The Epigenetics of human cancer

Hallmarks of human cancer

Cancer cells show Defined key features

Alterations in central biological pathways that Ensure immortality and invasiveness

This figure illustrates some of the many approaches employed in developing therapeutics targeted to the known and emerging hallmarks of cancer.

EGFR indicates epidermal growth factor receptor; CTLA4, cytotoxic T lymphocyte-associated antigen 4; mAb, monoclonal antibody; HGF, hepatocyte growth factor; VEGF, vascular endothelial growth factor; PARP, poly-(ADP ribose) polymerase.

Source: Hanahan D, Weinberg RA. Hallmarks of cancer: the next generation. Cell. 2011; 144:646-674. Reprinted with permission.

Hallmarks of human cancer – the genetics

Alterations in central biological pathways that ensure immortality and invasiveness

Cancer is caused by key oncogene activation and/or loss of tumorsuppression and paralleled by the acquisition of hundreds of mutations in different genes and also altered gene expression of mutated/non-mutated genes

Together this results up a cancer gene expression pattern/genomic instability

Genetics - Epigenetics

Table 1. Examples of key cellular pathways disrupted in human cancers by genetic and epigenetic mechanisms

TGF- β , transforming growth factor β ; *DAPK*, death-associated protein kinase.

Hallmarks of human cancer – the epigenetics

ALTERATION OF EPIGENETIC REGULATORS (Mutation/altered expression) CAN SET UP CANCER ASSOCIATED GENE EXPRESSION SIGNATURES

THE CANCER EPIGENOME

Epigenetic alterations:

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- Can be causal for the alteration of gene expression \rightarrow epigenetic writer or epigenetic readers are altered (mutated or changed expression)
- Can also be a indirect consequence \rightarrow another mechanism not directly related to epigenetics alters gene expression, epigenetic modifications change as a consequence

The epigenetic machinery

Figure 1. Genetic mutations of epigenetic modifiers in cancer. The drawing shows the input of epigenetic processes in specifying gene expression patterns. Recent whole-exome sequencing studies show that mutations in various classes of epigenetic modifiers are frequently observed in many types of cancers, further highlighting the cross talk between genetics and epigenetics. Examples of some, but not all, of these mutations are illustrated here and listed in Table 2. The mutations of epigenetic modifiers potentially cause genome-wide epigenetic alterations in cancer, but, save for isocitrate dehydrogenase (IDH) mutations as discussed in the text, these have yet to be shown on a genome-wide scale. Understanding the relationship of genetic and epigenetic changes in cancer will offer novel insights for cancer therapies. MBDs, methylcytosine-binding proteins; PTM, posttranslational modification. (Adapted from You and Jones 2012.)

GENETIC AND EPIGENTIC ALTERATIONS IN HUMAN CANCER

Classic genetic basis of cancer

Knudsen hypothesis: Two-hit or multiple hit hypothesis:

Both alleles of a gene or multiple gene mutations result in cancer formation

The extended (epi)-genetic basis of cancer

- Mutations in single alleles
- Loss of parts or entire genes
- Loss of one allele; second allele silenced by epigenetic alterations
- Epigenetic silencing of both alleles

Epigenetic alterations:

- **Mutations in epigenetic writers/chromatin components**
- **Reduced expression of epigenetic regulators**
- **Increased expression of epigenetic regulators**

The consequence of mutations in epigenetic regulators are difficult to assess

- **Multiple direct and indirect effects on gene expression (silencing of DNA repair gene MLH1 mediates further mutations, etc)**
- **Important to understand immortalization and transformation**
- **Targets for chemotherapeutic approaches –> effect on multiple genes**

The epigenetic machinery

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CHROMATIN STRUCTURE IS DIFFERENT IN CANCER CELLS

Staining of chromatin by hematoxylin –> standard procedure in pathology

Figure 2. Chromatin structural changes in cancer cells. These two photomicrographs were taken from a patient with a squamous cell carcinoma of the skin. The left panel shows normal epidermal cells within one millimeter of the contiguous tumor shown at the same magnification on the right. The chromatin, which stains purple as a result of its affinity to hematoxylin, appears much more coarse and granular in the cancer cells than in normal epidermis. Such changes in the staining characteristics of chromatin are used by pathologists as diagnostic criteria for cancer.

Definition of epigenetic determinants in human cancer

Classic visual inspection Detailed molecular analysis

Genome-wide epigenetic mapping Chromosome conformation capture RNA seq (lncRNA, siRNA, miRNA, etc) DNA methyaltion profiling Genome sequencing Proteiomics Metabolomics

Changes in chromatin conformation High resolution information on

Epigenetic alterations on the single nucleotide level

Detailed understanding of pathological processes in human cancer

Alteration of epigenetic modifiers in human cancer

Histone modifying complexes

Histone modifying complexes

MDS, myelodysplastic syndromes; AML, acute myeloid leukemia; ICF, immunodeficiency, centromere instability, and
facial anomalies; SNPs, single-nucleotide polymorphisms; TCC, transitional cell carcinoma; HCC, hepatocellular RCCC, renal clear cell carcinoma; TET, ten-eleven translocation; NuRD, nucleosome remodeling and deacetylation.

1. Alteration of DNA methylation in human cancer

Figure 3. Epigenetic alterations involving DNA methylation can lead to cancer by various mechanisms. Loss of DNA cytosine methylation (white hexagons) illustrated in the hypo column results in genome instability. Focal hypermethylation (pink hexagons) at gene promoters shown in the hyper column causes heritable silencing and, therefore, inactivation of tumor suppressors and other genes. Additionally, methylated CpG sites (pink hexagons) are prone to mutation: They are hot spots for C to T transition mutations caused by spontaneous hydrolytic deamination; or methylation of CpG sites can increase the binding of some chemical carcinogens to DNA; and it increases the rate of UV-induced mutations.

