## Daniele Ghezzi ScD

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Diagnostic Challenges in Late Onset Multiple Acyl-CoA Dehydrogenase Deficiency: Clinical, Morphological, and Genetic Aspects. Frontiers in Neurology, 2022, 13, 815523.	2.4	7
2	Biallelic Variants in ENDOG Associated with Mitochondrial Myopathy and Multiple mtDNA Deletions. Cells, 2022, 11, 974.	4.1	4
3	Clinical implementation of RNA sequencing for Mendelian disease diagnostics. Genome Medicine, 2022, 14, 38.	8.2	85
4	Response to: Phenotypic heterogeneity of Leigh syndrome due to <i>NDUFA12</i> variants is multicausal. Human Mutation, 2022, 43, 99-100.	2.5	0
5	A novel homozygous MSTO1 mutation in Ashkenazi Jewish siblings with ataxia and myopathy. Journal of Human Genetics, 2021, 66, 835-840.	2.3	3
6	A Clinical-Based Diagnostic Approach to Cerebellar Atrophy in Children. Applied Sciences (Switzerland), 2021, 11, 2333.	2.5	0
7	Novel <i>NDUFA12</i> variants are associated with isolated complex I defect and variable clinical manifestation. Human Mutation, 2021, 42, 699-710.	2.5	12
8	Impaired complex I repair causes recessive Leber's hereditary optic neuropathy. Journal of Clinical Investigation, 2021, 131, .	8.2	89
9	Exploiting pyocyanin to treat mitochondrial disease due to respiratory complex III dysfunction. Nature Communications, 2021, 12, 2103.	12.8	16
10	Current and New Next-Generation Sequencing Approaches to Study Mitochondrial DNA. Journal of Molecular Diagnostics, 2021, 23, 732-741.	2.8	21
11	Myopathic mitochondrial DNA depletion syndrome associated with biallelic variants in <i>LIG3</i> . Brain, 2021, 144, e74-e74.	7.6	5
12	Leber's Hereditary Optic Neuropathy: A Report on Novel mtDNA Pathogenic Variants. Frontiers in Neurology, 2021, 12, 657317.	2.4	13
13	Role of PITRM1 in Mitochondrial Dysfunction and Neurodegeneration. Biomedicines, 2021, 9, 833.	3.2	17
14	Clinical, imaging, biochemical and molecular features in Leigh syndrome: a study from the Italian network of mitochondrial diseases. Orphanet Journal of Rare Diseases, 2021, 16, 413.	2.7	16
15	Expanding the phenotypic spectrum of <i>BCS1L</i> â€related mitochondrial disease. Annals of Clinical and Translational Neurology, 2021, 8, 2155-2165.	3.7	11
16	Bi-allelic variants in OGDHL cause a neurodevelopmental spectrum disease featuring epilepsy, hearing loss, visual impairment, and ataxia. American Journal of Human Genetics, 2021, 108, 2368-2384.	6.2	12
17	Response to: "Heterogeneous phenotypic expression of C1QBP variants is attributable to variable heteroplasmy of secondary mtDNA deletions and mtDNA copy number― Human Mutation, 2020, 41, 2014-2015.	2.5	0
18	Homozygous mutations in <i>C1QBP</i> as cause of progressive external ophthalmoplegia (PEO) and mitochondrial myopathy with multiple mtDNA deletions. Human Mutation, 2020, 41, 1745-1750.	2.5	15

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19	A homozygous MRPL24 mutation causes a complex movement disorder and affects the mitoribosome assembly. Neurobiology of Disease, 2020, 141, 104880.	4.4	29
20	ATPase Domain <scp><i>AFG3L2</i></scp> Mutations Alter <scp>OPA1</scp> Processing and Cause Optic Neuropathy. Annals of Neurology, 2020, 88, 18-32.	5.3	31
21	Biâ€allelic pathogenic variants in <i>NDUFC2</i> cause earlyâ€onset Leigh syndrome and stalled biogenesis of complex I. EMBO Molecular Medicine, 2020, 12, e12619.	6.9	17
