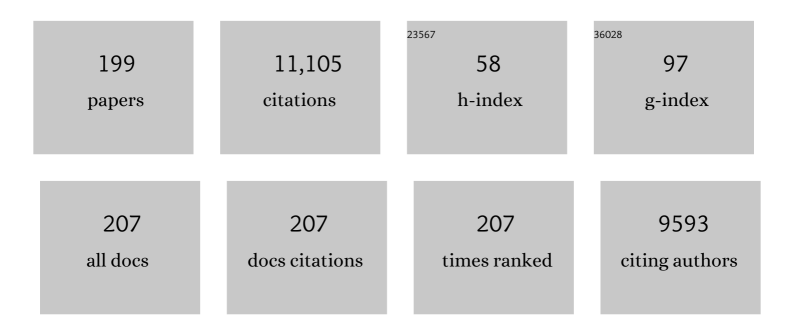
## Michael A Levine

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Genetic evidence that the human CYP2R1 enzyme is a key vitamin D 25-hydroxylase. Proceedings of the National Academy of Sciences of the United States of America, 2004, 101, 7711-7715.	7.1	630
2	Mutation in the Gene Encoding the Stimulatory G Protein of Adenylate Cyclase in Albright's Hereditary Osteodystrophy. New England Journal of Medicine, 1990, 322, 1412-1419.	27.0	396
3	FGF-23 Inhibits Renal Tubular Phosphate Transport and Is a PHEX Substrate. Biochemical and Biophysical Research Communications, 2001, 284, 977-981.	2.1	320
4	Paternally Inherited Inactivating Mutations of the <i>GNAS1</i> Gene in Progressive Osseous Heteroplasia. New England Journal of Medicine, 2002, 346, 99-106.	27.0	284
5	Resistance to multiple hormones in patients with pseudohypoparathyroidism. American Journal of Medicine, 1983, 74, 545-556.	1.5	277
6	Evaluating Children With Fractures for Child Physical Abuse. Pediatrics, 2014, 133, e477-e489.	2.1	232
7	Epidemiology and Diagnosis of Hypoparathyroidism. Journal of Clinical Endocrinology and Metabolism, 2016, 101, 2284-2299.	3.6	230
8	Diagnosis and management of pseudohypoparathyroidism and related disorders: first international Consensus Statement. Nature Reviews Endocrinology, 2018, 14, 476-500.	9.6	224
9	Differential Susceptibility to Hypertension Is Due to Selection during the Out-of-Africa Expansion. PLoS Genetics, 2005, 1, e82.	3.5	208
10	Localisation of mesenchymal tumours by somatostatin receptor imaging. Lancet, The, 2002, 359, 761-763.	13.7	198
11	Familial isolated hypoparathyroidism caused by a mutation in the gene for the transcription factor GCMB. Journal of Clinical Investigation, 2001, 108, 1215-1220.	8.2	183
12	Body Mass Index Differences in Pseudohypoparathyroidism Type 1a <i>Versus</i> Pseudopseudohypoparathyroidism May Implicate Paternal Imprinting of Gαs in the Development of Human Obesity. Journal of Clinical Endocrinology and Metabolism, 2007, 92, 1073-1079.	3.6	181
13	An Association between Neonatal Severe Primary Hyperparathyroidism and Familial Hypocalciuric Hypercalcemia in Three Kindreds. New England Journal of Medicine, 1982, 306, 257-264.	27.0	174
14	Diagnosis and Management of Osteopetrosis: Consensus Guidelines From the Osteopetrosis Working Group. Journal of Clinical Endocrinology and Metabolism, 2017, 102, 3111-3123.	3.6	170
15	Growth Hormone Deficiency in Pseudohypoparathyroidism Type 1a: Another Manifestation of Multihormone Resistance. Journal of Clinical Endocrinology and Metabolism, 2003, 88, 4059-4069.	3.6	156
16	A multinational study to develop universal standardization of whole-body bone density and composition using GE Healthcare Lunar and Hologic DXA systems. Journal of Bone and Mineral Research, 2012, 27, 2208-2216.	2.8	150
17	Characterization of GATA3 Mutations in the Hypoparathyroidism, Deafness, and Renal Dysplasia (HDR) Syndrome. Journal of Biological Chemistry, 2004, 279, 22624-22634.	3.4	145
18	Clinical Implications of Genetic Defects in G Proteins: The Molecular Basis of McCune-Albright Syndrome and Albright Hereditary Osteodystrophy. Medicine (United States), 1996, 75, 171-184.	1.0	144