Normal tissue: 80% of CpG Cancer: 40-60% of CpG

Global DNA methylation is lost from large blocks (28kb –

Local hypermethylation at specific locations such as tumor suppressors

A. HYPOMETHYLATION

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Hypomethylation distinguishes genes of some human cancers from their normal counterparts

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It has been suggested that cancer represents an alteration in DNA, heritable by progeny cells, that leads to abnormally regulated expression of normal cellular genes; DNA alterations such as mutations^{1,2}, rearrangements³⁻⁵ and changes in methylation⁶⁻⁸ have been proposed to have such a role. Because of increasing evidence that DNA methylation is important in gene expression (for review see refs 7, 9-11), several investigators have studied DNA methylation in animal tumours, transformed cells and leukaemia cells in culture^{8,12-30}. The results of these studies have varied; depending on the techniques and systems used, an increase $12-19$, decrease $20-24$, or no change $25-29$ in the degree of methylation has been reported. To our knowledge, however, primary human tumour tissues have not been used in such studies. We have now examined DNA methylation in human cancer with three considerations in mind: (1) the methylation pattern of specific genes, rather than total levels of methylation, was determined: (2) human cancers and adiacent analogous normal tissues, unconditioned by culture media, were analysed: and (3) the cancers were taken from patients who had received neither radiation nor chemotherapy. In four of five patients studied, representing two histological types of cancer, substantial hypomethylation was found in genes of cancer cells compared with their normal counterparts. This hypomethylation was progressive in a metastasis from one of the patients.

Landmark paper introducing differential DNA methylation in cancer research

ETTERSTONATURE

Prepare DNA from healthy and tumor tissue Digest with methylation sensitive restriction enzyme (does not cut metylated DNA) Probe southern blot with probes cDNA probes of different genes

Cancer tissue: smaller fragments à **methylation lost**

Fig. 1 Methylation pattern of adenocarcinoma of the colon and normal colonic mucosa from patient 1. Lanes C and N were prepared from DNA digests of the colon carcinoma and normal colonic mucosa. respectively. a HpaII digest, HGH probe: b. Hhal digest, HGH probe; c. MspI digest, HGH probe; d. HpaII digest, y-globin probe; e, Hhal digest, γ -globin probe; f, Hoall digest, α -globin probe: g. Hhal digest, α -globin probe. The asterisks on the left of $a - g$ indicate molecular weight markers (HindIII-digested bacteriophage λ DNA, of sizes 9,500, 6,700, 4,400, 2,000 and 570 bp from top to bottom, respectively).

Methods: Tissues were frozen in liquid nitrogen, then pulverized and the DNA extracted essentially as described else-

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where⁵¹. The DNA was cleaved with 50-fold excess of restriction endonuclease, as assessed by the amount of restriction endonuclease required to digest pBR322 DNA included in a matched duplicate digest of the human DNA. DNA digests (5 μ g per lane as assessed by a fluorimetric assay⁵²) were electrophoresed on 0.8–1.5% agarose gels at 70 V for 4 h, then transferred to nitrocellulose by the modification of Southern's³⁹ procedure described by Wahl et al.⁴⁰, pBR322 plasmids containing cDNA inserts of human growth hormone (chGH800/pBR322; ref. 41), human γ -globin (JW101; ref. 42) and human α -globin (JW101; ref. 42), were grown in L-broth, and plasmid DNA was isolated by standard techniques^{53,54}. The inserts from these plasmids were purified and the DNA labelled with ³²P-dCTP to a specific activity of 10⁹ d.p.m. μ g⁻¹ using a technique described elsewhere⁵⁵. The blots obtained by this method were identical to those obtained when probes were labelled by nick-translation, but exposure times were significantly reduced. The probes were hybridized to the filters for 36–60 h and then washed according to a protocol supplied by K. Peden⁵⁶. The autoradiographs shown were exposed for 2-4 days using pre-exposed Kodak XAR-5 film with DuPont Lightning Plus intensifying screens⁵⁷.

A. HYPOMETHYLATION –> alteration in gene expression

Figure 5

High variability of gene expression associated with blocks

(a) An example of hypervariably expressed genes contained within a block; note genes MMP7, MMP10, and MMP3 highlighted in red. Methylation values for cancer samples (red) and normal samples (blue) with hypomethylated block locations highlighted (pink shading) are plotted against genomic location. Grey bars are as in Fig. 2. (b) Standardized log expression values for 26 hypervariable genes in cancer located within hypomethylated block regions (normal samples in blue, cancer samples in red). Standardization was performed using the gene expression barcode. Genes with standardized expression values below 2.54, or the 99.5th percentile of a normal distribution (horizontal dashed line) are determined to be silenced by the barcode method $\frac{26}{1}$. Vertical dashed lines separate the values for the different genes. Note there is consistent expression silencing in normal samples compared to hypervariable expression in cancer samples. A similar plot drawn from an alternative GEO dataset is shown in Supplementary Figure 18.

Stochastic methylation variation of differentially methylated regions, distinguishing cancer from normal tissue, in colon, lung, breast, thyroid and Wilms' tumors.

Whole-genome bisulfite sequencing shows these variable DMRs are related to loss of sharply delimited methylation boundaries at CpG islands.

Furthermore, we find hypomethylation of discrete blocks encompassing half the genome, with extreme gene expression variability.

«We suggest a model for cancer involving loss of epigenetic stability of well-defined genomic domains that underlies increased methylation variability in cancer that may contribute to tumor heterogeneity.»

Hansen KD et al. Nat Genet. 2011. Increased methylation variation in epigenetic domains across cancer types.

A. HYPOMETHYLATION –> genomic instability

Mouse model for cancer:

Hypermorph of DNMT1 (DNMT^{chip}) reduced DNA methylation DNMT1-/Chip mice are viable; 80% develop aggressive lymphoma

CGH array shows genomic instability:

The increased fluorescence ratios observed for chromosomes 14 and 15 are consistent with singlecopy whole-chromosome gains throughout the tumor

(D) Array comparative genome hybridization (CGH) analyses of three Dnmt1chip/–tumors, showing clear single-copy, whole-chromosome gain of chromosome 15 (x, y, and z), whole-chromosome gains of 14and loss on distal 12 (x), andgains of chromosome 14and proximal 9 (y). The X gain (x) reflects a sex difference between tumor and control. Array CGH was performed as in (26). Fluorescence ratios (average of quadruplicate measurements) for each bacterial artificial chromosome are plotted as a function of genome location based onthe February 2002 freeze of the assembled mouse genome sequence (http://genome.ucsc.edu). Verticallines delimit chromosome boundaries.

