A mammalian methylation array for profiling methylation levels at conserved sequences

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SUMMARY

Infinium methylation arrays are widely used to robustly measure methylation of DNA in humans.

However, such arrays are not available for the vast majority of non-human mammals. Moreover,

even if species-specific arrays were available, probe differences between them would confound

cross-species comparisons. To address these challenges, we developed the Mammalian

Methylation Array, a single custom Infinium array that measures cytosine methylation levels of

over 35 thousand CpG sites that are well conserved across species within the mammalian class.

By design, the probes on the array tolerate cross-species mutations. To design the array, we

developed the Conserved Methylation Array Probe Selector (CMAPS) algorithm, which takes as

input a multi-species sequence alignment and probe design constraints. A greedy search

algorithm was used to identify oligonucleotide sequences (probes) with high coverage across

different mammalian species. We annotate the probes on the array with respect to genes in 159

different species and provide details on the sequence context including CpG island status and

chromatin states. Our calibration experiments demonstrate the high fidelity of this array in

humans, rats, and mice. The mammalian methylation array has several strengths: it applies to all

mammalian species even those that have not yet been sequenced, it provides deep coverage of

specific cytosines facilitating the development of highly robust epigenetic biomarkers, and it

covers highly conserved CpGs which greatly increases the probability that biological insights

gained in one species will readily translate to others. The mammalian methylation array is expected to find many applications in preclinical studies, comparative biology, and epigenetic studies of aging and development.

Introduction

Methylation of DNA by the attachment of a methyl group to cytosines is one of the most widely studied epigenetic modifications in vertebrates, due to its implications in regulating gene expression across many biological processes including disease (Ooi et al., 2007; Robertson, 2005; Smith and Meissner, 2013). A variety of different assays have been proposed for measuring DNA methylation including microarray based methylation arrays (Bibikova et al., 2009, 2011) and sequencing based assays such as whole genome bisulfite sequencing (WGBS)(Cokus et al., 2008; Lister et al., 2009) and reduced representation bisulfite sequencing (RRBS)(Meissner et al., 2005). Despite the availability of sequencing based assays, array based technology remains widely used for measuring DNA methylation due to its low-cost and high reproducibility and reliability(Pidsley et al., 2016).

The first human methylation array (Illumina Infinium 27K) was introduced by Illumina Inc in 2009 (Bibikova et al., 2009), which were followed by the 450K(Bibikova et al., 2011) and EPIC arrays with larger coverage(Pidsley et al., 2016). More recently, Illumina released a mouse methylation array (Infinium Mouse Methylation BeadChip) that profiles over 285k markers across diverse murine strains. It will probably not be economical to develop similar methylation arrays for less frequently studied mammalian species (e.g. elephants or marine mammals) due to insufficient demand. Moreover, even if costs were no impediment, species-specific arrays would likely be sub-optimal in comparative studies across different species as the measurement platforms would be different.

3

To address these challenges, we developed a single mammalian methylation array designed to be used to measure DNA methylation across mammals. The array targets CpGs for which the CpG and flanking sequence are highly conserved across many mammals so that the methylation of many of these CpGs can be measured in each mammal. The design repurposes the degenerate base technology (originally used by Illumina Infinium probes to tolerate withinhuman variation) to tolerate cross-species mutations across mammalian species. To select the specific probe sequences including tolerated mutations that appear on the array we developed the Conserved Methylation Array Probe Selector (CMAPS). CMAPS takes as input a multiple sequence alignment to a reference genome and a set of probe design constraints, and selects a set of probe sequences including tolerated mutations, which can be used to guery methylation in many species. We apply CMAPS to select over 35 thousand CpGs for the mammalian methylation array, which we complemented with close to two thousand known human biomarker CpGs. We characterize the CpGs on the mammalian methylation array with various genomic annotations. Further, we use calibration data to evaluate the fidelity of individual probes in humans, mice, and rats. CMAPS has led to the design of the mammalian methylation array, which will facilitate the study of cytosine methylation at conserved loci across all mammal species.

Results

Designing the Mammalian Methylation Array

The CMAPS algorithm is designed to select a set of Illumina Infinium array probes such that for a target set of species many probes are expected to work in each species (**Methods**). Array probes are sequences of length 50bp flanking a target CpG based on the human reference genome. Selecting sequences present in the human reference genome increases the likelihood that measurements in other species will transfer to human. The mammalian methylation array adapts the degenerate base technology for tolerating human SNPs so that probes can tolerate a

limited number of cross-species mutations. The CMAPS algorithm is provided as input a multiple-species sequence alignment to a reference genome. CMAP uses these inputs to then select the CpGs to target on the array. As part of selecting the CpGs, CMAP also selects the probe sequence design to target them including the specific set of degenerate bases. For designing the mammal methylation array, CMAPS was applied to the subset of 62 mammals within a 100-way alignment of 99 vertebrate genomes with human genome (Haeussler et al., 2019), but we note the CMAPS method is general.

In designing a probe for a CpG, CMAPS considers multiple different options. One option is the type of probe. Illumina's current methylation array technology allows up to two types of probes: Infinium I and Infinium II. The latter is newer technology requiring only one silica bead to query the methylation of a CpG, while the former requires two beads. By only requiring one bead Infinium II probes allow under fixed array capacity limits interrogating more CpGs, though Infinium I probes are better able to query CpGs in CpG rich regions (Bibikova et al., 2011). Another option for each of these two types of probes is whether the probe is on the forward or reverse genomic strand, giving four total combinations of options for probe type and strand for each CpG. In addition, CMAPS has options for the position and nucleotides identity of tolerated mutation across correspond to degenerate bases. The array degenerate base technology allows for potentially up to three degenerate bases per probe sequence, which are positions that can be designed to tolerate variation in the sequence being interrogated. For some probes fewer than three degenerate bases could be designed, which was determined based on a design score computed by Illumina for each probe and in the case of Infinium II probes also the number of CpGs within the probe sequence. CMAPS uses a greedy algorithm to select the tolerated mutations for each combination of probe type and strand. The algorithm aims to maximize the number of species in the alignment the probe is expected to work based on just local alignment information that is without considering how uniquely mappable the probe is across the genome. A probe for a CpG

is expected to work in a non-human species based on local alignment information if there are no differences in the alignment between the human genome sequence and the other species excluding those accounted for by the probe's degenerate bases (**Figure 1a, Methods**). For each CpG site in the human genome, CMAPS retained for further consideration the Infinium I probe out of the two options (forward or reverse of the CpG) which had the greater number of species for which the probe was expected to work, and likewise for Infinium II.

We next applied a series of rules to identify a reduced subset of candidate probes. First, we included all 36,133 Infinium II probes that were expected to work in mouse (based on the mm10 genome), which maximizes the expected array utility for one of the most widely used model organisms. For the remaining set of CpG not selected in the previous step, we sorted them in descending order of the number of species for which an Infinium II probe was expected to work. We then added the top 16,867 CpG sites for a total of 53,000 CpG sites. Next, we ranked the CpGs targeted on the Illumina EPIC array (Pidsley et al., 2016) in descending order of the number of species for which a probe targeting the CpG is expected to work. For this the probe was required to be of the same probe type and strand as on the EPIC array, but used the degenerate bases picked by the CMAPS algorithm. The probe was allowed to differ in terms of degenerate base positions, as EPIC probes typically do not account for degenerate bases across species. For this we selected the top 3,000 CpG sites ranked sites that had not already been picked based on the earlier criteria.