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19	Paternal imprinting of Gαs in the human thyroid as the basis of TSH resistance in pseudohypoparathyroidism type 1a. Biochemical and Biophysical Research Communications, 2002, 296, 67-72.	2.1	141
20	Activity of the Stimulatory Guanine Nucleotide-Binding Protein Is Reduced in Erythrocytes from Patients with Pseudohypoparathyroidism and Pseudopseudohypoparathyroidism: Biochemical, Endocrine, and Genetic Analysis of Albright's Hereditary Osteodystrophy in Six Kindreds*. Journal of Clinical Endocrinology and Metabolism, 1986, 62, 497-502.	3.6	139
21	Cherubism: best clinical practice. Orphanet Journal of Rare Diseases, 2012, 7, S6.	2.7	138
22	ARAF recurrent mutation causes central conducting lymphatic anomaly treatable with a MEK inhibitor. Nature Medicine, 2019, 25, 1116-1122.	30.7	136
23	Rat cerebral cortex corticotropin-releasing hormone receptors: evidence for receptor coupling to multiple G-proteins. Journal of Neurochemistry, 2001, 76, 509-519.	3.9	135
24	Hypocalcemia in the Critically III patient. Journal of Intensive Care Medicine, 2013, 28, 166-177.	2.8	134
25	Clinical Implications of Guanine Nucleotide–Binding Proteins as Receptor–Effector Couplers. New England Journal of Medicine, 1985, 312, 26-33.	27.0	130
26	Mutations in SLC34A3/NPT2c Are Associated with Kidney Stones and Nephrocalcinosis. Journal of the American Society of Nephrology: JASN, 2014, 25, 2366-2375.	6.1	124
27	A Mouse Model of Albright Hereditary Osteodystrophy Generated by Targeted Disruption of Exon 1 of the Gnas Gene. Endocrinology, 2005, 146, 4697-4709.	2.8	122
28	Maximal Urine-Concentrating Ability: Familial Hypocalciuric Hypercalcemia <i>Versus</i> Typical Primary Hyperparathyroidism. Journal of Clinical Endocrinology and Metabolism, 1981, 52, 736-740.	3.6	115
29	Familial Hypocalciuric Hypercalcemia. New England Journal of Medicine, 1982, 307, 416-426.	27.0	105
30	Perinatal calcium metabolism: physiology and pathophysiology. Seminars in Fetal and Neonatal Medicine, 2004, 9, 23-36.	2.7	102
31	Hypercalcemia in children and adolescents. Current Opinion in Pediatrics, 2010, 22, 508-515.	2.0	101
32	Obesity Decreases Hepatic 25-Hydroxylase Activity Causing Low Serum 25-Hydroxyvitamin D. Journal of Bone and Mineral Research, 2019, 34, 1068-1073.	2.8	100
33	Albright hereditary osteodystrophy and del(2)(q37.3) in four unrelated individuals. American Journal of Medical Genetics Part A, 1995, 58, 1-7.	2.4	98
34	An update on the clinical and molecular characteristics of pseudohypoparathyroidism. Current Opinion in Endocrinology, Diabetes and Obesity, 2012, 19, 443-451.	2.3	94
35	<i>CYP2R1</i> Mutations Impair Generation of 25-hydroxyvitamin D and Cause an Atypical Form of Vitamin D Deficiency. Journal of Clinical Endocrinology and Metabolism, 2015, 100, E1005-E1013.	3.6	94
36	Mapping of the gene encoding the α subunit of the stimulatory G protein of adenylyl cyclase (GNAS1) to 20q13.2 → q13.3 in human by in situ hybridization. Genomics, 1991, 11, 478-479.	2.9	92

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37	Primary hyperparathyroidism in children and adolescents. Journal of the Chinese Medical Association, 2012, 75, 425-434.	1.4	89
38	Suppression of Sclerostin Alleviates Radiation-Induced Bone Loss by Protecting Bone-Forming Cells and Their Progenitors Through Distinct Mechanisms. Journal of Bone and Mineral Research, 2017, 32, 360-372.	2.8	88
