## Feng Zhang

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/3051938/publications.pdf

Version: 2024-02-01

249 papers 25,678 citations

51 h-index 150 g-index

252 all docs

252 docs citations

times ranked

252

38970 citing authors

#	Article	IF	CITATIONS
1	Homozygous variants in <i>AKAP3</i> induce asthenoteratozoospermia and male infertility. Journal of Medical Genetics, 2023, 60, 137-143.	3.2	9
2	Haploinsufficiency in non-homologous end joining factor $1$ induces ovarian dysfunction in humans and mice. Journal of Medical Genetics, 2022, 59, 579-588.	3.2	3
3	Homozygous mutations in <i>CCDC34</i> cause male infertility with oligoasthenoteratozoospermia in humans and mice. Journal of Medical Genetics, 2022, 59, 710-718.	3.2	20
4	Rice GLUTATHIONE PEROXIDASE1-mediated oxidation of bZIP68 positively regulates ABA-independent osmotic stress signaling. Molecular Plant, 2022, 15, 651-670.	8.3	20
5	Homozygous mutation in SLO3 leads to severe asthenoteratozoospermia due to acrosome hypoplasia and mitochondrial sheath malformations. Reproductive Biology and Endocrinology, 2022, 20, 5.	3.3	11
6	Chromosome-Level Genome Assembly of Anthidium xuezhongi Niu & Zhu, 2020 (Hymenoptera: Apoidea:) Tj ETQ	q0 <u>0</u> 00 rgB	T <u> </u> Overlock 1
7	<i>HSD17B12</i> dosage insufficiency induced premature ovarian insufficiency in humans and mice. Clinical and Translational Medicine, 2022, 12, e737.	4.0	O
8	CEP128 is involved in spermatogenesis in humans and mice. Nature Communications, 2022, 13, 1395.	12.8	23
9	Mitogenomic features of three species of Entomobryoidea (Collembola) from China. Zootaxa, 2022, 5120, 283-288.	0.5	1
10	Testing the systematic status of <i>Homalictus</i> and <i>Rostrohalictus</i> with weakened crossâ€vein groups within Halictini (Hymenoptera: Halictidae) using lowâ€coverage wholeâ€genome sequencing. Insect Science, 2022, 29, 1819-1833.	3.0	7
11	Deficiency of X-linked TENT5D causes male infertility by disrupting the mRNA stability during spermatogenesis. Cell Discovery, 2022, 8, 23.	6.7	12
12	Phylogenomics of Elongate-Bodied Springtails Reveals Independent Transitions from Aboveground to Belowground Habitats in Deep Time. Systematic Biology, 2022, 71, 1023-1031.	5.6	10
13	A recurrent SHANK1 mutation implicated in autism spectrum disorder causes autistic-like core behaviors in mice via downregulation of mGluR1-IP3R1-calcium signaling. Molecular Psychiatry, 2022, 27, 2985-2998.	7.9	12
14	Molecular mechanisms underlying cTAGE5/MEA6-mediated cargo transport and biological functions. Journal of Genetics and Genomics, 2022, 49, 519-522.	3.9	2
15	Human pathogenic RNA viruses establish noncompeting lineages by occupying independent niches. Proceedings of the National Academy of Sciences of the United States of America, 2022, 119, .	7.1	9
16	Homozygous Variant in <i>KASH5</i> Causes Premature Ovarian Insufficiency by Disordered Meiotic Homologous Pairing. Journal of Clinical Endocrinology and Metabolism, 2022, 107, 2589-2597.	3.6	7
17	Epigenetic features drastically impact CRISPR–Cas9 efficacy in plants. Plant Physiology, 2022, 190, 1153-1164.	4.8	25
18	Diagnostic yield and clinical impact of exome sequencing in early-onset scoliosis (EOS). Journal of Medical Genetics, 2021, 58, 41-47.	3.2	40

#	Article	IF	Citations
19	The coordination of guard-cell autonomous ABA synthesis and DES1 function in situ regulates plant water deficit responses. Journal of Advanced Research, 2021, 27, 191-197.	9.5	28
20	A heterozygous hypomorphic mutation of Fanca causes impaired follicle development and subfertility in female mice. Molecular Genetics and Genomics, 2021, 296, 103-112.	2.1	9
21	Novel lossâ€ofâ€function variants in <scp><i>DNAH17</i></scp> cause multiple morphological abnormalities of the sperm flagella in humans and mice. Clinical Genetics, 2021, 99, 176-186.	2.0	26
22	Whole exome sequencing and trio analysis to broaden the variant spectrum of genes in idiopathic hypogonadotropic hypogonadism. Asian Journal of Andrology, 2021, 23, 288.	1.6	5
23	Deleterious variants in X-linked CFAP47 induce asthenoteratozoospermia and primary male infertility. American Journal of Human Genetics, 2021, 108, 309-323.	6.2	74
24	Perturbations of genes essential for Mýllerian duct and Wölffian duct development in Mayer-Rokitansky-Küster-Hauser syndrome. American Journal of Human Genetics, 2021, 108, 337-345.	6.2	41
25	A chromosome-level genome of the spider <i>Trichonephila antipodiana</i> reveals the genetic basis of its polyphagy and evidence of an ancient whole-genome duplication event. GigaScience, 2021, 10, .	6.4	187
26	A mitogenomic phylogeny of the Entomobryoidea (Collembola): A comparative perspective. Zoologica Scripta, 2021, 50, 658-666.	1.7	5
27	Genome-wide CRISPR screens reveal synthetic lethal interaction between CREBBP and EP300 in diffuse large B-cell lymphoma. Cell Death and Disease, 2021, 12, 419.	6.3	21
28	Novel bi-allelic variants in DNAH2 cause severe asthenoteratozoospermia with multiple morphological abnormalities of the flagella. Reproductive BioMedicine Online, 2021, 42, 963-972.	2.4	19
29	Temporal transcriptomic landscape of postnatal mouse ovaries reveals dynamic gene signatures associated with ovarian aging. Human Molecular Genetics, 2021, 30, 1941-1954.	2.9	8
30	Loss of DRC1 function leads to multiple morphological abnormalities of the sperm flagella and male infertility in human and mouse. Human Molecular Genetics, 2021, 30, 1996-2011.	2.9	26
31	Novel Mutations in X-Linked, USP26-Induced Asthenoteratozoospermia and Male Infertility. Cells, 2021, 10, 1594.	4.1	6
32	CFAP65 is required in the acrosome biogenesis and mitochondrial sheath assembly during spermiogenesis. Human Molecular Genetics, 2021, 30, 2240-2254.	2.9	18
33	Bi-allelic truncating variants in CFAP206 cause male infertility in human and mouse. Human Genetics, 2021, 140, 1367-1377.	3.8	23
34	Genome mapping coupled with CRISPR gene editing reveals a P450 gene confers avermectin resistance in the beet armyworm. PLoS Genetics, 2021, 17, e1009680.	3.5	44
35	Ongoing global and regional adaptive evolution of SARS-CoV-2. Proceedings of the National Academy of Sciences of the United States of America, 2021, 118, .	7.1	196
36	Bi-allelic mutations of DNAH10 cause primary male infertility with asthenoteratozoospermia in humans and mice. American Journal of Human Genetics, 2021, 108, 1466-1477.	6.2	50

