Alanna C Morrison

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226 15,107 57 120 h-index g-index citations papers 10.6 20,101 5.01 257 avg, IF L-index ext. citations ext. papers

#	Paper	IF	Citations
226	Genetic variants in novel pathways influence blood pressure and cardiovascular disease risk. <i>Nature</i> , 2011 , 478, 103-9	50.4	1564
225	A comprehensive 1,000 Genomes-based genome-wide association meta-analysis of coronary artery disease. <i>Nature Genetics</i> , 2015 , 47, 1121-1130	36.3	1290
224	Genome-wide association study of blood pressure and hypertension. <i>Nature Genetics</i> , 2009 , 41, 677-87	36.3	1065
223	Meta-analyses of genome-wide association studies identify multiple loci associated with pulmonary function. <i>Nature Genetics</i> , 2010 , 42, 45-52	36.3	467
222	Diagnostic Yield and Clinical Utility of Sequencing Familial Hypercholesterolemia Genes in Patients With Severe Hypercholesterolemia. <i>Journal of the American College of Cardiology</i> , 2016 , 67, 2578-89	15.1	458
221	Genetic analysis of over 1 million people identifies 535 new loci associated with blood pressure traits. <i>Nature Genetics</i> , 2018 , 50, 1412-1425	36.3	386
220	Genome-wide association study identifies six new loci influencing pulse pressure and mean arterial pressure. <i>Nature Genetics</i> , 2011 , 43, 1005-11	36.3	338
219	Genome-wide association and large-scale follow up identifies 16 new loci influencing lung function. <i>Nature Genetics</i> , 2011 , 43, 1082-90	36.3	313
218	Exome-wide association study of plasma lipids in >300,000 individuals. <i>Nature Genetics</i> , 2017 , 49, 1758-	1 76 .6	310
217	Inactivating mutations in NPC1L1 and protection from coronary heart disease. <i>New England Journal of Medicine</i> , 2014 , 371, 2072-82	59.2	307
216	Genetic associations at 53 loci highlight cell types and biological pathways relevant for kidney function. <i>Nature Communications</i> , 2016 , 7, 10023	17.4	295
215	Sequencing of 53,831 diverse genomes from the NHLBI TOPMed Program. <i>Nature</i> , 2021 , 590, 290-299	50.4	268
214	The genetics of blood pressure regulation and its target organs from association studies in 342,415 individuals. <i>Nature Genetics</i> , 2016 , 48, 1171-1184	36.3	251
213	Association Between Telomere Length and Risk of Cancer and Non-Neoplastic Diseases: A Mendelian Randomization Study. <i>JAMA Oncology</i> , 2017 , 3, 636-651	13.4	236
212	Association of low-frequency and rare coding-sequence variants with blood lipids and coronary heart disease in 56,000 whites and blacks. <i>American Journal of Human Genetics</i> , 2014 , 94, 223-32	11	233
211	Large-scale genomic analyses link reproductive aging to hypothalamic signaling, breast cancer susceptibility and BRCA1-mediated DNA repair. <i>Nature Genetics</i> , 2015 , 47, 1294-1303	36.3	226
210	Prediction of coronary heart disease risk using a genetic risk score: the Atherosclerosis Risk in Communities Study. <i>American Journal of Epidemiology</i> , 2007 , 166, 28-35	3.8	222

209	Identification of heart rate-associated loci and their effects on cardiac conduction and rhythm disorders. <i>Nature Genetics</i> , 2013 , 45, 621-31	36.3	219
208	Meta-analysis identifies common and rare variants influencing blood pressure and overlapping with metabolic trait loci. <i>Nature Genetics</i> , 2016 , 48, 1162-70	36.3	152
207	Genome Analyses of >200,000 Individuals Identify 58 Loci for Chronic Inflammation and Highlight Pathways that Link Inflammation and Complex Disorders. <i>American Journal of Human Genetics</i> , 2018 , 103, 691-706	11	151
206	KLB is associated with alcohol drinking, and its gene product EKlotho is necessary for FGF21 regulation of alcohol preference. <i>Proceedings of the National Academy of Sciences of the United States of America</i> , 2016 , 113, 14372-14377	11.5	150
205	Low-frequency and rare exome chip variants associate with fasting glucose and type 2 diabetes susceptibility. <i>Nature Communications</i> , 2015 , 6, 5897	17.4	147
204	Association of genome-wide variation with the risk of incident heart failure in adults of European and African ancestry: a prospective meta-analysis from the cohorts for heart and aging research in genomic epidemiology (CHARGE) consortium. <i>Circulation: Cardiovascular Genetics</i> , 2010 , 3, 256-66		147
203	Association of genetic variation with systolic and diastolic blood pressure among African Americans: the Candidate Gene Association Resource study. <i>Human Molecular Genetics</i> , 2011 , 20, 2273-	8 4 ⁶	146
202	Genome-wide association analysis of blood-pressure traits in African-ancestry individuals reveals common associated genes in African and non-African populations. <i>American Journal of Human Genetics</i> , 2013 , 93, 545-54	11	145
201	Genome-wide association and Mendelian randomisation analysis provide insights into the pathogenesis of heart failure. <i>Nature Communications</i> , 2020 , 11, 163	17.4	140
200	Genome-wide association studies identify CHRNA5/3 and HTR4 in the development of airflow obstruction. <i>American Journal of Respiratory and Critical Care Medicine</i> , 2012 , 186, 622-32	10.2	131
199	Exome sequencing of 20,791 cases of type 2 diabetes and 24,440 controls. <i>Nature</i> , 2019 , 570, 71-76	50.4	129
198	CD163+ macrophages promote angiogenesis and vascular permeability accompanied by inflammation in atherosclerosis. <i>Journal of Clinical Investigation</i> , 2018 , 128, 1106-1124	15.9	126
197	Directional dominance on stature and cognition indiverse human populations. <i>Nature</i> , 2015 , 523, 459-4	1 63 0.4	119
196	Whole-genome sequence-based analysis of high-density lipoprotein cholesterol. <i>Nature Genetics</i> , 2013 , 45, 899-901	36.3	117
195	Plasma MCP-1 level and risk for peripheral arterial disease and incident coronary heart disease: Atherosclerosis Risk in Communities study. <i>Atherosclerosis</i> , 2005 , 183, 301-7	3.1	117
194	Genome-wide joint meta-analysis of SNP and SNP-by-smoking interaction identifies novel loci for pulmonary function. <i>PLoS Genetics</i> , 2012 , 8, e1003098	6	108
193	Genome-wide association analysis identifies multiple loci related to resting heart rate. <i>Human Molecular Genetics</i> , 2010 , 19, 3885-94	5.6	106
192	Genome-wide association analysis identifies six new loci associated with forced vital capacity. Nature Genetics, 2014, 46, 669-77	36.3	104

