Epigenetic Modifiers: Basic Understanding and Clinical Development

Richard L. Piekarz and Susan E. Bates

Abstract

More than 60 years after the first description of differentiation in cell culture and 40 years after the synthesis of 5-azacytidine, epigenetic therapies have been added to the anticancer armamentarium. DNA methyltransferase (DNMT) inhibitors such as 5-aza-2'-deoxycytidine or 5-azacytidine have been approved in myelodysplastic syndrome (MDS) and acute myelogenous leukemia (AML), whereas the histone deacetylase inhibitors (HDIs) including vorinostat, romidepsin, panobinostat, belinostat, and entinostat have been shown to be active in cutaneous and peripheral T-cell lymphoma. Although the range of malignancies in which monotherapy with DNMT inhibitors or HDIs are effective has been limited to date, the possibility remains that a broader spectrum of activity will be identified as combination studies are completed. Meanwhile, basic science has provided a steadily increasing understanding of the complexity of the epigenome, including the histone code and triggers for aberrant methylation, and their contribution to oncogenesis. As our basic understanding of the epigenetics of cancer increases, the number of potential therapeutic targets will also increase, offering more hope in the quest to treat cancer by normalizing the epigenome. This issue of CCR Focus is dedicated to understanding the clinical and translational aspects of epigenetics research.

Epigenetics—the study of stable genetic modifications that result in changes in gene expression and function without a corresponding alteration in DNA sequence. The epigenome is a catalog of the epigenetic modifications that occur in the genome. Epigenetic changes have been associated with disease, but further progress requires the development of better methods to detect the modifications and a clearer understanding of factors that drive these changes. (National Institutes of Health Roadmap).1

The inclusion of epigenetics in the National Institutes of Health (NIH) roadmap has highlighted the need for research in both epigenetic mechanisms of oncogenesis and in epigenetic therapies. This decade has seen the addition of several epigenetic therapies to the anticancer armamentarium. The U.S. Food and Drug Administration (FDA) has approved three epigenetic targeting agents for oncology: Vidaza (Celgene, Summit, NJ, azacitidine, 2004), Dacogen (SuperGen, Inc., Dublin, CA, decitabine, 5-aza-2'-deoxycytidine, 2006), and Zolinza (Merck & Co., Inc., Whitehouse Station, NJ, vorinostat, 2006); and many more are in clinical and preclinical development (Table 1). Yet, we still do not fully understand the best schedule, the ideal dosing, or true mechanism of action of these agents. Their enzymatic targets are known; however, the downstream effectors have not been elucidated. This issue of CCR Focus addresses our current understanding of these matters and addresses the arrival of epigenetic therapies as conventional anticancer therapies.

Differentiating Agents: Epigenetic Targets

One interesting aspect of epigenetic therapies is their historical origin as differentiating agents. The spontaneous differentiation of leukemic cells was first noted in cell culture systems in the 1940s (1). Over time, it became clear that many different agents and culture conditions could promote such differentiation, although there were model-specific differences (2). Agents found to induce differentiation in leukemic cells included antifolates, anthracyclines, camptothecins, retinoids, phorbol esters, vitamins D3 and B12, and various cytokines. Among these agents were also compounds later understood to alter chromatin structure: hypomethylating agents and histone deacetylase inhibitors (HDIs). In 1977, Constantinides reported that 5-azacytidine was able to induce striated muscle cells from murine embryonic 10T1/2 cells and in 1979 this work was extended to show that pluripotency in these cells following exposure to 5-azacytidine could produce chondrocytes, adipocytes, and muscle cells (3, 4). Jones and Taylor then reported in 1980 that these differentiating effects were associated with alterations in DNA methylation (5). DNA hypomethylation was soon linked to activation of gene expression resulting in phenotypic cell differentiation in murine Friend erythroleukemia cells by observation of the

tight association between the activities following treatment with 5-azacytidine or its analog 5-aza-2’-deoxycytidine (6). In independent studies, erythroid differentiation following the addition of sodium butyrate was observed at the phenotypic level in the human erythroleukemic cell line, K562 (7, 8). A review of sodium butyrate in 1982 observed that the agent induced histone hyperacetylation and connected this property with the inhibition of cell growth and the differentiated phenotype in certain model systems, notably in erythroleukemia cells (9).

