

HumanMethylation27k probes in gene bodies

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27k probes by design and color

First we have to grab the probes that made it onto the 450k chip.

```
> library('IlluminaHumanMethylation450k.db')
> probes.27k <- IlluminaHumanMethylation450k_get27k()
> lapply(probes.27k, function(x) {
+   if( class(x) == 'list') lapply(x, head)
+   else head(x)
+ })
```

\$I

\$I\$R

```
      Probe_ID
1 cg04037732
2 cg04765675
3 cg04927982
4 cg05113908
5 cg08798116
6 cg09229960
```

\$I\$G

```
      Probe_ID
1 cg02004156
2 cg04344997
3 cg04702045
4 cg05935584
5 cg06700462
6 cg06791102
```

\$II

```
      Probe_ID
1 cg03515901
2 cg08455548
3 cg20401549
4 cg00029931
5 cg00032666
6 cg00060882
```

Annotation

How many probes align to UCSC gene bodies? That's a bit complicated, because each Illumina probe can be mapped to several transcripts, and each transcript to several probes. Normalizing the schema reduced the database size by 100MB.

```
> probes.27k <- unlist(probes.27k, recursive=T)
> head(mget(probes.27k, IlluminaHumanMethylation450kPROBELOCATION, ifnotfound=NA))

$cg04037732
[1] "NM_001166660:1stExon" "NM_181303:1stExon" "NM_018977:5'UTR"
[4] "NM_018977:1stExon" "NM_181303:5'UTR" "NM_001166660:5'UTR"

$cg04765675
[1] "NM_018360:TSS1500" "NM_001168683:TSS1500"

$cg04927982
[1] "NM_001146702:TSS200" "NM_004187:TSS200"

$cg05113908
[1] "NM_001079855:5'UTR" "NM_003918:5'UTR"

$cg08798116
[1] "NM_001448:1stExon"

$cg09229960
[1] "NM_000117:1stExon"
```

Location mapping is many-to-one and emerges as a list of concatenations, so we cannot simply set up a simpleBimap object... UNLESS the concatenation is part of a VIEW (across four or so different tables). So a VIEW is what we use.

```
> probeloc <- mget(probes.27k, IlluminaHumanMethylation450kPROBELOCATION,
+               ifnotfound=NA)
> body.or.exon <- function(x) length( grep('(Body|Exon)', x) ) > 0
> length(which(unlist(lapply(probeloc, body.or.exon))))

[1] 12173

> in.body <- function(x) length( grep('Body', x) ) > 0
> gene.body.probes <- names(which(unlist(lapply(probeloc, in.body))))
> length(which(unlist(lapply(probeloc, in.body))))

[1] 5700

> head(gene.body.probes)

[1] "cg16510010" "cg19963797" "cg20085077" "cg01860753" "cg03085637"
[6] "cg05795157"
```

They are not nearly as scarce as I initially thought.

Versioning

It's important to keep track of where information came from, and who is in charge of keeping it organized, when documenting phenomena like this.

```
> IlluminaHumanMethylation450k_dbInfo()[c(8:10,22:24,31,33),]
      name                                     value
8  EGSOURCEDATE                               2010-Sep7
9  EGSOURCENAME                               Entrez Gene
10 EGSOURCEURL                                ftp://ftp.ncbi.nlm.nih.gov/gene/DATA
22 GPSOURCENAME                               UCSC Genome Bioinformatics (Homo sapiens)
23 GPSOURCEURL                                ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19
24 GPSOURCEDATE                               2010-Mar22
31  MANIFEST                                  HumanMethylation450_15017482_v.1.1.csv
33 MANIFESTDATE                               2010-Dec7

> IlluminaHumanMethylation450kSVNID

[1] "$Id: zzz.R 1291 2011-05-04 23:49:22Z ttriche $"

> IlluminaHumanMethylation450kBLAME

[1] "$Author: ttriche $"
```

Writing to Bioconductor standards simply enforces this.

R session

```
> toLatex(sessionInfo())
```

- R version 2.13.0 Patched (2011-04-25 r55641),
x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8,
LC_COLLATE=C, LC_MONETARY=C, LC_MESSAGES=en_US.UTF-8,
LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C,
LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: AnnotationDbi 1.14.1, Biobase 2.12.1, DBI 0.2-5,
IlluminaHumanMethylation450k.db 1.4.6, RSQLite 0.9-4,
org.Hs.eg.db 2.5.0
- Loaded via a namespace (and not attached): tools 2.13.0