## Arti Pandya

## List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/6649061/publications.pdf

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713332 623574 1,372 25 14 21 citations g-index h-index papers 25 25 25 1540 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Prevalence and Evolutionary Origins of the del(GJB6-D13S1830) Mutation in the DFNB1 Locus in Hearing-Impaired Subjects: a Multicenter Study. American Journal of Human Genetics, 2003, 73, 1452-1458.	2.6	269
2	American College of Medical Genetics and Genomics guideline for the clinical evaluation and etiologic diagnosis of hearing loss. Genetics in Medicine, 2014, 16, 347-355.	1.1	207
3	Frequency and distribution of GJB2 (connexin 26) and GJB6 (connexin 30) mutations in a large North American repository of deaf probands. Genetics in Medicine, 2003, 5, 295-303.	1.1	138
4	Advances in hereditary deafness. Lancet, The, 2001, 358, 1082-1090.	6.3	136
5	Heterogenous Point Mutations in the Mitochondrial tRNA Ser(UCN) Precursor Coexisting with the A1555G Mutation in Deaf Students from Mongolia. American Journal of Human Genetics, 1999, 65, 1803-1806.	2.6	78
6	Coffin-Siris syndrome: Review and presentation of new cases from a questionnaire study. American Journal of Medical Genetics Part A, 2001, 99, 1-7.	2.4	71
7	Correlation between Waardenburg syndrome phenotype and genotype in a population of individuals with identified PAX3 mutations. Human Genetics, 1998, 102, 499-506.	1.8	69
8	Does Universal Newborn Hearing Screening Identify All Children with GJB2 (Connexin 26) Deafness? Penetrance of GJB2 Deafness. Ear and Hearing, 2006, 27, 732-741.	1.0	69
9	Natural history and genotypeâ€phenotype correlations in 72 individuals with <i>SATB2</i> â€essociated syndrome. American Journal of Medical Genetics, Part A, 2018, 176, 925-935.	0.7	57
10	Phenotypic variation in Waardenburg syndrome: mutational heterogeneity, modifier genes or polygenic background?. Human Molecular Genetics, 1996, 5, 497-502.	1.4	39
11	Longitudinal assessment of adaptive and maladaptive behaviors in fragile X males: Growth, development, and profiles., 1999, 83, 257-263.		38
12	Fine mapping of a gene responsible for regulating dietary cholesterol absorption; founder effects underlie cases of phytosterolaemia in multiple communities. European Journal of Human Genetics, 2001, 9, 375-384.	1.4	38
13	A Comparative Analysis of the Genetic Epidemiology of Deafness in the United States in Two Sets of Pedigrees Collected More than a Century Apart. American Journal of Human Genetics, 2008, 83, 200-207.	2.6	34
14	Age-related language characteristics of children and adolescents with fragile X syndrome. American Journal of Medical Genetics Part A, 1999, 83, 253-256.	2.4	29
15	Cancer Risk in Klippel–Trenaunay Syndrome. Lymphatic Research and Biology, 2019, 17, 630-636.	0.5	20
16	Clinical evaluation and etiologic diagnosis of hearing loss: A clinical practice resource of the American College of Medical Genetics and Genomics (ACMG). Genetics in Medicine, 2022, 24, 1392-1406.	1.1	18
17	Genetic hearing loss: the journey of discovery to destination – how close are we to therapy?. Molecular Genetics & Genomic Medicine, 2016, 4, 583-587.	0.6	15
18	Analysis of risk factors associated with unilateral hearing loss in children who initially passed newborn hearing screening. International Journal of Pediatric Otorhinolaryngology, 2018, 106, 100-104.	0.4	15

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#	Article	IF	CITATIONS
19	Analyses of del(GJB6â€D13S1830) and del(GJB6â€D13S1834) deletions in a large cohort with hearing loss: Caveats to interpretation of molecular test results in multiplex families. Molecular Genetics & Description (Genomic Medicine, 2020, 8, e1171.	0.6	10
20	Genetic Epidemiology of Deafness. , 2002, , 67-91.		6
21	Genetic Evaluation and Counseling in the Context of Early Hearing Detection and Intervention. Seminars in Hearing, 2006, 27, 205-212.	0.5	5
22	Confirmation of COL4A6 variants in X-linked nonsyndromic hearing loss and its clinical implications. European Journal of Human Genetics, 2021, , .	1.4	5
23	Oculodentodigital Dysplasia: A Case Report and Major Review of the Eye and Ocular Adnexa Features of 295 Reported Cases. Case Reports in Ophthalmological Medicine, 2020, 2020, 1-16.	0.3	4
24	Co-occurring Down syndrome and SUCLA2 -related mitochondrial depletion syndrome. , 2017, 173, 2720-2724.		2
25	Cover Image, Volume 176A, Number 4, April 2018. American Journal of Medical Genetics, Part A, 2018, 176,	0.7	0