191	Duffy antigen receptor for chemokines (Darc) polymorphism regulates circulating concentrations of monocyte chemoattractant protein-1 and other inflammatory mediators. <i>Blood</i> , 2010 , 115, 5289-99	2.2	96
190	Use of >100,000 NHLBI Trans-Omics for Precision Medicine (TOPMed) Consortium whole genome sequences improves imputation quality and detection of rare variant associations in admixed African and Hispanic/Latino populations. <i>PLoS Genetics</i> , 2019 , 15, e1008500	6	90
189	Association of the PHACTR1/EDN1 Genetic Locus With Spontaneous Coronary Artery Dissection. Journal of the American College of Cardiology, 2019 , 73, 58-66	15.1	86
188	Novel Blood Pressure Locus and Gene Discovery Using Genome-Wide Association Study and Expression Data Sets From Blood and the Kidney. <i>Hypertension</i> , 2017 ,	8.5	85
187	DNA Methylation Analysis Identifies Loci for Blood Pressure Regulation. <i>American Journal of Human Genetics</i> , 2017 , 101, 888-902	11	83
186	Pleiotropic genes for metabolic syndrome and inflammation. <i>Molecular Genetics and Metabolism</i> , 2014 , 112, 317-38	3.7	81
185	Gene-age interactions in blood pressure regulation: a large-scale investigation with the CHARGE, Global BPgen, and ICBP Consortia. <i>American Journal of Human Genetics</i> , 2014 , 95, 24-38	11	8o
184	Five common gene variants identify elevated genetic risk for coronary heart disease. <i>Genetics in Medicine</i> , 2007 , 9, 682-9	8.1	79
183	G-protein beta3 subunit and alpha-adducin polymorphisms and risk of subclinical and clinical stroke. <i>Stroke</i> , 2001 , 32, 822-9	6.7	78
182	Combined admixture mapping and association analysis identifies a novel blood pressure genetic locus on 5p13: contributions from the CARe consortium. <i>Human Molecular Genetics</i> , 2011 , 20, 2285-95	5.6	70
181	Analysis commons, a team approach to discovery in a big-data environment for genetic epidemiology. <i>Nature Genetics</i> , 2017 , 49, 1560-1563	36.3	68
180	Large-scale genome-wide analysis identifies genetic variants associated with cardiac structure and function. <i>Journal of Clinical Investigation</i> , 2017 , 127, 1798-1812	15.9	68
179	Sequencing of 53,831 diverse genomes from the NHLBI TOPMed Program		68
178	ADD1 460W allele associated with cardiovascular disease in hypertensive individuals. <i>Hypertension</i> , 2002 , 39, 1053-7	8.5	67
177	ACAT: A Fast and Powerful p Value Combination Method for Rare-Variant Analysis in Sequencing Studies. <i>American Journal of Human Genetics</i> , 2019 , 104, 410-421	11	66
176	Genomic variation associated with mortality among adults of European and African ancestry with heart failure: the cohorts for heart and aging research in genomic epidemiology consortium. <i>Circulation: Cardiovascular Genetics</i> , 2010 , 3, 248-55		66
175	Causal Effect of Plasminogen Activator Inhibitor Type 1 on Coronary Heart Disease. <i>Journal of the American Heart Association</i> , 2017 , 6,	6	65
174	Cystatin C and Cardiovascular Disease: A Mendelian Randomization Study. <i>Journal of the American College of Cardiology</i> , 2016 , 68, 934-45	15.1	65

(2015-2018)