These observations launched the search for compounds that could be used clinically in the differentiation therapy of hematologic disorders and malignancies, despite the differing mechanisms of action. Case reports described the first clinical use of 5-azacytidine to induce fetal hemoglobin in a patient with beta-thalassemia in 1982 (10) and in a patient with sickle cell anemia in 1983 (11). The approval of 5-azacytidine in 2004 for myelodysplastic syndrome (MDS) marked the end of a 40-year quest to find a clinical application for a compound originally synthesized in 1964 (12). Clinical experience with sodium butyrate was first reported with a partial response in a patient with acute myelogenous leukemia (AML) in 1983 (13), reviewed by Gore and Carducci (14). Although butyrate was never approved as an anticancer therapy, its evaluation and that of other short chain fatty acids led directly to the development of hydroxamic acids, notably vorinostat, now approved for the treatment of cutaneous T-cell lymphoma (CTCL) (ref. 15).

### DNA Methylation as a Therapeutic Target

Recognition of the extent of aberrant methylation found in human cancer has led to the understanding that DNA methylation, if not a first hit in oncogenesis, is at least a mediator of oncologic progression. In distilling the literature surrounding this topic for this issue of CCR Focus, McCabe, Brandes, and Vertino note that cancer cells exhibit widespread loss of intergenic DNA methylation and gain of DNA methylation in promoter-associated CpG islands, defined as clusters of the CpG dinucleotide, that are normally found in an unmethylated state (16). These methylated CpG islands are often found associated with the promoters of tumor suppressor genes, such as the retinoblastoma (Rb) gene (17) or p21 (18).

As new methods for examining genome-wide methylation patterns have become available, the extent to which aberrant

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**Table 1. Drugs with epigenetic targets either approved or in development**

<table>
<thead>
<tr>
<th>Drug target</th>
<th>Generic or trade name</th>
<th>Development name</th>
<th>Chemical class</th>
<th>Pharmaceutical sponsor</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNMT inhibitor</td>
<td>Azacitidine</td>
<td>5-azacytidine</td>
<td>Nucleoside analog</td>
<td>Celgene (Summit, NJ)</td>
<td>FDA approved, 2004</td>
</tr>
<tr>
<td>DNMT inhibitor</td>
<td>Vidaza</td>
<td>Decitabine</td>
<td>Nucleoside analog</td>
<td>Eisai Co., Ltd. (Tokyo, Japan)</td>
<td>EMEA approved, 2008</td>
</tr>
<tr>
<td>DNMT inhibitor</td>
<td>Azacitidine</td>
<td>Decitabine</td>
<td>Nucleoside analog</td>
<td>Eisai Co., Ltd. (Tokyo, Japan)</td>
<td>FDA approved, 2006</td>
</tr>
<tr>
<td>DNMT inhibitor</td>
<td>SGI-110</td>
<td>RG108</td>
<td>Nucleoside analog</td>
<td>SuperGen (Dublin, CA)</td>
<td></td>
</tr>
<tr>
<td>DNMT inhibitor</td>
<td>SGI-1036</td>
<td>DZNep</td>
<td>Nucleoside analog</td>
<td>SuperGen</td>
<td></td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Sodium phenylbutyrate</td>
<td>Bupropion</td>
<td>Small chain fatty acid</td>
<td>Ucyclyd Pharma (Scottsdale, AZ)</td>
<td>FDA approved, 2004</td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Valproic acid</td>
<td>Depakote</td>
<td>Small chain fatty acid</td>
<td>Abbott Laboratories (Abbott Park, IL)</td>
<td>FDA approved, 2006</td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Vorinostat</td>
<td>SAHA</td>
<td>Hydroxamic acid</td>
<td>Merck &amp; Co.</td>
<td>FDA approved, 2006</td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Panobinostat</td>
<td>LBHS89</td>
<td>Hydroxamic acid</td>
<td>Novartis Pharmaceuticals (East Hanover, NJ)</td>
<td>FDA approved, 2006</td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Belinostat</td>
<td>PXD101</td>
<td>Hydroxamic acid</td>
<td>TopoTarget (Rockaway, NJ)</td>
<td>FDA approved, 2006</td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Romidepsin</td>
<td>JNJ-2681585</td>
<td>Hydroxamic acid</td>
<td>Johnson &amp; Johnson (Langhorne, PA)</td>
<td>FDA approved, 2006</td>
</tr>
<tr>
<td>HDAC inhibitor</td>
<td>Entinostat</td>
<td>Dapsipetide, FK228</td>
<td>Cyclic peptide</td>
<td>Gloucester Pharmaceuticals (Cambridge, MA)</td>
<td>New drug application filed with FDA</td>
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<td>HDAC inhibitor</td>
<td>Entinostat</td>
<td>MS275, SNDX-275</td>
<td>Benzamide</td>
<td>Syndax Pharmaceuticals (Waltham, MA)</td>
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<td>HDAC inhibitor</td>
<td>MGCD-0103</td>
<td>Benzamide</td>
<td></td>
<td>MethylGene (Montreal, Quebec, Canada)</td>
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</tr>
</tbody>
</table>

