Koichi Nakanishi

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Oxford Classification of IgA nephropathy 2016: anÂupdate from the IgA Nephropathy Classification Working Group. Kidney International, 2017, 91, 1014-1021.	5.2	748
2	Rituximab for childhood-onset, complicated, frequently relapsing nephrotic syndrome or steroid-dependent nephrotic syndrome: a multicentre, double-blind, randomised, placebo-controlled trial. Lancet, The, 2014, 384, 1273-1281.	13.7	326
3	Alport syndrome: a unified classification of genetic disorders of collagen IV α345: a position paper of the Alport Syndrome Classification Working Group. Kidney International, 2018, 93, 1045-1051.	5.2	206
4	IPNA clinical practice recommendations for the diagnosis and management of children with steroid-resistant nephrotic syndrome. Pediatric Nephrology, 2020, 35, 1529-1561.	1.7	179
5	A review of clinical characteristics and genetic backgrounds in Alport syndrome. Clinical and Experimental Nephrology, 2019, 23, 158-168.	1.6	135
6	Steroid Treatment for Severe Childhood IgA Nephropathy: A Randomized, Controlled Trial. Clinical Journal of the American Society of Nephrology: CJASN, 2006, 1, 511-517.	4.5	114
7	Organoids from Nephrotic Disease-Derived iPSCs Identify Impaired NEPHRIN Localization and Slit Diaphragm Formation in Kidney Podocytes. Stem Cell Reports, 2018, 11, 727-740.	4.8	113
8	Immunohistochemical study of $\hat{l}\pm 1$ -5 chains of type IV collagen in hereditary nephritis. Kidney International, 1994, 46, 1413-1421.	5.2	111
9	A multicenter randomized trial indicates initial prednisolone treatment for childhood nephrotic syndrome for two months is not inferior to six-month treatment. Kidney International, 2015, 87, 225-232.	5.2	104
10	Essential points from Evidence-based Clinical Practice Guidelines for Chronic Kidney Disease 2018. Clinical and Experimental Nephrology, 2019, 23, 1-15.	1.6	99
11	Group A streptococcal antigen in the glomeruli of children with henoch-schönlein nephritis. American Journal of Kidney Diseases, 2003, 41, 366-370.	1.9	98
12	Genetic, Clinical, and Pathologic Backgrounds of Patients with Autosomal Dominant Alport Syndrome. Clinical Journal of the American Society of Nephrology: CJASN, 2016, 11, 1441-1449.	4.5	94
13	Validity of the Oxford classification of IgA nephropathy in children. Pediatric Nephrology, 2012, 27, 783-792.	1.7	92
14	Minimal change nephrotic syndrome associated with immune dysregulation, polyendocrinopathy, enteropathy, X-linked syndrome. Pediatric Nephrology, 2009, 24, 1181-1186.	1.7	85
15	Molecular analysis of digenic inheritance in Bartter syndrome with sensorineural deafness. Journal of Medical Genetics, 2007, 45, 182-186.	3.2	84
16	Analysis of NPHS1, NPHS2, ACTN4, and WT1 in Japanese patients with congenital nephrotic syndrome. Kidney International, 2005, 67, 1248-1255.	5.2	77
17	HNF1B alterations associated with congenital anomalies of the kidney and urinary tract. Pediatric Nephrology, 2010, 25, 1073-1079.	1.7	76
18	Long-Term Results of a Randomized Controlled Trial in Childhood IgA Nephropathy. Clinical Journal of the American Society of Nephrology: CJASN, 2011, 6, 1301-1307.	4.5	76

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19	Milder clinical aspects of X-linked Alport syndrome in men positive for the collagen IV α5 chain. Kidney International, 2014, 85, 1208-1213.	5.2	72
20	Differential diagnosis of Bartter syndrome, Gitelman syndrome, and pseudo–Bartter/Gitelman syndrome based on clinical characteristics. Genetics in Medicine, 2016, 18, 180-188.	2.4	67
