## Shahab Noorian

## List of Publications by Citations

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The third column is the impact factor (IF) of the journal, and the fourth column is the number of citations of the article.

14<br/>papers24<br/>citations3<br/>h-index4<br/>g-index18<br/>ext. papers38<br/>ext. citations1.4<br/>avg, IF1.24<br/>L-index

#	Paper	IF	Citations
14	Reliability of pubertal self assessment method: an Iranian study. <i>Iranian Journal of Pediatrics</i> , <b>2013</b> , 23, 327-32	1	7
13	Diagnostic Approach to the Patients with Suspected Primary Immunodeficiency. <i>Endocrine, Metabolic and Immune Disorders - Drug Targets</i> , <b>2020</b> , 20, 157-171	2.2	6
12	A novel pathogenic variant of gene causes rigidity and multifocal seizure syndrome, lethal neonatal. <i>International Journal of Neuroscience</i> , <b>2021</b> , 131, 875-878	2	3
11	GCK Mutation in a Child with Maturity Onset Diabetes of the Young, Type 2. <i>Iranian Journal of Pediatrics</i> , <b>2013</b> , 23, 226-8	1	3
10	A novel nonsense mutation in the WFS1 gene causes the Wolfram syndrome. <i>Journal of Pediatric Endocrinology and Metabolism</i> , <b>2016</b> , 29, 607-9	1.6	2
9	Whole-Exome Sequencing in Idiopathic Short Stature: Rare Mutations Affecting Growth. <i>Journal of Pediatric Genetics</i> , <b>2021</b> , 10, 284-291	0.7	1
8	The Prevalence of Selective and Partial Immunoglobulin A Deficiency in Patients with Autoimmune Polyendocrinopathy. <i>Immunological Investigations</i> , <b>2021</b> , 1-9	2.9	1
7	A Family With Novel X-Linked Recessive Homozygous Mutation in (c.628_629 del, p.1210fs*) in Kallmann Syndrome Associated Unilateral Ptosis: Case Report and Literature Review. <i>AACE Clinical Case Reports</i> , <b>2021</b> , 7, 216-219	0.7	1
6	A novel missense mutation of the HGD gene causes Alkaptonuria. <i>Meta Gene</i> , <b>2018</b> , 18, 174-176	0.7	O
5	The Role of Thyroid Function Tests in Diagnosing Allan-herndon-dudley Syndrome Revisited: A Novel Iran-based Mutation <i>Basic and Clinical Neuroscience</i> , <b>2021</b> , 12, 563-568	1.4	О
4	Dizygotic Twins Concordant for Down Syndrome: Implication for Establishing a National Birth Defect Registry in Iran. <i>Iranian Journal of Public Health</i> , <b>2016</b> , 45, 1667-1668	0.7	
3	Familial hypercholesterolemia in an Iranian family due to a mutation in the APOE gene (first case report). <i>Journal of Diabetes and Metabolic Disorders</i> ,1	2.5	
2	Compound Heterozygous Mutations Presented with Quadriparesis and Menopause. A Case Report <i>Twin Research and Human Genetics</i> , <b>2022</b> , 1-3	2.2	
1	COVID-19 and Diabetic Ketoacidosis in a Child: A Case Report. <i>International Journal of Enteric Pathogens</i> , <b>2021</b> , 9, 78-80	0.2	