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14. ABSTRACT
Our goals for this project are to explore LSD1 inhibition of protein-protein interactions as a potential therapy for ERα breast cancer, ameliorating iLCC for in vivo use, and using novel proteomics approaches to identify coregulatory proteins interacting with ERα and LSD1 in ERα+ cells and deduce how the complement of LSD1 associating proteins change in ERα cancers. We have made significant progress on the aims this project, specifically by examining the effects of small molecule, LSD1-selective and broad-spectrum histone demethylase inhibitors on breast cancer tumor growth as compared to iLCC. Using protein engineering, and H/D exchange MS, we identified the LSD1 alpha helical coiled-coil as a central mediator of ER signaling and coregulatory protein recruitment. Using siRNA knockdown and a newly identified highly selective inhibitor of the LSD1 demethylase active site chemistry, we showed that loss of LSD1 or inhibition of the LSD1-CoREST complex by iLCC inhibits breast cancer proliferation, but the antiproliferative effects of tranylcypromine or pargyline exhibit antiproliferative effects in ER+ and ER- breast cancers by a secondary mechanism requiring LSD1 as a scaffolding element.

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Introduction

Our goals for this project are to explore LSD1 inhibition of protein-protein interactions as a potential therapy for ERα breast cancer, ameliorating iLCC for in vivo use, and using novel proteomics approaches to identify coregulatory proteins interacting with ERα and LSD1 in ERα+ cells and deduce how the complement of LSD1 associating proteins change in ERα- cancers. We have made significant progress on the aims this project. There are no changes we wish to make to the Statement of Work.

Key Words
histone demethylase, estrogen receptor, CoREST, corepressor, estrogen-receptor positive breast cancer, estrogen receptor negative breast cancer, epigenetics, nuclear hormone receptor, estrogen

Overall Project Summary

Flavin-dependent, lysine-specific protein demethylases (KDM1s) are a subfamily of amine oxidases that catalyze the selective posttranslational oxidative demethylation of methyllysine side chains within protein and peptide substrates, a process that generates chemical cues for the regulation of gene transcription. KDM1 family human demethylases utilize a FAD cofactor to oxidize C-N bonds with subsequent production of a demethylated lysine residue and formaldehyde byproduct. The most well characterized KDM1-family demethylase, LSD1/KDM1A, is an 852 amino acid (aa) polypeptide composed of three domains: an amine oxidase catalytic domain (AOD) (residues 272-415 and 515-852), an ~100aa insert known as the ‘tower’ domain (residues 415-515), and a SWIRM domain (residues 172-272) (Figure 1). The three-dimensional structure of LSD1 has been solved by X-ray crystallographic methods in the unliganded form and in complex with varying ligands, including peptides and inhibitors.

The LSD1 demethylase typically associates with biomolecules and enzymes that link its catalytic activity to distinctive biological occupations in normal and disease states. Such interactions among members of transcriptional regulatory complexes have been shown to influence the degree of catalytic activity, substrate specificity, and/or localization of these enzymes within chromatin. CoREST is one of the earliest identified interaction partners of LSD1 and this protein complex assembly (i.e. LSD1/CoREST complex) is frequently found within several distinct, larger multi-protein complexes. Most likely due to this early association, the structural and functional characterization of this particular interaction far outstrips that available for any other LSD1 interaction.

LSD1 regulates the expression of many genes important in breast cancer progression and proliferation. Present various transcriptional complexes paired with tightly-associated nucleosome targeting protein CoREST and the histone deacetylases HDACs 1 and 2, the LSD1/CoREST/HDAC complex is the core enzymatic machinery responsible for removing methyl marks on histone H3. The context to which the LSD1 complex acts is controlled by its interaction with partnering co-regulatory proteins that may direct access to specific methylation sites or influence LSD1 specificity.

By mechanisms that are far from well understood, estrogen binding to the estrogen receptor alpha (ER) undergoes conformational changes to facilitate co-regulatory protein association, then is recruited by components of the LSD1 multi-protein complex to translate nuclear hormone signaling into productive transcriptional activation of estrogen-responsive genes (Figure 2). In hormone-responsive breast cancers, ER activity is a proven target for pharmacological inhibition, as ER controls many cancer-associated proliferation genes. However in highly aggressive breast cancers, ER production is abrogated, and thus by comparison there are limited therapeutic options for these aggressive cancers. In both breast cancers, LSD1 is highly expressed and activated and we have validated in cell culture that LSD1 is a potential target for both ER-positive and ER-negative cancers. In this proposal we wish to examine the molecular contributors that govern signal transmission between the ER complex and the LSD1/CoREST/HDAC1/2 complex to initiate transcriptional programs at ER target genes, to validate that LSD1 as a breast cancer target in both ER- and ER+ cancers target by examining the ability of our recently developed potent and selective inhibitor the LSD1-CoREST interaction to reduce tumor growth in a murine xenograft models of breast cancer.
Figure 2. Epigenetic transcriptional complexes that recruit LSD1 via interactions with estrogen and androgen nuclear hormone receptor transcription factors.

**Aim 1 Progress** - We wish to unequivocally confirm that iLCC functions as an inhibitor of both ERα+ and ERα− breast cancers in vivo in a clinically-relevant xenograft animal model of breast cancer.

**Evaluation of LSD1 Inhibition on Nuclear Hormone Transcription Factor Signaling** We initially turned to the estrogen-inducible ERα transcription factor to probe the involvement of the catalytic activity of the LSD1/CoREST, and later protein-protein interaction inhibitors. Using a combination of siRNA knockdown and LSD1 chemical inhibition by designed arylcyclopropylamine small molecule LSD1 inhibitors, we showed that LSD1 enzymatic activity is required for ER function (Figure 3). We also determined that inhibition of LSD1 demethylation prevents ER recruitment to estrogen-responsive elements (ERE) within promoters for target genes. LSD1 inhibition by small molecules showed strong antiproliferative effects on breast cancer cells and anticipated effects on H3K4 and H3K9 histone demethylation patterns. In ERα-negative cancers, LSD1 is highly overexpressed and presumably constitutively activated by stimulatory effects from coregulatory molecules. We recently also showed that in a dose-dependent manner, small molecule inhibitors of LSD1 exhibited marked antiproliferative effects in ER-negative cancer cell lines and disrupted transcriptional activation of ER target genes.

Figure 4. (A) Model of the CoREST-derived linker structure adapted from the structure of the LSD1-CoREST crystal structure; (B) LSD1 inhibitor iLCC (derived from the Linter region of CoREST) caused the dose-dependent inhibition of transcription at ER target genes in MCF7 cells when added exogenously; (C) Localization of iLCC to the nucleus of MCF7 cells following doxycycline induction; (D) Doxycycline mediated expression of iLCC causes a dose-dependent abrogation of methylation of histone H3; (E) iLCC stabilizes LSD1 during siRNA knockdown of CoREST; (F) Western blot indicating that CoREST Linker-Sant2 domain directly interacts with full length ER in an estrogen-dependent manner, but does not interact directly with LSD1.
Analysis of the Interaction of LSD1 with a CoREST Fragment

We subsequently probed the interaction of CoREST with LSD1 using a combination of mutagenesis, activity assays, epitope mapping, and biophysical methods including isothermal titration calorimetry, surface plasmon resonance and hydrogen/deuterium exchange mass spectrometry. We discovered that CoREST residues 293-380, form a central isolated helix that interacts very strongly (16 nM K_D) with the LSD1 coiled-coil tower domain to create an unusual offset dual triple-helical bundle interaction interface. To validate that this CoREST and LSD1 interacted only through this helix-helix interface, we used structure-driven design to produce a fully active deletion mutant of LSD1 lacking the tower domain, produced full length CoREST and confirmed that no additional sites of interaction exists between these two proteins.

In collaboration with Prof. Donald McDonnell (collaborator), we determined that a GST-CoREST293-380 fusion expressed in both HeLa and MCF7 breast cancer cells was capable of pulling down endogenous LSD1 and partially depleting the pool of the LSD1-CoREST complex. We subsequently created a cell permeable HIV-Tat-HA tagged version of the truncated CoREST linker (iLCC) and evaluated its effectiveness at inhibiting the LSD1-CoREST interaction in vitro and in vivo and its capacity to prevent ERα recruitment and to inhibit cell proliferation in ER-positive and ER-negative cancers cell lines. iLCC proved to be highly cell permeable and unusually stable for over 20h in cell culture (and >8 days with daily dosing). We subsequently attributed this stability to protection by LSD1 binding, since in a CoREST siRNA knockdown normally LSD1 is unstable and rapidly degrades, but addition of iLCC rescued LSD1 protein levels, confirming that the helix-helix interactions of CoREST with LSD1 are required for stability. At 100 nM, iLCC marginally increased global histone demethylation levels at but significantly increased methylation at H3K4 within the promoter of ER target genes. In the absence of iLCC, we used ChIP/qPCR recruitment analysis to determine that CoREST and LSD1 associated with the proximal ERE binding site of ERα-promoter targets such as pS2 and PR in MCF7 cells. Treatment of the MCF7 cells with iLCC at 100 nM disrupted the binding interaction between LSD1 and CoREST, leading to the decreased recruitment of ERα, CoREST, and LSD1 to those promoter binding sites.

In addition, iLCC treatment decreased transcriptional levels of ERα+ target genes (e.g. pS2, PR, CatD, GREB1, AmyB, and MCM2), but not 1DH3A or TIF2 which are not targets of ER- positive and reduced cellular proliferation in a dosage-dependent manner. Lastly, moving to the ER-negative cancer cell line MDA-MB-231, as was seen when LSD1 activity was chemically inhibited or the protein levels knocked down by siRNA interference, we showed that siRNA knockdown of CoREST protein levels decreased the expression of proliferation-associated genes like pS2 and PR and inhibited growth proliferation. However, viability of the cells was preserved and no toxic effects have been observed at these inhibitor concentrations.
**Induction of the EMT in Murine Breast Cancer Xenografts with iLCC Treatment** Recently, we have conducted initial evaluation of iLCC in a xenograft nude mouse model of ER+ breast cancer. In preparation for this, we prepared a doxycycline-inducible retroviral expression system for iLCC and after selection for iLCC-expressing clones, evaluated the ability of iLCC to be expressed proportional to Dox dosing. We validated that Dox-induced iLCC exhibited nearly identical behavior as exogenously administered iLCC, creating antiproliferative effects, loss of ER, LSD1 and CoREST at ER target promoters, increased methylation at these sites, and we confirmed its nuclear localization using immunofluorescence. Transplantation of Dox-inducible iLCC-expressing cells into nude mice resulted in marked phenotypic changes to the palpable tumors, in some cases producing satellite tumors, a possible sign of stimulation of the EMT in these mice, which is negatively regulated by LSD1.

**MS and Biophysical Analyses of LSD1 Complexation by CoREST and ER** In year 2 of this proposal period, progress toward examining molecular contributors to these protein-protein interactions has also been made. We also examined the role of the LSD1 tower domain in CoREST recruitment and ER capture. In addition, isoform-selective potent small molecule inactivators of LSD1 were synthesized and used to examine the contribution of coregulatory molecule interactions to demethylase catalysis and its effect on ER+ and ER-breast cancer cellular proliferation. We have also successfully expressed full length CoREST to provide a physiologically-relevant template to examine ER binding and LSD1 recruitment, and how these events are required for proliferation versus demethylation chemistry in preparation of follow up murine xenograft implantation proliferation experiments that are scheduled.

LSD1 contains domains known to mediate interactions with other biomolecules. LSD1 contains an α-helical, antiparallel coiled-coil ‘tower’ domain composed of two α-helices (termed TαA and TαB) bridged by a tight turn formed by a KPPRD sequence. This domain projects almost 100 Å from the AOD and functionally contributes to the association with the corepressor CoREST (aka RCOR1/KIAA0071) by binding to a subdomain within its sequence. CoREST consists of an N-terminal ELM2 domain followed by dual SANT domains (SANT1 and SANT2) with an intervening region colloquially known as the linker domain (Figure 4). The N-terminal region spanning the ELM2 domain and first SANT domain have been mapped as a binding region for HDAC1, and a crystal structure of HDAC1 bound to a similar ELM2/SANT region in the CoREST-related protein, MTA1, has recently been reported. Although known to associate with the MTAs 1-3 and SMRT, it is not clear if these interactions are strictly tower domain mediated, nor is it clear where the LSD1 binding site is within these co-repressors since they only share a single SANT domain with CoREST.

Shi and colleagues initially investigated the interaction of LSD1 with domains within CoREST, and concluded that CoREST may act as a scaffolding protein that joins deacetylase and demethylase activities into a single catalytic sub-complex. The CoREST SANT2 domain, and possibly the SANT1 domain, has been shown to facilitate demethylation of nucleosomes, presumably by acting as a bridge between LSD1 and its substrate. A similar mechanism is observed in the SMRT protein, where the C-terminal SANT2 domain stabilizes the protein complex on chromatin by interacting with histone tails. In the first crystal structure of LSD1 in complex with the linker and SANT2 domains of CoREST, the CoREST linker domain forms an L-shaped helical conformation, with the short helix contacting the base of the LSD1 tower domain, the longer helix extending up the TαB helix, and the SANT2 domain contacting the top turn of the tower (Figure 4). These interactions are mainly hydrophobic in nature, although a few electrostatic interactions may facilitate proper alignment of the CoREST linker with the LSD1 tower domain. A second crystal structure of LSD1 and CoREST with a histone substrate-like peptide bound (pK4M) showed little overall conformational change as compared to the non-peptide-bound structure. However, the model does illustrate that the L-shaped short arm formed by the CoREST linker may have an indirect effect on substrate binding by stabilizing the helix formed from LSD1 residues 372-395.

As noted above, although the CoREST SANT2 domain contacts the LSD1 tower in reported crystal structures, our laboratory recently demonstrated that the CoREST linker domain is chiefly responsible for high affinity binding with LSD1. Specifically, a CoREST fragment consisting of the linker domain (residues 293-380) exhibits low nanomolar binding affinity towards LSD1 ($K_d = 7.78$ nM), while inclusion of the SANT2 domain does not substantially alter binding affinity and in isolation lacks significant affinity towards LSD1. Our group also used mutagenesis and isothermal titration calorimetry to reveal that the energy of binding along the CoREST/LSD1 helix-helix interface is not concentrated into ‘hotspots,’ but is instead distributed almost uniformly along the interaction interface. As CoREST is required for nucleosomal demethylation, this argues that inhibitors of the CoREST linker/LSD1 tower interaction may inhibit LSD1 functionality in the cell.
Aim 2 Progress—Although iLCC is remarkably resistant to degradation in cellulo, we wish to also ameliorate its structure and develop formulations to that will reduce susceptibility to hydrolysis.

**Transcriptional Regulation by LSD1 Small Molecule Inhibition** Motivated by the relationship between the AOD of KDM1A to that of MAO-A/B and PAO, the first KDM1A small molecule inhibitors were discovered by our group in collaboration with R. Sheikhattar. Through screening a focused group of irreversible and reversible amine oxidase inhibitors, tranylcypromine (trans-2-phenylcyclopropylamine, 2-PCPA, Parnate™) was found to exhibit the highest KDM1A inhibitory activity towards methylated bulk histones as well as methylated nucleosomal substrates in vitro. Treatment of P19 embryonal carcinoma cells with parnate resulted in the global increase of H3K4 methylation as well as transcriptional derepression of two KDM1A target genes, Egr1 and the pluripotent stem cell marker Oct4. This was the first example of inhibition of histone demethylation and interruption of transcriptional programs regulated by KDM1A using small molecules. In a subsequent study, our group determined that tranylcypromine was a mechanism-based inactivator of KDM1A, forming a covalent adduct with the FAD cofactor and demonstrating inactivation kinetic parameters of $K_i = 242 \mu M$ and a $k_{\text{inact}} = 0.0106 \text{s}^{-1}$. Soon after, the laboratories of Cole, Yokohama, and Mattevi provided additional support for this inhibitory mechanism, including defining the structure of the tranylcypromine-KDM1A adduct using X-ray methods, which revealed the N5 of the FAD isoalloxazine ring as the site of covalent attachment of the inhibitor.

**Antiproliferative Effects of iLCC compared to LSD1-Selective Inhibitors** In Aim 2 we sought investigate the activity of KDM1A inhibitors on MCF7 proliferation versus monoamine oxidase inhibitors previously used in the literature to help evaluate whether LSD1 demethylase activity, or the interactions with coregulatory proteins like CoREST are contributing to the antiproliferative activity of LSD1-CoREST antagonists or siRNA knockdown of ER, CoREST, and/or LSD1 (Figures 5 and 6). First, we treated MCF7 breast cancer cells with a gradient of GSK-LSD1 to test its ability to alter histone methylation state in a dose-dependent manner. Cells were treated with 0, 0.05, 0.25, 0.5, 1, or 10 $\mu M$ GSK-LSD1 and incubated for 24 h. Cells were then collected and histones extracted. Each sample was analyzed by immunoblot such that each contained the same amount of total protein. Total histone H3 was analyzed as a loading control, while H3K4me2 was detected as a readout of KDM1A activity (Figure ).

![Figure 5](image). (A) siRNA and arylcyclopropylamines inhibit transcription of ER target genes; (B) Effect of arylcyclopropylamines inhibitors on cell proliferation of ER-positive and ER-negative breast cancer cell lines; (C) Knockdown of CoREST impacts the transcription of representative ER targets pS2 and PR.

**Figure 6.** Histone methylation as a function of GSK-LSD1 concentration. (a) An immunoblot of MCF7 cells treated with the designated concentrations of GSK-LSD1. The top pane represents H3K4me2 modification, while the bottom pane represents total H3 as a loading control. (b) Quantification of H3K4me2 modification. The relative intensities of H3K4me2 bands were normalized by H3 total protein loading.

H3K4me2 methylation appears to be significantly increased between the no treatment control and most other samples, although a decrease in methylation is also observed at the highest concentration of 10 $\mu M$. Although a dose-dependent response it not observed, this may be due to the concentration range chosen. It is
possible that the lowest concentration tested (50 nM) gave the maximum effect detectable. A lower concentration range may in fact demonstrate a dose-dependent response similar to that observed by Kruger et al. in a small cell lung carcinoma model (Cancer Cell, 2015). To test this, 96-well plates of MCF7 cells were treated with a five-fold dilution series of GSK2879552 (abbrev. GSK-LSD1), Parnate, and ICI182,780 (ICI). The KDM1A inhibitor GSK-LSD1 and monoamine oxidase inhibitor Parnate dilutions series began at 10 µM, while the ICI gradient began at 100 nM. Plates were treated with fresh media and compound every 2 days. On the final day, cell proliferation was measured and normalized to the initial seeding density. Proliferation curves were generated for each compound gradient (a-c). Additionally, a plot of the highest doses of each compound was generated for direct comparison (d) The proliferation assay indicated that both Parnate and ICI had a dose-dependent affect on MCF7 cell proliferation. Alternately, GSK-LSD1 did not appear to alter proliferation versus the control. At the highest doses, GSK-LSD1 and control cells reached nearly the same cell density, while ICI and Parnate both exhibited approximately 65% growth inhibition. This difference suggests that Parnate may be working through a mechanism other than KDM1A inhibition to slow breast cancer cell proliferation and throws into question recent reports that use the inhibitor to assign functionality to KDM1A. However, our results with iLCC and siRNA knockdown of LSD1 and CoREST support a clear relationship where LSD1 is required for proliferation, as such it suggests that LSD1 when in complex with the ER may be playing a scaffolding role in facilitating transcription.

Figure 7. Proliferation of MCF7 cells as a function of inhibitor. (a) Proliferation as a function of monoamine oxidase inhibitor Parnate concentration. (b) Proliferation as a function of KDM1A inhibitor GSK2879552 (i.e. GSK-LSD1) concentration. (c) Proliferation as a function of estrogen receptor alpha antagonist ICI concentration. (d) Summary of the highest concentrations of each treatment.

Aim 3- To assist in understanding LSD1 function in breast cancer, we wish to provide an innovative solution to identify ER targets involved in communication with LSD1 within complex protein mixtures by combining SPROX shotgun proteomics with chemical genetic probes of ER communication with the LSD1 complex (developed by the PI).