Lastly, we sorted the CpG sites in descending order of number of species they can target and picked the top 4,000 CpGs targeted by Infinium I probes that had not already been included. The Infinium I probes were selected to allow querying CpG dense regions such as CpG islands, as CpGs do not count towards the limited number of positions of variation as for Infinium II probes. This resulted in a set targeting 60,000 CpGs (**Figure 1b**).

For some of these 60,000 CpGs, the sequence of the probe targeting it can map to multiple locations in a genome, which could result in a confounded signal coming from multiple CpG sites. This issue is compounded by individual probes corresponding to multiple sequences reflecting different possible combinations of the degenerate bases. To identify a subset of probes less susceptible to such confounders, for 16 high quality genomes, we computed for each probe how many of its versions map uniquely in that genome (see Methods). We then filtered CpGs down by requiring all versions of a probe targeting it map uniquely in at least 80% of the species they are expected to target out of the 16 high quality genomes, unless the probe is expected to target at least 40 mammals from the alignment, in which case the mapping criterion was discarded. This reduced the set of candidate CpGs to 35,989 CpGs.

We added probes targeting 1986 CpGs to the mammalian methylation array based on their utility for human biomarker studies (Supplementary Data). These probes, which were previously implemented in human Illumina Infinium arrays (EPIC, 450K, 27K), were selected due to their utility for human biomarker studies estimating age, blood cell counts, or the proportion of neurons in brain tissue(Guintivano et al., 2013; Hannum et al., 2013; Horvath, 2013; Horvath and Levine, 2015; Horvath et al., 2018; Houseman et al., 2012; Levine et al., 2018). The final manufactured mammalian methylation array measures cytosine levels of 37,492 cytosines: 37,488 of these cytosines are followed by a guanine (CpGs) and 4 are followed by another nucleotide (non-CpGs). The probe identifiers (cg numbers) of 86 of these cytosines ends with either ".1" or ".2", i.e. these are duplicate probes for 43 genomic locations.

A detailed analysis of the Infinium probe context of the mammalian array and relation to human and mouse arrays is presented in **Supplementary Figure S1**. The mammalian methylation array focus on highly conserved regions led to a an array that is distinct from other currently available Infinium arrays that focus on specific species. For example, the mammalian array only shares

3107 probes with the Illumina MouseMethylation array and only 7111 CpGs with the Illumina EPIC array.

Mappability analysis

All 37488 CpGs profiled on the mammalian methylation array apply to humans, but only a subset of these CpGs applies to other species. When conducting analyses in a specific species it can thus be desirable to restrict analyses to the subset of CpG that apply in that species. One approach for doing this is simply omit CpGs whose detection p-values from normalization methods (Methods) are insignificant. This approach has the advantage of being applicable to species that have not yet been sequenced. Mapping sequences to genomes has the added benefit of providing a candidate position of the sequence in the target genome from which other information about the CpG can be inferred such as the nearest gene or CpG island status.

We have mapped the array CpGs to 159 species, which also provides a candidate position from which a gene for the CpG can be associated. As expected, the closer a species is to humans, the more CpGs map to the genome of this species. Over 30k CpGs on the array map to most placental mammalian genomes (eutherians, **Figure 2a, Supplementary Data**). Roughly 15K CpGs map to most non-placental mammalian genomes (marsupials), such as kangaroos or opossums. Far fewer CpGs map to egg laying mammalian genomes (monotremes), such as platypus (**Figure 2**). A CpG that is adjacent to a given gene in humans may not map to a position adjacent the corresponding (orthologous) gene in another species. Between 15k to 22k CpGs (over 70%) were assigned to human orthologous species based on their mapped position in most phylogenetic orders (rodents, bats, carnivores, **Figure 2b,c** and Supplementary Data). These numbers surrounding orthologous genes are probably overly conservative (i.e. lower than the true numbers) because we found the majority of CpGs (about 58%) that do not map to orthologous

genes in the non-human species are located in intergenic regions outside of promoters (Methods), which suggests that one of the gene assignments was inaccurate.

Chromosome and gene region coverage of array

We analyzed the chromosome and gene region coverage of the mammalian methylation array for human and mouse. The mammalian methylation has substantial coverage of all chromosomes (human, 235-3938; and mouse, 687-3179 probes per chromosome), with the exception of chrY that only has 2 probes in both species (**Supplementary Figure S2a**). When we assign the probes to the closest gene neighbor, around 80% of the probes are proximal to a gene in both of these species (**Supplementary Figure S2b**). The remaining 20% of probes are neither aligned to a promoter nor a gene body. The distribution of gene region and the distances to transcriptional start sites are comparable between human and mouse (**Supplementary Figure S2b**). CpGs on the mammalian array cover 6871 human and 5659 mouse genes when each CpGs is assigned uniquely to its closest gene neighbor (**Supplementary Figure S2c**). The gene coverage is uneven: while on average a gene is covered by 2 CpGs some genes are covered by as many as 150 CpGs. In mouse, 73% of CpGs (21,664) were assigned to a human orthologous genes (**Supplementary Figure S2d**), suggesting many CpG measurements from the array in mice will be informative to humans (and vice versa).

Gene sets represented in mammalian array

We analyzed gene set enrichments of all genes that are represented on the mammalian array using GREAT(McLean et al., 2010). Significant gene sets covered implicated gene sets that were found to play a role in development, growth, transcriptional regulation, metabolism, cancer, mortality, aging, and survival (**Supplementary Figure S3**). We also used the *TissueEnrich*(Jain and Tuteja, 2019) software to analyze gene expression (Methods). The majority of mammalian

methylation array probes (~65%) are adjacent to genes that are expressed in all considered human and mouse tissue (**Supplementary Figure S4a,b**). However, the mammalian array also contains CpGs that are adjacent to genes that are expressed in a tissue-specific manner, notably testis and cerebral cortex (**Supplementary Figure S4c**).

CpG island and methylation status

We analyzed the CpG island and DNA methylation properties of CpGs on the mammalian array. In general, an average of 5563 (18%) of probes in the mammalian array are located in CpG island depending on the species (**Figure 3a**). We used a CpG island detection algorithm (gCluster software (Li et al., 2020)) that additionally provided several species-level quantitative measures for each CpG island including the length, GC content, and CpG density that we provide as a resource (**Supplementary Data**). We also analyzed the DNA methylation levels in human for fractional methylation called from whole genome bisulfite sequencing data across 37 human tissues(Roadmap Epigenomics Consortium et al., 2015)⁵ (**Supplementary Figure 5**). This confirmed that the mammalian methylation array target CpGs across a wide range of fractional methylation levels.

