V21: Analysis of DNA methylation data

Epigenetics refers to **alternate phenotypic states** that are

not based on differences in genotype,

and are potentially reversible,

but are generally stably maintained during cell division.

Examples: imprinting, twins, cancer vs. normal cells, differentiation, ...

Narrow interpretation of this concept : stable differential states of gene expression.

Laird, Hum Mol Gen 14, R65 (2005)

What is epigenetics?

A much **more expanded view** of epigenetics has recently emerged in which multiple mechanisms interact to collectively establish

- alternate states of chromatin structure (open packed/condensed),
- histone modifications,
- composition of associated proteins (e.g. histones),
- transcriptional activity,
- activity of microRNAs, and
- in mammals, cytosine-5 DNA methylation at CpG dinucleotides.

Waddington epigenetic landscape for embryology



b) Later depiction of the elandscape. The ball repres

Slack, Nature Rev Genet 3, 889-895 (2002)

Waddington worked in embryology

a) is a painting by John Piper that was used as the frontispiece for Waddington's book *Organisers and Genes*.
It represents an epigenetic landscape.

Developmental pathways that could be taken by each cell of the embryo are metaphorically represented by the path taken by water as it flows down the valleys.

scape. could be o are he path the valleys. Conrad Hal Waddington (1905 – 1975) pictures.royalsociety.org

b) Later depiction of the epigenetic
landscape. The ball represents a cell, and
the bifurcating system of valleys represents
bundles of trajectories in state space.

Basic principles of epigenetics: DNA methylation and histone modfications

The human genome contains ~20 000 genes that must be expressed in specific cells at precise times.

In cells, DNA is wrapped around clusters (octamers) of globular **histone** proteins to form **nucleosomes**.

These nucleosomes of DNA and histones are organized into **chromatin**, the building block of a chromosome.



Fig. 1. Carriers of epigenetic information: DNA and nucleosome. The left panel shows a DNA double helix that is methylated symmetrically on both strands (orange spheres) at its center CpG (PDB structure: 329d). DNA methylation is the only epigenetic mechanism that directly targets the DNA. The right panel shows a nucleosome spindle consisting of eight histone proteins (center), around which two loops of DNA are wound (PDB structure: 1KX5). The nucleosome is subject to covalent modifications of its histones and to the binding of non-histone proteins.

Rodenhiser, Mann, CMAJ 174, 341 (2006)

Bock, Lengauer, Bioinformatics 24, 1 (2008)

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Epigenetic modifications



Reversible and site-specific **histone modifications** occur at multiple sites at the unstructured histone tails through **acetylation**, **methylation** and **phosphorylation**.

DNA methylation occurs at 5-position of cytosine residues within CpG pairs in a reaction catalyzed by DNA methyltransferases (DNMTs).

Cytosine methylation

<u>Observation</u>: 3-6 % of all cytosines are methylated in human DNA. This methylation occurs (almost) exclusively when cytosine is followed by a guanine base -> **CpG dinucleotide**.



This is typically explained in the following way:

As most CpGs serve as targets of DNA methyltransferases, they are usually methylated (see following page)

Ésteller, Nat. Rev. Gen. 8, 286 (2007) www.wikipedia.org

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Cytosine methylation

5-Methylcytosine can easily **deaminate** to **thymine**.



If this mutation is not repaired, the affected CpG is permanently converted to TpG (or CpA if the transition occurs on the reverse DNA strand).

Hence, methylCpGs represent **mutational hot spots** in the genome. If such mutations occur in the germ line, they become heritable.

A constant loss of CpGs over thousands of generations can explain the low frequency of this special dinucleotide in the genomes of human and mouse.

> Esteller, Nat. Rev. Gen. 8, 286 (2007) Bioinformatics III www.wikipedia.org

chromatin organization affects gene expression

B Gene "switched on" Active (open) chromatin Unmethylated cytosines (white circles) Acetylated histones Gene "switched off" Silent (condensed) chromatin Methylated cytosines (red circles) Deacetylated histones

Schematic of the reversible changes in chromatin organization that influence gene expression:

genes are expressed (switched on) when the chromatin is **open** (active), and they are inactivated (switched off) when the chromatin is **condensed** (silent).