39	Coupling of the PTH/PTHrP Receptor to Multiple G-Proteins: Direct Demonstration of Receptor Activation of G <sub>s</sub> , G <sub>q/11</sub> , and G <sub>i(1)</sub> by [α - <sup>32</sup> P]GTP-γ -Azidoanilide Photoaffinity Labeling. Endocrine, 1998, 8, 201-210.	2.2	86
40	National Health and Nutrition Examination Survey Whole-Body Dual-Energy X-Ray Absorptiometry Reference Data for GE Lunar Systems. Journal of Clinical Densitometry, 2014, 17, 344-377.	1.2	83
41	Vitamin D Status in Abused and Nonabused Children Younger Than 2 Years Old With Fractures. Pediatrics, 2011, 127, 835-841.	2.1	82
42	Autosomal Dominant Hypoparathyroidism Caused by Germline Mutation in <i>GNA11</i> : Phenotypic and Molecular Characterization. Journal of Clinical Endocrinology and Metabolism, 2014, 99, E1774-E1783.	3.6	79
43	Chromosomal localization of the genes encoding two forms of the G protein β polypeptide, β1 and β3, in man. Genomics, 1990, 8, 380-386.	2.9	77
44	The role of SH3BP2 in the pathophysiology of cherubism. Orphanet Journal of Rare Diseases, 2012, 7, S5.	2.7	77
45	CYP3A4 mutation causes vitamin D–dependent rickets type 3. Journal of Clinical Investigation, 2018, 128, 1913-1918.	8.2	77
46	Olfactory dysfunction in humans with deficient guanine nucleotide-binding protein. Nature, 1986, 322, 635-636.	27.8	75
47	Pathogenic variant in EPHB4 results in central conducting lymphatic anomaly. Human Molecular Genetics, 2018, 27, 3233-3245.	2.9	73
48	CYP3A4 Induction by Rifampin: An Alternative Pathway for Vitamin D Inactivation in Patients With CYP24A1 Mutations. Journal of Clinical Endocrinology and Metabolism, 2017, 102, 1440-1446.	3.6	72
49	Cinacalcet Monotherapy in Neonatal Severe Hyperparathyroidism: A Case Study and Review. Journal of Clinical Endocrinology and Metabolism, 2014, 99, 7-11.	3.6	71
50	Mutations in the ABCC6 Gene as a Cause of Generalized Arterial Calcification of Infancy: Genotypic Overlap with Pseudoxanthoma Elasticum. Journal of Investigative Dermatology, 2014, 134, 658-665.	0.7	70
51	Regulation of Corticotropin-Releasing Hormone Receptor Type 1α Signaling: Structural Determinants for G Protein-Coupled Receptor Kinase-Mediated Phosphorylation and Agonist-Mediated Desensitization. Molecular Endocrinology, 2005, 19, 474-490.	3.7	68
52	Clinical Implications of Genetic Defects in G Proteins. Archives of Medical Research, 1999, 30, 522-531.	3.3	67
53	Consensus Development for the Supplementation of Vitamin D in Childhood and Adolescence. Hormone Research in Paediatrics, 2002, 58, 39-51.	1.8	66
54	Selective Resistance to Parathyroid Hormone Caused by a Novel Uncoupling Mutation in the Carboxyl Terminus of Gαs. Journal of Biological Chemistry, 2001, 276, 165-171.	3.4	65

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55	Directional memory arises from long-lived cytoskeletal asymmetries in polarized chemotactic cells. Proceedings of the National Academy of Sciences of the United States of America, 2016, 113, 1267-1272.	7.1	65
56	Infantile hypothyroidism in two sibs: An unusual presentation of pseudohypoparathyroidism type Ia. Journal of Pediatrics, 1985, 107, 919-922.	1.8	60
57	Immunochemical Analysis of the <i>î±</i> -Subunit of the Stimulatory C-Protein of Adenylyl Cyclase in Patients with Albright's Hereditary Osteodystrophy*. Journal of Clinical Endocrinology and Metabolism, 1990, 71, 1208-1214.	3.6	59