173	GWAS and colocalization analyses implicate carotid intima-media thickness and carotid plaque loci in cardiovascular outcomes. <i>Nature Communications</i> , 2018 , 9, 5141	17.4	64
172	Common and rare von Willebrand factor (VWF) coding variants, VWF levels, and factor VIII levels in African Americans: the NHLBI Exome Sequencing Project. <i>Blood</i> , 2013 , 122, 590-7	2.2	60
171	Multi-ancestry genome-wide gene-smoking interaction study of 387,272 individuals identifies new loci associated with serum lipids. <i>Nature Genetics</i> , 2019 , 51, 636-648	36.3	59
170	Single-trait and multi-trait genome-wide association analyses identify novel loci for blood pressure in African-ancestry populations. <i>PLoS Genetics</i> , 2017 , 13, e1006728	6	58
169	Genetic determinants influencing human serum metabolome among African Americans. <i>PLoS Genetics</i> , 2014 , 10, e1004212	6	57
168	Replication of the Wellcome Trust genome-wide association study of essential hypertension: the Family Blood Pressure Program. <i>European Journal of Human Genetics</i> , 2008 , 16, 1507-11	5.3	57
167	A meta-analysis of 120 246 individuals identifies 18 new loci for fibrinogen concentration. <i>Human Molecular Genetics</i> , 2016 , 25, 358-70	5.6	54
166	Resequencing and clinical associations of the 9p21.3 region: a comprehensive investigation in the Framingham heart study. <i>Circulation</i> , 2013 , 127, 799-810	16.7	53
165	Genome-Wide Association Study for Incident Myocardial Infarction and Coronary Heart Disease in Prospective Cohort Studies: The CHARGE Consortium. <i>PLoS ONE</i> , 2016 , 11, e0144997	3.7	53
164	Effects of long-term averaging of quantitative blood pressure traits on the detection of genetic associations. <i>American Journal of Human Genetics</i> , 2014 , 95, 49-65	11	52
163	Inherited coding variants at the CDKN2A locus influence susceptibility to acute lymphoblastic leukaemia in children. <i>Nature Communications</i> , 2015 , 6, 7553	17.4	51
162	Genome-Wide Association Transethnic Meta-Analyses Identifies Novel Associations Regulating Coagulation Factor VIII and von Willebrand Factor Plasma Levels. <i>Circulation</i> , 2019 , 139, 620-635	16.7	51
161	Prospective Study of Epigenetic Age Acceleration and Incidence of Cardiovascular Disease Outcomes in the ARIC Study (Atherosclerosis Risk in Communities). <i>Circulation Genomic and Precision Medicine</i> , 2018 , 11, e001937	5.2	45
160	Multiethnic meta-analysis identifies ancestry-specific and cross-ancestry loci for pulmonary function. <i>Nature Communications</i> , 2018 , 9, 2976	17.4	45
159	Rare and low-frequency variants and their association with plasma levels of fibrinogen, FVII, FVIII, and vWF. <i>Blood</i> , 2015 , 126, e19-29	2.2	45
158	Causal Role of Alcohol Consumption in an Improved Lipid Profile: The Atherosclerosis Risk in Communities (ARIC) Study. <i>PLoS ONE</i> , 2016 , 11, e0148765	3.7	45
157	Efficient Variant Set Mixed Model Association Tests for Continuous and Binary Traits in Large-Scale Whole-Genome Sequencing Studies. <i>American Journal of Human Genetics</i> , 2019 , 104, 260-274	11	43
156	Secondary findings and carrier test frequencies in a large multiethnic sample. <i>Genome Medicine</i> , 2015 , 7, 54	14.4	42

155	Large-scale genome-wide association studies and meta-analyses of longitudinal change in adult lung function. <i>PLoS ONE</i> , 2014 , 9, e100776	3.7	42
154	Diabetes genes and prostate cancer in the Atherosclerosis Risk in Communities study. <i>Cancer Epidemiology Biomarkers and Prevention</i> , 2010 , 19, 558-65	4	42
153	Multi-ancestry study of blood lipid levels identifies four loci interacting with physical activity. <i>Nature Communications</i> , 2019 , 10, 376	17.4	41
152	Mining gold dust under the genome wide significance level: a two-stage approach to analysis of GWAS. <i>Genetic Epidemiology</i> , 2011 , 35, 111-8	2.6	41
151	Associations of autozygosity with a broad range of human phenotypes. <i>Nature Communications</i> , 2019 , 10, 4957	17.4	40
150	Genome-wide association study of cardiac structure and systolic function in African Americans: the Candidate Gene Association Resource (CARe) study. <i>Circulation: Cardiovascular Genetics</i> , 2013 , 6, 37-46		40
149	Multiancestry Genome-Wide Association Study of Lipid Levels Incorporating Gene-Alcohol Interactions. <i>American Journal of Epidemiology</i> , 2019 , 188, 1033-1054	3.8	39
148	Analysis of loss-of-function variants and 20 risk factor phenotypes in 8,554 individuals identifies loci influencing chronic disease. <i>Nature Genetics</i> , 2015 , 47, 640-2	36.3	39
147	Genetic association study of putative functional single nucleotide polymorphisms of genes in folate metabolism and spina bifida. <i>American Journal of Obstetrics and Gynecology</i> , 2009 , 201, 394.e1-11	6.4	38
146	Allelic variations in angiogenic pathway genes are associated with preeclampsia. <i>American Journal of Obstetrics and Gynecology</i> , 2010 , 202, 445.e1-11	6.4	38
145	LPL polymorphism predicts stroke risk in men. <i>Genetic Epidemiology</i> , 2002 , 22, 233-42	2.6	38
144	GWAS for male-pattern baldness identifies 71 susceptibility loci explaining 38% of the risk. <i>Nature Communications</i> , 2017 , 8, 1584	17.4	37
143	Genome-wide association study of gene by smoking interactions in coronary artery calcification. <i>PLoS ONE</i> , 2013 , 8, e74642	3.7	36
142	Rare Exome Sequence Variants in CLCN6 Reduce Blood Pressure Levels and Hypertension Risk. <i>Circulation: Cardiovascular Genetics</i> , 2016 , 9, 64-70		35
141	New Blood Pressure-Associated Loci Identified in Meta-Analyses of 475 000 Individuals. <i>Circulation: Cardiovascular Genetics</i> , 2017 , 10,		33
140	Sodium intake and cardiovascular disease. <i>Annual Review of Public Health</i> , 2011 , 32, 71-90	20.6	33
139	Dynamic incorporation of multiple in silico functional annotations empowers rare variant association analysis of large whole-genome sequencing studies at scale. <i>Nature Genetics</i> , 2020 , 52, 969-	9 3 633	33
138	New alcohol-related genes suggest shared genetic mechanisms with neuropsychiatric disorders. Nature Human Behaviour, 2019, 3, 950-961	12.8	32

