## Cheng-Qi Xu

## List of Publications by Year in descending order

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257101 276539 2,009 77 24 41 h-index citations g-index papers 78 78 78 2806 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Genome-wide association identifies a susceptibility locus for coronary artery disease in the Chinese Han population. Nature Genetics, 2011, 43, 345-349.	9.4	256
2	Assessment of association of rs2200733 on chromosome 4q25 with atrial fibrillation and ischemic stroke in a Chinese Han population. Human Genetics, 2009, 126, 843-849.	1.8	93
3	Regulation of <i>CARD8</i> Expression by <i>ANRIL</i> and Association of <i>CARD8</i> Single Nucleotide Polymorphism rs2043211 (p.C10X) With Ischemic Stroke. Stroke, 2014, 45, 383-388.	1.0	87
4	De novo loss-of-function KCNMA1 variants are associated with a new multiple malformation syndrome and a broad spectrum of developmental and neurological phenotypes. Human Molecular Genetics, 2019, 28, 2937-2951.	1.4	76
5	Functional dominant-negative mutation of sodium channel subunit gene SCN3B associated with atrial fibrillation in a Chinese GeneID population. Biochemical and Biophysical Research Communications, 2010, 398, 98-104.	1.0	75
6	Angiogenic Factor AGGF1 Activates Autophagy with an Essential Role in Therapeutic Angiogenesis for Heart Disease. PLoS Biology, 2016, 14, e1002529.	2.6	75
7	Significant association of SNP rs2106261 in the ZFHX3 gene with atrial fibrillation in a Chinese Han GenelD population. Human Genetics, 2011, 129, 239-246.	1.8	74
8	Identification of NPPA variants associated with atrial fibrillation in a Chinese GeneID population. Clinica Chimica Acta, 2010, 411, 481-485.	0.5	59
9	The Same Chromosome 9p21.3 Locus Is Associated With Type 2 Diabetes and Coronary Artery Disease in a Chinese Han Population. Diabetes, 2011, 60, 680-684.	0.3	51
10	The IL-33-ST2L Pathway Is Associated with Coronary Artery Disease in a Chinese Han Population. American Journal of Human Genetics, 2013, 93, 652-660.	2.6	48
11	Role of microRNA-27a in down-regulation of angiogenic factor AGGF1 under hypoxia associated with high-grade bladder urothelial carcinoma. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2014, 1842, 712-725.	1.8	48
12	Molecular Basis of Gene-Gene Interaction: Cyclic Cross-Regulation of Gene Expression and Post-GWAS Gene-Gene Interaction Involved in Atrial Fibrillation. PLoS Genetics, 2015, 11, e1005393.	1.5	47
13	Genomic variant in CAV1 increases susceptibility to coronary artery disease and myocardial infarction. Atherosclerosis, 2016, 246, 148-156.	0.4	44
14	Angiogenic Factor AGGF1 Promotes Therapeutic Angiogenesis in a Mouse Limb Ischemia Model. PLoS ONE, 2012, 7, e46998.	1.1	43
15	αB-Crystallin Interacts with Nav1.5 and Regulates Ubiquitination and Internalization of Cell Surface Nav1.5. Journal of Biological Chemistry, 2016, 291, 11030-11041.	1.6	41
16	MiR-144 regulates hematopoiesis and vascular development by targeting meis1 during zebrafish development. International Journal of Biochemistry and Cell Biology, 2014, 49, 53-63.	1.2	39
17	Minor Allele C of Chromosome 1p32 Single Nucleotide Polymorphism rs11206510 Confers Risk of Ischemic Stroke in the Chinese Han Population. Stroke, 2010, 41, 1587-1592.	1.0	35
18	A novel DSPPmutation is associated with type II dentinogenesis Imperfecta in a chinese family. BMC Medical Genetics, 2007, 8, 52.	2.1	33

