# **REVIEW**

# **Histone Deacetylase Inhibitors: Inducers of Differentiation or Apoptosis of Transformed Cells**

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Histone deacetylase (HDAC) inhibitors have been shown to be potent inducers of growth arrest, differentiation, and/or apoptotic cell death of transformed cells in vitro and in vivo. One class of HDAC inhibitors, hydroxamic acid-based hybrid polar compounds (HPCs), induce differentiation at micromolar or lower concentrations. Studies (x-ray crystallographic) showed that the catalytic site of HDAC has a tubular structure with a zinc atom at its base and that these HDAC inhibitors, such as suberoylanilide hydroxamic acid and trichostatin A, fit into this structure with the hydroxamic moiety of the inhibitor binding to the zinc. HDAC inhibitors cause acetylated histones to accumulate in both tumor and normal tissues, and this accumulation can be used as a marker of the biologic activity of the HDAC inhibitors. Hydroxamic acid-based HPCs act selectively to inhibit tumor cell growth at levels that have little or no toxicity for normal cells. These compounds also act selectively on gene expression, altering the expression of only about 2% of the genes expressed in cultured tumor cells. In general, chromatin fractions enriched in actively transcribed genes are also enriched in highly acetylated core histones, whereas silent genes are associated with nucleosomes with a low level of acetylation. However, HDACs can also acetylate proteins other than histones in nucleosomes. The role that these other targets play in the induction of cell growth arrest, differentiation, and/or apoptotic cell death has not been determined. Our working hypothesis is that inhibition of HDAC activity leads to the modulation of expression of a specific set of genes that, in turn, result in growth arrest, differentiation, and/or apoptotic cell death. The hydroxamic acid-based HPCs are potentially effective agents for cancer therapy and, possibly, cancer chemoprevention. [J Natl Cancer Inst 2000;92:1210-6]

Neoplastic transformation is characterized by inappropriate cell proliferation and/or altered patterns of cell death. However, neoplastic transformation does not necessarily destroy the potential for expression of differentiated characteristics, including cessation of proliferation under appropriate environmental conditions (1). For example, cells infected with temperaturesensitive transforming viruses (2) can display either normal or transformed properties, depending on the activity of a temperature-sensitive viral protein. Some malignant cells (e.g., from teratocarcinomas, neuroblastomas, or leukemias) can differentiate along apparently normal pathways when placed in a normal embryonic environment (3-7). In addition, various chemical agents [hybrid polar compounds (HPCs) (8–10), retinoids (11– 15), vitamin  $D_3$  (16), and several other agents (17–19)] can induce certain transformed cells in vitro to express differentiated characteristics and stop proliferating.

Histones are part of the core proteins of nucleosomes. Acety-

lation and deacetylation of these proteins play a role in the regulation of gene expression (20). There are two classes of enzymes involved in determining the state of acetylation of histones, histone acetyl transferases (HATs) and histone deacetylases (HDACs). There are several reports (21–24) that altered HAT or HDAC activity is associated with cancers.

During the last decade, a number of HDAC inhibitors have been identified that induce cultured tumor cells to undergo growth arrest, differentiation, and/or apoptotic cell death (25–35). These agents also inhibit the growth of cancer cells in animal models (32,35–40), and several agents, in particular, hydroxamic acid-based HDAC inhibitors, inhibit tumor growth in animals at doses that are apparently nontoxic and appear to be selective.

This review focuses on studies of HDAC inhibitors, especially on the hydroxamic acid-based HPCs. These compounds represent a class of agents that are potentially effective cancer therapies. (Studies were identified for this review by searching the MEDLINE® database for appropriate papers published in the last 10 years and by a review of bibliographies from articles identified through that search. In addition, we include some of our unpublished data.)

# HISTONE ACETYLATION AND DEACETYLATION AND GENE EXPRESSION

#### **Structure of Nucleosomes**

Analyses (x-ray and electron crystallographic) show that nucleosomes contain an average of 150 base pairs of DNA wrapped around the nucleosomal core of histones in 1.75 turns of left-handed superhelical DNA (41–43). Five classes of histones have been identified in chromatin: histones H1, H2A, H2B, H3, and H4. Each nucleosome contains two H2As, two H2Bs, two H3s, and two H4s in the core (Fig. 1). Histone H1 occurs in chromatin in about half the amount of the other types of histones and appears to lie on the outer portion of the nucleosome.

#### Role of Histone Acetylases and Deacetylases

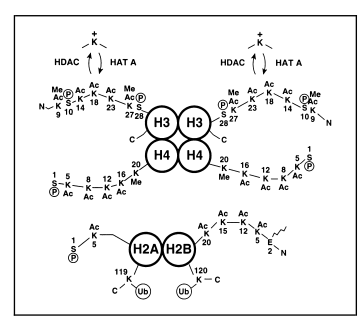
Histones of the nucleosomal core can be acetylated and deacetylated, and the amount of acetylation is controlled by the opposing activities of two types of enzymes, HATs and HDACs. Substrates for these enzymes include  $\varepsilon$ -amino groups of lysine

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**Fig. 1.** Histones in nucleosomes. Lysines (K) in the amino (N)-terminal tails of histones H3, H4, H2A, and H2B are potential acetylation/deacetylation sites for histone acetyltransferase (HAT) and histone deacetylase (HDAC).  $K^+ = positively$  charged lysine, Ub = ubiquitin, P = phosphate, Ac = acetyl, S = serine, E = glutamic acid, and Me = methyl. [Adapted with permission from Davie (44).]

residues located in the amino-terminal tails of the histones. When HDAC removes the acetyl group from histone lysine, it restores a positive charge to the lysine residue condensing the structure of nucleosomes (44).

