Methods for Activity Analysis of the Proteins that Regulate Histone Methylation

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Abstract: The enzymes that regulate histone methylation states and the protein domains that recognize methylated histone residues have been implicated in a number of human diseases, including cancer, as a result of their ability to affect transcriptional changes by altering chromatin structure. These proteins are recognized as potential therapeutic targets for the treatment of diseases associated with epigenetic disruption; however, few inhibitors of their activity have been identified. The majority of histone demethylase and methyltransferase enzyme inhibitors have been discovered on the basis of their structural similarity to substrates or known inhibitors of enzymes with analogous mechanisms. The general lack of potency and specificity of these compounds indicates that novel chemotypes are needed to address the large number of recently discovered histone-modifying enzymes. High-throughput screening (HTS) allows rapid testing of chemically diverse small molecule libraries, provided assays amenable to HTS exist. Here we review the biochemical and cellular assays available for testing the proteins and enzymes that regulate histone methylation. Progress in the development of high-throughput, sensitive, and robust assays will enable discovery of small molecules for epigenetic therapy.

Keywords: Epigenetics, lysine methyltransferase, arginine methyltransferase, lysine demethylase, reader domain, histone methylation, enzyme assays, protein-protein interactions, small molecule inhibitors.

INTRODUCTION

Post-translational modifications (PTMs) of histones, including acetylation, methylation, ubiquitination, and phosphorylation, affect the transcription of associated genes by altering local chromatin structure and the accessibility of transcription factors [1]. These epigenetic marks also act as beacons in the recruitment of enzymes that further modify the landscape of histone PTMs or DNA methylation, or that directly (de)stabilize chromatin structure [2]. Both the enzymes that regulate the addition or removal of histone modifications and the proteins harboring domains that recognize specific PTMs have been implicated in a variety of human diseases, including cancer.

The most thoroughly characterized histone modification is lysine acetylation which generally leads to an increase in transcriptional activity following the loss of lysine’s positive charge with acetylation and decreased electrostatic interaction with the negatively-charged DNA backbone [3]. Histone deacetylase (HDAC) inhibitors have been used in the clinic to promote hyperacetylation of histone tails and reactivate aberrantly silenced tumor suppressor genes [4]. The functional consequence of histone methylation is more complex, with methylation of both arginine and lysine possible, and transcriptional changes dependent on the site and degree of methylation. For example, methylation at H3K9, H3K27, and H4K20 typically leads to repressed transcription, whereas methylation at H3K4 and H3K36 results in enhanced transcriptional activation [5].

Research on the mechanisms and regulation of histone methylation has intensified in the last decade in response to the discovery that histone methylation is impermanent and the identification of histone demethylases. This surge of activity surrounding epigenetics has resulted in the discovery of greater than 50 methyltransferase enzymes, 30 histone demethylases, and numerous protein domains that recognize and bind methylated histone residues. To study these PTMs and the mechanisms of enzymes that regulate them has necessitated the development of various biochemical and cell-based assays. In addition, validating the involvement of many of these proteins in human disease has generated considerable interest in the discovery of small molecule inhibitors of histone-modifying enzymes and of compounds that disrupt the interactions between histones and the domains that bind specific PTMs, the so-called “readers” of the histone code.

Here we review recent advances in methods to study histone methylation with a focus on the discovery of small molecule modulators of histone methyltransferases, demethylases, and reader domains as potential therapeutic agents in the treatment of cancer and other diseases associated with epigenetic deregulation. To date, no fewer than six high-throughput screens of epigenetic targets have been performed with more than 200,000 compounds and published on the open-access PubChem database (http://pubchem.ncbi.nlm.nih.gov/), demonstrating the need for robust high-throughput assays and techniques for rapid characterization of the downstream effects of compounds that influence hist-
tome PTMs. While acknowledging the importance of histone acetylation, phosphorylation, and other PTMs in the development of a spectrum of human diseases, we direct our attention here to the rapidly-evolving field of histone methylation.