Chromatin state annotation of array probes

We analyzed the overlap of human CpG's targeted on the mammal methylation array with chromatin states for 127 cell and tissues. The CpGs cover all available chromatin states including different types of promoters (including bivalent promoters), regions repressed by polycomb group proteins, transcription start and end site, and enhancer regions (**Figure 3b**). Among enhancers, CpG's had greater overlap with brain and neurosphere than other tissue groups. In addition to analyzing the array CpG's overlap for cell and tissue specific chromatin states, we also analyzed them for a universal chromatin state annotation, which provides a single annotation to the genome

per position based on data from more than 100 cell and tissue types (Vu and Ernst, 2020) (**Supplementary Figure S6**). This revealed the greatest enrichment for bivalent promoter states and also strong enrichment for other promoter states and a state associated with polycomb repression.

While the mammalian methylation array was specifically designed to profile CpGs in highly conserved stretches of DNA based on sequence conservation, we assessed whether there was also evidence of conservation at the functional genomics level using human-mouse LECIF scores (Kwon and Ernst, 2020). The human-mouse LECIF quantifies evidence of conservation between human and mouse at the functional genomics level using chromatin state and other functional genomic annotations. In general, probes on the array had higher LECIF score than regions that align between human and mouse in general (**Figure 3c**).

Mammalian array study of calibration data

To validate the accuracy of the mammalian methylation array we applied it to synthetic DNA methylation samples for three species: human (n=10 arrays), mouse (n=20), and rat (n=15), where the methylation levels were known. The DNA samples from human, mouse and rat were engineered such that the fractional methylation at all CpG sites in their genomes approximately 0%, 25%, 50%, 75% and 100% (**Methods**). The calibration data thus allow us to define a benchmark annotation measure "ProportionMethylated" (with ordinal values 0, 0.25, 0.5, 0.75, 1). The distribution of the intensity of the probes in each human sample is roughly centered around the benchmark measure (ProportionMethylated) (**Figure 4a**). However, as expected, the distributions in the mouse and rat samples of all the probes show somewhat different patterns in these two species compared to the human samples likely because many probes in the design of our array do not map to these genomes (**Figure 4b-c**). We also evaluate these for each species after removing the probes that were not designed to map to that species, and normalizing the

array data using the SeSaMe package, which defines beta (relative intensity) values for each probe (Zhou et al., 2018). After this procedure, we see sharper peaks close to 0 and 1, though the quantification of absolute methylation levels are somewhat degraded around the beta value 0.75 as we move away from humans (**Figure 4d-f**).

Additionally, for each species, DNA methylation levels of each CpG we computed the correlation with the benchmark variable "ProportionMethylated" across the arrays. High positive correlations would be evidence for the accuracy of the array, which is indeed what we observe. CpGs that map to the human, mouse, and rat genome have a median Pearson correlation of r=0.986 with an interquartile range of [0.96,0.99], r=0.959 with IQR=[0.92,0.98], and r=0.956 with IQR=[0.91,0.98] with the benchmark variable ProportionMethylated in the respective species. The numbers of CpGs on the mammalian array that pass a given correlation threshold (irrespective of the mappability to a given species) are reported in Table 1. We also compare the SeSaMe normalization with the "noob" normalization that is implemented in the *minfi* R package (Aryee et al., 2014; Triche et al., 2013) (**Table 1**). We find that SeSaMe slightly outperforms *minfi* when it comes to the number of CpGs that exceed a given correlation threshold with ProportionMethylated.

Comparison with the human EPIC methylation array study in calibration data

We compared the mammalian methylation to the human EPIC methylation array, which profiles 866k CpGs in the human genome, for non-human samples. Some of the EPIC array probes are expected to apply to the mouse and rat genomes as well (Needhamsen et al., 2017). To facilitate a comparison between the mammalian methylation array and the human EPIC array for non-human samples we applied the latter to calibration data from mouse (*n*=15 arrays) and rat (*n*=10). The same engineered DNA data methylation data were analyzed on the human EPIC array as on the mammalian methylation array above. In particular, we were able to correlate each

CpG on the EPIC array with a benchmark measure (ProportionMethylated) in mice and rats (Table 1). Only 2356 (out of 866k) CpGs on the human EPIC exceed a correlation of 0.90 with ProportionMethylated in mice. By contrast, 24050 CpGs on the mammalian array exceed the same correlation threshold in mice. Similarly, the mammalian array outperforms the EPIC array in rats: only 6159 CpGs on the EPIC array exceed a correlation of 0.90 with ProportionMethylated compared with 22427 CpGs on the mammalian array. The results are similar for the correlation thresholds of 0.85 and 0.95 (Table 1).

The EPIC array contains 5574 CpGs that were also prioritized by the CMAPS algorithm based on high levels of conservation, excluding the 1986 CpGs from human biomarker studies. Out of these 5574 shared CpGs, 4341 and 3948 CpGs map to the mouse and rat genome, respectively. While human EPIC probes target the same CpG, the corresponding mammalian probe is typically different from EPIC probe due to differences in probe type (type I versus type II probe), DNA strand, or the handling of mutations across species degenerate bass. In the following comparison, we limited the analysis to the 4341 and 3948 probes when analyzing calibration data from mice or rats, respectively. We find that the mammalian array probes are better calibrated than the corresponding EPIC array probes when applied to mouse and rat calibration data according to two different analysis that focus on shared CpGs between the two platforms. First, the mammalian array outperforms the EPIC array when considering mean methylation levels across the shared CpGs (Figure 5). Second, when correlating each of the shared CpGs with the benchmark value ProportionMethylated we observe median correlation of 0.72 for both mice and rat calibration data generated on the EPIC array. For the same probes we observe median correlations of 0.94 and 0.93 for mice and rat calibration data generated on the mammalian array (SeSaMe normalization), respectively. We are distributing the methylation data and results from our calibration data analysis in three species (Supplementary Data). These calibration results will allow users to select cytosines whose methylation have a high correlation with the benchmark data in human, mice or rat.

DISCUSSION

The mammalian methylation array, which was enabled by the CMAPS algorithm for selecting conserved probes, is applicable to all mammals and hence drives down the cost per chip through economies of scale. The mammalian methylation array has unique strengths: it applies to all mammalian species even those that have not yet been sequenced, it provides deep coverage of specific cytosines which is a prerequisite for developing robust epigenetic biomarkers, and its focus on highly conserved CpGs increases the chances that findings in one species will translate to those in another species. We expect that the mammalian methylation array is particularly well suited for DNA methylation based biomarker studies in mammals.

Our calibration data demonstrate that the array largely leads to high quality measurements in three species: human, mouse and rat. Our calibration data shows that the mammalian methylation array greatly outperforms the human EPIC chip when it comes to high fidelity measurement applications to mice and rats. The array thus should be preferable for most non-human applications unless high-fidelity measurements are not needed in which case the larger content of the EPIC array may make it preferable.

The mammalian methylation array has several limitations. First, only a fraction of genes in a given species are represented by CpGs. Second, it focuses on CpGs in highly conserved stretches of DNA and hence does not cover parts that are specific to a given species. Third, it provides worse coverage in more distal species, particularly in marsupials than in placental mammals (eutherians). Finally, the calibration data suggests there are some shifts in the *absolute* methylation levels detected for intermediate methylation levels, but the relative order is preserved. The correct *relative* ordering of beta values is of primary importance in most statistical tests and analyses.