### **HATs and HDACs**

There are at least four groups of proteins with intrinsic HAT activity (45-50). The first group contains the GCN5 and P/CAF proteins, which are related to yeast HAT GCN5. The second group contains the closely related cyclic adenosine monophosphate response element-binding protein (CBP) and p300, which act as coactivators for a number of transcription factor complexes. The third group contains the TAF250 protein, part of the basic transcription complex TFIID that binds the TATA box. The fourth group contains the SRC-1 and ACTR proteins that are coactivators for ligand-activated nuclear receptors. In addition, there are probably several other proteins with HAT activity, such as BRCA2, that are part of transcription complexes. HATs play a role in activation of gene expression and may also be involved in gene repression, as suggested by the observation in Drosophila that acetylation of the transcription factor T-cell factor by CBP represses transcription (51).

Eight HDACs have been described in mammalian cells (45,52–59). The yeast RPD3 homologues are HDAC1, HDAC2, HDAC3, and HDAC8; the yeast HDA1 homologues are HDAC4 (also known as HDAC-A), HDAC5 (also known as mHDA1), HDAC6 (also known as mHDAC2), and HDAC7.

### **Regulation of Transcription**

Chromatin fractions enriched in actively transcribed genes are also enriched in highly acetylated core histones (20,42,45), whereas silent genes are associated with nucleosomes with a low level of acetylation. Allfrey (60) first suggested that histone acetylation was involved in the regulation of transcription. Dur-

ing the past decade, considerable evidence has accumulated to establish the role of acetylation and deacetylation of histones in the regulation of transcription (20,41–43,45). The following model describes a role for histone acetylation in regulating gene transcription. Nucleosomes containing highly charged hypoacetylated histones bind tightly to the phosphate backbone of DNA, inhibiting transcription, presumably, because transcription factors, regulatory complexes, and RNA polymerase do not have access to the DNA. Acetylation neutralizes the charge of the histones and generates a more open DNA conformation. Transcription factors and the transcription apparatus then have access to the DNA, and expression of the corresponding genes is promoted (Fig. 2).

In addition to HDACs and HATs, other factors are involved in the regulation of chromatin structure, including methyl-CpG-binding protein (61–63) and adenosine triphosphate (ATP)-dependent chromatin-remodeling complexes (64). These chromatin-modifying complexes interact with HAT and HDAC complexes to regulate transcriptional activity of genes [for a recent review of chromatin methylation, see (63); for reviews of the ATP-dependent chromating remodeling complexes, see (64,65).]

HDACs are bound to large protein complexes that regulate gene transcription. Mammalian HDAC1 and HDAC2 are associated with the Sin3 complex that includes NCo-R, SMRT, and several other, as yet, unidentified proteins and appear to repress gene expression by deacetylating core histones. In addition to deacetylation of histones, HDACs may also regulate gene expression by deacetylating transcription factors, such as p53, GATA-1, TFIIE, and TFIIF (66–68). HDACs may also participate in cell cycle regulation. The transcription repression mediated by RB binding to the transcription factor E2F involves recruitment of HDAC1 or HDAC2 by RB (69,70).

# Disruption of HAT and/or HDAC Activity and Development of Cancer

Mutations in the CBP gene, which encodes an HAT, are associated with leukemogenesis and the developmental disorder

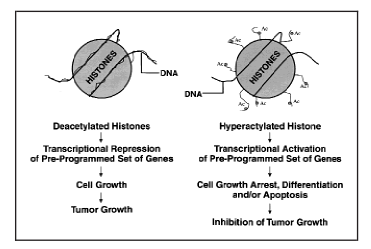


Fig. 2. Proposed mechanism of action of histone deacetylase (HDAC) inhibitors that induce tumor growth arrest, differentiation, and/or apoptotic cell death. With inhibition of HDAC, histones are acetylated (Ac), and the DNA that is tightly wrapped around a deacetylated histone core relaxes. We propose that the accumulation of acetylated histones in nucleosomes leads to expression of specific genes, which, in turn, lead to cell growth arrest, differentiation, and/or apoptotic cell death and, as a consequence, inhibition of tumor growth.

Rubinstein–Taybi syndrome (71). Patients with Rubinstein–Taybi syndrome have a propensity to develop cancer. Microdeletions, translocations, inversions, and various point mutations in the CBP gene have been identified in patients with Rubinstein–Taybi syndrome as well as in patients with some types of colorectal or gastric carcinomas (21). Gene fusions with CBP are associated with several leukemias. In therapy-related acute myeloid and lymphoid leukemias and in myelodysplasia, the CBP gene has been found fused to the MLL gene, and the CBP gene has been found fused in acute myeloid leukemia to the MOZ gene (72,73).

