

biotmle: Targeted Learning for Biomarker Discovery

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Software

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Summary

The `biotmle` package provides an implementation of a biomarker discovery methodology based on targeted minimum loss-Based estimation (TMLE) (van der Laan and Rose 2011) and a generalization of the moderated t-statistic of (Smyth 2004), designed for use with biological sequencing data (e.g., microarrays, RNA-seq). The statistical approach made available in this package relies on the use of TMLE to rigorously evaluate the association between a set of potential biomarkers and another variable of interest while adjusting for potential confounding from another set of user-specified covariates. The implementation is in the form of a package for the R language for statistical computing (R Core Team 2017).

There are two principal ways in which the biomarker discovery techniques in the `biotmle` R package can be used: to evaluate the association between (1) a phenotypic measure (say, environmental exposure) and a biomarker of interest, and (2) an outcome of interest (e.g., survival status at a given time) and a biomarker measurement, both while controlling for background covariates (e.g., BMI, age). By using an estimation procedure based on TMLE, the package produces results based on the Average Treatment Effect (ATE), a statistical parameter with a well-studied causal interpretation (see van der Laan and Rose (2011) for extended discussions), making the `biotmle` R package well-suited for applications in bioinformatics, epidemiology, and genomics.

After adjusting our data set to be consistent with the expect input format – please consult the vignette accompanying the R package for details – we would call the principal function of this R package: `biomarkertmle`.

We would perform a moderated test on the output of the `biomarkertmle` function using the function `modtest_ic`.

While the principal table of results produced by this R package matches those produced by the well-known `limma` R package (Smyth 2005), there are also several plot methods made available for the `bioTMLE` S4 class – subclassed from the popular `SummarizedExperiment` class – introduced by this package (Huber et al. 2015). For illustrative purposes, we demonstrate the output of two such functions on anonymized experimental data below:

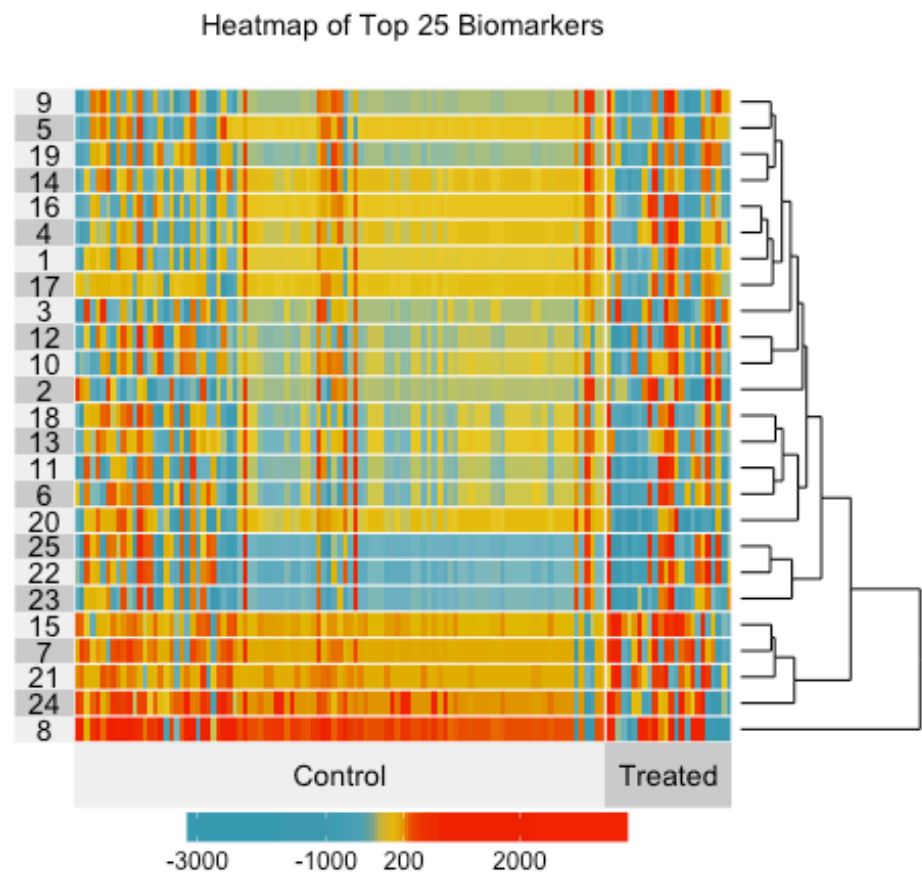


Figure 1: Heatmap visualizing the Average Treatment Effect contribution of a change in exposure to each biomarker of interest

