

Epigenetics and palms What's beyond the double helix...

Alain RIVAL, Estelle JALIGOT, Thierry BEULE, Frédérique RICHAUD, Pascal ILBERT

UMR DIADE Palm Development Group Cirad BioS Montpellier France.







What is Epigenetics?

« The interactions of genes with their environment, which bring the phenotype into being ».

Waddington, C. (1942) The Epigenoptype. Endeavour, 1, 18-20

"Changes in gene expression not attributable to nucleotide sequence variation".

'Things written in pen you can't change. That's DNA," says geneticist Danielle Reed. "Things written in pencil you can. That's epigenetics."





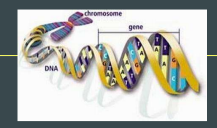
Several new sequencing technologies are emerging that have the potential to provide increases in throughput and reductions in cost

Companies such as 454 Life Sciences, Solexa, and Helicos Biosciences all have competing technologies, vying to be widely adopted for the next generation of sequencing machines

➢ The success of genome sequencing project has created wide-spread interest in exploring epigenomes in order to elucidate how the genome executes the information it holds.







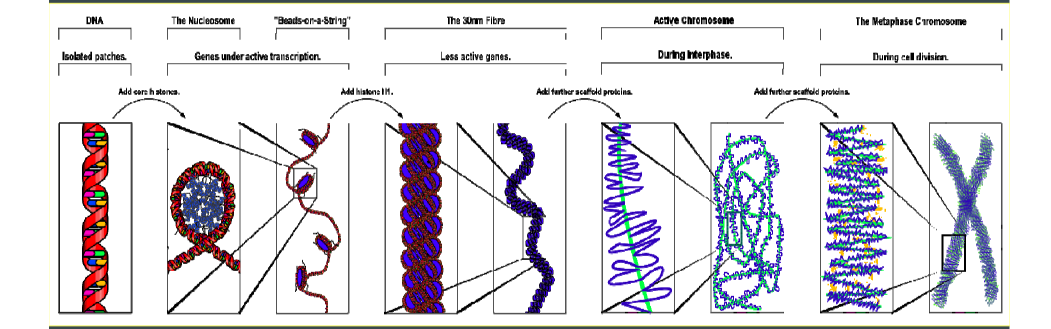
An organised DNA Packaging

- A mammal diploid cell contains in average 6000 Mb of DNA, corresponding to 1.8 meters, which is packed within a nucleus of 6 microns diameter, following a strongly organized packaging
- Each chromosome yields one single molecule of linear DNA which is folded several times and distributed over the centromere
- Within the nucleus, DNA is never encountered under a free form but associated to other molecules, principally histones. Histones are small basic proteins which are present with the same proportion as DNA.
- Other proteins associated to DNA are non histones acidic proteins, which account for 10 to 30% of the whole structure.
- This DNA-Protein complex is called chromatin (as it is easily stained) and it constitutes the chromosomes





DNA increasing orders of packing





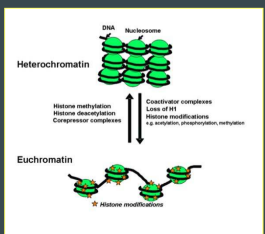


Chromatin structure and gene activity

Heterochromatin refers to regions of the genome that have low gene density, contain satellite repeat elements and are late replicating.

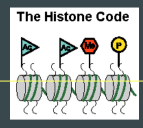
Euchromatin is a lightly packed form of chromatin that is rich in gene concentration, and is often under active transcription. Euchromatin comprises the most active portion of the genome within the cell nucleus.

Heterochromatin and euchromatin are associated with distinct DNA methylation and histone modification patterns that correlate with particular states of gene activity









Chromatine structure and gene activity

>Acetylation of histones loosens the chromatin, facilitating replication and transcription.

> Methylated histones hold the DNA more tightly, thus restricting access and impeding transcription.

>Methylation of Lysine 4 and Lysine-27 on Histone3 may be involved in development.