**Abbreviation:** EMEA, European Medicines Agency.
and varying methylation patterns exist in human cancer has begun to emerge. Neither the trigger, nor the initiating event, nor the sequence of events that follow and give rise to hypermethylation, nor the mechanisms underlying the selection of a high prevalence of tumor suppressor genes for methylation have been determined. McCabe and colleagues illustrate the multiple pathways involved in hypermethylation (16), leading to the sobering realization that, although we have been successful with one strategy for reducing hypermethylation with the nucleoside analogs 5-aza-2'-deoxycytidine or 5-aza-2'-cytidine (now understood to function as DNA methyltransferase (DNMT) inhibitors following their incorporation into DNA), we are still far from understanding how to normalize DNA methylation in cancer. However, the progress made in basic science has allowed the identification of multiple genes in which methylation has potential clinical application in diagnosis, early detection, prognosis, or therapy outcome.

Issa and Kantarjian offer a more sanguine outlook in the translation of these two agents to the clinic in the therapy of MDS (19). The authors describe the earlier clinical failure of agents that had clear ability to demethylate and reactivate genes in preclinical models, followed by the slow recognition that there was a biphasic dose response curve in which high doses of the nucleoside analog inhibited DNA synthesis whereas hypomethylation requires DNA synthesis. At lower doses there would be DNA incorporation followed by DNMT inhibition. This meant a prolonged dosing period for MDS, eventually yielding response rates in the 60% to 70% range for both 5-aza-2'-deoxycytidine and 5-aza-2'-cytidine. Having an effective agent offers the opportunity to develop biomarkers of response. Issa summarizes these efforts, noting that whereas reduced 5-methylcytosine content can be detected in correlative studies, change in gene expression has correlated better with response than has alteration in global methylation. This observation leads to the important issues of determining mechanisms of resistance to DNMT inhibitors (20) and how to improve on the clinical results already obtained.

**Histone Deacetylase as a Therapeutic Target**

HDIs were found to be remarkable and effective cytotoxic agents in vitro. Whereas the traditional mechanism of action underlying the antineoplastic activity of HDIs was considered the increased acetylation of lysine residues that form the octomeric histone core of chromatin, in recent studies a remarkable array of potential mechanisms has been proposed, suggesting the mechanisms underlying HDAC activity are pleiotropic and likely to be cell context-dependent. In this issue of CCR Focus, Schrump discusses the varied mechanisms of action proposed to underlie the activity of HDIs in vitro (21). These mechanisms include:

