Many mitochondrial disorders, resulting from mutations in both nuclear and mitochondrial DNA, have been characterised. Here, we focus on some of the diseases caused by mutations to mitochondrial DNA.

### LHON (Leber's hereditary optic neuropathy)
- Mutations: 3460G>A (complex I: ND1), 11778G>A (complex I: ND4), 14484T>C (complex I: ND6)
- Affects: Retinal ganglion cells
- Symptoms: Sudden, painless loss of central vision, initially in one eye; second eye affected after 2 – 3 months

### MELAS (Mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes)
- Mutations: 3243A>G (transfer RNA: leu), 3271T>C (transfer RNA: leu)
- Affects: Brain and nervous system
- Symptoms: Stroke-like episodes in the brain, migraines, vomiting, seizures; muscle weakness, exercise intolerance

### MERFF (Myoclonic Epilepsy with Ragged Red Fibers)
- Mutations: 8344A>G (transfer RNA: lys), 8356T>C (transfer RNA: lys)
- Affects: Muscles and nervous system
- Symptoms: Myoclonus (muscle jerks), seizures, ataxia and muscle weakness; deafness and short stature

### NARP (Neuropathy, ataxia, and retinitis pigmentosa)
- Mutation: 8993T>G (ATP synthase: ATP6)
- Affects: Nervous system
- Symptoms: NARP causes retinitis pigmentosa (blindness), neuropathy and ataxia

Other mitochondrial diseases involve large deletions of the mitochondrial genome:
- Kearns-Sayre syndrome: myopathy, ophthalmoplegia, cardiomyopathy
- CPEO: paralysis of eye movement