Sarah E Sheppard

List of Publications by Year in descending order

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1163117 888059 27 340 8 17 citations g-index h-index papers 29 29 29 548 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	What is new with 22q? An update from the 22q and You Center at the Children's Hospital of Philadelphia. American Journal of Medical Genetics, Part A, 2018, 176, 2058-2069.	1.2	106
2	Kaposiform lymphangiomatosis effectively treated with <scp>MEK</scp> inhibition. EMBO Molecular Medicine, 2020, 12, e12324.	6.9	51
3	Expanding the genotypic and phenotypic spectrum in a diverse cohort of 104 individuals with Wiedemannâ€6teiner syndrome. American Journal of Medical Genetics, Part A, 2021, 185, 1649-1665.	1.2	34
4	Genotype–phenotype specificity in Menke–Hennekam syndrome caused by missense variants in exon 30 or 31 of CREBBP. American Journal of Medical Genetics, Part A, 2019, 179, 1058-1062.	1.2	23
5	Bi-allelic Loss-of-Function Variants in NUP188 Cause a Recognizable Syndrome Characterized by Neurologic, Ocular, and Cardiac Abnormalities. American Journal of Human Genetics, 2020, 106, 623-631.	6.2	18
6	Androgenetic chimerism as an etiology for Beckwith–Wiedemann syndrome: diagnosis and management. Genetics in Medicine, 2019, 21, 2644-2649.	2.4	15
7	Pathogenic variants in <i>MDFIC</i> cause recessive central conducting lymphatic anomaly with lymphedema. Science Translational Medicine, 2022, 14, eabm4869.	12.4	14
8	Trisomy 9 mosaic syndrome: Sixteen additional patients with new and/or less commonly reported features, literature review, and suggested clinical guidelines. American Journal of Medical Genetics, Part A, 2021, 185, 2374-2383.	1.2	11
9	Further delineation of the phenotypic spectrum of nevus comedonicus syndrome to include congenital pulmonary airway malformation of the lung and aneurysm. American Journal of Medical Genetics, Part A, 2020, 182, 746-754.	1.2	9
10	Genetics etiologies and genotype phenotype correlations in a cohort of individuals with central conducting lymphatic anomaly. European Journal of Human Genetics, 2022, 30, 1022-1028.	2.8	9
11	Heterozygous recurrent <scp><i>HNF4A</i></scp> variant p. <scp>Arg85Trp</scp> causes Fanconi renotubular syndrome 4 with maturity onset diabetes of the young, an autosomal dominant phenocopy of Fanconi Bickel syndrome with colobomas. American Journal of Medical Genetics, Part A, 2021, 185, 566-570.	1.2	7
12	Expanded phenotypic spectrum of <scp><i>JAG1</i></scp> â€associated diseases: Central conducting lymphatic anomaly with a pathogenic variant in <scp><i>JAG1</i></scp> . Clinical Genetics, 2021, 99, 742-743.	2.0	7
13	Rare variants in KDR, encoding VEGF Receptor 2, are associated with tetralogy of Fallot. Genetics in Medicine, 2021, 23, 1952-1960.	2.4	7
14	Cerebrofacial vascular metameric syndrome is caused by somatic pathogenic variants in <i>PIK3CA</i> Journal of Physical Education and Sports Management, 2021, 7, a006147.	1.2	6
15	Proposed criteria for nevoid basal cell carcinoma syndrome in children assessed using statistical optimization. Scientific Reports, 2021, 11, 19791.	3.3	5
16	Clinical Effectiveness of Telemedicine-Based Pediatric Genetics Care. Pediatrics, 2022, 150, .	2.1	5
17	Congenital polyvalvular disease expands the cardiac phenotype of the RASopathies. American Journal of Medical Genetics, Part A, 2021, 185, 1486-1493.	1.2	3
18	Genetic skin disorders: The value of a multidisciplinary clinic. American Journal of Medical Genetics, Part A, 2021, 185, 1159-1167.	1.2	3

#	Article	IF	CITATIONS
19	Hyperinsulinism in an individual with an EP300 variant of Rubinsteinâ€√aybi syndrome. American Journal of Medical Genetics, Part A, 2021, 185, 1251-1255.	1.2	2
20	Chromosome 4q28.3q32.3 duplication in a patient with lymphatic malformations, craniosynostosis, and dysmorphic features. Clinical Dysmorphology, 2021, 30, 89-92.	0.3	2
21	Generalized, severe epidermolysis bullosa simplex caused by a Keratin 5 p.E477K mutation. Pediatric Dermatology, 2019, 36, 1007-1009.	0.9	1
22	Muenke syndrome: Medical and surgical comorbidities and longâ€term management. American Journal of Medical Genetics, Part A, 2019, 179, 1442-1450.	1.2	1
23	Misdiagnosis of capillary malformations in darker skin phototypes. Pediatric Dermatology, 2021, 38 Suppl 2, 137-139.	0.9	1
24	Cover Image, Volume 176A, Number 10, October 2018. , 2018, 176, i-i.		0
25	50 Years Ago in T J P. Journal of Pediatrics, 2021, 233, 211.	1.8	O
26	Elucidating the clinical spectrum and molecular basis of HYAL2 deficiency. Genetics in Medicine, 2022, 24, 631-644.	2.4	0
27	50 Years Ago in T J P. Journal of Pediatrics, 2022, 241, 195.	1.8	O