**METHYLTRANSFERASE ASSAYS**

**Radiotracer Assays**

Radioactivity-based assays to measure histone methyltransferase activity are sensitive, easy to configure, and applicable to a diverse range of enzymes. Filter binding assays measure transfer of tritiated methyl groups from $[^3H]$-SAM to a substrate, followed by removal of unreacted $[^3H]$-SAM and retention of radiolabeled-product by filtration and subsequent scintillation counting (Fig. 1A). The fungal mycotoxin metabolite chaetocin (3,6-epi-dithio-diketopiperazine) was one of the first methyltransferase inhibitors discovered by screening 3,000 compounds against the *Drosophila melanogaster* SU(VAR)3-9 lysine methyltransferase using a radioactive histone H3 peptide filter binding assay (Table 1) [6]. Filters assembled in 96- and 384-well vacuum plates have increased the throughput of this assay. Other plate-based techniques include immobilization of a biotinylated peptide substrate in an avidin/streptavidin-plate and removal of free $[^3H]$-SAM with washing steps. The washing steps

Table 1. Methods Used in the Discovery of Small Molecule Inhibitors of Histone Methyltransferase and Demethylase Enzymes

| Method | Compound | Target | Secondary Assays |
|--------|----------|--------|------------------|
| **Radiotracer Assays** | | | |
| Filter Binding | Chaetocin [6] | SU(VAR)3-9 | Biochemical and cellular immunoblot, MS |
| | Pyrazole Amide 7b [68] | PRMT4 | PRMT1, PRMT3 selectivity (filter binding) |
| $[^14C]$-Labeling | Sinefungin [69] | N.D. | N.D. |
| **Antibody-Based Assays** | | | |
| DELFIA | BIX-01294, BIX-01338 [13, 32] | G9a, GLP | MS, Cellular expression analysis, ChIP |
| ELISA | AMI-1, AMI-5 [12] | Hmt1p, PRMT1 | Filter binding, ERα-reporter gene assay |
| **Coupled Enzyme Assays** | | | |
| FDH-Coupled | Myricetin, β-lapachone [45] | JMJD2E | MS |
| | 8-hydroxyquinolines [43] | JMJD2E, 2A | MS, FRAP |
| **Mass Spectrometry (MS)** | | | |
| | Disulfiram or selenium derivatives [70] | JMJD2A | Zn(II)-ejection assay |
| **Fluorescence Polarization (FP)** | | | |
| | Stilbamidine, allantodapsone, RM-65 [14, 71] | RmtA, PRMT1 | TRF, ERα -reporter gene assay, cellular IHC |
| **Crystallography/Modeling/Virtual Screening** | | | |
| | UNC-0321 [17] | G9a, GLP | SAHH-coupled assay, AlphaScreen, FP, MCE |
| | E70 [72] | GLP | MS, ITC, cellular expression analysis |
| | Tranylcypromine-Lys conjugates [73] | LSD1 | HRP-coupled assay, cellular immunoblot |
| | 2,4-PDCA [33] | JMJD2E, 2A | FDH-coupled assay, MS, crystallography |
| | 4-dimethyl-amino-benzyl-NOG [74] | JMJD2A, 2C, 2D | Biochemical and cellular immunoblot |
| | N-oxalyl-D-tyrosine derivatives [75] | JMJD2A | MS, FDH-coupled assay, crystallography |
| **Structural or Mechanistic Similarity** | | | |
| | Tranylcypromine [48] | LSD1 | Immunoblot, ChIP, transcriptional analysis |
| | Lys4-cyclopropyl and propargylamine H3 peptide derivatives [50] | LSD1 | HRP-coupled assay, MS |
| | Bisguanidine and biguanide polyamine analogs [54] | LSD1 | Biochemical and cellular immunoblot, transcriptional analysis, ChIP |
| | N-oxalylglycine (NOG) [76] | JMJD2E, 2A, 2B | Immunoblot, FDH-coupled assay |
| | Succinic acid [77] | JMJD2D | Biochemical and cellular immunoblot |

N.D., not determined.
may be eliminated with the use of avidin/streptavidin-coated FlashPlates that contain a thin layer of scintillant bound to the walls of the microplate wells where the signal is based on isotope proximity [7].