Interface Mapping by MS Methods Mass spectrometry experiments to identify the physical basis for communication between the ER and LSD1 complex have been initiated. Unfortunately, because of the poor degree of fragmentation LSD1 exhibited using bottom up proteomics, insufficient coverage was achieved in initial MS analyses to facilitate SPROX studies with the ER. However, we have recently turned to using H/D exchange using high resolution ESI/MS methods which overcame the fragmentation issues. Based upon our preliminary analyses of LSD1 using binding to histone H3 and CoREST as probes, we have been able to easily visualize specific conformational changes within LSD1 and the LSD1 CoREST complex upon substrate binding. We are now pursuing similar analyses with LSD1 and CoREST fragments with the ligand binding domain of the ERα, aided as well by newly produced recombinant full length CoREST which serves as a template for these interactions.
Molecular Interactions Underlying LSD1 Communication with Coregulatory Proteins

Lysine-specific demethylase 1A (KDM1A/LSD1) removes mono- and dimethyl post-translational modifications on lysine residues of histones leading to both repression and activation of associated genes. Interestingly, KDM1A activity and its substrate specificity are regulated by both active site recognition and association with a number of coregulatory proteins and inclusion into multiprotein complexes through protein-protein interactions. CoREST is one such binding partner of KDM1A that facilitates nucleosomal demethylation through interacting with its unprecedented tower domain. Besides CoREST and paralogs, there are many homologous binding partners that utilize this domain as an oligomerization motif. As the interaction with the tower domain is a central theme in the elaborate control of KDM1A and direction of its activity, over the past year, we conducted analysis of the binding interface between KDM1A and CoREST in order to provide insight into their collaborative mechanism. Toward this end, initial steps have been taken to define the key regions implicated in the binding interaction. Using this information along with structure-aided protein engineering, we developed an LSD1 enzyme that possessed full demethylase enzymatic activity, yet was devoid of the tower domain and subsequently lost all affinity for CoREST (see the structure of the ‘towerless LSD1 in the inset).

Interfacial Binding Kinetics Govern Histone H3 Recognition and Specificity

Lysine-specific demethylase 1A (KDM1A/LSD1) is an FAD-dependent enzyme that catalyzes the oxidative demethylation of histone H3K4me1/2 and H3K9me1/2 repressing and activating transcription, respectively. Although the active site is expanded compared to the greater amine oxidase superfamily, it is too sterically restricted to encompass the minimal 21-mer peptide substrate footprint. The remainder of the substrate/product is therefore expected to extend along the surface of KDM1A. We show that full-length histone H3 which lacks any posttranslational modifications is a tight-binding, competitive inhibitor of KDM1A demethylation activity with a $K_i$ of 18.9 ± 1.2 nM; a value that is approximately 100-fold higher than the 21-mer peptide product (i.e H31-21). The relative H3 affinity is independent of preincubation time suggesting that H3 rapidly reaches equilibrium with KDM1A. Rapid dilution experiments confirmed the increased binding affinity of full-length H3 was at least partially due to a slow off-rate ($k_{off}$) of $1.2 \times 10^{-3}$ s$^{-1}$, a half-life ($t_{1/2}$) of 9.63 min, and residence time (t) of 13.9 min. Independent affinity capture surface plasmon resonance experiments confirmed the tight-binding nature of the H3/KDM1A interaction, revealing a $K_d$ of 9.02 ± 2.27 nM, a $k_{on}$ of $9.26 \times 10^4 ± 1.5 \times 10^5$ $M^{-1}$s$^{-1}$ and a $k_{off}$ of $8.35 \times 10^{-4} ± 3.4 \times 10^5$ s$^{-1}$. Additionally, no other core histones exhibited inhibition of KDM1A demethylation activity, which is consistent with H3 being the preferred histone substrate of KDM1A. Together these data suggest that KDM1A likely contains a histone H3 secondary specificity recognition element on the enzyme surface.

Key Research Accomplishments:
- Characterization of iLCC vs. the LSD1-selective inhibitor GSK2879552
- Validation of iLCC inhibition of LSD1 function in vivo and in vitro
- Identification of CoREST knockdown as a target for iLCC competition for LSD1
- Successful identification of LSD1-dependent protein-protein interactions with coregulatory molecules using H/D exchange MS methods and deletion mutagenesis (e.g. tower domain deletion mutant)
- Observation of mechanistic differences between iLCC, parnate the small molecule LSD1-selective inhibitor GSK2879552 that suggest a scaffolding function for LSD1 or additional target coregulatory molecules influence breast cancer proliferation

Conclusion

In short we have made substantial progress on the project and are excited to complete the experiments outlined in the statement of work for the remaining period of this proposal period. Thank you for your generous support of this research project.
Publications, Abstracts and Presentations
Published or in press manuscripts are attached as supporting information. Two additional manuscripts not included in this update are under review.

This work was presented as a research talk by Prof. McCafferty at the Enzymes Gordon Research Conference, UCSF, UNC-Chapel Hill and a GSK-sponsored research symposium in the Research Triangle Park.

Jennifer Link will present this work in an invited talk in San Diego for the ACS National Meeting in March 2016 (abstract accepted Nov 2015)

Inventions, Patents, and Licenses
Nothing to report

Reportable Outcomes
Nothing to report

Other Achievements
Nothing to report

References
Nothing to report

Appendices
Nothing to report
Flavin-dependent, lysine-specific protein demethylases (KDM1s) are a subfamily of amine oxidases that catalyze the selective posttranslational oxidative demethylation of methyllysine side chains within protein and peptide substrates. KDM1s participate in the widespread epigenetic regulation of both normal and disease state transcriptional programs. Their activities are central to various cellular functions, such as hematopoietic and neuronal differentiation, cancer proliferation and metastasis, and viral lytic replication and establishment of latency. Interestingly, KDM1s function as catalytic subunits within complexes with coregulatory molecules that modulate enzymatic activity of the demethylases and coordinate their access to specific substrates at distinct sites within the cell and chromatin. Although several classes of KDM1-selective small molecule inhibitors have been recently developed, these pan-active site inhibition strategies lack the ability to selectively discriminate between KDM1 activity in specific, and occasionally opposing, functional contexts within these complexes. Here we review the discovery of this class of demethylases, their structures, chemical mechanisms, and specificity. Additionally, we review inhibition of this class of enzymes as well as emerging interactions with coregulatory molecules that regulate demethylase activity in highly specific functional contexts of biological and potential therapeutic importance.

INTRODUCTION

The functional capacity of genetically encoded proteins can be powerfully expanded by reversible posttranslational modifications (PTMs). Protein side chain covalent modifications such as methylation, phosphorylation, acetylation, glycosylation, ubiquitination, sumoylation, ADP-ribosylation, nitrosylation, carboxylation, and sulfonation, among others, can potentiate local and global protein architecture and generate novel reactive functional groups for binding and catalysis. Additionally, these modifications can generate signals for cellular compartmentalization and confer alterations to physiochemical properties such as stability, solubility, and resistance to proteolytic degradation. Such modifications are intimately integrated into all aspects of modulating cellular proliferation and biological recognition.
Allfrey first linked histone acetylation and methylation to the control of RNA synthesis in 1964. Since that time, it has become clear that protein and histone methylation is a pervasive cellular regulatory mechanism. Specifically, protein lysine methylation status plays a critical role in biology and pathobiology by influencing transcriptional activation and repression, chromatin remodeling, normal and oncogenic signaling, viral pathogenesis, and protein and transcription factor recruitment, among other functions. S-adenosylmethionine-dependent methyltransferases (SAM-dependent MTases) catalyze the installation of lysine methyl marks, forming mono-, di-, and tri-methylated side chain PTMs. These marks are removed by two mechanistically-distinct classes of lysine-specific demethylases (KDMs) that utilize either a flavin adenine dinucleotide (FAD) cofactor (KDM subfamily 1) or iron(II) and α-ketoglutarate cofactors (KDM subfamily 2–6) to catalyze the oxidative demethylation of protein side chains.

For decades, methylation marks on proteins like histones were presumed to be immutable, reversed only through protein turnover and degradation. The first hints of a candidate histone demethylase enzyme emerged in 1998 and again in 2001, when several histone deacetylase (HDAC)-containing complexes were discovered to associate with KIAA0601 (aka KDM1A/LSD1/BHC110/AOF2), a protein with homology to human FAD-dependent oxidoreductases (MAO-A/B) and maize polyamine oxidase (mPAO) (Figure 1a), enzymes that catalyze the oxidative cleavage of C-N bonds. In 2003, Shiekhattar reported that BHC110/KIAA0601 formed a stable complex with several proteins, including HDACs 1 and 2, CoREST, BRCA2-associated factor, BCH80, TFII-1, KIAA0182/GSE-1, ZNF217, ZNF198/FIM and ZNF261/X-FIM.

In 2003 Amasino and coworkers independently discovered that the gene FLOWERING LOCUS D (FLD) encoded a plant homolog of KIAA0601. FLD is one of six genes in the
**Arabidopsis thaliana** autonomous floral-promotion pathway that initiates the transition from a vegetative to reproductive state by repression of the MADS-box transcription factor FLOWERING LOCUS C (FLC). FLD not only contains a KIAA0601-like amine oxidase domain, but also possesses an N-terminal SWI3p, Rsc8p, and Moira (SWIRM) domain similar to that associated with a range of proteins involved in chromatin remodeling, including KIAA0601. Deletion of FLD in *A. thaliana* results in hyperacetylation of histones in the FLC locus, upregulation of FLC expression, and extremely delayed flowering, suggesting that the autonomous pathway involves regulation of histone deacetylase activity by FLD. Deletion of FLD also resulted in increased histone methylation levels (R. Amsino, personal communication), implicating FLD (and by analogy KIAA0601) as the elusive human histone demethylase enzyme.

In early 2004, Shi and coworkers provided the first direct evidence that KIAA0601 (from here on referred to as KDM1A) functions as a histone demethylase and transcriptional corepressor. The authors reveal that KDM1A specifically demethylates mono- and di-methylated histone H3 lysine 4 (H3K4me1/2), a histone mark linked to active transcription, and that lysine demethylation occurs via an oxidative reaction that generates formaldehyde. In addition, loss of KDM1A through siRNA knockdown results in increased H3K4 methylation and concomitant derepression of several neuronal-associated target genes. Shi’s discovery of KDM1A activity implied that lysine methylation might be dynamically controlled.

The second human flavin-dependent histone demethylase, KDM1B (aka LSD2/AOF1) was identified by Shi and coworkers in 2004 through a domain homology search of genomic databases. Mattevi and coworkers first isolated and confirmed the flavin-dependent demethylation activity of KDM1B, noting specificity for H3K4me1/2, despite the relatively low sequence identity (<25%) with KDM1A. Unlike KDM1A, KDM1B does not form a biochemically-stable complex with the C-terminal domain of the corepressor CoREST, but does possess both CW-type and C4H2C2-type zinc finger motifs. This suggests that KDM1B may interact with different targets or coregulatory molecules and may be involved in transcriptional programs distinct from those of KDM1A.

Herein we review the biological function, biochemical characterization, and inhibition of this enzyme class to date. First, we briefly discuss the biological importance and therapeutic potential of KDM1s. This is followed by the structural organization, chemical mechanism, and substrate specificity of the KDM1 enzymes. We then outline the various inhibitor classes that have been developed for these demethylases, specifically highlighting the utility of peptide-based inhibitors. Finally, we describe the known interactions between the KDM1s and regulatory biomolecules, which direct their activity toward specific cellular pathways. Given the wide-ranging and ubiquitous functions of this enzyme class, probes with the ability to target KDM1 activity in a manner that is pathway-specific would be of heuristic and therapeutic value. We see these coregulatory molecules as a valuable starting point for the development of such probes, as underscored by the recent development of peptide inhibitors of this enzyme class.

**KDM1A AND KDM1B BIOLOGY AND THERAPEUTIC POTENTIAL**

KDM1A is involved in a wide variety of cellular processes and pathologies, including signal transduction, chromatin remodeling, transcriptional regulation, development, differentiation, viral pathogenesis, and cancer proliferation and metastasis. As such, KDM1 demethylases have emerged as potential therapeutic targets. Although their clinical value is only beginning to be explored, at the time of this review’s publication, five clinical trials involving KDM1A were either planned or currently recruiting subjects. Three trials will investigate KDM1A inhibition as a therapeutic strategy in acute myeloid leukemia (AML), one will evaluate the same strategy in small cell lung cancer, and one seeks to correlate the status of KDM1A with cardiovascular responses to changes in sodium intake (see http://clinicaltrials.gov). We expect the scope of clinical trials to expand with our growing knowledge of KDM1A function.

**Cellular Differentiation**

Epigenetic modifications are critical for the maintenance of pluripotent stem cells and their subsequent differentiation into specialized tissues. Specifically, KDM1A is an integral part of the cellular machinery that governs differentiation processes in various cell types, including embryonic, hematopoietic, neuronal, pituitary, and osteogenic. While KDM1A activity is critical for proper development, aberrant regulation can contribute to tissue-specific disease states. For example, KDM1A activity has been linked to neuronal cell proliferation and survival, and long-term memory formation, and is therefore an emerging therapeutic target for some neurological and cognitive disorders. Similarly, recent work has also identified KDM1A as a key regulator of leukemia stem cells, sparking interest in its potential as a therapeutic target in AML (see below), among other hematopoietic-related diseases.

**Cancer Development and Progression**

KDM1A has also been widely implicated in the development and progression of cancer. In addition to the malignancies
described in detail below, KDM1A is emerging as a potential therapeutic target in various other cancer types.46–56

Neuroblastoma. Overexpression of KDM1A in neuroblastic tumors correlates with poor prognosis57 and may be related to the aberrant downregulation of KDM1A-silencing micro-RNAs.58,59 Schulte and coworkers first noted in 2009 that KDM1A inhibition in neuroblastoma cells leads to increased H3K4 methylation, decreased proliferation in cell culture, and reduced tumor growth in a xenograft model.57 Similarly, KDM1A inhibitors synergize with retinoic acid, a currently approved treatment for neuroblastoma.60 A later report demonstrated similar results in the closely related medulloblastoma tumor type, where KDM1A inhibition decreases proliferation and initiates apoptosis.61

Colorectal Cancer. KDM1A is also an emerging target in colorectal cancers, where it is markedly overexpressed in tumor samples compared to matched normal tissue62 and exhibits a positive correlation with metastatic potential.63,64 Although genetic deletion of KDM1A does not result in a global increase of H3K4me2, it does alter gene expression programs and reduces proliferation in cell culture and in vivo.62,65 KDM1A may also promote colorectal cancer progression by derepressing the Wnt/b-catenin signaling pathway, a known regulator of tumorigenesis.62

Leukemia. KDM1A is also a promising target for pharmacological intervention in leukemia. For example, KDM1A inhibition alone provokes cytotoxic effects in AML cell lines,66 and when used in conjunction with HDAC inhibitors, reduces tumor engraftment and improves survival in mice.67 KDM1A inhibition also unlocks the anti-leukemic effects of all-trans-retinoic acid (ATRA), a differentiation-inducing agent that otherwise is ineffective for the treatment of AML.66 KDM1A has been implicated in other leukemia subtypes as well. In a model of MLL-AF9 leukemia cells, KDM1A cooperates with the MLL MTase fusion protein to maintain an oncogenic transcription program, and KDM1A inhibitors preferentially reduce the repopulation potential of MLL-AF9 cells over normal hematopoietic cells.44 Interestingly, Sprüssel and coworkers evaluated the potential side effects of KDM1A inhibition in leukemia and found that while a conditional KDM1A knockdown significantly alters existing pools of normal hematopoietic tissues, these effects are largely reversible following reinstatement of the demethylase.31 For a more detailed discussion of KDM1A’s function and therapeutic potential in leukemia, we refer readers to a recent review by Mould and colleagues.69

Breast Cancer. KDM1A has also been implicated as a regulator of breast cancer development and progression. The majority of the breast tumors are estrogen-dependent, making estrogen antagonists one of the most commonly employed therapies. However, circumvention of estrogen signaling, such as through loss of estrogen-receptor alpha (ERα), can lead to hormone-resistant tumors, reduced treatment options, and worse prognosis. KDM1A is generally associated with a more aggressive breast cancer phenotype and may be overexpressed in both ERα-positive and -negative cells,70–73 possibly due to aberrant stabilization of the demethylase.74 Correspondingly, several groups have reported that KDM1A inhibition decreases proliferation of breast cancer cell lines.70,72,75 Notably, at least two groups have reported contradicting results that suggest KDM1A may be downregulated in breast cancer tissue and that inhibition affects metastasis with no observable effects on proliferation.76,77 KDM1A inhibition may also have therapeutic effects when used in conjunction with anti-estrogen treatment79 and HDAC inhibitors.80

Prostate Cancer. A similarly vague relationship exists between KDM1A and prostate cancer. One study reported a correlation between KDM1A expression and risk of recurrence,81 while at least two other studies have noted little to no overexpression of KDM1A in prostate cancer cell lines and tumor samples.82,83 However, Metzger and coworkers have linked KDM1A activity to transcriptional regulation with the androgen receptor (AR), a nuclear hormone receptor closely linked with prostate cancer.84 Likewise, KDM1A inhibition appears to impede AR-mediated transcription and prostate cancer proliferation.85–87 A review detailing further the roles of demethylases in prostate cancer has recently been published.88

The Epithelial-to-Mesenchymal Transition
In addition to these roles in specific cancer types, KDM1A is also intimately linked with the epithelial-to-mesenchymal transition (EMT), the process by which a polarized, adhesive epithelial cell transitions into a motile, invasive mesenchymal phenotype. The importance of epigenetic mechanisms during the EMT is only beginning to be appreciated.89 KDM1A is an established functional partner of the EMT master regulator SNAI1 and cooperates to silence epithelial genes.90 KDM1A also regulates upstream inducers of the EMT,91 including TGFβ,69,72,73 Wnt,52,94 NF-κB,80 and Notch.74,75 Through these and potentially other mechanisms, KDM1A participates in large-scale chromatin remodeling events that accompany the EMT.92

The EMT is a hallmark of aggressive cancers with a high risk of metastasis, potentially implicating KDM1A as a
Viral Pathogenesis

Lastly, KDM1A is associated with herpes simplex virus (HSV) infection and replication, as HSV hijacks KDM1A-containing machinery required for human neuronal gene repression during host infection. Coopposition of KDM1A demethylase activity helps promote early stage (α) viral gene activation, regulates late stage (βγ) viral gene repression, and has been linked to repression during the development of HSV latency, the process by which viruses enter and emerge from dormancy in infected host cells.99 Kristie and coworkers recently demonstrated that interruption of KDM1A activity potently represses HSV immediate-early (IE) gene expression and genome replication, resulting in suppression of primary infections and subsequent reactivation from latency in murine infection models.100–102 In addition, KDM1A activity may play a similar role in the infection processes of several other viruses.101,103,104 Collectively, these studies underscore the emerging importance of KDM1A as an antiviral target.

Emerging Biological Roles for KDM1B

The biological roles of KDM1B are beginning to emerge; however, our current understanding of this enzyme lags far behind that of KDM1A. In general, KDM1B is believed to function as an enhancer of gene transcription that functions in highly transcribed coding regions of chromatin.105 Although mechanistic data is not yet available, KDM1B has been linked to genomic imprinting,106 liver development,107 somatic cell reprogramming,108,109 and NF-κB signaling.110 Future research will likely reveal a broader set of KDM1B-dependent biological processes.

**DOMAIN ORGANIZATION AND STRUCTURAL FEATURES OF KDM1 DEMETHYLASES**

**KDM1A Demethylase**

KDM1A is an 852 amino acid (aa) polypeptide composed of three domains: an amine oxidase catalytic domain (AOD) (residues 272–415 and 515–852), an ~100 aa insert known as the “tower” domain (residues 415–515), and a SWIRM domain (residues 172–272) (Figure 1b,c).11,111,112 The N-terminal region of the enzyme (~170 aa) has no predicted conserved structural elements, but does contain a nuclear localization signal (RRKRAK; residues 112–117) for nuclear import via importin-α translocases.113,114 The three-dimensional structure of KDM1A has been solved by X-ray crystallographic methods in the unliganded form and in complex with varying ligands, including peptides and inhibitors.111,112,115–126 The AOD of the enzyme contains a single non-covalently bound FAD molecule and bears similarity to human MAO-A/B and mPAO (17.6%, 17.6%, and 22.4% sequence identity, respectively; Figure 1a).127 Consistent with other flavin-dependent amine oxidases,128,129 the AOD can be further subdivided into two separate lobes, where residues 357–415, 515–558, and 658–769 are involved in substrate binding and residues 272–356, 559–657, and 770–852 form an expanded Rossmann fold used to bind FAD.112,115

The tower domain of KDM1A is an antiparallel coiled-coil composed of two α-helices (termed TzA and TzB) bridged by a tight turn formed by a KPPRD sequence.112 This domain projects almost 100 Å from the AOD and functionally is the site of association with co-repressor CoREST (aka RCOR1/KIAA0071), among other proteins.112,115,130 The SWIRM domain of KDM1A is predominantly a bundle of α-helices, with a long central α-helix that separates two smaller helix-turn-helix motifs (Figure 1c).111 This domain also engages the C-terminal tail of the AOD with a highly hydrophobic interface that buries a surface area of ~1,700 Å².111 Mutations of conserved residues along this interface have been shown to reduce the catalytic activity of KDM1A and its stability.111

Although SWIRM domains are highly conserved amongst chromatin associating proteins and have been shown to bind DNA,112,113 this is not the case in KDM1A. The residues that compose the typical DNA-binding interface (26; residues 247–259) are not conserved and are partially blocked by their interaction with the AOD.111,115 Accordingly, neither KDM1A nor the isolated SWIRM domain binds DNA.115,132

In addition, Luka and coworkers showed that KDM1A binds to the small molecule tetrahydrofolate with a Kᵦ of 2.8 μM, and recently reported the crystal structure of the KDM1A-CoREST-tetrahydrofolate complex.126,133 In this complex, the folate binds in the active site in close proximity to the FAD cofactor, and overlaps with the peptide-binding site observed by Forneris.117 Luka and coworkers also detected the production of 5,10-methylene-tetrahydrofolate by mass spectrometry following incubation of KDM1A with a H3K4-derived methylated peptide substrate and tetrahydrofolate. From this data, it was postulated that tetrahydrofolate binding...
might protect the cell from potentially toxic formaldehyde or regulate access of the substrate to the FAD within the active site. It is not known at this time if KDM1B exhibits a similar ability to bind tetrahydrofolate.