58	Recombinant Human Parathyroid Hormone Effect on Health-Related Quality of Life in Adults With Chronic Hypoparathyroidism. Journal of Clinical Endocrinology and Metabolism, 2018, 103, 722-731.	3.6	59
59	Pseudohypoparathyroidism. Endocrinology and Metabolism Clinics of North America, 2018, 47, 865-888.	3.2	59
60	Thresholds for Surgery and Surgical Outcomes for Patients with Primary Hyperparathyroidism: A National Survey of Endocrine Surgeons. Journal of Clinical Endocrinology and Metabolism, 1998, 83, 2658-2665.	3.6	59
61	Hormonal Tolerance to Ethanol is Associated with Decreased Expression of the GTP-Binding Protein, Gsalpha, and Adenylyl Cyclase Activity in Ethanol-Treated LS Mice. Alcoholism: Clinical and Experimental Research, 1991, 15, 705-710.	2.4	58
62	A Novel Loss-of-Function Mutation, Gln459Arg, of the Calcium-Sensing Receptor Gene Associated with Apparent Autosomal Recessive Inheritance of Familial Hypocalciuric Hypercalcemia. Journal of Clinical Endocrinology and Metabolism, 2009, 94, 4372-4379.	3.6	58
63	Hypophosphatemic Rickets with Hypercalciuria due to Mutation in <i>SLC34A3</i> /Type IIc Sodium-Phosphate Cotransporter: Presentation as Hypercalciuria and Nephrolithiasis. Journal of Clinical Endocrinology and Metabolism, 2009, 94, 4433-4438.	3.6	57
64	Short-Term Safety of Zoledronic Acid in Young Patients With Bone Disorders: An Extensive Institutional Experience. Journal of Clinical Endocrinology and Metabolism, 2015, 100, 4163-4171.	3.6	57
65	A new missense mutation in the growth hormone-releasing hormone receptor gene in familial isolated CH deficiency. European Journal of Endocrinology, 2003, 148, 25-30.	3.7	55
66	Probing the Bimolecular Interactions of Parathyroid Hormone and the Human Parathyroid Hormone/Parathyroid Hormone-Related Protein Receptor. 2. Cloning, Characterization, and Photoaffinity Labeling of the Recombinant Human Parathyroid Hormone/Parathyroid Hormone-Related Protein Receptor. Biochemistry, 1995, 34, 10553-10559.	2.5	53
67	The etiology and significance of fractures in infants and young children: a critical multidisciplinary review. Pediatric Radiology, 2016, 46, 591-600.	2.0	52
68	CYP2R1 mutations causing vitamin D-deficiency rickets. Journal of Steroid Biochemistry and Molecular Biology, 2017, 173, 333-336.	2.5	52
69	Identification of Signaling Molecules Mediating Corticotropin-Releasing Hormone-R1α-Mitogen-Activated Protein Kinase (MAPK) Interactions: The Critical Role of Phosphatidylinositol 3-Kinase in Regulating ERK1/2 But Not p38 MAPK Activation. Molecular Endocrinology. 2006. 20. 3179-3195.	3.7	51
70	A phase I study of cediranib in combination with cilengitide in patients with recurrent glioblastoma. Neuro-Oncology, 2015, 17, 1386-1392.	1.2	50
71	Multidisciplinary guidelines for initial evaluation of complicated lymphatic anomalies—expert opinion consensus. Pediatric Blood and Cancer, 2020, 67, e28036.	1.5	50
72	Inhibition of Clucose-Stimulated Insulin Release in the Perfused Rat Pancreas by Parathyroid Secretory Protein-I (Chromogranin-A)*. Endocrinology, 1989, 124, 1235-1238.	2.8	48

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73	Protein Kinase A-Induced Negative Regulation of the Corticotropin-Releasing Hormone R1α Receptor-Extracellularly Regulated Kinase Signal Transduction Pathway: The Critical Role of Ser301for Signaling Switch and Selectivity. Molecular Endocrinology, 2004, 18, 624-639.	3.7	48