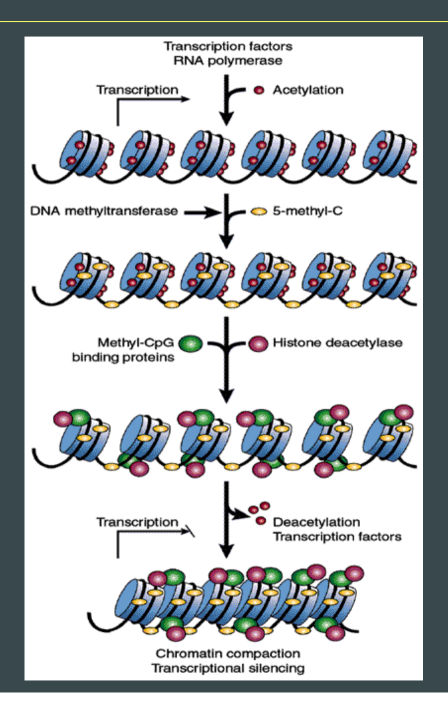
> Specifically, there is fewer methylated Lysine 27 in the chromatin of differentiated cells.

>Lysine-4 methylation acts to promote transcription by recruiting nucleosome remodeling enzymes and histone acetylase.





Epigenetic prints on chromatin



Transcriptionally active chromatin regions tend to be hyperacetylated and hypomethylated.

If a region of DNA or a gene is destined for silencing, chromatin remodeling enzymes such as histone deacetylases and ATP-dependent chromatin remodelers likely begin the gene silencing process.

One or more of these activities may recruit DNA methyltransferase resulting in DNA methylation, followed finally by recruitment of the methyl-CpG binding proteins.

The region of DNA will then be heritably maintained in an inactive state. Epigenomics is the large scale study of epigenetic marks on the genome including:

Covalent modifications of histone tails (acetylation, methylation, phosphorylation, ubiquitination)
DNA methylation.
Small RNAs

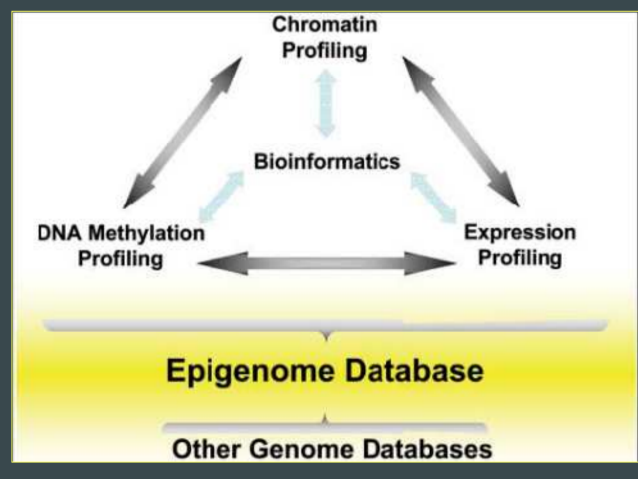
Epigenetic components are all amenable to genome-wide studies

Integrated approaches that correlate gene expression with DNA methylation and chromatin profiles are being designed.





From genome to epigenome

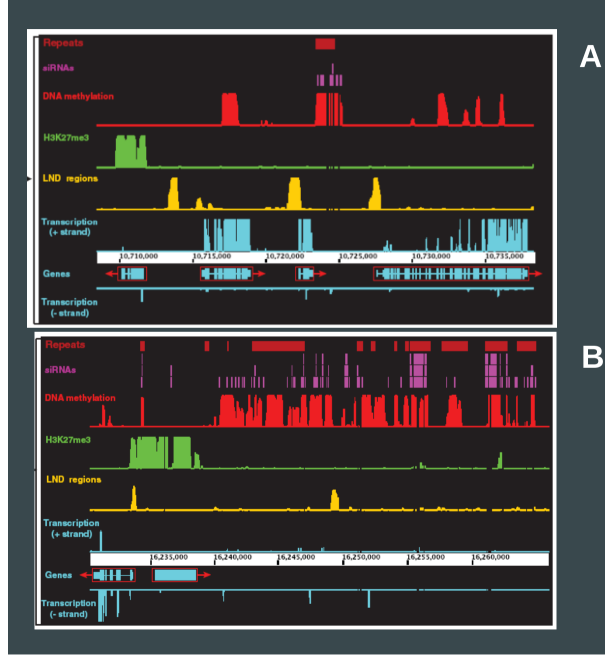


Murell et al, Human Molecular Genetics 2005 14 (Review Issue 1):





