Leonardo Salviati

List of Publications by Year in descending order

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38742 34986 10,907 154 50 98 citations g-index h-index papers 160 160 160 15388 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Genotype–phenotype correlation in Gordon's syndrome: report of two cases carrying novel heterozygous mutations. Journal of Nephrology, 2022, 35, 859-862.	2.0	5
2	The Splicing of the Mitochondrial Calcium Uniporter Genuine Activator MICU1 Is Driven by RBFOX2 Splicing Factor during Myogenic Differentiation. International Journal of Molecular Sciences, 2022, 23, 2517.	4.1	2
3	Heterozygous frameshift variants in HNRNPA2B1 cause early-onset oculopharyngeal muscular dystrophy. Nature Communications, 2022, 13, 2306.	12.8	20
4	Distinct Phenotypic and microRNA Expression in X-Linked Charcot–Marie–Tooth Correlated with a Novel Mutation in the GJB1 Gene. , 2022, 1, 66-74.		0
5	A novel RRM2B mutation associated with mitochondrial DNA depletion syndrome. Molecular Genetics and Metabolism Reports, 2022, 32, 100887.	1.1	2
6	Two unusual cases of Gitelman's syndrome with a complex inheritance: how the phenotype can help interpret the genotype: lesson for the clinical nephrologist. Journal of Nephrology, 2021, 34, 1327-1330.	2.0	0
7	Toll-like receptor 3 pathway deficiency, herpes simplex encephalitis, and anti-NMDAR encephalitis: more questions than answers. Pediatric Research, 2021, 89, 1043-1043.	2.3	3
8	Motor axonal neuropathy associated with <scp><i>GNE</i></scp> mutations. Muscle and Nerve, 2021, 63, 396-401.	2.2	12
9	Deficit of human ornithine aminotransferase in gyrate atrophy: Molecular, cellular, and clinical aspects. Biochimica Et Biophysica Acta - Proteins and Proteomics, 2021, 1869, 140555.	2.3	22
10	Cytochrome c Defects in Human Disease. , 2021, , 191-200.		O
11	The displacement of frataxin from the mitochondrial cristae correlates with abnormal respiratory supercomplexes formation and bioenergetic defects in cells of Friedreich ataxia patients. FASEB Journal, 2021, 35, e21362.	0.5	9
12	Hybrid Minigene Assay: An Efficient Tool to Characterize mRNA Splicing Profiles of NF1 Variants. Cancers, 2021, 13, 999.	3.7	7
13	New pathogenic variants in COQ4 cause ataxia and neurodevelopmental disorder without detectable CoQ10 deficiency in muscle or skin fibroblasts. Journal of Neurology, 2021, 268, 3381-3389.	3.6	17
14	The multiple roles of coenzyme Q in cellular homeostasis and their relevance for the pathogenesis of coenzyme Q deficiency. Free Radical Biology and Medicine, 2021, 166, 277-286.	2.9	24
15	Craniosynostosis is a feature of <scp><i>CHD7</i></scp> â€related <scp>CHARGE</scp> syndrome. American Journal of Medical Genetics, Part A, 2021, 185, 2160-2163.	1.2	2
16	The pyruvate kinase activator mitapivat reduces hemolysis and improves anemia in a \hat{l}^2 -thalassemia mouse model. Journal of Clinical Investigation, 2021, 131, .	8.2	39
17	Molecular and Cellular Studies Reveal Folding Defects of Human Ornithine Aminotransferase Variants Associated With Gyrate Atrophy of the Choroid and Retina. Frontiers in Molecular Biosciences, 2021, 8, 695205.	3.5	2
18	Correlation between $\hat{l}\pm 1$ -Antitrypsin Deficiency and SARS-CoV-2 Infection: Epidemiological Data and Pathogenetic Hypotheses. Journal of Clinical Medicine, 2021, 10, 4493.	2.4	4

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19	Crosstalk between Long-Term Sublethal Oxidative Stress and Detrimental Inflammation as Potential Drivers for Age-Related Retinal Degeneration. Antioxidants, 2021, 10, 25.	5.1	11
