Markéta HavlovicovÃ;

List of Publications by Year in descending order

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1937685 1872680 8 58 4 6 citations h-index g-index papers 8 8 8 84 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	<scp>Zimmermann–Laband</scp> syndrome in monozygotic twins with a mild neurobehavioral phenotype lacking gingival overgrowth—A case report of a novel <scp> <i>KCNN3</i> </scp> gene variant. American Journal of Medical Genetics, Part A, 2022, 188, 1083-1087.	1.2	2
2	Co-designing models for the communication of genomic results for rare diseases: a comparative study in the Czech Republic and the United Kingdom. Journal of Community Genetics, 2022, 13, 313-327.	1.2	3
3	Expanding the phenotype spectrum associated with pathogenic variants in the <i>COL2A1</i> and <i>COL11A1</i> genes. Annals of Human Genetics, 2020, 84, 380-392.	0.8	11
4	The Key Role of Purine Metabolism in the Folate-Dependent Phenotype of Autism Spectrum Disorders: An In Silico Analysis. Metabolites, 2020, 10, 184.	2.9	7
5	Assisted Reproductive Techniques and Pregnancy Results in Women with Mayer-Rokitansky-Kýster-Hauser Syndrome Undergoing Uterus Transplantation: the Czech Experience. Journal of Pediatric and Adolescent Gynecology, 2020, 33, 410-414.	0.7	18
6	Schinzelâ€"Giedion Syndrome: First Czech Patients Confirmed by Molecular Genetic Analysis. Journal of Pediatric Neurology, 2019, 17, 125-127.	0.2	0
7	Modeling ageâ€specific facial development in Williams–Beurenâ€; Noonanâ€; and 22q11.2 deletion syndromes in cohorts of Czech patients aged 3–18 years: A crossâ€sectional threeâ€dimensional geometric morphometry analysis of their facial gestalt. American Journal of Medical Genetics, Part A, 2018, 176, 2604-2613.	1.2	2
8	Identification of likely associations between cerebral folate deficiency and complex genetic―and metabolic pathogenesis of autism spectrum disorders by utilization of a pilot interaction modeling approach. Autism Research, 2017, 10, 1424-1435.	3.8	15