | 270400 | Smith-Lemli-Opitz syndrome | Autosomal recessive |
| X | 300384 | EMD | | 310300 | Emery-Dreifuss muscular dystrophy, type 1, X-linked | X-linked |
| X | 309550 | FMRI | | 300624 | Fragile X syndrome | X-linked |
| 17 | 606800 | GAA | | 232300 | Glycogen storage disease, type 2 | Autosomal recessive |
| 9 | 606999 | GALT | | 230400 | Galactosemia | Autosomal recessive |
| X | 300644 | GLA | | 301500 | Fabry disease | X-linked |
| 2 | 600890 | HADHA | | 609016; 609015 | Long-chain 3-hydroxyl-CoA dehydrogenase (LCHAD) deficiency; Mitochondrial trifunctional protein deficiency | Autosomal recessive |
| 11 | 141900 | HBB | | 603903 | HBB-related hemoglobinopathy | Autosomal recessive |
| 1 | 609831 | MMACHC | | 277400 | Methylmalonic aciduria and homocystinuria, cbIC type | Autosomal recessive, digenic inheritance (PRDX1 gene) |
| 12 | 612349 | PAH | | 261600 | Phenylketonuria | Autosomal recessive |
| 16 | 601785 | PMM2 | | 212065 | Congenital disorder of glycosylation, type 1A | Autosomal recessive |
| 5 | 606718 | SLC26A2 | | 600972 | Achondrogenesis, type 1B (diastrophic dysplasia) | Autosomal recessive |
| 5 | 600354 | SMN1 | | 253300 | Spinal muscular atrophy | Autosomal recessive |