Several leukemogenic transcription factors repress expression of specific genes because of aberrant recruitment of HDACs. This repression of gene expression appears to be an important step in the leukemogenic action of these transcription factors. For example, aberrant recruitment of HDAC activity has been reported in cell lines derived from patients with acute promyelocytic leukemia (APL) (22-24). The oncoprotein encoded by the translocation-generated fusion gene in APL (promyelocytic leukemia [PML]-retinoic acid receptor-α) represses transcription by recruitment of HDAC1. Furthermore, resistance to the differentiating actions of all-trans-retinoic acid in a patient with APL was overcome by cotreatment with an inhibitor of HDAC (74). [In a further study (75), four other patients with APL failed to respond.] HDAC-dependent aberrant transcriptional repression is implicated as the main oncogenic mechanism in specific types of myeloid leukemia and lymphoma. For example, in non-Hodgkin's lymphoma, the transcriptional repressor BCL6 is inappropriately overexpressed within the lymphoid compartment, resulting in aberrant transcriptional repression and lymphoid oncogenic transformation (76). Another example is acute myelogenous leukemia of the M2 subtype associated with the t(8;21) chromosomal translocation involving the AML1 and ETO genes (77). The AML1–ETO fusion protein, unlike the AML-1 protein (a transcriptional activator), is a potent dominant transcriptional repressor. In both of these cases, transcriptional repression appears to be mediated by recruitment of HDAC to the transcriptional repressor complex.

### **HDAC Inhibitors**

Compounds that inhibit HDAC activity are shown in Fig. 3. Several structural classes of HDAC inhibitors have been identified including the following: 1) short-chain fatty acids [e.g., butyrates (28,31)]; 2) hydroxamic acids [e.g., trichostatin A (TSA) (25,26), suberoylanilide hydroxamic acid (SAHA) (34), and oxamflatin (35)]; 3) cyclic tetrapeptides containing a 2-amino-8-oxo-9,10-epoxy-decanoyl (AOE) moiety [e.g., trapoxin A) (27)]; 4) cyclic peptides not containing the AOE moiety [e.g., FR901228 and apicidin (33,78)]; and 5) benzamides [e.g., MS-27-275 (32)]. HDAC inhibitors invariably inhibit proliferation of transformed cells in culture, and a subset has been shown to inhibit tumor growth in animal models (26,32,35-40). The butyrates represent the only class that is approved currently for use in the clinic. The butyrates are not ideal agents because of the high concentrations required (millimolar) to achieve inhibition of HDAC activity and multiple effects on other enzyme systems (28,31). TSA, originally developed as an antifungal agent (25,26,29), is a potent inhibitor of HDAC that is active at nanomolar concentrations. The finding that TSA-resistant cell lines have an altered HDAC is evidence that this enzyme is an important target for TSA. Oxamflatin

Name	Structure	
Butyric Acid	УТОН	
MS-27-275	NH NH2	
SAHA		
Trichostatin A	-N-ОН	
Oxamflatin	Д-он	
Apicidin		
Depsipeptide	NH N	
Depudecin	○ OH	
Trapoxin		

**Fig. 3.** Histone deacetylase inhibitors (*see* text for references to these inhibitors). SAHA = suberoylanilide hydroxamic acid.

(35), a hydroxamic acid-based compound, and the benzamide MS-27-275 (32) inhibit HDAC activity at micromolar concentrations. Apicidin is a fungal metabolite that exhibits potent, broad-spectrum antiprotozoal activity and inhibits HDAC activity at nanomolar concentrations (78). Depsipeptide (FR901228), isolated from *Chromobacterium violaceum* (33), inhibits HDAC activity at micromolar concentrations. Trapoxin (27) and depudecin (30) irreversibly bind to HDAC and inhibit its activity at nanomolar and micromolar concentrations, respectively.

In our laboratory, a series of hydroxamic acid-based HPCs have been synthesized that inhibit HDACs at micromolar concentrations or lower *in vitro* and *in vivo* (34,36,38,79) (Fig. 4), and extensive structure–activity studies have been done with these compounds (34,79). The essential characteristics of hydroxamic acid-based HPCs are a polar site, the hydroxamic group, a six-carbon hydrophobic methylene spacer, a second polar site, and a terminal hydrophobic group (Fig. 4). Substitution of the hydroxamic acid with a carboxylic acid or amide oxime group results in inactive compounds. Modification of the hydroxamic acid, such as introduction of a methyl group on an adjacent carbon or *N*-methylation, results in inactive compounds. The benzene ring in the hydrophobic moiety can be

	Name	Structure	Opt. Conc.	% Diff.		
	SBHA	но Туон	30 μM	90%		
	SAHA	THO H	2.5 μΜ	68%		
	СВНА	HN-OH H	4.0 μΜ	73%		
	Pyroxamide	THE STATE OF THE S	4.0 μΜ	51%		
	Hydrophobic Polar site					

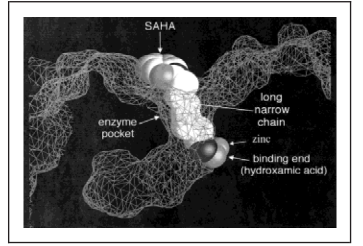
Fig. 4. Hydroxamic acid-based hybrid polar compounds. The optimal concentration to induce murine erythroleukemia cells to differentiate (% Diff) was determined from the percent of differentiated cells [detected as benzidine-stained cells (benzidine binds to the iron-containing heme of hemoglobin); for details of methods, see~(34)]. SBHA = suberic bishydroxamic acid; SAHA = suberoylanilide hydroxamic acid; CBHA = m-carboxy-cinnamic acid bishydroxamic acid

modified in the meta and para positions without loss of activity; however, in general, larger substituents are associated with loss of activity. The optimal methylene spacer is six methylenes, five- and seven-carbons spacers being less active.