Transfer of tritiated methyl groups from $[^3]H$-SAM to a bead-bound substrate provides a homogeneous, sensitive method for rapidly measuring methyltransferase activity (Fig. 1B). Scintillation proximity assays (SPA) utilize microscopic beads containing scintillant. The interaction of these beads with β-particles generated by the radioactive decay of tritium releases photons that may be measured with scintillation counters or charge-coupled device (CCD) imagers. A variety of SPA bead formats allows utilization of various methyltransferase substrates, from biotinylated histone peptides to nucleosomés. The lack of a separation step and intrinsic sensitivity of the assay make SPA techniques amenable for HTS. However, the use of radioactivity is undesirable due to health risks as well as the cost of radiolabeled substrate and waste disposal. Thus the development and use of non-radioactive assays should be utilized whenever possible.

**Coupled Enzyme Assays**

Simple substrate-to-product conversion is difficult to monitor in histone methyltransferase reactions where the spectral differences in SAM and SAH are insufficient for direct monitoring. Several coupled assays have been used to measure the SAH product of methyltransferases (Fig. 1C). The cysteine-free SAH hydrolase of *Sulfolobus solfataricus* generates a free thiol in homocysteine that may be measured with thiol-sensitive fluorophores [8], including the maleimide derivatives ThioGlo 1 [methyl-10-(2,5-dioxo-2,5-dihydro-1H-pyrrol-1-yl)-9-methoxy-3-oxo-3H-benzo[f]chromene-2-carboxylate] and CPM (7-diethylamino-3-(4′-maleimidylphenyl)-4-methylcoumarin) [8, 9]. The free thiol of homocysteine may also be measured by reaction with DTNB (5,5′-dithiobis-2-nitrobenzoic acid) to form the colored product TNB (5-thio-2-nitrobenzoic acid). Homocysteine may also be generated from SAH through the consecutive actions of SAH nucleosidase (SAHN) to generate adenine and $S$-ribosylhomocysteine and LuxS ($S$-ribosylhomocysteinate) to cleave the latter intermediate to form Hcy [10]. Alternatively, the adenine product of SAHN may be utilized by adenine deaminase to form hypoxanthine, where a decrease in UV-absorbance is monitored at 265 nm [11]. The SAHH/ThioGlo assay described by Collazo et al. [8] is likely the most straightforward and sensitive of these methods to measure SAH production, considering the single coupling enzyme and fluorescence detection. However, assay components must be carefully selected to minimize background fluorescence due to the reaction of free thiols with ThioGlo 1, and chemical blocking of free thiols of the substrate or enzyme may be considered. Coupled enzyme assays are inherently more difficult to optimize and susceptible to false positives when seeking enzyme inhibitors. Nonetheless, the removal of SAH in the coupling assays described here is advantageous for relief of SAH product inhibition, a characteristic common to histone methyltransferases.

**Antibody-Based Techniques**

Several biochemical assays have been developed that rely on antibody recognition of specific post-translational modifications. Enzyme-linked immunosorbent assays (ELISA) for histone methyltransferases involve immobilization of biotinylated histone peptide substrate on an avidin-coated microplate. Following enzymatic methylation of Lys or Arg residues on the immobilized peptide, a primary antibody specific to the PTM is captured on the plate. A secondary antibody conjugated with horseradish peroxidase (HRP) is used for chemiluminescence detection. The first protein arginine methyltransferase (PRMT) inhibitor AMI-1, a symmetrical sulfonated urea, was identified in an ELISA-based HTS assay of 9,000 diverse compounds using fungal Hmt1p (also known as Rmt1p) and the hNRNP protein Npl3p (Table 1) [12]. AMI-1 is a non-specific arginine methyltransferase inhibitor, although it does not affect methylation of lysine.

A modern version of the ELISA is the dissociation-enhanced lanthanide fluorescent immunoenassay (DELFIA) where the secondary antibody is labeled with a lanthanide chelate (typically europium) instead of HRP, and product formation is measured with time-resolved fluorescence (TRF). This technique was used to identify the first inhibitor of a histone lysine methyltransferase that was not competitive with the SAM cosubstrate. BIX-01294 was discovered in a screen of 125,000 compounds as a selective inhibitor of the G9a lysine methyltransferase (IC$_{50}$ 2.7 μM) and was competitive with the histone H3 peptide substrate (Table 1) [13]. BIX-01338 was also identified in this screen as a G9a inhibitor (IC$_{50}$ 4.7 μM), but it also inhibited other Lys and Arg methyltransferases and was SAM-competitive. Inhibitors of the arginine methyltransferase PRMT1 were also identified using a DELFIA assay to screen compounds selected by iterative virtual screening of the National Cancer Institute (NCI) diversity collection [14]. Two of the compounds active in the PRMT1 in vitro assay, allantoadapson and stilbamidine, were validated by demonstrating hypomethylation of H4R3 in HepG2 cells and inhibition of ERα-transcriptional activation in a cell-based reporter assay.