Gaudet et al Science 2003

Comparative genomic hybridization (CGH)

Technological approaches to array CGH

Array CGH has been implemented using a wide variety of techniques. Therefore, some of the advantages and limitations of array CGH are dependent on the technique chosen. The initial approaches used arrays produced from large insert genomic DNA clones, such as BACs. The use of BACs provides sufficient intense signals to detect single-copy changes and to locate aberration boundaries accurately. However, initial DNA yields of isolated BAC clones are low and DNA amplification techniques are necessary. These techniques include ligation-mediated polymerase chain reaction (PCR), degenerate primer PCR using one or several sets of primers, and rolling circle amplification.[20] Arrays can also be constructed using cDNA. These arrays currently yield a high spatial resolution, but the number of cDNAs is limited by the genes that are encoded on the chromosomes, and their sensitivity is low due to cross-hybridization.[15] This results in the inability to detect single copy changes on a genome wide scale.[21] The latest approach is spotting the arrays with short oligonucleotides. The amount of oligos is almost infinite, and the processing is rapid, cost-effective, and easy. Although oligonucleotides do not have the sensitivity to detect single copy changes, averaging of ratios from oligos that map next to each other on the chromosome can compensate for the reduced sensitivity.[22] It is also possible to use arrays which have overlapping probes so that specific breakpoints may be uncovered. **ACGH profile of the IMR32 neuroblastoma cell line**

Methodology

Array CGH is based on the same principle as conventional CGH. In both techniques, DNA from a reference (or control) sample and DNA from a test (or patient) sample are differentially labelled with two different fluorophores and used as probes that are cohybridized competitively onto nucleic acid targets. (((In conventional CGH, the target is a reference metaphase spread))). In array CGH, these targets can be genomic fragments cloned in a variety of vectors (such as BACs or plasmids), cDNAs, or oligonucleotides.[18]Figure 2.[15] is a schematic overview of the array CGH technique. DNA from the sample to be tested is labeled with a red fluorophore (Cyanine 5) and a reference DNA sample is labeled with green fluorophore (Cyanine 3). Equal quantities of the two DNA samples are mixed and cohybridized to a DNA microarray of several thousand evenly spaced cloned DNA fragments or oligonucleotides, which have been spotted in triplicate on the array. After hybridization, digital imaging systems are used to capture and quantify the relative fluorescence intensities of each of the hybridized fluorophores.[18] The resulting ratio of the fluorescence intensities is proportional to the ratio of the copy numbers of DNA sequences in the test and reference genomes. If the intensities of the flurochromes are equal on one probe, this region of the patient's genome is interpreted as having equal quantity of DNA in the test and reference samples; if there is an altered Cy3:Cy5 ratio this indicates a loss or a gain of the patient DNA at that specific genomic region.[19]

A. HYPOMETHYLATION –> genomic instability, altered gene expression

Coloretal cancer + liver metastasis:

Reduced DNA methylation in LINE1 retrotransposons

LINE1 elements contain normal promoter but also an additional promoters that can activate downstream genes (i.e. MET oncogene)

LINE1 elements are normally epigenetically silenced by DNA methylation

Metastasis \rightarrow loss of LINE1 methylation \rightarrow increases expression Of MET onocgene.

A consolidated model for the activation mechanism of the MET proto-oncogene by LINE-1 hypomethylation in colorectal cancer (CRC) metastasis. In this model, hypomethylation of CpG sites is characterised by enhanced conversion of 5-mc (5-methylcytosine) to 5-c (5-cytosine), with an intermediary generation of 5-hmc (5-hydroylated methylcytosine). Enhanced CpG demethylation with consequent accumulation of 5-hmc prevents the binding of various DNA methyltransferases (DNMTs) and methyl binding proteins (MBPs) to CpG sites, triggering a cascade of events that favour DNA hypomethylation. Occurrence of DNA hypomethylation events in the context of LINE-1 elements can potentially permit activation of certain oncogenes, including the MET proto-oncogene which may result in accelerated CRC metastasis.

A. HYPOMETHYLATION –> genomic instability

Prepare DNA from healthy and tumor tissue and sperm Digest with methylation sensitive restriction enzyme (does not cut metylated DNA) Probe southern blot with probes cDNA probes of different genes Cancer tissue: smaller fragments \rightarrow **methylation lost**

FIGURE 4 - Analysis of methylation of Chr1 Sat2 DNA sequences in malignant breast tissue samples. Breast adenocarcinoma DNA samples, in the first 12 lanes, and sperm and lung DNA samples, in the last 2 lanes, were analyzed for Chr1 Sat2 DNA hypomethylation. The 5 lowest bands in the sperm DNA samples, which were also seen in the breast cancer samples showing considerable hypomethylation $(21, 25, 73, 01, 46)$, had DNA fragments of about 0.8, 1.3, 1.8, 2.2 and 2.7 kb. In addition, a hybridizing fragment migrating at the position for 4.0-kb DNA is seen above the 2.7-kb fragment in samples 21, 25, O2 and 46.

Narayan et al 1998

1. Alteration of DNA methylation in human cancer

Figure 3. Epigenetic alterations involving DNA methylation can lead to cancer by various mechanisms. Loss of DNA cytosine methylation (white hexagons) illustrated in the hypo column results in genome instability. Focal hypermethylation (pink hexagons) at gene promoters shown in the hyper column causes heritable silencing and, therefore, inactivation of tumor suppressors and other genes. Additionally, methylated CpG sites (pink hexagons) are prone to mutation: They are hot spots for C to T transition mutations caused by spontaneous hydrolytic deamination; or methylation of CpG sites can increase the binding of some chemical carcinogens to DNA; and it increases the rate of UV-induced mutations.

Normal tissue: 80% of CpG Cancer: 40-60% of CpG

Global DNA methylation is lost from large blocks (28kb –

Local hypermethylation at specific locations such as tumor suppressors

B. UV damage at 5mC

Figure 7-F-4. Pyrimidine dimer induced by UV light. This figure uses thymine as an example. Cytosine may form a similar dimer.

B. UV damage at 5mC

Formation of T5mC dimeric photoproducts. CPD, cyclobutane pyrimidine dimers (c,s stereo isomers); 64PP, pyrimidine (6-4) pyrimidone photoproducts, sugars are represented by triangles, phosphate groups by a circle.

B. INCREASED MUTATION RATE RELATED TO 5mC

5mC transition mutation to T is most frequent type of transition mutation (for example 50% of p53 mutations in colorectal cancer occur at sites of DNA methylation)

5mC →T is frequent in exons of tumorsuppressor gene bodies

5mC → promotes the formation of adducts with benzo(a)pyrene diol epoxide à **increased mutation rate in smokers**

B. INCREASED MUTATION RATE RELATED TO 5mC

5mC à**T is frequent in exons of the tumorsuppressor gene p53 in colorectal cancer and lung cancer**

B. INCREASED MUTATION RATE RELATED TO 5mC

5mC à**promotes the formation of adducts with benzo(a)pyrene diol epoxide** à **increased mutation rate in smokers**

1. Alteration of DNA methylation in human cancer

Figure 3. Epigenetic alterations involving DNA methylation can lead to cancer by various mechanisms. Loss of DNA cytosine methylation (white hexagons) illustrated in the hypo column results in genome instability. Focal hypermethylation (pink hexagons) at gene promoters shown in the hyper column causes heritable silencing and, therefore, inactivation of tumor suppressors and other genes. Additionally, methylated CpG sites (pink hexagons) are prone to mutation: They are hot spots for C to T transition mutations caused by spontaneous hydrolytic deamination; or methylation of CpG sites can increase the binding of some chemical carcinogens to DNA; and it increases the rate of UV-induced mutations.