(2012-2015)

137	Direct Estimates of the Genomic Contributions to Blood Pressure Heritability within a Population-Based Cohort (ARIC). <i>PLoS ONE</i> , 2015 , 10, e0133031	3.7	32	
136	Novel genetic associations for blood pressure identified via gene-alcohol interaction in up to 570K individuals across multiple ancestries. <i>PLoS ONE</i> , 2018 , 13, e0198166	3.7	31	
135	Multi-ancestry sleep-by-SNP interaction analysis in 126,926 individuals reveals lipid loci stratified by sleep duration. <i>Nature Communications</i> , 2019 , 10, 5121	17.4	31	
134	Association of exome sequences with plasma C-reactive protein levels in >9000 participants. <i>Human Molecular Genetics</i> , 2015 , 24, 559-71	5.6	31	
133	Large-scale whole-exome sequencing association studies identify rare functional variants influencing serum urate levels. <i>Nature Communications</i> , 2018 , 9, 4228	17.4	31	
132	Association of folate receptor (FOLR1, FOLR2, FOLR3) and reduced folate carrier (SLC19A1) genes with meningomyelocele. <i>Birth Defects Research Part A: Clinical and Molecular Teratology</i> , 2010 , 88, 689-5	94	30	
131	Practical Approaches for Whole-Genome Sequence Analysis of Heart- and Blood-Related Traits. American Journal of Human Genetics, 2017 , 100, 205-215	11	29	
130	Association of Rare Loss-Of-Function Alleles in HAL, Serum Histidine: Levels and Incident Coronary Heart Disease. <i>Circulation: Cardiovascular Genetics</i> , 2015 , 8, 351-5		29	
129	Parental history of stroke predicts subclinical but not clinical stroke: the Atherosclerosis Risk in Communities Study. <i>Stroke</i> , 2000 , 31, 2098-102	6.7	29	
128	Loss-of-function variants influence the human serum metabolome. <i>Science Advances</i> , 2016 , 2, e1600800	14.3	27	
127	Meta-analysis of 49 549 individuals imputed with the 1000 Genomes Project reveals an exonic damaging variant in ANGPTL4 determining fasting TG levels. <i>Journal of Medical Genetics</i> , 2016 , 53, 441-	9 ^{5.8}	27	
126	Discovery of Genetic Variation on Chromosome 5q22 Associated with Mortality in Heart Failure. <i>PLoS Genetics</i> , 2016 , 12, e1006034	6	26	
125	Discovery of rare variants associated with blood pressure regulation through meta-analysis of 1.3 million individuals. <i>Nature Genetics</i> , 2020 , 52, 1314-1332	36.3	26	
124	Characteristics of a spina bifida population including North American Caucasian and Hispanic individuals. <i>Birth Defects Research Part A: Clinical and Molecular Teratology</i> , 2008 , 82, 692-700		25	
123	The power of genetic diversity in genome-wide association studies of lipids. <i>Nature</i> , 2021 ,	50.4	24	
122	Metabolomic Pattern Predicts Incident Coronary Heart Disease. <i>Arteriosclerosis, Thrombosis, and Vascular Biology</i> , 2019 , 39, 1475-1482	9.4	23	
121	Rare coding variants and X-linked loci associated with age at menarche. <i>Nature Communications</i> , 2015 , 6, 7756	17.4	23	
120	Chromosome 9p21 single nucleotide polymorphisms are not associated with recurrent myocardial infarction in patients with established coronary artery disease. <i>Circulation Journal</i> , 2012 , 76, 950-6	2.9	23	