The structure of the catalytic core of HDACs has been determined by x-ray crystallography (80). HDACs share an approximately 390-amino acid region of homology, referred to as the deacetylase core. Residues that form the active site are conserved across all HDACs. The deacetylase core identifies a gene superfamily that includes an HDAC homologue in the hyperthermophilic bacterium Aquifex aeolicus (termed "HDLP"), which was used for x-ray crystallography studies. There is a 35.2% base-pair identity between sequences of the catalytic core of the HDLP and of the mammalian HDAC1. HDLP deacetylates histones in vitro, its activity is inhibited by TSA and SAHA, but its specific activity is equal to about 7.5% of that of partially purified HDAC1. From x-ray crystallographic analyses of HDLP, an HDLP-TSA complex, and an HDLP-SAHA complex, the active catalytic site in the HDLP was shown to be formed by a tubular pocket, a zinc-binding site, and two asparagine-histidine charge-relay systems (Fig. 5). The hydroxamic acid moieties of TSA and SAHA bind to the zinc in the tubular pocket and the carbon-ring group projects out of the pocket on the surface of the protein.

### Activity of HDAC Inhibitors In Vitro

The hydroxamic acid-based HPCs (e.g., *m*-carboxy-cinnamic acid bishydroxamic acid [CBHA], suberic bishydroxamic acid

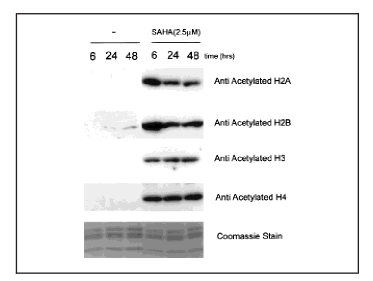


**Fig. 5.** SAHA (suberoylanilide hydroxamic acid) binds to the pocket of the catalytic site of a histone deacetylase-like protein, schematically represented by the netting. SAHA makes contact with residues at the rim, walls, and bottom of the pocket (enzyme pocket). The hydroxamic acid moiety of SAHA binds to the zinc at the bottom of the pocket (80). (The figure is courtesy of Michael S. Finnin and Nikola P. Pavletich.)

[SBHA], SAHA, and pyroxamide) (Fig. 4) inhibit partially purified HDAC1 and HDAC3 at concentrations of  $0.01-1.0 \mu M$  (34). Furthermore, the optimal concentrations of various HPCs that induce murine erythroleukemia (MEL) cell differentiation as assayed by the proportion of cells that become benzidine positive (a stain for heme of hemoglobin) are correlated directly with the concentration required to inhibit the activity of partially purified HDAC1 or HDAC3 over a wide concentration range.

With the use of MEL cells and T24 human bladder carcinoma cells in culture, the effects of SAHA and related hydroxamic acid-based HPCs on the acetylation of histones have been examined (34). SAHA, pyroxamide, SBHA, and CBHA (Fig. 4) cause accumulation of acetylated histones. Acetylated histone type-specific antibodies were used to show that, when cells were cultured with hydroxamic acid-based HPCs, the level of acetylation in histones H2A, H2B, H3, and H4 increased (Fig. 6). Increased histone acetylation could be detected as early as 1 hour after MEL or T24 cells were cultured with SAHA or other hydroxamic acid-based HPCs. The level of acetylated histones reached a maximum 6–12 hours after the addition of HPCs and remained elevated as long as the HPC was present (34).

HDAC inhibitors can induce growth arrest, differentiation, and/or apoptotic cell death in a wide variety of cultured transformed cells, including neuroblastoma, melanoma, and leukemia cells, as well as cells from breast, prostate, lung, ovary, and colon cancers (25–30,36,40,78,81). For example, SAHA induces terminal cell differentiation in several cell lines, including MEL, T24 human bladder carcinoma, and MCF-7 human breast adenocarcinoma. Differentiation was evaluated by parameters that included morphology, arrest in G<sub>1</sub> phase of the cell cycle, and developmental markers, such as hemoglobin in MEL cells, milk proteins in MCF-7 cells, and gelsolin in T24 cells. SAHA induces apoptotic death of human multiple myeloma cells (ARP-1), human prostate cell lines (LNCaP), and myelomonocytic leukemia cells (U937). CBHA induced apoptotic cell death of several human neuroblastoma cell lines, LAI-55n, KCN-69n, and SK-N-ER. Apoptosis was assayed by DNA fragmentation analysis and the deletion of a sub- $G_1$  (<2N ploidy) population by flow cytometry.



**Fig. 6.** Effect of SAHA (suberoylanilide hydroxamic acid) on histone acetylation in MEL cells. Cells were cultured without (–) or with 2.5  $\mu$ M SAHA for the times indicated. The acetylation of the histones was analyzed by use of antibodies specific for acetylated H2A, H2B, H3, and H4. The Coomassie-stained gel, **at the bottom,** indicates that the amount of protein loaded in each lane was similar [for details of methods, see~(36)].

Van Lint et al. (82) have shown that the action of HDAC inhibitors on gene expression is selective. In cells cultured with TSA, the expression of only about 2% of expressed genes is changed (increased or decreased) twofold or more compared with untreated control cells. Our laboratory has obtained comparable results with transformed cells cultured with SAHA. The basis for the gene selectivity of SAHA or TSA is not known.

One gene most consistently induced by HDAC inhibitors is the cyclin-dependent kinase inhibitor p21<sup>WAF1</sup>, which plays an important, if not determinant, role in the arrest of cell growth. Butyrate, TSA, depsipeptide, oxamflatin, MS-27-275, and the hydroxamic acid-based HPCs (28,31,32,34,83) induce p21<sup>WAF1</sup> transcription. The relation between SAHA-mediated histone hyperacetylation and increased p21<sup>WAF1</sup> gene expression has been studied in T24 human bladder carcinoma cells (84). Increased transcription of the p21<sup>WAF1</sup> gene is associated with an increased level of acetylation on histones associated with the p21<sup>WAF1</sup> gene.