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Local hypermethylation at specific locations such as tumor suppressors

C. HYPERMETHYLATION

In human cancer, DNA methylation occurs at CpG islands that are located in and around the transcriptional start site of abnormally silenced genes (in cancer cells) Typically 5-10% of these CpG island promoters are methylated in cancer.

In an individual tumor hundreds of genes can be subjected to this type of epigenetic alteration.

Apparently, the number of genes with hypermethylated promoters outnumbers the number of genes mutated in the same cancer.

$→$ **Gene silencing of tumorsuppressors**

Figure 4. Chromatin structural changes in cancer cells. (A) In a typical cell, a CpG-island-containing active gene can be recognized by virtue of a nucleosome-depleted promoter, absence of promoter DNA methylation, but marked by H3K4me3 surrounding the promoter and histone acetylation along the locus. Gene body CpG methylation often can be observed. Nongenic regions flanking an active gene are frequently marked by repressive epigenetic marks, such as H3K9me3 and 5mC. (B) The cancer epigenome is characterized by simultaneous global losses in DNA methylation (gray shading), interspersed with silenced genes that have abnormal gains of DNA methylation and repressive histone modifications in CpG island promoter regions. These silenced genes may be hypomethylated in their gene body, similar to surrounding chromatin. The hypomethylated regions can have an abnormally open nucleosome configuration and acetylated histone lysines. Conversely, abnormal DNA hypermethylation in promoter CpG islands of silenced genes is associated with nucleosomes positioned over the transcription start sites.

2. HYPERMETHYLATION OF GENE PROMOTERS of TSGs (tumor suppressor genes) IN CANCER

NORMAL DEVELOPMENT: Promoters of 60% of genes contain CpG islands \rightarrow vast majority is never DNA methylated during development

CANCER: 5–10% of the thousands of promoter CpG islands that never normally contain DNA methylation from embryonic development onwards show increased methylation in human cancer

Silencing of tumor suppressors ("driver") but also potential "passengers"

^a A classic tumor-suppressor gene (TSG) is known to be mutated in the germline of families with hereditary cancer syndromes.

2. HYPERMETHYLATION OF GENE PROMOTERS OF TSGs: MLH1 in colorectal cancer

DNA missmatch repair

MLH1: DNA mismatch repair pathway

- Defect in DNA damage repair
- Accumulation of mutation
- Genomic instability (microsatellite instability)
- Driver mutation for colon cancer

A subclass of colorectal cancer is characterized by a hypermethylation of CpG islands at the MLH1 promoter

2. HYPERMETHYLATION OF GENE PROMOTERS OF TSGs: MLH1 in colorectal cancer

MLH1 is not frequently mutated

Somatic Mutation Frequencies in Colorectal Cancer

Pathways to tumorigenesis in Colorectal cancer

Lynch syndrome (HNPCC or hereditary nonpolyposis colorectal cancer) is an autosomal dominant genetic condition that has a high risk of colon cancer[1] as well as other cancers including endometrial cancer (second most common), ovary, stomach, small intestine, hepatobiliary tract, upper urinary tract, brain, and skin. The increased risk for these cancers is due to inherited mutations that impair DNA mismatch repair. **It is a type of cancer syndrome**. MUTATIONS in MLH1, MSH2, MSH6, PMS2

Serrated adenocarcinoma is a recently described subset of colorectal cancer (CRC), which account for about 10% of all CRCs and follows an alternative pathway in which serrated polyps replace the traditional adenoma as the precursor lesion to CRC. Serrated polyps form a heterogeneous group of colorectal lesions that includes hyperplastic polyps (HPs), sessile serrated adenoma (SSA), traditional serrated adenoma (TSA) and mixed polyps.

EPIGENETIC DRUGS: DNMT INHIBITORS à **loss of DNA methylation**

Azacitidine is a **chemical analogue of the nucleoside cytidine**, which is present in DNA and RNA. It is thought to have antineoplastic activity via two mechanisms – at low doses, by **inhibiting of DNA methyltransferase, causing hypomethylation of DNA, and at high doses, by its direct cytotoxicity to abnormal hematopoietic cells** in the bone marrow through its incorporation into DNA and RNA, resulting in cell death. Azacitidine is a ribonucleoside, so it is incorporated into RNA to a larger extent than into DNA. In contrast, **decitabine (5-aza-2'-deoxycytidine) is a deoxyribonucleoside, so it can only incorporate into DNA**. Azacitidine's incorporation into RNA leads to the dissembly of polyribosomes, defective methylation and acceptor function of transfer RNA, and inhibition of the production of proteins**. Its incorporation into DNA leads to covalent binding with DNA methyltransferases, which prevents DNA synthesis and subsequent leads to cytotoxicity.**

After azanucleosides such as **azacitidine have been metabolized to 5-aza-2ʹ-deoxycytidinetriphosphate** (aka, decitabine-triphosphate), they can be incorporated into DNA and azacytosine can be **substituted for cytosine**. Azacytosine-guanine dinucleotides are recognized as substrate by the DNA methyltransferases, which catalyze the methylation reaction by a nucleophilic attack. This **results in a covalent bond between the carbon-6 atom of the cytosine ring and the enzyme. The bond is normally resolved by beta-elimination through the carbon-5 atom, but this latter reaction does not occur with azacytosine because its carbon-5 is substituted by nitrogen, leaving the enzyme remains covalently bound to DNA and blocking its** DNA methyltransferase function.

In addition, the **covalent protein adduction also compromises the functionality of DNA and triggers DNA damage signaling, resulting in the degradation of trapped DNA methyltransferases. As a consequence, methylation marks become lost during DNA replication.**

2. HYPERMETHYLATION OF GENE PROMOTERS OF TSGs: MLH1 in colorectal cancer

Reversibly of DNA damage repair defect caused by MLH1 silencing

Treatment of colon cancer cell line (with silenced MLH1) with 5- Aza-2ʹ-deoxycytidine

Re-expression of MLH1 and rescue of DNA damage repair

2. HYPERMETHYLATION OF GENE PROMOTERS OF TSGs: WHAT ARE THE IMPORTANT GENES?

^a A classic tumor-suppressor gene (TSG) is known to be mutated in the germline of families with hereditary cancer syndromes.