20	The Kidney in Mitochondrial Diseases. , 2021, , 1-13.		0
21	Biallelic mutations in the TOGARAM1 gene cause a novel primary ciliopathy. Journal of Medical Genetics, 2020, 58, jmedgenet-2020-106833.	3.2	12
22	Multiple acyl-COA dehydrogenase deficiency in elderly carriers. Journal of Neurology, 2020, 267, 1414-1419.	3.6	23
23	The combined use of enzyme activity and metabolite assays as a strategy for newborn screening of mucopolysaccharidosis type I. Clinical Chemistry and Laboratory Medicine, 2020, 58, 2063-2072.	2.3	12
24	Vanillic Acid Restores Coenzyme Q Biosynthesis and ATP Production in Human Cells Lacking <i>COQ6</i> . Oxidative Medicine and Cellular Longevity, 2019, 2019, 1-11.	4.0	35
25	Implementation of Second-Tier Tests in Newborn Screening for Lysosomal Disorders in North Eastern Italy. International Journal of Neonatal Screening, 2019, 5, 24.	3.2	45
26	ADCK2 Haploinsufficiency Reduces Mitochondrial Lipid Oxidation and Causes Myopathy Associated with CoQ Deficiency. Journal of Clinical Medicine, 2019, 8, 1374.	2.4	27
27	Photosensitive epilepsy and long QT: expanding Timothy syndrome phenotype. Clinical Neurophysiology, 2019, 130, 2134-2136.	1.5	6
28	DRP1-mediated mitochondrial shape controls calcium homeostasis and muscle mass. Nature Communications, 2019, 10, 2576.	12.8	274
29	FSHD1 and FSHD2 form a disease continuum. Neurology, 2019, 92, e2273-e2285.	1.1	50
30	Vitamin K2 cannot substitute Coenzyme Q10 as electron carrier in the mitochondrial respiratory chain of mammalian cells. Scientific Reports, 2019, 9, 6553.	3.3	18
31	The Arg1038Gly missense variant in the <i>NF1</i> gene causes a mild phenotype without neurofibromas. Molecular Genetics & Enomic Medicine, 2019, 7, e616.	1.2	26
32	Dominant Noonan syndrome-causing <i>LZTR1</i> mutations specifically affect the Kelch domain substrate-recognition surface and enhance RAS-MAPK signaling. Human Molecular Genetics, 2019, 28, 1007-1022.	2.9	58
33	A new Italian family with HTRA1 mutation associated with autosomal-dominant variant of CARASIL: Are we pointing towards a disease spectrum?. Journal of the Neurological Sciences, 2019, 396, 108-111.	0.6	11
34	Sarcopenia: Aging-Related Loss of Muscle Mass and Function. Physiological Reviews, 2019, 99, 427-511.	28.8	767
35	The idebenone metabolite QS10 restores electron transfer in complex I and coenzyme Q defects. Biochimica Et Biophysica Acta - Bioenergetics, 2018, 1859, 901-908.	1.0	31
36	COX16 is required for assembly of cytochrome c oxidase in human cells and is involved in copper delivery to COX2. Biochimica Et Biophysica Acta - Bioenergetics, 2018, 1859, 244-252.	1.0	25

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37	In mammalian skeletal muscle, phosphorylation of TOMM22 by protein kinase CSNK2/CK2 controls mitophagy. Autophagy, 2018, 14, 311-335.	9.1	51
38	Prenatal detection of trisomy 8 mosaicism: Pregnancy outcome and follow up of a series of 17 consecutive cases. European Journal of Obstetrics, Gynecology and Reproductive Biology, 2018, 221, 23-27.	1.1	15
39	Mutations in COQ8B (ADCK4) found in patients with steroid-resistant nephrotic syndrome alter COQ8B function. Human Mutation, 2018, 39, 406-414.	2.5	43
40	Newborn screening for lysosomal storage disorders by tandem mass spectrometry in North East Italy. Journal of Inherited Metabolic Disease, 2018, 41, 209-219.	3.6	114
41	Drug Repurposing for Duchenne Muscular Dystrophy: The Monoamine Oxidase B Inhibitor Safinamide Ameliorates the Pathological Phenotype in mdx Mice and in Myogenic Cultures From DMD Patients. Frontiers in Physiology, 2018, 9, 1087.	2.8	11
