NPDR Supplementary Material

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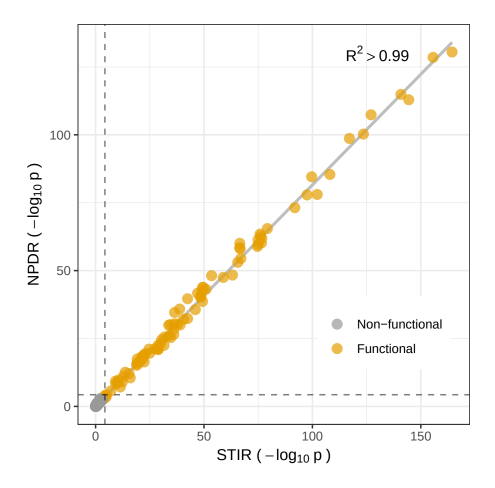


Figure S1: Similarity between NPDR and STIR for one simulation of m=200 samples and p=1000 attributes. In 100 replications, R^2 ranges from 0.9827 to 0.9994. STIR is based on a t-test of projected distances and NPDR is based on a logistic regression of projected distances.

References

- [1] Trang T. Le, Jonathan Savitz, Hideo Suzuki, Masaya Misaki, T. Kent Teague, Bill C. White, Julie H. Marino, Graham Wiley, Patrick M. Gaffney, Wayne C. Drevets, Brett A. McKinney, and Jerzy Bodurka. Identification and replication of RNA-Seq gene network modules associated with depression severity. *Translational Psychiatry*, 8(1):180, September 2018.
- [2] Trang T Le, Ryan J Urbanowicz, Jason H Moore, and Brett A McKinney. Statistical inference relief (stir) feature selection. *Bioinformatics*, 2018.

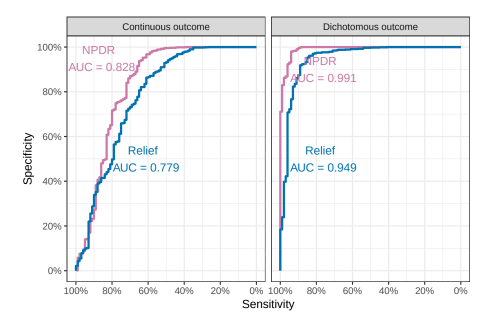


Figure S2: Receiver Operating Characteristics (ROC) curves for Relief-F and NPDR for simulated case-control data with interactions (left) and RRelief and NPDR for simulated continuous outcome data with main effects (right). Simulation uses m=200 samples and p=1000 attributes with 100 functional.

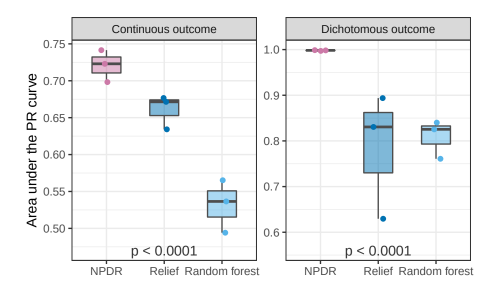


Figure S3: NPDR and Relief comparison of area under the PRC for 100 replicate simulations of case-control (left) and continuous (right) data. All simulations use m=200 samples and p=1000 attributes with 100 functional. NPDR yields significantly higher auPRC.

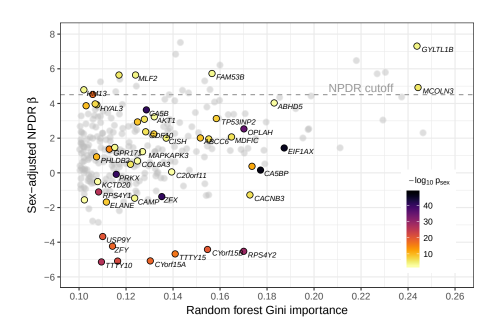


Figure S4: NPDR vs random forest feature importance. Gene scatter plot of importance score for association with major depressive disorder using random forest and NPDR with correction for sex. Most genes highly associated with sex are removed by adjustment in NPDR (above horizontal dashed line). Many genes with high random forest importance score also have high association with sex (dark genes). Genes are labeled if they are strongly associated with sex (Bonferroni P value < 0.001).

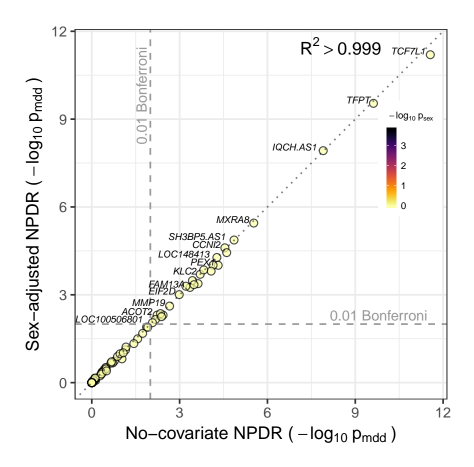


Figure S5: NPDR with and without sex adjustment for analysis of MDD-associated genes in Le et al.'s RNASeq dataset [1]. Adjustment for the sex covariate has a negligible effect on the resulting P values for each important gene because of the balanced study design. Both methods yield consistent results with STIR from previous study (Fig. 4 of Ref. [2]), not shown.

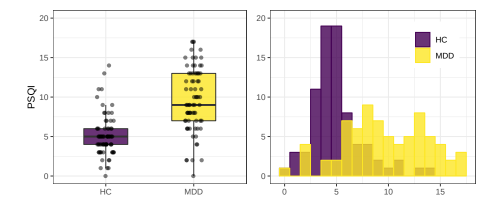


Figure S6: The distribution of the Pittsburgh Sleep Quality Index (PSQI) among individuals with and without MDD in Le et al.'s RNASeq dataset [1].