## Henry J Lin

## List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/847510/publications.pdf

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430874 345221 1,609 41 18 36 h-index citations g-index papers 45 45 45 3452 citing authors all docs docs citations times ranked

#	Article	IF	CITATIONS
1	Multi-ethnic genome-wide association study for atrial fibrillation. Nature Genetics, 2018, 50, 1225-1233.	21.4	552
2	Genetic loci associated with heart rate variability and their effects on cardiac disease risk. Nature Communications, 2017, 8, 15805.	12.8	95
3	Altered cofactor regulation with disease-associated p97/VCP mutations. Proceedings of the National Academy of Sciences of the United States of America, 2015, 112, E1705-14.	7.1	87
4	Genetic Risk Prediction of Atrial Fibrillation. Circulation, 2017, 135, 1311-1320.	1.6	87
5	PR interval genome-wide association meta-analysis identifies 50 loci associated with atrial and atrioventricular electrical activity. Nature Communications, 2018, 9, 2904.	12.8	71
6	A multiancestry genome-wide association study of unexplained chronic ALT elevation as a proxy for nonalcoholic fatty liver disease with histological and radiological validation. Nature Genetics, 2022, 54, 761-771.	21.4	68
7	Multi-ancestry GWAS of the electrocardiographic PR interval identifies 202 loci underlying cardiac conduction. Nature Communications, 2020, $11,2542$ .	12.8	59
8	Hematopoietic Prostaglandin D Synthase Suppresses Intestinal Adenomas in ApcMin/+ Mice. Cancer Research, 2007, 67, 881-889.	0.9	55
9	Exstrophy of the cloaca in a 47,XXX child: Review of genitourinary malformations in tripleâ€X patients. American Journal of Medical Genetics Part A, 1993, 45, 761-763.	2.4	50
10	Anomalous inferior and superior venae cavae with oculoauriculovertebral defect: Review of Goldenhar complex and malformations of left-right asymmetry., 1998, 75, 88-94.		32
11	Intrachromosomal triplication of $2q11.2$ - $q21$ in a severely malformed infant: Case report and review of triplications and their possible mechanism. American Journal of Medical Genetics Part A, 1999, 82, 312-317.	2.4	32
12	Genome-wide association study of iron traits and relation to diabetes in the Hispanic Community Health Study/Study of Latinos (HCHS/SOL): potential genomic intersection of iron and glucose regulation?. Human Molecular Genetics, 2017, 26, 1966-1978.	2.9	31
13	Asymptomatic Maternal Combined Homocystinuria and Methylmalonic Aciduria (cblC) Detected through Low Carnitine Levels on Newborn Screening. Journal of Pediatrics, 2009, 155, 924-927.	1.8	29
14	Mosaic tetrasomy 8q: Inverted duplication of 8q23.3qter in an analphoid marker. American Journal of Medical Genetics Part A, 2000, 92, 69-76.	2.4	27
15	Glutathione transferase GSTT1, broccoli, and prevalence of colorectal adenomas. Pharmacogenetics and Genomics, 2002, 12, 175-179.	5 <b>.</b> 7	27
16	ExomeChip-Wide Analysis of 95 626 Individuals Identifies 10 Novel Loci Associated With QT and JT Intervals. Circulation Genomic and Precision Medicine, 2018, 11, e001758.	3.6	27
17	Effects of Calcium, Magnesium, and Potassium Concentrations on Ventricular Repolarization in Unselected Individuals. Journal of the American College of Cardiology, 2019, 73, 3118-3131.	2.8	27
18	A multi-ethnic polygenic risk score is associated with hypertension prevalence and progression throughout adulthood. Nature Communications, 2022, 13, .	12.8	27