42	Molecular and cellular basis of ornithine $\hat{\Gamma}$ -aminotransferase deficiency caused by the V332M mutation associated with gyrate atrophy of the choroid and retina. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2018, 1864, 3629-3638.	3.8	12
43	Molecular diagnosis of coenzyme Q ₁₀ deficiency: an update. Expert Review of Molecular Diagnostics, 2018, 18, 491-498.	3.1	33
44	Transcriptional programming of lipid and amino acid metabolism by the skeletal muscle circadian clock. PLoS Biology, 2018, 16, e2005886.	5.6	107
45	Increased mitophagy in the skeletal muscle of spinal and bulbar muscular atrophy patients. Human Molecular Genetics, 2017, 26, ddx019.	2.9	37
46	MCM5: a new actor in the link between DNA replication and Meier-Gorlin syndrome. European Journal of Human Genetics, 2017, 25, 646-650.	2.8	60
47	Age-Associated Loss of OPA1 in Muscle Impacts Muscle Mass, Metabolic Homeostasis, Systemic Inflammation, and Epithelial Senescence. Cell Metabolism, 2017, 25, 1374-1389.e6.	16.2	388
48	Mitochondrial cytopathies and the kidney. Nephrologie Et Therapeutique, 2017, 13, S23-S28.	0.5	31
49	A synonymous splicing mutation in the SF3B4 gene segregates in a family with highly variable Nager syndrome. European Journal of Human Genetics, 2017, 25, 371-375.	2.8	20
50	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
51	Secondary coenzyme Q 10 deficiencies in oxidative phosphorylation (OXPHOS) and non-OXPHOS disorders. Mitochondrion, 2016, 30, 51-58.	3.4	70
52	Coenzyme Q biosynthesis in health and disease. Biochimica Et Biophysica Acta - Bioenergetics, 2016, 1857, 1079-1085.	1.0	185
53	Exome sequencing analysis in a pair of monozygotic twins re-evaluates the genetics behind their intellectual disability and reveals a CHD2 mutation. Brain and Development, 2016, 38, 590-596.	1.1	11
54	The <i>COQ2</i> genotype predicts the severity of coenzyme Q ₁₀ deficiency. Human Molecular Genetics, 2016, 25, 4256-4265.	2.9	53

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55	The chaperone co-inducer BGP-15 alleviates ventilation-induced diaphragm dysfunction. Science Translational Medicine, 2016, 8, 350ra103.	12.4	53
56	Severe encephalopathy associated to pyruvate dehydrogenase mutations and unbalanced coenzyme Q10 content. European Journal of Human Genetics, 2016, 24, 367-372.	2.8	17
57	Mitochondrial dysfunction in inherited renal disease and acute kidney injury. Nature Reviews Nephrology, 2016, 12, 267-280.	9.6	276
58	<i>ALDH18A1</i> gene mutations cause dominant spastic paraplegia SPG9: loss of function effect and plausibility of a dominant negative mechanism. Brain, 2016, 139, e3-e3.	7.6	42
59	Clinical and molecular study in a long-surviving patient with MLASA syndrome due to novel PUS1 mutations. Neurogenetics, 2016, 17, 65-70.	1.4	29
60	<i>FBXO28</i> is a critical gene of the 1q41q42 microdeletion syndrome. American Journal of Medical Genetics, Part A, 2015, 167, 1418-1420.	1.2	15
61	Haploinsufficiency of the NOTCH1 Receptor as a Cause of Adams–Oliver Syndrome With Variable Cardiac Anomalies. Circulation: Cardiovascular Genetics, 2015, 8, 572-581.	5.1	84
62	Is there a link between COQ6 and schwannomatosis?. Genetics in Medicine, 2015, 17, 312-313.	2.4	7
63	Loss-of-function variants of SETD5 cause intellectual disability and the core phenotype of microdeletion 3p25.3 syndrome. European Journal of Human Genetics, 2015, 23, 753-760.	2.8	73
64	Primary coenzyme Q10 deficiency presenting as fatal neonatal multiorgan failure. European Journal of Human Genetics, 2015, 23, 1254-1258.	2.8	42