Antibody detection has also been used in a homogeneous assay utilizing AlphaScreen technology to measure histone G9a methyltransferase activity [15]. This dual bead-based technology utilizes laser excitation of donor beads, releasing a flow of singlet oxygen to generate chemiluminescence emission from acceptor beads in close proximity (Fig. 1D). Acceptor beads coated with secondary antibody are coupled to specific antibody detection of enzyme-modified substrates. AlphaScreen was used in the structural optimization of the BIX-01294 inhibitor of G9a, and IC$_{50}$ values obtained tracked closely with those measured in a SAHH/ThioGlo 1 coupled assay [16, 17]. These studies led to the discovery of UNC0224 and UNC0321, the most potent G9a inhibitors identified to date (Table 1). The advantage of this assay compared to ELISA-based methods is the homogeneous nature of AlphaScreen where elimination of washing and substrate immobilization steps leads to an increased assay throughput. Importantly, the AlphaScreen assay was readily miniaturized to 1,536-well plate format allowing for its use in screening large compound collections [15].

AlphaScreen assays to measure histone methyltransferase activity may utilize direct coupling of the acceptor bead to the primary antibody in place of a secondary anti-immunoglobulin donor bead plus primary antibody. Some methyl mark antibodies directly conjugated to acceptor
beads are commercially available (http://las.perkinelmer.com/). In vitro AlphaScreen assays most commonly use streptavidin-coated donor beads to bind biotinylated peptide substrates. However, donor beads may be conjugated with antibodies that recognize histone substrates at a site distinct from that modified by the methyltransferase or demethylase enzyme. This sandwich-based approach is suitable for enzymes with no or low activity towards peptide substrates as well as for cell-based quantitation of histone-modifying enzyme activity.

Time-resolved fluorescence resonance energy transfer (TR-FRET), like AlphaScreen, is a proximity based technique that may be used to measure enzymatic activity of histone-modifying enzymes. TR-FRET combines time-resolved fluorescence (TRF) and FRET in a homogeneous format where fluorescence emission of a donor dye overlaps with the excitation spectrum of an acceptor dye. Whereas AlphaScreen donor and acceptor molecules may be separated by up to 200 nm, the distance of interaction is typically limited to 1-10 nm for TR-FRET. To date, FRET has only been used to measure inhibition of HDAC activity in cells using a baculovirus-expressed GFP-p53 fusion protein [18]. A terbium-labeled acetyl-p53 (Lys382) antibody was used to measure activity of the p300 histone acetyltransferase (HAT) using the non-histone GFP-p53 substrate where GFP is the
FRET acceptor. We note that if an endpoint assay based on antibody detection is desired, the traditional method of stopping the enzymatic reaction by addition of acid will not be usable: instead, an addition of EDTA at a millimolar concentration can be used as an alternative.

The limitations of these techniques lie in the availability of specific antibodies, particularly in assays that require discrimination between mono- and dimethylated or di- and trimethylated residues [19]. Examination of more than 200 antibodies raised against 57 different histone modifications found that 25% of these failed specificity testing by Western blot and dot blot analysis, and 20% of the specific antibodies were deemed unsuitable for chromatin immunoprecipitation (ChIP) analysis [20]. The high failure rate prompted the authors to create a website for posting these and new results of antibody specificity tests for the detection of epigenetic marks (http://combio.med.harvard.edu/antibodies/). Where specific antibodies are not available, the use of tagged chromatin binding domains to measure changes in levels of specific post-translationally modified residues has been suggested (Wigle, T.J., personal communication), although it is unclear whether the micromolar affinities typical of reader domains for their cognate epigenetic mark would be sufficient for such applications. Additional complications in attempting to quantify PTMs with reader domains in a cellular environment may result from the observation by Fuechs et al. that neighboring PTMs can affect the affinity of chromatin-associated domains for a particular epigenetic mark [21]. Using a library of histone peptides with single or multiple PTMs immobilized in a peptide array, both enhanced and diminished binding of several reader domains were observed for combinations of PTMs. Using the same peptide array, the authors also found the specificity of some commercial antibodies raised against a specific epigenetic mark were affected by other PTMs present on the peptide bait [21].

