Depression GWAS meta-analysis results excluding PGC2 and 23andMe, April 2023 release (iPSYCH2015+UKB+FinnGen+MVP data)

The file

'daner_MDDwoBP_20201001_HRC_MDDwoBP_UKBtransformed_FinnGen_MVPaf _2_HRC_MAF01.gz' contains results from a GWAS meta-analysis of depression corresponding to the primary meta-analysis reported in the paper '*Depression pathophysiology, risk prediction of recurrence and comorbid psychiatric disorders using genome-wide analyses*' published in Nature Medicine, 2023 (DOI), but with the GWAS meta-analysis reported in Howard et al. 2019¹ (excluding all Danish samples) replaced by the UKB depression summary stats from Howard et al. 2019¹, thus not including any PGC or 23andMe summary statistics. To access the summary statistics from the meta-analysis of all cohorts, including 23andMe, a data transfer agreement is required from 23andMe (dataset-request@23andMe.com) before a request is made to the corresponding authors. See

https://research.23andme.com/collaborate/#dataset-access/ for more information and to apply for access to the data.

Citation for studies using these data

T. D. Als, M. Kurki, J. Grove, G. Voloudakis, K. Therrien, E. Tasanko, T. T. Nielsen, J. Naamanka, K. Veerapen, D. Levey, J. Bendl, J. Bybjerg-Grauholm, B. Zheng, D. Demontis, A. Rosengren, G. Athanasiadis, M. Bækved-Hansen, P. Qvist, G. B. Walters, T. Thorgeirsson, H. Stefánsson, K. L. Musliner, V. M. Rajagopal, L. Farajzadeh, J. Thirstrup, B. J. Vilhjálmsson, J. J. McGrath, M. Mattheisen, S. Meier, E. Agerbo, K. Stefánsson, M. Nordentoft, T. Werge, D. M. Hougaard, P. B. Mortensen, M. B. Stein, J. Gelernter, I. Hovatta, P. Roussos, M. J. Daly, O. Mors, A. Palotie, and A. D. Børglum (April 2023). *Depression pathophysiology, risk prediction of recurrence and comorbid psychiatric disorders using genome-wide analyses*. Nature Medicine (NMED-A120965C).

Disclaimer

These data are provided "as is", and without warranty, for scientific and educational use only. If you download these data, you acknowledge that these data will be used only for non-commercial research purposes; that the investigator is in compliance

with all applicable state, local, and federal laws or regulations and institutional policies regarding human subjects and genetics research; that secondary distribution of the data without registration by secondary parties is prohibited; and that the investigator will cite the publication in any communications or publications arising directly or indirectly from these data.

Data Use Agreement

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- Investigators will use these results for scientific research and educational use only;
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- Investigators certify that they are in compliance with all applicable local, state, and federal laws or regulations and institutional policies regarding human subjects and genetics research;
- Investigators will cite the appropriate publication in any communications or publications arising directly or indirectly from these data;
- Investigators will never attempt to identify any participant who contributed to these data;
- 7) Investigators may not use these data to develop any type of risk or predictive test for an unborn individual;
- For any risk or predictive test for a child or adult, investigators must acknowledge that this is an experimental use of these data and that essentially all psychiatric disorders have important non-genetic etiological components;
- 9) When these data are made available prior to publication, investigators agree to respect and not compete with the scientific priorities of the iPSYCH team according to the Fort Lauderdale principles.

Appropriate use of these data requires considerable attention to detail, prior experience, and technical skill. Errors are easy to make. If investigators use these data, any and all consequences are entirely their responsibility.

File Description

daner_MDDwoBP_20201001_HRC_MDDwoBP_UKBtransformed_FinnGen_MVP

af_2_HRC_MAF01.gz: A subset of the primary depression GWAS meta-analysis of samples of European ancestry, excluding 23andMe and PGC, thus with the GWAS meta-analysis reported in Howard et al. 2019¹ (excluding all Danish samples) replaced by the UKB depression summary stats from Howard et al. 2019¹.

CHR: Chromosome (hg19)

SNP: Marker name

BP: Base pair location (hg19)

A1: Reference allele for OR (may or may not be minor allele)

A2: Alternative allele

FRQ_A_268615 : allele frequency in 268615 cases

FRQ_U_667123: allele frequency in 667123 controls

INFO: Imputation information score (the reported imputation INFO score is a

weighted average across the cohorts, for which INFO score was available,

contributing to the meta-analysis for that variant)

OR: Odds ratio for the effect of the A1 allele

SE: Standard error of the log(OR)

P: P-value for association test in the meta-analysis

Direction: direction of effect in the included cohorts

Additional Notes

MD5

(daner_MDDwoBP_20201001_HRC_MDDwoBP_UKBtransformed_FinnGen_MVPaf _2_HRC_MAF01.gz) = b9948582ad612897628e8e2d2d34edf9

The reported imputation INFO score is a weighted average across the cohorts contributing to meta-analysis for that variant.

 Howard, D. M. *et al.* Genome-wide meta-analysis of depression identifies 102 independent variants and highlights the importance of the prefrontal brain regions. *Nat Neurosci* 22, 343-352 (2019). <u>https://doi.org:10.1038/s41593-018-0326-7</u>