119	Variants in CXADR and F2RL1 are associated with blood pressure and obesity in African-Americans in regions identified through admixture mapping. <i>Journal of Hypertension</i> , 2012 , 30, 1970-6	1.9	23
118	Impact of Rare and Common Genetic Variants on Diabetes Diagnosis by Hemoglobin A1c in Multi-Ancestry Cohorts: The Trans-Omics for Precision Medicine Program. <i>American Journal of Human Genetics</i> , 2019 , 105, 706-718	11	22
117	Comparison of HapMap and 1000 Genomes Reference Panels in a Large-Scale Genome-Wide Association Study. <i>PLoS ONE</i> , 2017 , 12, e0167742	3.7	21
116	Integrative pathway genomics of lung function and airflow obstruction. <i>Human Molecular Genetics</i> , 2015 , 24, 6836-48	5.6	20
115	Effects of rare and common blood pressure gene variants on essential hypertension: results from the Family Blood Pressure Program, CLUE, and Atherosclerosis Risk in Communities studies. <i>Circulation Research</i> , 2013 , 112, 318-26	15.7	20
114	Genetic association of the glycine cleavage system genes and myelomeningocele. <i>Birth Defects Research Part A: Clinical and Molecular Teratology</i> , 2016 , 106, 847-853		18
113	Targeted sequencing in candidate genes for atrial fibrillation: the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Targeted Sequencing Study. <i>Heart Rhythm</i> , 2014 , 11, 452-	. / 9.7	18
112	Hypertension Susceptibility Loci are Associated with Anthracycline-related Cardiotoxicity in Long-term Childhood Cancer Survivors. <i>Scientific Reports</i> , 2017 , 7, 9698	4.9	18
111	The impact of data quality on the identification of complex disease genes: experience from the Family Blood Pressure Program. <i>European Journal of Human Genetics</i> , 2006 , 14, 469-77	5.3	18
110	Use of wrapper algorithms coupled with a random forests classifier for variable selection in large-scale genomic association studies. <i>Journal of Computational Biology</i> , 2009 , 16, 1705-18	1.7	17
109	Genetic loci associated with ideal cardiovascular health: A meta-analysis of genome-wide association studies. <i>American Heart Journal</i> , 2016 , 175, 112-20	4.9	17
108	A genome-wide association study identifies new loci for factor VII and implicates factor VII in ischemic stroke etiology. <i>Blood</i> , 2019 , 133, 967-977	2.2	17
107	Genome-wide association study of 1,5-anhydroglucitol identifies novel genetic loci linked to glucose metabolism. <i>Scientific Reports</i> , 2017 , 7, 2812	4.9	16
106	FastSKAT: Sequence kernel association tests for very large sets of markers. <i>Genetic Epidemiology</i> , 2018 , 42, 516-527	2.6	16
105	Whole-genome sequencing study of serum peptide levels: the Atherosclerosis Risk in Communities study. <i>Human Molecular Genetics</i> , 2017 , 26, 3442-3450	5.6	16
104	Single nucleotide polymorphisms associated with coronary heart disease predict incident ischemic stroke in the atherosclerosis risk in communities study. <i>Cerebrovascular Diseases</i> , 2008 , 26, 420-4	3.2	16
103	Meta-analysis of exome array data identifies six novel genetic loci for lung function. <i>Wellcome Open Research</i> , 2018 , 3, 4	4.8	16
102	Supplemental Association of Clonal Hematopoiesis With Incident Heart Failure. <i>Journal of the American College of Cardiology</i> , 2021 , 78, 42-52	15.1	16

(2016-2014)

101	in Genomic Epidemiology (CHARGE) Consortium Targeted Sequencing Study. <i>Circulation:</i> Cardiovascular Genetics, 2014 , 7, 335-43		15
100	Association of copper-zinc superoxide dismutase (SOD1) and manganese superoxide dismutase (SOD2) genes with nonsyndromic myelomeningocele. <i>Birth Defects Research Part A: Clinical and Molecular Teratology</i> , 2012 , 94, 762-9		15
99	ESR1 polymorphism is associated with plasma lipid and apolipoprotein levels in Caucasians of the Rochester Family Heart Study. <i>Journal of Lipid Research</i> , 2008 , 49, 1701-6	6.3	15
98	Sequence analysis of six blood pressure candidate regions in 4,178 individuals: the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) targeted sequencing study. <i>PLoS ONE</i> , 2014 , 9, e109155	3.7	15
97	An Empirical Comparison of Joint and Stratified Frameworks for Studying G IE Interactions: Systolic Blood Pressure and Smoking in the CHARGE Gene-Lifestyle Interactions Working Group. <i>Genetic Epidemiology</i> , 2016 , 40, 404-15	2.6	15
96	Leveraging linkage evidence to identify low-frequency and rare variants on 16p13 associated with blood pressure using TOPMed whole genome sequencing data. <i>Human Genetics</i> , 2019 , 138, 199-210	6.3	14
95	A multi-ancestry genome-wide study incorporating gene-smoking interactions identifies multiple new loci for pulse pressure and mean arterial pressure. <i>Human Molecular Genetics</i> , 2019 , 28, 2615-2633	5.6	14
94	Metabolomics Identifies Novel Blood Biomarkers of Pulmonary Function and COPD in the General Population. <i>Metabolites</i> , 2019 , 9,	5.6	14
93	Common Coding Variants in Are Associated With the Nav1.8 Late Current and Cardiac Conduction. Circulation Genomic and Precision Medicine, 2018 , 11, e001663	5.2	14
92	Mutations in folate transporter genes and risk for human myelomeningocele. <i>American Journal of Medical Genetics, Part A</i> , 2017 , 173, 2973-2984	2.5	13
91	Dynamic Scan Procedure for Detecting Rare-Variant Association Regions in Whole-Genome Sequencing Studies. <i>American Journal of Human Genetics</i> , 2019 , 104, 802-814	11	13
90	GOSR2 Lys67Arg is associated with hypertension in whites. <i>American Journal of Hypertension</i> , 2009 , 22, 163-8	2.3	13
89	Pharmacogenomics of statin-related myopathy: Meta-analysis of rare variants from whole-exome sequencing. <i>PLoS ONE</i> , 2019 , 14, e0218115	3.7	12
88	Association of retinoic acid receptor genes with meningomyelocele. <i>Birth Defects Research Part A:</i> Clinical and Molecular Teratology, 2011 , 91, 39-43		12
87	Evaluating the context-dependent effect of family history of stroke in a genome scan for hypertension. <i>Stroke</i> , 2003 , 34, 1170-5	6.7	12
86	Genome scan for hypertension in nonobese African Americans: the National Heart, Lung, and Blood Institute Family Blood Pressure Program. <i>American Journal of Hypertension</i> , 2004 , 17, 834-8	2.3	12
85	Population genomic analysis of 962 whole genome sequences of humans reveals natural selection in non-coding regions. <i>PLoS ONE</i> , 2015 , 10, e0121644	3.7	12
84	Whole genome sequence analysis of serum amino acid levels. <i>Genome Biology</i> , 2016 , 17, 237	18.3	12

