

Kalman Tory

List of Publications by Year in descending order

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Version: 2024-02-01

19
papers

571
citations

759233

12
h-index

888059

17
g-index

20
all docs

20
docs citations

20
times ranked

977
citing authors

#	ARTICLE	IF	CITATIONS
1	Mutation-dependent recessive inheritance of NPHS2-associated steroid-resistant nephrotic syndrome. <i>Nature Genetics</i> , 2014, 46, 299-304.	21.4	134
2	Genotype-phenotype associations in WT1 glomerulopathy. <i>Kidney International</i> , 2014, 85, 1169-1178.	5.2	113
3	A molecular mechanism explaining albuminuria in kidney disease. <i>Nature Metabolism</i> , 2020, 2, 461-474.	11.9	99
4	Pseudouridylation defect due to <i>DKC1</i> and <i>NOP10</i> mutations causes nephrotic syndrome with cataracts, hearing impairment, and enterocolitis. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2020, 117, 15137-15147.	7.1	32
5	Selective measurement of β -smooth muscle actin: why α -actin can not be used as a housekeeping gene when tissue fibrosis occurs. <i>BMC Molecular Biology</i> , 2017, 18, 12.	3.0	31
6	An international cohort study of autosomal dominant tubulointerstitial kidney disease due to mutations identifies distinct clinical subtypes. <i>Kidney International</i> , 2020, 98, 1589-1604.	5.2	27
7	Comprehensive genetic testing in children with a clinical diagnosis of ARPKD identifies phenocopies. <i>Pediatric Nephrology</i> , 2018, 33, 1713-1721.	1.7	25
8	NPHS2 homozygous p.R229Q variant: potential modifier instead of causal effect in focal segmental glomerulosclerosis. <i>Pediatric Nephrology</i> , 2013, 28, 2061-2064.	1.7	21
9	The mutation-dependent pathogenicity of <i>NPHS2</i> p.R229Q: A guide for clinical assessment. <i>Human Mutation</i> , 2018, 39, 1854-1860.	2.5	17
10	Endoplasmic reticulum-retained podocin mutants are massively degraded by the proteasome. <i>Journal of Biological Chemistry</i> , 2018, 293, 4122-4133.	3.4	16
11	C-terminal oligomerization of podocin mediates interallelic interactions. <i>Biochimica Et Biophysica Acta - Molecular Basis of Disease</i> , 2018, 1864, 2448-2457.	3.8	15
12	Autonomic dysfunction in uremia assessed by heart rate variability. <i>Pediatric Nephrology</i> , 2003, 18, 1167-1171.	1.7	13
13	Signs of autonomic neuropathy in childhood uremia. <i>Pediatric Nephrology</i> , 2001, 16, 25-28.	1.7	10
14	<i>EPG5</i> c.1007A>G mutation in a sibling pair with rapidly progressing Vici syndrome. <i>Annals of Human Genetics</i> , 2020, 84, 80-86.	0.8	6
15	Clinical and genetic findings in Hungarian pediatric patients carrying chromosome 16p copy number variants and a review of the literature. <i>European Journal of Medical Genetics</i> , 2020, 63, 104027.	1.3	4
16	Identification of incompletely penetrant variants and interallelic interactions in autosomal recessive disorders by a population-genetic approach. <i>Human Mutation</i> , 2021, 42, 1473-1487.	2.5	4
17	Critical Considerations in Genetic Counseling of Patients With the NPHS2 R229Q Variant. <i>American Journal of Kidney Diseases</i> , 2019, 73, 576.	1.9	1
18	MO032PODOCIN REGULATES THE SIZE OF THE GLOMERULAR PORE. <i>Nephrology Dialysis Transplantation</i> , 2020, 35, .	0.7	0

#	ARTICLE	IF	CITATIONS
19	MO044: Cellular mechanism of the exceptional dominant transmission in NPHS2-associated glomerulopathy. Nephrology Dialysis Transplantation, 2022, 37, .	0.7	0