Drivers: genes that are causal for tumor formation

Passengers: genes that are not causal for tumor formation, but support the tumorformation and progression

2. HYPERMETHYLATION OF GENE PROMOTERS OF TSGs: WHAT ARE THE IMPORTANT GENES?

Approaches to identify relevance of potential TSG silenced in human cancer

Table 5. Steps in documenting the importance of a hypermethylated gene for tumorigenesis

- 1. Document CpG island promoter methylation and correlate with transcriptional silencing of the gene and ability to reverse the silencing with demethylating drugs in culture.
- 2. Document correlation of promoter hypermethylation with specificity for this change in tumor cells (cell culture and primary tumors) versus normal cell counterparts and incidence for the hypermethylation change in primary tumors.
- 3. Document the position of the hypermethylation change for tumor progression of given cancer Fine mapping of critical position types.
- 4. Document the potential significance for the gene silencing in tumorigenesis through gene reinsertion studies in culture and effects on soft agar cloning, growth of tumor cells in nude mouse explants, etc.
- 5. Establish function of the protein encoded by the silenced gene, through either known characteristics of the gene or testing for activity of recognized protein motifs in culture systems, etc.
- 6. Document tumor-suppressor activity and functions of the gene for cell renewal, etc., especially for totally unknown genes, through mouse knockout studies.

First observation in a caner cell line in vitro

 \rightarrow Bisulfite sequencing, gene expression, 5-aza-2'-deoxycytidine treatment

Validation of first observation in different type of cancer cells (primary or cancer cell lines and cancer tissue and normal tissue)

 \rightarrow Bisulfite sequencing, gene expression, 5-aza-2'-deoxycytidine treatment, large datasets

- \rightarrow More detailled methyaltion analysis using cancer progression models (cancer specimen, in vitro models for cancer progression (EMT, oncogenes…)
- Functional experiments
- \rightarrow Reconstitute the expression of the silenced gene (cell models, mouse models), asses key BIOLOGICAL features of cancer cells (proliferation, apoptosis, etc…)
- Functional experiments
- \rightarrow Identify the biological pathways the gene is involved

Functional experiments

 \rightarrow More complex model systems

Clinical data

 \rightarrow Patient survival according to eventual mutations in the gene of interest, gene expression levels….

Figure 5. The position of abnormal DNA methylation in tumor progression. This is depicted in the classic model (Kinzler and Vogelstein 1997) for genetic alterations during the evolution of colon cancer. Altered DNA methylation is shown to occur from very early on in tumorigenesis (red arrow), as discussed in the text, during the conversion of normal to hyperplastic epithelium, accruing during the progression from noninvasive to invasive and, ultimately, metastatic tumors. This places it in a strategic position for channeling stem cells into abnormal clonal expansion (illustrated in Fig. 6) by cooperating with key genetic alterations. These epigenetic abnormalities also have connotations for cancer treatment and markers of prognosis.

Experiments have specifically shown that this leads to the buildup of repressive histone marks in gene promoter regions and, subsequently, DNA hyper-methylation in what may constitute a molecular progression

Engineering the mutations/epigenetic alterations into mice or cells, in vitro, appears to trap stem progenitor cells in states of abnormal self-renewal and/or diminishes their capacity for lineage commitment and differentiation.

Then, inducing DNA de-methylation can partially restore the capacity of cells with the mutations to respond to differentiation cues

Figure 6. Epigenetic gene-silencing events and tumorigenesis. The earliest steps in tumorigenesis are depicted as abnormal clonal expansion, which evolves during the stress of cell renewal. This is caused by factors, such as aging and chronic injury from, for example, inflammation. These cell clones are those at risk of subsequent genetic and epigenetic events that could drive tumor progression. Abnormal epigenetic events, such as the aberrant gene silencing focused on in this chapter, could be the earliest heritable causes, in many instances, for a potential role in inducing the abnormal clonal expansion from within stem/ progenitor cell compartments in a renewing adult cell system. The gene silencing is triggered by chromatin modifications that repress transcription, and the DNA hypermethylation of this chromatin serves as the tight lock to stabilize the heritable silencing. The gene silencing, in turn, disrupts normal homeostasis, preventing stem and progenitor cells from moving properly along the differentiation pathway for a given epithelial cell system (blue arrow) and channels them into the abnormal clonal expansion (red arrow).

Loss of imprinting (LOI): EXAMPLE H19-IGF2 locus

Proposed model of imprinting at the H19–IGF2 locus: the H19–IGF2 locus consists of a paternally methylated ICR (Imprinting Control Region). On the maternal allele, this ICR is unmethylated and is bound by the insulator protein CTCF that prevents access of the IGF2 promoter to

downstream enhancers.

H19–IGF2 **LOCUS**

The *H19–IGF2* locus resides adjacent to the *KCNQ1* region on human Chr. 1p15.5. The region consists of the **paternally expressed** *insulin-like growth factor 2* **(***IGF2***) gene** and **maternally expressed** *H19* **lncRNA**. The *IGF2* protein functions as a growth factor essential for embryonic development, whereas *H19* may function as a tumor suppressor.

Imprinting at this locus is maintained by an ICR, which is selectively methylated on the paternal allele. The insulator protein, CCCTC- binding factor (*CTCF*), binds to critical regulatory regions in the unmethylated ICR (Imprinting control region) on the maternal allele, thus blocking access of downstream enhancers to the *IGF2* promoter.

Loss of methylation at the *H19/IGF2* **ICR results in short body length and low birth weight, both in rodent models as well as in humans**

IGF2 knock-out allele was inherited from the father of the mouse

Loss of imprinting (LOI): EXAMPLE H19-IGF2 locus

methylation of ICR on maternal allele; increased IGF2 expression; increased proliferation potential

In mouse loss of IGF2 imprinting may be sufficient to initiate tumorigenesis

IDH1 mutations in glioblastoma lead to epigenetic deregulation in oncogenesis

Classic oncogenic driver mutations in glioblastoma IDH1 mutations are highly frequent in glioblastoma

CNS, central nervous system; WHO, World Health Organisation; AML, acute myeloid leukemia; p-2-HGA, p-2-hydroxyglutaric aciduria; MDS, myelodysplastic syndrome; MPN, myeloproliferative neoplasms.