### In Vivo Studies With HDAC Inhibitors

The butyrate analogue phenylbutyrate gave mixed results when tested as an HDAC inhibitor in animals and in a patient with APL. It was ineffective to moderately effective in inhibiting growth of solid tumors or leukemias, and that activity was observed only at relatively high doses (28). A 13-year-old girl with relapsed APM who no longer responded to treatment with retinoic acid alone was treated with retinoic acid plus phenylbutyrate and had a complete clinical remission that was sustained for 7 months, during five treatment courses, before relapsing and becoming resistant to this treatment (74). The acetylation of histones in her mononuclear blood cells was elevated during the period of administration of the phenylbutyrate. No remissions were induced in four other patients with APL (75).

Several other HDAC inhibitors, including depsipeptide (32), oxamflatin (35), MS-27-275 (32), and the hydroxamic acid-based HPCs (37–39), inhibit tumor growth in animal models (Figs. 3 and 4). TSA did not inhibit the growth of a human

melanoma xenograft in nude mice, but azeloic bishydroxamate did (40). Treatment with HDAC inhibitors can increase the accumulation of acetylated histone in tumor tissue and/or normal tissues (e.g., spleen, bone marrow cells, and peripheral mononuclear cells). Thus, the level of acetylated histones is a useful intermediary marker of HDAC inhibitor activity.

Hydroxamic acid-based HPCs (Fig. 4) have been tested extensively in animal studies. One study (37) used rats with Nmethylnitrosourea-induced mammary carcinomas. When these rats were fed SAHA (900 parts/million, continuously, beginning 7 days before the administration of N-methylnitrosourea), the incidence of mammary tumors was reduced by 40%, and the mean tumor volume was reduced by 78%—without side effects. Another study (39) used mice in which the tobacco-specific carcinogen 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanone induces lung tumors. When these mice were fed SAHA (900 parts/ million, continuously, beginning 7 days before administration of the carcinogen to the end of the studies), the formation of lung tumors was substantially inhibited—also without toxic effects. A third rodent study (38) used nude mice bearing transplanted CWR22 androgen-dependent human prostate tumors. When these mice were given SAHA (25, 50, or 100 mg/kg per day) daily by intraperitoneal injection for 3 weeks, starting as soon as palpable tumors were detected, SAHA suppressed tumor growth at all three doses. With doses of 50 and 100 mg/kg per day, the mean tumor volume was reduced by 97%. Acetylation of histones H3 and H4 increased in the CWR22 tumor cells within 6 hours after SAHA was injected. Pyroxamide had similar effects on CWR22 tumor growth and the accumulation of acetylated histones (Fig. 4). When SAHA or pyroxamide was given at doses that markedly inhibited tumor growth, no toxicity, as evaluated by weight gain and histologic examination of multiple tissues at necropsy, was detected.

### **CONCLUSIONS**

The studies summarized in this review indicate that the hydroxamic acid-based HPCs, in particular, SAHA and pyroxamide—are potent inhibitors of HDAC *in vitro* and *in vivo* and induce growth arrest, differentiation, or apoptotic cell death of transformed cells. We suggest that inhibition of HDAC activity leads to relaxation of the structure of chromatin associated with a specific set of programmed genes. The relaxed chromatin structure allows these genes to be expressed, which, in turn, arrests tumor cell growth. SAHA and pyroxamide are lead compounds among the family of hydroxamic acid-based HPCs and are currently in phase I clinical trials.

## REFERENCES

- Marks PA, Rifkind RA. Erythroleukemic differentiation. Annu Rev Biochem 1978;47:419–48.
- (2) Beug H, Blundell PA, Graft T. Reversibility of differentiation and proliferative capacity in avian myelomonocytic cells transformed by tsE26 leukemia virus. Genes Dev 1987;1:277–86.
- (3) Brinster RL. The effect of cells transferred into the mouse blastocyst on subsequent development. J Exp Med 1974;140:1049–56.
- (4) DeCosse JJ, Gossens CL, Kuzma JF, Unsworth BR. Breast cancer: induction of differentiation by embryonic tissue. Science 1973;181:1057–8.
- (5) Gootwine E, Webb CG, Sachs L. Participation of myeloid leukaemic cells injected into embryos in haematopoietic differentiation in adult mice. Nature 1982;299:63–5.
- (6) Illmensee K, Mintz B. Totipotency and normal differentiation of single