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19	Intestinal tumor suppression in Apc Min/+ mice by prostaglandin D 2 receptor PTGDR. Cancer Medicine, 2014, 3, 1041-1051.	2.8	26
20	Occipital encephalocele and MURCS association: Case report and review of central nervous system anomalies in MURCS patients., 1996, 61, 59-62.		22
21	Common and Rare Coding Genetic Variation Underlying the Electrocardiographic PR Interval. Circulation Genomic and Precision Medicine, 2018, 11, e002037.	3.6	19
22	DOOR syndrome (deafness, onycho-osteodystrophy, and mental retardation): A new patient and delineation of neurologic variability among recessive cases. American Journal of Medical Genetics Part A, 1993, 47, 534-539.	2.4	14
23	Monogenic and Polygenic Contributions to QTc Prolongation in the Population. Circulation, 2022, 145, 1524-1533.	1.6	14
24	Naturally occurring Phe151Leu substitution near a conserved folding module lowers stability of glutathione transferase P1–1. Biochimica Et Biophysica Acta - Proteins and Proteomics, 2003, 1649, 16-23.	2.3	13
25	Evaluating p97 Inhibitor Analogues for Potency against p97–p37 and p97–Npl4–Ufd1 Complexes. ChemMedChem, 2016, 11, 953-957.	3.2	13
26	Omphalocele with absent radial ray (ORR): A case with diploid-triploid mixoploidy. American Journal of Medical Genetics Part A, 1998, 75, 235-239.	2.4	12
27	An Outbreak of Polygenic Scores for Coronary Artery Disease. Journal of the American College of Cardiology, 2020, 75, 2781-2784.	2.8	11
28	Use of HLA marker associations and HLA haplotype linkage to estimate disease risks in families with gluten-sensitive enteropathy. Clinical Genetics, 2008, 28, 185-198.	2.0	10
29	Lego bricks and the octet rule: Molecular models for biochemical pathways with plastic, interlocking toy bricks. Biochemistry and Molecular Biology Education, 2018, 46, 54-57.	1.2	10
30	GWAS of QRS duration identifies new loci specific to Hispanic/Latino populations. PLoS ONE, 2019, 14, e0217796.	2.5	8
31	Home use of a compact, 12‑lead ECG recording system for newborns. Journal of Electrocardiology, 2019, 53, 89-94.	0.9	7
32	Rare Coding Variants Associated With Electrocardiographic Intervals Identify Monogenic Arrhythmia Susceptibility Genes: A Multi-Ancestry Analysis. Circulation Genomic and Precision Medicine, 2021, 14, e003300.	3.6	7
33	Community Partnership in Precision Medicine: Themes from a Community Engagement Conference. Ethnicity and Disease, 2018, 28, 503-510.	2.3	6
34	Identification of Functional Genetic Determinants of Cardiac Troponin T and I in a Multiethnic Population and Causal Associations With Atrial Fibrillation. Circulation Genomic and Precision Medicine, 2021, 14, CIRCGEN121003460.	3.6	5
35	Glucose-6-phosphate dehydrogenase deficiency presenting with rhabdomyolysis in a patient with coronavirus disease 2019 pneumonia: aÂcase report. Journal of Medical Case Reports, 2022, 16, 106.	0.8	5
36	Disease risk estimates from marker association data: Application to individuals at risk for hemochromatosis. Clinical Genetics, 1985, 27, 127-133.	2.0	3

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#	Article	lF	CITATIONS
37	High Dietary Niacin May Increase Prostaglandin Formation but Does Not Increase Tumor Formation in <i>Apc</i> <sup>Min/+</sup> Mice. Nutrition and Cancer, 2011, 63, 950-959.	2.0	2
38	A tortuous proximal urethra in urorectal septum malformation sequence?. American Journal of Medical Genetics, Part A, 2014, 164, 1298-1303.	1.2	2
39	Infantile Marfan's Syndrome. Circulation, 1998, 97, 1103-1104.	1.6	0
40	Neonatal Marfan Syndrome. Fetal and Pediatric Pathology, 2001, 20, 235-239.	0.3	0
41	NEONATAL MARFAN SYNDROME. Fetal and Pediatric Pathology, 2001, 20, 235-239.	0.3	0