Capillary Electrophoresis

The principles of capillary electrophoresis applied to a microfluidic chip (MCE) allow separation of nanoliter-sized samples based on the charge-to-mass ratio of the components of interest. The loss of lysine’s positive charge with acetylation results in facile electrophoretic separation of substrate and product peptides and offers a direct method for measuring HAT or HDAC activity. This type of electrophoretic mobility shift assay has been used to profile inhibitors of HDAC and HAT enzymes, including PCAF and GCN5L2 [22, 23]. A similar approach may be employed for changes in phosphorylation of histone peptides.

Methylation of lysine, however, does not change the positive charge of this amino acid, rendering a direct electrophoretic separation impossible. To address this limitation, methods originally developed for methyllysine mapping of histones by mass spectrometry (MS) have recently been applied to MCE. These techniques use the endoproteinase LysC to selectively cleave at the carboxyl side of unmodified lysine residues, leaving methylated lysine polypeptides intact (Fig. 1E). Such a histone mark-dependent cleavage allows for the production of fluorescent peptides of different charge-to-mass ratios. The utility of this method was demonstrated using LysC and fluorescent-labeled histone peptides for inhibitor screening and mechanism of action studies of the G9a methyltransferase and the LSD1 lysine demethylase [24]. The G9a MCE assay was used to determine the Morrison K i (63 pM) of UNC0321, the 2,4-diamino-7-aminoalkoxy-quinazoline compound designed from the BIX-01294 chemical scaffold (Table 1) [17]. The multiplicity of Lys residues in the N-terminal tails of histones, however, complicates generation of suitable peptides upon LysC cleavage and may require mutation or chemical modification of non-substrate Lys residues in substrate peptides. In addition to measuring lysine methyltransferase activity, protease-coupled MCE may be applicable to monitoring Arg methylation with the use of the ArgC endopeptidase [24].

Mass Spectrometry

Mass spectrometry (MS) has been an important tool in identifying novel PTMs as well as characterizing activities of histone-modifying enzymes [25]. The majority of this work has been performed using a bottom-up approach where proteins are digested into small peptides prior to MS analysis. Tandem mass spectrometry (MS/MS) is used to map the site of modification and reconstruct protein identity using sequence information. The major disadvantage to this approach is that information on the combinatorial nature of histone modifications is lost. A top-down approach, on the other hand, refers to MS analysis of intact proteins to yield information on combinations of modifications and their relative abundance [26]. This is particularly useful in the analysis of heterogeneous mixtures of histones, for example in tissues and cell culture.

An intermediate approach to these techniques was constructed to mitigate the need for a high-resolution mass analyzer in top-down MS while retaining information on the interplay between PTMs on a single histone polypeptide. In middle-down MS, limited proteolysis of histones with endopeptidases such as GluC or AspN generates polypeptides of up to 50 residues [27]. As the majority of PTMs are located on the N-terminal tails of these proteins, information on the complex spectrum of histone PTMs is gained with middle-down MS. This MS proteomics method has been used for global assessment of combinatorial PTMs and their tissue-specific abundance [28].

Quantitative MS has been used to measure enzymatic activity of histone-modifying enzymes. In vitro approaches generally target specific PTMs and use MS peak intensities of high resolution, well-defined peptides for relative quantitation of product formation (Fig. 2A). This label-free method has been used to measure activities of numerous histone-modifying enzymes [29-31] and to determine inhibitor potency [32, 33]. An internal standard for normalization that consists of a nonvarying, unmodified peptide may be used in conjunction with peptide intensity profiling to measure changes in histone modifications. Advances in the throughput of MS-based quantitation assays have increased the capacity for inhibitor screening in these sensitive and selective assays [34]. For example, inhibition of the LSD1 histone demethylase has been measured using an eight-way parallel, staggered flow injection MS from samples in 384-well plate format [35]. Other quantitative methods for measuring histone modifying enzymes utilize various labeling techniques, including stable isotope labeling with amino acids in cell culture (SILAC) [36], enzymatic incorporation of 18O [37],
iTRAQ [38], and chemical tagging of reactive side chains or groups with stable isotopes [39]. Stable isotopes may also be incorporated into methylated histone residues using [13CD3]-labeled SAM in place of the SAM methyl donor [40].