Distribution patterns and transcription activity

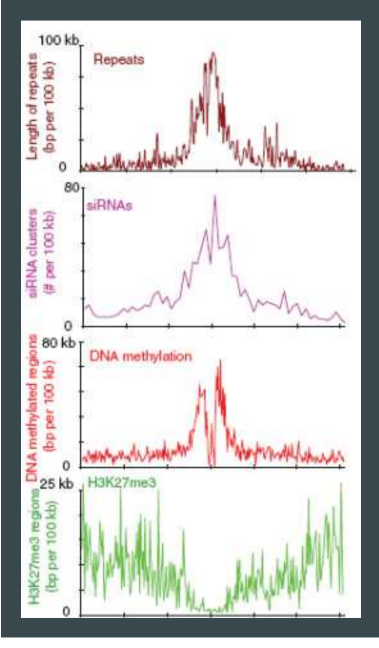


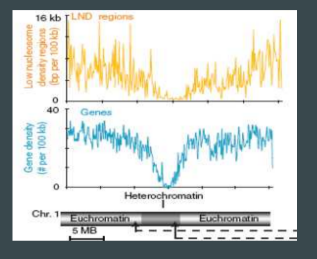
Detailed distribution patterns and transcription activity (vertical blue bars) in a gene-rich region (A) and a repeat-rich region (B).

Red boxes: genes; Arrows indicate the direction of transcription.

Zhang, Science 320, 489 (2008)

Motiv density along chromosome





Distribution of genes, repetitive sequences, DNA methylation, siRNAs, H3K27me3, and low nucleosome density (LND) regions in Arabidopsis.

The chromosomal distributions use chromosome 1 as an example. The x axis shows chromosomal position.

The small RNAs world

Breakthrough Online For an expanded version of this section, with references and links, see www. sciencernag.org/content/ vol/298/issue5502/#speck



The Winner staged by its more gamorous skiling.DNA, it turning valety of guess.RNA, long upstaged by its more gamorous skiling.DNA, is turning valety of audits.RNA long up-

Small RNAs Make Big Splash

to the or

Life cycle. With a helping hand from proteins

RISC and Diger small RNAs are been We new

know that these molecules keep DNA in line

their colleagues injected stretches of double-

stranded RNA into worms. Double-stranded

RNA forms when a familiar single strand

kinks back in a hairpin bend, putting two

complementary sequences alongside each

other. To the researchers' surprise, double-

stranded RNA dramatically inhibited genes.

that had helped generate the RNA in the first

place. This inhibition, which was later seen

in flies and other organisms, came to be

known as RNA interference (RNAi). It

helped prove that RNA molecules were be-

Another crucial step came last year,

when Gregory Hannon of Cold Spring Har-

bor Laboratory in New York and his col-

leagues identified an enzyme, appropriately

dubbed Dicer, that generates the small RNA

molecules by chopping double-stranded

RNA into little pieces. These bits belong to

20 DECEMBER 2002 VOL 298 SCIENCE www.sciencemag.org

hind some gene silencing.

and ensure a cell's good health.

missed as little more than drones, taking orders from DNA and converting genetic information into proteins. But a string of recent discoveries indicates that a class of RNA molecules called small RNAs operate many of the cell's controls. They can turn the tables on DNA, shutting down genes or altering their levels of expression. Remarkably in some species, truncated RNA molecules literally shape genomes, carving out chunks to keep and discarding others. RIDC const plan up the fi There are even hints that certain small RNAs might help chart a cell's destiny by directing genes to turn on or off during development, which could have profound implications for coaxing cells to form one type of tissue or another. Science hails these electrifying discoveries, which are prompting biologists to overhaul their vision of the cell and its evolution, as 2002's Breakthrough of the Year.

For decades, RNA molecules were dis-

These astonishing feats are performed by short stretches of RNA ranging in length from 21 to 28 nucleotides. Their role had gone unnoticed until recently, in part because researchers, focused on the familiar larger RNA molecules, tossed out the crucial small ones during experiments. As a result, RNA has long been viewed primarily as an essential but rather dull molecule hat ferries the genetic code from the nucleus to the ribosomes, the cell's protein factories, and helps assemble amino acids in the correct order during protein synthesis.