Several software tools have been adapted for use with the mammalian methylation array that range from normalization to higher level gene enrichment analysis. Software tools for generating normalized data include SeSaMe and the minfi R package (Aryee et al., 2014; Zhou et al., 2018). The eFORGE software (Breeze et al., 2019), which has been adapted for the use with the mammalian array, facilitates chromatin state analysis and transcription factor binding site analysis. Many researchers will be interested in genome coordinates of the mammalian CpGs in different species. Toward this end, we provide genome coordinates in 159 species. This list of species will increase as more high quality genomes become available. Detailed gene annotations in many species are available including details on gene region (e.g. exon, promoter, 5 prime untranslated region) and CpG island status (Supplementary Data). For human and mice we provide chromatin state annotations (Ernst and Kellis, 2012; Gorkin et al., 2020; Roadmap Epigenomics Consortium et al., 2015; Vu and Ernst, 2020) and the LECIF score on evidence of conservation at the functional genomics level between human and mouse(Kwon and Ernst, 2020).

In other articles, we will describe the application of the mammalian methylation array to many different mammalian species. These upcoming studies will demonstrate that the mammalian methylation array is useful for many applications that involve mammalian species.

Methods

Conserved Methylation Array Probe Selector (CMAPS)

Given a multi-species sequence alignment and reference genome, for each CG site and each of the four different possible probe designs, CMAPS computes an estimate of the number of species from the alignment that could be targeted if the use of degenerate base technology is optimized for tolerated mutations. The four probe designs involve each combination of probe type (Infinium I vs. Infinium II), and whether the probe sequence is on the forward or reverse DNA strand. For

each probe option, CMAPS conducts a greedy search to select tolerated mutations, including position and allele that maximize species coverage for the probe. The maximum number of degenerate bases that can be included in a probe is a function of a design score provided by Illumina Inc. For Infinium II probes only, CpGs present in the probe sequence count as if they are a degenerate base. More specifically, the algorithm for determining the number of species and selecting the mutations to handle performs the following steps for each probe design:

- Let M be the maximum number of degenerate bases that can be designed into a specific probe, based on the design score
- 2. For each species s in the alignment, let M_s be the number of mismatches in the alignment between that species and the human reference sequence of the probe
 - a. If $M_s > M$ or the species does not have the target CpG, continue to next species
 - b. If $M_s \ll M$,
 - i. For each mismatch in species s, add each degenerate position to a multiset
 - ii. add the species to a set F of feasible species to target with this probe
- 3. For all |P| choose M combinations of possible degenerate positions:
 - a. For each unique position in the combination
 - i. For each possible alternate nucleotide count the number of species in F
 that contain that alternate nucleotide
 - ii. Pick the top k alternate nucleotides based on the count in i., where k is the number of occurrences of the current position in S
 - b. Compute the number of species that match the human reference when accounting for the degenerate substitutions handled in a
- 4. Select the combination of positions in S that maximizes 3.b

Our procedure for selecting the specific targeted CpG and probe designs are described in the

main text. We note that 30 of the CpGs selected for the mammalian methylation array based on

the conservation criteria (using the sequence alignment) overlap with the 1986 human biomarker

CpGs. The design of the probes targeting them could differ however. The probe names of different

probes targeting the same CpG are distinguished by extensions ".1" and ".2". For example

cq00350702.1 and cq00350702.2 target the same cytosine but use different probe chemistry. The

array contains four probes that measure cytosines that are not followed by a guanine selected by

human biomarkers, which are indicated with a "ch" instead of a "cg".

The CMAPS algorithm was applied with human hg19 as the reference genome and using

the Multiz alignment of 99 vertebrates with the hg19 human genome downloaded from the UCSC

Genome Browser (Haeussler et al., 2019; Rosenbloom et al., 2015). For the purpose of designing

the mammalian array, only the 62 mammalian species in this alignment were considered and 16

for the mappability analysis. However, the current version of the mappability analysis provides

genome coordinates for 159 species.

The mammalian methylation array includes an additional 62 human SNP markers (whose

probe names start with "rs" for human studies), which can be used to detect plate map errors

when dealing with multiple tissue samples collected from the same person. Finally, the

mammalian array also adopted a standard suite of probes from the Illumina EPIC array for

measuring bisulfite conversion efficiency in humans.

Mapping probes to genomic coordinates

We used two different approaches for mapping probes to genomes. The first approach (BSbolt

software) was primarily used in designing the array. Subsequently, we adopted a second

mappability approach (QUASR software) that allowed us to map more probes to more species.

17

Mappability Approach 1: BSbolt

For version 1 of our mappability analysis (i.e. for designing the array), we applied the BSbolt mapping approach to 16 high quality genomes from: Baboon (papHam1), Cat (felCat5), Chimp (panTro4), Cow (bosTau7), Dog(canFam3), Gibbon(nomLeu3), Green Monkey (chlSab1), Horse, (equCab2), Human (hg19), Macacque (macFas5), Marmoset(calJac3), Mouse (mm10), Rabbit (oryCun2), Rat (rn5), Rhesus Monkey (rheMac3), Sheep (oviAri3).

We utilized the BSBolt software (Farrell et al., 2020) package from https://github.com/NuttyLogic/BSBolt

to perform the alignments. For each species' genome sequence, BSBolt creates an 'in silico' bisulfite-treated version of the genome. As many of the currently available genomes are in a low quality assembly state (e.g. thousands of contigs or scaffolds), we used the utility "Threader" (which can be found in BSBolt's forebear BSseeker2(Guo et al., 2013) as a standalone executable) to reformat these fasta files into concatenated and padded pseudo-chromosomes. The set of nucleotide sequences of the designed probes, which includes degenerate base positions, was explicitly expanded into a larger set of nucleotide sequence representing every possible combination of those degenerate bases. For Infinium I probes, which have both a methylated and an unmethylated version of the probe sequence, only the methylated version was used as BSBolt's version of the genome treats all CG sites as methylated. The initial 37K probe sequences resulted in a set of 184,352 sequences to be aligned against the various species genomes. We then ran BSBolt with parameters Align -M 0 -DB [path to bisulfite-treated genome] -BT2 bowtie2 -BT2-p 4 -BT2-k 8 -BT2-L 20 -F1 [Probe Sequence File] -O [Alignment Output File] -S to align the enlarged set of probe sequences to each prepared genome.

As we were not interested in the final BSBolt style output, we made a small modification to the code to retain its temporary output of alignment results in "sam" format. From these files, we collected only alignments where the entire length of the probe perfectly matched to the genome

sequence (i.e. the CIGAR string '50M' and flag XM=0"). Then, for each genome we collapsed all

the sequence variant alignments for each probeID down to a list of loci for that genome and for

that probe.