**KDM1B Demethylase**

The closely related KDM1B is an 822 aa flavoenzyme amine oxidase that contains a predicted unordered N-terminus and a nuclear localization signal (~50 aa) responsible for its residence within the nucleus of human cells. KDM1B is organized into three domains: an N-terminal dual zinc finger domain (residues 50–264), a SWIRM domain (residues 264–372), and a C-terminal catalytic AOD (residues 372–822; Figure 1b,d). In the substrate-free structure of KDM1B, the zinc fingers and catalytic AOD are packed closely against a central SWIRM domain (Figure 1d), collectively resembling a boot. Comparison of the AOD between KDM1A and KDM1B (33% sequence identity) shows that many secondary structural elements are conserved, including strong conservation of the catalytic architecture around the FAD-binding site. However, when comparing the AODs through structural alignment, there is a 2.0 Å RMSD difference in the overall polypeptide backbone conformation. These structural differences are mostly attributed to the length and conformation of solvent exposed loops within the domain. Despite these differences, the similarity of the AODs of KDM1A and KDM1B argues for a common chemical mechanism for catalysis.

**Structural Differences Between KDM1A and KDM1B**

Despite catalyzing identical chemical reactions and sharing significant AOD structural similarity, KDM1A and KDM1B have several structurally important differences. For example, the SWIRM domains of both enzymes only share 24% sequence identity. Although helices within this domain are qualitatively positioned, significant differences have been observed at both solvent exposed interfaces along the helical termini. Since the entrance to the AOD active site lies at the interface of the AOD and SWIRM domains, structural and sequence differences at this interface are suspected to impact substrate specificity of each isoform. Indeed, at this interface the SWIRM-AOD subdomain interactions differ in several aspects when comparing KDM1A to KDM1B. Notably, the SWIRM domain of KDM1B closely packs against the N-terminal zinc finger domain, burying a surface area of 1,872 Å². As compared to KDM1A, the SWIRM domain of KDM1B lacks a C-terminal helix (KDM1A; residues 171–181), but is replaced by an extended, coiled loop (KDM1B; residues 271–281). This substitution in KDM1B leads to the formation of two additional hydrogen bonds (E452 with N266, and Y268 with D571) and one electrostatic interaction (E323 with K323) that may influence the degree of association among these domains. Not only does this loop provide additional intramolecular interactions, but also forms a second binding site for the N-terminal tail of histone H3.

The most striking structural difference between KDM1A and KDM1B is the inclusion of distinct domains that mediate interactions with other biomolecules (Figures 1b,e). KDM1A contains an α-helical, antiparallel coiled-coil tower domain that is absent in KDM1B (Figure 1c–e). DALI structural similarity analyses indicate that the tower domain fold appears infrequently among known eukaryotic protein structures, with the PI3Kz-p85z subunit complex (PDB 3HHM) and the DNA double-strand break repair ATPase RAD50 (PDB 1L8D) bearing the closest structural similarity. In contrast, the tower fold was heavily represented among a subset of prokaryotic proteins known to utilize such motifs as part of intermolecular protein recruitment, membrane docking, and membrane translocation functions.

Although KDM1B does not contain a tower domain, it does contain two individual zinc fingers and two linker domains that are not reflected in the structure of KDM1A (Figure 1b,e). At the N-terminus of KDM1B, a CH2C2-type zinc finger is joined to a second CW-type zinc finger (Zf-CW) by a linker formed from two α-helices (Figure 1bd). Immediately following the Zf-CW domain, a second linker formed by four α-helices joined through several large loops bridges these domains to the AOD. The surface of the CH2C2-type zinc finger shows a marked concentration of basic residues, and thus may impact demethylase substrate specificity or positioning within nucleosomal DNA. Additionally, these residues may facilitate interactions with coregulatory molecules or serve to recruit transcriptional machinery, such as phosphorylated RNA polymerase II. Interestingly, it appears that the zinc finger domain of KDM1B is required for histone demethylation activity. Wong and coworkers suggest that mutations which disrupt the zinc finger domains or relays of interactions among the zinc finger-SWIRM-AOD likely lead to subtle conformational alterations in the AOD that in turn impair the incorporation of FAD, and consequently its enzymatic activity.

Furthermore, although zinc fingers are common folding motifs, the Cys2A-Cys2B-His2A'-Cys2B' sequence within KDM1B folds into a cross-brace topology that has not been previously observed, raising the possibility that this domain may participate in biomolecular interactions that are exclusive to KDM1B. On the other hand, the Zf-CW of KDM1B bears significant sequence similarity to CW-type zinc finger domains, a class linked to histone H3 binding. Despite this, as
an isolated recombinant domain, the KDM1B Zf-CW fails to bind histone H3-derived peptides (residues 1–21 containing H3K4me1/2 or unmethylated H3K4), which is atypical to behavior exhibited by other isolated Zf-CW domains. However, since ligand binding within Zf-CW domains is mediated by interactions within a well-conserved hydrophobic cleft, which happens to be absent in the KDM1B Zf-CW domain structure, this may partially explain its inability to bind histone H3 peptides.

Taken together, the exclusive presence of the tower domain in KDM1A and the zinc finger motifs in KDM1B strongly suggest that these two demethylases share a related chemical mechanism, but likely participate in associations with non-identical sets of interacting partner biomolecules. These partners may influence individual catalytic activity, substrate specificity, or localization of these enzymes within genetic loci.

CHEMICAL MECHANISM OF KDM1-FAMILY FLAVIN-DEPENDENT DEMETHYLASES

KDM1A and KDM1B are Mechanistically Distinct from Iron(II)-Dependent Lysine-Demethylases

KDM1 family human demethylases utilize a FAD cofactor to oxidize C-N bonds with subsequent production of a demethylated lysine residue and formaldehyde byproduct. By contrast, KDM subfamilies 2–6 (Jumonji-C domain-containing demethylases) are mechanistically distinct from the KDM1s since they utilize non-heme iron(II) and \( \alpha \)-ketoglutarate (\( \alpha \)-KG) cofactors to demethylate lysine residues with the concomitant production of succinate and formaldehyde. KDM families 2–6 are also distinguished from KDM1A and KDM1B by their unique ability to demethylate trimethylated lysine residues. However, KDM1 flavoenzymes have a mechanistic requirement of a protonated iminium intermediate, achievable only with mono- and dimethylated lysines.

Involvement of Flavin Adenine Dinucleotide as a Cofactor

Experimental and theoretical mechanistic studies have been conducted on flavin-utilizing KDMs, focusing almost exclusively on KDM1A. Mattevi and coworkers first confirmed the participation of the non-covalent FAD in the demethylation reaction using two truncated forms of KDM1A lacking the first N-terminal 157 or 184 aa. They visualized production of the two-electron fully reduced flavin, noting the absence of intermediate one-electron reduced forms of the cofactor. This suggested that KDM1A was likely related by mechanism to the broader family of flavoenzyme oxidases that catalyze two-electron oxidation of substrates with concomitant cofactor reduction. Once the substrate is oxidized, hydrolysis by bulk solvent of the iminium intermediate produces an unstable hemiaminal that spontaneously decomposes into formaldehyde and the mono-demethylated product. Additionally, reoxidation of the FAD cofactor by molecular oxygen produces a molar equivalent of H\(_2\)O\(_2\) and the fully oxidized quinone (Scheme 1).

Evidence for a Direct Hydride Transfer Mechanism

McCafferty, Fitzpatrick, and coworkers conducted a detailed investigation of the chemical mechanism of KDM1A. The mechanism was examined using the effects of pH and isotopic substitution on steady-state and pre-equilibrium kinetic parameters. Using a 21-mer derived from histone H3
substrate. Thus, one may argue that the release of the demethylated product occurs before reoxidation. Interestingly, in cell culture KDM1A remove a single methyl mark from p53-K370me2 to produce K370me1; however, in vitro it completely demethylates this residue. Thus, one may argue that the release of the demethylated product may be influenced by coregulatory molecules or other contributing factors capable of tuning its catalytic proficiency or substrate specificity.

Using classical molecular dynamics (MD) and quantum mechanics/molecular mechanics (QM/MM) approaches, Karasulu and coworkers provided theoretical support for a hydride transfer mechanism via adduct formation were found to be crucial for facilitating the oxidative C–H bond cleavage. Similarly, Truhral and coworkers provided theoretical support for a hydride transfer mechanism, but also noted that because of the magnitude of the calculated free energy, a mechanism involving concerted transfer of a hydrogen atom and an electron could not be ruled out. Lastly, Kong and coworkers used theoretical calculations to provide support for a hydride transfer mechanism, but also used MD to implicate a conserved Y761 and K661-water-flavin motif in orienting FAD with respect to the substrate. Collectively, these theoretical analyses are in good agreement with the experimentally determined hydride transfer mechanism for KDM1A.

KDM1B Chemical Mechanism

Tentatively, we suspect that KDM1B also adopts a similar hydride transfer chemical mechanism given the overall structural homology it maintains with the AOD of KDM1A, the conservation of active site residues, and subsequent, nearly identical, substrate specificity towards peptide substrates in vitro. Additional support for this hypothesis is reflected by the similar pH-rate profile behavior of the kinetic parameters as compared to KDM1A, non-covalent nature of the FAD cofactor, and highly similar catalytic efficiency parameters, with KDM1B being slightly less efficient in vitro.

Chemical Mechanism Implications

When comparing KDM1A and KDM1B to other amine oxidases, it is important to note that the rate constant of substrate amine oxidation is 2–5 orders of magnitude slower than that of the MAOs and PAOs. This suggests that KDM1s likely evolved for substrate specificity rather than catalytic proficiency. Given the physiological role of KDM1 demethylases in the regulation of gene expression, high catalytic efficiency, which is necessary for metabolic enzymes to maintain balance among metabolic pathways, would be much less critical than very high specificity.
SUBSTRATE SPECIFICITY

Specificity for Histone Substrates

Shi and coworkers first reported that KDM1A was capable of demethylating methyllysine peptides derived from the highly conserved N-terminus of histone H3, as well as full-length H3 in vitro.\textsuperscript{13} No cross-reactivity for polyamine substrates was observed, despite the similarity of KDM1A to the PAO superfamily.\textsuperscript{13} They also reported that KDM1A is specific for histone H3K4me1/2, as no other methyllysine sites were processed.\textsuperscript{13} Forneris and coworkers demonstrated that histone H3 tail peptides greater than 16 aa in length are necessary to achieve demethylase efficiency in vitro. Additionally, they revealed that KDM1A demethylated H3K4me1 and H3K4me2 with similar kinetic parameters in vitro, illustrating a lack of a strong kinetic preference for either substrate.\textsuperscript{147}

Mattevi and coworkers have also examined the contribution of residues within the H3 N-terminal tail to the efficiency of KDM1A demethylase activity, as well as the effects of epigenetic PTMs within this sequence.\textsuperscript{144,147,152} For example, Mattevi showed that methylation of H3K9 does not affect enzyme catalysis, yet acetylation at H3K9 increases the \( K_m \) 6-fold for H3 peptide substrates methylated on K4. Similarly, phosphorylation of H3S9 abolishes demethylase activity, suggesting that KDM1A 'reads' PTMs along the histone tail.\textsuperscript{147,152} It was also proposed that electrostatic interactions likely contribute significantly to substrate binding, since demethylation activity was shown to be sensitive to ionic strength as well as hyperacetylation in vitro.\textsuperscript{147,152} Additionally, Mattevi showed that unmethylated H3-derived product peptides exhibit inhibition, which may serve to regulate KDM1A activity or localization.\textsuperscript{147}

Parallel experiments conducted with KDM1B revealed a similar substrate specificity profile,\textsuperscript{14} which was anticipated given the significant active site structural similarity shared between the two enzymes in the AOD.\textsuperscript{112,134,136} Mutations of residues in this cleft in KDM1A abolish or abrogate demethylation activity, suggesting that interactions with substrates within this region may provide an additional specificity determinant.\textsuperscript{111} Additionally, surface plasmon resonance interaction studies have demonstrated that the SWIRM domain of KDM1A interacts with H3-derived peptides.\textsuperscript{152} In KDM1B, the surface landscape of this region is quite different and binds the coregulatory protein Nuclear Protein 60kDa/Glyoxylate Reductase 1 Homolog (NPAC/GLYR1).\textsuperscript{135,136}

In order to dissect the molecular basis of KDM1A and KDM1B substrate specificity, several three-dimensional structures have been determined of enzyme-substrate like complexes where short peptides occupy the active site in two general conformations (Figure 2).\textsuperscript{116,117,123,134–136} The structure of KDM1A bound to a substrate-like peptide inhibitor containing a N-methylpropargyl derivatized lysine residue at position four revealed density for residues 1–7 of the peptide and adopted an unusually compacted backbone with three consecutive \( \gamma \)-turns (Figure 2a).\textsuperscript{116} Subsequently, a H3 peptide inhibitor containing a K4M mutation (H3K4M, pK4M) was co-crystallized with KDM1A and CoREST,\textsuperscript{117} which revealed density for the first 16 residues of the peptide with the first five adopting an \( \alpha \)-helical conformation (Figure 2b). It is important to note that the C-termini of each of these peptides points in opposing directions upon exiting the active site. Additionally, the N-terminal region of SNAI1 and related peptide derivatives have been co-crystallized with KDM1A and display a nearly identical conformation to that of the H3K4M peptide (Figure 2c), thus suggesting some form of molecular mimicry.

As a member of the flavin-dependent oxidases, the KDM1 demethylases exhibit a strict requirement for proper substrate positioning in relation to the FAD cofactor to promote catalysis.\textsuperscript{129} However, KDM1s differ from other amine oxidases because their active sites are significantly expanded in order to accommodate the N-terminal residues of the histone H3 peptide (i.e. 1,245 Å\textsuperscript{3} in KDM1A).\textsuperscript{112} Within KDM1A, for example, there are four major invaginations lined with distinct groupings of residues to form complimentary interactions with H3 substrate side chains as well as for recognition of PTMs.\textsuperscript{111} No structural features within the active site of either enzyme have been found that might suggest that the methylation state of H3K4 can be distinguished sterically or electronically, and discrimination against H3K4me3 substrates is due to the inherent chemical mechanism of the enzymes.\textsuperscript{111,134}

Although more than 16 N-terminal residues of H3 are required for efficient catalysis, a significant portion of the C-terminus of substrates likely extends from the active site and potentially interact along a cleft formed at the interface of the AOD and SWIRM domain.\textsuperscript{111,112,115,141} Mutations of residues in this cleft in KDM1A abolish or abrogate demethylation activity, suggesting that interactions with substrates within this region may provide an additional specificity determinant.\textsuperscript{111} Additionally, surface plasmon resonance interaction studies have demonstrated that the SWIRM domain of KDM1A interacts with H3-derived peptides.\textsuperscript{152} In KDM1B, the surface landscape of this region is quite different and binds the coregulatory protein Nuclear Protein 60kDa/Glyoxylate Reductase 1 Homolog (NPAC/GLYR1).\textsuperscript{135,136}

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in KDM1A inhibition (Figure 2e). Finally, when KDM1B was co-crystallized in the presence of the same H3K4M peptide, it exhibited a similar binding conformation to that found in KDM1A (Figure 2d). Interestingly, elucidation of the structure of KDM1B bound to the cofactor NPAC/GLYR1 revealed a secondary binding site occupied by the C-terminal region of the H3 peptide tail.

Due to the above conformational differences and identification of the folate binding site within the KDM1A active site, there is some controversy as to which binding mode positions the K4 residue in the proper geometry that allows catalysis to proceed. Despite this discrepancy, both conformations reveal that the N-terminus of the peptides bind in an anionic pocket with a maximum of three residues N-terminal to the ethyllysine residue. Additionally, both conformations make extensive contacts with a large set of conserved, negatively charged residues in the active site cavity. Overall, these structures exemplify the apparent plasticity of the active sites of KDM1A and KDM1B and only hint at the richness of substrate specificity mechanisms available to this enzyme class.

**Table I  KDM1 Demethylase Family**

<table>
<thead>
<tr>
<th>Name</th>
<th>Synonyms</th>
<th>Histone Substrates</th>
<th>Non-Histone Substrates</th>
</tr>
</thead>
<tbody>
<tr>
<td>KDM1A</td>
<td>LSD1, AOF2, BHC110, KIAA0601</td>
<td>H3K4me1/2, H3K9me1/2</td>
<td>p53 (K370), DNMT1 (K1096), E2F1 (K185), ERz (K266), HIV Tat (K51), HSP90 (K615h/K614m), MTAT1 (K532), MYPT1 (K442), STAT3 (K140)</td>
</tr>
<tr>
<td>KDM1B</td>
<td>LSD2, AOF1, C6orf193</td>
<td>H3K4me1/2</td>
<td></td>
</tr>
</tbody>
</table>

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Based on these data, it is speculated that KDM1A and KDM1B can recognize and demethylate different substrates in vastly different binding modes.4

**Specificity for Non-Histone Substrates**

In addition to histone H3, non-histone proteins have been shown to be substrates for KDM1 family enzymes. KDM1A has been shown to demethylate p53, DNMT1, E2F1, MYPT1, STAT3, HIV Tat, HSP90, MTA1, and potentially ERz (Table I).103,154,155,168–174 To our knowledge, no non-histone substrates have yet been identified for KDM1B; however, we suspect that future work will also define multiple, albeit different substrates and roles for this enzyme. The existence of multiple non-histone substrates for KDM1A has strong implications in apoptosis, cell cycle progression, and the transcriptional activation or repression of associated genes.

The first non-histone substrate of KDM1A identified was the tumor-suppressor p53.154,155 Like histone proteins, p53 is subject to regulation by various PTMs. In particular, dimethylation of p53 at K370 (K370me2) promotes its association with 53BP1 (p53-binding protein 1), a protein that coactivates p53 transcriptional programs.175–177 Although in vitro KDM1A completely removes dimethylation at K370, in cell culture it preferentially removes a single methylation mark, generating the monomethylated species. This mark abrogates transcription through modulating p53-DNA interactions that subsequently repress the pro-apoptotic functions of p53. Further complicating this interaction, Barton and coworkers observed that KDM1A localizes only to genes where p53 represses transcription.154 This suggests some type of inherent regulation of the KDM1A-p53 interaction and may ensure repression of desired target genes.178

In contrast, in p53-deficient tumors, Set7/9 and KDM1A regulate DNA damage-induced cell death in a manner opposite to that observed in p53+/− cells, via modulation of the stability of the transcription factor E2F1.169 Kontaki and Talianidis illustrated that Set7/9 methylates E2F1 at K185 and prevents E2F1 accumulation during DNA damage and subsequent activation of its pro-apoptotic target gene, p73. This PTM is removed by KDM1A, and is required for E2F1 stabilization and downstream pro-apoptotic function. Interestingly, the molecular mechanism of these events involves crosstalk between lysine methylation and other PTMs that affect E2F1 stability and turnover.

Recently, Sakane and coworkers have identified KDM1A as a HIV Tat K51me1-specific demethylase that is required for the activation of HIV transcription in latently infected T cells.103 The RNA-binding domain of Tat is a known region containing PTMs179 in which K51 is methylated by Set7/9.180 This PTM corresponds to an early event in the Tat transactivation cycle and strengthens the interaction of Tat with TAR RNA.180 On the other hand, acetylation of K50 is mediated by p300/KAT3B, which dissociates the complex formed by Tat, TAR RNA, and the cyclin T1 subunit of the positive transcription elongation factor b (P-TEFb).181,182 Subsequent to K50 acetylation, the histone acetyltransferase, PCAF/KAT2B, is recruited to the elongating RNA polymerase II complex.183–185 Interestingly, the association of the KDM1A/CoREST complex with the HIV promoter subsequently activated Tat transcriptional activity in a K51-dependent manner. In accordance with Tat transcriptional activity, shRNAs directed at KDM1A or its inhibition by phenelzine suppressed the activation of HIV transcription in latently infected cells. The study again illustrates the dual nature of KDM1A and its ability to function as both a transcriptional suppressor and activator depending on the context of the substrate and interacting partners.