Signs that RNA might be more versatile came in the early 1990s, when biologists determined that some small RNAs could quash the expression of various genes in plant and, later, aminal cells. But they didn't appreciate the molecules' true powers until 1998. That's when Andrew Fire of the Camegie Institution of Washington in Baltimore, Maryland, Craig Mello of the University of Masschusets Medical School in Worcester, and

one of two small RNA classes produced by different types of genes: microRNAs (miRNAs) and small interfering RNAs (siRNAs). SiRNAs are considered to be the main players in RNAi, although miRNAs, which inhibit translation of RNA into protein, were recently implicated in this machinery as well. To bring about RNAi, small RNAs degrade the messenger RNA that transports a DNA sequence to the ribosome. Exactly how this degradation occurs isn't known, but scientists believe that Dicer delivers small RNAs to an enzyme complex called RISC. which uses the sequence in the small RNAs to identify and degrade messenger RNAs with a complementary sequence.

Such degradation ratchets down the expression of the gene into a protein, Although quashing expression might not sound particularly useful, biologists now believe that in plants, RNAi acts like a genome "immune system," protecting against harmful DNA or viruses that could disrupt the genome. Similar hints were unearthed in anirules this year. In labs studying gene function, RNAi is now commonly used in place of gene "Incocousts": Rather than delete a gene, a laborious process, double-stranded

RNA is applied to ramp down its expression. The year's most stumning revelations emerged in the fall, in four papers examining how RNA interference helps pilot a peculiar—and pervasive—genetic phenomenon known as epigenetics. Epigenetics refers to changes in gene expression that persist across at least one generation but are not caused by changes in the DNA code.

In recent years, researchers have found that one type of epigenetic regulation is caused by adjustments in the shape of complexes known as chromatin, the bundles of DNA and certain fundamental proteins that make up the chromosomes. By changing shape—becoming either more or less compact—chromatin can alter which genes ere expressed. But what prompts this shape• These tiny RNAs (~21-26 nt) induce silencing through homologous sequence interactions

• They can control mRNA stability or translation, or target epigenetic modifications to specific regions of the genome.

• Small RNAs and evolutionarily conserved RNA-mediated silencing pathways have established a new paradigm for understanding eukaryotic gene regulation and revealed novel host defenses to viruses and transposons.



2296



MicroRNAs (miRNAs) and transacting siRNAs (tasiRNAs) are primarily involved in regulating gene expression and plant development,

siRNAs play a major role in defending the genome against the proliferation of invading viruses and endogenous transposable elements.

The function of the fourth type of sRNAs, naturalantisense siRNAs (nat-siRNAs), is not entirely clear but is likely related to plant stress responses
Zhang et al., PNAS 104, 4536 (2007)





Deciphering the small RNAs machinery

E. Jean Finnegan and Marjori A. Matzke hp Transgene Rdp. Ago Nucleus miRNA gene Cytoplasm tone methylation (HsK9) Histone methylation (H3K9) **DNA** methylation DNA methylation Heterochromatic IRNA hp RNA -----TITITITI miRNA dsRNA Cleavage of mRNA target dsRNA hp RNA тинит Inhibition of translation of viral RNA

The Small BNA World

© Journal of Cell Science 2003 (116, pp. 4689-4693)

RNA-directed DNA methylation (RdDM), first discovered in plants requires a dsRNA that is processed to 21-24 nt small RNAs.

In Arabidopsis, links between locus specific small RNAs, DNA methyltransferases, and histone modifications, including deacetylation and histone H3 lysine 9 (H3K9) methylation have been identified (see red

Jatiway).

Some small RNAs might target native promoters of endogenous genes (see dashed blue pathway).

