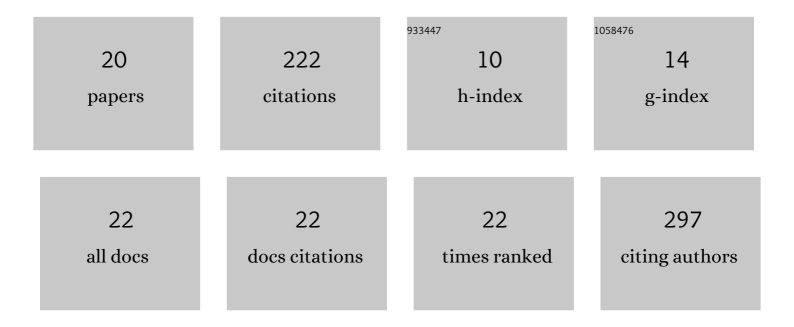
Nasser A Elhawary

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Molecular Updating of β-Thalassemia Mutations in the Upper Egyptian Population. Hemoglobin, 2010, 34, 538-547.	0.8	23
2	The <i>MTHFR</i> 677T Allele May Influence the Severity and Biochemical Risk Factors of Alzheimer's Disease in an Egyptian Population. Disease Markers, 2013, 35, 439-446.	1.3	22
3	Association between <i>ANKK1</i> (rs1800497) and <i>LTA</i> (rs909253) Genetic Variants and Risk of Schizophrenia. BioMed Research International, 2015, 2015, 1-8.	1.9	18
4	Genetic etiology and clinical challenges of phenylketonuria. Human Genomics, 2022, 16, .	2.9	18
5	Molecular characterization of exonic rearrangements and frame shifts in the dystrophin gene in Duchenne muscular dystrophy patients in a Saudi community. Human Genomics, 2018, 12, 18.	2.9	17
6	The Impact of Common Tumor Necrosis Factor Haplotypes on the Development of Asthma in Children: An Egyptian Model. Genetic Testing and Molecular Biomarkers, 2011, 15, 293-299.	0.7	16
7	TNFâ€238 polymorphism may predict bronchopulmonary dysplasia among preterm infants in the Egyptian population. Pediatric Pulmonology, 2013, 48, 699-706.	2.0	16
8	Frameshift deletion mechanisms in Egyptian Duchenne and Becker muscular dystrophy families. Molecules and Cells, 2004, 18, 141-9.	2.6	15
9	Combined Genetic Biomarkers Confer Susceptibility to Risk of Urothelial Bladder Carcinoma in a Saudi Population. Disease Markers, 2017, 2017, 1-11.	1.3	13
10	Mutations in Transglutaminase 1 Gene in Autosomal Recessive Congenital Ichthyosis in Egyptian Families. Disease Markers, 2004, 20, 325-332.	1.3	12
11	<p>Methylenetetrahydrofolate Reductase Gene Variants Confer Potential Vulnerability to Autism Spectrum Disorder in a Saudi Community</p> . Neuropsychiatric Disease and Treatment, 2019, Volume 15, 3569-3581.	2.2	10
12	Transporter <i>TAP1</i> -637G and Immunoproteasome <i>PSMB9</i> -60H Variants Influence the Risk of Developing Vitiligo in the Saudi Population. Disease Markers, 2014, 2014, 1-8.	1.3	9
13	Genetic biomarkers predict susceptibility to autism spectrum disorder through interactive models of inheritance in a Saudi community. Cogent Biology, 2019, 5, 1606555.	1.7	9
14	High-precision DNA microsatellite genotyping in Duchenne muscular dystrophy families using ion-pair reversed-phase high performance liquid chromatography. Clinical Biochemistry, 2006, 39, 758-761.	1.9	5
15	Association between β+252 tumour necrosis factor polymorphism and asthma in western Saudi children. Saudi Journal of Biological Sciences, 2011, 18, 107-111.	3.8	4
16	Risk of Colorectal Carcinoma May Predispose to the Genetic Variants of the GST, CYP450, and TP53 Genes Among Nonsmokers in the Saudi Community. International Journal of General Medicine, 2021, Volume 14, 1311-1323.	1.8	4
17	Common Tag STSs in the AZF Region Associated with Azoospermia and Severe Oligospermia in Infertile Egyptian Men~!2009-12-21~!2010-04-13~!2010-06-18~!. The Open Andrology Journal, 2010, 2, 11-18.	0.2	4
18	Null genetic risk of ACE gene polymorphisms with nephropathy in type 1 diabetes among Egyptian population. Egyptian Journal of Medical Human Genetics, 2011, 12, 187-192.	1.0	2

#	Article	IF	CITATIONS
19	Azoospermia factor microdeletions: common tag STSs in infertile men with azoospermia and sever oligospermia from Egypt. BMC Genomics, 2014, 15, .	2.8	2
20	Variations in TAP1 and PSMB9 Genes Involved in Antigen Processing and Presentation Increase the Risk of Vitiligo in the Saudi Community. International Journal of General Medicine, 2021, Volume 14, 10031-10044.	1.8	2