In addition to p53 and E2F1, Chen and coworkers have identified DNA methyltransferase 1 (DNMT1) as a novel substrate of KDM1A.168 Targeted deletion of the gene encoding KDM1A (a02) in mouse embryonic stem cells induces the progressive loss of DNA methylation in addition to an increase in the methylation of DNMT1 and a decrease in the DNMT1 protein level. Chen and coworkers illustrated that Set7/9 methylation of K1096 of DNMT1 can be reversed by KDM1A in vitro. Despite this evidence, it remains to be determined if K1096 is the major targeted site of KDM1A, since DNMT1 is methylated on multiple sites in vivo. By acting on and demethylating DNMT1, KDM1A may play a role in the coordination of DNA and histone methylation.

In another example, like E2F1, the transcription factor STAT3 is subject to PTMs that modulate downstream events in response to different cytokines and growth factors.186 Following phosphorylation, STAT3 is methylated on K140 by SET7/9 and subsequently demethylated by KDM1A when it is bound to a subset of promoters that it activates.171 Timing for this process occurs in an ordered sequence on the SOC3 promoter, illustrating the underlying temporal and spatial control of the methylation and demethylation events. The authors conclude that lysine methylation of promoter-bound STAT3 leads to biologically important downregulation of the dependent responses. This work illustrates the ability of methyltransferases and demethylases to control the methylation status of a transcription factor in a promoter-specific manner and provide a way to modulate the time of residence and promoter occupancy of the protein. Additionally, the transcription factors or the proteins recruited during these events may also provide a docking site for other proteins to carry out important functions, a role suggested for KDM1A by Mattevi and coworkers.152
Myosin phosphatase target subunit 1 (MYPT1) is an example of a KDM1A non-histone substrate that has a downstream affect on cell cycle progression.\textsuperscript{170} MYPT1, a known regulator of phosphorylation of the transcription factor retinoblastoma protein 1 (RB1), is methylated \textit{in vitro} and in cell culture by SET7/9 and subsequently demethylated by KDM1A, in which K442 is the target residue. The methylation status of MYPT1 influences the affinity the protein maintains for the ubiquitin-proteasome pathway and subsequent protein turnover. Interestingly, overexpression of KDM1A, which is prevalent in many cancer subtypes (see above), could activate RB1 phosphorylation by inducing a destabilization of the MYPT1 protein. Subsequently, the release of E2F could activate transcription of genes required for S phase to enhance cell cycle progression.

An extremely unique target of KDM1A is metastatic tumor antigen 1 (MTA1), a core subunit of the NURD complex (discussed in detail later). Interestingly, it is the only known dual function coregulator with an expected corepressor activity, but discussed in detail later). Interestingly, it is the only known dual function coregulator with an expected corepressor activity, but unusual ability to also stimulate transcription.\textsuperscript{187,188} MTA1 is a core subunit of the NURD complex (dissected in detail later). Interestingly, it is the only known dual function coregulator with an expected corepressor activity, but critical for S phase to enhance cell cycle progression.\textsuperscript{170} MYPT1, a known regulator of phosphorylation of the transcription factor retinoblastoma protein 1 (RB1), is methylated \textit{in vitro} and in cell culture by SET7/9 and subsequently demethylated by KDM1A, in which K442 is the target residue. The methylation status of MYPT1 influences the affinity the protein maintains for the ubiquitin-proteasome pathway and subsequent protein turnover. Interestingly, overexpression of KDM1A, which is prevalent in many cancer subtypes (see above), could activate RB1 phosphorylation by inducing a destabilization of the MYPT1 protein. Subsequently, the release of E2F could activate transcription of genes required for S phase to enhance cell cycle progression.

INHIBITION OF FLAVIN-DEPENDENT KDM1 DEMETHYLASES

Discovery and Optimization of Mechanism-Based Inhibitors

The discovery and development of KDM1 family inhibitors has been the subject of several recent comprehensive reviews; therefore, an abbreviated overview of KDM1 inhibitors is presented, which includes many recent developments.\textsuperscript{15,21,69,142,195–199} Motivated by the relationship between the AOD of KDM1A to that of MAO-A/B and PAO, the first KDM1A inhibitors were discovered by McCafferty and coworkers.\textsuperscript{200} Through screening a focused group of irreversible and reversible amine oxidase inhibitors, tranylcypromine (\textit{trans}-2-phenylcyclopropylamine, 2-PCPA, Parnate\textsuperscript{TM}, 1) was found to exhibit the highest KDM1A inhibitory activity towards methylated bulk histones as well as methylated nucleosomal substrates \textit{in vitro}. Treatment of P19 embryonal carcinoma cells with 1 resulted in the global increase of H3K4 methylation as well as transcriptional derepression of two KDM1A target genes, Egr1 and the pluripotent stem cell marker Oct4. This was the first example of small molecule inhibition of histone demethylation and interruption of transcriptional programs regulated by KDM1A.\textsuperscript{200}

In a subsequent study, our group determined that tranylcypromine was a mechanism-based inactivator of KDM1A, forming a covalent adduct with the FAD cofactor and demonstrating inactivation kinetic parameters of $k_i = 242$ $\mu$M and $k_{\text{inac}} = 0.0106$ s$^{-1}$.\textsuperscript{127} Soon after, the laboratories of Cole\textsuperscript{118} Yokohama,\textsuperscript{119,122} and Mattevi\textsuperscript{121} provided additional support for this inhibitory mechanism, including defining the structure of the tranylcypromine-KDM1A adduct using X-ray methods, which revealed the NS of the FAD isoalloxazine ring as the site of covalent attachment of the inhibitor.

In addition to tranylcypromine (1), our study revealed less potent KDM1A inhibition from the hydrazine-containing phenelzine (2) and the propargylamines, represented by pargylene.
KDM1A inhibitors 1 and 2 achieved complete inhibition in vitro at significantly lower concentrations than are physiologically relevant. However, propargylamines were not inhibitory at 5 mM, and only partial inhibition of KDM1A was observed at 30 mM. Exposure of cells to such non-physiologically relevant concentrations of 3 causes inhibition of proliferation and alteration of transcriptional programs via mechanisms that do not necessarily involve KDM1A activity. As such, one should approach the use of members of this inhibitor class with caution when attempting to attribute phenotypic effects exclusively to KDM1A inhibition.

Analysis of the crystal structures of KDM1A, MAOs, and PAOs, lead our group to conclude that KDM1A may be distinguished from related amine oxidases since MAO-A/B and PAO have significantly restricted active site volume when compared to KDM1A. As such, we designed, synthesized, and characterized analogues of 1 for the purpose of increasing potency and specificity towards KDM1A through aromatic substitutions and heteroaromatic exchanges. Within this initial group of inhibitors, improved selectivity and potency was observed, and these analogues were subsequently utilized to probe the role of KDM1A activity in estrogen receptor signaling. Since that time, numerous research groups, in addition to our own, have further optimized the potency and selectivity of arylcyclopropylamines for KDM1A inhibition. In addition to the small molecules discussed above, unmethylated product peptides (7) of the KDM1A reaction as well as a related substrate-like inhibitor containing a methionine point mutant (8; H3K4M; pK4M) have been shown to be competitive inhibitors of KDM1A with enhanced binding over methylated substrates. Recently, Kumarasignhe and Woster have also developed a series of cyclic peptides based on the substrate-like inhibitor pK4M that were more hydrolytically stable than the acyclic analogues. Additionally, moderate anti-tumor effects were observed in MCF7 and Calu-6 cancer cell lines. On the other hand, Mattevi and coworkers have developed a series of short peptide reversible inhibitors based on the N-terminal region of SNAI1, some of which exhibited anti-

**Peptide-Based Inhibitors**

Protein-protein and protein-ligand interactions provide auspicious starting points for the development of high affinity peptide and peptidomimetic inhibitors. Although this class of molecules has traditionally suffered from poor stability and cell permeability, modifications to the native peptide structure can greatly increase their pharmacokinetic properties and therapeutic value. This inhibition strategy is especially appealing in regard to KDM1A, as it is involved in multiprotein complexes, several of which are beginning to be biophysically characterized in detail (see below). Here we outline the existing peptide-based active site inhibitors of the KDM1 class demethylases with the hope that they will catalyze the development of peptide-based inhibitors that target a wider range of KDM1 interfaces.

In addition to the small molecules discussed above, unmethylated product peptides (7) of the KDM1A reaction as well as a related substrate-like inhibitor containing a methionine point mutant (8; H3K4M; pK4M) have been shown to be competitive inhibitors of KDM1A with enhanced binding over methylated substrates. Additionally, moderate anti-tumor effects were observed in MCF7 and Calu-6 cancer cell lines. On the other hand, Mattevi and coworkers have developed a series of short peptide reversible inhibitors based on the N-terminal region of SNAI1, some of which exhibited anti-
proliferative activity.123,124 Both classes of peptides show promise and may serve to lay the foundation for the development of peptidomimetic small molecule inhibitors.

In addition to reversible peptide inhibitors, several groups have also adopted an inhibitor design strategy whereby flavin-reactive moieties are grafted as ‘warheads’ onto peptides derived from histone H3, including reactive N-propargyl, -cyclopropyl, -aziridine, -phenelzine, -vinylchloride, or -tranylcypromine substituents (compounds 9–17; Figure 4).201,208,209 Within this series, phenelzine-containing H3K4-derived peptide (15) exhibited the most potent inhibitory activity in vitro;201 however, none of these compounds exhibited significant KDM1A inhibitor activity in cellular models, presumably due to poor cellular uptake or susceptibility to proteolytic degradation.210 Additionally, although alone parargylamine is a very poor inhibitor of KDM1A,147,200,201 presentation of the propargylamine group as a pendant substituent within a peptide derived from histone H3 (9, 10) significantly enhanced its activity as a mechanism-based inactivator (Figure 4).116,201,208

**Emerging KDM1A Inhibitor Classes**

Recently, Jung and coworkers have reported the development of lysine mimetics that possess a propargyl warhead for covalent modification of KDM1A (18,19; Figure 5).211 Although these compounds were inhibitory at low micromolar concentrations, they did not exhibit selectivity over the MAOs and PAO in this first generation series. This class of molecules may likely gain selectivity through future optimization efforts.

Similarly, Suzuki and coworkers have reported a series of KDM1A inactivators in which tranylcypromine (1) is coupled to a lysine carrier moiety at the nitrogen atom (20; Figure 5).212,213 The nonpeptidic small molecules selectively and efficiently delivered 1 to the KDM1A active site creating the inactivated, FAD-adduct in a time-dependent manner. Additionally, the molecules exhibited potent cell growth inhibitory activities against cancer cell lines. Together, the high potency and selectivity towards KDM1A allows this class of molecules to be considered good candidates for further optimization and implementation as chemical biology probes.

The considerable homology KDM1A shares with PAOs, including spermine oxidase (SMOX/SMO/PAO/PAO-1/PAOH1) and others, spurred Woster and Casero to develop a class of reversible KDM1A antagonists capable of derepressing tumor suppressor genes, as well as inducing other beneficial anticancer activities. The first class of polyamine analogues contained bisguanide and bisguanidine functional groups (21, 22) that inhibited KDM1A activity in cell and animal models of cancer.214–216 Since that time, they have expanded the KDM1A inhibitor repertoire to also include (bis)urea and (bis)thiourea analogues (23–25) that have exhibited excellent potency, cell permeability, and selectivity for KDM1A inhibition.217 In a recent example,66 the polyamine (1,15-bis(N5-[3,3-(diphenyl)propyl]-N1-biguanido)-4,12-diazapentadecane) (22; Figure 6) induced cytotoxicity and inhibited KDM1A activity in human AML cell lines. Exposure to this agent increased global levels of H3K4me1/2 and exhibited reexpression of the tumor suppressor E-cadherin.

Polyamine analogues have enormous potential as therapeutics for many classes of human cancers, including breast, prostate, and AML. Also, there is potential for these molecules as probes for understanding the contribution of KDM1A to

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**FIGURE 4** Structures of substrate-based peptide inhibitors.
specific epigenetic demethylation events that contribute to cancer pathobiology. As such, the discovery and development of polyamine analogues as reversible KDM1A inhibitors has been the subject of several recent seminal reviews.19,197,218–222

Merging key pharmacophoric features of reported KDM1A inhibitors, Dulla and coworkers recently reported the development of 3-amino/guanidine substituted phenyl oxazole-containing inhibitors.223 Following treatment of cultured cells with nanomolar inhibitor concentrations, viability decreased as compared to the low micromolar range observed for inhibition of KDM1A in vitro. Although the basis of the enhanced reactivity in cell culture and the degree of KDM1A selectivity remains to be fully explored, one inhibitor (26; Figure 7) avoids zebrafish embryo apoptosis and toxicity and may constitute a lead for further optimization. Additionally, it remains to be determined if this class is selective for KDM1A since these compounds were designed based on existing broad spectrum amine oxidase inhibitors.

Other examples of bifunctional KDM1A inhibitors have also recently been described. First, Zheng and coworkers reported a novel triazole-dithiocarbamate scaffold that inhibits KDM1A more potently than 1.54 The 1,2,3-triazole moiety within these inhibitors is readily accessible through the Cu(I)-catalyzed Huisgen 1,3-dipolar cycloaddition of azides with alkynes (i.e. Sharpless “click” chemistry).224,225 Additionally, this pharmacophore has been shown to possess numerous biological activities,226,227 including inhibitory activity towards MAO-A.228,229 On the other hand, the dithiocarbamate pharmacophore was chosen for its intrinsic inhibitory activity against fungi, bacteria, and malignant cancer.230–232 One compound synthesized in the series (27; Figure 7) effectively reduced tumor growth of human gastric cancer cells in vivo.

In another example, Rotili and coworkers synthesized hybrid KDM1A/JmjC bifunctional inhibitors. This was achieved by coupling the skeleton of the KDM1A inhibitor tranylcypromine (1), with 4-carboxy-4’-carbomethoxy-2,2'-bipyridine or 5-carboxy-8-hydroxyquinoline chelating groups that were developed as competitive inhibitors for the iron(II)/α-KG-dependent JmjC demethylases.86 Two compounds in this series (28, 29; Figure 7) increase H3K4 and H3K9 methylation levels in cells and cause growth arrest and substantial apoptosis in LNCaP prostate and HCT116 colon cancer cells.86

Fortuitously, Yang and coworkers decided to test natural polyphenols as potential inhibitors of KDM1A.233 Dietary polyphenols are known to have beneficial effects against diabetes, cancer, and cardiovascular disease, and exposure has been linked to both antioxidant activity and modulation of signaling pathways.234–236 Yang and coworkers demonstrated that resveratrol (30), curcumin (31), and quercetin (32) displayed inhibitory activity against KDM1A in vitro, which was independent of their antioxidant properties (Figure 8). However, since quercetin forms colloid-like aggregates in aqueous solution it may inhibit KDM1A in vitro non-specifically due to aggregate formation.237 Based on this precedence, further insight into its mode of action as well as structurally-related compounds is needed.

Screening has also been a means for identifying novel scaffolds against KDM1A and KDM1B and has yielded several anti-KDM1A inhibitor candidates. By combining protein structure similarity clustering and in vitro compound screening, Beuttner and coworkers discovered the KDM1A inhibitory activity of the γ-pyrene, Namoline, in vivo and in vitro (33, Figure 9).85 The IC₅₀ of this compound was reported to be 51 μM against KDM1A and was fully reversible. Namoline

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**FIGURE 5** Structures of nonpeptidic, warhead small molecule inhibitors that are substrate derived mimics.

**FIGURE 6** Structures of bisguanide and bisguanidine polyamine inhibitors.

*Biopolymers (Peptide Science)*
administration to prostate cancer cells led to silencing of AR-regulated gene expression and impairment of androgen-dependent proliferation in vitro and in vivo. Namolene may be a promising starting point (i.e. core structure) for the development of therapeutics to treat androgen-dependent prostate cancer.

On the other hand, Sharma and coworkers recently reported a structure-based virtual screen of a compound library containing ~2 million small molecules against KDM1A. Computational docking and scoring followed by biochemical screening led to the identification of a novel N’-(1-phenylethylidene)-benzohydrazide series of inhibitors. Hit-to-lead optimization and structure-activity relationship studies aided the discovery of a specific, reversible inhibitor (34) of KDM1A with a reported IC\textsubscript{50} of 31 nM. Compound 34 inhibits proliferation and survival in breast and colorectal cancer cell lines.

Zha and coworkers have also garnered recent success with a pharmacophore-based virtual screening approach to identify novel inhibitors of KDM1A. Using a moderate database of a commercial compound library and molecular docking tools, the group was able to identify nine candidate KDM1A inhibitors that lacked structural similarity to previously identified inhibitor classes. These compounds were subsequently tested in vitro against KDM1A and showed IC\textsubscript{50} values in the low micromolar range. One compound, XZ09 (35), which showed the highest potency against KDM1A, was also tested versus MAO-A/B and showed moderate selectivity. The XZ09 core structure represents a lead compound that, with further modification, could serve as another tool in probing KDM1A function.

Finally, Cole and coworkers recently reported the design, synthesis, and characterization of the KDM1A inhibitory properties of phenelzine analogues. Bizine (36; Figure 10), a novel phenelzine analogue containing a phenyl-butyrylamide appendage, exhibited good in vitro activity against KDM1A (K\textsubscript{i} = 0.06 μM; k\textsubscript{inact} = 0.15 min\textsuperscript{-1}; k\textsubscript{inact}/K\textsubscript{i} = 2.5 μM\textsuperscript{-1} min\textsuperscript{-1}) and was selective versus MAO-A/B (26- and 60-fold, respectively) and KDM1B (>100-fold). Bizine exhibited anti-proliferative effects in some cancer cells and was found to be effective at modulating bulk histone methylation. Additionally, these compounds synergistically inhibited cellular proliferation in combination with several histone deacetylase inhibitors. Interestingly, neurons exposed to oxidative stress were protected by the presence of bizine.

**FUNCTIONALLY IMPORTANT INTERACTIONS OF KDM1A AND KDM1B DEMETHYLASES WITH BIOMOLECULES**

KDM1 family demethylases typically operate as catalytic subunits within specific stable complexes formed with additional biomolecules and enzymes that perform coregulatory or scaffolding functions. Such interactions link the catalytic activity of KDM1s to distinctive biological occupations, and have been shown to influence the degree of catalytic activity, substrate specificity, and/or localization of these enzymes within chromatin.

**Interaction of KDM1A with CoREST and CoREST-like Proteins in Heteromeric Complexes**

CoREST is one of the earliest identified interaction partners of KDM1A and this protein complex assembly (i.e. KDM1A/CoREST complex) is frequently found within several distinct, larger multi-protein complexes. Most likely due to this early association, the structural and functional characterization of this particular interaction far outstrips that available for any other KDM1A interaction. CoREST consists of an N-terminal ELM2 domain followed by dual SANT domains (SANT1s and SANT2) with an intervening region colloquially known as the linker domain (Figure 11f). The N-terminal region spanning the ELM2 domain and first SANT domain have been mapped as a binding region for HDAC1, and a
The crystal structure of HDAC1 bound to a similar ELM2/SANT region in the CoREST-related protein, MTA1, has recently been reported. Shi and colleagues first mapped the interaction of KDM1A to the CoREST linker domain, indicating that CoREST likely acts as a scaffolding protein that joins deactylase and demethylase activity into a single catalytic sub-complex. The CoREST SANT2 domain, and possibly the SANT1 domain, has been shown to facilitate demethylation of nucleosomes, presumably by acting as a bridge between KDM1A and its substrate. A similar mechanism is observed in the SMRT protein, where the C-terminal SANT2 domain stabilizes the protein complex on chromatin by interacting with histone tails. As a point of further clarification, Yang and coworkers highlighted the similarity between the SANT2 domains of CoREST and the viral DNA-binding protein v-Myb, and demonstrated that an isolated CoREST SANT2 domain binds directly to DNA with modest affinity (\( K_d = 84 \text{mM} \)).

Seeking more detailed molecular characterization, Yang and colleagues reported the first crystal structure of KDM1A in complex with the linker and SANT2 domains of CoREST in 2006 (Figure 11a). In this model, the CoREST linker domain forms an L-shaped helical conformation, with the short helix contacting the base of the KDM1A tower domain, the longer helix extending up the TaB helix, and the SANT2 domain contacting the top turn of the tower (Figure 11a, insert). These interactions are mainly hydrophobic in nature, although a few electrostatic interactions may facilitate proper alignment of the CoREST linker with the KDM1A tower domain. A second crystal structure of KDM1A and CoREST with a histone substrate-like peptide bound (pK4M) showed little overall conformational change as compared to the non-peptide-bound structure. However, the model does illustrate that the L-shaped short arm formed by the CoREST linker may have an indirect effect on substrate binding by stabilizing the helix formed from KDM1A residues 372–395.