#	Article	IF	CITATIONS
37	First record of Seira dowlingi (Wray, 1953) (Collembola, Entomobryidae, Seirinae) from China and mitogenome comparison with the New World specimens. Zootaxa, 2021, 5020, 191-196.	0.5	4
38	Molecular basis of cerium oxide nanoparticle enhancement of rice salt tolerance and yield. Environmental Science: Nano, 2021, 8, 3294-3311.	4.3	36
39	A recurrent rare intronic variant in CAPN3 alters mRNA splicing and causes autosomal recessive limbâ€girdle muscular dystrophy‶ in three Pakistani pedigrees. American Journal of Medical Genetics, Part A, 2021, , .	1.2	0
40	Whole exome sequencing identified a rare WT1 lossâ€ofâ€function variant in a nonâ€syndromic POI patient. Molecular Genetics & Cenomic Medicine, 2021, , e1820.	1.2	2
41	Familial Translocation t(2;4) (q37.3;p16.3), Resulting in a Partial Trisomy of 2q (or 4p) and a Partial Monosomy of 4p (or 2q), Causes Dysplasia. Frontiers in Genetics, 2021, 12, 741607.	2.3	0
42	Persulfidation of Nitrate Reductase 2 Is Involved in I-Cysteine Desulfhydrase-Regulated Rice Drought Tolerance. International Journal of Molecular Sciences, 2021, 22, 12119.	4.1	18
43	Homozygous mutations in <i>SPEF2</i> ii>induce multiple morphological abnormalities of the sperm flagella and male infertility. Journal of Medical Genetics, 2020, 57, 31-37.	3.2	57
44	Biallelic mutations in <i>CFAP65 </i> cause male infertility with multiple morphological abnormalities of the sperm flagella in humans and mice. Journal of Medical Genetics, 2020, 57, 89-95.	3.2	55
45	Joint utilization of genetic analysis and semi-cloning technology reveals a digenic etiology of Müllerian anomalies. Cell Research, 2020, 30, 91-94.	12.0	10
46	Xq22 deletions and correlation with distinct neurological disease traits in females: Further evidence for a contiguous gene syndrome. Human Mutation, 2020, 41, 150-168.	2.5	15
47	The Nuclear Matrix Protein SAFB Cooperates with Major Satellite RNAs to Stabilize Heterochromatin Architecture Partially through Phase Separation. Molecular Cell, 2020, 77, 368-383.e7.	9.7	104
48	<i>TBX6</i> missense variants expand the mutational spectrum in a nonâ€Mendelian inheritance disease. Human Mutation, 2020, 41, 182-195.	2.5	27
49	MircroRNA-10b Promotes Human Embryonic Stem Cell-Derived Cardiomyocyte Proliferation via Novel Target Gene LATS1. Molecular Therapy - Nucleic Acids, 2020, 19, 437-445.	5.1	14
50	Hydrogen sulfide promotes rice drought tolerance via reestablishing redox homeostasis and activation of ABA biosynthesis and signaling. Plant Physiology and Biochemistry, 2020, 155, 213-220.	5.8	48
51	Rationally Designed APOBEC3B Cytosine Base Editors with Improved Specificity. Molecular Cell, 2020, 79, 728-740.e6.	9.7	104
52	Rare deleterious BUB1B variants induce premature ovarian insufficiency and early menopause. Human Molecular Genetics, 2020, 29, 2698-2707.	2.9	13
53	Bi-allelic Loss-of-function Variants in CFAP58 Cause Flagellar Axoneme and Mitochondrial Sheath Defects and Asthenoteratozoospermia in Humans and Mice. American Journal of Human Genetics, 2020, 107, 514-526.	6.2	71
54	Molecular structure and evolution mechanism of two populations of double minutes in human colorectal cancer cells. Journal of Cellular and Molecular Medicine, 2020, 24, 14205-14216.	3.6	9