RNAi and epigenetic alterations of the genome, such as DNA methylation and histone modifications

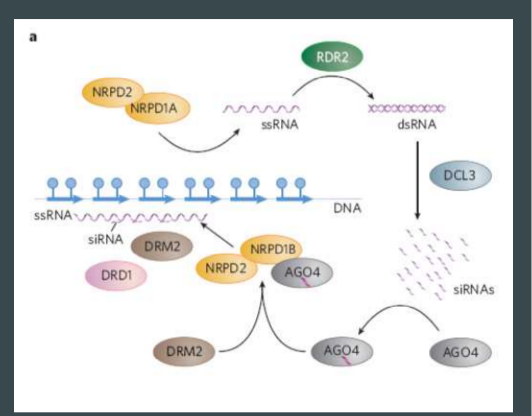


Cell Science



RNA-directed DNA methylation

Putative pathway for RNA directed DNA methylation in *A. thaliana*. Target loci (in this case tandemly repeated sequences; coloured arrows) recruit an RNA polymerase IV complex consisting of NRPD1A and NRPD2 through an unknown mechanism, and this results in the generation of a single-stranded RNA (ssRNA) species. This ssRNA is converted to double-stranded RNA (dsRNA) by the RNA-dependent RNA polymerase RDR2.



The dsRNA is then processed into 24-nucleotide siRNAs by DCL3. The siRNAs are subsequently loaded into the protein AGO4, which associates with another form of the RNA polymerase IV complex, NRPD1B-NRPD2. AGO4 that is 'programmed' with siRNAs can then locate homologous genomic sequences and guide the protein DRM2, which has *de novo* cytosine methyltransferase activity. Targeting of DRM2 to DNA sequences also involves the chromatin remodelling protein DRD1.





Conclusions

 \checkmark The 'epigenetic code' considerably extends the information potential of the genetic code.

 \checkmark Thus, one genome can generate many 'epigenomes' as the fertilised egg progresses through development and translates its information into a multitude of cell fates.

 \checkmark The transcriptomes of an organism are continually changing in response to developmental and environmental cues.

✓ The epigenome is not static and can be molded by developmental signals, environmental perturbations, and disease states.

✓ Therefore, many epigenomes will need to be sequenced for a single organism, making epigenome sequencing perhaps even more challenging than genome sequencing.





The search for epigenetic factors of flower development in oil palm



Alain Rival (Cirad)

Sylvie Doulbeau (IRD), Frédérique Aberlenc (IRD), James Tregear (IRD), Estelle Jaligot (Cirad), Pascal Ilbert (Cirad), Thierry Beulé (Cirad)





Rationale

 The commercial development of large scale propagation of oil palm is hampered by the generation of somaclonal variants affecting the floral architecture: the « mantled » genotype.

 Field observation of clonal plantings on the long term and results from DNA markers analysis (Rival et al, 1998) are consistent with an epigenetic origin for somaclonal variation.

 The "mantled" phenotype is correlated with a global hypomethylation of genomic DNA, which was revealed through a whole genome approach (Jaligot et al, 2000) and the use of Methylation-Sensitive DNA markers, such as MS-RFLPS and MSAPs (Jaligot et al, 2002; 2004).





Epigenetic factors of flower development

Journal of Experimental Botany, Vol. 59, No. 12, pp. 3271–3281, 2008 doi:10.1093/jxb/ern178 Advance Access publication 17 July, 2008



RESEARCH PAPER

Isolation and expression analysis of genes encoding MET, CMT, and DRM methyltransferases in oil palm (*Elaeis guineensis* Jacq.) in relation to the 'mantled' somaclonal variation

Alain Rival^{1,*,†}, Estelle Jaligot^{1,*}, Thierry Beulé¹ and E. Jean Finnegan² ¹ CIRAD, IRD, UMR DIAPC, BP 64501, F-34394 Montpellier, Cedex 5, France ² CSIRO Plant Industry, GPO Box 1600, ACT 2602, Canberra, Australia