White circles = unmethylated cytosines;

```
red circles = methylated cytosines.
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Rodenhiser, Mann, CMAJ 174, 341 (2006)

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Enzymes that control DNA methylation and histone modfications

These dynamic chromatin states are controlled by reversible epigenetic patterns of **DNA methylation** and **histone modifications**.

Enzymes involved in this process include

- DNA methyltransferases (DNMTs),
- histone deacetylases (HDACs),
- histone acetylases,
- histone methyltransferases and the
- methyl-binding domain protein MECP2.



For example, **repetitive** genomic sequences

(e.g. human endogenous retroviral sequences

= HERVs) are heavily methylated,

which means transcriptionally silenced.

Rodenhiser, Mann, CMAJ 174, 341 (2006) Feinberg AP & Tycko P (2004) Nature Reviews: 143-153

DNA methylation

Typically, unmethylated clusters of CpG pairs are located in **tissue-specific genes** and in essential **housekeeping genes**.

(House-keeping genes are involved in routine maintenance roles and are expressed in most tissues.)

These clusters, or **CpG islands**, are targets for proteins that bind to unmethylated CpGs and initiate gene transcription.

In contrast, **methylated CpGs** are generally associated with silent DNA, can block methylation-sensitive proteins and can be easily mutated.

The loss of normal DNA methylation patterns is the best understood epigenetic cause of disease.

In animal experiments, the removal of genes that encode DNMTs is lethal; in humans, overexpression of these enzymes has been linked to a variety of cancers.

Rodenhiser, Mann, CMAJ 174, 341 (2006)

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CpG islands

CpG islands are characterized by a high density of CpG dinucleotides that can be targeted by DNA methylation.

CpG islands are regulatory elements and are often located in the promoter region of genes.

Criteria to define CpG islands:

Gardiner-Garden and Frommer:

≥ 200 bp length, G + C ≥ 50% CpG_{obs}/CpG_{exp} ≥ 0.6

Takai and Jones:

≥ 500 bp length G + C ≥ 55% CpG_{obs}/CpG_{exp} ≥ 0.65.

Hutter, Helms, Paulsen, Genomics 88, 323 (2006)



CpG islands

Average total length of CpG islands per gene in repeat-masked sequences at five different locations in (A) Mouse, (B) human.

Imprinted genes are monoallelically expressed, the other allele is silenced by DNA methylation. About 100 imprinted genes are experimentally confirmed.

Ctrl1, ctrl2: groups of randomly selected (most likely biallelic) control genes

Takai and Jones parameters

-> CpG islands frequent in promoters and in the gene body of imprinted genes.

Hutter, Helms, Paulsen, Genomics 88, 323 (2006) Bioinformatics III

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Differentiation linked to alterations of chromatin structure



(B) Upon
differentiation,
inactive genomic
regions may be
sequestered by
repressive chromatin
enriched for
characteristic histone
modifications.

(A) In pluripotent cells,chromatin is hyperdynamic and globally accessible.

ML Suva et al. Science 2013; 339:1567-1570

Altered DNA methylation upon cancerogenesis

Normal cell 9 Tumour-suppressor gene with promoter CpG island Locus with methylated 5'-regulatory region,
 Repetitive sequences, 'Open' chromatin conformation e.g. germline-specific gene e.g. transposable element Cancer cell F1 CpG-island hypermethylation DNA hypomethylation 'Closed' chromatin conformation 'Open' or 'relaxed' chromatin conformation Entry into cell cycle Loss of imprinting and overgrowth Avoidance of apoptosis Inappropiate cell-type expression Defects in DNA repair Genome fragility Angiogenesis Activation of endoparasitic sequences. Loss of cell adhesion Tumorigenesis Unmethylated CpG P Methylated CpG