74	Comparison of metabolism of vitamins D2 and D3 in children with nutritional rickets. Journal of Bone and Mineral Research, 2010, 25, 1988-1995.	2.8	48
75	Is McCune-Albright syndrome overlooked in subjects with fibrous dysplasia of bone?. Journal of Pediatrics, 2003, 142, 532-538.	1.8	47
76	Isolated growth hormone (GH) deficiency due to compound heterozygosity for two new mutations in the GHâ€releasing hormone receptor gene. Clinical Endocrinology, 2001, 54, 681-687.	2.4	46
77	Discordance between Genetic and Epigenetic Defects in Pseudohypoparathyroidism Type 1b Revealed by Inconsistent Loss of Maternal Imprinting at GNAS1. American Journal of Human Genetics, 2003, 73, 314-322.	6.2	46
78	Reduction in Gs?? Induces Osteogenic Differentiation in Human Mesenchymal Stem Cells. Clinical Orthopaedics and Related Research, 2005, &NA, 231-238.	1.5	46
79	Safety and Efficacy of 5 Years of Treatment With Recombinant Human Parathyroid Hormone in Adults With Hypoparathyroidism. Journal of Clinical Endocrinology and Metabolism, 2019, 104, 5136-5147.	3.6	46
80	Inhibition of Tissue-Nonspecific Alkaline Phosphatase Attenuates Ectopic Mineralization in the Abcc6 Mouse Model of PXE but Not in the Enpp1 Mutant Mouse Models of GACI. Journal of Investigative Dermatology, 2019, 139, 360-368.	0.7	46
81	Enhanced Expression of the Inhibitory Protein Gi2alpha and Decreased Activity of Adenylyl Cyclase in Lymphocytes of Abstinent Alcoholics. Alcoholism: Clinical and Experimental Research, 1993, 17, 315-320.	2.4	44
82	Analysis of the <i>GCM2</i> Gene in Isolated Hypoparathyroidism: A Molecular and Biochemical Study. Journal of Clinical Endocrinology and Metabolism, 2008, 93, 1426-1432.	3.6	44
83	Genetic Basis for Resistance to Parathyroid Hormone. Hormone Research in Paediatrics, 2003, 60, 87-95.	1.8	43
84	A Highly Sensitive Polymerase Chain Reaction Method Detects Activating Mutations of the <it>GNAS</it> Gene in Peripheral Blood Cells in McCune-Albright Syndrome or Isolated Fibrous Dysplasia. Journal of Bone and Joint Surgery - Series A, 2005, 87, 2489.	3.0	41
85	Resting Energy Expenditure Is Decreased in Pseudohypoparathyroidism Type 1A. Journal of Clinical Endocrinology and Metabolism, 2016, 101, 880-888.	3.6	41
86	MRâ€assisted PET motion correction in simultaneous PET/MRI studies of dementia subjects. Journal of Magnetic Resonance Imaging, 2018, 48, 1288-1296.	3.4	41
87	Inherited non-alcoholic fatty liver disease and dyslipidemia due to monoallelic ABHD5 mutations. Journal of Hepatology, 2019, 71, 366-370.	3.7	41
88	Madelung-Like Deformity in Pseudohypoparathyroidism Type 1b. Journal of Clinical Endocrinology and Metabolism, 2011, 96, E1507-E1511.	3.6	40
89	Dual Effects of Bisphosphonates onÂEctopicÂSkin and Vascular Soft TissueÂMineralization versus Bone Microarchitecture in a Mouse Model of Generalized Arterial Calcification of Infancy. Journal of Investigative Dermatology, 2016, 136, 275-283.	0.7	40
90	Bone Mineral Density in Pseudohypoparathyroidism Type 1a. Journal of Clinical Endocrinology and Metabolism, 2010, 95, 4465-4475.	3.6	38

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91	Pseudohypoparathyroidism: From Bedside to Bench and Back. Journal of Bone and Mineral Research, 1999, 14, 1255-1260.	2.8	37