21	Cyclosporine and steroid therapy in children with steroid-resistant nephrotic syndrome. Pediatric Nephrology, 2009, 24, 2177-2185.	1.7	66
22	Clinical practice guideline for pediatric idiopathic nephrotic syndrome 2013: medical therapy. Clinical and Experimental Nephrology, 2015, 19, 6-33.	1.6	62
23	Natural History and Genotype–Phenotype Correlation in Female X-Linked Alport Syndrome. Kidney International Reports, 2017, 2, 850-855.	0.8	62
24	Genetic disorders of human congenital anomalies of the kidney and urinary tract (CAKUT). Pediatrics International, 2003, 45, 610-616.	0.5	61
25	Spontaneous remission in children with IgA nephropathy. Pediatric Nephrology, 2013, 28, 71-76.	1.7	58
26	Natural history of genetically proven autosomal recessive Alport syndrome. Pediatric Nephrology, 2014, 29, 1535-1544.	1.7	55
27	Genotype-phenotype correlations influence the response to angiotensin-targeting drugs in Japanese patients with male X-linked Alport syndrome. Kidney International, 2020, 98, 1605-1614.	5.2	55
28	Morbidity in children with frequently relapsing nephrosis: 10-year follow-up of a randomized controlled trial. Pediatric Nephrology, 2015, 30, 459-468.	1.7	54
29	Strong Association of the HLA-DR/DQ Locus with Childhood Steroid-Sensitive Nephrotic Syndrome in the Japanese Population. Journal of the American Society of Nephrology: JASN, 2018, 29, 2189-2199.	6.1	54
30	The Pharmacological Characteristics of Molecular-Based Inherited Salt-Losing Tubulopathies. Journal of Clinical Endocrinology and Metabolism, 2010, 95, E511-E518.	3.6	52
31	Pre-dialysis chronic kidney disease in children: results of a nationwide survey in Japan. Nephrology Dialysis Transplantation, 2013, 28, 2345-2355.	0.7	52
32	Comprehensive genetic diagnosis of Japanese patients with severe proteinuria. Scientific Reports, 2020, 10, 270.	3.3	50
33	Improved renal survival in Japanese children with IgA nephropathy. Pediatric Nephrology, 2008, 23, 905-912.	1.7	49
34	Combination therapy with mizoribine for severe childhood IgA nephropathy: a pilot study. Pediatric Nephrology, 2008, 23, 757-763.	1.7	49
35	Detection of mutations in the COL4A5 gene in over 90% of male patients with x-linked Alport's syndrome by RT-PCR and direct sequencing. American Journal of Kidney Diseases, 1999, 34, 854-862.	1.9	47
36	Treatment strategies for Henoch-Schönlein purpura nephritis by histological and clinical severity. Pediatric Nephrology, 2011, 26, 563-569.	1.7	47

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37	Association between low birth weight and childhood-onset chronic kidney disease in Japan: a combined analysis of a nationwide survey for paediatric chronic kidney disease and the National Vital Statistics Report. Nephrology Dialysis Transplantation, 2016, 31, 1895-1900.	0.7	47
38	Clinical and Genetic Characteristics in Patients With Gitelman Syndrome. Kidney International Reports, 2019, 4, 119-125.	0.8	47
39	Epithelial-to-mesenchymal transition in cyst lining epithelial cells in an orthologous PCK rat model of autosomal-recessive polycystic kidney disease. American Journal of Physiology - Renal Physiology, 2011, 300, F511-F520.	2.7	44
40	Detection of Splicing Abnormalities and Genotype-Phenotype Correlation in X-linked Alport Syndrome. Journal of the American Society of Nephrology: JASN, 2018, 29, 2244-2254.	6.1	43
41	Two-Year Follow-Up of a Prospective Clinical Trial of Cyclosporine for Frequently Relapsing Nephrotic Syndrome in Children. Clinical Journal of the American Society of Nephrology: CJASN, 2012, 7, 1576-1583.	4.5	42