#	Article	IF	CITATIONS
55	Unsupervised Inference of Developmental Directions for Single Cells Using VECTOR. Cell Reports, 2020, 32, 108069.	6.4	11
56	A <i>DNAH17</i> missense variant causes flagella destabilization and asthenozoospermia. Journal of Experimental Medicine, 2020, 217, .	8.5	88
57	Human Pluripotent Stem Cell-Derived Neural Cells and Brain Organoids Reveal SARS-CoV-2 Neurotropism Predominates in Choroid Plexus Epithelium. Cell Stem Cell, 2020, 27, 937-950.e9.	11.1	314
58	Disruption in <i>ACTL7A</i> causes acrosomal ultrastructural defects in human and mouse sperm as a novel male factor inducing early embryonic arrest. Science Advances, 2020, 6, eaaz4796.	10.3	50
59	Bi-allelic SHOC1 loss-of-function mutations cause meiotic arrest and non-obstructive azoospermia. Journal of Medical Genetics, 2020, 58, jmedgenet-2020-107042.	3.2	20
60	The First Draft Genome of the Plasterer Bee Colletes gigas (Hymenoptera: Colletidae: Colletes). Genome Biology and Evolution, 2020, 12, 860-866.	2.5	12
61	Homology of labial chaetae in Entomobryoidea (Collembola). Zootaxa, 2020, 4766, zootaxa.4766.3.8.	0.5	6
62	Bi-allelic DNAH8 Variants Lead to Multiple Morphological Abnormalities of the Sperm Flagella and Primary Male Infertility. American Journal of Human Genetics, 2020, 107, 330-341.	6.2	111
63	DVL mutations identified from human neural tube defects and Dandy-Walker malformation obstruct the Wnt signaling pathway. Journal of Genetics and Genomics, 2020, 47, 301-310.	3.9	6
64	Streamlining universal singleâ€copy orthologue and ultraconserved element design: A case study in Collembola. Molecular Ecology Resources, 2020, 20, 706-717.	4.8	10
65	Patients with severe asthenoteratospermia carrying SPAG6 or RSPH3 mutations have a positive pregnancy outcome following intracytoplasmic sperm injection. Journal of Assisted Reproduction and Genetics, 2020, 37, 829-840.	2.5	30
66	Mitochondrial genome of Brachystomella parvula (Collembola: Brachystomellidae). Mitochondrial DNA Part B: Resources, 2020, 5, 104-105.	0.4	1
67	Phylogeny of Neotropical Seirinae (Collembola, Entomobryidae) based on mitochondrial genomes. Zoologica Scripta, 2020, 49, 329-339.	1.7	11
68	A novel homozygous mutation in WDR19 induces disorganization of microtubules in sperm flagella and nonsyndromic asthenoteratospermia. Journal of Assisted Reproduction and Genetics, 2020, 37, 1431-1439.	2.5	35
69	Rare mutations in the autophagyâ€regulating gene <i>AMBRA1</i> contribute to human neural tube defects. Human Mutation, 2020, 41, 1383-1393.	2.5	15
70	A novel hemizygous loss-of-function mutation in ADGRG2 causes male infertility with congenital bilateral absence of the vas deferens. Journal of Assisted Reproduction and Genetics, 2020, 37, 1421-1429.	2.5	11
71	Different Gene Networks Are Disturbed by Zika Virus Infection in A Mouse Microcephaly Model. Genomics, Proteomics and Bioinformatics, 2020, 18, 737-748.	6.9	12
72	Human and mouse studies establish TBX6 in Mendelian CAKUT and as a potential driver of kidney defects associated with the 16p11.2 microdeletion syndrome. Kidney International, 2020, 98, 1020-1030.	5.2	17

#	Article	IF	CITATIONS
73	A fertile male with a single sY86 deletion on the Y chromosome. Asian Journal of Andrology, 2020, 22, 333.	1.6	2
74	Novel compound heterozygous variants in dynein axonemal heavy chain 17 cause asthenoteratospermia with sperm flagellar defects. Journal of Genetics and Genomics, 2020, 47, 713-717.	3.9	6
75	A putative rice l-cysteine desulfhydrase encodes a true l-cysteine synthase that regulates plant cadmium tolerance. Plant Growth Regulation, 2019, 89, 217-226.	3.4	20
76	The discovery of Neotropical Lepidosira (Collembola, Entomobryidae) and its systematic position. Zoologica Scripta, 2019, 48, 783-800.	1.7	6
77	Sequence characterization of RET in 117 Chinese Hirschsprung disease families identifies a large burden of de novo and parental mosaic mutations. Orphanet Journal of Rare Diseases, 2019, 14, 237.	2.7	10
78	Bi-allelic Mutations in TTC29 Cause Male Subfertility with Asthenoteratospermia in Humans and Mice. American Journal of Human Genetics, 2019, 105, 1168-1181.	6.2	62
79	Rare variants in FANCA induce premature ovarian insufficiency. Human Genetics, 2019, 138, 1227-1236.	3.8	56
80	A novel approach to remove the batch effect of single-cell data. Cell Discovery, 2019, 5, 46.	6.7	37
81	A High-quality Draft Genome Assembly of Sinella curviseta: A Soil Model Organism (Collembola). Genome Biology and Evolution, 2019, 11, 521-530.	2.5	13
82	Loss-of-function mutations in QRICH2 cause male infertility with multiple morphological abnormalities of the sperm flagella. Nature Communications, 2019, 10, 433.	12.8	108
83	Bi-allelic Mutations in ARMC2 Lead to Severe Astheno-Teratozoospermia Due to Sperm Flagellum Malformations in Humans and Mice. American Journal of Human Genetics, 2019, 104, 331-340.	6.2	113
84	A de novo mutation in DHD domain of SKI causing spina bifida with no craniofacial malformation or intellectual disability. American Journal of Medical Genetics, Part A, 2019, 179, 936-939.	1.2	3
85	Cytosine, but not adenine, base editors induce genome-wide off-target mutations in rice. Science, 2019, 364, 292-295.	12.6	491
86	<i>BRCA2</i> in Ovarian Development and Function. New England Journal of Medicine, 2019, 380, 1086-1087.	27.0	38
87	Bi-allelic Mutations in TTC21A Induce Asthenoteratospermia in Humans and Mice. American Journal of Human Genetics, 2019, 104, 738-748.	6.2	103
88	PhenoPro: a novel toolkit for assisting in the diagnosis of Mendelian disease. Bioinformatics, 2019, 35, 3559-3566.	4.1	27
89	Homozygous loss-of-function mutations in FSIP2 cause male infertility with asthenoteratospermia. Journal of Genetics and Genomics, 2019, 46, 53-56.	3.9	31
90	NovelCFAP43 and CFAP44 mutations cause male infertility with multiple morphological abnormalities of the sperm flagella (MMAF). Reproductive BioMedicine Online, 2019, 38, 769-778.	2.4	26