Mappability Approach 2: QUASR

For version 2 of our mappability analysis, we aligned the probe sequences to all available

mammalian genomes in ENSEMBL and NCBI Refseq databases using the QUASR package

(Gaidatzis et al., 2015). The fasta sequence files for each genome were downloaded from these

public databases. The alignment assumed that the DNA has been subjected to a bisulfite

conversion treatment. For each species' genome sequence, QUASR creates an in-silico-bisulfite-

treated version of the genome. The probes were aligned to these bisulfite treated genome

sequences, which does not consider C-T as a mismatch. The alignment was ran with QUASR (a

wrapper for Bowtie2) with parameters -k 2 --strata --best -v 3 and bisulfite =

"undir" to align the enlarged set of probe sequences to each prepared genome. From these

files, we collected the best candidate unique alignment to the genome. Additionally, the estimated

CpG coordinates at the end of each probe was used to extract the sequence from each genome

fasta files and exclude any probes with mismatches in the target CpG location.

Genomic loci annotations

Gene annotations (gff3) for each genome considered were also downloaded from the

same sources as the genome. Following the alignment, the CpGs were annotated to genes based

on the distance to the closest transcriptional start site using the Chipseeker package(Yu et al.,

2015). Genomic location of each CpG was categorized as either intergenic region, 3' UTR, 5'

UTR, promoter (minus 10 kb to plus 100 bp from the nearest TSS), exon, or intron. The unique

19

region assignment is prioritized as follows: exons, promoters, introns, 5' UTR, 3' UTR, and intergenic.

Additional genomic annotations, including human ortholog ENSEMBL ID, were extracted from the BioMart ENSEMBL database(Yates et al., 2020). The candidate gene for each probe was compared with human orthologous ENSEMBL ID to examine the similarity of the alignment with the human. For each probe, we examined if the assigned species ENSEMBL ID is identical to human-to-other-species-orthologous ENSEMBL ID in human mappability file. Orthologous comparison with human was done for genomes that could be matched to human genome by "targetSpecies_homolog_associated_gene_name" in Biomart using getLDS() function.

Cell and tissue specific chromatin state annotations were based on the 25-state ChromHMM model based on imputed data for 12-marks (Ernst and Kellis, 2015; Roadmap Epigenomics Consortium et al., 2015). The chromatin state annotations from a ChromHMM model that was not specific to a single cell or tissue type were from (Vu and Ernst, 2020). We also provide in the annotation files of the array ChromHMM chromatin state annotations for mouse from (Gorkin et al., 2020). The human-mouse LECIF score was from (Kwon and Ernst, 2020).

CpG island annotation

We called CpG islands using the "gCluster" algorithm(Gómez-Martín et al., 2018). This algorithm uses clustering methods to identify the sequences that have high G+C content and CpG density with the default parameters. Besides CpG island status, this algorithm calculated several other attributes including length, GC content, and CpG density for each defined island. The outcome of this algorithm was a BED file that was used to annotate the probes using the "annotatr" package in R by checking the overlap of the aligned probes and CpG island genomic coordinates.

Human DNA methylation distribution

We downloaded the fraction methylated values based on whole genome bisulfite

sequencing data from 37 different cells and tissues types from the Roadmap Epigenomics

Consortium

(http://egg2.wustl.edu/roadmap/data/byDataType/dnamethylation/WGBS/FractionalMethylation.t

ar.gz)(Roadmap Epigenomics Consortium et al., 2015). For each CpG, we averaged the fractional

methylation values across the Roadmap samples.

GREAT analysis

We applied the GREAT analysis software tool(McLean et al., 2010) to conduct gene set

enrichments for genes near CpGs on the array in human and mouse. The GREAT software

performs both a binomial test (over genomic regions) and a hypergeometric test over genes when

using a whole genome background. We performed the enrichment based on default settings

(Proximal: 5.0 kb upstream, 1.0 kb downstream, plus Distal: up to 1,000 kb) for gene sets

associated with GO terms, MSigDB, PANTHER and KEGG pathway. To avoid large numbers of

multiple comparisons, we restricted the analysis to the gene sets with between 10 and 3,000

genes. We report nominal P values and two adjustments for multiple comparisons: Bonferroni

correction and the Benjamini-Hochberg false discovery rate.

Tissue enrichment analysis

The enrichment of tissue specific genes was done by TissueEnrich R package(Jain and Tuteja,

2019) using teEnrichment() function limited to human protein atlas(Uhlén et al., 2015) and mouse

21

ENCODE(Yue et al., 2014) databases.

Normalization methods

R software scripts implementing normalization methods can be accessed through our webpage (see the section on Data availability). Two software scripts are currently available for extracting beta values from raw signal intensities, based on Minfi and SeSAMe, respectively. Both methods use the noob method (Triche et al., 2013) for background subtraction. The two scripts evaluate each probe's hybridization and extension performance using normalization control probes and Infinium-I probe out-of-band measurements (the pOOBAH method (Zhou et al. 2018), respectively. Users can use the detection p-values for each CpG to filter out non-significant

Calibration data

We generated methylation data on two different platforms: the mammalian methylation array (HorvathMammalMethylChip40) and the human EPIC methylation array.

methylation readouts from probes unlikely to work in the target species.

The DNA samples from each species were enzymatically manipulated so that they would exhibit 0%, 25%, 50%, 75% and 100% percent methylation at each CpG location, respectively. We purchased premixed DNA standards from EpigenDx Inc (products 80-8060H-PreMixHuman, 80-8060M-PreMixMouse, and Standard80-8060R-PreMixRat Premixed Calibration Standard).

The variable "ProportionMethylated" (with ordinal values 0, 0.25, 0.5, 0.75, 1) can be interpreted as a benchmark for each CpG that maps to the respective genome. Thus, the DNA methylation levels of each CpG are expected to have a high positive correlation with ProportionMethylated across the arrays measurement from a given species. The mammalian array was applied to synthetic DNA data from 3 species: human (n=10 mammalian arrays), mouse (n=20), and rat (n=15).

Similarly, the human EPIC array was applied to calibration data from of mouse (n=15 EPIC arrays) and rat (n=10). Thus, we applied 3 EPIC arrays and 2 EPIC arrays per value (0, 0.25, 0.5, 0.75,

22

1) of ProportionMethylated in our mouse and rat studies, respectively. The EPIC array data were

normalized using the noob method (R function preprocessNoob in minfi).

Data availability

The mammalian methylation array (HorvathMammalMethylChip40) is registered at the NCBI

Gene Expression Omnibus (GEO) as platform GPL28271. The chip manifest file, calibration data,

supplementary data, and R software scripts are or will be available from available

https://github.com/shorvath/MammalianMethylationConsortium/ or the Gene Expression

Omnibus.

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Award (JE).

Conflict of Interest Statement

The Regents of the University of California is the sole owner of a provisional patent application

directed at this invention for which AA, JE and SH are named inventors. SH is a founder of the

non-profit Epigenetic Clock Development Foundation, which plans to license several patents from

his employer UC Regents, and distributes the mammalian methylation array. Bret Barnes is an

employee for Illumina Inc which manufactures the mammalian methylation array. The other

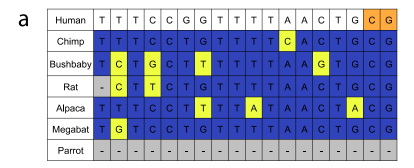
23

authors declare no conflicts of interest.