Figure 1 | Altered DNA-methylation patterns in tumorigenesis. The hypermethylation of CpG islands of tumoursuppressor genes is a common alteration in cancer cells, and leads to the transcriptional inactivation of these genes and the loss of their normal cellular functions. This contributes to many of the hallmarks of cancer cells. At the same time, the genome of the cancer cell undergoes global hypomethylation at repetitive sequences, and tissue-specific and imprinted genes can also show loss of DNA methylation. In some cases, this hypomethylation is known to contribute to cancer cell phenotypes, causing changes such as loss of imprinting, and might also contribute to the genomic instability that characterizes tumours. E, exon. Esteller, Nat. Rev. Gen. 8, 286 (2007)

DNA methylation is typically only weakly correlated with gene expression!



Left: different states of hematopoiesis (blood cell differentiation). HSC: hematopoietic stem cell MPP1/2: multipotent progenitor cell

Right: skin cell differentiation

Bioinformatics III

Bock et al. , Mol. Cell. 47, 633 (2012)

Promoter methylation vs. gene-body methylation

The relationship between methylation and gene expression is complex.

High levels of gene expression are often associated with low **promoter methylation** but elevated **gene body methylation**.

The **causality relationships** between expression levels and DNA methylation have not yet been determined.



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Detect DNA methylation by bisulfite conversion



www.wikipedia.org

Processing of DNA methylation data with RnBeads



Left stages: processing of raw data (sequencing reads e.g. from bisulfite conversion)

Assenov et al. Nature Methods 11, 1138–1140 (2014)

DNA methylation analysis with RnBeads



Top: read coverage of CpGs

Distribution of beta-values

Bottom: "Volcano" plot x-axis – difference of methylation site between 2 probes, y-axis – statistical significance of the difference;

Assenov et al. Nature Methods 11, 1138–1140 (2014)

Require enough variation and enough significance

DNA fiber forms B-DNA

Z-DNA

Requires more methylation, higher concentration of physiological salts



Dry Environment

A-DNA

Most prominent in cellular conditions

Equilibrium shift with specific conditions

Protein-DNA^{Me} interaction (R.DpnI from *E.coli***)**



Left: structural transitions of DNA affect accessibility of the base pairs

Right: recognition of 6-methylated adenine (common form of DNA methylation in bacteria)

Siwek et al. Nucl. Acids Res. (2012) 40 (15): 7563-7572.



Protein-DNA^{Me} interaction (R.DpnI from *E.coli***)**

Binding of bacterial restriction enzyme R.DpnI to adeninemethylated or unmethylated target sequence -> methylation has clear effects on width of major groove Binding of MeCP2 to cytosinemethylated or unmethylated target sequence -> methylation has smaller effects on width of major groove

PhD thesis Siba Shanak (2015)

Beta-values measure fractional DNA methylation levels

After analysis of raw sequencing data + filtering of problematic regions etc

the degree of methylation is typically expressed as fractional **beta value:** %mCG(i) / (%mCG(i) + %CG(i))

A beta value for CpG position *i* takes on values between 0 (position *i* not methylated) and 1 (position *i* fully methylated)

Methylation levels of neighboring sites are correlated

- Observation: methylation levels of neighboring CpG positions within 1000 bp are often correlated;
- distance between neighboring CpGs is ca. 100 bp (1% frequency)
- Idea: exploit this effect to "smoothen" experimental data,
 e.g. when this is obtained at low coverage

Master thesis of Junfang Chen (February 2014):

Journal of Bioinformatics and Computational Biology Vol. 12, No. 6 (2014) 1442005 (16 pages) © Imperial College Press DOI: 10.1142/S0219720014420050



AKSmooth: Enhancing low-coverage bisulfite sequencing data via kernel-based smoothing

Junfang Chen^{*,†,‡}, Pavlo Lutsik[†], Ruslan Akulenko^{*}, Jörn Walter^{†,§} and Volkhard Helms^{*,§}

*Center for Bioinformatics, Saarland University Saarbrücken 66123, Germany

[†]Department of Genetics, Saarland University Saarbrücken 66123, Germany [‡]s9juchen@stud.uni-saarland.de