#	Article	IF	CITATIONS
91	Noncoding rare variants of TBX6 in congenital anomalies of the kidney and urinary tract. Molecular Genetics and Genomics, 2019, 294, 493-500.	2.1	8
92	Whole-exome sequencing reveals SALL4 variants in premature ovarian insufficiency: an update on genotype–phenotype correlations. Human Genetics, 2019, 138, 83-92.	3.8	27
93	A novel multiplex fluorescent competitive PCR for copy number variation detection. Genomics, 2019, 111, 1745-1751.	2.9	1
94	The copy number variation landscape of congenital anomalies of the kidney and urinary tract. Nature Genetics, 2019, 51, 117-127.	21,4	144
95	Novel homozygous <i>CFAP69</i> mutations in humans and mice cause severe asthenoteratospermia with multiple morphological abnormalities of the sperm flagella. Journal of Medical Genetics, 2019, 56, 96-103.	3.2	70
96	Phylogenomics from lowâ€coverage wholeâ€genome sequencing. Methods in Ecology and Evolution, 2019, 10, 507-517.	5.2	59
97	TBX6-associated congenital scoliosis (TACS) as a clinically distinguishable subtype of congenital scoliosis: further evidence supporting the compound inheritance and TBX6 gene dosage model. Genetics in Medicine, 2019, 21, 1548-1558.	2.4	60
98	TBX6 compound inheritance leads to congenital vertebral malformations in humans and mice. Human Molecular Genetics, 2019, 28, 539-547.	2.9	46
99	Biallelic mutations of CFAP251 cause sperm flagellar defects and human male infertility. Journal of Human Genetics, 2019, 64, 49-54.	2.3	56
100	Molecular phylogeny of Entomobrya (Collembola: Entomobryidae) from China: Color pattern groups and multiple origins. Insect Science, 2019, 26, 587-597.	3.0	15
101	Phenotypic heterogeneity of intellectual disability in patients with congenital insensitivity to pain with anhidrosis: A case report and literature review. Journal of International Medical Research, 2018, 46, 2445-2457.	1.0	11
102	Species delimitation in the morphologically conserved <i>Coecobrya</i> (Collembola:) Tj ETQq0 0 0 rgBT /Overlo taxonomy. Zoologica Scripta, 2018, 47, 342-356.	ck 10 Tf 50 1.7	0 307 Td (Ent 33
103	The nuclear matrix protein HNRNPU maintains 3D genome architecture globally in mouse hepatocytes. Genome Research, 2018, 28, 192-202.	5.5	91
104	RET somatic mutations are underrecognized in Hirschsprung disease. Genetics in Medicine, 2018, 20, 770-777.	2.4	24
105	Colouration, chaetotaxy and molecular data provide species-level resolution in a species complex of Dicranocentrus (Collembola: Entomobryidae). Invertebrate Systematics, 2018, 32, 1298.	1.3	9
106	Elevated microRNA-520d-5p in the serum of patients with Parkinson's disease, possibly through regulation of cereloplasmin expression. Neuroscience Letters, 2018, 687, 88-93.	2.1	14
107	Deletion of exon 4 in LAMA2 is the most frequent mutation in Chinese patients with laminin $\hat{l}\pm 2$ -related muscular dystrophy. Scientific Reports, 2018, 8, 14989.	3.3	17
108	cTAGE5/MEA6 plays a critical role in neuronal cellular components trafficking and brain development. Proceedings of the National Academy of Sciences of the United States of America, 2018, 115, E9449-E9458.	7.1	18

#	Article	IF	CITATIONS
109	The coexistence of copy number variations (CNVs) and single nucleotide polymorphisms (SNPs) at a locus can result in distorted calculations of the significance in associating SNPs to disease. Human Genetics, 2018, 137, 553-567.	3.8	57
110	A new species of Dicranocentrus SchÃ $\P$ tt from Hainan (China) with a key to the Chinese species of the genus (Collembola, Entomobryidae). ZooKeys, 2018, 762, 59-68.	1.1	3
111	A revision of the genus Lepidobrya Womersley (Collembola: Entomobryidae) based on morphology and sequence data of the genotype. Zootaxa, 2017, 4221, zootaxa.4221.5.2.	0.5	5
112	An Organismal CNV Mutator Phenotype Restricted to Early Human Development. Cell, 2017, 168, 830-842.e7.	28.9	66
113	Copy number variants of ABCF1, IL17REL, and FCGR3A are associated with the risk of gout. Protein and Cell, 2017, 8, 467-470.	11.0	11
114	Biallelic Mutations in CFAP43 and CFAP44 Cause Male Infertility with Multiple Morphological Abnormalities of the Sperm Flagella. American Journal of Human Genetics, 2017, 100, 854-864.	6.2	220
115	Efficient CNV breakpoint analysis reveals unexpected structural complexity and correlation of dosage-sensitive genes with clinical severity in genomic disorders. Human Molecular Genetics, 2017, 26, 1927-1941.	2.9	20
116	A single mutation in the prM protein of Zika virus contributes to fetal microcephaly. Science, 2017, 358, 933-936.	12.6	399
117	Delimiting species of Protaphorura (Collembola: Onychiuridae): integrative evidence based on morphology, DNA sequences and geography. Scientific Reports, 2017, 7, 8261.	3.3	16
118	CNVbase: Batch identification of novel and rare copy number variations based on multi-ethnic population data. Journal of Genetics and Genomics, 2017, 44, 367-370.	3.9	1
119	SPRINT: an SNP-free toolkit for identifying RNA editing sites. Bioinformatics, 2017, 33, 3538-3548.	4.1	64
120	A TBX5 3â€2UTR variant increases the risk of congenital heart disease in the Han Chinese population. Cell Discovery, 2017, 3, 17026.	6.7	23
121	<i>cTAGE5</i> deletion in pancreatic $\hat{l}^2$ cells impairs proinsulin trafficking and insulin biogenesis in mice. Journal of Cell Biology, 2017, 216, 4153-4164.	5.2	32
122	American Strain of Zika Virus Causes More Severe Microcephaly Than an Old Asian Strain in Neonatal Mice. EBioMedicine, 2017, 25, 95-105.	6.1	47
123	Assessing genome-wide copy number variation in the Han Chinese population. Journal of Medical Genetics, 2017, 54, 685-692.	3.2	7
124	Interaction between Y chromosome haplogroup O3* and 4-n-octylphenol exposure reduces the susceptibility to spermatogenic impairment in Han Chinese. Ecotoxicology and Environmental Safety, 2017, 144, 450-455.	6.0	3
125	Three new species of Coecobrya (Collembola: Entomobryidae) from caves in the Thai Peninsula. Zootaxa, 2017, 4286, 187.	0.5	10
126	Generation of special autosomal dominant polycystic kidney disease iPSCs with the capability of functional kidney-like cell differentiation. Stem Cell Research and Therapy, 2017, 8, 196.	5.5	4