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19	Aggf1 acts at the top of the genetic regulatory hierarchy in specification of hemangioblasts in zebrafish. Blood, 2014, 123, 501-508.	0.6	33
20	NINJ2– A novel regulator of endothelial inflammation and activation. Cellular Signalling, 2017, 35, 231-241.	1.7	32
21	Genomic Variant in IL-37 Confers A Significant Risk of Coronary Artery Disease. Scientific Reports, 2017, 7, 42175.	1.6	31
22	Identification of a molecular signaling gene-gene regulatory network between GWAS susceptibility genes ADTRP and MIA3/TANGO1 for coronary artery disease. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2017, 1863, 1640-1653.	1.8	31
23	Candidate Pathway-Based Genome-Wide Association Studies Identify Novel Associations of Genomic Variants in the Complement System Associated With Coronary Artery Disease. Circulation: Cardiovascular Genetics, 2014, 7, 887-894.	5.1	30
24	SUMOylation of Vps34 by SUMO1 promotes phenotypic switching of vascular smooth muscle cells by activating autophagy in pulmonary arterial hypertension. Pulmonary Pharmacology and Therapeutics, 2019, 55, 38-49.	1,1	28
25	Endothelial cell metabolic memory causes cardiovascular dysfunction in diabetes. Cardiovascular Research, 2022, 118, 196-211.	1.8	26
26	Significant Association Between $<$ i>CAV1 $<$ /i> Variant rs3807989 on 7p31 and Atrial Fibrillation in a Chinese Han Population. Journal of the American Heart Association, 2015, 4, .	1.6	25
27	BRG1 variant rs1122608 on chromosome 19p13.2 confers protection against stroke and regulates expression of pre-mRNA-splicing factor SFRS3. Human Genetics, 2014, 133, 499-508.	1.8	24
28	Significant genetic association of a functional TFPI variant with circulating fibrinogen levels and coronary artery disease. Molecular Genetics and Genomics, 2018, 293, 119-128.	1.0	24
29	Meta-analysis identifies robust association between SNP rs17465637 in MIA3 on chromosome 1q41 and coronary artery disease. Atherosclerosis, 2013, 231, 136-140.	0.4	22
30	Significant association of rare variant p.Gly8Ser in cardiac sodium channel β4â€subunit SCN4B with atrial fibrillation. Annals of Human Genetics, 2019, 83, 239-248.	0.3	22
31	Genomic Variants in NEURL, GJA1 and CUX2 Significantly Increase Genetic Susceptibility to Atrial Fibrillation. Scientific Reports, 2018, 8, 3297.	1.6	21
32	Angiotensin II increases angiogenesis by NFâ€Pâê"mediated transcriptional activation of angiogenic factor AGGF1. FASEB Journal, 2018, 32, 5051-5062.	0.2	21
33	Small GTPases SAR1A and SAR1B regulate the trafficking of the cardiac sodium channel Nav1.5. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2018, 1864, 3672-3684.	1.8	20
34	Targeting AGGF1 (angiogenic factor with G patch and FHA domains 1) for Blocking Neointimal Formation After Vascular Injury. Journal of the American Heart Association, 2017, 6, .	1.6	19
35	Angiogenic Factor AGGF1-Primed Endothelial Progenitor Cells Repair Vascular Defect in Diabetic Mice. Diabetes, 2019, 68, 1635-1648.	0.3	19
36	Statistical and Functional Studies Identify Epistasis of Cardiovascular Risk Genomic Variants From Genomeâ€Wide Association Studies. Journal of the American Heart Association, 2020, 9, e014146.	1.6	19

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37	Defective circulating CD4+LAP+ regulatory T cells in patients with dilated cardiomyopathy. Journal of Leukocyte Biology, 2015, 97, 797-805.	1.5	18
38	UBC9 regulates cardiac sodium channel Nav1.5 ubiquitination, degradation and sodium current density. Journal of Molecular and Cellular Cardiology, 2019, 129, 79-91.	0.9	16