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Correlated methylation of neighboring CpGs

 $\hat{f}_{h}(t) = \frac{\sum_{i}^{N} K_{h}(t, i) C_{t}(i) y_{i}}{\sum_{i}^{N} K_{h}(t, i) C_{t}(i)}, \quad h: \text{"band-width": size of window}$

$$K_h(t,i) = K\left(\frac{|i-t|}{h}\right),$$

$$C_t(i) = \begin{cases} g_t & \text{if } i = t; \\ 1 & \text{if } i \neq t. \end{cases}$$

(# of neighboring CpGs around *t*)

y_i: methylation level of *i*-th CpG site within window of given size

 C_t (i): weighting factor to consider read coverage of neighboring CpG sites relative to that of target site

 $K_{h}(t, i)$: Kernel function that considers the distance between positions t and i.

-> more distant positions get smaller weight.

Choice of kernel function

The kernel K

$$K_h(t,i) = D\left(\frac{|i-t|}{h}\right),$$

is either a standard Gaussian function

$$D(\mu) = \frac{1}{h\sqrt{2\pi}} e^{-\frac{1}{2}\mu^2}$$

or the Epanechnikov kernel

$$D(\mu) = \begin{cases} \frac{3}{4}(1-\mu^2) & \text{if } |\mu| \le 1; \\ 0 & \text{otherwise} \end{cases}$$

or the tricubic kernel

$$D(\mu) = \begin{cases} \frac{70}{81} (1 - |\mu|^3)^3 & \text{if } |\mu| \le 1; \\ 0 & \text{otherwise.} \end{cases}$$



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Correlation of low-coverage and high-coverage data



Three Cancer Samples on Autosome

C1, C2, C3 are three different samples.

Best results for window considering nearby 10-20 CpGs.

Gaussian kernel ("hg") more robust with distance (exponential weighting).

Tricubic and Epanechikov kernels show stronge decrease for large windows.

Every method was tested for including neighboring

5, 10, 15, ... 70 CpGs.

Red symbols "hl": low-coverage data (unsmoothened)

Brown symbols "hb": low-coverage data processed with (another) Bsmooth-program

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DNA methylation in breast cancer



DNA methylation in cancer



The Cancer Genome Atlas

doi:10.1038/nature11412

Comprehensive molecular portraits of human breast tumours

The Cancer Genome Atlas Network*

ARTICLE



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The Cancer Genome Atlas

DNA methylation

Illumina Infinium DNA methylation arrays were used to assay 802 breast tumours. Data from HumanMethylation27 (HM27) and HumanMethylation450 (HM450) arrays were combined and filtered to yield a common set of 574 probes used in an unsupervised clustering analysis, which identified five distinct DNA methylation groups (Supplementary Fig. 8). Group 3 showed a hypermethylated phenotype and was significantly enriched for luminal B mRNA subtype and under-represented for *PIK3CA*, *MAP3K1* and *MAP2K4* mutations. Group 5 showed the lowest levels of DNA methylation, overlapped with the basal-like mRNA subtype, and showed a high frequency of *TP53* mutations. HER2-positive (HER2⁺) clinical status, or the HER2E mRNA subtype, had only a modest association with the methylation subtypes.

A supervised analysis of the DNA methylation and mRNA expression data was performed to compare DNA methylation group 3 (N = 49) versus all tumours in groups 1, 2 and 4 (excluding group 5, which consisted predominantly of basal-like tumours). This analysis identified 4,283 genes differentially methylated (3,735 higher in group 3 tumours) and 1,899 genes differentially expressed (1,232 downregulated); 490 genes were both methylated and showed lower expression in group 3 tumours (Supplementary Table 4). A DAVID (database for annotation, visualization and integrated discovery) functional annotation analysis identified 'extracellular region part' and 'Wnt signalling pathway' to be associated with this 490-gene set; the group 3 hypermethylated samples showed fewer *PIK3CA* and *MAP3K1* mutations, and lower expression of Wnt-pathway genes.