As noted above, although the CoREST SANT2 domain contacts the KDM1A tower in reported crystal structures, Mccafferty and coworkers have demonstrated that the CoREST linker domain is chiefly responsible for high affinity binding with KDM1A. Specifically, a CoREST fragment consisting of the linker domain (residues 293–380) exhibits low nanomolar binding affinity towards KDM1A \( (K_d = 7.78 \text{nM}) \), while inclusion of the SANT2 domain does not substantially alter binding affinity and in isolation lacks significant affinity towards KDM1A. Our group also used mutagenesis and isothermal titration calorimetry to reveal that the energy of binding along the CoREST/KDM1A helix-helix interface is not concentrated into "hotspots," but is instead distributed almost uniformly along the interaction interface. As CoREST is required for nucleosomal demethylation, this argues that inhibitors of the CoREST linker/KDM1A tower interaction may inhibit KDM1A functionality in the cell.

In addition, MD studies by Baron and colleagues have suggested that the KDM1A/CoREST interaction may be dynamic. Specifically, their simulations indicate that the KDM1A SWIRM and CoREST SANT2 domains oscillate in distance and rotational angle in relation to each other, and that substrate binding allosterically rigidifies this fluctuation, favoring an "open" state. They further suggest that blocking these "hinge sites" may prevent KDM1A/CoREST association with chromatin or other protein partners and therefore provide another potential strategy for modulating KDM1A function.

In addition to the regulatory mechanisms enforced on KDM1A by CoREST, two CoREST paralogs, CoREST2 (RCOR2, 523 aa) and CoREST3 (RCOR3, four isoforms of 436, 449, 495, and 553 aa), have recently been revealed as further governors of KDM1A activity. These paralogs share a similar domain organization, including conserved ELM2 and dual SANT domains. Not surprisingly, CoREST2 and CoREST3 both interact with KDM1A and are capable of incorporation into larger protein complexes. However, CoREST2 displays a reduced ability to facilitate KDM1A nucleosomal demethylation and transcriptional repression as compared to CoREST. This weakening of repressive activity is attributed to residue L165 in the CoREST2 SANT1 domain, which

FIGURE 9 Structures of screen identified and validated inhibitors.

FIGURE 10 Structures of mechanism-based phenelzine analogs.
diminishes its affinity for HDACs 1 and 2. However, CoREST3 plays an even more antagonistic role, as it competitively inhibits KDM1A nucleosomal demethylation. A CoREST3 chimera with the SANT2 domain replaced by the corresponding domain in CoREST reverses this phenotype and reinstates KDM1A nucleosomal demethylation. Collectively, these studies clearly indicate that CoREST and its paralogs play important roles in a subset of KDM1A-dependent transcriptional regulatory events by acting as a scaffold for the assembly and recruitment of deacetylase/demethylase-containing complexes, and by serving as a bridge to link epigenetic enzymes to nucleosomal substrates.

**REST Complex.** RE1-silencing transcription factor (REST) is a supervisor of cell fate decisions during neuronal differentiation, and functions as a transcription repressor that localizes to RE1 motifs in the promoters of neuron-specific genes (Figure 11b). In multipotent progenitor cells, these genes are held in a 'primed-repressed' state, in which REST recruits corepressor complexes mSin3 and CoREST to its N- and C-terminal repressor domains, respectively, while simultaneously maintaining an activating chromatin environment and basal-level gene transcription. Upon differentiation into a neuron cell, REST is largely degraded to activate a neuronal expression program, although the CoREST complex sustains a role in tempering the expression of certain genes. Alternately, upon differentiation into a non-neuronal cell type, the REST/mSin3/CoREST repressor machinery persists on RE1-containing promoters and adjusts chromatin in the vicinity of neuron-specific genes to a repressive state. Specifically, H3K4 demethylation is executed by KDM1A, and H3K9 methylation is likely mediated by the histone methyltransferases (HMTs) G9a.
and EHMT1, which are bridged to REST by the corepressor protein CDYL. The HMTs SUV39H1 and SETDB1 have also been linked to the CoREST complex and may play a role in H3K9 methylation. Increased CpG methylation (mCpG) in RE1 motifs has also been observed in differentiated cells compared to embryonic stem cells, although the DNA methyltransferase responsible has not yet been identified.

The H3K4 and H3K9 methylation status within chromatin encompassing repressed neuron-specific genes further recruits additional corepressors. For example, demethylated H3K4 is bound by PHF21A (BHC80), which stabilizes KDM1A on chromatin and enhances repression. However, PHF21A-bound by PHF21A (BHC80), which stabilizes KDM1A on additional corepressors. For example, demethylated H3K4 is encompassing repressed neuron-specific genes further recruits methyltransferase responsible has not yet been identified.

The H3K4 and H3K9 methylation status within chromatin encompassing repressed neuron-specific genes further recruits additional corepressors. For example, demethylated H3K4 is bound by PHF21A (BHC80), which stabilizes KDM1A on chromatin and enhances repression. However, PHF21A exists as multiple isoforms, is expressed tissue-specifically, and has been reported to paradoxically inhibit KDM1A activity and REST-mediated repression, indicating that its activity may be context-specific. Similarly, REST-associated G9a recruits CBX3 (HP1) to methylated H3K9me2 residues, where it is known to bind through its chromodomain and maintain a heterochromatric state. In another example, the mCpG-binding protein MeCP2 is also recruited to the larger corepressor complex for stable gene silencing, although it also appears to play a role in transient gene repression between neuron depolarization events.

Other accessory proteins that may not directly interact with chromatin are also recruited to regulate gene repression. For example, REST repression is reportedly dependent upon sumoylation of HMG20B (BRAF35) and its recruitment to the CoREST complex. Previous studies have suggested that sumoylated regulators may bind the CoREST/KDM1A interaction surface, although it is unclear if this is the case for HMG20B. In contrast, the repressive activity of the REST complex is antagonized by HMG20A (iBRAF), which forms a heterodimer with HMG20B to prevent its sumoylation and incorporation into the CoREST complex. In fact, HMG20A is a critical regulator of neuronal differentiation that promotes neuron-specific gene expression by recruitment of the H3K4 HMT, MLL.

**TLX Orphan Nuclear Receptor Complex.** KDM1A also regulates neuronal processes through its interaction with the orphan nuclear hormone receptor TLX (NR2E1). In line with KDM1A’s role in neuronal differentiation, TLX maintains the proliferative and self-renewal properties of adult neural stem cells and regulates neurogenesis. Yokoyama and coworkers first reported that TLX interacts directly with the AOD and SWIRM domains of KDM1A and further demonstrated that TLX recruits the core CoREST/KDM1A/HDAC1 complex (Figure 11c). Additionally, recruitment of the complex occurs in a KDM1A-dependent manner for H3K4 demethylation, H3 deacetylation, and downstream target gene repression. A subsequent report revealed that HDAC5 also associates with this complex. Interestingly, immunoprecipitation of TLX also coprecipitates several other proteins, including ZMYM3 and two that are also found in the SNAI1 complex (see below), namely GSE1 and ZMYM2. Despite these results, it is currently unclear whether any crosstalk exists between these complexes.

Within these complexes, Sun and coworkers also reported that the TLX/KDM1A ensemble regulates neural stem cell proliferation. Specifically, the microRNA, miR-137, antagonizes this proliferative activity and encourages differentiation by downregulating KDM1A. On the other hand, while in complex with KDM1A, TLX represses miR-137 expression. This feedback regulation likely contributes to timely neuronal differentiation events in which KDM1A is intimately involved.

**TAL1 Complex.** TAL1 is a transcription factor that is involved in normal hematopoiesis and leukemogenesis and can act as both a transcriptional activator and repressor, depending upon the genomic context and stage of cellular differentiation. Specifically, TAL1 promotes transcription of erythroid-specific genes through association with p300 and PCAF but can also associate with the mSin3A and ETO2 complexes to induce gene repression. KDM1A is recruited by TAL1 along with the core CoREST/HDAC complex to repress erythroid-specific genes in progenitor cells prior to differentiation (Figure 11d), but is dismissed from TAL1 promoters during the early stages of differentiation. The PTM of TAL1 may regulate this gene activation, as phosphorylation of S172 by protein kinase A (PKA) abolishes TAL1 interaction with KDM1A and results in recruitment of the H3K4 HMT, hSET1, activation of target genes, and initiation of erythroid differentiation. It is currently unclear if the KDM1A/CoREST core complex cooperates with any of the other known TAL1-interacting corepressor complexes to stimulate gene repression.

**CtBP Complex.** C-terminal binding proteins (CtBPs) are implicated widely as repressors of mammalian gene expression, and likely function in these varied contexts by recruiting different subsets of corepressors. KDM1A was first coprecipitated with the CtBP complex prior to discovery of its demethylase activity and has since been associated with several CtBP functions, including pituitary gland development, suppression of the tumor suppressor BRCA1, and activation of tissue-specific genes in gastrointestinal endocrine cells. However, the best-established role of KDM1A in the CtBP complex is its control of the EMT through suppression of the epithelial gene E-cadherin.

On the molecular level, CtBPs interact with coregulators (Figure 11e) mainly through two interaction domains: a
hydrophobic pocket that canonically binds proteins containing a Pro-X-Asp-Leu-Ser (PXDLS) sequence motif and a structurally distinct surface crevice that binds an Arg-Arg-Thr (RRT) motif. Additionally, CtBP proteins form extensive scaffolds by dimerization in a manner that positions the PXDLS binding pockets at opposite ends of the dimer, and in such a manner that the PXDLS pocket of one CtBP protein is in relatively close proximity to the RRT binding crevice of the second CtBP molecule. Initial studies indicated that the PXDLS-binding region mediates CtBP genomic localization by associating with DNA-binding transcription factors containing PXDLS motifs. However, later studies revealed that this site also recruits accessory proteins with histone-modification activity. For example, deacetylase activity is recruited by PXDLS motifs in class II HDACs and possibly by class I HDACs bridged through other PXDLS-containing corepressor proteins, such as NRP140 and LCoR. Similarly, PXDLS-like motifs in Wiz recruit the HMTs G9a and EHMT1, and possibly CDYL and HDAC1/2 to the CtBP complex. Several other proteins co-precipitate with CtBP, although not all have well-defined functions.

Interestingly, direct binding of CtBP1 to CoREST and HDAC2, respectively, has been reported despite the lack of PXDLS motifs in these proteins, and seem to bridge KDM1A to the complex. However, KDM1A recruitment to the CtBP complex also appears to be mediated by ZNF217,288,292 a large DNA-binding zinc finger protein that spans the CtBP dimer interface to interact with both PXDLS and RRT domains simultaneously. ZNF217 localizes the CtBP complex for transcriptional repression of several disease-related genes, such as E-cadherin and BRCA1. Other DNA-binding proteins have also been reported to bridge KDM1A to the PXDLS-binding pocket and appear to dramatically alter the demethylase's biological function. For example, recruitment by ZEB1 can extend gene repression programs associated with pituitary differentiation, while recruitment by RREB1 can actually promote CtBP/KDM1A-mediate gene activation. Clearly, KDM1A activity within the CtBP complex may have far-reaching biological implications that are most likely determined by the presence of specific coregulatory subunits.

NuRD Complex. The nucleosome remodeling and histone deacetylase (NuRD) complex is a large, multi-protein complex that has been implicated as a regulator of a wide variety of cellular processes, from human development to the progression of several cancer types. Since its discovery, the majority of its core protein constituents have been well defined: the ATP-dependent chromatin remodelers Mi-2 α/β (CHD3/4), histone deacetylases HDAC1/2, methyl-CpG-binding proteins MDB2/3, retinoblastoma-binding proteins RBBP4/7, corepressors GATAD2A/B (p66x/β), and metastasis-associated proteins MTA1/2/3 (Figure 11g).

Interestingly, the MTA proteins share striking similarities with the CoREST proteins. Specifically, the MTA isoforms possess, from N- to C-termini, BAH, ELM2, SANT and GATA-like zinc finger domains (Figure 11f). As previously mentioned, the ELM2 and SANT domains serve to recruit HDACs 1 and 2, presumably in similar fashion to CoREST. Wang and coworkers first established that the MTA2 protein, like CoREST, is sufficient to stimulate KDM1A demethylation of nucleosome substrates and mapped an interaction interface involving the KDM1A tower domain and MTA1/2/3 SANT domains. The authors further demonstrated that this interaction recruits H3K4 demethylase activity to the NuRD complex for enhanced gene repression, and that KDM1A association with each MTA isoform corresponds to a unique transcriptional outcome. Functionally, the authors linked KDM1A in the NuRD complex to silencing of the TGFβ signaling pathway and repression of the EMT and metastatic potential in a breast cancer cell line. Since this initial study, KDM1A has been associated as a corepressor of the NuRD complex in a subset of its biological functions, such as lipid homeostasis, aberrant gene repression in Ewing sarcoma, and enhancer decommissioning during embryonic stem cell differentiation. It is, however, noteworthy that a recent mass spectrometry-proteomics coupled study did not identify KDM1A as a stable component of the NuRD complex in HeLa cell extracts, indicating that it likely interacts in a context-specific manner.

While the MTA proteins resemble CoREST in their ability to recruit HDAC/KDM1A, several domains unique to MTA suggest a unique functionality, possibly through altered interactions with chromatin or coregulatory proteins. In fact, the KDM1A/NuRD complex has been shown to also associate with the x-KG dependent histone demethylase KDM5B. In this context, KDM5B and KDM1A cooperate to sequentially demethylate H3K4 and repress the metastasis and angiogenesis-associated CCL14 chemokine pathway. However, it remains unclear if KDM5B recruitment is dependent upon a specific MTA isoform. A recent report from Nair and colleagues suggests a regulatory role of KDM1A in NuRD activity. The authors show that G9a methylation of MTA K532 is required for its incorporation into the repressive NuRD complex, while KDM1A-mediated demethylation of the same residue triggers its dissociation from the NuRD complex and recruitment to the coactivator complex, NURF. Although these studies provide much needed insight into the function of KDM1A within the NuRD complex, further investigation into the demethylase activity of KDM1A within the heteromeric complex is needed.
KDM1A Cooption During Herpes Simplex Virus Infection

Cooption of host KDM1A activity plays a role in infection and latency of herpes simplex virus (HSV) and has recently been reviewed. Briefly, HSV establishes an active infection at the portal of entry (typically a mucosal membrane), where the host cell initially silences the inserted viral genome. Through a series of coordinated derepression events, viral immediate-early (IE) stage (α) genes are first transcribed and eventually give rise to transcription of late-stage (β and γ) genes. Although the primary infection produces massive viral titers that are destructive toward the host tissue, the virus also establishes a latent infection in neuronal cells, in which viral genes are present but silenced. This state is non-destructive to the host cell and allows for periodic reemergence of the virus through poorly understood mechanisms.

The HSV genome is parsimonious, lacking much of the cellular machinery required for transcription and regulation of viral genes. The virus instead appropriates host cell transcriptional regulators, including KDM1A, to establish active and latent infections. Gu and coworkers first associated the REST/CoREST/HDAC repressor complex, and later KDM1A, with HSV infection (Figure 12). Consistently, Pinnoji and colleagues identified REST-binding RE1 motifs in the viral genome, suggesting a potential mechanism for cooption of the complex. Indeed, REST appears to aid in repression of viral genes during latent infection, as genomic insertion of a REST mutant incapable of binding corepressor proteins produces a more virulent strain compared to wild-type REST. Similarly, HDAC inhibitors reactivate silenced viral gene expression, while REST antagonizes reactivation.

KDM1A also plays a role in viral gene activation. In order to induce transcription, the viral protein VP16 recruits host proteins HCF1 and OCT1 to target gene promoters. Liang and coworkers report that HCF1 recruits KDM1A and HMTs, SET1 and MLL1, for H3K9 demethylation and H3K4 methylation, respectively (Figure 12). Accordingly, the authors also report that inhibition of KDM1A prevents activation of viral genes and viral reemergence from latency. Interestingly, the JMJD2 family H3K9 HDMs work synergistically with viral activation and emergence from latency. Addi-

KDM1A Involvement in Nuclear Hormone Signaling AR Complexes.

The AR is a nuclear hormone receptor that is intimately linked to prostate function, regulating processes ranging from normal tissue development to the genesis and progression of prostate cancers, including castration-resistant tumors. Metzger and colleagues first reported KDM1A association with AR, suggesting that a large portion of the two proteins are involved in their mutual interaction and that the demethylase facilitates hormone-dependent AR-mediated transcription by demethylating H3K9 residues. Subsequent studies demonstrated that the Jumonji-domain demethylase, KDM4C, also interacts with KDM1A/AR and works in concert with KDM1A to demethylate H3K9me3 residues. Concurrently, hybrid inhibitors that simultaneously target both KDM1A and KDM4C exhibit anti-proliferative effects in prostate cancer cells. Although KDM1A canonically demethylates H3K4 residues, Metzger and colleagues proposed a mechanism for the observed switch in substrate specificity by which protein kinase Cβ1 (PKCβ1) associates with AR/KDM1A at target genes following hormone treatment and phosphorylates H3T6, a modification that prevents KDM1A demethylation of H3K4 on peptide substrates (Figure 13a).

KDM1A may also play a role in repression of AR target genes. Unlike AR, KDM1A resides on AR target gene promoters even in the absence of androgen treatment, when these genes are presumably repressed. Furthermore, the KDM1A/AR complex forms a negative feedback loop at high androgen levels, in which AR recruits KDM1A to an enhancer of the AR gene and mediates gene repression by demethylation of H3K4 (Figure 13a). Although the role of KDM1A as a coregulator of AR signaling is fairly well established, further work will hopefully better delineate its opposing roles as both an activator and repressor.

ERα Complexes.

Paralleling the function of the AR, the ERα is the main modulator of estrogen (E2) signaling in estrogen-responsive tissues. As such, its dysregulation results in the development and progression of several cancer types. Recently, much attention has focused on the epigenetic
mechanisms that accompany ERα signaling. As observed for KDM1A participation in AR signaling, the demethylase seems to function as both an activator and repressor of ERα signaling. Perillo and coworkers reported that in a ‘resting state’ with no hormone stimulation, KDM1A is constitutively bound to ERα target gene promoters and demethylates H3K4. However, E2 addition causes the recruitment of ERα to these promoters and KDM1A-dependent gene activation, most likely by catalyzing the demethylation of inhibitory H3K9 methyl marks. HMTs such as SET9 and SUV39H1 may also participate in E2-stimulated gene activation by methylating H3K4 (Figure 13b).

The molecular mechanisms governing the activating and repressive roles of KDM1A are beginning to be described. For example, PELP1, an ERα coregulator with no known enzymatic activity, can interact directly with KDM1A, localizes on the KDM1A/KDM1A complex causes dissociation from the promoter, accumulation of H3K9me2 marks, and gene repression.
known KDM1A/ERα-bound genes, and aids in gene activation.\textsuperscript{321} When associated with KDM1A/ERα, the histone binding preference of PELP1 switches from H3K9me2 to H3K4me2, presumably to allow the subsequent demethylation of H3K9me2 by KDM1A (Figure 13b).\textsuperscript{321} Interestingly, a recent report suggests that targeting this KDM1A/PELP1 activity may be of therapeutic benefit in breast cancer.\textsuperscript{78} KDM1A may also demethylate ERα to promote transcriptional activation, although further validation of this activity is required.\textsuperscript{323} More indirectly, KDM1A demethylation events and the accompanying production hydrogen peroxide byproducts also reportedly induces controlled DNA damage that attracts chromatin remodelers to enhance chromatin plasticity and promote target gene transcription.\textsuperscript{194} In opposition to PELP1-mediated activation, KDM1A also interacts with the ERα corepressor CAC1, which binds KDM1A/ERα, reduces occupancy of the complex on target genes, allows H3K9me2 accumulation, and represses gene transcription (Figure 13b).\textsuperscript{322} Additionally, retinoic acid may interfere with the KDM1A-mediated E2 response, as it appears to prevent activation of protein kinase A (PKA) and prohibits H3K9 demethylation, although the mechanism is not yet clear.\textsuperscript{323}

In addition to these roles in ERα signaling, KDM1A may also perform ERα-independent functions in breast cancer. For example, inhibition of the demethylase resensitizes hormone-refractory breast cancer cells to anti-estrogen treatment.\textsuperscript{78} In ERα-negative breast cancer, an interplay between KDM1A and HDACs is apparently necessary for tumor progression, as both affect the activity of the other towards histone substrates and simultaneous administration of KDM1A and HDAC inhibitors synergistically inhibits proliferation.\textsuperscript{79,80} Also, overexpression of the ERα corepressor, CAC1, induces sensitivity to paclitaxel in ERα-positive, paclitaxel-resistant breast cancer cells.\textsuperscript{322} These studies collectively imply that KDM1A functions in a wide range of ERα activities, and is additionally involved in various resistance mechanisms.

