## Cheat sheet for recountmethylation

TERMINOLOGY			Download DNAm compilations	
CpG locus	: DNA with covalently bound methyl (CH <sub>3</sub> ) groups; ation here specify cytosine-bound methyl groups in a		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 right-click -> download from within your browser.  Note, it may help to increase your timeout period for long downloads.  • Use recountmethylation: e.g. getdb_h5se_rg("hm450k") to download the HM450k RGChannelSet data as an HDF5-SummarizedExperiment object (see ?getdb for more info).  • Use wget: From command line, enter wget -r <filepath>, replacing <filepath> with an address from</filepath></filepath>	
DNA methylation (DNAm)				
CpG Island	oG Island : DNA regions enriched for cytosine, guanine, and CpG loci.			
	E	BeadArray terms	https://www.recount.bio/data.Note the -r is needed for h5se	
CpG : Class of DNAm array technology using bead- brobe bounded probes to quantify DNAm at specific CpG loci.			objects, which are directories.	
BeadArray: Type of microarray using BeadArray probes, manufactured by Illumina, to quantify DNAm.			DelayedArray operations and pipes	
HM27k : Type of DNAm array, introduced around 2005, targeted roughly 27,000 CpG loci in humans.		-	The following operations make use of <code>DelayedArray</code> backend for efficient memory usage, delayed large operations, and automatic large file management.	
HM450k :	Type of DNAm array, introduced around 2011, targeting roughly 450,000 CpG loci in humans. Probes primarily cover gene and CpG island regions.		<ul> <li>Perform summary operations with dim() / nrow() / ncol()</li> </ul>	
			<ul> <li>Rapidly update the sample metadata in an h5se object with quickResaveHDF5SummarizedExperiment().</li> </ul>	
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.		<ul> <li>Automatically process data in blocks using functions like colsums(), colMeans(), colVars, rowSums, rowMeans, and rowVars(), etc.</li> </ul>	
			RECAST DATASET OBJECTS	
HDF5 / h5	Da	eta object classes  : Hierarchial database format 5, a type of database syntax implementing compression and chunking.	If operations on datasets throw errors due to their class, you may attempt to recast and rerun them without the DelayedArray backend.  Recast an RGChannelSet	
se		: Short for SummarizedExperiment, a multifaceted object class in R/Bioconductor containing slots for assay measurements and metadata for the platform, samples, and experiment.	<pre>library(minfi) rg &lt;- loadHDF5SummarizedExperiment(rg.path) green.matrix &lt;- as.matrix(getGreen(rg)) red.matrix &lt;- as.matrix(getRed(rg)) anno &lt;- annotation(rg) metadata &lt;- DataFrame(pData(rg)) rg.new &lt;- RGChannelSet(Green = green.matrix,</pre>	
h5se		: Short for  HDF5-SummarizedExperiment, a hybrid object class that uses	Red = red.matrix, annotation = anno, colData = metadata)	
		DelayedArray for caching.	Percet o granulare l'agus	
RGChannelSet/rg :		: Type of se object containing dual color channel data for red and green channels on Illumina's BeadArray DNAm array platforms.	<pre>Recast a GenomicRatioSet  library(minfi) gr &lt;- loadHDF5SummarizedExperiment(gr.path) bval.matrix &lt;- as.matrix(getBeta(gr)) anno &lt;- annotation(gr)</pre>	
GenomicMethylSet/gm :		: Type of se object containing the methylated (a.k.a. 'M') and unmethylated (a.k.a. 'U') signals calculated from red and green dual channel intensity data.	metadata <- DataFrame(pData(gr)) ranges <- granges(gr) gr.new <- GenomicRatioSet(gr = ranges, Beta = bval.matrix, annotation = anno, colData = metadata)	
GenomicRadioSet/gr :		: Type of se 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		E	recount methylation	
Once you've selected a DNAm array olatform (e.g. HM450k or EPIC), you			Data format	

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.