42	Molecular Analysis of Patients With Type III Bartter Syndrome: Picking Up Large Heterozygous Deletions With Semiquantitative PCR. Pediatric Research, 2007, 62, 364-369.	2.3	41
43	Treatment with microemulsified cyclosporine in children with frequently relapsing nephrotic syndrome. Nephrology Dialysis Transplantation, 2010, 25, 3956-3962.	0.7	41
44	Two-Year Outcome of the ISKDC Regimen and Frequent-Relapsing Risk in Children with Idiopathic Nephrotic Syndrome. Clinical Journal of the American Society of Nephrology: CJASN, 2013, 8, 756-762.	4.5	41
45	High incidence of idiopathic nephrotic syndrome in East Asian children: a nationwide survey in Japan (JP-SHINE study). Clinical and Experimental Nephrology, 2017, 21, 651-657.	1.6	41
46	The effect of aldosterone blockade in patients with Alport syndrome. Pediatric Nephrology, 2006, 21, 1824-1829.	1.7	40
47	Identification of mutations in FN1 leading to glomerulopathy with fibronectin deposits. Pediatric Nephrology, 2016, 31, 1459-1467.	1.7	40
48	Common risk variants in NPHS1 and TNFSF15 are associated with childhood steroid-sensitive nephrotic syndrome. Kidney International, 2020, 98, 1308-1322.	5.2	39
49	Cryopreservation of four valuable strains of microalgae, including viability and characteristics during 15Âyears of cryostorage. Journal of Applied Phycology, 2012, 24, 1381-1385.	2.8	38
50	Prospective 5-year follow-up of cyclosporine treatment in children with steroid-resistant nephrosis. Pediatric Nephrology, 2013, 28, 765-771.	1.7	37
51	Evidence-based clinical practice guidelines for polycystic kidney disease 2014. Clinical and Experimental Nephrology, 2016, 20, 493-509.	1.6	37
52	In vivo and in vitro splicing assay of SLC12A1 in an antenatal salt-losing tubulopathy patient with an intronic mutation. Human Genetics, 2009, 126, 533-538.	3.8	36
53	Progression to end-stage kidney disease in Japanese children with chronic kidney disease: results of a nationwide prospective cohort study. Nephrology Dialysis Transplantation, 2014, 29, 878-884.	0.7	36
54	A Deep Intronic Mutation in the SLC12A3 Gene Leads to Gitelman Syndrome. Pediatric Research, 2009, 66, 590-593.	2.3	35

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55	Clinical practice guideline for pediatric idiopathic nephrotic syndrome 2013: general therapy. Clinical and Experimental Nephrology, 2015, 19, 34-53.	1.6	35
56	Long-term outcome of childhood-onset complicated nephrotic syndrome after a multicenter, double-blind, randomized, placebo-controlled trial of rituximab. Pediatric Nephrology, 2017, 32, 2071-2078.	1.7	35
57	Efficacy and safety of lisinopril for mild childhood IgA nephropathy: a pilot study. Pediatric Nephrology, 2009, 24, 845-849.	1.7	34
58	Risk factors for developing severe clinical course in HUS patients: a national survey in Japan. Pediatrics International, 2008, 50, 441-446.	0.5	33
59	Disappearance of glomerular IgA deposits in childhood IgA nephropathy showing diffuse mesangial proliferation after 2 years of combination/prednisolone therapy. Nephrology Dialysis Transplantation, 2011, 26, 163-169.	0.7	31
60	X-Linked Alport Syndrome Caused by Splicing Mutations in COL4A5. Clinical Journal of the American Society of Nephrology: CJASN, 2014, 9, 1958-1964.	4.5	31
61	Recurrent EIARF and PRES With Severe Renal Hypouricemia by Compound Heterozygous SLC2A9 Mutation. Pediatrics, 2011, 127, e1621-e1625.	2.1	29
62	Somatic mosaicism for a mutation of the COL4A5 gene is a cause of mild phenotype male Alport syndrome. Nephrology Dialysis Transplantation, 2008, 23, 2525-2530.	0.7	28
