

Cheat sheet for recountmethylation

TERMINOLOGY		Download DNAm compilations	
DNA methylation (DNAm) terms		<p>The following command line options show how to download database files from the server (http://www.recount.bio/data) (http://www.recount.bio/data)), or you may try to <code>right-click -> download</code> from within your browser. Note, it may help to increase your timeout period for long downloads.</p> <ul style="list-style-type: none">Use <code>recountmethylation</code> : e.g. <code>getdb_h5se_rg("hm450k")</code> to download the HM450k <code>RGChannelSet</code> data as an <code>HDF5-SummarizedExperiment</code> object (see <code>?getdb</code> for more info).Use <code>wget</code> : From command line, enter <code>wget -r <filepath></code>, replacing <code><filepath></code> with an address from https://www.recount.bio/data . Note the <code>-r</code> is needed for <code>h5se</code> objects, which are directories.	
CpG locus	: Cytosine-guanine dinucleotide pair		
DNA methylation (DNAm)	: DNA with covalently bound methyl (CH ₃) groups; here specify cytosine-bound methyl groups in a CpG locus.		
CpG Island	: DNA regions enriched for cytosine, guanine, and CpG loci.		
BeadArray terms			
CpG probe	: Class of DNAm array technology using bead-bounded probes to quantify DNAm at specific CpG loci.	DelayedArray operations and pipes	
BeadArray	: Type of microarray using BeadArray probes, manufactured by Illumina, to quantify DNAm.		
HM27k	: Type of DNAm array, introduced around 2005, targeted roughly 27,000 CpG loci in humans.		<p>The following operations make use of <code>DelayedArray</code> backend for efficient memory usage, delayed large operations, and automatic large file management.</p> <ul style="list-style-type: none">Perform summary operations with <code>dim()</code> / <code>nrow()</code> / <code>ncol()</code>Rapidly update the sample metadata in an <code>h5se</code> object with <code>quickResaveHDF5SummarizedExperiment()</code> .Automatically process data in blocks using functions like <code>colSums()</code> , <code>colMeans()</code> , <code>colVars</code> , <code>rowSums</code> , <code>rowMeans</code> , and <code>rowVars()</code> , etc.
HM450k	: Type of DNAm array, introduced around 2011, targeting roughly 450,000 CpG loci in humans. Probes primarily cover gene and CpG island regions.		
EPIC	: Type of DNAm array, introduced around 2015, targeting roughly 855,000 CpG loci in humans. Shares 93% (about 453,000) of probes with the HM450k platform. Probes primarily cover genes, CpG islands, and intergenic regulatory features.		
Data object classes		RECAST DATASET OBJECTS	
HDF5 / h5	: Hierarchical database format 5, a type of database syntax implementing compression and chunking.	<p>If operations on datasets throw errors due to their class, you may attempt to recast and rerun them without the <code>DelayedArray</code> backend.</p> <div>Recast an <code>RGChannelSet</code></div> <pre>library(minfi) rg <- loadHDF5SummarizedExperiment(rg.path) green.matrix <- as.matrix(getGreen(rg)) red.matrix <- as.matrix(getRed(rg)) anno <- annotation(rg) metadata <- DataFrame(pData(rg)) rg.new <- RGChannelSet(Green = green.matrix, Red = red.matrix, annotation = anno, colData = metadata)</pre> <div>Recast a <code>GenomicRatioSet</code></div> <pre>library(minfi) gr <- loadHDF5SummarizedExperiment(gr.path) bval.matrix <- as.matrix(getBeta(gr)) anno <- annotation(gr) metadata <- DataFrame(pData(gr)) ranges <- granges(gr) gr.new <- GenomicRatioSet(gr = ranges, Beta = bval.matrix, annotation = anno, colData = metadata)</pre>	
se	: Short for <code>SummarizedExperiment</code> , a multifaceted object class in R/Bioconductor containing slots for assay measurements and metadata for the platform, samples, and experiment.		
h5se	: Short for <code>HDF5-SummarizedExperiment</code> , a hybrid object class that uses <code>DelayedArray</code> for caching.		
RGChannelSet / rg	: Type of <code>se</code> object containing dual color channel data for red and green channels on Illumina's BeadArray DNAm array platforms.		
GenomicMethylSet / gm	: Type of <code>se</code> object containing the methylated (a.k.a. 'M') and unmethylated (a.k.a. 'U') signals calculated from red and green dual channel intensity data.		
GenomicRadioSet / gr	: Type of <code>se</code> object containing the DNAm fractions (a.k.a. Beta-values) and/or logit2-transformed fractions (a.k.a. M-values) calculated from the M and U signals.		

DECISION TREE

Once you've selected a DNAm array platform (e.g. HM450k or EPIC), you can use the following decision tree to determine which database compilation object type to download.