1. Histone acetylation with alterations in gene expression that effect cell cycle arrest and limit cell growth, including up-regulation of genes such as p21, p27, and other genetic markers of differentiation; and down-regulation of genes involved in growth such as cyclin D (22–24);
2. Acetylation of nonhistone proteins such as p53, HIF-1alpha, pRB, STAT-3, Rel A/p65, or estrogen receptor that may impair their function and thereby influence cell growth or survival (25–27);
3. Acetylation of Hsp90, with its attendant loss of ability to chaperone client proteins resulting in their ubiquitinylination and proteasomal degradation (28, 29);
4. A prometaphase cell cycle arrest that results from reduced premitotic phosphorylation of pericentromeric histone H3 and disruption of kinetochore assembly (30);
5. An antiangiogenic effect potentially mediated by impairing HIF-1α stability (31);
6. Direct activation of apoptotic pathways through reduction of antiapoptotic proteins such as Bcl-2 and increased expression of proapoptotic proteins such as BAX and BAK (32, 33);
7. Enhanced production of reactive oxygen species (ROS) (refs. 34, 35);
8. Disruption of aggresome formation through acetylation of tubulin (36);
9. Enhanced antitumor immunity through enhancement of TRAIL or up-regulation of antigen expression that could facilitate cancer cell recognition (37–40);
10. Disruption of DNA repair through acetylation or down-regulation of proteins such as Ku70, Ku86, BRCA1, and RAD51 (41–43).

One question that such a wide-ranging list raises is whether histone acetylation is actually an indispensable component of HDI activity. If differentiation is important, then histone acetylation per se may not be needed, as other agents induce differentiation. On the other hand, any one of the activities above could be critical in a given cell type. One of the complicating aspects of the laboratory evaluations that elucidated the multiple mechanisms of action itemized above is the use of different cell lines and the focus on only one or two aspects within each study. It is important that, going forward, in vitro and in vivo studies of HDIs embrace the possibility that multiple activities can occur concurrently and that these are assessed simultaneously. Furthermore, some consideration for evaluating these various mechanisms should be incorporated into clinical trials. Some investigators have championed one effect over another as being more important in limiting cell growth or inducing cytotoxicity. On the other hand, the multiplicity of these activities has raised the question of whether more specific HDIs should be developed. The current HDIs target class I (HDACs 1, 2, 3, and 8), class II (HDAC 4, 5, 6, 7, 9, and 10), and HDAC 11 (generally considered class IV) enzymes to varying degrees. It is quite possible that different HDIs may have a spectrum of activities much as we have already observed for different agents that belong to a general drug class such as anthracyclines, vinca alkaloids, or platinums.

Although, as noted above, the HDIs have a range of pathways by which to inhibit cell growth or trigger cell death, the clinical activity to date has been largely confined to hematologic malignancies, and particularly T-cell lymphomas. Prince and coauthors examine the clinical experience with HDIs to date, including a systematic review of results from the early phase...
trials for HDIs in development, including both outcome and toxicities (44). The authors note the dramatic activity in T-cell lymphoma, both peripheral T-cell lymphoma (PTCL) and CTCL, as a class effect observed with several HDIs and then go on to describe examples of activity in patients with B-cell lymphoma, Hodgkin lymphoma, and multiple myeloma. They note dose- and schedule-dependent activity in AML, suggesting that in the absence of the ability to markedly increase dosing of the HDI, effective combinations will be needed to exploit the observed activity.

**HDAC Inhibitors in T-Cell Lymphomas**

The demonstration that HDIs are active in T-cell lymphoma, and the approval of vorinostat for CTCL has raised a number of questions:

What is the mechanism of the marked efficacy of HDIs in T-cell lymphoma? With what seems to be a class effect, is there a basis for differences among the HDIs? And finally, with the approval of vorinostat, should other HDIs be developed?

The marked efficacy of HDIs in T-cell lymphoma is not understood. It is tempting to speculate that the responsive subset of T-cell lymphomas has its origin in an as-yet unknown chromosomal rearrangement that recruits the class 1 HDACs to the promoter of a gene normally limiting cell proliferation. However, chromosomal alterations such as those described in AML have not been described in T-cell lymphoma. Further, the lymphomas in general have proven to be a tumor type distinctly susceptible to different therapeutic interventions. It seems equally likely that HDIs trigger apoptosis in an apoptosis-prone environment.