#	Article	IF	Citations
127	New Australian Paronellidae (Collembola) reveal anomalies in existing tribal diagnoses. Invertebrate Systematics, 2017, 31, 375.	1.3	13
128	SIX2 haploinsufficiency causes conductive hearing loss with ptosis in humans. Journal of Human Genetics, 2016, 61, 917-922.	2.3	12
129	MAZ mediates the cross-talk between CT-1 and NOTCH1 signaling during gliogenesis. Scientific Reports, 2016, 6, 21534.	3.3	16
130	Dental scales could occur in all scaled subfamilies of Entomobryidae (Collembola): new definition of Entomobryinae with description of a new genus and three new species. Invertebrate Systematics, 2016, 30, 598.	1.3	9
131	A Novel c-Jun N-terminal Kinase (JNK) Signaling Complex Involved in Neuronal Migration during Brain Development. Journal of Biological Chemistry, 2016, 291, 11466-11475.	3.4	33
132	Copy Number Variation of HLA-DQA1 and APOBEC3A/3B Contribute to the Susceptibility of Systemic Sclerosis in the Chinese Han Population. Journal of Rheumatology, 2016, 43, 880-886.	2.0	4
133	New cave species of Sinella Brook, 1882 from China (Collembola: Entomobryidae). Zootaxa, 2016, 4161, 523.	0.5	1
134	Supplementary descriptive notes of the Sinella and Coecobrya (Collembola: Entomobryidae) species from North America, Hawaii and Japan. Zootaxa, 2016, 4085, 536-56.	0.5	0
135	The B-cell receptor BR3 modulates cellular branching via Rac1 during neuronal migration. Journal of Molecular Cell Biology, 2016, 8, 363-365.	3.3	1
136	New insight into the systematics of Tomoceridae (Hexapoda, Collembola) by integrating molecular and morphological evidence. Zoologica Scripta, 2016, 45, 286-299.	1.7	25
137	Mea6 controls VLDL transport through the coordinated regulation of COPII assembly. Cell Research, 2016, 26, 787-804.	12.0	34
138	Association of a TDRD1 variant with spermatogenic failure susceptibility in the Han Chinese. Journal of Assisted Reproduction and Genetics, 2016, 33, 1099-1104.	2.5	8
139	Autism-like behaviours and germline transmission in transgenic monkeys overexpressing MeCP2. Nature, 2016, 530, 98-102.	27.8	260
140	Two novel copy number variations involving the $\hat{l}_{\pm}$ -globin gene cluster on chromosome 16 cause thalassemia in two Chinese families. Molecular Genetics and Genomics, 2016, 291, 1443-1450.	2.1	22
141	Clinical and molecular genetic analysis of a family with late-onset LAMA2-related muscular dystrophy. Brain and Development, 2016, 38, 242-249.	1.1	16
142	Progress and perspective of <i>TBX6</i> gene in congenital vertebral malformations. Oncotarget, 2016, 7, 57430-57441.	1.8	24
143	New blind species and new records of Sinella from Nanjing, China (Collembola, Entomobryidae). ZooKeys, 2016, 604, 31-40.	1.1	2
144	Cave-dwelling Coecobrya from southern China with a survey of clypeal chaetae in Entomobryoidea (Collembola). European Journal of Taxonomy, 2016, , .	0.6	5

#	Article	IF	CITATIONS
145	Contribution to the Willowsia species having body scales of the long basal rib type: four new species and a redescription of W. qui (Collembola: Entomobryidae). European Journal of Taxonomy, 2016, , .	0.6	0
146	Genome-wide copy number variant analysis for congenital ventricular septal defects in Chinese Han population. BMC Medical Genomics, 2015, 9, 2.	1.5	20
147	Molecular phylogeny supports S-chaetae as a key character better than jumping organs and body scales in classification of Entomobryoidea (Collembola). Scientific Reports, 2015, 5, 12471.	3.3	27
148	MacroH2A1 associates with nuclear lamina and maintains chromatin architecture in mouse liver cells. Scientific Reports, 2015, 5, 17186.	3.3	44
149	Fifteen Novel EIF2B1-5 Mutations Identified in Chinese Children with Leukoencephalopathy with Vanishing White Matter and a Long Term Follow-Up. PLoS ONE, 2015, 10, e0118001.	2.5	17
150	<strong>A revision of <em>Pseudoparonella</em>, <em>Plumachaetas</em>, <em>Parachaetoceras</em> and <em>Lawrenceana</em> (Collembola: Paronellidae), with description of three new species from New Caledonia</strong> . Zootaxa, 2015, 4058, 561.	0.5	2
151	<strong>Some <em>Willowsia</em> from Nepal and Vietnam (Collembola: Entomobryidae) and description of one new species</strong> . Zootaxa, 2015, 3905, 489.	0.5	3
152	Two closely related Homidia species (Entomobryidae, Collembola) revealed by morphological and molecular evidence. Zootaxa, 2015, 3918, 285-94.	0.5	11
153	<strong>First instar tibiotarsal chaetotaxy supports the Entomobryidae and Symphypleona (Collembola) forming a cluster in a phylogenetic tree</strong> . Zootaxa, 2015, 3955, 487.	0.5	8
154	Contribution to the eyed Sinella from China: two new species and additional reports on nine known species (Collembola: Entomobryidae). Zootaxa, 2015, 3973, 474-90.	0.5	3
155	<i>TBX6</i> Null Variants and a Common Hypomorphic Allele in Congenital Scoliosis. New England Journal of Medicine, 2015, 372, 341-350.	27.0	239
156	Systematic revision of Entomobryidae ( <scp>C</scp> ollembola) by integrating molecular and new morphological evidence. Zoologica Scripta, 2015, 44, 298-311.	1.7	67
157	Non-coding genetic variants in human disease: Figure 1 Human Molecular Genetics, 2015, 24, R102-R110.	2.9	466
158	Two new species of Coecobrya (Collembola, Entomobryidae) from China, with an updated key to the Chinese species of the genus. ZooKeys, 2015, 498, 17-28.	1.1	5
159	CNV instability associated with DNA replication dynamics: evidence for replicative mechanisms in CNV mutagenesis. Human Molecular Genetics, 2015, 24, 1574-1583.	2.9	15
160	Correlation between frequency of non-allelic homologous recombination and homology properties: evidence from homology-mediated CNV mutations in the human genome. Human Molecular Genetics, 2015, 24, 1225-1233.	2.9	14
161	Single-nucleotide polymorphisms and haplotypes of non-coding area in the CP gene are correlated with Parkinson's disease. Neuroscience Bulletin, 2015, 31, 245-256.	2.9	7
162	Assessing structural variation in a personal genomeâ€"towards a human reference diploid genome. BMC Genomics, 2015, 16, 286.	2.8	153