22	Epileptic phenotypes in children with earlyâ€onset mitochondrial diseases. Acta Neurologica Scandinavica, 2019, 140, 184-193.	2.1	15
23	Mutations in <i>ELAC2</i> associated with hypertrophic cardiomyopathy impair mitochondrial tRNA 3′â€end processing. Human Mutation, 2019, 40, 1731-1748.	2.5	31
24	The zebrafish orthologue of the human hepatocerebral disease gene <i>MPV17</i> plays pleiotropic roles in mitochondria. DMM Disease Models and Mechanisms, 2019, 12, .	2.4	21
25	Clinical-genetic features and peculiar muscle histopathology in infantile <i>DNM1L</i> -related mitochondrial epileptic encephalopathy. Human Mutation, 2019, 40, 601-618.	2.5	31
26	Homozygous variant in <i>OTX2</i> and possible genetic modifiers identified in a patient with combined pituitary hormone deficiency, ocular involvement, myopathy, ataxia, and mitochondrial impairment. American Journal of Medical Genetics, Part A, 2019, 179, 827-831.	1.2	4
27	APOPT 1/ COA 8 assists COX assembly and is oppositely regulated by UPS and ROS. EMBO Molecular Medicine, 2019, 11, .	6.9	19
28	Benign hereditary chorea and deletions outside NKX2-1: What's the role of MBIP?. European Journal of Medical Genetics, 2018, 61, 581-584.	1.3	9
29	Neonatal mitochondrial leukoencephalopathy with brain and spinal involvement and high lactate: expanding the phenotype of ISCA2 gene mutations. Metabolic Brain Disease, 2018, 33, 805-812.	2.9	20
30	Clinical, biochemical, and genetic features associated with <i>VARS2</i> -related mitochondrial disease. Human Mutation, 2018, 39, 563-578.	2.5	22
31	SLC25A10 biallelic mutations in intractable epileptic encephalopathy with complex I deficiency. Human Molecular Genetics, 2018, 27, 499-504.	2.9	37
32	KARS-related diseases: progressive leukoencephalopathy with brainstem and spinal cord calcifications as new phenotype and a review of literature. Orphanet Journal of Rare Diseases, 2018, 13, 45.	2.7	32
33	Compound heterozygous missense and deep intronic variants in NDUFAF6 unraveled by exome sequencing and mRNA analysis. Journal of Human Genetics, 2018, 63, 563-568.	2.3	15
34	Neurologic Phenotypes Associated With Mutations in <i>RTN4IP1</i> ( <i>OPA10</i> ) in Children and Young Adults. JAMA Neurology, 2018, 75, 105.	9.0	26
35	Clinical and Biochemical Features in a Patient With Mitochondrial Fission Factor Gene Alteration. Frontiers in Genetics, 2018, 9, 625.	2.3	34
36	Concurrent <i>AFG3L2</i> and <i>SPG7</i> mutations associated with syndromic parkinsonism and optic atrophy with aberrant OPA1 processing and mitochondrial network fragmentation. Human Mutation, 2018, 39, 2060-2071.	2.5	32

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37	Mutations in <i>TIMM50</i> compromise cell survival in OxPhosâ€dependent metabolic conditions. EMBO Molecular Medicine, 2018, 10, .	6.9	23
38	Human diseases associated with defects in assembly of OXPHOS complexes. Essays in Biochemistry, 2018, 62, 271-286.	4.7	75
39	Clinical, biochemical and genetic spectrum of 70 patients with ACAD9 deficiency: is riboflavin supplementation effective?. Orphanet Journal of Rare Diseases, 2018, 13, 120.	2.7	61
40	Clinicopathologic and molecular spectrum of <i>RNASEH1</i> -related mitochondrial disease. Neurology: Genetics, 2017, 3, e149.	1.9	19
41	Recessive mutations in <i>MSTO1</i> cause mitochondrial dynamics impairment, leading to myopathy and ataxia. Human Mutation, 2017, 38, 970-977.	2.5	44