**Binding Assays**

Given the low turnover of many histone methyltransferases, assays that measure displacement of SAM or a peptide substrate may be used to further characterize inhibitors or to select potential inhibitors. Fluorescence polarization (FP) assays measure changes in fluorescence anisotropy in the presence of compounds that displace fluorescently-labeled cognate peptides from the methyltransferase active site. Exploration of the SAR around the G9a inhibitor BIX-01294 was aided by measuring displacement of a fluorescently labeled histone H3 peptide upon compound binding [16]. A similar FP assay was developed to measure binding of histone H4 peptides to PRMT1 [41], and was used for the mechanistic analysis and comparison of binding affinities for published PRMT1 inhibitors (Table 1) [42]. FP assays, however, require large amounts of enzyme and may not be appropriate for large-scale discovery of peptide-competitive inhibitors.

**Cell-Based Methyltransferase Assays**

The options for cell-based assays to measure changes in posttranslational modifications of histone tails are limited and primarily semi-quantitative. The multiplicity of methyltransferases and a lack of characterization of their substrates make it difficult to assign activity to a particular enzyme. Transcriptional changes of genes known to be affected by histone methylation at a particular site are frequently used as surrogate markers of methyltransferase activity in cells [14]. ChIP at methyltransferase target genes may also be used to measure methylation changes at promoter-proximal sites. Measurements compared before and after knockdown of the methyltransferase by genetic or RNAi-based methods assist in validation of the inhibitor’s target.

Cell-based assays to measure changes in a particular epigenetic mark include fluorescence immunostaining with specific antibodies in live cells [43]. A similar technique is standard ELISA with fixed, permeabilized cells using a primary antibody directed against a specific PTM and a europium-labeled secondary antibody for TRF detection [14]. As discussed above, quantitation of methylation at a specific site is complicated by the availability of specific antibodies.

**HISTONE DEMETHYLASES**

The JmjC domain demethylases utilize non-heme iron (II) in a 2-oxoglutarate (2-OG)–dependent oxygenation mechanism to demethylate lysine residues of histone tails [29]. Their aberrant activities are associated with cancer, inflammation, and X-linked mental retardation [44]. Reported inhibitors of the JmjC demethylases include 2,4-pyridinedicarboxylic acid (PDCA) and hydroxamic acid derivatives, small compounds of micromolar potency that are competitive with 2-OG (Table 1) [33]. The coupling enzyme formaldehyde dehydrogenase (FDH) has been used to oxidize the demethylase product formaldehyde with concomitant reduction of its NAD⁺ cofactor to monitor the change in UV absorbance or fluorescence of NADH (Fig. 2B), thus producing a convenient real time kinetic assay for JmjC activity [45, 46]. AlphaScreen has also been used to measure JMJD2E H3K9me3 demethylase activity in vitro. Using a panel of 2,4-PDCA analogs, the AlphaScreen assay yielded similar IC₅₀ values for JMJD2E inhibition compared to those obtained with the FDH-coupled assay [19].

The continuous and homogenous nature of the FDH-coupled assay allowed its straightforward miniaturization to 1,536-well plate format [45] and its subsequent use in a high-throughput screen of 236,000 compounds for inhibitors of the JMJD2E enzyme using an H3K9me3 peptide substrate [43]. Of the chemical scaffolds designated as active in the screen, optimization of the 8-hydroxyquinolines was performed to generate a 5-carboxy-8HQ cell-permeable compound with 200 nM potency in the enzymatic assay [43]. Compared to 2,4-PDCA, the 5-carboxy-8HQ was more selective for JMJD2 enzymes than the 2-OG–dependent hypoxia inducible factor prolyl hydroxylases PHD2 and FIH. The crystal structure of this compound with the homologous JMJD2A protein indicates that 5-carboxy-8HQ disrupts the binding of the active site Fe(II).