- teratocarcinoma cells cloned by injection into blastocysts. Proc Natl Acad Sci U S A 1976;73:549–53.
- (7) Brinster RL. The effects of cells transferred into mouse blastocyst on subsequent development. J Exp Med 1974;140:1049–56.
- (8) Friend C, Scher W, Holland JG, Sato T. Hemoglobin synthesis in murine virus-induced leukemic cells in vitro: stimulation of erythroid differentiation by dimethyl sulfoxide. Proc Natl Acad Sci U S A 1971;68:378–82.
- (9) Reuben RC, Wife RL, Breslow R, Rifkind RA, Marks PA. A new group of potent inducers of differentiation in murine erythroleukemia cells. Proc Natl Acad Sci U S A 1976;73:862–6.
- (10) Marks PA, Richon VM, Kiyokawa H, Rifkind RA. Inducing differentiation of transformed cells with hybrid polar compounds: a cell cycle-dependent process. Proc Natl Acad Sci U S A 1994;91:10251–4.
- (11) Lotan R. Different susceptibilities of human melanoma and breast carcinoma cell lines to retinoic acid-induced growth inhibition. Cancer Res 1979;39:1014–9.
- (12) Reiss M, Pitman SW, Sartorelli AC. Modulation of the terminal differentiation of human squamous carcinoma cells in vitro by all-trans-retinoic acid. J Natl Cancer Inst 1985;74:1015–23.
- (13) Strickland S, Mahdavi V. The induction of differentiation in teratocarcinoma stem cells by retinoic acid. Cell 1978;15:393–403.
- (14) Sidell N. Retinoic acid-induced growth inhibition and morphologic differentiation of human neuroblastoma cells in vitro. J Natl Cancer Inst 1982; 68:589–96.
- (15) Takenaga K, Hozumi M, Sakagami Y. Effects of retinoids on induction of differentiation of cultured mouse myeloid leukemia cells. Cancer Res 1980; 40:914–9.
- (16) Zhou JG, Norman AW, Lubbert M, Collins ED, Uskokovic MR, Koeffler HP. Novel vitamin D analogs that modulate leukemic cell growth and differentiation with little effect on either intestinal calcium absolution or bone mobilization. Blood 1989;74:82–93.
- (17) Michaeli J, Marks PA, Rifkind RA. Differentiating agents in cancer therapy. In: Pinedo HM, Longo D, Chabner BA, editors. Cancer chemotherapy and biological response modifiers. Annual 13. Amsterdam (The Netherlands): Elsevier; 1992. p. 286–307.
- (18) Huberman E, Callabaum MF. Induction of terminal differentiation in human promyelocytic leukemia cells by tumor-promoting agents. Proc Natl Acad Sci U S A 1979;76:1293–7.
- (19) Pierce GB, Speers WC. Tumors as caricatures of the process of tissue renewal: prospects for therapy by directing differentiation. Cancer Res 1988;48:1996–2004.
- (20) Grunstein M. Histone acetylation and chromatin structure and transcription. Nature 1997;389:349–52.
- (21) Muraoka M, Konishi M, Kikuchi-Yanoshita R, Tanaka K, Shitara N, Chang JM, et al. p300 gene alterations in colorectal and gastric carcinomas. Oncogene 1996;12:1565–9.
- (22) He LZ, Guidez F, Tribioli C, Peruzzi D, Ruthardt M, Zelent A, et al. Distinct interactions of PML-RARalpha and PLZF-RARalpha with corepressors determine differential responses to RA in APL. Nat Genet 1998; 18:126–35
- (23) Grignani F, De Matteis S, Nervi C, Tomassoni L, Gelmetti V, Cioce M, et al. Fusion proteins of the retinoic acid receptor-alpha recruit histone deacetylase in promyelocytic leukaemia. Nature 1998;391:815–8.
- (24) Lin RJ, Nagy L, Inoue S, Shao W, Miller WH Jr, Evans RM. Role of the histone deacetylase complex in acute promyelocytic leukaemia. Nature 1998;391:811–4.
- (25) Tsuji N, Kobayashi M, Nagashima K, Wakisaka Y, Koizumi K. A new antifungal antibiotic, trichostatin. J Antibiot (Tokyo) 1976;29:1–6.
- (26) Yoshida M, Kijima M, Akita M, Beppu T. Potent and specific inhibition of mammalian histone deacetylase both in vivo and in vitro by trichostatin A. J Biol Chem 1990;265:17174–9.
- (27) Kijima M, Yoshida M, Sugita K, Horinouchi S, Beppu T. Trapoxin, an antitumor cyclic tetrapeptide, is an irreversible inhibitor of mammalian histone deacetylase. J Biol Chem 1993;268:22429–35.
- (28) Newmark HL, Lupton JR, Young CW. Butyrate as a differentiating agent: pharmacokinetics, analogues and current status. Cancer Lett 1994;78:1–5.
- (29) Yoshida M, Horinouchi S, Beppu T. Trichostatin A and trapoxin: novel chemical probes for the role of histone acetylation in chromatin structure and function. Bioessays 1995;17:423–30.
- (30) Kwon HJ, Owa T, Hassig CA, Shimada J, Schreiber SL. Depudecin in-