42	Genetic diagnosis of Mendelian disorders via RNA sequencing. Nature Communications, 2017, 8, 15824.	12.8	432
43	A novel de novo dominant mutation in <i>ISCU</i> associated with mitochondrial myopathy. Journal of Medical Genetics, 2017, 54, 815-824.	3.2	25
44	Functionally pathogenic <i>EARS2</i> variants in vitro may not manifest a phenotype in vivo. Neurology: Genetics, 2017, 3, e162.	1.9	11
45	Not only dominant, not only optic atrophy: expanding the clinical spectrum associated with OPA1 mutations. Orphanet Journal of Rare Diseases, 2017, 12, 89.	2.7	39
46	Novel mutations in IBA57 are associated with leukodystrophy and variable clinical phenotypes. Journal of Neurology, 2017, 264, 102-111.	3.6	38
47	Biallelic Mutations in <i>DNM1L</i> are Associated with a Slowly Progressive Infantile Encephalopathy. Human Mutation, 2016, 37, 898-903.	2.5	64
48	<i><scp>ADCK</scp>3</i> mutations with epilepsy, strokeâ€like episodes and ataxia: a <scp>POLG</scp> mimic?. European Journal of Neurology, 2016, 23, 1188-1194.	3.3	42
49	<i>COA7</i> ( <i>Clorf163/RESA1</i> ) mutations associated with mitochondrial leukoencephalopathy and cytochrome c oxidase deficiency. Journal of Medical Genetics, 2016, 53, 846-849.	3.2	40
50	Clinical findings in a patient with <i>FARS2</i> mutations and earlyâ€infantileâ€encephalopathy with epilepsy. American Journal of Medical Genetics, Part A, 2016, 170, 3004-3007.	1.2	24
51	Riboflavin-Responsive and -Non-responsive Mutations in FAD Synthase Cause Multiple Acyl-CoA Dehydrogenase and Combined Respiratory-Chain Deficiency. American Journal of Human Genetics, 2016, 98, 1130-1145.	6.2	118
52	<i>LYRM7</i> mutations cause a multifocal cavitating leukoencephalopathy with distinct MRI appearance. Brain, 2016, 139, 782-794.	7.6	51
53	Disease-Causing SDHAF1 Mutations Impair Transfer of Fe-S Clusters to SDHB. Cell Metabolism, 2016, 23, 292-302.	16.2	89
54	New genes and pathomechanisms in mitochondrial disorders unraveled by NGS technologies. Biochimica Et Biophysica Acta - Bioenergetics, 2016, 1857, 1326-1335.	1.0	87

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55	A novel AIFM1 mutation expands the phenotype to an infantile motor neuron disease. European Journal of Human Genetics, 2016, 24, 463-466.	2.8	51
56	A nonsense mutation of human <scp>XRCC</scp> 4 is associated with adultâ€onset progressive encephalocardiomyopathy. EMBO Molecular Medicine, 2015, 7, 918-929.	6.9	24
57	Mitochondrial leukoencephalopathy and complex II deficiency associated with a recessive SDHB mutation with reduced penetrance. Molecular Genetics and Metabolism Reports, 2015, 5, 51-54.	1.1	16
58	COQ4 Mutations Cause a Broad Spectrum of Mitochondrial Disorders Associated with CoQ10 Deficiency. American Journal of Human Genetics, 2015, 96, 309-317.	6.2	86
59	RNASEH1 Mutations Impair mtDNA Replication and Cause Adult-Onset Mitochondrial Encephalomyopathy. American Journal of Human Genetics, 2015, 97, 186-193.	6.2	91
60	Mitochondrial dysfunction in Parkinson disease: evidence in mutant PARK2 fibroblasts. Frontiers in Genetics, 2015, 6, 78.	2.3	77
61	Mitochondrial Complex III Deficiency Caused by TTC19 Defects: Report of a Novel Mutation and Review of Literature. JIMD Reports, 2015, 22, 115-120.	1.5	15