#	Article	IF	CITATIONS
163	Two new species of Willowsia from New Caledonia (Collembola: Entomobryidae). Zootaxa, 2014, 3872, 381-6.	0.5	3
164	Screening of Duchenne Muscular Dystrophy (DMD) Mutations and Investigating Its Mutational Mechanism in Chinese Patients. PLoS ONE, 2014, 9, e108038.	2.5	28
165	<strong>Three new species of <em>Coecobrya</em> (Collembola: Entomobryidae) </strong> /> <strong>from southern and northwest China</strong> . Zootaxa, 2014, 3760, 260.	0.5	6
166	<p><strong>New species of <em>Sinella</em> and <em>Coecobrya</em> (Collembola: Entomobryidae) from New Caledonia</strong></p> . Zootaxa, 2014, 3814, 553.	0.5	3
167	<strong>New species of <em>Monodontocerus </em> (Collembola:) Tj ETQq1 1 0.7843 taxonomic characters</strong> . Zootaxa, 2014, 3768, 557.	314 rgBT 0.5	/Overlock 10 7
168	A remarkable new genus of Oncopoduridae (Collembola) from China. Journal of Natural History, 2014, 48, 2069-2082.	0.5	2
169	A new species of Dicranocentrus (Collembola, Entomobryidae) from China with comments onÂtheÂsystematic position of the genus. ZooKeys, 2014, 417, 1-8.	1.1	4
170	A peculiar cave species of Tomocerus (Collembola, Tomoceridae, Tomocerinae) from Vietnam, with a discussion of the postantennal organ and Aprelabral Achaetae in Tomocerinae. ZooKeys, 2014, 408, 61-70.	1.1	14
171	Epigenetic regulation of Atrophin1 by lysine-specific demethylase 1 is required for cortical progenitor maintenance. Nature Communications, 2014, 5, 5815.	12.8	46
172	Evaluation of copy number variation detection for a SNP array platform. BMC Bioinformatics, 2014, 15, 50.	2.6	34
173	Microcephaly-Associated Protein WDR62 Regulates Neurogenesis through JNK1 in the Developing Neocortex. Cell Reports, 2014, 6, 104-116.	6.4	71
174	Genome-wide CNV analysis in mouse induced pluripotent stem cells reveals dosage effect of pluripotent factors on genome integrity. BMC Genomics, 2014, 15, 79.	2.8	8
175	NIPA2 mutations are correlative with childhood absence epilepsy in the Han Chinese population. Human Genetics, 2014, 133, 675-676.	3.8	7
176	Highâ€resolution molecular genomic autopsy reveals complex sudden unexpected death in epilepsy risk profile. Epilepsia, 2014, 55, e6-12.	5.1	76
177	Cryptic diversity, diversification and vicariance in two species complexes of <i><scp>T</scp>omocerus</i> ( <scp>C</scp> ollembola, <scp>T</scp> omoceridae) from <scp>C</scp> hina. Zoologica Scripta, 2014, 43, 393-404.	1.7	36
178	Disjunct distribution of Szeptyckiellagen. nov. from New Caledonia and South China undermines the monophyly of Willowsiini (Collembola: Entomobryidae). Journal of Natural History, 2014, 48, 1299-1317.	0.5	9
179	Rearrangement structure-independent strategy of CNV breakpoint analysis. Molecular Genetics and Genomics, 2014, 289, 755-763.	2.1	5
180	Association between HLA-DQA1 gene copy number polymorphisms and susceptibility to rheumatoid arthritis in Chinese Han population. Journal of Genetics, 2014, 93, 215-218.	0.7	4

