



Core Project R03OD032630





Details








Projects	Name	Award	Publications	Repositories	Analytics
1R03OD032630-01	Methods to maximize the utility of common fund functional genomic data in multi-ethnic genetic studies	\$335,407.00	9 publications	0 repositories	0 properties










Publications

Published works associated with this project.

ID	Title	Authors	RC R	SJR	Cita tion s	Cit./ yea r	Journal	Publ ishe d	Upda ted
30643251  DOI 	Association studies of up to 1.2 million individuals yield new insights into the	Liu, Mengzhen	65. 27	0	1,41 1	235. 167	Nat Genet	2019	Dec 5,

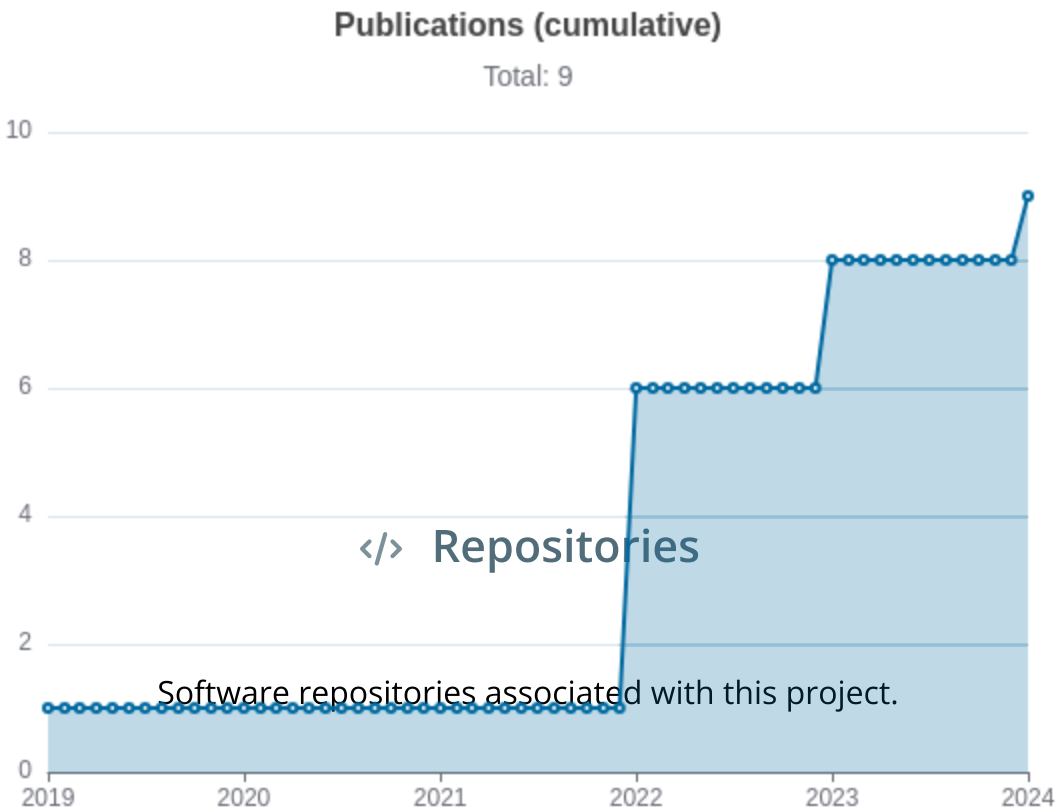
	genetic etiology...	...123 more... Vrieze, Scott	2							2025 (just now)
36477530  DOI 	Genetic diversity fuels gene discovery for tobacco and alcohol use.	Saunders, Gretchen R B ...217 more... Vrieze, Scott	30.774	18.288	336	112	Nature	2022		Dec 5, 2025 (just now)
36750564  DOI 	Multi-ancestry and multi-trait genome-wide association meta-analyses inform clinical risk predict...	Khunsrirak sakul, Chachrit ...16 more... Liu, Dajiang J	3.95	0	37	18.5	Nat Comm un	2023		Dec 5, 2025 (just now)
36702996  DOI 	Multi-ancestry transcriptome-wide association analyses yield insights into tobacco use biology an...	Chen, Fang ...88 more... Liu, Dajiang J	3.786	0	37	18.5	Nat Genet	2023		Dec 5, 2025 (just now)
35672318  DOI 	Integrating 3D genomic and epigenomic data to enhance target gene discovery and drug repurposing ...	Khunsrirak sakul, Chachrit ...12 more...	2.281	0	33	11	Nat Comm un	2022		Dec 5, 2025

		Liu, Dajiang J							(just now)
35927319  DOI 	Rare genetic variants explain missing heritability in smoking.	Jang, Seon-Kyeong ...88 more... Vrieze, Scott	1.4 55	0	19	6.33 3	Nat Hum Behav	2022	Dec 5, 2025 (just now)
35833142  DOI 	Construction and Application of Polygenic Risk Scores in Autoimmune Diseases.	Khunsrirak sakul, Chachrit ...4 more... Liu, Dajiang J	1.0 35	0	13	4.33 3	Front Immunol	2022	Dec 5, 2025 (just now)
38918381  DOI 	Dissecting heritability, environmental risk, and air pollution causal effects using > 50 milli...	McGuire, Daniel ...8 more... Jiang, Bibo	0.9 73	0	2	2	Nat Commun	2024	Dec 5, 2025 (just now)
35178743  DOI 	Assessing reproducibility of high-throughput experiments in the case of missing data.	Singh, Roopali ...1 more... Li, Qunhua	0.5 24	0	6	2	Stat Med	2022	Dec 5, 2025 (just now)

Notes

RCR [Relative Citation Ratio](#)

SJR [Scimago Journal Rank](#)



Name	Description	Stars	Watchers	Forks	Issues	PRs	Commits	Contrib.
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No data

Name	Tags	Last Commit	Avg Issue	Avg PR	Languages	License	Readme	Contributing	Dependencies
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No data

Notes

Repository For storing, tracking changes to, and collaborating on a piece of software.

PR "Pull request", a draft change (new feature, bug fix, etc.) to a repo.

Closed/Open Resolved/unresolved.

Avg Issue/PR Average time issues/pull requests stay open for before being closed.

Only the `main` /default branch is considered for metrics like # of commits.

of dependencies is totaled from all manifests in repo, direct and transitive, e.g. `package.json` + `package-lock.json`.

Analytics

Traffic metrics of websites associated with this project.

Notes

Active Users [Distinct users who visited the website](#) .

New Users [Users who visited the website for the first time](#) .

Engaged Sessions [Visits that had significant interaction](#) .

"Top" metrics are measured by number of engaged sessions.