39	<i>Mog1</i> knockout causes cardiac hypertrophy and heart failure by downregulating <i>tbx5â€cryabâ€hspb2</i> signalling in zebrafish. Acta Physiologica, 2021, 231, e13567.	1.8	14
40	Angiogenic factor AGGF1 acts as a tumor suppressor by modulating p53 post-transcriptional modifications and stability via MDM2. Cancer Letters, 2021, 497, 28-40.	3.2	14
41	Identification and functional analysis of two new de novo <i>KCNMA1</i> variants associated with Liangâ€"Wang syndrome. Acta Physiologica, 2022, 235, e13800.	1.8	14
42	Identification of rare variants in TNNI3 with atrial fibrillation in a Chinese GeneID population. Molecular Genetics and Genomics, 2016, 291, 79-92.	1.0	13
43	Loss of heterozygosity detected at three short tandem repeat locus commonly used for human DNA identification in a case of paternity testing. Legal Medicine, 2017, 24, 7-11.	0.6	13
44	Identification of a new <i>adtrp1â€tfpi</i> regulatory axis for the specification of primitive myelopoiesis and definitive hematopoiesis. FASEB Journal, 2018, 32, 183-194.	0.2	13
45	<i>&gt;phlda3</i> overexpression impairs specification of hemangioblasts and vascular development. FEBS Journal, 2018, 285, 4071-4081.	2.2	13
46	Association of SNP Rs9943582 in APLNR with Left Ventricle Systolic Dysfunction in Patients with Coronary Artery Disease in a Chinese Han GenelD Population. PLoS ONE, 2015, 10, e0125926.	1.1	12
47	IL-13 may be involved in the development of CAD via different mechanisms under different conditions in a Chinese Han population. Scientific Reports, 2018, 8, 6182.	1.6	12
48	Genetic Regulation of the Thymic Stromal Lymphopoietin (TSLP)/TSLP Receptor (TSLPR) Gene Expression and Influence of Epistatic Interactions Between IL-33 and the TSLP/TSLPR Axis on Risk of Coronary Artery Disease. Frontiers in Immunology, 2018, 9, 1775.	2.2	12
49	Functional rare variant in a C/EBP beta binding site in NINJ2 gene increases the risk of coronary artery disease. Aging, 2021, 13, 25393-25407.	1.4	11
50	Role of epigenetic m 6 A RNA methylation in vascular development: mettl3 regulates vascular development through PHLPP2/mTORâ€AKT signaling. FASEB Journal, 2021, 35, e21465.	0.2	10
51	Analysis of the genetic association between IL27 variants and coronary artery disease in a Chinese Han population. Scientific Reports, 2016, 6, 25782.	1.6	9
52	ADTRP regulates TFPI expression via transcription factor POU1F1 involved in coronary artery disease. Gene, 2020, 753, 144805.	1.0	9
53	Ubiquitination-activating enzymes UBE1 and UBA6 regulate ubiquitination and expression of cardiac sodium channel Nav1.5. Biochemical Journal, 2020, 477, 1683-1700.	1.7	9
54	DOCK2 regulates antifungal immunity by regulating RAC GTPase activity. Cellular and Molecular Immunology, 2022, 19, 602-618.	4.8	9

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55	Angiogenic factor AGGF1 blocks neointimal formation after vascular injury via interaction with integrin $\hat{l}\pm7$ on vascular smooth muscle cells. Journal of Biological Chemistry, 2022, 298, 101759.	1.6	9
56	Alpha-actin-2 mutations in Chinese patients with a non-syndromatic thoracic aortic aneurysm. BMC Medical Genetics, 2016, 17, 45.	2.1	8
57	Exomic and Epigenomic Analyses in a Pair of Monozygotic Twins Discordant for Cryptorchidism. Twin Research and Human Genetics, 2017, 20, 349-354.	0.3	8