Differences between the structurally very different HDIs currently in development include schedule, potency, and pharmacology. Examples of scheduling differences include vorinostat administered on a daily oral schedule; panobinostat administered orally three times weekly; romidepsin with an intravenous day 1, 8, and 15 schedule; and belinostat administered by short intravenous infusion on days 1 through 5. There are marked differences in potency, when observed from an in vitro standpoint. Romidepsin is active at nanomolar concentrations whereas vorinostat is active at micromolar concentrations. In pharmacokinetics, romidepsin has been found to have a 2.5-hour half-life, only slightly longer than that of vorinostat. In contrast, panobinostat has a longer half-life at 8 hours, and entinostat a much longer half-life of more than 30 hours (45, 46). Differences in pharmacology are incompletely understood. One mechanism of protection from normal tissue toxicity is the efflux of compounds by P-glycoprotein. As a substrate for P-glycoprotein–mediated drug efflux, romidepsin would likely not penetrate into the central nervous system, whereas other HDAC inhibitors may have central nervous system effects (47, 48). As one example, Depakote (Abbott Laboratories, Abbott Park, IL, valproic acid), is in clinical use as an antiseizure medication by P-glycoprotein. As a substrate for P-glycoprotein–mediated drug efflux, romidepsin would likely not penetrate into the central nervous system, whereas other HDAC inhibitors may have central nervous system effects (47, 48). As one example, Depakote (Abbott Laboratories, Abbott Park, IL, valproic acid), is in clinical use as an antiseizure medication. As one example, Depakote (Abbott Laboratories, Abbott Park, IL, valproic acid), is in clinical use as an antiseizure medication not thought to be susceptible to P-glycoprotein–mediated drug efflux, important components in the blood brain barrier (49). With variations in structure, schedule, and pharmacology, differences in spectrum of activity are likely to emerge over time.
Recognizing that the authors have a bias about the answer to the last question above, on the basis of a decade of effort in the development of romidepsin, we would argue in the affirmative. We have noted dramatic, durable activity of romidepsin in the National Cancer Institute multicenter trial in patients with T-cell lymphoma (Fig. 1) (ref. 50). As discussed elsewhere in this issue of *CCR Focus*, a comparable level of activity was noted in an independent registration trial (44), and romidepsin is currently under review by the FDA for this indication. One need only think of current antineoplastic agents to conclude that agents with identical targets may have very different antineoplastic activities. Daunorubicin and doxorubicin have very similar structures, differing only by a hydroxyl group, and have a very different spectrum of activity, in leukemias and solid tumors, respectively. The most recent addition to the microtubule targeting armamentarium, the epothilones, and their predecessors the taxanes, provide an example of drugs with differing structures, but identical targets.

**Understanding the Restricted Clinical Efficacy**

One observation is clear—the extraordinary activity of DNMT inhibitors and HDIs *in vitro* has not translated to clinical efficacy in solid tumors.

As regards the DNMT inhibitors, Issa and Kantarjian address the question of why hypomethylating agents were able to succeed in MDS and yet have failed to benefit patients with other tumors, at least to date (19). One possible answer lies in the high doses of the drug used in earlier studies, typically the maximally tolerated dose, likely resulting in inhibition of DNA synthesis rather than DNMT inhibition and hypomethylation. Alternatively, the agents may be susceptible to the drug resistance mechanisms prevalent in the patient populations enrolled in earlier studies, suggesting that a patient population without a significant degree of prior therapy should be tested (19, 20).

Improved clinical outcomes in nonhematologic malignancies may be observed if lower doses of demethylating agents are used, and if rational combination therapies are studied.

For currently available HDIs this raises the general question of whether the lack of activity in solid tumors is due to incorrect dosing or a lack of potency. An earlier assumption that resistance to HDIs could result simply from the slower growth rate of solid tumors has been discounted to some extent. *In vitro* studies have shown sensitivity to HDAC inhibition in culture conditions in which growth is inhibited, such as in tumor spheroids or in serum starvation, in which resistance to classical cytotoxic agents is observed (51, 52). This stands in contrast to DNMT inhibitors, which require DNA synthesis. Despite the *in vitro* data arguing against drug resistance due to slow growth, the fact remains that responses to HDIs in solid tumors in monotherapy trials have been rare, almost individually reportable. Figure 2 shows CT scans obtained in a patient with renal cell carcinoma who enrolled in a phase I study of romidepsin (53) and experienced a partial remission lasting 5 months following therapy with romidepsin. Unfortunately, a follow-up phase II trial failed to confirm activity in renal cancer (54).