#	Article	IF	Citations
181	Mechanism, Prevalence, and More Severe Neuropathy Phenotype of the Charcot-Marie-Tooth Type 1A Triplication. American Journal of Human Genetics, 2014, 94, 462-469.	6.2	42
182	TAK1 is activated by TGF- $\hat{l}^2$ signaling and controls axonal growth during brain development. Journal of Molecular Cell Biology, 2014, 6, 349-351.	3.3	9
183	Molecular phylogeny reveals independent origins of body scales in Entomobryidae (Hexapoda:) Tj ETQq1 1 0.784	1314 rgBT 2.7	/Oyerlock 10 47
184	Increased genome instability in human DNA segments with self-chains: homology-induced structural variations via replicative mechanisms. Human Molecular Genetics, 2013, 22, 2642-2651.	2.9	22
185	Combinational analysis of linkage and exome sequencing identifies the causative mutation in a Chinese family with congenital cataract. BMC Medical Genetics, 2013, 14, 107.	2.1	10
186	Replicative mechanisms of CNV formation preferentially occur as intrachromosomal events: evidence from Potocki–Lupski duplication syndrome. Human Molecular Genetics, 2013, 22, 749-756.	2.9	13
187	Multiplex Genome Engineering Using CRISPR/Cas Systems. Science, 2013, 339, 819-823.	12.6	12,725
188	Molecular Analysis of a Deletion Hotspot in the NRXN1 Region Reveals the Involvement of Short Inverted Repeats in Deletion CNVs. American Journal of Human Genetics, 2013, 92, 375-386.	6.2	42
189	Two new species ofPogonognathellus(Collembola: Tomoceridae) from China, with a discussion of East Asian species. Journal of Natural History, 2013, 47, 1243-1255.	0.5	4
190	First record of Acrocyrtus Yosii, 1959 (Collemobla, Entomobryidae) from Chinese mainland. ZooKeys, 2013, 260, 1-16.	1.1	8
191	Zinc-finger-nucleases mediate specific and efficient excision of HIV-1 proviral DNA from infected and latently infected human T cells. Nucleic Acids Research, 2013, 41, 7771-7782.	14.5	146
192	DAZ duplications confer the predisposition of Y chromosome haplogroup K* to non-obstructive azoospermia in Han Chinese populations. Human Reproduction, 2013, 28, 2440-2449.	0.9	13
193	<p><strong>Five new eyed species of <em>Sinella</em> (Collembola:) Tj ETQq1 1 0.78433 Zootaxa, 2013, 3736, 549.</strong></p>	14 rgBT /O\ 0.5	verlock 10 Tf 6
194	Two new species of i>Onychiurus i>(Collembola: Onychiuridae) from Eastern China. Journal of Natural History, 2012, 46, 1895-1904.	0.5	8
195	Efficient typing of copy number variations in a segmental duplication-mediated rearrangement hotspot using multiplex competitive amplification. Journal of Human Genetics, 2012, 57, 545-551.	2.3	63
196	POSH Localizes Activated Rac1 to Control the Formation of Cytoplasmic Dilation of the Leading Process and Neuronal Migration. Cell Reports, 2012, 2, 640-651.	6.4	63
197	Novel Androgen Receptor Gene Mutation in Patient With Complete Androgen Insensitivity Syndrome. Urology, 2012, 80, 216-218.	1.0	3
198	Two new species of Coecobrya (Collembola: Entomobryidae: Entomobryinae) from China, with a key to the Chinese species of the genus. Zootaxa, 2012, 3399, 61.	0.5	5

#	Article	IF	Citations
199	Contribution to study of eyed species of Sinella (Collembola: Entomobryidae), with a key to this group of species from China. Zootaxa, 2012, 3180, 35.	0.5	6
200	NIPA2 located in $15q11.2$ is mutated in patients with childhood absence epilepsy. Human Genetics, 2012, 131, 1217-1224.	3.8	33
201	Cave <i>Sinella</i> (Collembola: Entomobryidae) from China. Journal of Natural History, 2011, 45, 1213-1231.	0.5	5
202	Structural variation of the human genome: mechanisms, assays, and role in male infertility. Systems Biology in Reproductive Medicine, 2011, 57, 3-16.	2.1	32
203	Chromosome Catastrophes Involve Replication Mechanisms Generating Complex Genomic Rearrangements. Cell, 2011, 146, 889-903.	28.9	391
204	Transformational homology of the tergal setae during postembryonic development in the Sinella-Coecobrya group (Collembola: Entomobryidae). Contributions To Zoology, 2011, 80, 213-230.	0.5	27
205	A review of the boneti-group of the genus Coecobrya (Collembola: Entomobryidae). Zootaxa, 2011, 2748, .	0.5	5
206	New insight into the systematics of the <i>Willowsia </i> complex (Collembola: Entomobryidae). Annales De La Societe Entomologique De France, 2011, 47, 1-20.	0.9	16
207	Potocki-Lupski Syndrome: A Microduplication Syndrome Associated with Oropharyngeal Dysphagia and Failure to Thrive. Journal of Pediatrics, 2011, 158, 655-659.e2.	1.8	36
208	X-Linked Congenital Hypertrichosis Syndrome Is Associated with Interchromosomal Insertions Mediated by a Human-Specific Palindrome near SOX3. American Journal of Human Genetics, 2011, 88, 819-826.	6.2	87
209	Frequency of Nonallelic Homologous Recombination Is Correlated with Length of Homology: Evidence that Ectopic Synapsis Precedes Ectopic Crossing-Over. American Journal of Human Genetics, 2011, 89, 580-588.	6.2	104
210	Alu-specific microhomology-mediated deletion of the final exon of SPAST in three unrelated subjects with hereditary spastic paraplegia. Genetics in Medicine, 2011, 13, 582-592.	2.4	53
211	Additional genomic duplications in AZFc underlie the b2/b3 deletion-associated risk of spermatogenic impairment in Han Chinese population. Human Molecular Genetics, 2011, 20, 4411-4421.	2.9	33
212	New species of Homidia (Collembola, Entomobryidae) from eastern China with description of the first instar larvae. ZooKeys, 2011, 152, 21-42.	1.1	13
213	Identification of Uncommon Recurrent Potocki-Lupski Syndrome-Associated Duplications and the Distribution of Rearrangement Types and Mechanisms in PTLS. American Journal of Human Genetics, 2010, 86, 462-470.	6.2	79
214	Mechanisms for Nonrecurrent Genomic Rearrangements Associated with CMT1A or HNPP: Rare CNVs as a Cause for Missing Heritability. American Journal of Human Genetics, 2010, 86, 892-903.	6.2	125
215	Identification of Copy Number Variation Hotspots in Human Populations. American Journal of Human Genetics, 2010, 87, 494-504.	6.2	42
216	GJB1/Connexin 32 whole gene deletions in patients with X-linked Charcot–Marie–Tooth disease. Neurogenetics, 2010, 11, 465-470.	1.4	27