IDH1 mutations in glioblastoma lead to epigenetic deregulation in oncogenesis

IDH1 mutation results result the abberrant production of high levels of **2-hydroxyglutarate (2-HG),** a possible oncometabolite that is thought to influence a range of cellular programs involved in epigenetic control and various processes leading to tumor development.

IDH1 mutations are heterozygous, typically involving an amino acid substitution in the active site of the enzyme in codon 132. The mutation results in a loss of normal enzymatic function and the abnormal production of 2 hydroxyglutarate (2-HG)

This abnormal **2-hydroxy-glutarate** metabolite, which increases to mM levels in cells with the IDH mutations, constituting a biomarker in and of itself, competes with the necessary **α-ketoglutarate metabolite needed by the TET and lysine (K) demethylase (KDM) enzymes**, which regulate either chromatin demethylase function or levels of DNA methylation.

α-ketoglutarate metabolite reduced because processed to R-2HG

Figure 2. Multiple aKG-dependent dioxygenases family are dependent on aKG for their enzymatic activities to modify gene expression. When associated with aKG, r spure at manple announcement and spendes animy are cependent on announcement accrete and through years explosive manner based and the based of the based on the based on oxidized to form derivatives 5-formylcytosine and 5-carboxylcytosine. While this activity requires a KG, TET methylcytosine hydroxylation is strongly inhibited by R-2HG
production following IDH1/2 mutation. Similarly, the Man and the engine manner in the manner of the manner of the manner of the manner of the and the manner of the manner of the manner of the state of the manner of the www.musumer.com and the margin of the margin of the margin of the margin of the consequences of reduced FTO activity in mutant IDH musumer weak of the consequences of reduced FTO activity in mutant IDH musumer in the conse ally data DNA lesions containing 1-methyledenine and 3-methylcytosine by oxidative demethylation. R-2HG production inhibits ALKBH2/3 activity, thus interfering with normal DNA repair processes. Inhibition of KDM4A and KDM4B by R-2HG leads to DNA repair defects by impairing homologous recombination (HR). In addition. IDH numa unique accumulation of the repressive H3K9ms3 histone mark at the promoter of gene encoding the DNA damage response kinase ATM, thus decreasing the
Invisions cause accumulation of the repressive H3K9ms3 histone mark a is the endogenous negative regulator of mTOR, by inhibiting its polyubiquitylation, thus decreasing both mTORC1 and mTORC2 activation. In presence of R-2HG,
The endogenous negative regulator of mTOR, by inhibiting its poly DEPTOR is rapidly degraded, resulting in a massive increase in mTORC1/2 activation in a PISK/PTEN-Independent fashion, thus favoring the proliferation and survival of
cancer cells harboring IDH1/2 mutations. In addition, R production by (DH1/2 multations engages a complex network that likely regulates mTCR activity through regulatory feedback loops. [HIF-1 Signaling] Hypoxia-inducible
factor 1a (HIF-1a) is a major regulator of cell prolliera stating and compare interaction of the "unit unit of the compare that statistize the collagen the likely the condigent by sixe of the product of the condigent by the condigent of the condigent of the condigent of the condi maturation.

Table 1. Enzymes Regulated by Mutated IDH1/2 and the Associated Oncometabolite R-2HG

Table 1. Examples of key cellular pathways disrupted in human cancers by genetic and epigenetic mechanisms

TGF- β , transforming growth factor β ; *DAPK*, death-associated protein kinase.

SILENCING OF p16

Table 1. Examples of key cellular pathways disrupted in human cancers by genetic and epigenetic mechanisms

TGF- β , transforming growth factor β ; *DAPK*, death-associated protein kinase.

3. The importance of epigenetic gene regulation in early tumorprogression

SILENCING OF p16

Because of its importance in cell proliferation, the human INK4 gene locus, encoding p16, is a frequent target of inactivation by deletion or aberrant DNA methylation in a wide variety of human cancers

- p16 silencing is an early event in tumorformation
- \rightarrow Increased activity of cyclinD/CDK4
- \rightarrow pRB hyperphoslphorylation
- \rightarrow E2F activates genes required for the induction of S-Phase

SILENCING OF MLH1

Table 1. Examples of key cellular pathways disrupted in human cancers by genetic and epigenetic mechanisms

TGF- β , transforming growth factor β ; *DAPK*, death-associated protein kinase.

Pathways to tumorigenesis in Colorectal cancer

SILENCING OF Chfr

Table 1. Examples of key cellular pathways disrupted in human cancers by genetic and epigenetic mechanisms

TGF- β , transforming growth factor β ; *DAPK*, death-associated protein kinase.

Chfr function

CHFR is an early mitotic checkpoint gene that functions as a key player in controlling chromosomal integrity.

CHFR is expressed in the cytoplasm of all normal tissues and accumulates in the nucleus in response to **microtubule poisoning, carcinogen or radiation damaging stress**.

After localization into the nucleus, CHFR becomes phosphorylated by protein kinase B (PKB/AKT), a member of the PI3K signaling pathway.

Microtubule stress/Radiation leads to an elevation of CHFR expression levels and a mitotic arrest.

 \rightarrow In conditions with increased risk of genomic instability CHFR blocks mitotic progression to allow damage repair

 \rightarrow Absence of CHFR: mitosis precedes, DNA damage will not be repaired \rightarrow accumulation of mutations \rightarrow genomic instability/cancer.

CHFR is a E3 ubiquitin-protein ligase that delays entry into metaphase in response to microtubular stress/radiation by effecting target proteins in a proteosome-dependent and a proteosome-independent manner \rightarrow *maintains genomic stability*

Chfr knock out mouse model Skin Chfr knock out **mouse model Skin Chfr Skin carcinogenesis**

(mice are treated with carcinogen (DMBA = DNA damaging agent, TPA cell cycle stimulator

Figure 3 Loss of Chfr leads to chromosome instability. (a) Metaphase spreads with normal karyotype in wild-type MEFs (left) and aneuploidy in *Chfr^{-/-}* MEFs (right). (b) *Chfr^{-/-}* cells have increased aneuploidy. We counted the chromosome numbers of 100 metaphase spreads per sample from P4 MEFs. (c) Cell cycle profile of unsynchronized MEFs (at P4) was determined by fluorescence-activated cell-sorting analysis. Aneuploidy and polyploidy were observed in Chfr^{-/-} MEFs. (d) Multinucleated cells observed in Chfr^{-/-} MEFs. We examined 2,500 cells for each genotype by DAPI staining and fluorescence microscopy. The percentages of binucleated and multinucleated cells is indicated $(P < 0.001)$. (e.f) Chfr^{-/-} MEFs have errors in mitotic progression. MEFs were infected with retrovirus encoding histone H2B-EGFP. The mitotic progress of Chfr^{+/+} ($n = 40$) and Chfr^{-/-} ($n = 51$) MEFs was followed by time-lapse microscopy. (e) Mean time of each mitotic phase is shown. Prophase and anaphase were prolonged in Chfr^{-/-} MEFs compared with wild-type MEFs (* $P < 0.005$). (f) Mitotic errors were recorded. Differential interference contrast images are also included.