**SNAG Family Proteins Interact with KDM1A Using Product Mimicry**

The Snail/Gfi1 (SNAG) family of proteins encompasses over 70 different members characterized by a conserved SNAG domain and between four to six C\textsubscript{2}H\textsubscript{2} zinc finger domains.\textsuperscript{324} Zinc finger domains are generally located near the C-terminal regions of SNAG proteins and function as DNA-recognition domains that frequently localize to E-box sequences (CANNTG), although recruitment of individual proteins is likely tailored by minor variations in zinc fingers domains and DNA sequences.\textsuperscript{324} In all but one known example, SNAG domains begin at the most N-terminal residue and are composed of a conserved “PRSFLV” sequence, where only the fourth residue is variable (Figure 14a).\textsuperscript{324} Proteins in the SNAG family of transcription factors have been reported to regulate KDM1A activity through product-like interactions, in which the SNAG domain mimics the histone H3 N-terminal tail sequence and binds competitively to the KDM1A active site.

Lin and colleagues first identified an interaction between the SNAG domain of SNAI1 and the AOD of KDM1A and demonstrated that it could be disrupted with small molecule inhibitors.\textsuperscript{97} Soon after, they showed that SNAI1 recruits KDM1A to target promoters where demethylation of H3K4 is required for gene repression.\textsuperscript{90} Following this discovery, Baron and colleagues determined the crystal structure of KDM1A and CoREST in complex with the 20 N-terminal residues of SNAI1 encompassing the SNAG domain, revealing density for the first nine residues (Figure 14b).\textsuperscript{123} In this report, the authors indicate that the SNAG domain mimics histone H3 through several conserved, positively charged residues and binds in the active site of KDM1A, resulting in competitive inhibition. Similar to the H3 tail, the first three SNAI1 residues form hydrogen-bonds deep in the active site, while Phe4 (F4), corresponding to Lys4 on H3 (H3K4), forms edge-to-face interactions with the flavin cofactor and KDM1A residue Y761. On the outer rim of the active site, Arg7 and Lys8 (R7 and K8) of SNAI1 seem to occupy the same positions as Arg8 and Lys9 of H3 (H3R8 and H3K9), respectively. Although direct evidence has not yet been provided, KDM1A presumably interacts with other SNAG proteins, including SNAI2, Gfi1, and Gfi1b, in a similar manner. In fact, treatment with either the KDM1A inhibitor tranylcypromine (1) or a cell-permeable SNAG peptide phenocopies down-regulation of either KDM1A or SNAI2 in decreasing the EMT.\textsuperscript{325}

However, these studies raise an interesting question: if the SNAG domain is inhibitory toward KDM1A, how does it recruit the enzyme for demethylation of its target genes? One possible scenario is that the SNAG domain recruits KDM1A, but then relinquishes its hold in order to transfer binding to the H3 tail. In a recent study by Tortorici and colleagues, the first six residues of the SNAI1 SNAG domain were found to be sufficient for KDM1A binding, while residues 10–20 increased affinity only marginally and decreased ligand efficiency.\textsuperscript{124} On the other hand, the corresponding residues of H3 form multiple interactions with KDM1A,\textsuperscript{117} and hence may allow H3 to displace the SNAG domain.

**SNAI1/2 Complexes.** SNAI1 (Snail, Snail1) and SNAI2 (Slug) are master regulators of the EMT and bind to E-box motifs in the promoters of epithelial genes, such as E-cadherin, to repress transcription and provoke an invasive, mesenchymal phenotype.\textsuperscript{91,326} KDM1A recruitment to target genes and
demethylation of H3K4me2 is required for SNAI1-mediated gene repression. As discussed above, the SNAI1/2 SNAG domains recruit KDM1A via its active site. As such, small molecule inhibitors or isolated SNAG domain peptides can displace SNAI1, derepress target genes, and decrease invasiveness and motility in colon and neuroblastoma cancer cell lines without affecting proliferation. Interestingly, the chemotherapeutic drug doxorubicin has the opposite effect, enhancing the interaction between SNAI1 and KDM1A by inducing poly(ADP-ribosyl)ation of SNAI1. The authors further suggest that this increased interaction causes repression of the tumor-suppressor gene, PTEN, and may facilitate drug-resistance in cancer.

SNAI1 also coprecipitates the core KDM1A/CoREST/HDAC complex, as well as several auxiliary proteins such as HMG20A/B and PHF21A, which are associated with the REST complex, and CtBP, GSE1, and several zinc finger proteins, which are associated with the CtBP complex. It is currently unclear if SNAI1 recruits these larger KDM1A-containing complexes to further regulate gene expression. Recent studies have also identified the chromatin ‘reader’ SFMBT1, previously identified with the CoREST complex and other repressive complexes, as a member of the SNAI1/KDM1A complex. SFMBT1 binds directly to the CoREST linker region and mediates recruitment of KDM1A to SNAI1, gene repression, and induction of the EMT. Additionally, it recognizes H3K4me2/3 marks, and therefore may help to target the complex to active genes for subsequent repression (Figure 14c).

**Gfi1/1b Complexes.** Gfi1 and Gfi1b are SNAG-containing transcription factors that are critical for hematopoiesis. While
Gfi1 regulates the differentiation programs of hematopoietic stem cells, small lymphocytes (B and T cells), dendritic cells, granulocytes and macrophages, Gfi1b is involved in the development of megakaryocytes and erythrocytes. As might be expected, both proteins are also involved in hematopoietic-derived cancers, although their roles as promoters or repressors of cancer progression seem to vary significantly depending upon tissue origin.

As with the SNAI1 and 2 proteins, Gfi1 and Gfi1b recruit the core KDM1A/CoREST/HDAC complex via their SNAG domains, presumably by binding the KDM1A active site. Furthermore, Gfi1b recruits KDM1A in this manner to the majority of its target genes for demethylation of H3K4me2 residues and transcriptional repression. However, paralleling the tissue-dependent roles of Gfi1/1b, KDM1A differentially affects proliferation depending on the hematopoietic cell type and differentiation status. For example, splice variant Gfi1b P32, is required for erythroid differentiation and recruits the CoREST/KDM1A complex more efficiently than the major Gfi1b isoform when dimethylated on Lys8 (K8) in its SNAG domain. Similarly, the three CoREST isoforms all interact with Gfi1b but have varying effects on its activity (see above). Gfi1b has also been associated with the H3K9 HMTs SUV29H1 and G9a, although these proteins have not been explicitly shown to co-localize with Gfi1b/CoREST.

**KDM1A Association with Long Non-Coding RNAs**

A new class of interactions has recently been identified between KDM1A and long noncoding RNAs (lncRNAs). lncRNAs, broadly defined as non-protein coding RNA transcripts greater than 200 nucleotides, are differentially expressed in a number of human cancers and can regulate transcription by acting as scaffolds for chromatin modifying complexes. The HOTAIR transcript, which is dramatically overexpressed in several cancers and can be induced by estradiol, recruits the PRC2 and KDM1A/CoREST/REST complexes for downregulation of tumor suppressor genes at the HOXD locus. Binding interactions for EZH2 and KDM1A were observed at the 5'- and 3'-terminal domains of HOTAIR, respectively, and HOTAIR expression was required for co-immunoprecipitation and genomic co-localization of PRC2 protein complex and KDM1A (Figure 15a). These interactions and the corresponding changes in histone methylation states were also observed in vivo. The HOTAIR/PRC2/KDM1A complex has also been implicated as a positive regulator of NFAT5, a transcription factor associated with enhanced metastasis and angiogenesis in breast cancer. In contrast, glioblastoma cell cycle progression appears to be dependent upon HOTAIR and PRC2 but not KDM1A, implying these corepressors may play specific, non-interdependent roles in conjunction with HOTAIR. More extensive reviews of HOTAIR and its role in cancer have recently been published.

In 2013, a second breast cancer-associated lncRNA, SRA (steroid receptor RNA activator), was reported to associate with KDM1A-containing repressive complexes, though in this case the interaction was mediated by progesterone receptor (PR). Most recently, Porro and colleagues have demonstrated an interaction between KDM1A and the telomeric lncRNA TERRA. The authors report that TERRA, which consists of UUAGGG repeats and has been
implicated in the progressive shortening of telomeres, recruits KDM1A to telomeric DNA in the absence of the protective telomere repeat-binding factor 2 (TRF2). KDM1A does not seem to affect histone methylation levels in this context but instead promotes the removal of telomere 3′ guanine overhangs by enhancing the nuclease activity of MRE11, an enzyme involved in telomere 3′ end processing. Both TERRA and MRE11 interact with the SWIRM and N-terminal AOD region of KDM1A, but do not interact directly themselves. The authors therefore speculate that TERRA enhances the affinity of KDM1A for MRE11 and hence promotes telomeric shortening, although a direct substrate of KDM1A was not identified (Figure 15b). Given that the biological significance of lncRNAs is just beginning to be appreciated, future reports of KDM1A interactions with lncRNAs are anticipated.

Interaction of KDM1B with Proteins and Multi-Protein Complexes

Compared to the identified roles of KDM1A described above, much less is known about the interactions and biological significance of KDM1B. In contrast to KDM1A, KDM1B does not possess a coiled-coil tower domain, and hence does not interact with CoREST14 nor cooperate with HDACs in breast cancer.79 However, it has been reported as an effector of genomic imprinting,106 and its activity is required for somatic cell reprogramming108,109 and liver development.107 KDM1B has more recently been implicated as a regulator of breast cancer proliferation, where inhibition works synergistically with DNMT and KDM1B inhibitors to derepress aberrantly silenced genes and decrease cell growth.359

KDM1B has been reported to function as a positive regulator of gene transcription by binding to chromatin in the highly transcribed, H3K36me3-enriched coding regions down-stream of gene promoters.105 Here it helps to maintain H3K4 and H3K9 methylation by associating with a larger complex that includes Pol II and other elongation factors, as well as the SET-family HMTs NSD3 and G9a, which methylate histone H3K36 and H3K9 sites, respectively (Figure 16a).105 Later studies indicated that the H3K36me3 reader NPAC/GLYR1 is likely part of this complex as well, and enhances KDM1B demethylation of H3K4me1/2 by binding at its AOD/SWIRM interface (Figure 16b).136 Finally, KDM1B is recruited to the promoters of inflammatory genes by the NF-κB subunit c-Rel and is required for subsequent H3K9 demethylation, recruitment of NF-κB, and gene expression.110 Given the intricate mechanisms involved in regulating and modulating KDM1A activity, it is likely that future work will uncover an analogous but unique set of protein partners that govern KDM1B activity.

CONCLUSIONS

KDM1s function as oxidative demethylases, removing methyl marks on protein substrates that provide chemical cues for the regulation of transcription. As such, KDM1A and KDM1B are central to numerous transcriptional programs in normal and disease-associated biology. These include regulating differentiation processes of both hematopoietic and neuronal tissues, cancer proliferation and metastasis, and viral replication and emergence from latency. Within these enzymes exist interaction motifs unique to each KDM1 isofrom that facilitate interactions with biomolecules that regulate catalysis, promote KDM1 enzyme recruitment, and direct access to specific
substrates at explicit sites within the cell. These coregulatory molecules include proteins, product-like inhibitors, and lncRNAs. Recently, transcriptional outcomes have been correlated to specific interactions of KDM1A or KDM1B with coregulatory biomolecules. In fact, interactions with differing subsets of coregulators can lead to KDM1 involvement in opposing transcriptional programs. Although several classes of KDM1-selective small molecule inhibitors have been recently developed, pan-inhibition strategies lack the selectivity capable of discriminating between KDM1 activities in these specific functional contexts. The development of next-generation chemical biology probes and therapeutics that modulate these interactions may further deconvolute the complex biology of epigenetic regulation. We suspect that due to the sheer size of these multiprotein complexes that these probes will either be peptides, peptoids, or peptide-like in nature.

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Biopolymers (Peptide Science)
A rationally-designed chimeric KDM1A/KDM1B histone demethylase tower domain deletion mutant retaining enzymatic activity

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A target with therapeutic potential, lysine-specific demethylase 1A (KDM1A) is a regulator of gene expression whose tower domain is a protein–protein interaction motif. This domain facilitates the interaction of KDM1A with coregulators and multiprotein complexes that direct its activity to nucleosomes. We describe the design and characterization of a chimeric ‘towerless’ KDM1A, termed nD150 KDM1A(Tower) KDM1B chimera (chKDM1A(Tower)), which incorporates a region from the paralog lysine-specific demethylase 1B (KDM1B). This chimera copurifies with FAD and displays demethylase activity, but fails to bind the partner protein corepressor of the RE1-silencing transcription factor (CoREST). We conclude that KDM1A catalysis can be decoupled from tower-dependent interactions, lending chKDM1A(Tower) useful for dissecting molecular contributions to KDM1A function.

1. Introduction

In 2004, lysine-specific demethylase 1A (KDM1A also known as LSD1/BHC110/AOF2/KIAA0601/p110b) became the first histone demethylase to be isolated and characterized [1] and has subsequently been identified as a potential therapeutic target [2–4]. KDM1A is a flavin-dependent amine oxidase that demethylates mono- and dimethylated lysine residues at positions 4 and 9 on histone H3 (H3K4me1/2 and H3K9me1/2), leading to gene activation and repression, respectively [1,5,6]. This 852 amino acid (aa) polypeptide is composed of three structured domains (Fig. 1A). The catalytic amine oxidase domain (AOD) houses a single, non-covalently bound flavin adenine dinucleotide (FAD) cofactor required for catalysis while the N-terminal SWI3p, Rsc8p, and M0ira (SWIRM) domain is commonly found in chromatin-associating proteins (Fig. 1B) [7–10]. The ‘tower’ domain is a nearly 100 aa AOD insert that forms an approximately 90 Å long antiparallel coiled–coil from two α-helices (termed TaA and TaB) linked by a tight turn (Fig. 1B) [7,8]. Unprecedented among related amine oxidases [1], the tower domain serves as a protein interaction motif and facilitates KDM1A incorporation into multiprotein regulatory complexes that dictate its cellular function [11].

One of the best-studied tower domain interaction partners is the corepressor of the RE1-silencing transcription factor (CoREST), which facilitates demethylation of nucleosomal substrates by bridging KDM1A to chromatin through its DNA-binding SANT domains [8,12–17]. Additionally, CoREST orchestrates the inclusion of enzymes such as KDM1A and histone deacetylases as catalytic subunits within modular multiprotein complexes [18,19]. KDM1A activity is further governed by other homologous proteins that bind the tower domain and perform similar functions in different molecular contexts, such as the

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Abbreviations: AOD, amine oxidase domain; chKDM1AA(Tower), nD150 KDM1AA(Tower) KDM1B chimera; CoREST, corepressor of the RE1-silencing transcription factor; KDM1A, lysine-specific demethylase 1A; KDM1B, lysine-specific demethylase 1B; SWIRM, SWI3p, Rsc8p, and Moira

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CoREST paralogs RCOR2 and RCOR3 and the MTA proteins [17,20–22]. Therefore, one of the most significant challenges is understanding the impact of these tower-dependent interactions on KDM1A specificity, catalysis, and positioning within chromatin.

In order to evaluate these tower-dependent interactions independent of demethylase activity, a KDM1A tower domain deletion mutant is highly desired. Indeed, several groups have generated such mutants, but were limited by a lack of enzymatic activity [7,8,21,23]. These studies suggest that either the tower domain is inherently required for KDM1A catalysis or that the methods chosen to bridge the tower region were not conducive to proper active site folding. Interestingly the recent characterization of the KDM1A paralog KDM1B provides a unique opportunity to investigate this dichotomy [24]. KDM1B lacks a tower domain (Fig. 1A) but conserves the AOD architecture, sharing a 2.0 Å RMSD compared to KDM1A, despite a modest sequence similarity (<25%) (Fig. 1B and C). As this paralog overcomes exclusion of the tower domain in a manner that preserves a KDM1A-like active site conformation, we sought to use it as a template for engineering a chimeric KDM1A enzyme.

Here we report the rational design and characterization of a KDM1A tower domain deletion chimera. KDM1A and KDM1B sequence and structural alignments suggested that KDM1B residues V494–L531 could replace the KDM1A tower domain (residues T389–R524) with minimal active site disturbance. This construct, termed chKDM1AΔTower, copurified from Escherichia coli cellular lysates with a stoichiometric equivalent of FAD. As expected, chKDM1AΔTower failed to bind CoREST. However, unlike previous tower deletion mutants, our chimera exhibits kinetic parameters nearly identical to those of unaltered KDM1A and KDM1B, suggesting that the tower domain is not required for catalytic activity. The chKDM1AΔTower chimera therefore decouples tower-dependent protein interactions from catalysis and provides a tool to assess the effects of KDM1A mistargeting and orphanization.

**Fig. 1.** Domain maps and structural overview of KDM1A and KDM1B. (A) Domain maps of KDM1A and KDM1B. SWIRM domains are shown in green, AODs are shown in magenta, tower domain is shown in blue, C4H2C2 domain is shown in orange, Zf-CW domain is shown in cyan, and linker is shown in purple. Structural overview of (B) KDM1A (PDB 2IW5) and (C) KDM1B (PDB 4HSU). Coloring scheme follows that of domain map above, FAD cofactor is shown in yellow, and N and C-termini are labeled. Zinc ions in KDM1B structure are shown as gray spheres.
2. Materials and methods

2.1. Reagents and materials

Clones of genes encoding *Homo sapiens* nA150 KDM1A (UniProtKB accession No. O60341) and full-length KDM1B (UniProtKB accession No. Q8NB78) were codon optimized for *E. coli* by GenScript (Piscataway, NJ) and subcloned into pET-15b (Novagen) with Ndel and Xhol (New England Biolabs; NEB). The pDB-HisGST vector was obtained from the DNASU Plasmid Repository. Buffer salts were obtained from Sigma, EMD Millipore, and JT Baker. Tween 20 was obtained from AMRESCO. Protein purification was conducted using an ÄKTA FPLC (Amersham Biosciences).

2.2. Alignment of KDM1A and KDM1B and generation of a chimera model

Primary amino acid sequences of KDM1A and KDM1B were aligned using Clustal Omega [25]. A sequence alignment figure (Fig. 2A) was generated utilizing ESPript 3.0 [26]. Structural alignment of PDB files 2IW5 and 4HSU (KDM1A [13] and KDM1B [27], respectively) was conducted with PyMOL [28] (Fig. 2B). To generate a composite model of the chimera, the headers of PDB files 2IW5 and 4HSU were deleted and coordinates superimposed using Coot [29]. KDM1A tower domain residues T389–R524 were replaced with KDM1B residues V494–L531. The resultant chain was renumbered and coordinates of the chimera composite model exported into PyMOL (Fig. 2C). All-atom contacts were validated with MolProbity [30].

2.3. Cloning of chKDM1A.Tower

Joining KDM1A fragments L151–A388 and D525–M852 to KDM1B residues V494–L531 formed the chimera sequence. A previously described pET-15b vector containing 6×His nA150 KDM1A (residues 151–852) [31] was used as a template for construction of the chimera. This entire vector was amplified with exclusion of the KDM1A tower domain (residues T389–R524) using primers that incorporated Sall and KpnI restriction sites and PFU Turbo DNA polymerase (Agilent) under the following conditions: an initial denaturation step for 2 min at 95 °C, 30 cycles of denaturation for 30 s at 95 °C, annealing for 30 s over a gradient from 54 to 65 °C, elongation for 8 min at 72 °C, and a final elongation step for 10 min at 72 °C. The KDM1B insert (residues V494–L531) was amplified with complementary restriction sites under similar conditions, but with an elongation of 1 min at 72 °C. The resulting insert and vector were digested and ligated utilizing a Quick Ligation Kit (NEB). Restriction sites were removed using a Q5 Site-Directed Mutagenesis Kit (NEB) under the following conditions: an initial denaturation step for 1 min at 98 °C, 30 cycles of denaturation for 30 s at 98 °C, annealing for 30 s over a gradient from 59 to 70 °C, elongation for 4 min at 72 °C, and a final elongation step for 2 min at 72 °C. For primers, see Table S1.

2.4. Expression and purification of KDM1A and chKDM1A.Tower

Expression and purification of wild type nA150 KDM1A was conducted as previously described [15,31]. The 6×His-tagged nA150 KDM1A.Tower chimera in pET-15b (i.e. chKDM1A.Tower) was expressed in BL21 Star (DE3) *E. coli* (Novagen) at 15 °C. chKDM1A.Tower was purified under similar conditions to wild type nA150 KDM1A with minor modifications. Approximately 4.5 mg of chKDM1A.Tower per liter of culture were obtained at >95% purity. For more details, please see the Supplementary Material.

2.5. Cofactor analysis of chKDM1A.Tower

The method of Aliverti et al. was employed to determine the FAD molar extinction coefficient [32]. chKDM1A.Tower in gel filtration buffer at 1.5–2.0 mg/mL was incubated with SDS to a final concentration of 0.3% (w/v) and monitored until the UV trace stabilized. 15 μM FAD disodium salt hydrate (Sigma) in gel filtration buffer was used as a standard. The chimera was calculated to carry FAD in 1:1:1 ratio (FAD:protein) and to have a molar extinction coefficient of 10350 M⁻¹ cm⁻¹ at 455 nm using ε450 = 11300 M⁻¹ cm⁻¹ for free FAD (FADfree). This value was routinely used to determine protein concentration.