65	Validation of CFTR intronic variants identified during cystic fibrosis population screening by a minigene splicing assay. Clinical Chemistry and Laboratory Medicine, 2015, 53, 1719-23.	2.3	12
66	Molecular diagnosis of coenzyme Q ₁₀ deficiency. Expert Review of Molecular Diagnostics, 2015, 15, 1049-1059.	3.1	16
67	Ichthyosis and Kallmann syndrome: Not always a contiguous gene syndrome. Journal of Dermatological Science, 2015, 78, 158-160.	1.9	4
68	Characterization of 14 novel deletions underlying Rubinstein–Taybi syndrome: an update of the CREBBP deletion repertoire. Human Genetics, 2015, 134, 613-626.	3.8	38
69	Heterologous Expression in Yeast of Human Ornithine Carriers ORNT1 and ORNT2 and of ORNT1 Alleles Implicated in HHH Syndrome in Humans. JIMD Reports, 2015, 28, 119-126.	1.5	4
70	Genetic bases and clinical manifestations of coenzyme Q ₁₀ (CoQ ₁₀) deficiency. Journal of Inherited Metabolic Disease, 2015, 38, 145-156.	3.6	301
71	Facioscapulohumeral muscular dystrophy. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2015, 1852, 607-614.	3.8	47
72	Genetics of Coenzyme Q10 Deficiency. Molecular Syndromology, 2014, 5, 156-162.	0.8	102

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73	Impaired copper and iron metabolism in blood cells and muscles of patients affected by copper deficiency myeloneuropathy. Neuropathology and Applied Neurobiology, 2014, 40, 888-898.	3.2	12
74	Neurofibromatosis type 1 in two siblings due to maternal germline mosaicism. Clinical Genetics, 2014, 85, 386-389.	2.0	18
75	Molecular characterization of the human COQ5 C-methyltransferase in coenzyme Q10 biosynthesis. Biochimica Et Biophysica Acta - Molecular and Cell Biology of Lipids, 2014, 1841, 1628-1638.	2.4	48
76	Atrio-ventricular block requiring pacemaker in patients with late onset Pompe disease. Neuromuscular Disorders, 2014, 24, 648-650.	0.6	15
77	Mutations of cytochrome c identified in patients with thrombocytopenia THC4 affect both apoptosis and cellular bioenergetics. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2014, 1842, 269-274.	3.8	65
78	Molecular Analysis, Pathogenic Mechanisms, and Readthrough Therapy on a Large Cohort of <scp>K</scp> abuki Syndrome Patients. Human Mutation, 2014, 35, 841-850.	2.5	87
79	Complexity of the 5′UTR region of the CLCN5gene: eleven 5′UTR ends are differentially expressed in the human kidney. BMC Medical Genomics, 2014, 7, 41.	1.5	8
80	Effect of vanillic acid on COQ6 mutants identified in patients with coenzyme Q10 deficiency. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2014, 1842, 1-6.	3.8	64
81	Coenzyme Q10 deficiency in mitochondrial DNA depletion syndromes. Mitochondrion, 2013, 13, 337-341.	3.4	51
82	Mitochondrial Cristae Shape Determines Respiratory Chain Supercomplexes Assembly and Respiratory Efficiency. Cell, 2013, 155, 160-171.	28.9	955
83	The FSHD2 Gene SMCHD1 is a Modifier of Disease Severity in Families Affected by FSHD1. American Journal of Human Genetics, 2013, 93, 744-751.	6.2	154
84	Pontocerebellar hypoplasia type 6 caused by mutations in <i>RARS2</i> : definition of the clinical spectrum and molecular findings in five patients. Journal of Inherited Metabolic Disease, 2013, 36, 43-53.	3.6	70
85	Functional Analysis of Missense Mutations of <i>OAT </i> , Causing Gyrate Atrophy of Choroid and Retina. Human Mutation, 2013, 34, 229-236.	2.5	23
86	Whole exome sequencing in dominant cataract identifies a new causative factor, CRYBA2, and a variety of novel alleles in known genes. Human Genetics, 2013, 132, 761-770.	3.8	72