Chfr is frequently inactivated in human cancer

CHFR inactivation in multiple cancers

CHFR is more frequently inactivated in cancer than all other mitotic checkpoint control genes together.

Absence of CHFR in these cell lines resulted in a **high mitotic index** when exposed to microtubule stress compared to wild-type cancer cells, which was restored by reintroduction of functional CHFR.

In a breast cancer cell line model, **decreased CHFR expression resulted in an accelerated growth rate, enhanced invasiveness and amplified colony formation.**

In order to study the physiological role of CHFR and its function in tumorigenesis, Chfr knockout mice have been generated.

Chfr knockout mice develop invasive lymphomas and solid tumors (lung, liver, gastrointestinal) after 40 weeks and have an increased susceptibility to chemical carcinogenesis.

Embryonic fibroblasts from Chfr-deficient mice show substantial aneuploidy and polyploidy.

IHC, immunohistochemistry; (MS)-MLPA, (methylation-specific) multiplex ligation-dependent probe amplification; MSP, methylation-specific PCR; COBRA, combined bisulfite restriction analys

PML – RAR fusion and aberrant use of HDAC

Acute promyelocytic leukemia (APML, APL) is the M3 subtype of acute myelogenous leukemia (AML), a cancer of the white blood cells. In APL, there is an abnormal accumulation of immature granulocytes called promyelocytes.

The disease is characterized by a chromosomal translocation involving the retinoic acid receptor alpha (RARα or RARA) gene and is distinguished from other forms of AML by its responsiveness to all-trans retinoic acid (ATRA; also known as tretinoin) therapy.

Note: retinoic acid receptor activation by retinoic acid drives differentiation

Acute promyelocytic leukemia

PML – RAR fusion and aberrant use of HDAC

PML–RAR acts as an HDAC-dependent inhibitor of pathways regulated by wild-type PML and RAR.

PML–RAR acts as a bifunctional protein: (i) it inhibits the transcription of RA target genes in the presence of physiological concentrations of ligand, thus blocking myeloid differentiation; and (ii) it inhibits p53 function (in a manner requiring wild-type PML), allowing enhanced survival of APL blasts.

Both differentiation block and enhanced survival are mediated by recruitment of HDAC by the fusion protein.

Acute promyelocytic leukemia

PML – RAR fusion and aberrant use of HDAC

It is apparent that the multisubunit complexes that are responsible for covalently modifying histones and DNA during chromatin remodelling can contribute to cancer. Histone acetyltransferase (HAT) complexes acetylate histones (leading to relaxation of chromatin structure) and, typically, increase transcription of target genes. CREB-binding protein, which possesses intrinsic HAT activity, is fused — by chromosomal translocation — to the product of the mixed-lineage leukaemia gene (*MLL*) in aggressive paediatric leukaemias and to the histone acetyltransferases MOZ and MORF in other leukaemias.

Histone deacetylase (HDAC) complexes have also been implicated in oncogenesis. In acute promyelocytic leukaemia, a recurrent chromosomal translocation results in fusion of the promyelocytic leukaemia gene product (PML), which directly recruits SIN3A–HDAC, to the retinoic-acid receptor (RAR). During normal development of myeloid cells, the RAR transcriptionally activates target genes that promote differentiation into mature myeloid cells. In acute promyelocytic leukaemia, the PML–RAR fusion protein recruits SIN3A–HDAC to RAR targets (see figure). SIN3A– HDAC deacetylates histones, leading to transcriptional repression and to a block in differentiation that contributes to leukaemic growth.

Treatment of these patients with high-dose retinoic acid (RA) causes dissociation of SIN3A– HDAC and differentiation of immature leukemic blast cells into mature, and benign, myeloid cells.

Acute promyelocytic leukemia

Nature Reviews | Cancer

MLL fusion with other genes:

MLL gene on 11q23 is often rearranged with other partner genes in ALL (Acute lymphoblastic leukemia) and AML (Adult Acute Myeloid Leukemia):

accounting for 8% of pediatric and 10% of adult ALL, 15-20% of pediatric AML and <3% of adult AML

Chromosomal translocations lead to the fusion of 5'-end portion of MLL to one of >60 different partner genes, resulting in the formation of different fusion genes, such as MLL-AF4 (4q21), MLL-AF9 (9p22), MLL-ENL (19p13.3), MLL-AF10 (10p12), MLL-AF6 (6q27), MLL-ELL (19p13.1). All MLL fusion proteins retain N-terminal AT-hooks, DNA methyltransferase homology domain, thus preserving DNA binding activity whereas the **trithorax PHD domains, TAD and SET domains** are always replaced by the partners. In these fusions, the original MLL H3K4 methyltransferase activity is replaced by the partners which play a critical role in MLL oncoproteins

MLL fusion with other genes:

Aberrant expression of genes that promote cancer formation …classic example: MLL-AF9 (9p22)

MLL—AF9 fusion:

Schematic picture showing the tumor promoting effect of MLL translocations via recruitment of DOT1L. In normal cells, the MLL1 protein binds to its target genes and introduces H3K4 methylation via its catalytic SET domain (highlighted in dark blue).

In the tumor cells MLL1 forms fusions with other protein like AF9, which leads to the loss of the SET domain.

Although MLL fusion proteins lose the activity of H3K4 methylation, these fusion proteins gain the activity of H3K79 methylation via recruiting the H3K79 methyltransferase hDOT1L which can cause dysregulation of whole genomic expression and is associated with MLL leukemogenesis. Since hDOT1L plays a key role in the development of MLL leukemia, hDOT1L is an ideal target for MLL leukemia. Several hDOT1L inhibitors are underdeveloped. In particular, EPZ004777, **a special hDOT1L inhibitor, seems to be a promising drug for leukemia with MLL gene translocation**

MLL—AF9 fusion:

Fig. 3

Model for leukemogenesis mediated by aberrant DOT1L H3K79 methylation. (A) MLL gene undergoes chromosomal translocations and fusion with a number of fusion proteins such as AF9, AF10, and hDot1L. This leads to mistargeting of hDot1L to MLL-regulated genes such as Hox genes. Aberrant H3K79 methylation and the subsequent activation of the target genes result in leukemic transformation. (B) Inhibition of hDot1L HMTase activity with small molecules would prevent aberrant H3K79 methylation and inhibition of MLLrelated leukemogenesis.