63	Membranous nephropathy associated with thyroid-peroxidase antigen. Pediatric Nephrology, 2009, 24, 605-608.	1.7	28
64	Cyclosporine C2 Monitoring for the Treatment of Frequently Relapsing Nephrotic Syndrome in Children. Clinical Journal of the American Society of Nephrology: CJASN, 2014, 9, 271-278.	4.5	27
65	Biopsy timing and Oxford classification variables in Childhood/Adolescent IgA nephropathy. Pediatric Nephrology, 2015, 30, 293-299.	1.7	27
66	Updating the International IgA Nephropathy Prediction Tool for use in children. Kidney International, 2021, 99, 1439-1450.	5.2	26
67	A-20C angiotensinogen gene polymorphism and proteinuria in childhood IgA nephropathy. Pediatric Nephrology, 2004, 19, 144-147.	1.7	25
68	Comparison between conventional and comprehensive sequencing approaches for genetic diagnosis of Alport syndrome. Molecular Genetics & Genomic Medicine, 2019, 7, e883.	1.2	25
69	Segmental Membranous Glomerulonephritis in Children: Comparison with Global Membranous Glomerulonephritis. Clinical Journal of the American Society of Nephrology: CJASN, 2006, 1, 723-729.	4.5	24
70	Long-term outcome of childhood IgA nephropathy with minimal proteinuria. Pediatric Nephrology, 2015, 30, 2121-2127.	1.7	24
71	Combination therapy with or without warfarin and dipyridamole for severe childhood IgA nephropathy: an RCT. Pediatric Nephrology, 2018, 33, 2103-2112.	1.7	24
72	Mycophenolate Mofetil after Rituximab for Childhood-Onset Complicated Frequently-Relapsing or Steroid-Dependent Nephrotic Syndrome. Journal of the American Society of Nephrology: JASN, 2022, 33, 401-419.	6.1	24

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73	Telmisartan Ameliorates Fibrocystic Liver Disease in an Orthologous Rat Model of Human Autosomal Recessive Polycystic Kidney Disease. PLoS ONE, 2013, 8, e81480.	2.5	23
74	Lisinopril versus lisinopril and losartan for mild childhood IgA nephropathy: a randomized controlled trial (JSKDC01 study). Pediatric Nephrology, 2019, 34, 837-846.	1.7	23
75	A Novel System for Spinal Muscular Atrophy Screening in Newborns: Japanese Pilot Study. International Journal of Neonatal Screening, 2019, 5, 41.	3.2	22
76	Renal biopsy criterion in children with asymptomatic constant isolated proteinuria. Nephrology Dialysis Transplantation, 2012, 27, 3186-3190.	0.7	21
77	Diglycerol esters of fatty acids promote severe coalescence between protein-stabilized oil droplets by emulsifier–protein competitive interactions. Food Hydrocolloids, 2014, 42, 397-402.	10.7	21
78	Somatic mosaicism and variant frequency detected by next-generation sequencing in X-linked Alport syndrome. European Journal of Human Genetics, 2016, 24, 387-391.	2.8	21
79	Prognosis and acute complications at the first onset of idiopathic nephrotic syndrome in children: a nationwide survey in Japan (JP-SHINE study). Nephrology Dialysis Transplantation, 2021, 36, 475-481.	0.7	21
80	Hydrogenated Amorphous Silicon Carbide Optical Waveguide for Telecommunication Wavelength Applications. Applied Physics Express, 2010, 3, 122201.	2.4	20
81	Characterization of contiguous gene deletions in COL4A6 and COL4A5 in Alport syndrome-diffuse leiomyomatosis. Journal of Human Genetics, 2017, 62, 733-735.	2.3	20
82	lgA nephropathy with presentation of nephrotic syndrome at onset in children. Pediatric Nephrology, 2017, 32, 457-465.	1.7	20
83	Heterozygous missense variant of the proteasome subunit Î ² -type 9 causes neonatal-onset autoinflammation and immunodeficiency. Nature Communications, 2021, 12, 6819.	12.8	20