Supplemental Figure 8. DNA methylation subtypes and comparison to normal breast tissues. DNA methylation cluster membership was determined by a Recursively Partitioned Mixture Model (RPMM) for 466 breast tumors using 574 selected probes and compared to 122 breast normal samples in the same probe order. DNA methylation levels (beta value) are shown with a color spectrum; blue, no methylation to yellow, full methylation. Cluster memberships are indicated by the horizontal color bar: black Cluster 1 (n=80); red Cluster 2 (n=123); green Cluster 3 (n=44) blue Cluster 4 (n=128); cyan Cluster 5 (n=91). Molecular and clinical features as indicated in the color key. P-values for association with molecular and clinical features were calculated using a Chi-square test or Fisher's exact test, wherever applicable.

Idea: identify co-methylation of genes in TCGA samples



Co-methylation of genes 1 and 3 across samples

Tumor data

National Cancer Institute The Cancer Genome Atlas Understanding genomics Data Portal Understanding genomics to improve cancer care					
Data Type (Base- Specific)	Level 1 (Raw Data)	Level 2 (Normalized/ Processed)	Level 3 (Segmented/ Interpreted)	Level 4 (Summary Finding/ROI)	
DNA Methylation	Raw signals per probe	Normalized signals per probe or probe set and allele calls	Methylated sites/genes per sample	Statistically significant methylated sites/genes across samples	

- 183 tumor samples deposited in Sept 2011 (tumor group 1);
- 134 tumor samples deposited in Oct 2011 (tumor group 2) and
- 27 matched normal samples from Oct 2011.

Difficulties: batch effect



Filter 1: delete genes affected by batch effect

Difficulties: outliers



Filter 2: require zero outliers

Difficulties: low variance



Filter 3: delete genes with low variance

$$quartile3(beta_i) - quartile1(beta_i) \ge 0,1$$
$$_{i \in T}$$

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Comparison against randomized data



Known breast cancer genes in OMIM: mostly unmethylated



These 19 genes are associated with breast cancer in the Online version of the Mendelian Inheritance in Man (OMIM) database.

They are not involved in co-methylation because most of them show little changes of their (low) methylation levels

top 10 co-methylated gene pairs

First gene	Second gene	Pearson correlation	Related genes?
SPRR1B	SPRR1A	0,872	Yes
FCN2	FCN1	0,870	Yes
CD244	CD48	0,866	Yes
SPRR1B	SPRR4	0,862	Yes
TAS2R13	PRB4	0,859	No
F7	TFF1	0,856	Νο
SH3TC2	SPARCL1	0,853	Νο
ABCE1	SC4MOL	0,849	Νο
REG1B	REG1P	0,846	Yes
SPRR3	SPRR4	0,843	Yes

Some genes have related names -> co-methylation may be expected

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Are all co-methylated genes neighbors?

Less than half of all co-methylated gene pairs lie on the same chromosome



Functional similarity of co-methylated genes



Co-methylated gene pairs on the same chromosome have higher functional similarity (determined by FunSimMat) than that between random pairs of genes

Not the case for co-methylated gene pairs on different chromosomes

Enriched pathways in co-methylated gene clusters

Cluster				
ID	KEGG pathways	p-value	Genes involved in pathways	FDR
	hsa04950:Maturity onset diabetes of			
8	the young	0.003	HNF1B, FOXA2, NEUROD1	2.622
9	hsa04640:Hematopoietic cell lineage	0.009	CD1A, CD1E, CD1D	6.229
15	hsa04730:Long-term depression	0.004	GRM5, C7ORF16, PRKG2	2.952

	hsa04060:Cytokine-cytokine receptor			
22	interaction	0.047	EGF, TNFSF18, IL20	31.263
27	hsa04512:ECM-receptor interaction	0.005	COL5A2, COL11A1, SPP1	3.500
27	hsa04510:Focal adhesion	0.029	COL5A2, COL11A1, SPP1	17.498

Table S2. The results of pathway enrichment analysis of 29 gene clusters obtained using DAVID. These clusters were formed by applying Affinity Propagation clustering to 779 genes, which were left after three-stage filtered of all 13,313 genes from methylation data samples.

Further modifications of cytosine bases



Further modifications were discovered in the last few years. They are present in cells in much smaller fractions than 5-mC.

Tet enzymes catalyze the conversions. Biological roles mostly unclear.

http://he-group.uchicago.edu

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