

Title: MEGDEL Syndrome *GeneReview* – Table 3 with Additional Protein Information
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Table 3a. New Classification for Inborn Errors of Metabolism with 3-Methylglutaconic Aciduria as Discriminative Feature

| Patho-Mechanism | Disease Name | Former Designation | Additional Hallmarks ¹ of Phenotype | Affected Gene / Protein / Subcellular Localization / Proposed Function |
|-----------------------------------|--|----------------------------|---|---|
| Primary 3-MGA-uria | | | | |
| Organic aciduria | 3-methylglutaconyl-CoA hydratase deficiency (<i>AUH</i> defect) | 3-MGCA type I (3-MGCA-1) | Adult onset leukoencephalopathy, dementia, progressive spasticity | <i>AUH</i> / 3-methylglutaconyl-CoA hydratase / Mitochondrial matrix / Leucine catabolism |
| Secondary 3-MGA-uria | | | | |
| Defective phospholipid remodeling | <i>TAZ</i> defect (Barth syndrome) | 3-MGCA type II (3-MGCA-2) | (Cardio)myopathy, short stature, neutropenia, hypocholesterolemia, cognitive phenotype, mild dysmorphic features, OXPHOS dysfunction | <i>TAZ</i> / Tafazzin / (Inner) mitochondrial membrane / Cardiolipin remodeling |
| | <i>SERAC1</i> defect (MEGDEL syndrome) | 3-MGCA type IV (3-MGCA-4) | Progressive spasticity, dystonia, deafness, Leigh syndrome-like MRI, severe psychomotor retardation, hypocholesterolemia, OXPHOS dysfunction | <i>SERAC1</i> / <i>SERAC1</i> / Mitochondria-associated membranes fraction / Phosphatidylglycerol remodeling, cardiolipin composition |
| Mitochondrial membrane disorder | <i>OPA3</i> defect (Costeff syndrome) | 3-MGCA type III (3-MGCA-3) | Ataxia/extrapyramidal dysfunction, optic atrophy | <i>OPA3</i> / <i>OPA3</i> / Outer mitochondrial membrane / ?Protective function for respiratory chain |
| | <i>TMEM70</i> defect | 3-MGCA type IV (3-MGCA-4) | Broad phenotype, hypertrophic cardiomyopathy, myopathy, dysmorphic features, cataracts, psychomotor retardation, ATPase deficiency, lactic acidosis, hyperammonemia | <i>TMEM70</i> / <i>TMEM70</i> / Inner mitochondrial membrane / ?Complex V assembly and inner mitochondrial membrane |
| | <i>DNAJC19</i> defect (DCMA) | 3-MGCA type V | Dilated cardiomyopathy, ECG abnormalities, non-progressive cerebellar ataxia, small | <i>DNAJC19</i> / <i>DNAJC19</i> / ?Inner mitochondrial membrane |

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|---------|----------------|---------------------------|--|--|
| | syndrome) | (3-MGCA-5) | atrophic testes, cryptorchidism, growth failure, anemia, steatosis hepatitis (i.e., fatty liver) | membrane / ?Mitochondrial protein import |
| Unknown | NOS 3-MGA-uria | 3-MGCA type IV (3-MGCA-4) | Variable, mostly progressive neurologic disease | Unknown |

From Wortmann et al [2013]

OXPHOS = oxidative phosphorylation

DCMA = dilated cardiomyopathy with ataxia

NOS= not otherwise specified

1. In addition to 3-MGA-uria

References

Wortmann SB, Duran M, Anikster Y, Barth PG, Sperl W, Zschocke J, Morava E, Wevers RA. Inborn errors of metabolism with 3-methylglutaconic aciduria as discriminative feature: proper classification and nomenclature. J Inherit Metab Dis. 2013;36:923–8.