84	Molecular Background of Urate Transporter Genes in Patients with Exercise-Induced Acute Kidney Injury. American Journal of Nephrology, 2013, 38, 316-320.	3.1	19
85	Cyclosporin A absorption profiles in children with nephrotic syndrome. Pediatric Nephrology, 2005, 20, 910-913.	1.7	18
86	Evaluation of the inferior vena cava in potential pediatric renal transplant recipients. Pediatric Nephrology, 2004, 19, 1062-4.	1.7	17
87	Severe Alport syndrome in a young woman caused by a t(X;1)(q22.3;p36.32) balanced translocation. Pediatric Nephrology, 2010, 25, 2165-2170.	1.7	17
88	X-linked Alport syndrome associated with a synonymous p.Gly292Gly mutation alters the splicing donor site of the type IV collagen alpha chain 5 gene. Clinical and Experimental Nephrology, 2016, 20, 699-702.	1.6	17
89	The relationship between arginine vasopressin levels and hyponatremia following a percutaneous renal biopsy in children receiving hypotonic or isotonic intravenous fluids. Pediatric Nephrology, 2011, 26, 99-104.	1.7	16
90	lsovaleric acidemia: Therapeutic response to supplementation with glycine, l -carnitine, or both in combination and a 10-year follow-up case study. Molecular Genetics and Metabolism Reports, 2017, 11, 2-5.	1.1	16

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91	Detailed clinical manifestations at onset and prognosis of neonatal-onset Denys–Drash syndrome and congenital nephrotic syndrome of the Finnish type. Clinical and Experimental Nephrology, 2019, 23, 1058-1065.	1.6	16
92	Pathogenic evaluation of synonymous <i>COL4A5</i> variants in Xâ€linked Alport syndrome using a minigene assay. Molecular Genetics & Genomic Medicine, 2020, 8, e1342.	1.2	16
93	Highâ€dose mizoribine treatment for adolescents with systemic lupus erythematosus. Pediatrics International, 2006, 48, 152-157.	0.5	15
94	Destabilization of protein-based emulsions by diglycerol esters of fatty acids – The importance of chain length similarity between dispersed oil molecules and fatty acid residues of the emulsifier. Food Hydrocolloids, 2011, 25, 773-780.	10.7	15
95	Growth impairment in children with pre-dialysis chronic kidney disease in Japan. Clinical and Experimental Nephrology, 2015, 19, 1142-1148.	1.6	15
96	Insignificant impact of VUR on the progression of CKD in children with CAKUT. Pediatric Nephrology, 2016, 31, 105-112.	1.7	15
97	A comparison of splicing assays to detect an intronic variant of the OCRL gene in Lowe syndrome. European Journal of Medical Genetics, 2017, 60, 631-634.	1.3	15
98	Endoplasmic reticulum stress with low-dose cyclosporine in frequently relapsing nephrotic syndrome. Pediatric Nephrology, 2013, 28, 903-909.	1.7	14
99	Alport syndrome caused by a <i>COL4A5</i> deletion and exonization of an adjacent <i>AluY</i> . Molecular Genetics & amp; Genomic Medicine, 2014, 2, 451-453.	1.2	14
100	Determination of the pathogenicity of known COL4A5 intronic variants by in vitro splicing assay. Scientific Reports, 2019, 9, 12696.	3.3	14
101	Impaired NEPHRIN localization in kidney organoids derived from nephrotic patient iPS cells. Scientific Reports, 2021, 11, 3982.	3.3	14
102	Last Nucleotide Substitutions of COL4A5 Exons Cause Aberrant Splicing. Kidney International Reports, 2022, 7, 108-116.	0.8	14
103	Female X-linked Alport syndrome with somatic mosaicism. Clinical and Experimental Nephrology, 2017, 21, 877-883.	1.6	13
104	Study protocol: multicenter double-blind, randomized, placebo-controlled trial of rituximab for the treatment of childhood-onset early-stage uncomplicated frequently relapsing or steroid-dependent nephrotic syndrome (JSKDC10 trial). BMC Nephrology, 2019, 20, 293.	1.8	13