A second family of histone demethylases is the lysine specific demethylase (LSD) family, comprised of LSD1 and LSD2. These are flavin adenine dinucleotide (FAD)-dependent amine oxidases that demethylate di- or mono-methylated H3K4 [47]. The rapid turnover of LSD enzymes compared to JMJ enzymes enables easier detection of demethylase activity. While LSD enzymes generate formaldehyde and may be coupled to FDH, as outlined above, more robust assays exist that measure the hydrogen peroxide product (Fig. 2B). Horseradish peroxidase (HRP)-coupled assays use H₂O₂ to oxidize a number of HRP substrates including 4-aminoantipyrine, AmplexRed (10-acetyl-3,7-dihydroxyphenoxazine), and luminol for detection using absorbance, fluorescence, or chemiluminescence, respectively.

Of the reported LSD inhibitors, the majority are monoamine oxidase (MAO) inhibitors used in the clinic for antidepressant therapy, including pargyline and tranylcypromine [48, 49]. These are mechanism-based inhibitors that form covalent adducts with the FAD cofactor of LSD1/2 [50, 51]. Libraries constructed around these MAO inhibitor scaffolds have improved inhibitor potency and selectivity for LSD over MAO enzymes [51, 52]. Biguanide and bisguanidine polyamine analogues have also been described as LSD1 inhibitors and result in reexpression of aberrantly silenced tumor suppressor genes in a human colon cancer cell line [53, 54].

**HISTONE CODE READER DOMAINS**

Protein domains that bind to and recognize posttranslational modifications of histones are referred to as readers of the histone code. These domains include the Royal family of proteins (chromo-, tudor-, malignant brain tumor-domains, others) that recognize PTMs of Lys residues and bromodomains that recognize acetylated Lys [1]. Histone code readers function as adaptors to recruit proteins for further modification of chromatin structure [2]. In addition to recruitment of histone-modifying enzymes, domains that bind specific epigenetic marks link histone modification to changes in DNA methylation [55]. The important role of these recognition motifs and their increasing association with disease
make compounds that disrupt these protein-protein interactions a novel strategy for targeting epigenetic diseases.

The multiplicity of histone code readers suggests diversity in substrate specificity and, indeed, the interactions of these protein domains with the large spectrum of histone tail PTMs demonstrate a high level of specificity [56]. Crystal structures of bromodomains, a recognition domain family comprised of at least 57 members, indicate diversity in loop regions that dictate substrate specificity despite a conserved binding channel [57]. Thus, characterization of disease-associated epigenetic marks and reader domain specificity is crucial in targeting these binding domains. Peptide arrays coupled with mass spectrometry are used to bait, capture, and identify chromatin-associated domain binding partners [58]. Combining this technique with SILAC to expose “heavy” cell extracts to post-translationally modified peptide bait and “light” cell extracts to unmodified peptide, specific binders are distinguished from background binders [36]. Characterization of transcriptional target genes may be achieved with ChIP-Seq and MS-based sequencing, where the histone marks bound by these reader domains should overlap with the target gene mapping [59].

Two small molecule compounds that disrupt binding of a BRD4 bromodomain to acetylated histones were recently published in parallel, providing an excellent proof of concept for targeting bromodomains [57, 60]. The very similar triazole-diazepine-fused ring compounds, (+)-JQ1 and I-BET, bind stereoselectively to the bromodomain in the acetyl-lysine binding channel with equilibrium affinities in the range of 50 – 100 nM. Filippakopoulos et al. demonstrated (+)-JQ1-dependent cellular displacement of BRD4 from chromatin and antitumor activity in a mouse xenograft model of midline carcinoma [57]. Nicodeme et al. investigated the immunomodulatory effects of I-BET and found that it inhibited transcription of inflammatory genes in activated macrophages and attenuated the death of mice induced by septic shock [60]. Despite the remarkable success in developing these small molecule inhibitors of a bromodomain protein-protein interaction, very little is known about the chemical space surrounding other histone code readers, particularly regarding methyllysine or methylarginine marks, and high-throughput assays are needed to identify novel chemical scaffolds.