62	A slowly progressive mitochondrial encephalomyopathy widens the spectrum of <i>AIFM1</i> disorders. Neurology, 2015, 84, 2193-2195.	1.1	47
63	Cerebrospinal Fluid Monoamine Metabolite Analysis in Pediatric Movement Disorders. Journal of Child Neurology, 2015, 30, 1800-1805.	1.4	3
64	Loss of apoptosis-inducing factor critically affects MIA40 function. Cell Death and Disease, 2015, 6, e1814-e1814.	6.3	77
65	A family with paroxysmal nonkinesigenic dyskinesias (PNKD): Evidence of mitochondrial dysfunction. European Journal of Paediatric Neurology, 2015, 19, 64-68.	1.6	13
66	Mutations in GTPBP3 Cause a Mitochondrial Translation Defect Associated with Hypertrophic Cardiomyopathy, Lactic Acidosis, and Encephalopathy. American Journal of Human Genetics, 2014, 95, 708-720.	6.2	123
67	Common and Novel TMEM70 Mutations in a Cohort of Italian Patients with Mitochondrial Encephalocardiomyopathy. JIMD Reports, 2014, 15, 71-8.	1.5	23
68	A novel mutation in TTC19 associated with isolated complex III deficiency, cerebellar hypoplasia, and bilateral basal ganglia lesions. Frontiers in Genetics, 2014, 5, 397.	2.3	17
69	Cavitating leukoencephalopathy with multiple mitochondrial dysfunction syndrome and NFU1 mutations. Frontiers in Genetics, 2014, 5, 412.	2.3	49
70	The impairment of HCCS leads to MLS syndrome by activating a non anonical cell death pathway in the brain and eyes. EMBO Molecular Medicine, 2014, 6, 849-849.	6.9	0
71	The Mitochondrial Aminoacyl tRNA Synthetases: Genes and Syndromes. International Journal of Cell Biology, 2014, 2014, 1-11.	2.5	117
72	Mutations in APOPT1, Encoding a Mitochondrial Protein, Cause Cavitating Leukoencephalopathy with Cytochrome c Oxidase Deficiency. American Journal of Human Genetics, 2014, 95, 315-325.	6.2	64

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73	<i>VARS2</i> and <i>TARS2</i> Mutations in Patients with Mitochondrial Encephalomyopathies. Human Mutation, 2014, 35, 983-989.	2.5	86
74	The isolated carboxyâ€ŧerminal domain of human mitochondrial leucylâ€ <scp>tRNA</scp> synthetase rescues the pathological phenotype of mitochondrial <scp>tRNA</scp> mutations in human cells. EMBO Molecular Medicine, 2014, 6, 169-182.	6.9	43
75	Novel (ovario) leukodystrophy related to <i>AARS2</i> mutations. Neurology, 2014, 82, 2063-2071.	1.1	172
76	Adult-onset Alexander disease, associated with a mutation in an alternative GFAP transcript, may be phenotypically modulated by a non-neutral HDAC6 variant. Orphanet Journal of Rare Diseases, 2013, 8, 66.	2.7	21
77	Mutations in FBXL4, Encoding a Mitochondrial Protein, Cause Early-Onset Mitochondrial Encephalomyopathy. American Journal of Human Genetics, 2013, 93, 482-495.	6.2	138
78	<i>MTO1</i> Mutations are Associated with Hypertrophic Cardiomyopathy and Lactic Acidosis and Cause Respiratory Chain Deficiency in Humans and Yeast. Human Mutation, 2013, 34, 1501-1509.	2.5	67
79	SURF1 deficiency causes demyelinating Charcot-Marie-Tooth disease. Neurology, 2013, 81, 1523-1530.	1.1	53
80	A Homozygous Mutation in <i><scp>LYRM</scp>7/<scp>MZM</scp>1<scp>L<scp></scp></scp></i> Associated with Early Onset Encephalopathy, Lactic Acidosis, and Severe Reduction of Mitochondrial Complex <scp>III</scp> Activity. Human Mutation, 2013, 34, 1619-1622.	2.5	60
81	The impairment of HCCS leads to MLS syndrome by activating a nonâ€canonical cell death pathway in the brain and eyes. EMBO Molecular Medicine, 2013, 5, 280-293.	6.9	33