105	Examination of the predicted prevalence of Gitelman syndrome by ethnicity based on genome databases. Scientific Reports, 2021, 11, 16099.	3.3	13
106	Immunoglobulin A Nephropathy. , 2009, , 757-781.		13
107	Expression of type IV collagen in the developing human kidney. Pediatric Nephrology, 1998, 12, 554-558.	1.7	12
108	Culture of a high-chlorophyll-producing and halotolerant Chlorella vulgaris. Journal of Bioscience and Bioengineering, 2014, 117, 617-619.	2.2	12

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109	Cryptic exon activation in SLC12A3 in Gitelman syndrome. Journal of Human Genetics, 2017, 62, 335-337.	2.3	12
110	Clinical spectrum of male patients with OFD1 mutations. Journal of Human Genetics, 2019, 64, 3-9.	2.3	12
111	Systematic Review of Genotype-Phenotype Correlations in Frasier Syndrome. Kidney International Reports, 2021, 6, 2585-2593.	0.8	12
112	Focal Segmental Glomerulosclerosis in Patients With Complete Deletion of One <i>WT1</i> Allele. Pediatrics, 2012, 129, e1621-e1625.	2.1	11
113	Development of method for evaluating cell hardness and correlation between bacterial spore hardness and durability. Journal of Nanobiotechnology, 2012, 10, 22.	9.1	11
114	Renal biopsy criterion in idiopathic nephrotic syndrome with microscopic hematuria at onset. Pediatric Nephrology, 2015, 30, 445-450.	1.7	11
115	Risk factors for persistent proteinuria after a 2-year combination therapy for severe childhood IgA nephropathy. Pediatric Nephrology, 2015, 30, 961-967.	1.7	11
116	A novel nonsense SMC1A mutation in a patient with intractable epilepsy and cardiac malformation. Human Genome Variation, 2019, 6, 23.	0.7	11
117	How to resolve confusion in the clinical setting for the diagnosis of heterozygous COL4A3 or COL4A4 gene variants? Discussion and suggestions from nephrologists. Clinical and Experimental Nephrology, 2020, 24, 651-656.	1.6	11
118	16q12 microdeletion syndrome in two <scp>J</scp> apanese boys. Pediatrics International, 2014, 56, e75-8.	0.5	10
119	Clinical characteristics and long-term outcome of diarrhea-associated hemolytic uremic syndrome: a single center experience. Clinical and Experimental Nephrology, 2017, 21, 889-894.	1.6	10
120	Aberrant Smad3 phosphoisoforms in cyst-lining epithelial cells in the <i>cpk</i> mouse, a model of autosomal recessive polycystic kidney disease. American Journal of Physiology - Renal Physiology, 2017, 313, F1223-F1231.	2.7	10
121	An in vitro splicing assay reveals the pathogenicity of a novel intronic variant in ATP6V0A4 for autosomal recessive distal renal tubular acidosis. BMC Nephrology, 2017, 18, 353.	1.8	10
122	Development of ultra-deep targeted RNA sequencing for analyzing X-chromosome inactivation in female Dent disease. Journal of Human Genetics, 2018, 63, 589-595.	2.3	10
123	Study protocol: mycophenolate mofetil as maintenance therapy after rituximab treatment for childhood-onset, complicated, frequently-relapsing nephrotic syndrome or steroid-dependent nephrotic syndrome: a multicenter double-blind, randomized, placebo-controlled trial (JSKDC07). BMC Nephrology, 2018, 19, 302.	1.8	10
124	Crescentic IgA nephropathy in children. Pediatric Nephrology, 2020, 35, 1005-1014.	1.7	10
125	Rare renal ciliopathies in non-consanguineous families that were identified by targeted resequencing. Clinical and Experimental Nephrology, 2017, 21, 136-142.	1.6	9
126	Long-term outcome of congenital nephrotic syndrome after kidney transplantation in Japan. Clinical and Experimental Nephrology, 2018, 22, 719-726.	1.6	9