		No. CpGs whose correlation with the ProportionMethylation > threshold		
Species	Threshold	Mammal+Sesame	Mammal+Minfi	EPIC+Minfi
Mouse	0.85	27,868	26,944	4,550
Mouse	0.90	24,050	22,207	2,356
Mouse	0.95	16,444	12,797	604
Rat	0.85	26,425	25,779	17,650
Rat	0.90	22,427	20,989	6,159
Rat	0.95	15,101	12,848	819
Human	0.85	36,438	35,761	NA
Human	0.90	34,547	33,402	NA
Human	0.95	30,327	28,445	NA

Table 1. Correlating DNA methylation levels with calibration data. We evaluated the Mammalian Methylation Array with two different software methods for normalization: SeSaMe and Minfi (noob normalization). The EPIC array data were only normalized with the noob normalization method in Minfi. As indicated in the first column, the DNA samples came from three species: human (n=10 arrays), mouse (n=20), and rat (n=15). For each species, the "artificial" chromosomes exhibited on average 0%, 25%, 50%, 75% and 100% percent methylation at each CpG location. Thus, the variable "ProportionMethylated" (with ordinal values 0, 0.25, 0.5, 0.75, 1) can be considered as benchmark/gold standard. The table reports the number of CpGs for which the Pearson correlation with the ProportionMethylation was greater than the correlation threshold (second column).

Figures



b

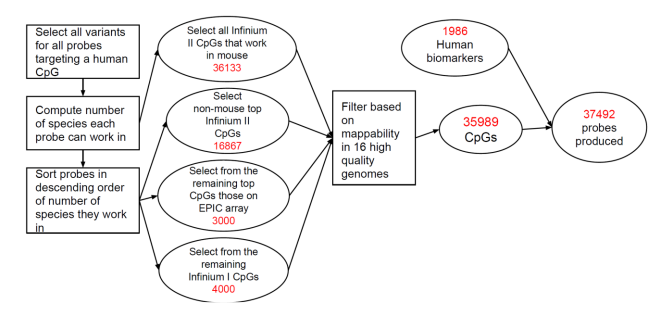


Figure 1. Overview of mammalian methylation array design process.

(a) Toy example of multiple sequence alignment at a CpG site considered by the CMAPS algorithm. The orange coloring highlights the CpG being targeted. Positions where other species have alignment that matches the human sequence are in dark blue; positions where other species have alignment that does not match the human sequence are in neon yellow; positions where other species have no alignment are in grey. (b) Flowchart detailing the selection of probes on the array by the CMAPS algorithm. A small fraction of probes designed were dropped during the manufacturing process.

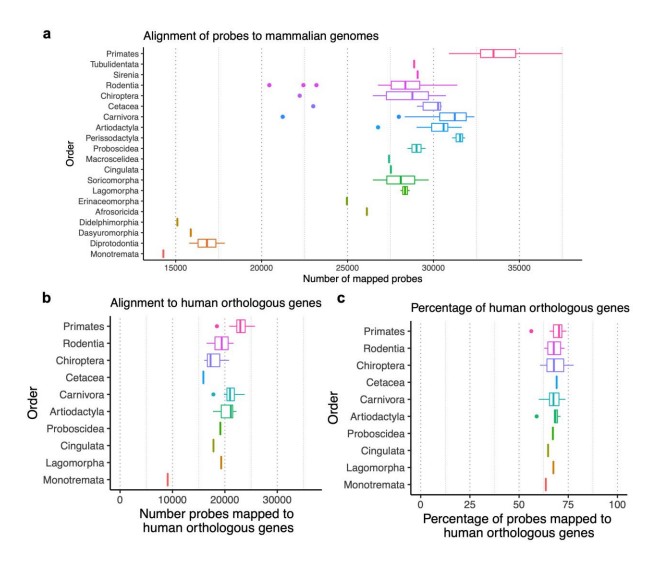


Figure 2. CpG and gene coverage of probes on the mammalian methylation array across different phylogenetic orders.

(a) Probe localization based on the QUASR package (Gaidatzis et al., 2015). The rows correspond to different phylogenetic orders. The phylogenetic orders are ordered based on the phylogenetic tree and increasing distance to human. The boxplots report the median number of mapped probes across species from the given phylogenetic order. (b) The number of probes mapped to human orthologous genes for a subset of genomes (Methods). (c) Percentage of the probes associated with human orthologous genes among mapped probes in these species.

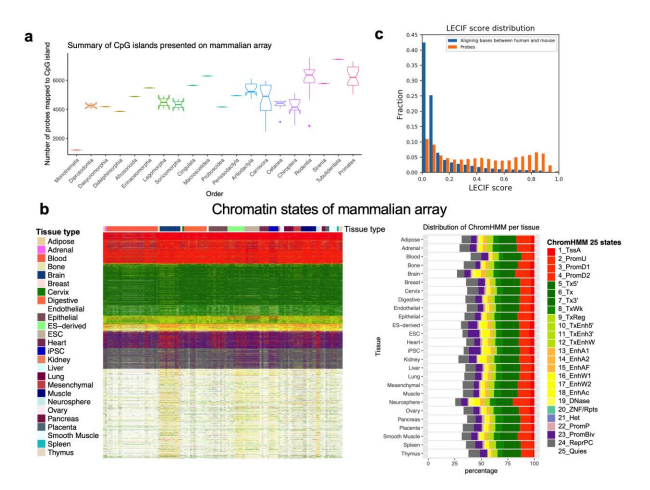


Figure 3. CpG island and chromatin state analysis of mammalian methylation probes.

We characterize the CpGs located on the mammalian methylation array regarding (a) CpG island status in different phylogenetic orders, (b) chromatin state analysis, and (c) LECIF score of evidence of human-mouse conservation at the functional genomics level. (a) The boxplots report the median number (and interquartile range) of CpGs that map to CpG islands in mammalian species of a given phylogenetic order (x-axis). The notch around the median depicts the 95% confidence interval. (b) The heatmap visualizes the ChromHMM chromatin state annotations of the location of the CpGs on the array (rows) in different human tissues (columns)(Ernst and Kellis, 2012, 2015). The colors correspond to 25 human chromatin states as detailed in the right panel. The probes in the left panel heatmap are ordered by the chromatin state with the maximum median frequency across 127 human cell and tissue types. The right panel indicates the distribution of chromatin states in each tissue type represented on the mammalian methylation array. (c) Comparison of distribution of LECIF score for probes on the array and aligning bases between human and mouse. The LECIF score has been binned as shown on the x-axis, and the fraction of probes or aligning bases with scores in that bin are shown on the y-axis.

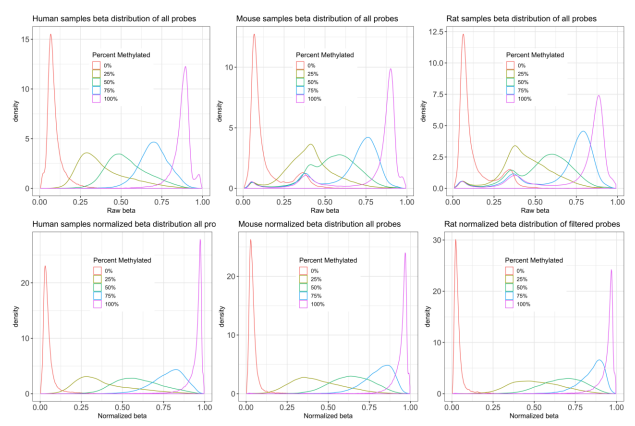


Figure 4. Distribution of probe intensities within sample, colored by the expected percentage of methylation at each site.