2.6. Expression and purification of His-GST-CoREST-C and His-GST

The His-GST-CoREST-C construct was expressed in BL21 (DE3) *E. coli* (Novagen) at 19 °C. Protein was purified by immobilized metal affinity chromatography (IMAC) and ion exchange chromatography (IEC). Approximately 2.0 mg protein per liter of culture were obtained at >75% purity. The His-GST construct was expressed and purified in a similar manner. For more details, see the Supplementary Material.

2.7. Steady-state demethylase assay

A peptide corresponding to the N-terminal 21 amino acids of histone H3 with a dimethylated K4 residue (H3K4me2–21) was prepared as previously described [31,33]. A continuous, fluorescence-based, steady-state kinetic assay was employed as previously described with slight modifications [31,33]. An HRP-coupled (1 U/mL of HRP) assay monitored enzymatic peroxide production in 50 mM Tris–HCl (pH 7.85) with 0.01% (w/v) CHAPS using Ampex Red (50 μM) as the fluorogenic electron acceptor. All measurements were performed in a final volume of 60 μL at 25 °C in 96-well format (Corning 3693). Reactions were initiated with the addition of enzyme (0.35 μM final concentration). The product, resorufin, was monitored with 535 nm excitation and 597 nm emission wavelengths. Data were analyzed using GraphPad Prism 6.0 (GraphPad Software, San Diego, CA). All data sets were corrected for background and zero by subtraction of the no enzyme control and the initial time point, respectively. Initial velocities within the 10% product conversion limit were fit to the Henri–Michaelis–Menten equation with non-linear regression analysis.

2.8. His-GST-CoREST-C pull-down interaction assay

A total of 15 μg of purified His-GST-CoREST-C (residues 286–482) or 9 μg His-GST were incubated with 12 μg nA150 KDM1A or 10 μg chKDM1A.Tower at 4 °C for 16 h in 100 μL of binding buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4 (pH 7.4), 10 mM DTT, and 0.05% (v/v) Tween 20) containing 20 μL of Glutathione Sepharose 4 Fast Flow beads (GE Healthcare). Flow-through (supernatant) was removed and beads were washed 3 times with 400 μL of binding buffer, resuspended in 80 μL of 2× SDS sample buffer, and denatured for 2 min at 100 °C. Input, beads (bound), and flow-through were analyzed by SDS–PAGE (4–20% gradient, Bio-Rad) and visualized with Coomassie Brilliant Blue.

3. Results

3.1. Design of the chKDM1A.Tower chimera from KDM1A and KDM1B alignments

To rationally design a tower domain deletion mutant of KDM1A, we looked to its ‘towerless’ paralog KDM1B, which performs
identical chemical reactions but interacts with a non-identical set of biomolecules. We therefore used sequence and structural alignments to define a sequence from KDM1B that could replace the tower domain of KDM1A and maintain active site geometry. As predicted, a large gap in the sequence alignment of KDM1A and KDM1B reflects the KDM1A tower domain with areas of considerable sequence conservation on either side (Fig. 2A). We should note that a small internal area of apparent homology is likely an artifact of shared α-helical secondary structure. Subsequent structural alignment and analysis indicated that while the region C-terminal to the tower domain is highly similar between the two enzymes, the corresponding N-terminal region in KDM1B contains a solvent exposed loop not conserved in KDM1A (Fig. 2B). In order to maintain the interhelical contacts in this region (e.g. KDM1B W497 with K512 and I516 with V493), we extended the tower replacement sequence to include this entire loop and also supplanted a portion of the KDM1A α-helix that contacts the base of the tower domain (termed αCox or Sα1) [7,8]. The final chKDM1A Tower design incorporated these KDM1B elements into KDM1A, with a final sequence of KDM1A fragments L151–A388 and D525–M852 linked by KDM1B fragment V494–L531 (Fig. S1).

Next, a preliminary evaluation of this design was performed by generating a composite structural model of chKDM1A Tower using Coot (Fig. 2C). Subsequent visual inspection of the model in PyMOL revealed no gross structural abnormalities. Additionally, analysis of all-atom contacts with MolProbity [30] supported this observation, as no steric clashes in the replaced tower region were visible. We therefore predicted that replacement of the KDM1A tower with the KDM1B loop would result in a properly folded, active enzyme.

3.2. Construction, expression, and purification of the chKDM1A ΔTower chimera

We next sought to generate the chKDM1A ΔTower chimera in which the KDM1B loop was inserted into the KDM1A sequence in place of the tower domain using restriction digestion cloning. After insertion, restriction sites were removed to yield a ‘seamless’ chimera that contained only residues from the primary amino acid sequences of KDM1A and KDM1B (Fig. S2). The chimeric clone was then overexpressed in E. coli under nearly identical conditions as reported for wild type KDM1A [15,31] resulting in a visible band at the expected molecular weight of approximately 70kDa (Fig. S3A). Purification of chKDM1A ΔTower from E. coli cellular lysates using a series of chromatographic separations resulted in a >95% homogenous sample as assessed by SDS–PAGE (Fig. S3B).

Our purified chKDM1A ΔTower solution appears yellow in color (Fig. S3C), suggesting it retains the essential FAD cofactor as previously observed for KDM1A and KDM1B. Indeed, a UV–visible spectrum of the chKDM1A ΔTower produced characteristic flavin cofactor absorbance maxima at 380 and 455 nm (Fig. 3) [34]. A s compares to an authentic FAD standard, the molar extinction coefficient (FADbound) of the chKDM1A ΔTower construct was determined to be 10350 M⁻¹ cm⁻¹ at 455 nm [32,34] (Fig. 3).

Furthermore, full cofactor occupancy of chKDM1A ΔTower was noted, as a stoichiometric amount of FAD was released upon SDS treatment. This FAD cofactor was also released into solution as a function of heat or TCA denaturation (data not shown). These data therefore suggest that, like KDM1A and KDM1B, the FAD cofactor of the chimera is non-covalently bound and potentially poised to initiate catalytic demethylation.
3.3. *chKDM1A* Tower exhibits similar demethylase activity as *KDM1A* and *KDM1B*

As the *chKDM1A* Tower-bound flavin chromophore is suggestive of properly assembled catalytic machinery, we next tested for bona fide enzymatic activity. Steady-state kinetic parameters of *chKDM1A* Tower were evaluated by monitoring enzymatic production of *H₂O₂* using a peptide substrate. As shown in Fig. 4, *chKDM1A* Tower exhibited well-behaved linear velocities and demethylated H3K4me21–21 with a *kcat* of 3.21 ± 0.16 μM, a *kcat/kM* of 0.57 ± 0.01 min⁻¹, and a catalytic efficiency of 0.18 ± 0.01 μM⁻¹ min⁻¹. These values compare very favorably to that of wild type KDM1A and KDM1B enzymes (Table 1). In fact, the *kcat/kM* of *chKDM1A* Tower was identical to that of KDM1B, and only approximately 10-fold lower than that of KDM1A [24,31]. Interestingly, values of both *kcat* and *kcat/kM* of the chimera were situated neatly near those of the two enzymes (Table 1). These data suggest that *chKDM1A* Tower preserves elements from both KDM1A and KDM1B as anticipated, and clearly maintains integrity of the amine oxidase active site. Thus, by grafting the KDM1B loop onto KDM1A, we generated a tower domain deletion mutant that retains wild-type-like demethylase activity.

3.4. *chKDM1A* Tower does not interact with *KDM1A* binding partner CoREST

Having achieved one of our goals in engineering an active KDM1A tower deletion mutant, we next determined if a known tower domain interacting protein was precluded from binding the chimera. The tower domain of wild type KDM1A is a well-known protein recruitment motif for CoREST [8,12,13]. We therefore hypothesized that this interaction would be abrogated in the chimera. The tower domain interacting protein was precluded from binding the chimera.* A GST pull-down assay previously used to evaluate CoREST in vitro is mediated mainly through the tower domain and does not receive significant contributions from the AOD and SWIRM domains [8,13]. Hence our chimera effectively abrogates association with CoREST, and presumably any other protein that relies predominantly on the tower domain for binding.

4. Discussion

There is currently a remarkable lack of tools for studying the influence of tower-dependent interactions on KDM1A recruitment into modular multiprotein complexes, substrate specificity, and chromatin localization. To address this issue, we here report what is, to the best of our knowledge, the first KDM1A tower deletion enzyme that retains catalytic activity comparable to wild type KDM1A and KDM1B. Using sequence and structure-driven design principles, we have rationally selected boundaries for excision of the KDM1A tower domain and replacement with a bridging segment from KDM1B. This chimera, termed *chKDM1A* Tower, copurifies with a stoichiometric equivalent of FAD and exhibits catalytic activity directly overlays with a FAD standard (solid black line).

**Fig. 3.** *chKDM1A* Tower copurifies with a non-covalently bound FAD cofactor. *chKDM1A* Tower was evaluated by UV–Vis and has absorption maxima at 380 and 455 nm (solid green line). The chimera was denatured by treatment with SDS, releasing the cofactor (dashed blue line). The spectrum of the denatured sample directly overlays with a FAD standard (solid black line).

**Table 1**

<table>
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<th>Enzyme Name</th>
<th><em>kcat</em> (μM)</th>
<th><em>kcat</em> (min⁻¹)</th>
<th><em>kcat/kM</em> (μM⁻¹ min⁻¹)</th>
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<tr>
<td>nA150 KDM1A wt</td>
<td>2.60 ± 0.2</td>
<td>5.97 ± 0.78</td>
<td>2.2 ± 0.12</td>
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<tr>
<td>nA25 mKDM1B wt</td>
<td>1.13 ± 1.3</td>
<td>2.00 ± 0.60</td>
<td>0.18 ± 0.04</td>
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<tr>
<td><em>chKDM1ATower</em></td>
<td>3.21 ± 0.16</td>
<td>5.07 ± 0.01</td>
<td>0.18 ± 0.01</td>
</tr>
</tbody>
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* In 50 mM Tris–HCl (pH 7.85), 0.01% CHAPS (w/v), 50 μM Amplex Red & 1 U/mL HRP at 25 °C in air saturated buffer; peptide titrated from 50 μM to 0 in 2-fold dilution series (n = 3). *Values reported by Gaweska et al. [31].
* Values reported by Karytinos et al. [24].

These data are in agreement with previous reports that KDM1A interaction with the C-terminal fragment of CoREST in vitro is mediated mainly through the tower domain and does not receive significant contributions from the AOD and SWIRM domains [8,13]. Hence our chimera effectively abrogates association with CoREST, and presumably any other protein that relies predominantly on the tower domain for binding.
in their coregulated promoter regions, thereby functionally disrupting the transcriptional programs for this subset of genes. Beyond KDM1A, this strategy of splicing related regions between functionally related proteins may prove to be an invaluable tool for investigating domains for which standard domain deletion proves intractable. This strategy is specifically useful when investigating enzymes that exist as subunits of multimeric complexes.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.febslet.2015.07.028.

References

[12] Shi, Y.J., Matson, C., Lan, F., Iwase, S., Baba, T. and Shi, Y. (2005) Regulation of downstream genetic programming. For example, we suspect that chKDM1AATower will no longer localize with CoREST


The PyMOL Molecular Graphics System, Version 1.5.0.4, Schrödinger, LLC.


Lysine-Specific Demethylase 1A (KDM1A/LSD1): Product Recognition and Kinetic Analysis of Full-Length Histones

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Lysine-Specific Demethylase 1A
(KDM1A/LSD1): Product Recognition and Kinetic Analysis of Full-Length Histones

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ABBREVIATIONS

3,5-DCHBS, sodium 3,5-dichloro-2-hydroxybenzenesulfonate; 4-AAP, 4-
aminoantipyrine; AOD, amine oxidase domain; CHAPS, 3-[(3-
cholamidopropyl)dimethylammonio]-1-propanesulfonate; FAD, flavin adenine
dinucleotide; HRP, horseradish peroxidase; IMAC, immobilized metal affinity
chromatography; IPTG, isopropyl β-D-thiogalactoside; KDM1A, lysine-specific
demethylase 1A; KDM1B, lysine-specific demethylase 1B; LB, lysogeny broth; LSD1,
lysine-specific demethylase 1A; PMT, photomultiplier tube; PTM, posttranslational
modification; RU, response units; SDS, sodium dodecyl sulfate; SWIRM, SWI3p, Rsc8p,
and Moira; TB, Terrific Broth; THF, tetrahydrofolate; U, units
ABSTRACT: Lysine-specific demethylase 1A (KDM1A/LSD1) is a FAD-dependent enzyme that catalyzes the oxidative demethylation of histone H3K4me1/2 and H3K9me1/2 repressing and activating transcription, respectively. Although the active site is expanded compared to the greater amine oxidase superfamily, it is too sterically restricted to encompass the minimal 21-mer peptide substrate footprint. The remainder of the substrate/product is therefore expected to extend along the surface of KDM1A. We show that full-length histone H3 which lacks any posttranslational modifications is a tight-binding, competitive inhibitor of KDM1A demethylation activity with a $K_i$ of $18.9 \pm 1.2 \, \text{nM}$; a value that is approximately 100-fold higher than the 21-mer peptide product. The relative H3 affinity is independent of preincubation time suggesting that H3 rapidly reaches equilibrium with KDM1A. Jump dilution experiments confirmed the increased binding affinity of full-length H3 was at least partially due to a slow off-rate ($k_{off}$) of $1.2 \times 10^{-3} \, \text{s}^{-1}$, a half-life ($t_{1/2}$) of 9.63 min, and residence time ($\tau$) of 13.9 min. Independent affinity capture surface plasmon resonance experiments confirmed the tight-binding nature of the H3/KDM1A interaction, revealing a $K_d$ of $9.02 \pm 2.27 \, \text{nM}$, a $k_{on}$ of $9.26 \times 10^4 \pm 1.5 \times 10^4 \, \text{M}^{-1} \text{s}^{-1}$ and a $k_{off}$ of $8.35 \times 10^{-4} \pm 3.4 \times 10^5 \, \text{s}^{-1}$. Additionally, no other core histones exhibited inhibition of KDM1A demethylation activity, which is consistent with H3 being the preferred histone substrate of KDM1A versus H2A, H2B, and H4. Together these data suggest that KDM1A likely contains a histone H3 secondary specificity element on the enzyme surface that contributes significantly to its recognition of substrates and products.
Chromatin condensation and relaxation and the dynamic regulation of gene expression are regulated by posttranslational modifications (PTMs) of histone proteins by epigenetic enzyme complexes.\textsuperscript{1, 2} Dysregulation of histone-modifying enzymes within these complexes underscores several disease pathologies, including cancers.\textsuperscript{3} As such, this class of enzymes has become the subject of intense inquiry for their potential role as targets for therapeutic intervention. Among these PTMs is histone lysine methylation, a mark that is intimately linked to both transcriptional activation and repression.\textsuperscript{4}

Although evidence of enzymatic demethylation of histones would arise with the work of Paik and Kim in 1973,\textsuperscript{5} it would not be until almost four decades later that an enzyme directly responsible for this activity would be uncovered. The isolation and characterization of lysine-specific demethylase 1A (KDM1A also known as LSD1/AOF2/BHC110/KIAA0601/p110b) by Shi and coworkers provided direct evidence that histone lysine methylation was, in fact, a reversible mark.\textsuperscript{6} KDM1A is a flavin-dependent amine oxidase that catalyzes the removal of methyl groups of mono- and dimethylated lysine residues at position 4 and 9 on histone H3 (H3K4me1/2 and H3K9me1/2, respectively).\textsuperscript{6-8} From N- to C-terminus, KDM1A is composed of an unstructured region that contains the nuclear localization signal\textsuperscript{9, 10} and three structured domains: a SWI3p, Rsc8p, and Moira (SWIRM) domain, an amine oxidase domain (AOD), and an unprecedented intervening domain within the AOD colloquially known as the tower (Figure 1A).\textsuperscript{11, 12}

More specifically, the tower domain KDM1A is composed of two antiparallel α-helices (termed T\textalpha{}A and T\textalpha{}B) that form a coiled-coil and extend approximately 100 Å away from the AOD (Figure 1B).\textsuperscript{11, 12} Functionally, this domain is the site of association
with the corepressor of the RE1-silencing transcription factor (CoREST also known as CoREST1/RCOR1/KIAA0071) as well as several homologous proteins (Figure 1B).\textsuperscript{12-15} Interestingly, CoREST endows KDM1A with the ability to demethylate nucleosomal substrates as a non-catalytic domain donor with DNA binding ability bridging the gap between KDM1A and its substrates.\textsuperscript{15-19} On the other hand, the SWIRM domain of KDM1A, which is found in multiple chromatin-associated proteins,\textsuperscript{20} is a bundle of $\alpha$-helices with a central helix that is joined by two smaller helix-turn-helix motifs (Figure 1B).\textsuperscript{11} Moreover, consistent with other flavin-dependent oxidoreductases, the AOD of KDM1A houses a non-covalently bound flavin adenine dinucleotide (FAD) cofactor and can be further subdivided into an expanded Rossman fold for cofactor binding and the substrate-binding lobe (Figure 1B).\textsuperscript{12, 15} Although, these characteristics group KDM1A into the larger family of flavoenzymes, the structure of the AOD does contain unique features due to the functional activity of the enzyme.

As compared to other amine oxidases, KDM1A contains a more expansive substrate-binding cavity (~1,245Å$^3$) that is suggested to compliment its broad substrate specificity, including non-histone proteins.\textsuperscript{12, 21} Additionally, although KDM1A requires a peptide substrate of at least 21 residues representing the N-terminal portion of histone H3 for efficient catalysis,\textsuperscript{22} the active site cavity is too sterically restricted to accommodate this entire fragment.\textsuperscript{12} In efforts to decode substrate specificity, several groups have co-crystallized peptides with KDM1A that revealed two main conformations.\textsuperscript{23-25} Interestingly, the conformations result in the C-terminus of the peptide exiting the active site in different orientations. Further confounding these results, Luka and coworkers recently solved a structure of KDM1A in complex with
tetrahydrofolate (THF) and suggested that the cofactor may sterically disrupt one of the binding conformations.\textsuperscript{26} Prior to this finding, several groups have suggested that KDM1A substrates exit the active site and extend along a conspicuous surface groove along the SWIRM-AOD interface aiding in substrate binding and recognition.\textsuperscript{11, 15, 27, 28} Consistent with this hypothesis, mutational analyses of residues that lie in this groove greatly reduce or abrogate the catalytic efficiency of KDM1A.\textsuperscript{11} Additionally, Tochio and coworkers demonstrated that an isolated SWIRM domain of KDM1A is able to bind the N-terminal tail of histone H3 via surface plasmon resonance (SPR) studies.\textsuperscript{29} Confounding these results, however, a 30-mer H3 peptide substrate showed no apparent change in catalytic efficiency when compared to a 21-mer.\textsuperscript{22} Additionally, none of the structural studies listed above revealed electron density in the cleft along the SWIRM-AOD interface, raising the question as to whether this phenomenon exists and if so where the secondary binding site resides.

Herein we describe the kinetic analysis of full-length histone products against KDM1A as a preliminary investigation as to whether or not the enzyme recognizes additional residues of its products. We now show that histone H3 is a tight-binding, competitive inhibitor of KDM1A demethylation activity and has a 100-fold higher affinity as compared to the 21-mer peptide product. Additionally, the potency of H3 inhibition is independent of preincubation time suggesting that the enzyme and product reach rapid equilibrium. Through jump dilution experiments, we illustrate that the increased affinity is at least partially due to the slow off-rate of the full-length product. Additionally, the tight-binding nature of the histone H3 product interaction was independently confirmed via SPR. As suspected, no other histone proteins inhibit
KDM1A demethylation activity. Together, these results suggest that KDM1A contains a secondary histone H3 binding site on the enzyme surface that contributes significantly to recognition of its histone H3 substrates and products. Additionally, our work further validates the suggested role of KDM1A as a contributor to the stabilization of associated multiprotein complexes within genetic loci.

EXPERIMENTAL PROCEDURES

Reagents and Materials. The clone of the *Homo sapiens* protein nΔ150 KDM1A (UniProtKB accession no. O60341; residues 151-852) was codon optimized for expression in *Escherichia coli* and synthesized by GenScript (Piscataway, NJ). The pDB-HisGST expression vector was obtained from the DNASU Plasmid Repository. Buffer salts were obtained from Sigma, EMD Millipore, and JT Baker. Trans-2-Phenylcyclopropylamine hydrochloride salt (2-PCPA), 4-AAP, and 3,5-DCHBS were obtained from Sigma. Protein purification was conducted using an ÄKTA FPLC (Amersham Biosciences).

