Supplemental Information for Human methylome variation across Infinium 450K data on the Gene Expression Omnibus

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Overview

This document contains supplemental information for the manuscript *Human methylome variation across Infinium 450K data on the Gene Expression Omnibus*, including details about data access, aggregation, and analyses, with links to specific scripts where applicable.

Gene Expression Omnibus (GEO) data

DNA methylation (DNAm) array samples were identified as published to the Gene Expression Omnibus (GEO) and available in the GEO Data Sets database as of March 31, 2019.

GEO queries and data summaries

The Entrez Utilities software v10.9 was used to quantify DNAm array sample availability by year and platform for 3 major Illumina BeadArray platforms ($\rm HM450K$, $\rm HM27K$, and $\rm EPIC/HM850K$).

Samples and studies were identified for download using the script $https://github.com/metamaden/recountmethylation_server/blob/master/src/edirect_query.py.$

 $\label{lem:decomposition} Data\ availability\ by\ year\ was\ determined\ using\ the\ script\ https://github.com/metamaden/recountmethylatio\ nManuscriptSupplement/blob/main/inst/python/eqplot.py.$

Plots for figures 1A and S1 were generated using the scripts https://github.com/metamaden/recountmethyl ationManuscriptSupplement/blob/main/inst/scripts/figures/fig1a.R and https://github.com/metamaden/recountmethylationManuscriptSupplement/blob/main/inst/scripts/figures/figS1.R.

Data acquisition

We used the Python v3.7.0 programming language to develop a download management system to handle and version file downloads from GEO. We used Celery v5.0.0 to handle job management and logging. We used this job management system to obtain GSM IDATs, GSE SOFT files, and other data using batch queries to the GEO Data Sets repository. IDATs and SOFT files were acquired using the script https://github.com/m etamaden/recountmethylation_server/blob/master/src/dl.py and download jobs were managed using the script https://github.com/metamaden/recountmethylation_server/blob/master/src/server.py

Processing DNAm data from IDATs

Signals were read from sample IDATs into an R session using the minfi v1.29.3 R/Bioconductor package (Aryee et al. 2014). DNAm assay data were read from IDATs using the script functions from the script https://github.com/metamaden/recountmethylation.pipeline/blob/master/R/rmpipeline.R.

Sample metadata

Scripts for metadata processing can be found in the directory https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/metadata.

Extracting sample/GSM metadata from GSE SOFT files in JSON format

After obtaining the GSE SOFT files from GEO, we extracted GSM-specific metadata as JSON-formatted files using the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/metadata/soft2json.R. Before further processing, we filtered the sample JSON-formatted metadata using the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/metadata/jsonfilt.R. This step attempted to remove any GSE-specific metadata while retaining sample-identifying characteristics.

Processing the sample metadata

The filtered JSON metadata were read into a list of tables, organized by GSE ID, using the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/met adata/make_gse_annolist.R. We then preprocessed these tables by moving the available metadata under common variables as shown in the script https://github.com/metamaden/recountmethylationManuscriptSup plement/tree/main/inst/scripts/metadata/metadata_preprocessing.R. This step was coordinated manually and separately for each respective GSE ID. We additionally obtained GSM record titles from JSON files using the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/metadata/get_gsm_titles.R. We postprocessed available metadata by mapping available data to controlled term vocabularies using the script https://github.com/metamaden/recountmethylationManuscrip

tSupplement/tree/main/inst/scripts/metadata/metadata_preprocessing.R. Controlled vocabularies were heavily inspired by terms used by the Marmal-aid resource, and we attempted to map the most frequently available and informative metadata we observed in the original SOFT files.

Predicting sample types from filtered sample JSON files

We ran the MetaSRA-pipeline software on the filtered versions of sample JSON metadata files (Bernstein et al. 2017). For this, we used a forked the original software available at https://github.com/metamaden/MetaSRA-pipeline. For each sample record, the MetaSRA-pipeline mapped a series of curated ontology terms and predicted the likelihoods for each of 6 sample type categories (Figure S2). We extracted the pipeline-mapped files into tables using the script https://github.com/metamaden/recountmethylationManuscriptSupelment/tree/main/inst/scripts/metadata/get_msrap_mdmap.R. We retained the most likely predicted sample types and their likelihoods as shown in the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/metadata/make_md_final.R.

DNAm model-based predictions for age, sex, and blood cell types

Model-based predictions of sex, age, and cell type fractions were obtained from DNAm data as described in https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/metad ata/metadata_model_predictions.R. Age predictions were obtained by running the agep function from the wateRmelon v1.28.0 package on noob-normalized DNAm Beta-values (Horvath 2013). Sex predictions were obtained by running the getSex function from minfi on genome-mapped MethylSet data (Aryee et al. 2014). Cell fraction predictions were obtained by running the estimateCellCounts on RGChannelSet-formatted data (Houseman et al. 2012).

Quality and summary metrics

We calculated quality signals for 17 BeadArray controls, methylated and unmethylated signals, and likely sample replicates by study, as described in the table script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/tables/tables2.R. BeadArray signals for 17 controls were calcuated using the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/R/beadarray_cgctrlmetrics.R. Our method was developed in consultation with Illumina's platform documentation and functions from the ewastools R package ("Illumina Genome Studio Methylation Module V1.8" 2010; "BeadArray Controls Reporter Software Guide" 2015; Heiss and Just 2018). Sample genetic identities were predicted using the functions call_genotypes() and check_snp_agreement() from the ewastools R package, which uses high-frequency SNP signals from HM450K arrays (Heiss and Just 2018). We predicted and recorded likely genotype-based replicates within each GSE record.

Statistical analyses

Tests, summaries, and plots

Sample data were obtained, read, and analyzed programmatically using the R v4.0.0 and Python v3.7.0 languages in a CentOS 7 remote server environment. IDAT signals were read into SummarizedExperiment objects using the minfi package. Summary statistics were generated using base R functions. Scripts to reproduce manuscript analyses are stored at https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/scripts/analyses. Statistical tests were performed using the stats v4.0.0 R package. Correlation tests used the Spearman method by setting method = "spearman" in the cor.test function. Analyses of variance (ANOVAs) were performed using the anova function. Label enrichments were

tested using Binomial with the binom.test function, and T-tests used the t.test function. Unless noted otherwise, p-value adjustments used the Benjamini-Hotchberg method by setting method = "BH" in the p.adjust function. Plots used base R functions and the R packages ggplot2 v3.1.0 and ComplexHeatmap v1.99.5. Scripts to reproduce manuscript figures are stored at https://github.com/metamaden/recountmeth ylationManuscriptSupplement/tree/main/inst/scripts/figures.

PCAs of autosomal DNAm

Approximate array-wide PCA was performed using noob-normalized Beta-values from autosomal probes and the prcomp from the stats R package. We condensed autosomal DNAm across 35,360 samples into 1,000 hashed feature columns, the performed cluster analysis. As a dimensionality reduction step, feature hashing or the "hashing trick" mapped CpG probe data into a smaller intermediate dimension while approximately preserving between-sample variation (Weinberger et al. 2010). Feature hashing was performed as shown in the script https://github.com/metamaden/recountmethylationManuscriptSupplement/tree/main/inst/pytho n/dnamhash.py. The hashed features data is available at https://recount.bio/data/recountmethylation_man uscript_supplement/data/pca_fh1k_all_gsm35k.zip.

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