(a-c) Distribution of beta values (relative intensity) of all probes on the array before normalization for (a) human samples, (b) mouse samples, and (c) rat samples. (d-f) Distribution of probe intensity after Sesame normalization and restricting probes to those that CMAPS designed to (d) the human genome in human samples, (e) the mouse genome in mouse samples, and (f) the rat genome in rat samples.

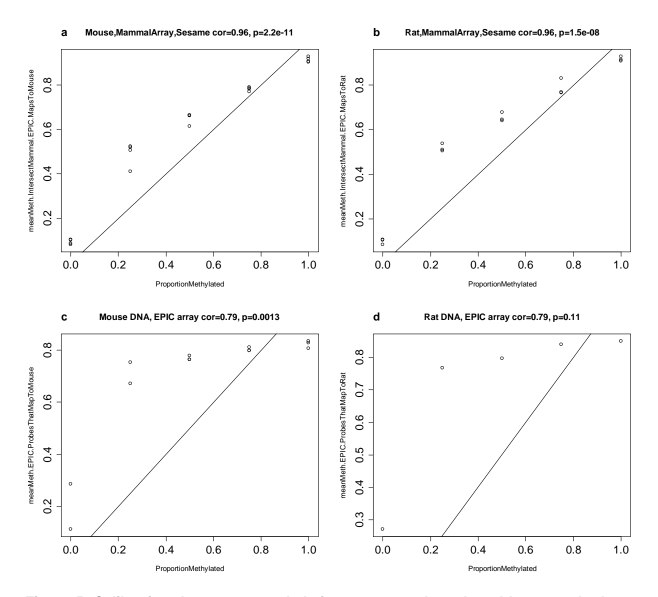
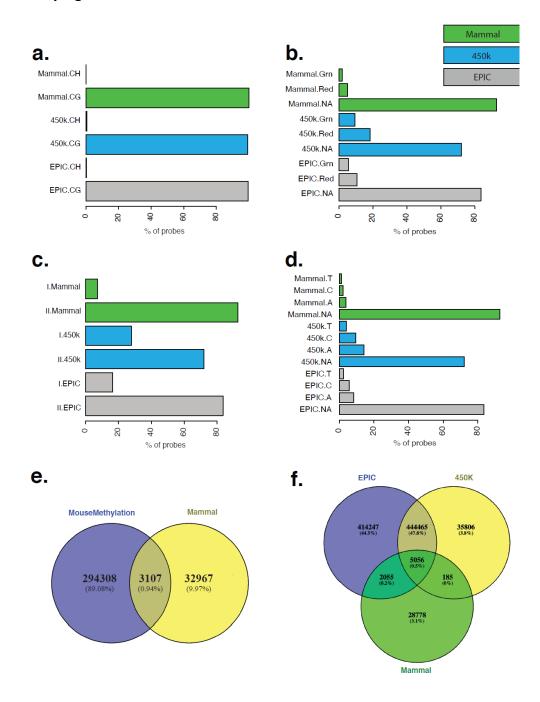


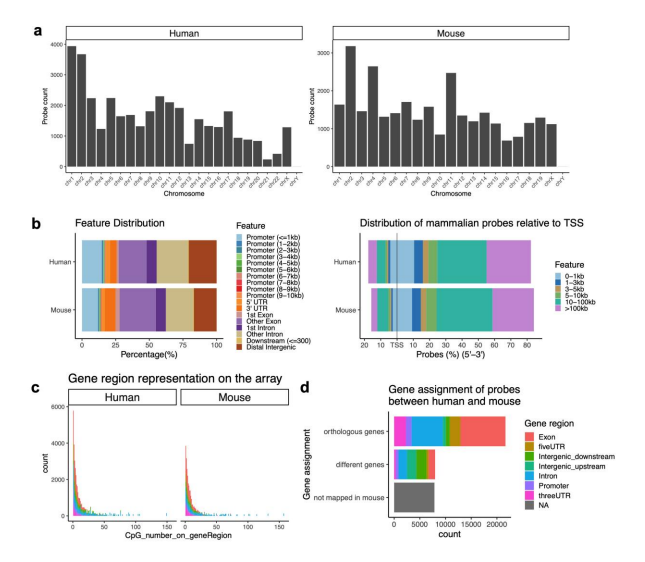
Figure 5. Calibration data: mean methylation across probes shared between the human EPIC array and the mammalian array. The mammalian methylation array contained 5574 probes targeting the same CpG that can also be found on the human EPIC array that were not included based on being human biomarkers. However, the mammalian array probes were engineered differently than EPIC probes so that they would more likely work across mammals. By applying both array types to calibration data, we are able to compare the calibration of the overlapping probes in mice (a,b) and rats (c,d). Upper panels (a,b) and lower panels (c,d) present the results for the mammalian array and the EPIC array, respectively. The benchmark measure (ProportionMethylated, x-axis) versus the mean value across roughly 4341 CpGs that map to mice (a,c) and roughly 3948 CpGs that map to rats (b,d). The mean methylation (y-axis) was formed across a subset of CpGs that i) are present on the human EPIC array, ii) present on the mammalian array, and iii) apply to the respective species according to the mappability analysis genome coordinate file.

Supplementary Figures

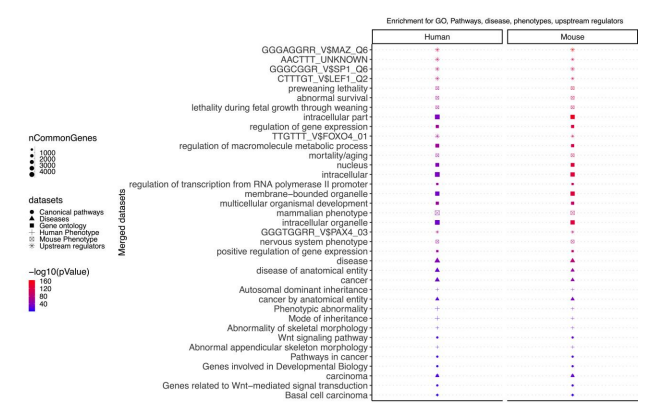


Supplementary Figure S1: Comparison of probe context between the Illumina EPIC, 450K and the Mammalian Methylation array: (a) Analysis of CpG and non-CpG (CH) probes, (b) color channel assignment, (c) type I and type II probes, and (d) next base reveals similar percentages across probes from these three array platforms. Color channel assignment and probe basepair context are important for DNA methylation array analysis and the similarity between these different arrays can facilitate extension of published analysis and normalization methods. Analysis of type I and type II probes shows a slightly lower percentage of type I probes for the mammalian methylation array. Type I probes assay DNA methylation using one color channel and two bead