82	Leukoencephalopathy with thalamus and brainstem involvement and high lactate â€~LTBL' caused by EARS2 mutations. Brain, 2012, 135, 1387-1394.	7.6	187
83	A novel homozygous mutation in SUCLA2 gene identified by exome sequencing. Molecular Genetics and Metabolism, 2012, 107, 403-408.	1.1	38
84	Cowchock Syndrome Is Associated with a Mutation in Apoptosis-Inducing Factor. American Journal of Human Genetics, 2012, 91, 1095-1102.	6.2	134
85	MELAS-like encephalomyopathy caused by a new pathogenic mutation in the mitochondrial DNA encoded cytochrome c oxidase subunit I. Neuromuscular Disorders, 2012, 22, 990-994.	0.6	18
86	Assembly Factors of Human Mitochondrial Respiratory Chain Complexes: Physiology and Pathophysiology. Advances in Experimental Medicine and Biology, 2012, 748, 65-106.	1.6	116
87	Mutations of the Mitochondrial-tRNA Modifier MTO1 Cause Hypertrophic Cardiomyopathy and Lactic Acidosis. American Journal of Human Genetics, 2012, 90, 1079-1087.	6.2	164
88	Infantile mitochondrial encephalopathy. Seminars in Fetal and Neonatal Medicine, 2011, 16, 205-215.	2.3	36
89	Mutations in TTC19 cause mitochondrial complex III deficiency and neurological impairment in humans and flies. Nature Genetics, 2011, 43, 259-263.	21.4	148
90	Severe X-Linked Mitochondrial Encephalomyopathy Associated with a Mutation in Apoptosis-Inducing Factor. American Journal of Human Genetics, 2010, 86, 639-649.	6.2	199

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91	Sym1, the yeast ortholog of the MPV17 human disease protein, is a stress-induced bioenergetic and morphogenetic mitochondrial modulator. Human Molecular Genetics, 2010, 19, 1098-1107.	2.9	69
92	Paroxysmal non-kinesigenic dyskinesia is caused by mutations of the MR-1 mitochondrial targeting sequence. Human Molecular Genetics, 2009, 18, 1058-1064.	2.9	70
93	SDHAF1, encoding a LYR complex-II specific assembly factor, is mutated in SDH-defective infantile leukoencephalopathy. Nature Genetics, 2009, 41, 654-656.	21.4	233
94	Myoclonus–dystonia syndrome: Clinical presentation, disease course, and genetic features in 11 families. Movement Disorders, 2008, 23, 28-34.	3.9	75
95	PINK1heterozygous rare variants: prevalence, significance and phenotypic spectrum. Human Mutation, 2008, 29, 565-565.	2.5	74
96	FASTKD2 Nonsense Mutation in an Infantile Mitochondrial Encephalomyopathy Associated with Cytochrome C Oxidase Deficiency. American Journal of Human Genetics, 2008, 83, 415-423.	6.2	107
97	Parkin analysis in early onset Parkinson's disease. Parkinsonism and Related Disorders, 2008, 14, 326-333.	2.2	42
98	Frequency and phenotypes of LRRK2 G2019S mutation in Italian patients with Parkinson's disease. Movement Disorders, 2006, 21, 1232-1235.	3.9	28
99	POLG1 in idiopathic Parkinson disease. Neurology, 2006, 67, 1698-1700.	1.1	46
100	Mitochondrial DNA haplogroup K is associated with a lower risk of Parkinson's disease in Italians. European Journal of Human Genetics, 2005, 13, 748-752.	2.8	197
101	High frequency stimulation of the subthalamic nucleus is efficacious in Parkin disease. Journal of Neurology, 2005, 252, 208-211.	3.6	40
102	GTP-cyclohydrolase I gene mutations in patients with autosomal dominant and recessive GTP-CH1 deficiency: Identification and functional characterization of four novel mutations. Journal of Inherited Metabolic Disease, 2004, 27, 455-463.	3.6	27