- duces morphological reversion of transformed fibroblasts via the inhibition of histone deacetylase. Proc Natl Acad Sci U S A 1998;95:3356–61.
- (31) Carducci M, Bowling MK, Eisenberger M, Sinibaldi V, Chen T, Nor D, et al. Phenylbutyrate (PB) for refractory solid tumors: phase I clinical and pharmacologic evaluation of intravenous and oral PB. Anticancer Res 1997;17:3972–3.
- (32) Saito A, Yamashita T, Mariko Y, Nosaka Y, Tsuchiya K, Ando T, et al. A synthetic inhibitor of histone deacetylase, MS-27-275, with marked in vivo antitumor activity against human tumors. Proc Natl Acad Sci U S A 1999; 96:4592–7.
- (33) Nakajima H, Kim YB, Terano H, Yoshida M, Horinouchi S. FR901228, a potent antitumor antibiotic, is a novel histone deacetylase inhibitor. Exp Cell Res 1998;241:126–33.
- (34) Richon VM, Emiliani S, Verdin E, Webb Y, Breslow R, Rifkind RA, et al. A class of hybrid polar inducers of transformed cell differentiation inhibits histone deacetylases. Proc Natl Acad Sci U S A 1998;95:3003–7.
- (35) Kim YB, Lee KH, Sugita K, Yoshida M, Horinouchi S. Oxamflatin is a novel antitumor compound that inhibits mammalian histone deacetylase. Oncogene 1999;18:2461–70.
- (36) Glick RD, Swendeman SL, Coffey DC, Rifkind RA, Marks PA, Richon VM, et al. Hybrid polar histone deacetylase inhibitor induces apoptosis and CD95/CD95 ligand expression in human neuroblastoma. Cancer Res 1999; 59:4392–9.
- (37) Cohen LA, Amin S, Marks PA, Rifkind RA, Desai D, Richon VM. Chemoprevention of carcinogen-induced mammary tumorigenesis by the hybrid polar cytodifferentiation agent, suberanilohydroxamic acid (SAHA). Anticancer Res 1999;19:4999–5005.
- (38) Butler LM, Higgins B, Fox WD, Agus DB, Cordon-Cardo C, Scher HJ, et al. Hybrid polar inhibitors of histone deacetylase suppress the growth of the CWR22 human prostate cancer xenograft. Proc Am Assoc Cancer Res 2000;41:abstract 289.
- (39) Desai D, El-Bayoumy K, Amin S. Chemopreventive efficacy of suberanilohydroxamic acid (SAHA), a cytodifferentiating agent, against tobaccospecific nitrosamine 4-(methylnitros-amino)-1-(3-pyridyl)-1butanone (NNK)-induced lung tumorigenesis [abstract]. Proc Am Assoc Cancer Res 1999;40:abstract 2396.
- (40) Qui L, Kelso MJ, Hansen C, West ML, Fairlie DP, Parsons PG. Antitumour activity *in vitro* and *in vivo* of selective differentiating agents containing hydroxamate. Br J Cancer 1999;80:1252–8.
- (41) Kornberg RD. Chromatin structure: a repeating unit of histones and DNA. Science 1974;184:868–71.
- (42) Kornberg RD, Lorch Y. Twenty-five years of the nucleosome, fundamental particle of the eukaryote chromosome. Cell 1999;98:285–94.
- (43) Luger K, Mader AW, Richmond RK, Sargent DF, Richmond TJ. Crystal structure of the nucleosome core particle at 2.8 A resolution. Nature 1997; 389:251–60.
- (44) Davie JR. Covalent modifications of histones: expression from chromatin templates. Curr Opin Genet Dev 1998;8:173–8.
- (45) Kouzarides T. Histone acetylases and deacetylases in cell proliferation. Curr Opin Genet Dev 1999;9:40–8.
- (46) Bannister AJ, Kouzarides T. The CBP co-activator is a histone acetyltransferase. Nature 1996;384:641–3.
- (47) Ogryzko VV, Schiltz RL, Russanova V, Howard BH, Nakatani Y. The transcriptional coactivators p300 and CBP are histone acetyltransferases. Cell 1996;87:953–9.
- (48) Mizzen CA, Yang XJ, Kokubo T, Brownell JE, Bannister AJ, Owen-Hughes T, et al. The TAF(II)250 subunit of TFIID has histone acetyltransferase activity. Cell 1996;87:1261–70.
- (49) Spencer TE, Jenster G, Burcin MM, Allis CD, Zhou J, Mizzen CA, et al. Steroid receptor coactivator-1 is a histone acetyltransferase. Nature 1997; 389:194–8.
- (50) Siddique H, Zou JP, Rao VN, Reddy ES. The BRCA2 is a histone acetyltransferase. Oncogene 1998;16:2283–5.
- (51) Waltzer L, Bienz M. Drosophila CBP represses the transcription factor TCF to antagonize Wingless signalling. Nature 1995;395:521–5.
- (52) Taunton T, Hassig CA, Schreiber SL. A mammalian histone deacetylase related to the yeast transcriptional regulator Rpd3p. Science 1996;272:408–11.
- (53) Emiliani S, Fischle W, Van Lint C, Al-Abed Y, Verdin E. Characterization of a human RPD3 ortholog, HDAC3. Proc Natl Acad Sci U S A 1998;95: 2795–800.