92	Differential Responses of Corticotropin-Releasing Hormone Receptor Type 1 Variants to Protein Kinase C Phosphorylation. Journal of Pharmacology and Experimental Therapeutics, 2006, 319, 1032-1042.	2.5	37
93	SH3BP2 is an activator of NFAT activity and osteoclastogenesis. Biochemical and Biophysical Research Communications, 2008, 371, 644-648.	2.1	37
94	Cost implications of different surgical management strategies for primary hyperparathyroidism. Surgery, 1998, 124, 1028-1036.	1.9	36
95	Digenic Inheritance of PROKR2 and WDR11 Mutations in Pituitary Stalk Interruption Syndrome. Journal of Clinical Endocrinology and Metabolism, 2017, 102, 2501-2507.	3.6	36
96	Isolation and Characterization of Myostatin Complementary Deoxyribonucleic Acid Clones from Two Commercially Important Fish: Oreochromis mossambicus and Morone chrysops. Endocrinology, 2001, 142, 1412-1418.	2.8	36
97	A Meta-Analysis Comparing the Biochemistry of Primary Hyperparathyroidism in Youths to the Biochemistry of Primary Hyperparathyroidism in Adults. Journal of Clinical Endocrinology and Metabolism, 2014, 99, 4555-4564.	3.6	35
98	Exome Sequencing Reveals Mutations in AIRE as a Cause of Isolated Hypoparathyroidism. Journal of Clinical Endocrinology and Metabolism, 2017, 102, 1726-1733.	3.6	35
99	Heterotopic Ossifications in a Mouse Model of Albright Hereditary Osteodystrophy. PLoS ONE, 2011, 6, e21755.	2.5	34
100	The McCune–Albright Syndrome. New England Journal of Medicine, 1991, 325, 1738-1740.	27.0	32
101	Risk factors for reduced skin thickness and bone density: Possible clues regarding pathophysiology, prevention, and treatment. Journal of the American Academy of Dermatology, 1998, 38, 248-255.	1.2	32
102	Expression of GCMB by Intrathymic Parathyroid Hormone-Secreting Adenomas Indicates Their Parathyroid Cell Origin. Journal of Clinical Endocrinology and Metabolism, 2004, 89, 8-12.	3.6	31
103	Late-onset hereditary hypophosphatemic rickets with hypercalciuria (HHRH) due to mutation of SLC34A3/NPT2c. Bone, 2017, 97, 15-19.	2.9	30
104	Effects of pravastatin, a new HMG-CoA reductase inhibitor, on vitamin D synthesis in man. Metabolism: Clinical and Experimental, 1991, 40, 524-528.	3.4	29
105	Genetic Disorders of Parathyroid Development and Function. Endocrinology and Metabolism Clinics of North America, 2018, 47, 809-823.	3.2	29
106	Structural Determinants Critical for Localization and Signaling within the Seventh Transmembrane Domain of the Type 1 Corticotropin Releasing Hormone Receptor: Lessons from the Receptor Variant R1d. Molecular Endocrinology, 2008, 22, 2505-2519.	3.7	28
107	Mendelian randomization analysis demonstrates that low vitamin D is unlikely causative for pediatric asthma. Journal of Allergy and Clinical Immunology, 2016, 138, 1747-1749.e4.	2.9	28
108	Intraoperative Measurements of Urinary Cyclic Amp to Guide Surgery for Primary Hyperparathyroidism. New England Journal of Medicine, 1980, 303, 1457-1460.	27.0	27

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109	McCune-Albright syndrome. Trends in Endocrinology and Metabolism, 1993, 4, 238-242.	7.1	27
110	Absence of mutations in the growth hormone (GH)-releasing hormone receptor gene in GH-secreting pituitary adenomas. Clinical Endocrinology, 2001, 54, 301-307.	2.4	26
111	Proteasome inhibitor bortezomib is a novel therapeutic agent for focal radiationâ€induced osteoporosis. FASEB Journal, 2018, 32, 52-62.	0.5	26