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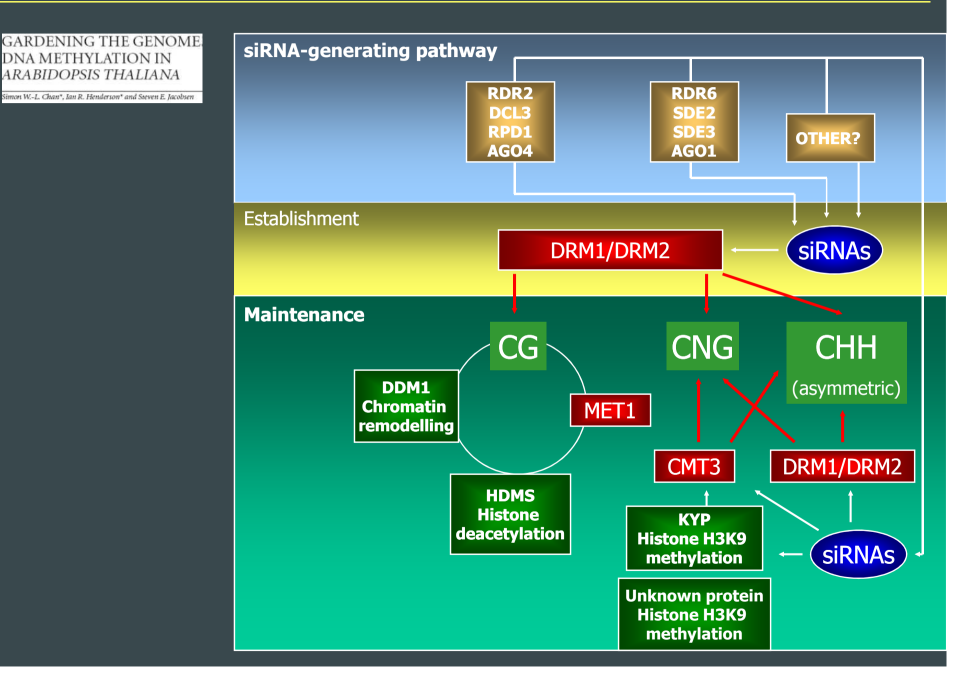




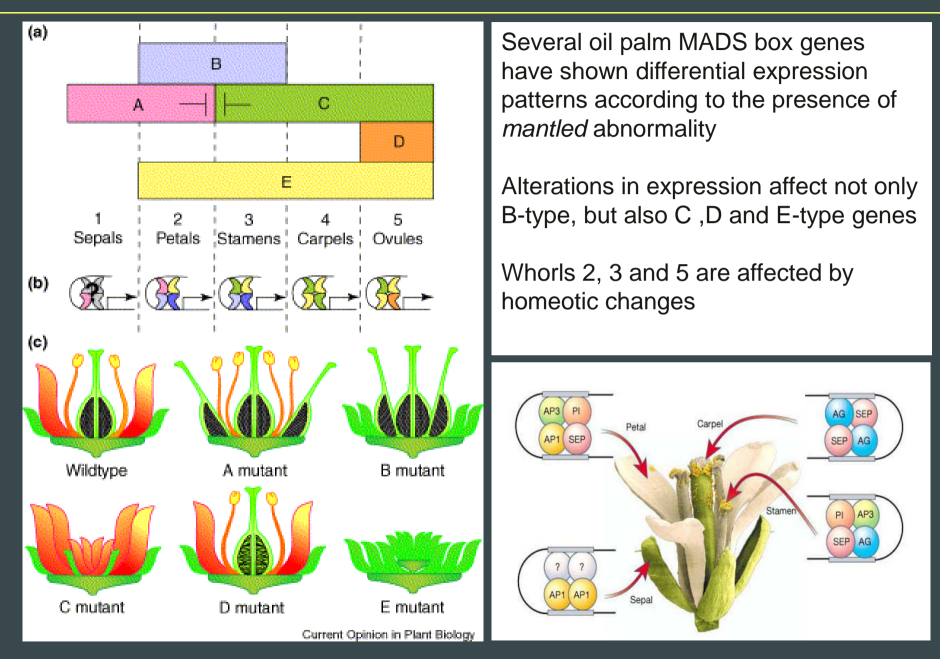
Real Time qPCR analysis on embryogenic calli



Establishment and Maintenance of DNA methylation



Methylation around MADS Box candidate genes



Methylation around MADS Box candidate genes

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Reduced expression of genes Eg DEF1 and EgGLO2 (B type), EgAG2 (C and D type) and EgS1 (type E) in abnormal oil palm flowers

Am. J. Bot.92(11):1836-1852.2005







Thank you for your kind attention ...

alain.rival@cirad.fr