- (54) Grozinger CM, Hassig CA, Schreiber SL. Three proteins define a class of human histone deacetylases related to yeast Hda1p. Proc Natl Acad Sci U S A 1999;96:4868–73.
- (55) Verdel A, Khochbin S. Identification of a new family of higher eukaryotic histone deacetylases. Coordinate expression of differentiation-dependent chromatin modifiers. J Biol Chem 1999;274:2440–5.
- (56) Fischle W, Emiliani S, Hendzel MJ, Nagase T, Nomura N, Voetler W, et al. A new family of human histone deacetylases related to *Saccharomyces* cerevisiae HDA1p. J Biol Chem 1999;274:11713–20
- (57) Miska EA, Karlsson C, Langley E, Nielsen SJ, Pines J, Kouzarides T. HDAC4 deacetylase associates with and represses the MEF2 transcription factor. EMBO J 1999;18:5099–107.
- (58) Kao HY, Downes M, Ordentlich P, Evans RM. Isolation of a novel deacetylase reveals that class I and class II deacetylases promote SMRTmediated repression. Genes Dev 2000;14:55–66.
- (59) Hu E, Chen Z, Frederickson T, Zhu Y, Kirkpatrick R, Zhang GF, et al. Cloning and characterization of a novel human class I histone deacetylase that functions as a transcription repressor. J Biol Chem 2000;275:254–64.
- (60) Allfrey VG. Post synthetic modifications of histone: a mechanism for the control of chromosome structure by the modulation of histones—DNA interactions In: Li HJ, Eckhardt RA, editors. Chromatin and chromosome structure. New York (NY): Academic Press; 1977. p. 167–91.
- (61) Cameron EE, Bachman KE, Myohanen S, Herman JG, Baylin SB. Synergy of demethylation and histone deacetylase inhibition in the re-expression of genes silenced in cancer. Nat Genet 1999;21:103–7.
- (62) Jones PL, Veenstra GJ, Wade PA, Vermaak D, Kass SU, Landsberger N, et al. Methylated DNA and MeCP2 recruit histone deacetylase to repress transcription. Nat Genet 1998;19:187–91.
- (63) Bird AP, Wolffe AP. Methylation-induced repression—belts, braces, and chromatin. Cell 1999;99:451–4.
- (64) Vignali M, Hassan AH, Neely KE, Workman JL. ATP-dependent chromatin-remodeling complexes. Mol Cell Biol 2000;20:1899–910.
- (65) Tyler JK, Kadonaga JT. The "dark side" of chromatin remodeling: repressive effects of transcription. Cell 1999;99:443–6.
- (66) Gu W, Roeder RG. Activation of p53 sequence-specific DNA binding by acetylation of the p53 C-terminal domain. Cell 1997;90:595–606.
- (67) Boyes J, Byfield P, Nakatani Y, Ogryzko V. Regulation of activity of the transcription factor GATA-1 by acetylation. Nature 1998;396:594–8.
- (68) Imhof A, Yang XJ, Ogryzko VV, Nakatani Y, Wolffe AP, Ge H. Acetylation of general transcription factors by histone acetyltransferases. Curr Biol 1997;7:689–92.
- (69) Luo RX, Postigo AA. Dean DC. Rb interacts with histone deacetylase to repress transcription. Cell 1998;92:463–73.
- (70) Brehm A, Miska EA, McCance DJ, Reid JL, Bannister AJ, Kouzarides T. Retinoblastoma protein recruits histone deacetylase to repress transcription. Nature 1998;391:597–601.
- (71) Petrij F, Giles RH, Dauwerse HG, Saris JJ, Hennekam RC, Masuno M, et al. Rubinstein–Taybi syndrome caused by mutations in the transcriptional co-activator CBP. Nature 1995;376:348–51.
- (72) Sobulo OM, Borrow J, Tomek R, Reshmi S, Harden A, Schlegelberger B,

- et al. MLL is fused to CBP, a histone acetyltransferase, in therapy-related acute myeloid leukemia with a t(11;16)(q23;p13.3). Proc Natl Acad Sci U S A 1997;94:8732–7.
- (73) Giles RH, Peters DJ, Breuning MH. Conjunction dysfunction: CBP/p300 in human disease. Trends Genet 1998;14:178–83.
- (74) Warrell RP Jr, He LZ, Richon V, Calleja E, Pandolfi PP. Therapeutic targeting of transcription in acute promyelocytic leukemia by use of an inhibitor of histone deacetylase. J Natl Cancer Inst 1998;90:1621–5.
- (75) Novich S, Camacho L, Gallagher R, Chanel S, Ho R, Tolentino T, et al. Initial clinical evaluation of "transcription therapy" for cancer: all-trans retinoic acid plus phenylbutyrate. Blood 1999;94:Suppl 1. p61a.
- (76) Dhordain P, Lin RJ, Quief S, Lantoine D, Kerckaert JP, Evans RM, et al. The LAZ3(BCL-6) oncoprotein recruits a SMRT/mSIN3A/histone deacetylase containing complex to mediate transcriptional repression. Nucleic Acids Res 1998:26:4645–51.
- (77) Wang J, Hoshino T, Redner RL, Kajigaya S, Liu JM. ETO, fusion partner in t(8;21) acute myeloid leukemia, represses transcription by interaction with the human N-CoR/mSin3/HDAC1 complex. Proc Natl Acad Sci U S A 1998:95:10860–5.
- (78) Darkin-Rattray SJ, Gurnett AM, Myers RW, Dulski PM, Crumley TM, Allocco JJ, et al. Apicidin: a novel antiprotozoal agent that inhibits parasite histone deacetylase. Proc Natl Acad Sci U S A 1996;93:13143–7.
- (79) Breslow R, Jursic B, Yan ZF, Friedman E, Leng L, Ngo L, et al. Potent cytodifferentiating agents related to hexamethylenebisacetamide. Proc Natl Acad Sci U S A 1991;88:5542–6.
- (80) Finnin MS, Donigian JR, Cohen A, Richon VM, Rifkind RA, Marks PA, et al. Structures of a histone deacetylase homologue bound to TSA and SAHA. Nature 1999;401:188–93.
- (81) Riggs MG, Whittaker RG, Neumann JR, Ingram VS. n-Butyrate causes histone modification in HeLa and Friend erythroleukaemia cells. Nature 1997;268:462–4.
- (82) Van Lint C, Emiliani S, Verdin E. The expression of a small fraction of cellular gene is changed in response to histone hyperacetylation. Gene Expr 1996;5:245–4.
- (83) Xiao H, Hasegawa T, Isobe KI. Both Sp1 and Sp3 are responsible for p21 waf1 promoter activity induced by histone deacetylase inhibitor in NIH3T3 cells. J Cell Biochem 1999;73:291–302.
- (84) Richon VM, Sandhoff TW, Rifkind RA, Marks PA. SAHA induced histone hyperacetylation of the p21<sup>waf1</sup> gene precedes transcription. Proc Am Assoc Cancer Res 2000;40:abstract 5131.

#### Notes

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