112	Etidronate prevents, but does not reverse, ectopic mineralization in a mouse model of pseudoxanthoma elasticum ( <i>Abcc6â^'/â^'</i> ). Oncotarget, 2018, 9, 30721-30730.	1.8	26
113	Association of Mutations in SLC12A1 Encoding the NKCC2 Cotransporter With Neonatal Primary Hyperparathyroidism. Journal of Clinical Endocrinology and Metabolism, 2016, 101, 2196-2200.	3.6	25
114	NAD+-mediated stimulation of adenylate cyclase in cardiac membranes. Biochemical and Biophysical Research Communications, 1987, 142, 631-637.	2.1	22
115	Structural domains determining signalling characteristics of the CRH-receptor type 1 variant R1Î <sup>2</sup> and response to PKC phosphorylation. Cellular Signalling, 2008, 20, 40-49.	3.6	22
116	A comprehensive study of long-term skeletal changes after spinal cord injury in adult rats. Bone Research, 2015, 3, 15028.	11.4	22
117	Low bone mineral density is a common finding in patients with homocystinuria. Molecular Genetics and Metabolism, 2016, 117, 351-354.	1.1	22
118	Decreased Expression of the GHRH Receptor Gene Due to a Mutation in a Pit-1 Binding Site. Molecular Endocrinology, 2002, 16, 450-458.	3.7	22
119	The Pseudohypoparathyroidism Type 1b Locus Is Linked to a Region Including <i>GNAS1</i> at 20q13.3. Journal of Bone and Mineral Research, 2003, 18, 424-433.	2.8	20
120	Thyroid-Specific Expression of Cholera Toxin A1 Subunit Causes Thyroid Hyperplasia and Hyperthyroidism in Transgenic Mice. Endocrinology, 1997, 138, 3133-3140.	2.8	20
121	Preimplantation Genetic Diagnosis for Severe Albright Hereditary Osteodystrophy. Journal of Clinical Endocrinology and Metabolism, 2008, 93, 901-904.	3.6	19
122	Outcomes of minimally invasive parathyroidectomy in pediatric patients with primary hyperparathyroidism owing to parathyroid adenoma: A single institution experience. Journal of Pediatric Surgery, 2017, 52, 188-191.	1.6	19
123	Teriparatide as a Systemic Treatment for Lower Extremity Nonunion Fractures: A Case Series. Endocrine Practice, 2015, 21, 136-142.	2.1	18
124	Digenic Heterozygous Mutations in SLC34A3 and SLC34A1 Cause Dominant Hypophosphatemic Rickets with Hypercalciuria. Journal of Clinical Endocrinology and Metabolism, 2020, 105, 2392-2400.	3.6	18
125	Rapid parathyroid hormone measurement during venous localization. Clinica Chimica Acta, 2000, 295, 193-198.	1.1	17
126	SH3BP2 Is Rarely Mutated in Exon 9 in Giant Cell Lesions Outside Cherubism. Clinical Orthopaedics and Related Research, 2007, 459, 22-27.	1.5	17

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127	Determination of Reference Intervals for Serum Total Calcium in the Vitamin D-Replete Pediatric Population. Journal of Clinical Endocrinology and Metabolism, 2013, 98, E1946-E1950.	3.6	17
128	Premature Epiphyseal Closure of the Lower Extremities Contributing to Short Stature after <b><i>cis</i></b> -Retinoic Acid Therapy in Medulloblastoma: A Case Report. Hormone Research in Paediatrics, 2016, 85, 69-73.	1.8	17
129	Decreased Serum 25-Hydroxyvitamin D in Aging Male Mice Is Associated With Reduced Hepatic Cyp2r1 Abundance. Endocrinology, 2018, 159, 3083-3089.	2.8	17
130	Bones and Joints: The Effects of Cannabinoids on the Skeleton. Journal of Clinical Endocrinology and Metabolism, 2019, 104, 4683-4694.	3.6	17