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127	Prognosis and pathological characteristics of five children with non-Shiga toxin-mediated hemolytic uremic syndrome. Pediatrics International, 2007, 49, 196-201.	0.5	8
128	Detection by multiplex ligation-dependent probe amplification of large deletion mutations in the COL4A5 gene in female patients with Alport syndrome. Pediatric Nephrology, 2009, 24, 1773-1774.	1.7	8
129	Membranous nephropathy secondary to Graves' disease with deposits of thyroid peroxidase in an adult. CEN Case Reports, 2014, 3, 90-93.	0.9	8
130	Pathogenesis of hypokalemia in autosomal dominant hypocalcemia type 1. Clinical and Experimental Nephrology, 2016, 20, 253-257.	1.6	8
131	Diagnostic strategy for inherited hypomagnesemia. Clinical and Experimental Nephrology, 2017, 21, 1003-1010.	1.6	8
132	Glomerular galactose-deficient IgA1 expression analysis in pediatric patients with glomerular diseases. Scientific Reports, 2020, 10, 14026.	3.3	8
133	Molecular mechanisms determining severity in patients with Pierson syndrome. Journal of Human Genetics, 2020, 65, 355-362.	2.3	8
134	A digest from evidence-based Clinical Practice Guideline for Polycystic Kidney Disease 2020. Clinical and Experimental Nephrology, 2021, 25, 1292-1302.	1.6	8
135	A digest from evidence-based clinical practice guideline for IgA nephropathy 2020. Clinical and Experimental Nephrology, 2021, 25, 1269-1276.	1.6	8
136	Immunoglobulin A Nephropathies in Children (Includes HSP). , 2016, , 983-1033.		8
137	Alport-like glomerular basement membrane changes with renal-coloboma syndrome. Pediatric Nephrology, 2012, 27, 1189-1192.	1.7	7
138	With respect to coefficient of linear thermal expansion, bacterial vegetative cells and spores resemble plastics and metals, respectively. Journal of Nanobiotechnology, 2013, 11, 33.	9.1	7
139	First Japanese case of Pierson syndrome with mutations in <i>LAMB2</i> . Pediatrics International, 2013, 55, 229-231.	O.5	7
140	Congenital nephrotic syndrome with a novel <i><scp>NPHS</scp>1</i> mutation. Pediatrics International, 2016, 58, 1211-1215.	0.5	7
141	Clinical trial recommendations for potential Alport syndrome therapies. Kidney International, 2020, 97, 1109-1116.	5.2	7
142	Long-term follow-up of atypical membranoproliferative glomerulonephritis: are steroids indicated?. Pediatric Nephrology, 2006, 21, 194-200.	1.7	6
143	Increased chymase-positive mast cells in children with crescentic glomerulonephritis. Pediatric Nephrology, 2009, 24, 1071-1075.	1.7	6
144	Mild prominence of the Sylvian fissure in a Bainbridgeâ€Ropers syndrome patient with a novel frameshift variant in <i><scp>ASXL</scp>3</i> . Clinical Case Reports (discontinued), 2018, 6, 330-336.	0.5	6

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145	Study protocol: high-dose mizoribine with prednisolone therapy in short-term relapsing steroid-sensitive nephrotic syndrome to prevent frequent relapse (JSKDC05 trial). BMC Nephrology, 2018, 19, 223.	1.8	6
146	A novel homozygous missense SLC25A20 mutation in three CACT-deficient patients: clinical and autopsy data. Human Genome Variation, 2020, 7, 11.	0.7	6
147	Clinicopathological significance of glomerular capillary IgA deposition in childhood IgA nephropathy. Pediatric Nephrology, 2021, 36, 899-908.	1.7	6
148	Growth of Alicyclobacillus acidoterrestris in the Hypoxic Environment of Bottled Fruit Juice. Biocontrol Science, 2014, 19, 85-88.	0.8	5
149	Clinicopathological characteristics and renal outcomes of childhood-onset lupus nephritis with acute kidney injury: A multicenter study. Modern Rheumatology, 2019, 29, 970-976.	1.8	5