One explanation for the drug resistance and the trademark failure of HDI activity in adult solid tumors may be that the changes in gene expression, as mentioned above, are associated with phenotypic and differentiation effects in solid tumors that may promote survival, rather than apoptosis. Several studies have suggested that cells that undergo p21-mediated growth arrest are rescued from the more cytotoxic effects of the HDIs (24, 55). Similarly, HDIs may activate nuclear factor-κB (NF-κB), which is believed to contribute to resistance to HDAC-induced apoptosis through its prosurvival activities. Prevention of NF-κB activation through siRNA knockdown or proteasome inhibition sensitizes head and neck cancer cells to HDIs (56). Finally, in addition to direct activation of apoptotic pathways.
following HDAC exposure, autophagy has been noted in cells treated with HDIs (57). Although autophagy can enhance apoptosis, it may also be a cell survival mechanism (58). Although these activities can be seen as a mechanism of drug resistance or an adaptive process as found following exposure to anticancer agents, it seems more likely at this writing that they are among the many downstream effects of the HDIs.

Another possible explanation for the difference in sensitivity to HDAC inhibition between solid tumors and hematologic malignancies is that the latter may have fewer mutations and more intact apoptotic pathways. One lesson derived from the genomic sequencing of breast, colon, and pancreatic cancer is that the number of mutations in these solid tumors is quite large and often not overlapping between cancers. Individual tumors contained an average of 90 mutant genes, 11 of which seemed to be oncogenic, among the 13,023 genes sequenced in 11 breast and 11 colorectal cancers (59). Perhaps such an explanation underlies sensitivity to epigenetic therapies in general as well as the more successful outcome of classical cancer chemotherapies in lymphomas and other hematologic malignancies. If solid tumors have intrinsic mechanisms of resistance, then increasing the dose or adding a second agent that exploits any of the molecular effects of the HDI might help overcome resistance in solid tumors.

### Combination Therapy

Combination therapies employing DNMT inhibitors and HDIs together or with other agents are being pursued clinically. The possibility to optimally re-express methylated genes following treatment with a DNMT inhibitor followed by an HDI has been the basis for combined epigenetic therapies. Given the in vitro evidence for synergy in such combinations, this remains an area of active study, with initial trials focusing on hematologic malignancies (44, 60). Eventually, randomized trials will be required to establish whether the clinical outcomes correlate with preclinical synergy for these agents. Whether combination epigenetic therapy will improve efficacy in solid tumors remains to be determined.

In addition to combined epigenetic therapies, regimens using HDIs in combination with established agents are actively being pursued. Rationally conceived combination therapies with HDIs can be considered to fit one of several strategies: (1) to counter the molecular effects of HDIs that abrogate their efficacy, (2) to extend the molecular effects of the HDI, or (3) to exploit the molecular effects of the HDI. Bots and Johnstone elaborate on the development of combination therapies (60). An example of the first strategy is the addition of flavopiridol to prevent the induction of p21, which causes cell cycle arrest and to some extent limits the activity of the HDI (61, 62). An example of the second type of combination is to add HDIs to death receptor ligands such as TRAIL that can synergize to increase apoptosis (63). Several examples of the third strategy involve the depletion of Hsp90 client proteins that follows acetylation of Hsp90 and impairment of its chaperone function. Hsp90 acetylation results in polyubiquitinylation of client proteins, such as ABL, Bcr-Abl, EGFR, ErbB2, c-Raf, c-kit, FLT3, ER, and AKT, and their degradation by the proteasome (64, 65).

One could predict that this last strategy will be the most promising, allowing selection of tumor type based on pre-existing dysregulation of client proteins. Thus, combining an HDI with trastuzumab should generate a synergistic anticancer effect in HER2 over-expressing breast cancer, a concept that has been confirmed in vitro and is undergoing clinical testing in breast cancer (66, 67). A similar concept should hold true for EGFR inhibitors in lung cancer, in which gefitinib and erlotinib effectively block signaling from mutant EGFR, but from which resistance eventually emerges (68). Other tyrosine kinase inhibitors could synergize with HDIs by this mechanism: imatinib, nilotinib and dasatinib could be potentiated in chronic myelogenous leukemia by loss of Bcr-Abl (69). The list of potentially active combinations grows very long employing this strategy.

