

# Hugo H Abarca-Barriga

## List of Publications by Year in descending order

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Version: 2024-02-01

24  
papers

290  
citations

1477746

6  
h-index

887659

17  
g-index

27  
all docs

27  
docs citations

27  
times ranked

640  
citing authors

#	ARTICLE	IF	CITATIONS
1	Variantes en el número de copias y consanguinidad parental en neonatos de altura con anomalías congénitas en Perú. Revista De La Facultad De Ciencias Medicas De Cordoba, 2022, 79, 132-140.	0.1	1
2	Management of genetic diseases: Present and future. Revista De La Facultad De Medicina Humana, 2021, 21, 399-416.	0.1	2
3	Avances genómicos de la última década y su influencia en el enfoque diagnóstico de la discapacidad intelectual.. Revista De Neuro-psiquiatria, 2021, 84, 33-50.	0.0	1
4	Importancia de la determinación de variantes en el número de copias en neonatos con aneuploidías autosómicas. Biomedica, 2021, 41, 282-292.	0.3	0
5	MLPA followed by targeted NGS to detect mutations in the dystrophin gene of Peruvian patients suspected of DMD/DMB. Molecular Genetics & Genomic Medicine, 2021, 9, e1759.	0.6	5
6	Phenotypic expansion in <i>KIF1A</i> -related dominant disorders: A description of novel variants and review of published cases. Human Mutation, 2020, 41, 2094-2104.	1.1	8
7	Peruvian Newborn Male with 3p13 Deletion Syndrome Encompassing the FOXP1 Gene: Review of the Literature. Journal of Pediatric Genetics, 2020, 09, 270-278.	0.3	2
8	High prevalence of congenital generalized lipodystrophy in Piura, Peru. Intractable and Rare Diseases Research, 2020, 9, 58-60.	0.3	1
9	Variantes en el número de copias en trastornos del neurodesarrollo, síndrome malformativo y talla baja en Perú. Acta Medica Peruana, 2020, 37, .	0.3	1
10	Ampliación del tamizaje de errores innatos del metabolismo en Perú: reporte de caso con trastorno del metabolismo de cobalamina. Acta Medica Peruana, 2020, 37, .	0.3	1
11	Genetic basis of pulmonary arterial hypertension. Revista De La Facultad De Medicina Humana, 2020, 20, 670-681.	0.1	0
12	Homozygous Deletion of the CFTR Gene Caused by Interstitial Maternal Isodisomy in a Peruvian Child with Cystic Fibrosis. Journal of Pediatric Genetics, 2019, 08, 147-152.	0.3	2
13	GATAD2B Gene Microdeletion Causing Intellectual Disability Autosomal Dominant Type 18: Case Report and Review of the Literature. Molecular Syndromology, 2019, 10, 186-194.	0.3	2
14	Williams-Buren syndrome in diverse populations. American Journal of Medical Genetics, Part A, 2018, 176, 1128-1136.	0.7	55
15	Novel contiguous gene deletion in peruvian girl with Trichothiodystrophy type 4 and glutaric aciduria type 3. European Journal of Medical Genetics, 2018, 61, 388-392.	0.7	7
16	Genetics and genomics in Peru: Clinical and research perspective. Molecular Genetics & Genomic Medicine, 2018, 6, 873-886.	0.6	12
17	A novel <i>ASPH</i> variant extends the phenotype of Shawafaraboulsi syndrome. American Journal of Medical Genetics, Part A, 2018, 176, 2494-2500.	0.7	17
18	Cover Image, Volume 176A, Number 5, May 2018. American Journal of Medical Genetics, Part A, 2018, 176, .	0.7	0

#	ARTICLE	IF	CITATIONS
19	Factores de riesgo en las enfermedades genéticas. Acta Medica Peruana, 2018, 35, 43-50.	0.3	4
20	Cover Image, Volume 173A, Number 9, September 2017. American Journal of Medical Genetics, Part A, 2017, 173, i.	0.7	0
21	Noonan syndrome in diverse populations. American Journal of Medical Genetics, Part A, 2017, 173, 2323-2334.	0.7	68
22	A Peruvian Child with 18p-/18q+ Syndrome and Persistent Microscopic Hematuria. Journal of Pediatric Genetics, 2017, 06, 258-266.	0.3	7
23	Microduplications encompassing the Sonic hedgehog limb enhancer <scp>ZRS</scp> are associated with Haas-type polysyndactyly and Laurin-Sandrow syndrome. Clinical Genetics, 2014, 86, 318-325.	1.0	72
24	Ocular pterygium-Digital keloid dysplasia. American Journal of Medical Genetics, Part A, 2014, 164, 2901-2907.	0.7	7