87	Recessive MYL2 mutations cause infantile type I muscle fibre disease and cardiomyopathy. Brain, 2013, 136, 282-293.	7.6	48
88	Survival transcriptome in the coenzyme Q $<$ sub $>$ 10 $<$ /sub $>$ deficiency syndrome is acquired by epigenetic modifications: a modelling study for human coenzyme Q $<$ sub $>$ 10 $<$ /sub $>$ deficiencies. BMJ Open, 2013, 3, e002524.	1.9	19
89	ADCK4 mutations promote steroid-resistant nephrotic syndrome through CoQ10 biosynthesis disruption. Journal of Clinical Investigation, 2013, 123, 5179-5189.	8.2	275
90	Haploinsufficiency of <i>COQ4</i> causes coenzyme Q ₁₀ deficiency. Journal of Medical Genetics, 2012, 49, 187-191.	3.2	95

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91	Patients with a phenotype consistent with facioscapulohumeral muscular dystrophy display genetic and epigenetic heterogeneity. Journal of Medical Genetics, 2012, 49, 41-46.	3.2	55
92	A novel CRYAB mutation resulting in multisystemic disease. Neuromuscular Disorders, 2012, 22, 66-72.	0.6	84
93	Copper and bezafibrate cooperate to rescue cytochrome c oxidase deficiency in cells of patients with sco2 mutations. Orphanet Journal of Rare Diseases, 2012, 7, 21.	2.7	29
94	Assessment of mitochondrial respiratory chain enzymatic activities on tissues and cultured cells. Nature Protocols, 2012, 7, 1235-1246.	12.0	732
95	Yeast complementation is sufficiently sensitive to detect the residual activity of ASL alleles associated with mild forms of argininosuccinic aciduria. Journal of Inherited Metabolic Disease, 2012, 35, 557-558.	3.6	4
96	Nitric-Oxide Supplementation for Treatment of Long-Term Complications in Argininosuccinic Aciduria. American Journal of Human Genetics, 2012, 90, 836-846.	6.2	73
97	Genetic susceptibility to teratogens: State of the art. Reproductive Toxicology, 2012, 34, 186-191.	2.9	27
98	Renal involvement in mitochondrial cytopathies. Pediatric Nephrology, 2012, 27, 539-550.	1.7	109
99	Optimization of respiratory chain enzymatic assays in muscle for the diagnosis of mitochondrial disorders. Mitochondrion, 2011, 11, 893-904.	3.4	50
100	Coenzyme Q deficiency in muscle. Current Opinion in Neurology, 2011, 24, 449-456.	3.6	73
101	Challenges in diagnosis and treatment of late-onset Pompe disease. Current Opinion in Neurology, 2011, 24, 443-448.	3.6	11
102	Alport syndrome and leiomyomatosis: the first deletion extending beyond COL4A6 intron 2. Pediatric Nephrology, 2011, 26, 717-724.	1.7	27
103	Polymorphisms of the SCN1A gene in children and adolescents with primary headache and idiopathic or cryptogenic epilepsy: is there a linkage?. Journal of Headache and Pain, 2011, 12, 435-441.	6.0	5
104	6q27 subtelomeric deletions: Is there a specific phenotype?. American Journal of Medical Genetics, Part A, 2011, 155, 1213-1214.	1.2	6
105	Renal Mitochondrial Cytopathies. International Journal of Nephrology, 2011, 2011, 1-10.	1.3	27
106	COQ6 mutations in human patients produce nephrotic syndrome with sensorineural deafness. Journal of Clinical Investigation, 2011, 121, 2013-2024.	8.2	343
107	Abnormalities of cerebral arteries are frequent in patients with late-onset Pompe disease. Journal of Neurology, 2010, 257, 1730-1733.	3.6	68
108	Comorbidity between headache and epilepsy in a pediatric headache center. Journal of Headache and Pain, 2010, 11, 235-240.	6.0	77

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109	Treatment of CoQ10 Deficient Fibroblasts with Ubiquinone, CoQ Analogs, and Vitamin C: Time- and Compound-Dependent Effects. PLoS ONE, 2010, 5, e11897.	2.5	92