In vitro data also suggest synergy with traditional cytotoxic chemotherapeutics (60). It has been previously noted that HDIs sensitize cancer cells to topoisomerase targeted therapies (70). Initially this was related to a direct interaction of HDAC 1 and 2 and topoisomerase II enzymes (70), but it was later noted that HDIs also increase expression of topoisomerase II (71). Moreover, it was found that acetylation of Ku70 (such as that following treatment of cells with an HDI) results in a functional impairment of DNA repair leading to an increased sensitivity to DNA damaging agents (72). An alternate effect of the acetylation of the cytoplasmic form of Ku70 is the release of Bax, a pro-apoptotic protein, which when released from Ku70 translocates to the mitochondria and triggers apoptosis through the release of cytochrome c and caspase-activation (73).

### Caveats about Combination Therapy

An important question is whether combination studies can succeed in which there has been no single agent activity. Few examples of anticancer therapy can be identified in which an agent has failed as monotherapy but has succeeded in combination. However, DNMT inhibitors and HDIs may be different in that a molecular effect that is insufficient to cause cell death could nonetheless set up an environment in which another agent could exert a more potent effect. Thus, the critical point for development of these agents in combination therapy will be to obtain clinical material to test for the molecular effect that is to be exploited in the combination. Only with this information can the results of combination trials be correctly interpreted.

Arguing for combination therapy as a means to fully exploit the activity of epigenetic therapy leads to a related issue, that of scheduling: determining how best to schedule the combined therapy. A chemotherapeutic-inducing cell cycle arrest could not be combined with a DNMT inhibitor. Similarly, the p21 induction and cell cycle arrest that results from HDAC inhibition could lead to resistance with most conventional cytotoxic agents that require cell division for activity. Theoretically considerable synergy from an HDI can be considered to fit one of several strategies: (1) to counter the molecular effects of HDIs that abrogate their efficacy, (2) to extend the molecular effects of the HDI, or (3) to exploit the molecular effects of the HDI. Bots and Johnstone elaborate on the development of combination therapies (60). An example of the first strategy is the addition of flavopiridol to prevent the induction of p21, which causes cell cycle arrest and to some extent limits the activity of the HDI (61, 62). An example of the second type of combination is to add HDIs to death receptor ligands such as TRAIL that can synergize to increase apoptosis (63). Several examples of the third strategy involve the depletion of Hsp90 client proteins that follows acetylation of Hsp90 and impairment of its chaperone function. Hsp90 acetylation results in polyubiquitinylation of client proteins, such as ABL, Bcr-Abl, EGFR, ErbB2, c-Raf, c-kit, FLT3, ER, and AKT, and their degradation by the proteasome (64, 65).

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report greater sensitivity is only observed when the HDI is given first (76). These results may again be cell-type specific.

Finally, when considering combination therapy with conventional chemotherapy agents, it should be recalled that HDIs have direct effects on the promoters of ABC transporters. MDR-1/ABCB1, which encodes P-glycoprotein, is uniformly increased at the promoter level following HDI exposure (47, 77, 78). ABCG2, a drug efflux transporter with a differing spectrum of activity, is also frequently up-regulated following HDI exposure in in vitro models (79). To the extent these multidrug transporters can reduce intracellular drug concentrations in vivo, then efficacy of a chemotherapy substrate could be reduced in combination with the HDI.

**Epigenetic Therapies Reach Main Street**

Although the cancer community has become acutely aware of epigenetic aberrations in cancer, and therapies aimed at normalizing the epigenetic profile of the cancer cell are now available and in the clinic, much is yet to be discovered. Although we know the cellular target for current epigenetic therapy, we do not really understand how these targets play a role in cancer, or what we could actually expect if we had a drug that fully normalized the epigenetic profile. Would this in effect be the differentiation therapy that was imagined from the 1970s onward? Would we, in fact, completely reverse the malignant phenotype, especially in solid tumors? It seems doubtful given the array of oncogenic mutations now understood to be involved in human cancer.