58	Lack of association between the APLNR variant rs9943582 with ischemic stroke in the Chinese Han GenelD population. Oncotarget, 2017, 8, 107678-107684.	0.8	8
59	Rod genesis driven by mafba in an nrl knockout zebrafish model with altered photoreceptor composition and progressive retinal degeneration. PLoS Genetics, 2022, 18, e1009841.	1.5	8
60	Significant Association between OPG/TNFRSF11B Variant and Common Complex Ischemic Stroke. Journal of Stroke and Cerebrovascular Diseases, 2018, 27, 1683-1691.	0.7	7
61	Receptor and Molecular Mechanism of AGGF1 Signaling in Endothelial Cell Functions and Angiogenesis. Arteriosclerosis, Thrombosis, and Vascular Biology, 2021, 41, 2756-2769.	1.1	7
62	Mutation p.Arg954Trp of KIF21A Causes Congenital Fibrosis of the Extraocular Muscles in a Chinese Family. Journal of Genetics and Genomics, 2006, 33, 685-691.	0.3	6
63	Mutational analysis of <i>HOXA10</i> gene in Chinese patients with cryptorchidism. Andrologia, 2017, 49, e12592.	1.0	6
64	Mechanistic insights into the interaction of cardiac sodium channel Nav1.5 with MOG1 and a new molecular mechanism for Brugada syndrome. Heart Rhythm, 2022, 19, 478-489.	0.3	6
65	Identification of rare variants in cardiac sodium channel $\hat{l}^2$ 4-subunit gene SCN4B associated with ventricular tachycardia. Molecular Genetics and Genomics, 2019, 294, 1059-1071.	1.0	5
66	Identification of a p.Trp403* nonsense variant in PHEX causing Xâ€linked hypophosphatemia by inhibiting p38 MAPK signaling. Human Mutation, 2019, 40, 879-885.	1.1	5
67	SNP rs2243828 in MPO associated with myeloperoxidase level and atrial fibrillation risk in Chinese Han population. Journal of Cellular and Molecular Medicine, 2020, 24, 10263-10266.	1.6	5
68	Feedback regulation of coronary artery disease susceptibility gene ADTRP and LDL receptors LDLR/CD36/LOX-1 in endothelia cell functions involved in atherosclerosis. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2021, 1867, 166130.	1.8	5
69	A novel <i><scp>FBN</scp>2</i> mutation cosegregates with congenital contractural arachnodactyly in a fiveâ€generation Chinese family. Clinical Case Reports (discontinued), 2018, 6, 1612-1617.	0.2	4
70	A lower complexity antenna selection for DF relay mimo systems. Journal of Electronics, 2011, 28, 320-327.	0.2	3
71	A genomeâ€wide association study identifies novel association between genetic variants in ⟨i⟩GGT7⟨ i⟩ and ⟨i⟩LINC00944⟨ i⟩ and hypertension. Clinical and Translational Medicine, 2021, 11, e388.	1.7	3
72	Annotation of susceptibility SNPs associated with atrial fibrillation. Aging, 2020, 12, 16981-16998.	1.4	3

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73	Genetic association analysis between IL9 and coronary artery disease in a Chinese Han population. Cytokine, 2022, 150, 155761.	1.4	3
74	Hyperlipidemia patients carrying LDLR splicing mutation c.1187-2A> Grespond favorably to rosuvastatin and PCSK9 inhibitor evolocumab. Molecular Genetics and Genomics, 2022, 297, 833-841.	1.0	3
75	Identification and characterization of a special type of subnuclear structure: AGGF1â€coated paraspeckles. FASEB Journal, 2022, 36, .	0.2	3
76	A family with Liddle's syndrome caused by a new c.1721 deletion mutation in the epithelial sodium channel $\hat{l}^2\hat{a}$ subunit. Experimental and Therapeutic Medicine, 2019, 17, 2777-2784.	0.8	2
77	Abstract 501: Angiotensin II Increases Angiogenesis by Nf-κB-mediated Transcriptional Activation of Angiogenic Factor Aggf1. Arteriosclerosis, Thrombosis, and Vascular Biology, 2018, 38, .	1.1	0