110	The Conserved Mitochondrial Twin Cx9C Protein Cmc2 Is a Cmc1 Homologue Essential for Cytochrome c Oxidase Biogenesis. Journal of Biological Chemistry, 2010, 285, 15088-15099.	3.4	34
111	Prevalence of AIP mutations in a large series of sporadic Italian acromegalic patients and evaluation of CDKN1B status in acromegalic patients with multiple endocrine neoplasia. European Journal of Endocrinology, 2010, 163, 369-376.	3.7	53
112	Reactive oxygen species, oxidative stress, and cell death correlate with level of CoQ ₁₀ deficiency. FASEB Journal, 2010, 24, 3733-3743.	0.5	142
113	Dandy-Walker Malformation Masking the Molar Tooth Sign: An Illustrative Case With Magnetic Resonance Imaging Follow-up. Journal of Child Neurology, 2010, 25, 1419-1422.	1.4	14
114	Is CFTR 621+3 A>G a cystic fibrosis causing mutation?. Journal of Human Genetics, 2010, 55, 23-26.	2.3	14
115	Novel mutations in the L1CAM gene support the complexity of L1 syndrome. Journal of the Neurological Sciences, 2010, 294, 124-126.	0.6	10
116	Coenzyme Q10 is frequently reduced in muscle of patients with mitochondrial myopathy. Neuromuscular Disorders, 2010, 20, 44-48.	0.6	84
117	Mutation analysis of COX18 in 29 patients with isolated cytochrome c oxidase deficiency. Journal of Human Genetics, 2009, 54, 419-421.	2.3	7
118	Coenzyme Q deficiency triggers mitochondria degradation by mitophagy. Autophagy, 2009, 5, 19-32.	9.1	179
119	Functional Complementation in Yeast Allows Molecular Characterization of Missense Argininosuccinate Lyase Mutations. Journal of Biological Chemistry, 2009, 284, 28926-28934.	3.4	30
120	Xâ€linked brachytelephalangic chondrodysplasia punctata: A simple trait that is not so simple. American Journal of Medical Genetics, Part A, 2009, 149A, 2464-2468.	1.2	10
121	Coenzyme Q10 distribution in blood is altered in patients with Fibromyalgia. Clinical Biochemistry, 2009, 42, 732-735.	1.9	60
122	Coenzyme Q10 Deficiencies in Neuromuscular Diseases. Advances in Experimental Medicine and Biology, 2009, 652, 117-128.	1.6	21
123	Analysis of Coenzyme Q10 in muscle and fibroblasts for the diagnosis of CoQ10 deficiency syndromes. Clinical Biochemistry, 2008, 41, 697-700.	1.9	65
124	Increased level of N-acetylaspartylglutamate (NAAG) in the CSF of a patient with Pelizaeus-Merzbacher-like disease due to mutation in the GJA12 gene. European Journal of Paediatric Neurology, 2008, 12, 348-350.	1.6	16
125	Early Coenzyme Q10 Supplementation in Primary Coenzyme Q10 Deficiency. New England Journal of Medicine, 2008, 358, 2849-2850.	27.0	217
126	Functional characterization of human COQ4, a gene required for Coenzyme Q10 biosynthesis. Biochemical and Biophysical Research Communications, 2008, 372, 35-39.	2.1	49

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127	Respiratory chain dysfunction and oxidative stress correlate with severity of primary CoQ ₁₀ deficiency. FASEB Journal, 2008, 22, 1874-1885.	0.5	150
128	A novel deletion in the GTPase domain of OPA1 causes defects in mitochondrial morphology and distribution, but not in function. Human Molecular Genetics, 2008, 17, 3291-3302.	2.9	91
129	A functionally dominant mitochondrial DNA mutation. Human Molecular Genetics, 2008, 17, 1814-1820.	2.9	104
130	Renal hypoplasia without optic coloboma associated with PAX2 gene deletion. Nephrology Dialysis Transplantation, 2007, 22, 2076-2078.	0.7	25
131	LETM1, deleted in Wolf Hirschhorn syndrome is required for normal mitochondrial morphology and cellular viability. Human Molecular Genetics, 2007, 17, 201-214.	2.9	163
132	Missense mutation of the COQ2 gene causes defects of bioenergetics and de novo pyrimidine synthesis. Human Molecular Genetics, 2007, 16, 1091-1097.	2.9	129