Several approaches have been taken to identify small molecule disruptors of histone code readers. Fluorescence polarization has been used to measure binding affinities of histone code readers to fluorescein-labeled peptides (Fig. 3A) [58, 61, 62]. However, these low affinity micromolar interactions make an FP assay format unsuitable for high-throughput identification of small molecule inhibitors of peptide-protein binding. An AlphaScreen assay was developed using biotinylated histone peptides synthesized with an epigenetic mark of interest incorporated to quantify binding.

Fig. (2). Histone demethylase assay formats. (A) Quantitative mass spectrometry measures demethylation of histone peptide substrates by identification and relative quantitation of peptides by MS/MS. Here, LSD1 demethylates dimethyl-H3K9 to consecutively generate mono-methylated and unmodified H3K9 peptides. (B) Activity of JmjC lysine demethylases may be measured by coupling formaldehyde production to NAD⁺ reduction using the coupling enzyme FDH and monitoring NADH formation. (C) Lysine demethylation catalyzed by LSD1 can be measured using the coupling enzymes HRP or FDH to quantitate hydrogen peroxide or formaldehyde products, respectively. FDH-coupled formaldehyde detection is also applicable to the testing of 2-OG-dependent JmjC-domain histone demethylases.
to an epitope-tagged reader domain protein (Fig. 3B) [63, 64]. This assay format allows facile miniaturization to 1,536-well plate density and is modular in nature with simple substitution of the reader domain or peptide modification of interest. For example, an AlphaScreen assay measuring binding of the MBT domain protein L3MBTL1 to a monomethylated histone peptide was successfully used to assess inhibitor potency of virtual screening hits identified through pharmacophore modeling and a methyllysine similarity search [65]. A similar approach to measuring these peptide/protein interactions may be taken using a TR-FRET readout, for example using europium-labeled streptavidin and an acceptor dye in close proximity where the emission spectrum of the donor fluorophore overlaps the absorption spectrum of the acceptor molecule. Here, Eu³⁺-labeled streptavidin excites anti-His₆ antibody conjugated with an acceptor dye when the peptide is bound to the reader domain protein.

**SUMMARY**

We have described a number of methods to measure histone methyltransferase and demethylase activity, as well as techniques to quantify binding of chromatin-associated domains to post-translationally modified peptides. While radioactivity-based assays remain the gold standard for evaluating histone-modifying enzymes, a number of robust and sensitive assays have been developed to address the large number of recently-identified histone methyltransferases and demethylases. Despite intense efforts to identify small molecule modulators of these enzymes, few inhibitors have been described. Of these, the majority are non-specific SAM competitors or metal ion chelators of methyltransferase or demethylase enzymes, respectively. The need exists for identification of novel scaffolds as a starting point for drug discovery related to epigenetic therapeutic targets. Robust high-throughput assays are necessary for screening large compound collections to enable detection of these chemical chemotypes and their downstream cellular effects. Continued improvements in assay technologies and novel methodologies will enable understanding of the biological role of histone methylation and our ability to mediate the epigenetic changes that affect human disease.

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search and the Intramural Research Program of NHGRI, NIH.

CONFLICT OF INTEREST

None declared.

ABBREVIATIONS

ADA = adenosine deaminase
ChIP = chromatin immunoprecipitation
DELFIA = dissociation-enhanced lanthanide fluorescent immunoassay
DTNB = 5,5’-dithiobis-2-nitrobenzoic acid
ELISA = enzyme-linked immunosorbent assay
FDH = formaldehyde dehydrogenase
FP = fluorescence polarization
FRAP = fluorescence recovery after photobleaching
FRET = fluorescence resonance energy transfer
HAT = histone acetyltransferase
Hcy = homocysteine
HDAC = histone deacetylase
HDM = histone demethylase
HMT = histone methyltransferase
HRP = horseradish peroxidase
HTS = high-throughput screening
LuxS = 3-ribosyl-homocysteinase
MCE = microfluidic capillary electrophoresis
MS = mass spectrometry
2-OG = 2-oxoglutarate
PTM = posttranslational modification
SAH = S-adenosyl-homocysteine
SAHH = SAH-hydrolase
SAHN = SAH-nucleosidase
SAM = S-adenosyl-methionine
SPA = scintillation proximity assay
TNB = 5-thio-2-nitrobenzoic acid
TRF = time-resolved fluorescence
TR-FRET = time-resolved FRET

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