83	Genome-Wide Association Study of Serum Fructosamine and Glycated Albumin in Adults Without Diagnosed Diabetes: Results From the Atherosclerosis Risk in Communities Study. <i>Diabetes</i> , 2018 , 67, 1684-1696	0.9	12
82	Mendelian randomization evaluation of causal effects of fibrinogen on incident coronary heart disease. <i>PLoS ONE</i> , 2019 , 14, e0216222	3.7	11
81	Variants for HDL-C, LDL-C, and triglycerides identified from admixture mapping and fine-mapping analysis in African American families. <i>Circulation: Cardiovascular Genetics</i> , 2015 , 8, 106-13		11
80	RNA sequence analysis of inducible pluripotent stem cell-derived cardiomyocytes reveals altered expression of DNA damage and cell cycle genes in response to doxorubicin. <i>Toxicology and Applied Pharmacology</i> , 2018 , 356, 44-53	4.6	11
79	DNA methylation age is associated with an altered hemostatic profile in a multiethnic meta-analysis. <i>Blood</i> , 2018 , 132, 1842-1850	2.2	11
78	Genetic architecture of lipid traits in the Hispanic community health study/study of Latinos. <i>Lipids in Health and Disease</i> , 2017 , 16, 200	4.4	11
77	Genome-wide linkage study of erythrocyte sodium-lithium countertransport. <i>American Journal of Hypertension</i> , 2005 , 18, 653-6	2.3	11
76	Folate metabolism gene 5,10-methylenetetrahydrofolate reductase (MTHFR) is associated with ADHD in myelomeningocele patients. <i>PLoS ONE</i> , 2012 , 7, e51330	3.7	11
75	Association of levels of fasting glucose and insulin with rare variants at the chromosome 11p11.2-MADD locus: Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium Targeted Sequencing Study. <i>Circulation: Cardiovascular Genetics</i> , 2014 , 7, 374-382		9
74	Genetic variation in solute carrier genes is associated with preeclampsia. <i>American Journal of Obstetrics and Gynecology</i> , 2010 , 203, 491.e1-491.e13	6.4	9
73	A whole-genome scan for stroke or myocardial infarction in family blood pressure program families. <i>Stroke</i> , 2008 , 39, 1115-20	6.7	9
72	Aptamer-Based Proteomic Platform Identifies Novel Protein Predictors of Incident Heart Failure and Echocardiographic Traits. <i>Circulation: Heart Failure</i> , 2020 , 13, e006749	7.6	8
71	Role of Rare and Low-Frequency Variants in Gene-Alcohol Interactions on Plasma Lipid Levels. <i>Circulation Genomic and Precision Medicine</i> , 2020 , 13, e002772	5.2	8
70	A Genome-wide Association Study Discovers 46 Loci of the Human Metabolome in the Hispanic Community Health Study/Study of Latinos. <i>American Journal of Human Genetics</i> , 2020 , 107, 849-863	11	8
69	Rare coding variants associated with blood pressure variation in 15 914 individuals of African ancestry. <i>Journal of Hypertension</i> , 2017 , 35, 1381-1389	1.9	7
68	Genetic variants in microRNA genes and targets associated with cardiovascular disease risk factors in the African-American population. <i>Human Genetics</i> , 2018 , 137, 85-94	6.3	7
67	Maternal hypertension and risk for hypospadias in offspring. <i>American Journal of Medical Genetics, Part A</i> , 2016 , 170, 3125-3132	2.5	7
66	Sequencing of SCN5A identifies rare and common variants associated with cardiac conduction: Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium. <i>Circulation: Cardiovascular Genetics</i> , 2014 , 7, 365-73		7