133	High frequency of mosaic CREBBP deletions in Rubinstein–Taybi syndrome patients and mapping of somatic and germ-line breakpoints. Genomics, 2007, 90, 567-573.	2.9	42
134	Glycogen synthase binds to sarcoplasmic reticulum and is phosphorylated by CaMKII in fast-twitch skeletal muscle. Archives of Biochemistry and Biophysics, 2007, 459, 115-121.	3.0	18
135	Argininosuccinate lyase deficiency: mutational spectrum in Italian patients and identification of a novelASLpseudogene. Human Mutation, 2007, 28, 694-702.	2.5	46
136	A novel deletion in the GJA12 gene causes Pelizaeus–Merzbacher-like disease. Neurogenetics, 2007, 8, 57-60.	1.4	42
137	A Mutation in Para-Hydroxybenzoate-Polyprenyl Transferase (COQ2) Causes Primary Coenzyme Q10 Deficiency. American Journal of Human Genetics, 2006, 78, 345-349.	6.2	322
138	Deletion of PTEN and BMPR1A on Chromosome 10q23 Is Not Always Associated with Juvenile Polyposis of Infancy. American Journal of Human Genetics, 2006, 79, 593-596.	6.2	40
139	Molecular analysis of two uncharacterized sequence variants of the VHL gene. Journal of Human Genetics, 2006, 51, 964-968.	2.3	17
140	hCOX18 and hCOX19: Two human genes involved in cytochrome c oxidase assembly. Biochemical and Biophysical Research Communications, 2005, 337, 832-839.	2.1	28
141	Post-natal developmental expression of \hat{l} ±KAP splice variants in rabbit fast-twitch and slow-twitch skeletal muscle. Journal of Muscle Research and Cell Motility, 2004, 25, 309-314.	2.0	0
142	NovelSURF1 mutation in a child with subacute encephalopathy and without the radiological features of Leigh Syndrome. American Journal of Medical Genetics Part A, 2004, 128A, 195-198.	2.4	29
143	Hepatocerebral Mitochondrial DNA Depletion Syndrome: Clinical and Morphologic Features of a Nuclear Gene Mutation. Journal of Pediatric Gastroenterology and Nutrition, 2004, 38, 216-220.	1.8	25
144	Two splice variants of CaMKII-anchoring protein are present in the sarcoplasmic reticulum of rabbit fast-twitch muscle. Biochemical and Biophysical Research Communications, 2003, 302, 73-83.	2.1	11

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145	Mutation Screening in Patients With Isolated Cytochrome c Oxidase Deficiency. Pediatric Research, 2003, 53, 224-230.	2.3	62
146	Mutation Screening in Patients With Isolated Cytochrome c Oxidase Deficiency. Pediatric Research, 2003, 53, 224-230.	2.3	44
147	Cytochrome c Oxidase Deficiency Due to a Novel SCO2 Mutation Mimics Werdnig-Hoffmann Disease. Archives of Neurology, 2002, 59, 862-5.	4.5	95
148	Complex Neurologic Syndrome Associated With the G1606A Mutation of Mitochondrial DNA. Archives of Neurology, 2002, 59, 1013.	4.5	32
149	Copper supplementation restores cytochrome c oxidase activity in cultured cells from patients with SCO2 mutations. Biochemical Journal, 2002, 363, 321.	3.7	55
150	Copper supplementation restores cytochrome c oxidase activity in cultured cells from patients with SCO2 mutations. Biochemical Journal, 2002, 363, 321-327.	3.7	66
151	Mitochondrial myopathy and ophthalmoplegia in a sporadic patient with the G12315A mutation in mitochondrial DNA. Neuromuscular Disorders, 2002, 12, 865-868.	0.6	29
152	Mitochondrial DNA depletion and <i>dGK</i> gene mutations. Annals of Neurology, 2002, 52, 311-317.	5.3	152
153	Acute Disseminated Encephalomyelitis Associated With Hepatitis C Virus Infection. Archives of Neurology, 2001, 58, 1679.	4.5	57
154	Acute Quadriplegic Myopathy in a 17-Month-Old Boy. Journal of Child Neurology, 2000, 15, 63-66.	1.4	13