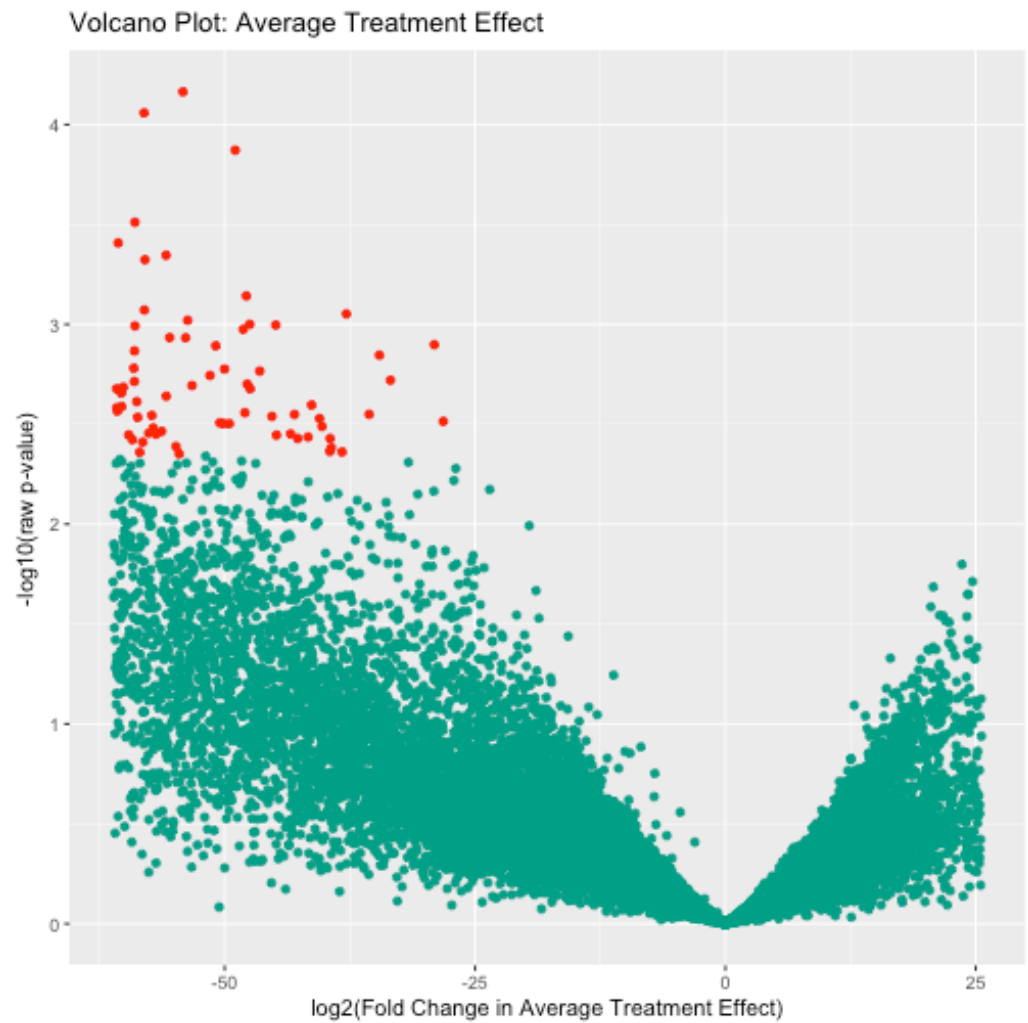


Figure 2: Volcano plot visualizing the log fold change in the Average Treatment Effect against the raw p-value from the moderated t-test performed on each biomarker

References

- Huber, Wolfgang, Vincent J Carey, Robert Gentleman, Simon Anders, Marc Carlson, Benilton S Carvalho, Hector Corrada Bravo, et al. 2015. “Orchestrating High-Throughput Genomic Analysis with Bioconductor.” *Nature Methods* 12 (2). Nature Research: 115–21.
- R Core Team. 2017. *R: A Language and Environment for Statistical Computing*. Vienna, Austria: R Foundation for Statistical Computing. <https://www.R-project.org/>.
- Smyth, Gordon K. 2004. “Linear Models and Empirical Bayes Methods for Assessing Differential Expression in Microarray Experiments.” *Statistical Applications in Genetics and Molecular Biology* 3 (1): 1–25.
- . 2005. “Limma: Linear Models for Microarray Data.” In *Bioinformatics and Computational Biology Solutions Using R and Bioconductor*, 397–420. Springer.
- van der Laan, Mark J, and Sherri Rose. 2011. *Targeted Learning: Causal Inference for Observational and Experimental Data*. Springer Science & Business Media.