## Ratna Dua Puri

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/8471188/publications.pdf

Version: 2024-02-01

41 papers

280 citations

1040056 9 h-index 14 g-index

42 all docs

42 docs citations

42 times ranked 504 citing authors

#	Article	IF	CITATIONS
1	<i>GALNS</i> mutations in Indian patients with mucopolysaccharidosis IVA. American Journal of Medical Genetics, Part A, 2014, 164, 2793-2801.	1.2	31
2	A data set of variants derived from 1455 clinical and research exomes is efficient in variant prioritization for earlyâ€onset monogenic disorders in Indians. Human Mutation, 2021, 42, e15-e61.	2.5	25
3	Novel and recurrent mutations in <i>WISP3</i> and an atypical phenotype. American Journal of Medical Genetics, Part A, 2015, 167, 2481-2484.	1.2	21
4	Do parental perceptions and motivations towards genetic testing and prenatal diagnosis for deafness vary in different cultures?. American Journal of Medical Genetics, Part A, 2013, 161, 76-81.	1.2	18
5	Molecular Diagnosis of Hereditary Fructose Intolerance: Founder Mutation in a Community from India. JIMD Reports, 2014, 19, 85-93.	1.5	18
6	Spectrum of severe skeletal dysplasias in North India. Indian Journal of Pediatrics, 2007, 74, 995-1002.	0.8	15
7	Clinical heterogeneity and molecular profile of triple A syndrome: a study of seven cases. Journal of Pediatric Endocrinology and Metabolism, 2018, 31, 799-807.	0.9	13
8	The first case of antenatal presentation in COG8â€congenital disorder of glycosylation with a novel splice site mutation and an extended phenotype. American Journal of Medical Genetics, Part A, 2019, 179, 480-485.	1.2	12
9	Clinical and genetic spectrum of 104 Indian families with central nervous system white matter abnormalities. Clinical Genetics, 2021, 100, 542-550.	2.0	12
10	<i>KCNQ1</i> mutations associated with Jervell and Lange–Nielsen syndrome and autosomal recessive Romano–Ward syndrome in India—expanding the spectrum of long QT syndrome type 1. American Journal of Medical Genetics, Part A, 2016, 170, 1510-1519.	1.2	11
11	Phenotype guided characterization and molecular analysis of Indian patients with long QT syndromes. Indian Pacing and Electrophysiology Journal, 2016, 16, 8-18.	0.6	11
12	Expanding the Phenotype of the Founder South Asian Mutation in the Nuclear Encoding Mitochondrial RMND1 Gene. Indian Journal of Pediatrics, 2018, 85, 87-92.	0.8	9
13	NGS-based expanded carrier screening for genetic disorders in North Indian population reveals unexpected results – a pilot study. BMC Medical Genetics, 2020, 21, 216.	2.1	9
14	Expanding the phenotypic and genotypic spectrum of Wiedemann–Steiner syndrome: First patient from India. American Journal of Medical Genetics, Part A, 2020, 182, 953-956.	1.2	9
15	Genetic Approach to Diagnosis of Intellectual Disability. Indian Journal of Pediatrics, 2016, 83, 1141-1149.	0.8	8
16	Sialidosis type II: Expansion of phenotypic spectrum and identification of a common mutation in seven patients. Molecular Genetics and Metabolism Reports, 2020, 22, 100561.	1.1	8
17	Lysosomal storage disorders: Novel and frequent pathogenic variants in a large cohort of Indian patients of Pompe, Fabry, Gaucher and Hurler disease. Clinical Biochemistry, 2021, 89, 14-37.	1.9	6
18	Late onset Pompe Disease in India – Beyond the Caucasian phenotype. Neuromuscular Disorders, 2021, 31, 431-441.	0.6	6

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19	Prenatal presentation of a rare genetic disorder: a clinical, autopsy and molecular correlation. Autopsy and Case Reports, 2019, 9, e2019124.	0.6	6
20	Hotspots in PTPN11 gene among Indian children with Noonan syndrome. Indian Pediatrics, 2017, 54, 638-640.	0.4	5
21	Challenges in Chronic Genetic Disorders: Lessons From the COVID-19 Pandemic. Indian Pediatrics, 2021, 58, 391-392.	0.4	4
22	Is the diagnostic yield influenced by the indication for fetal autopsy?. American Journal of Medical Genetics, Part A, 2016, 170, 2119-2126.	1.2	3
23	Next Generation Sequencing in the Clinic. Indian Journal of Pediatrics, 2016, 83, 281-282.	0.8	3
24	Genotype-phenotype spectrum of 130 unrelated Indian families with Mucopolysaccharidosis type II. European Journal of Medical Genetics, 2022, 65, 104447.	1.3	3
25	The Role of Radiographs in Fetal Autopsy. Journal of Fetal Medicine, 2014, 1, 7-9.	0.1	2
26	Fetal Dysmorphology. Journal of Fetal Medicine, 2015, 2, 151-159.	0.1	2
27	<code><scp> <i>COASY</i> </scp> related pontocerebellar hypoplasia type 12: A common Indian mutation with expansion of the phenotypic spectrum. American Journal of Medical Genetics, Part A, 2022, , .</code>	1.2	2
28	Prenatal Diagnosis and Elucidation of a Novel Molecular Mechanism in Carpenter Syndrome. Journal of Fetal Medicine, 2014, 1, 89-93.	0.1	1
29	Editorial: New Horizons in Genetic Diagnosis in Pediatric Practice: The Excitement and Challenges!. Indian Journal of Pediatrics, 2016, 83, 1131-1132.	0.8	1
30	ALG9 Associated Gillessen-Kaesbach–Nishimura Syndrome (GIKANIS): An Uncommon Aetiology of Enlarged Foetal Kidneys. Journal of Fetal Medicine, 2018, 05, 237-239.	0.1	1
31	Gaucher Disease in Fetus: The Usual and the Unusual Presentations in a Family. Journal of Fetal Medicine, 2019, 6, 147-150.	0.1	1
32	Mutation and Phenotypic Spectrum of Patients With RASopathies. Indian Pediatrics, 2021, 58, 30-33.	0.4	1
33	Mutation and Phenotypic Spectrum of Patients With RASopathies. Indian Pediatrics, 2021, 58, 30-33.	0.4	1
34	Prenatal phenotypic spectrum of full trisomy 18 in an Indian cohort. American Journal of Medical Genetics, Part A, 2022, , .	1.2	1
35	Indian Undiagnosed Diseases Program (I-UDP) — The Unmet Need. Indian Pediatrics, 2022, 59, 198-200.	0.4	1
36	Is Molecular Diagnosis Necessary for Children with Duchenne Muscular Dystrophy?. Indian Pediatrics, 2019, 56, 549-550.	0.4	0

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37	Filaggrin Null-Mutation in Asthma in an Indian Cohort: One Link in a Polygenic Trait. Indian Journal of Pediatrics, 2020, 87, 583-584.	0.8	O
38	Levels of Lyso GL-1 in Gaucher and Lyso GL-3 in Fabry patients from India: Diagnostic aids for these lysosomal storage disorders. Clinica Chimica Acta, 2021, 521, 177-190.	1.1	0
39	Is Molecular Diagnosis Necessary for Children with Duchenne Muscular Dystrophy?. Indian Pediatrics, 2019, 56, 549-550.	0.4	O
40	Mutation and Phenotypic Spectrum of Patients With RASopathies. Indian Pediatrics, 2020, , .	0.4	0
41	Indian Undiagnosed Diseases Program (I-UDP) - The Unmet Need Indian Pediatrics, 2022, , .	0.4	0