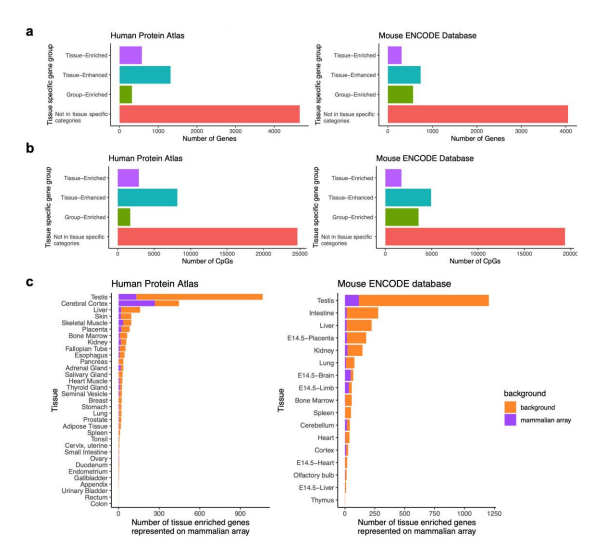
types, i.e. one unmethylated bead type and one methylated bead type. Conversely, type II probes assay DNA methylation using one bead type and two color channels indicating methylated and unmethylated cytosines. Adjustment for DNA methylation signal detected by these different probe types is one of the most important steps in DNA methylation array normalization, and a sufficient number of type I probes were included in the Mammalian Methylation array to facilitate the extension of published data normalization methods. (e) Comparison of shared and non-shared probes between the Mammalian Methylation array and MouseMethylation array loci reveals 3107 shared probes. (f) Comparison of shared and non-shared probes between the EPIC, 450k and the Mammalian methylation array. Comparative analysis was performed using Illumina probe IDs, which are unique to each probe. Intersection of IDs between arrays reveals over 5,000 probes that are common to all platforms (center). These probes can be used to follow up published human epigenome-wide association study (EWAS) results in model organisms such as mouse (*Mus musculus*) or rat (*Rattus norvegicus*), or across a range of other species, including all primates and other mammals.



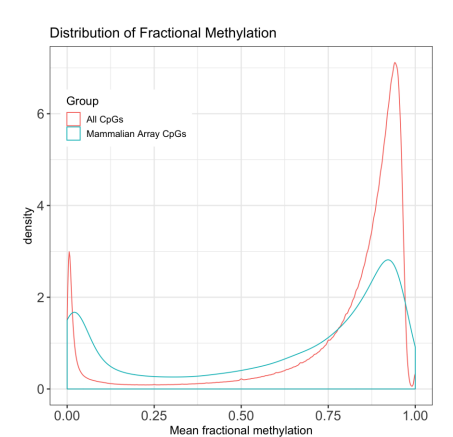
Supplementary Figure S2. Chromosome and gene region analysis of mammalian methylation probes in humans and mice. The analysis is based on mapping probes on the mammalian methylation array to the human (hg19) and mouse (mm10) genome using QUASR package(Gaidatzis et al., 2015). (a) The number of probes per human and mouse chromosome. (b) The left panel reports the percentage of probes that are located in different gene regions (promoters, 5' UTR, 3' UTR, introns, exons) in humans and mice. The right panel reports the distribution of the probes relative to the nearest transcriptional start site. (c) Histogram of CpG number in different gene regions in human and mouse genomes (as defined in the legend of panel d). (d) Alignment to orthologous genes between humans and mice. The colors indicate the mapped gene region in the mouse genome. The unique region assignment are prioritized as follows: exons, promoters, introns, 5' UTR, 3' UTR.



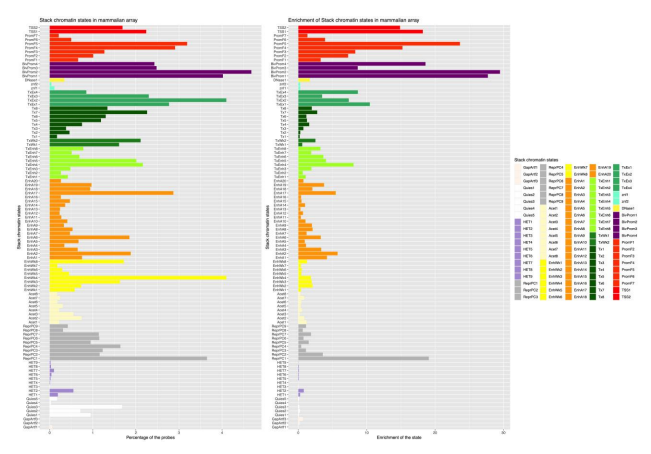
Summary Figure S3. GREAT gene set enrichment analysis of all probes on the mammalian methylation array. The figure shows the top enriched pathway based on gene-level enrichment analysis for genes proximal to probes using GREAT 7 . The two columns correspond to enrichment analysis for human (hg19) and mouse (mm10) genomes, respectively, using the whole genome as background. The top five enriched datasets from each category (Canonical pathways, diseases, gene ontology, human and mouse phenotypes, and upstream regulators) were selected and further filtered for significance at p < 10^{-5} . The category is indicated by the shape, the number of genes by the size of the shape, and the significance of the enrichment is indicated by the color scale.



Supplementary Figure S4. Human and mouse tissue-specific probes on mammalian methylation array. Characterization of the tissue specificity of CpG probes on the mammalian methylation array using the human protein atlas(Uhlén et al., 2015) and mouse ENCODE gene expression data(Yue et al., 2014). The left and right panels report results for human and mouse genomes, respectively. Each probe is mapped to the closest gene while other genes in the flanking region are ignored in this analysis. The number of genes (a) and the number of CpG probes (b) versus a categorical measure of tissue specificity. The categories on the y-axis have the following definitions. The following categories are defined in the TissueEnrich software "Tissue Enriched" labels genes with an expression level greater than 1 (TPM or FPKM) that also have at least five-fold higher expression levels in a particular tissue compared to all other tissues. "Group Enriched" labels genes with an expression level greater than 1 (TPM or FPKM) that also have at least five-fold higher expression levels in a group of 2-7 tissues compared to all other tissues, and that are not considered Tissue Enriched. "Tissue Enhanced" labels genes with an expression level greater than 1 (TPM or FPKM) that also have at least five-fold higher expression levels in a particular tissue compared to the average levels in all other tissues, and that are not considered Tissue Enriched or Group Enriched. (c) The number of tissue-enriched genes represented on mammalian array vs background in human and mouse transcriptome.



Supplementary Figure S5. Distribution of DNA methylation levels. Distribution of average fractional methylation across 37 cell and tissue types(Roadmap Epigenomics Consortium et al., 2015) at CpG sites on the array (blue) and all sites in the genome (red).



Supplementary Figure S6: Mammalian Methylation Array enrichment for Universal Chromatin State Annotations. (Left) Distribution of probe overlap with a universal chromatin state annotation by the stacked modeling approach of ChromHMM applied to data from more than 100 cell or tissue types(Vu and Ernst, 2020). (Right) The same as left, but showing the fold enrichments of the state relative to a uniform background. The strongest enrichment is seen for some bivalent promoter states. A full characterization of the states can be found in (Vu and Ernst, 2020).

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