150	Comparison of clinical and genetic characteristics between Dent disease 1 and Dent disease 2. Pediatric Nephrology, 2020, 35, 2319-2326.	1.7	5
151	Use of renin-angiotensin system inhibitors as initial therapy in children with Henoch–Schönlein purpura nephritis of moderate severity. Pediatric Nephrology, 2022, 37, 1845-1853.	1.7	5
152	Comprehensive genetic analysis using next-generation sequencing for the diagnosis of nephronophthisis-related ciliopathies in the Japanese population. Journal of Human Genetics, 2022, 67, 427-440.	2.3	5
153	Detection of large deletion mutations in the COL4A5 gene of female Alport syndrome patients. Pediatric Nephrology, 2008, 23, 2085-2090.	1.7	4
154	Cryo-preserved porcine kidneys are feasible for teaching and training renal biopsy: "the bento kidney― Transplantation Research, 2012, 1, 5.	1.5	4
155	A birth of bipartite exon by intragenic deletion. Molecular Genetics & Genomic Medicine, 2017, 5, 287-294.	1.2	4
156	Possible involvement of ILâ€6â€producing tissueâ€resident macrophages in earlyâ€onset pericardial effusion pathogenesis after hematopoietic stem cell transplantation. Pediatric Blood and Cancer, 2018, 65, e26982.	1.5	4
157	Early parenteral nutrition in neonates with congenital diaphragmatic hernia. Pediatrics International, 2020, 62, 200-205.	0.5	4
158	Influenza virus vaccination in pediatric nephrotic syndrome significantly reduces rate of relapse and influenza virus infection as assessed in a nationwide survey. Scientific Reports, 2021, 11, 23305.	3.3	4
159	Bartter syndrome type 3 in an elderly complicated with adrenocorticotropin-deficiency. Endocrine Journal, 2014, 61, 855-860.	1.6	3
160	Extracellular signalâ€regulated kinase activation of selfâ€healing Langerhans cell histiocytosis: A case report. Journal of Dermatology, 2019, 46, 812-815.	1.2	3
161	Non-immunosuppressive therapies for childhood IgA nephropathy. Pediatric Nephrology, 2021, 36, 3057-3065.	1.7	3
162	Evaluation of suspected autosomal Alport Syndrome synonymous variants. Kidney360, 2022, 3, 10.34067/KID.0005252021.	2.1	3

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163	Primary diffuse leptomeningeal atypical teratoid rhabdoid tumor (AT/RT) demonstrating atypical imaging findings in an adolescent patient. Radiology Case Reports, 2022, 17, 485-488.	0.6	3
164	A patient with autosomal recessive Alport syndrome due to segmental maternal isodisomy. Human Genome Variation, 2014, 1, 14006.	0.7	2
165	Importance of clinical practice guidelines to practicing pediatric nephrologists and IPNA survey. Pediatric Nephrology, 2021, 36, 3493-3497.	1.7	2
166	Clinical features of autosomal recessive polycystic kidney disease in the Japanese population and analysis of splicing in PKHD1 gene for determination of phenotypes. Clinical and Experimental Nephrology, 2022, 26, 140-153.	1.6	2
167	Unilateral nephrectomy for young infants with congenital nephrotic syndrome of the Finnish type. Clinical and Experimental Nephrology, 2022, 26, 162-169.	1.6	2
168	Heterozygous Urinary Abnormality–Causing Variants of COL4A3 and COL4A4 Affect Severity of Autosomal Recessive Alport Syndrome. Kidney360, 2020, 1, 936-942.	2.1	2
169	High serum cystatin C levels in juvenile myelomonocytic leukemia patients without abnormal kidney function. Pediatric Nephrology, 2022, 37, 1687-1691.	1.7	2
170	Long-term follow-up of juvenile acute nonproliferative glomerulitis (JANG). Pediatric Nephrology, 2007, 22, 1957-1961.	1.7	1
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