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65	Association of NOS3 Glu298Asp SNP with hypertension and possible effect modification of dietary fat intake in the ARIC study. <i>Hypertension Research</i> , 2010 , 33, 165-9	4.7	7	
64	Clonal Hematopoiesis Is Associated With Higher Risk of Stroke. <i>Stroke</i> , 2021 , STROKEAHA121037388	6.7	7	
63	ADAM19 and HTR4 variants and pulmonary function: Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium Targeted Sequencing Study. <i>Circulation: Cardiovascular Genetics</i> , 2014 , 7, 350-8		6	
62	Association between NEDD4L gene and sodium lithium countertransport. <i>American Journal of Hypertension</i> , 2011 , 24, 145-8	2.3	6	
61	Comprehensive linkage and linkage heterogeneity analysis of 4344 sibling pairs affected with hypertension from the Family Blood Pressure Program. <i>Genetic Epidemiology</i> , 2007 , 31, 195-210	2.6	6	
60	Genome scan for hypertension in nonobese African Americans: The national heart, lung, and blood institute family blood pressure program. <i>American Journal of Hypertension</i> , 2004 , 17, 834-838	2.3	6	
59	American Heart Association Life Simple 7: Lifestyle Recommendations, Polygenic Risk, and Lifetime Risk of Coronary Heart Disease <i>Circulation</i> , 2022 ,	16.7	6	
58	Meta-analysis of exome array data identifies six novel genetic loci for lung function. <i>Wellcome Open Research</i> , 3, 4	4.8	6	
57	Whole genome sequence analysis of pulmonary function and COPD in 19,996 multi-ethnic participants. <i>Nature Communications</i> , 2020 , 11, 5182	17.4	6	
56	A Mendelian randomization of B and total fibrinogen levels in relation to venous thromboembolism and ischemic stroke. <i>Blood</i> , 2020 , 136, 3062-3069	2.2	6	
55	Fine mapping the region reveals a common intronic insertion associated to HDL-C. <i>Npj Aging and Mechanisms of Disease</i> , 2015 , 1, 15011	5.5	5	
54	Sequence variation in TMEM18 in association with body mass index: Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium Targeted Sequencing Study. <i>Circulation: Cardiovascular Genetics</i> , 2014 , 7, 344-9		5	
53	Association of SLC34A2 variation and sodium-lithium countertransport activity in humans and baboons. <i>American Journal of Hypertension</i> , 2009 , 22, 288-93	2.3	5	
52	Linkage disequilibrium structure and its impact on the localization of a candidate functional mutation. <i>Genetic Epidemiology</i> , 2001 , 21 Suppl 1, S620-5	2.6	5	
51	Efficient gene-environment interaction tests for large biobank-scale sequencing studies. <i>Genetic Epidemiology</i> , 2020 , 44, 908-923	2.6	5	
50	Chromosome Xq23 is associated with lower atherogenic lipid concentrations and favorable cardiometabolic indices. <i>Nature Communications</i> , 2021 , 12, 2182	17.4	5	
49	Whole-genome sequencing association analysis of quantitative red blood cell phenotypes: The NHLBI TOPMed program. <i>American Journal of Human Genetics</i> , 2021 , 108, 874-893	11	5	
48	Determinants of penetrance and variable expressivity in monogenic metabolic conditions across 77,184 exomes. <i>Nature Communications</i> , 2021 , 12, 3505	17.4	5	

47	Discovery and systematic characterization of risk variants and genes for coronary artery disease in over a million participants		5
46	Functionally oriented analysis of cardiometabolic traits in a trans-ethnic sample. <i>Human Molecular Genetics</i> , 2019 , 28, 1212-1224	5.6	5
45	Contributions of rare coding variants in hypotension syndrome genes to population blood pressure variation. <i>Medicine (United States)</i> , 2018 , 97, e11865	1.8	5
44	Sequence-Based Analysis of Lipid-Related Metabolites in a Multiethnic Study. <i>Genetics</i> , 2018 , 209, 607-	646	4
43	Association of the IGF1 gene with fasting insulin levels. <i>European Journal of Human Genetics</i> , 2016 , 24, 1337-43	5.3	4
42	Regional association-based fine-mapping for sodium-lithium countertransport on chromosome 10. <i>American Journal of Hypertension</i> , 2008 , 21, 117-21	2.3	4
41	Genetic loci associated with prevalent and incident myocardial infarction and coronary heart disease in the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium. <i>PLoS ONE</i> , 2020 , 15, e0230035	3.7	4
40	Genetic analysis of over one million people identifies 535 novel loci for blood pressure		4
39	Mendelian Randomization Analysis of Hemostatic Factors and Their Contribution to Peripheral Artery Disease-Brief Report. <i>Arteriosclerosis, Thrombosis, and Vascular Biology</i> , 2021 , 41, 380-386	9.4	4
38	Association between ABO haplotypes and the risk of venous thrombosis: impact on disease risk estimation. <i>Blood</i> , 2021 , 137, 2394-2402	2.2	4
37	Maternal gene-micronutrient interactions related to one-carbon metabolism and the risk of myelomeningocele among offspring. <i>Birth Defects Research</i> , 2017 , 109, 99-105	2.9	3
36	Gene-educational attainment interactions in a multi-ancestry genome-wide meta-analysis identify novel blood pressure loci. <i>Molecular Psychiatry</i> , 2021 , 26, 2111-2125	15.1	3
35	Effects of Gender-Specific Differences, Inflammatory Response, and Genetic Variation on the Associations Among Depressive Symptoms and the Risk of Major Adverse Coronary Events in Patients With Acute Coronary Syndrome. <i>Biological Research for Nursing</i> , 2018 , 20, 168-176	2.6	3
34	Gene-by-Psychosocial Factor Interactions Influence Diastolic Blood Pressure in European and African Ancestry Populations: Meta-Analysis of Four Cohort Studies. <i>International Journal of Environmental Research and Public Health</i> , 2017 , 14,	4.6	3
33	Burden of rare deleterious variants in WNT signaling genes among 511 myelomeningocele patients. <i>PLoS ONE</i> , 2020 , 15, e0239083	3.7	3
32	Multi-ancestry genome-wide gene-sleep interactions identify novel loci for blood pressure. Molecular Psychiatry, 2021,	15.1	3
31	Mendelian randomization supports bidirectional causality between telomere length and clonal hematopoiesis of indeterminate potential <i>Science Advances</i> , 2022 , 8, eabl6579	14.3	3
30	Association of Variants with Blood Pressure in the Atherosclerosis Risk in Communities Study. International Journal of Hypertension, 2019, 2019, 2137629	2.4	2