MLL—AF9 fusion:

A

LSD1: H3K4 de-methylase

Interaction of LSD1 with MLL-AR9 ensures leukemia stem cells features \rightarrow oncogenic function

LSD1 inhibitors work in MLL-AR9 fusions models

6. MAJOR EPIGENETIC DRUGS IN CURRENTLY IN DEVELOPMENT

Figure 5. The development status of epigenetic drugs. KDMi, lysine demethylase inhibitor; KMTi, lysine methyltransferase inhibitor; HATi, histone acetyltransferase inhibitor; HDACi, histone deacetylase inhibitor; BETi, bromodomain and extracarboxy terminal domain inhibitor; DNMTi, DNA methyltransferase inhibitor.

Figure 6. Representative structures of drugs targeting DNA methylation, histone modifications, and histone readers in oncology.

EPIGENETIC DRUGS: DNMT INHIBITORS

Figure 8. Structures of selected epigenetic drugs. Three nucleoside analogs are known that can inhibit DNA methylation after incorporation into DNA. 5-aza-CR (Vidaza) and 5-aza-CdR (decitabine) have been FDA approved for the treatment of the preleukemic disorder, myelodysplasia. Two HDAC inhibitors are also FDA approved for cutaneous Tcell lymphoma and several others are in clinical trials.

EPIGENETIC DRUGS: DNMT INHIBITORS

Azacitidine is a chemical analogue of the nucleoside cytidine, which is present in DNA and RNA. It is thought to have antineoplastic activity via two mechanisms – at low doses, by **inhibiting of DNA methyltransferase, causing hypomethylation of DNA, and at high doses, by its direct cytotoxicity to abnormal hematopoietic cells** in the bone marrow through its incorporation into DNA and RNA, resulting in cell death. Azacitidine is a ribonucleoside, so it is incorporated into RNA to a larger extent than into DNA. In contrast, **decitabine (5-aza-2'-deoxycytidine) is a deoxyribonucleoside, so it can only incorporate into DNA**. Azacitidine's incorporation into RNA leads to the dissembly of polyribosomes, defective methylation and acceptor function of transfer RNA, and inhibition of the production of proteins. Its incorporation into DNA leads to covalent binding with DNA methyltransferases, which prevents DNA synthesis and subsequent leads to cytotoxicity.

After azanucleosides such as **azacitidine have been metabolized to 5-aza-2ʹ-deoxycytidinetriphosphate** (aka, decitabine-triphosphate), they can be incorporated into DNA and azacytosine can be **substituted for cytosine**. Azacytosine-guanine dinucleotides are recognized as substrate by the DNA methyltransferases, which catalyze the methylation reaction by a nucleophilic attack. This **results in a covalent bond between the carbon-6 atom of the cytosine ring and the enzyme. The bond is normally resolved by beta-elimination through the carbon-5 atom, but this latter reaction does not occur with azacytosine because its carbon-5 is substituted by nitrogen, leaving the enzyme remains covalently bound to DNA and blocking its** DNA methyltransferase function.

In addition, the **covalent protein adduction also compromises the functionality of DNA and triggers DNA damage signaling, resulting in the degradation of trapped DNA methyltransferases. As a consequence, methylation marks become lost during DNA replication.**

EPIGENETIC DRUGS: DNMT INHIBITORS

Western blot of DNMT1 expression levels in human leukemia cells (Kasumi-1) and breast cancer cell lines (MCF7) after 72 hr daily treatment of 10 nM (leukemia) DACOGEN.

Special effect in leukemia:

Treatment with DAC for 72hrs \rightarrow loss of DNMT1 is mainainted \rightarrow Memory effect «reprogramming of cancer cell

Effect of azanucleosides:

- de-repression of genes
- Other effects: immuneresponse \rightarrow important anti-tumor effect!

Quantitative real-time PCR analyses of CDKN2B gene expression over time in Kasumi-1 after 72 hr daily treatment of 10 nM DAC. Expression levels are adjusted to GAPDH for each sample and graphed as fold changes relative to mock.

CDKN2B: This gene encodes a cyclin-dependent kinase inhibitor, which forms a complex with CDK4 or CDK6, and prevents the activation of the CDK kinases, thus the encoded protein functions as a cell growth regulator that controls cell cycle G1 progression

Tsai et al 2012 Cancer Cell

EPIGENETIC DRUGS: HDAC INHIBITORS

Two HDAC inhibitors are also FDA approved for cutaneous Tcell lymphoma and several others are in clinical trials. Drugs targeting other epigenetic processes are in earlier stages of clinical development (see also Figs. 5 and 6 of Ch. 35 [Audia and Campbell 2014]).

Vorinostat has been shown to bind to the active site of histone deacetylases and act as a chelator for zinc ions also found in the active site of histone deacetylases. Vorinostat's inhibition of histone deacetylases results in the accumulation of acetylated histones and acetylated proteins, including transcription factors crucial for the expression of genes needed to induce cell differentiation. It acts on class I, II and IV of histone deacetylase.

EPIGENETIC DRUGS: HDAC INHIBITORS

Romidepsin acts as a prodrug with the disulfide bond undergoing reduction within the cell to release a **zinc-binding thiol**. The thiol **binds to a zinc atom in the binding pocket of Zn-dependent histone deacetylase to block its activity**.

Romidepsin (2) is a prodrug, with a disulfide bridge reduced by glutathione upon uptake into the

EPIGENETIC DRUGS: HDAC INHIBITORS

Work well in lymphoma (T cells) ì but low efficiency in solid tumors when given alone.

Tests: HDAC treatment sensitizes tumors to conventional chemotherapy and newer targeted therapies i.e. NSCLC: HDACi (entinostat) + EGFRi i.e. NSCLC: HDACi (vorionostat) + carboplatin and paxitaxel

HDAC inhibiotors increase acetylation levels of genes and cause altered gene expresstion

 \rightarrow Hypothesis: TSG re-expressed (to be validated)

Many pathways altered \rightarrow difficult to identify a DIRECT controbution of HDACi