It is interesting to note that both of our clinically active classes of epigenetic therapies involve inhibition of enzymes that work to silence DNA. As enzyme inhibitors, the target is clear; only their specificity for a given HDAC or DNMT remains to be worked out. However, the critical downstream events that follow enzyme inhibition and determine cell death are not elucidated, with HDIs even less so than the DNMT inhibitors. Both hypomethylating agents and HDIs invoke multiple changes in cells that sometimes cause cell death and sometimes promote cell survival. Downstream events resulting from these therapies may provoke disparate results. Both agents have numerous potential avenues of synergy in treating cancer. One of the most compelling strategies is to use the epigenetic therapies as radiation sensitizers, as we consider DNA to be a target of both (80).

It must be recognized that any discussion of epigenetic therapies for the future must go beyond DNA methylation and histone deacetylase inhibition. Because epigenetic modifications have many components, it is exceedingly likely that other as yet unidentified therapeutic targets exist. It is critical that investigators evaluate such targets, develop active compounds, and then establish proof of concept clinical trials in a much shorter time frame than it took to develop our current epigenetic therapies. Potential “other” epigenetic therapies include histone acetyltransferase (HAT) inhibitors (recall that as many genes are down-regulated following HDAC inhibition as up-regulated), inhibitors of HDAC class III enzymes such as sirtuin inhibitors, EZH2 antagonists to prevent perpetuation of DNA methylation, and histone methyltransferase inhibitors (81–83). Although HAT activity provides the acetyl groups that allow gene expression following inhibition of HDACs, it is interesting to note that in some cancer models increased HAT activity has been associated with tumor progression and inhibition with an anticancer effect (81).

Determining and exploiting clinically beneficial effects is a major goal for the future. The pleiotropic nature of the molecular effects of chromatin modifying agents is perhaps not surprising, given the ubiquitous nature of the methyl transferase and histone deacetylase enzymes they target. It remains to be determined if more specific agents will be more or less beneficial, or enjoy a greater therapeutic window. We do know that most of our successful anticancer agents have multiple mechanisms of action.

Further progress also requires the development of better methods to detect the epigenetic modifications and a clearer understanding of factors that drive these changes. The inclusion of epigenetics in the NIH roadmap indicates recognition of its role in human disease including but certainly not limited to malignancy. At least two large consortia have taken the lead in attempting to map the epigenetic modifications in detail over a defined region of the human genome. The American Association of Cancer Research has launched the Human Epigenome Task Force, whereas the Human Epigenome Project (HEP) has its origins in the United Kingdom. HEP has focused initially on DNA methylation profiling whereas the Human Epigenome Task Force will also examine histone modifications associated with histone acetylation patterns in addition to DNA methylation. In 2006, HEP released data including 1.9 million CpG methylation values, obtained from the analysis of 2,524 amplicons across chromosomes 6, 20, and 22 in 43 samples (84). This work is in its infancy but has the potential to give us many more clues to the role of epigenetics in cancer, beyond a lifetime of individual gene promoter studies.

With the approval of three epigenetic agents and additional ones in the pipeline, we have the beginnings of a toolbox for manipulating the epigenome. Welcome to Main Street.

**Disclosure of Potential Conflicts of Interest**

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Epigenetic Therapies Reach Main Street

Although the cancer community has become acutely aware of epigenetic aberrations in cancer, and therapies aimed at normalizing the epigenetic profile of the cancer cell are now available and in the clinic, much is yet to be discovered. Although we know the cellular target for current epigenetic therapy, we do not really understand how these targets play a role in cancer, or what we could actually expect if we had a drug that fully normalized the epigenetic profile. Would this in effect be the differentiation therapy that was imagined from the 1970s onward? Would we, in fact, completely reverse the malignant phenotype, especially in solid tumors? It seems doubtful given the array of oncogenic mutations now understood to be involved in human cancer.

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Epidigenetic Modifiers: Basic Understanding and Clinical Development


Clinical Cancer Research

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Richard L. Piekarz and Susan E. Bates


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