#	Article	IF	Citations
217	Deletion and duplication of 15q24: Molecular mechanisms and potential modification by additional copy number variants. Genetics in Medicine, 2010, 12, 573-586.	2.4	31
218	Two syntopic and remarkably similar new species of <i>Sinella </i> and <i>Coecobrya </i> from South China (Collembola, Entomobryidae). Zoosystema, 2010, 32, 469-477.	0.6	7
219	The Suppression of CRMP2 Expression by Bone Morphogenetic Protein (BMP)-SMAD Gradient Signaling Controls Multiple Stages of Neuronal Development. Journal of Biological Chemistry, 2010, 285, 39039-39050.	3.4	49
220	Whole-Genome Sequencing in a Patient with Charcot–Marie–Tooth Neuropathy. New England Journal of Medicine, 2010, 362, 1181-1191.	27.0	698
221	Two new species of the genus Sinella Brook, (Collembola: Entomobryidae) from East China. Journal of Natural History, 2010, 44, 2535-2541.	0.5	5
222	Genomic disorders: A window into human gene and genome evolution. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 1765-1771.	7.1	60
223	A new species and a new record of the genus Homidia BÃ $\P$ rner, 1906 from East China (Collembola:) Tj ETQq $1\ 1\ C$	).784314 0.5	rgBT /Overlo
224	A new cave Sinella species from South China (Collembola: Entomobryidae). Zootaxa, 2009, 2009, 35-40.	0.5	7
225	<strong>A new species of the genus <em>Lepidocyrtus </em>Bourlet and a new record of <em>Seira</em> <em>delamarei </em>Jacquemart (Collembola: Entomobryidae) from the east coast of India</strong> . Zootaxa, 2009, 2310, 43-50.	0.5	7
226	New species and rediagnosis of <i>Coecobrya </i> (Collembola: Entomobryidae), with a key to the species of the genus. Journal of Natural History, 2009, 43, 2597-2615.	0.5	32
227	Complex rearrangements in patients with duplications of MECP2 can occur by fork stalling and template switching. Human Molecular Genetics, 2009, 18, 2188-2203.	2.9	165
228	Complex human chromosomal and genomic rearrangements. Trends in Genetics, 2009, 25, 298-307.	6.7	239
229	Autism and other neuropsychiatric symptoms are prevalent in individuals with <i>MeCP2</i> duplication syndrome. Annals of Neurology, 2009, 66, 771-782.	5.3	271
230	Association of polymorphisms in four bilirubin metabolism genes with serum bilirubin in three Asian populations. Human Mutation, 2009, 30, 609-615.	2.5	34
231	Increased LIS1 expression affects human and mouse brain development. Nature Genetics, 2009, 41, 168-177.	21.4	199
232	The DNA replication FoSTeS/MMBIR mechanism can generate genomic, genic and exonic complex rearrangements in humans. Nature Genetics, 2009, 41, 849-853.	21.4	382
233	The b2/b3 subdeletion shows higher risk of spermatogenic failure and higher frequency of complete AZFc deletion than the gr/gr subdeletion in a Chinese population. Human Molecular Genetics, 2009, $18$ , $1122-1130$ .	2.9	86
234	Copy Number Variation in Human Health, Disease, and Evolution. Annual Review of Genomics and Human Genetics, 2009, 10, 451-481.	6.2	1,026

#	Article	IF	CITATIONS
235	Y chromosome evidence of earliest modern human settlement in East Asia and multiple origins of Tibetan and Japanese populations. BMC Biology, 2008, 6, 45.	3.8	129
236	The clinical spectrum associated with a chromosome 17 short arm proximal duplication (dup 17p11.2) in three patients. American Journal of Medical Genetics, Part A, 2008, 146A, 917-924.	1.2	10
237	A spatial analysis of genetic structure of human populations in China reveals distinct difference between maternal and paternal lineages. European Journal of Human Genetics, 2008, 16, 705-717.	2.8	45
238	The Genus <i>Willowsia </i> and Its Mexican Species (Collembola: Entomobryidae). Annals of the Entomological Society of America, 2007, 100, 36-40.	2.5	7
239	Genetic studies of human diversity in East Asia. Philosophical Transactions of the Royal Society B: Biological Sciences, 2007, 362, 987-996.	4.0	61
240	Partial deletions are associated with an increased risk of complete deletion in AZFc: a new insight into the role of partial AZFc deletions in male infertility. Journal of Medical Genetics, 2007, 44, 437-444.	3.2	82
241	A new species of Willowsia (Collembola: Entomobryidae) from South China. Zootaxa, 2007, 1645, 63-68.	0.5	0
242	Rapid Evolution, Genetic Variations, and Functional Association of the Human Spermatogenesis-Related Gene NYD-SP12. Journal of Molecular Evolution, 2007, 65, 154-161.	1.8	11
243	The association of Y chromosome haplogroups with spermatogenic failure in the Han Chinese. Journal of Human Genetics, 2007, 52, 659-663.	2.3	12
244	Roles of springtails in soil ecosystem. Biodiversity Science, 2007, 15, 154.	0.6	7
245	A New Chinese Species of the Genus Willowsia from Tibet (Collembola: Entomobryidae). Journal of the Kansas Entomological Society, 2006, 79, 261-266.	0.2	0
246	Genetic Structure of Hmong-Mien Speaking Populations in East Asia as Revealed by mtDNA Lineages. Molecular Biology and Evolution, 2005, 22, 725-734.	8.9	105
247	Genetic evidence supports demic diffusion of Han culture. Nature, 2004, 431, 302-305.	27.8	398
248	Revision of Acanthocyrtus (Collembola: Entomobryidae), with description of a new genus from eastern Asia. Zoological Journal of the Linnean Society, 0, 157, 495-514.	2.3	12
249	$\ddot{\text{i}}$ »¿A new species of Arrhopalites Börner (Collembola, Symphypleona, Arrhopalitidae) from China, with a key to the Asian species of the caecus group. ZooKeys, 0, 1102, 163-176.	1.1	0