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29	The impact of multiple single day blood pressure readings on cardiovascular risk estimation: The Atherosclerosis Risk in Communities study. <i>European Journal of Preventive Cardiology</i> , 2016 , 23, 1529-3	6 ^{3.9}	2
28	Association between SLC20A1 and sodium-lithium countertransport. <i>American Journal of Hypertension</i> , 2011 , 24, 1069-72	2.3	2
27	Genome-wide association study provides new insights into the genetic architecture and pathogenesis of heart failure		2
26	Rare Non-coding Variation Identified by Large Scale Whole Genome Sequencing Reveals Unexplained Heritability of Type 2 Diabetes		2
25	Rare coding variants in 35 genes associate with circulating lipid levels & multi-ancestry analysis of 170,000 exomes		2
24	Genetic discovery and translational decision support from exome sequencing of 20,791 type 2 diabetes cases and 24,440 controls from five ancestries		2
23	Efficient variant set mixed model association tests for continuous and binary traits in large-scale whole genome sequencing studies		2
22	Identification of novel candidate risk genes for myelomeningocele within the glucose homeostasis/oxidative stress and folate/one-carbon metabolism networks. <i>Molecular Genetics & Mamp; Genomic Medicine</i> , 2020 , 8, e1495	2.3	2
21	Genetic susceptibility, obesity and lifetime risk of type 2 diabetes: The ARIC study and Rotterdam Study. <i>Diabetic Medicine</i> , 2021 , 38, e14639	3.5	2
20	Identifying blood pressure loci whose effects are modulated by multiple lifestyle exposures. <i>Genetic Epidemiology</i> , 2020 , 44, 629-641	2.6	1
19	Meta-analysis of exome array data identifies six novel genetic loci for lung function. <i>Wellcome Open Research</i> , 3, 4	4.8	1
18	FastSKAT: Sequence kernel association tests for very large sets of markers		1
17	Novel blood pressure locus and gene discovery using GWAS and expression datasets from blood and the kidney		1
16	Multi-ancestry analysis of gene-sleep interactions in 126,926 individuals identifies multiple novel blood lipid loci that contribute to our understanding of sleep-associated adverse blood lipid profile		1
15	CLUE: Exact maximal reduction of kinetic models by constrained lumping of differential equations. <i>Bioinformatics</i> , 2021 ,	7.2	1
14	FGL1 as a modulator of plasma D-dimer levels: Exome-wide marker analysis of plasma tPA, PAI-1, and D-dimer. <i>Journal of Thrombosis and Haemostasis</i> , 2021 , 19, 2019-2028	15.4	1
13	Leveraging a health information exchange for analyses of COVID-19 outcomes including an example application using smoking history and mortality. <i>PLoS ONE</i> , 2021 , 16, e0247235	3.7	1
12	Maternal Hypertension-Related Genotypes and Congenital Heart Defects. <i>American Journal of Hypertension</i> , 2021 , 34, 82-91	2.3	1

11	BinomiRare: A robust test for association of a rare genetic variant with a binary outcome for mixed models and any case-control proportion. <i>Human Genetics and Genomics Advances</i> , 2021 , 2,	0.8	1
10	Identification of novel and rare variants associated with handgrip strength using whole genome sequence data from the NHLBI Trans-Omics in Precision Medicine (TOPMed) Program. <i>PLoS ONE</i> , 2021 , 16, e0253611	3.7	1
9	The genomics of heart failure: design and rationale of the HERMES consortium. <i>ESC Heart Failure</i> , 2021 ,	3.7	1
8	Whole-genome sequencing in diverse subjects identifies genetic correlates of leukocyte traits: The NHLBI TOPMed program. <i>American Journal of Human Genetics</i> , 2021 , 108, 1836-1851	11	1
7	Rare and low-frequency exonic variants and gene-by-smoking interactions in pulmonary function. <i>Scientific Reports</i> , 2021 , 11, 19365	4.9	O
6	Epidemiologic and Genetic Associations of Erythropoietin With Blood Pressure, Hypertension, and Coronary Artery Disease. <i>Hypertension</i> , 2021 , 78, 1555-1566	8.5	0
5	Elucidating mechanisms of genetic cross-disease associations at the PROCR vascular disease locus <i>Nature Communications</i> , 2022 , 13, 1222	17.4	O
4	Large-Scale Genomic Analyses Link Reproductive Aging to Hypothalamic Signaling, Breast Cancer Susceptibility, and BRCA1-Mediated DNA Repair. <i>Obstetrical and Gynecological Survey</i> , 2015 , 70, 758-76	2 ^{2.4}	
3	Association of SERPINA9 gene variants with carotid artery atherosclerosis: the Atherosclerosis Risk in Communities (ARIC) Carotid MRI Study. <i>International Journal of Molecular Epidemiology and Genetics</i> , 2013 , 4, 258-67	0.9	
2	Reply to RMisestimation of heritability and prediction accuracy of male-pattern baldnessR <i>Nature Communications</i> , 2018 , 9, 2538	17.4	
1	Rare coding variants in RCN3 are associated with blood pressure <i>BMC Genomics</i> , 2022 , 23, 148	4.5	