131	25-Hydroxyvitamin D Can Interfere With a Common Assay for 1,25-Dihydroxyvitamin D in Vitamin D Intoxication. Journal of Clinical Endocrinology and Metabolism, 2015, 100, 2883-2889.	3.6	16
132	Single Gland, Ectopic Location: Adenomas are Common Causes of Primary Hyperparathyroidism in Children and Adolescents. World Journal of Surgery, 2020, 44, 1518-1525.	1.6	16
133	Balanced rearrangement of chromosomes 2, 5, and 13 in a family with duplication 5q and fetal loss. American Journal of Medical Genetics Part A, 1984, 19, 783-790.	2.4	15
134	Persistent Hypercalcemia After Parathyroidectomy in an Adolescent and Effect of Treatment With Cinacalcet HCl. Clinical Chemistry, 2006, 52, 2286-2293.	3.2	15
135	Transmission imaging for integrated PET-MR systems. Physics in Medicine and Biology, 2016, 61, 5547-5568.	3.0	15
136	Heterozygous Mutations in TBX1 as a Cause of Isolated Hypoparathyroidism. Journal of Clinical Endocrinology and Metabolism, 2018, 103, 4023-4032.	3.6	15
137	Short Stature is Progressive in Patients with Heterozygous NPR2 Mutations. Journal of Clinical Endocrinology and Metabolism, 2020, 105, 3190-3202.	3.6	15
138	Stress alters adenylyl cyclase activity in the pituitary and frontal cortex of the rat. Life Sciences, 1993, 53, 1719-1727.	4.3	14
139	A novel mutation in the GCM2 gene causing severe congenital isolated hypoparathyroidism. Journal of Pediatric Endocrinology and Metabolism, 2012, 25, 741-6.	0.9	14
140	Mapping Structural Determinants within Third Intracellular Loop That Direct Signaling Specificity of Type 1 Corticotropin-releasing Hormone Receptor. Journal of Biological Chemistry, 2012, 287, 8974-8985.	3.4	14
141	Ketotic Hypercalcemia: A Case Series and Description of a Novel Entity. Journal of Clinical Endocrinology and Metabolism, 2014, 99, 1531-1536.	3.6	14
142	Generation of mice encoding a conditional null allele of Gcm2. Transgenic Research, 2014, 23, 631-641.	2.4	13
143	Exome sequencing reveals a nonsense mutation in MMP13 as a new cause of autosomal recessive metaphyseal anadysplasia. European Journal of Human Genetics, 2015, 23, 264-266.	2.8	13
144	ENPP1 variants in patients with GACI and PXE expand the clinical and genetic heterogeneity of heritable disorders of ectopic calcification. PLoS Genetics, 2022, 18, e1010192.	3.5	13

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145	Reduced adenylyl cyclase activation with no decrease in β-adrenergic receptors in basenji greyhound leukocytes: Relevance to β-adrenergic responses in airway smooth muscle. Journal of Allergy and Clinical Immunology, 1995, 95, 860-867.	2.9	12
146	Gestational Gigantomastia Complicated by PTHrP-Mediated Hypercalcemia. Journal of Clinical Endocrinology and Metabolism, 2018, 103, 3124-3130.	3.6	12
147	High-throughput Molecular Analysis of Pseudohypoparathyroidism 1b Patients Reveals Novel Genetic and Epigenetic Defects. Journal of Clinical Endocrinology and Metabolism, 2021, 106, e4603-e4620.	3.6	12
148	Hypocalcemia in Nonwhite Breast-Fed Infants. Clinical Pediatrics, 1992, 31, 695-698.	0.8	11
149	Three Novel Mutations in the PHEX Gene in Chinese Subjects with Hypophosphatemic Rickets Extends Genotypic Variability. Calcified Tissue International, 2011, 88, 370-377.	3.1	11
150	Molecular Basis of Primary Hyperparathyroidism. , 2015, , 279-296.		11
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