Kiyomitsu Nara

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Phenotype–genotype correlation in patients with typical and atypical branchio-oto-renal syndrome. Scientific Reports, 2022, 12, 969.	3.3	6
2	Whole exome analysis of patients in Japan with hearing loss reveals high heterogeneity among responsible and novel candidate genes. Orphanet Journal of Rare Diseases, 2022, 17, 114.	2.7	3
3	Investigation of the hearing levels of siblings affected by a single GJB2 variant: Possibility of genetic modifiers. International Journal of Pediatric Otorhinolaryngology, 2021, 149, 110840.	1.0	2
4	Clinical and genetic analysis of children with hearing loss and bilateral enlarged vestibular aqueducts. International Journal of Pediatric Otorhinolaryngology, 2021, 152, 110975.	1.0	0
5	Differences in hearing levels between siblings with hearing loss caused by GJB2 mutations. Auris Nasus Larynx, 2020, 47, 938-942.	1.2	2
6	Variants encoding a restricted carboxy-terminal domain of SLC12A2 cause hereditary hearing loss in humans. PLoS Genetics, 2020, 16, e1008643.	3.5	36
7	ClinGen expert clinical validity curation of 164 hearing loss gene–disease pairs. Genetics in Medicine, 2019, 21, 2239-2247.	2.4	67
8	Expert specification of the ACMG/AMP variant interpretation guidelines for genetic hearing loss. Human Mutation, 2018, 39, 1593-1613.	2.5	312