Expression and Purification of nΔ150 KDM1A. The gene encoding nΔ150 KDM1A was subcloned into a pET-15b vector (Novagen) containing an N-terminal 6xHis-tag with an intervening thrombin cleavage site, utilizing NdeI and XhoI restriction enzymes. The clone was expressed and purified as previously reported with minor modifications (See Supporting Information for detailed procedure). The final concentration of KDM1A was routinely determined by UV absorption spectroscopy.
using an extinction coefficient of 10,790 M⁻¹ cm⁻¹ at 458 nm, which was determined after
protein denaturation using 0.3% (w/v) SDS.²²,³¹

Expression and Purification of His-GST-CoREST-C. The gene encoding CoREST-C (CoREST1/RCOR1: UniProtKB accession no. Q9UKL0; residues 286-482) from *H. sapiens* was subcloned into a pDB-HisGST expression vector, containing an N-terminal 6xHis-tag and GST-tag with an intervening tobacco etch virus protease (TEV) cleavage site, utilizing NdeI and XhoI restriction sites. The clone was expressed and purified as previously reported with minor modifications.³² Following IMAC, fractions containing His-GST-CoREST-C were extensively dialyzed against GST-PBS dialysis buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.8 mM KH₂PO₄ (pH 7.4), 5 mM βME) overnight in 3.5 kDa MWCO SnakeSkin dialysis tubing (Thermo Scientific) and applied at 0.1 mL/min to 10 mL Glutathione Sepharose 4 Fast Flow resin (GE Healthcare). The column was washed at 2.0 mL/min with 5 CV of GST-PBS dialysis buffer followed by 10 CV of TEV cleavage buffer (50 mM Tris-HCl (pH 7.8), 150 mM NaCl, and 1 mM DTT). GST-tag cleavage was performed by incubating column-bound protein in a 0.1 to 1 ratio of 7xHis TEV (S219V), prepared by previously described methods,³³ to column bound protein overnight. Eluted CoREST-C was pooled and supplemented with 5 mM imidazole and passed over IMAC resin equilibrated with 50 mM Tris-HCl (pH 7.8), 150 mM NaCl, and 5 mM imidazole to remove TEV protease. Samples containing CoREST-C were pooled and concentrated using 5 kDa MWCO Vivaspin 20 centrifugal filters at 2000 × g. Samples were aliquoted, flash frozen in liquid N₂ and stored at −80°C. This method routinely provides protein at > 90% purity as
assessed by SDS-PAGE and with typical yields of 1.5 mg per liter of culture as determined by Bradford method.

Expression and Purification of His-GST-CoREST-Linker. The coding sequence of the linker region of CoREST (residues 293-380) was extracted and amplified by using PCR with a forward primer (5’-GCGCATATGGTCAAAAAAGAAAAACATAGC-3’) and a reverse primer (5’-GCGCTCGAGTTAATTACATTTCTGAATGACC-3’) under the following conditions: an initial denaturation step for 2 min at 95°C, 30 cycles of denaturation for 30 sec at 95°C, annealing for 30 min over a gradient from 54 to 65°C, elongation for 1 min at 72°C, and a final elongation step for 10 min at 72°C. The primers were designed to contain NdeI and XhoI restriction sites at the N- and C-termini, respectively, to allow for facile ligation into pDB-HisGST expression vector. The resulting His-GST-CoREST-Linker construct was expressed and purified as described above for His-GST-CoREST-C.

Purification of KDM1A/CoREST Complexes. Purified CoREST constructs were mixed with KDM1A in a 1.5:1 molar ratio and allowed to incubate for 1 h on ice. Incubated samples were then purified using a HiPrep 26/60 Sephacryl S-200 HR (GE Healthcare) column equilibrated with 25 mM HEPES-Na (pH 7.4), 200 mM NaCl, 1 mM βME and eluted at a flow rate of 0.7 mL/min. The elution peaks had retention volumes that corresponded to the molecular weight of the complexes. Formation was further verified by SDS-PAGE. Complexes were stored at −20°C in gel filtration buffer with a final concentration of 40% glycerol.
**Expression and Purification of Histone Proteins.** Expression and purification of core histone proteins was conducted by combining two previously reported methods with minor modifications (See Supporting Information for detailed procedure).34, 35

**Synthesis and Purification of Histone H3 Substrate and Product Peptides.** A peptide corresponding to the first 21 amino acids of the N-terminal tail of histone H3 that incorporated a dimethylated lysine at residue 4 (H3K4me21-21) was synthesized and purified as previously described.30, 36 The unmodified 21-mer histone H3 product peptide (H31-21) was synthesized and purified in a similar fashion.

**Steady-State Kinetic Assays.** Steady-state kinetic assays employing continuous fluorescence or absorbance-based methods for detection of enzymatic peroxide production (Figure S1) were performed as previously described with slight modifications (See Supporting Information for details; Figures S2 & S3).30, 36

KDM1A tolerance for 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate (CHAPS) and dimethyl sulfoxide (DMSO) was assessed in order to confirm usable range of additives (Figure S4). DMSO seemingly increases initial velocity of KDM1A within 2-fold under 10% DMSO (v/v). CHAPS concentration for disruption of non-specific aggregation affects was kept at 0.01% (w/v) in inhibitor analyses. For all IC50 titrations, unless otherwise noted, substrate concentration was held at or near the apparent $K_M$ ($K_{M, app}$) in order to ensure an equal population of the free enzyme and $ES$ complex was available for inhibitor interactions.37, 38
Steady-State Kinetic Data Analysis. Steady-state kinetic data were analyzed using GraphPad Prism 6.0 (GraphPad Software, San Diego, CA) or Grafit 4.0 (Erithacus Software, London, UK). Briefly, the no enzyme control (n=3) was subtracted from all data sets. Data was then forced through the origin by subtraction of the initial time point and responses converted to concentration units of H$_2$O$_2$ (µM). Initial velocities were calculated via linear regression and responses were limited to within 10% total product conversion. Initial velocities were then plotted versus substrate concentrations and fit to the Henri-Michaelis-Menten equation (eq 1) utilizing non-linear regression analysis:

\[
\nu_0 = \frac{V_{max}[S]}{K_m + [S]}
\]

where \( \nu_0 \) is the experimental initial velocity, \([S]\) is the substrate concentration, \( V_{max} \) is the maximal velocity at the utilized enzyme concentration, and \( K_m \) is the substrate concentration at half maximal velocity.

Dose-response curves were prepared by comparing initial rate data at specified inhibitor concentrations to a no inhibitor control and fit directly to eq 2:

\[
\frac{\nu_i}{\nu_0} = \frac{1}{1 + ([I]/IC_{50})^h}
\]

where \( \nu_0 \) is the initial velocity of the control, \( \nu_i \) is the inhibited initial velocity, \([I]\) is the inhibitor concentration, \( IC_{50} \) is the inhibitor concentration at which the rate of
demethylation is half that of the control, and \( h \) is the Hill coefficient of the curve (slope factor).

The \( K_i \) of reversible, competitive inhibitors was determined by a titration of inhibitor versus substrate followed by global fitting of the data to eq 3:

\[
V_0 = \frac{V_{max} [S]}{K_m (1 + \frac{[I]}{K_i}) + [S]} \tag{3}
\]

where \( K_i \) is the equilibrium inhibitory constant.

As a secondary determination of \( K_i \), we used the Cheng-Prusoff relationship for competitive inhibition:

\[
IC_{50} = K_i \left( 1 + \frac{[S]}{K_m} \right) \tag{4}
\]

where \( IC_{50} \) is the inhibitor concentration at which the rate of demethylation is half that of the control as defined above in eq 2.

In order to define the apparent \( K_i \) (\( K_i^{app} \)) value of inhibitors that fall in the tight-binding regime (i.e. \( \leq 10[E]_T \)), dose-response data was fit to the Morrison’s quadratic equation for tight-binding inhibition:

\[
\frac{v_i}{v_0} = 1 - \frac{([E]_T + [I]_T + K_i^{app}) - \sqrt{([E]_T + [I]_T + K_i^{app})^2 - 4[E]_T[I]_T}}{2[E]_T} \tag{5}
\]
where \([I]_T\) is the total inhibitor concentration, \(K_{i}^{\text{app}}\) is the apparent equilibrium dissociation constant for the enzyme-inhibitor complex, and \([E]_T\) is the total enzyme concentration.

As a secondary determination of \(K_{i}^{\text{app}}\), data was transformed for analysis via a Henderson plot and fit to the following linear expression:

\[
\frac{[I]}{1 - \frac{v_i}{v_0}} = K_{i}^{\text{app}} \left( \frac{v_0}{v_i} \right) + [E]
\]  

(6)

\(IC_{50}\) dependence on enzyme concentration for confirmation of tight-binding nature of inhibitor and as an additional means to determine \(K_{i}^{\text{app}}\) was fit to the following equation:

\[
IC_{50} = K_{i}^{\text{app}} + m[E]_T
\]  

(7)

where \(m\) is the linear slope.

In order to determine the equilibrium inhibitory constant \((K_i)\) of tight-binding competitive inhibitors, data was fit to following equation:

\[
IC_{50} = K_i \left( 1 + \frac{[S]}{K_m} \right) + \frac{1}{2}[E]_T
\]  

(8)

**Determination of Dissociation Rates of H3 from KDM1A.** Rates of dissociation for the H3/KDM1A binary complex were determined by monitoring the recovery of activity following a 1 h preincubation that included 50× \(K_{i}^{\text{app}}\) of histone H3.
(3.0 μM) and 100× [E] (2.5 μM KDM1A) to ensure saturation of enzyme \(^{43}\) (Note: H3\(^{1-21}\) and 2-PCPA were incubated at 10× \(IC_{50}\) since they do not display tight-binding characteristics; 50 μM and 200 μM, respectively). Preincubated samples were diluted 100-fold into assay buffer containing 5× \(K_M\) of H3K4me2\(^{1-21}\) peptide substrate (25 μM), 50 μM Amplex Red, and 1 U/mL HRP. Product was monitored continuously using absorbance maximum at 572 nm for resorufin. Progress curve data was fit to eq 9 to generate \(k_{obs}\):

\[
y = v_s t + \frac{v_i - v_s}{k_{obs}} (1 - e^{-k_{obs}t}) + \text{background} \tag{9}
\]

where \(v_i\) and \(v_s\) represent the initial and steady-state velocities respectively, \(k_{obs}\) is the apparent first-order rate constant for the transition from \(v_i\) to \(v_s\) and \(t\) is time. Under the experimental conditions used, \(k_{obs}\) approximates the dissociation rate constant (\(k_{off}\)) of the enzyme-inhibitor complex. The enzyme-inhibitor dissociation half-life (\(t_{1/2}\)) was calculated using the formula \(t_{1/2} = 0.693/k_{off}\) and residence time (\(\tau\)) was calculated using the equation \(\tau = 1/k_{off}\).

**Surface Plasmon Resonance (SPR) Measurements.** All SPR measurements were made using a BIAcore 3000 instrument, and data analyses were performed using BIAevaluation version 4.1 (BIAcore). An anti-histone H3 antibody (Abcam, ab1791) was immobilized [•3000 response units (RU)] on a research grade BIAcore CM5 chip using EDC-sNHS coupling chemistry with reagents obtained from BIAcore according to manufacturer’s protocol. In a parallel flow cell, an anti-lysozyme antibody (Abcam,
ab2408) was immobilized [•2800 response RU]. During the screening experiments with
degassed 50 mM sodium phosphate (pH 7.4) and 1 mM DTT as a running buffer, H3 and
lysozyme were injected over immobilized antibodies in respective flow cells for 2 min at
a flow rate of 30 µL/min and washed with running buffer until sensograms equilibrated.
KDM1A was then injected for 2 min at a flow rate of 30 µL/min to
monitor the binding interaction. The surfaces were regenerated via injection of 10 µL of
glycine (pH 2.0) at a flow rate of 50 µL/min. The response from the KDM1A-lysozyme-
anti-lysozyme antibody surface was used to subtract out the background (nonspecific)
signal. The $K_d$ value of the H3/KDM1A interaction was measured via injection of
KDM1A for 2 min at various concentrations ranging from 5.0 to 50.0 nM. Global curve
fitting to the 1:1 Langmuir binding model was used to determine the association ($k_{on}$) and
dissociation ($k_{off}$) rates and the apparent equilibrium dissociation constant ($K_d$).

RESULTS

CoREST (CoREST1/RCOR1) Does Not Alter the Catalytic Efficiency of
KDM1A Toward Minimal Peptide Substrates. KDM1A requires a minimal substrate
corresponding to the first 21 residues of the N-terminal histone H3 tail for efficient
demethylation activity.\textsuperscript{22} Although the enzyme itself is sufficient for demethylation of
peptide and histone substrates; activity toward nucleosomes \textit{in vitro} is regulated by its
interaction with CoREST, the minimal portion of which contains the linker and SANT2
domain or possibly the SANT1 domain.\textsuperscript{15, 16, 18} As such, we wanted to evaluate KDM1A
activity toward a peptide substrate in the presence and absence of various CoREST
constructs to determine the degree to which the partner influenced its catalytic activity and/or affinity for its substrates. Kinetic parameters and representative data for KDM1A activity in the presence of equimolar CoREST are listed in Table 1. Of the complexes tested, KDM1A/CoREST-Linker (residues 293-380) and KDM1A/CoREST-C (residues 286-452), showed only a very modest 1.5-fold increase in the initial velocity as compared to KDM1A. Despite this enhanced velocity, the complexes maintained nearly identical catalytic efficiencies because of a proportional rise in the apparent $K_M$. We should note that our derived kinetic values are in reasonable agreement with previously reported values with and without CoREST.$^{13, 22, 23, 30}$

**H3$^{1-21}$ is a Competitive Inhibitor of KDM1A Demethylation Activity.** In order to provide a basis for comparison for full-length products, we first reconfirmed the inhibition modality and potency of the peptide products of the enzymatic reaction (Figure S5). The unmodified peptide product representing the first 21 residues of histone H3 (H3$^{1-21}$) has previously been reported to be a competitive inhibitor of KDM1A activity with a $K_i$ of $1.8 \pm 0.6 \mu$M.$^{22}$ In order to eliminate any potential deleterious non-specific aggregation effects, we utilized 0.01% (w/v) CHAPS in our assay buffers.$^{44}$ Dose-response curves of H3$^{1-21}$ showed an average $IC_{50}$ of $4.84 \pm 0.16 \mu$M for KDM1A. Potency and mechanism of inhibition of H3$^{1-21}$ toward KDM1A was determined through a double titration of inhibitor ([I] that gives 30% to 75% activity as determined by eq 2) versus the substrate followed by global fitting of the data. The data were fit to models of competitive, uncompetitive, and non-competitive inhibition and were best described by a competitive model, as determined by $F$-test analysis. The double titration revealed a $K_i$ of
1.77 ± 0.08 µM. Subsequent reciprocal transformation and Lineweaver-Burk plot analysis of the data also fit the pattern to competitive inhibition, resulting in a series of nested lines intersecting at a single y-intercept. Additionally, a secondary plot of apparent $K_M$ versus $[I]$ also revealed a $K_i$ of 1.84 ± 0.14 µM, while the Cheng-Prusoff relationship for competitive inhibition yielded a value of 2.07 ± 0.07 µM. In our hands, the competitive nature of the product peptide and potency is nearly identical to that previously reported.

The H3/KDM1A Interaction Reaches Rapid Equilibrium. With the relative affinity and competitive nature of the unmodified peptide product reassessed, we looked to investigate the full-length H3 product (H3$_{1-135}$). As an initial test of potency, we assessed the dose-response of H3 inhibition of KDM1A-mediated demethylation of a peptide substrate. The initial rates as a function of H3 do not appear to be curvilinear in nature and are best fit by linear regression (Figure 2A). This suggests that the degree of inhibition at a fixed concentration of H3 does not vary over time and the interaction is not slow binding in nature. However, in some cases the establishment of the enzyme-inhibitor equilibrium is much slower than the course of the assay.

In order to assess this potential explanation for the increase in observed binding affinity, we preincubated KDM1A with H3 for specified times and then initiated the assay. We observed no significant change in the $IC_{50}$ as a function of time even with long preincubation times of up to an hour. Analysis of the dose-response curves resulted in an average $IC_{50}$ of 153.3 ± 26.0 nM ($[E]_T = 175$ nM) (Figure 2B), a value within the theoretical tight-binding limit of the assay. As there was no change in the average $IC_{50}$ as
a function of time, and the initial velocities as a function of $[I]$ appear linear, we suggest that H3 and KDM1A reach rapid equilibrium under the assay parameters utilized. However, one cannot rule out that the time-dependent transition occurs at a rate that is faster than the detection limits of the assay used.

We should note that as controls we tested the activity of HRP in the presence of increasing concentrations of full-length histone H3. Hydrogen peroxide (2.5 µM) and HRP (1 U/mL) were set at fixed concentrations and titrated against H3. In this assay, HRP retained all of its activity even at the highest concentrations tested (Figure S6A). Additionally, H3 was tested for its ability to both quench resorufin derived fluorescence and absorbance as well as quinoneimine dye derived absorbance (Figure S6B-C). In this assay, peroxide was titrated against fixed HRP and H3 to establish standard curves of responses versus peroxide concentration. Even at the highest concentrations of H3 tested, these assays showed no statistical difference between standard curves prepared in the presence and absence of H3. These controls strongly suggest that the inhibitory activity realized in the demethylation assays were solely due to the H3/KDM1A interaction.

Full-Length Histone H3 is a Tight-Binding, Competitive Inhibitor of KDM1A. As noted above, the average IC$_{50}$ of full-length H3 toward KDM1A, appears to be within the theoretical tight-binding limit (i.e. ≤ 10-fold $[E]_T$). Therefore, in order to assess the apparent affinity of H3 toward KDM1A we first conducted a titration of H3 at a starting point of 30-fold $[E]_T$ in a 1.5-fold dilution series. Data were fit to the Morrision quadratic equation for concentration-response data for tight-binding inhibitors (eq 5) resulting in a $K_i^{app}$ of 59.1 ± 5.95 nM (Figure 3A). Data were additionally analyzed
via Henderson plot revealing a $K_{i}^{\text{app}}$ of 103.4 ± 4.47 nM that was consistent with the tight-binding nature of the H3 interaction (Figure S7).\textsuperscript{42}

One of the hallmarks of tight-binding ligands is the dependence of $IC_{50}$ on enzyme concentration (eq 7).\textsuperscript{46} Therefore, in order to further confirm the tight-binding nature of the full-length H3 we assessed the effect of enzyme concentration on $IC_{50}$. Here, enzyme at differential concentrations was titrated against histone H3 with a fixed concentration of peptide substrate held at the apparent $K_{M}$ (Figure 3B). Linear regression of the data resulted in a $K_{i}^{\text{app}}$ of 61.2 ± 5.78 nM. This analysis again confirmed the tight-binding nature of the H3/KDM1A interaction that leads to complications when assessing ligand modality.

Since the potency of H3 is near the concentration of KDM1A used in the assays; standard mechanism of inhibition through double titration of inhibitor followed by global fitting of the data is precluded. Instead, the $IC_{50}$ determined at several substrates concentrations were plotted against the $[S]/K_{M}$ ratio (0.156 to 10x $K_{M}$) (Figure 3C). The data were fit to models of competitive, uncompetitive, and non-competitive inhibition and were best fit by a competitive model as determined by $F$-test analysis. A $K_{i}$ of 18.9 ± 1.2 nM was observed. This $K_{i}$ is nearly a 100-fold increase in binding affinity towards the free enzyme as compared to the peptide product described above.

In order to determine if the binding affinity of H3\textsuperscript{1-135} was either primarily electrostatic or hydrophobic in nature, an $IC_{50}$ shift assay was completed with increasing NaCl concentrations. At higher ionic strength, the potency of H3\textsuperscript{1-135} toward KDM1A in the demethylation assay decreased (Figure 3D). Based on this data, we suggest that the interaction of H3\textsuperscript{1-135} with KDM1A is likely to be mostly electrostatic in nature. It is
interesting to note that KDM1A showed negligible activity over background at 300 mM NaCl. Therefore, as a control experiment, we repeated the titration of H3\textsuperscript{1-135} against HRP in the presence of H\textsubscript{2}O\textsubscript{2} and 300 mM NaCl and activity was unaffected (data not shown). We can therefore conclude that the binding of KDM1A to the peptide substrate H3K4me1/2\textsuperscript{1-21} is also electrostatic in nature, which was also postulated by Forneris and coworkers.\textsuperscript{22}

**Histone H3/KDM1A Binary Complex has an Observable Residence Time.** To confirm reversibility of inhibition and to determine the off-rate ($k_{\text{off}}$) of full-length H3 binding, rapid dilution experiments were conducted where enzyme and inhibitor were preincubated for 1 h. The enzyme/inhibitor complex was then diluted 100-fold into assay buffer containing the peptide substrate, coupling enzyme, and chromophore. Reaction progress curve data show that KDM1A activity is slowly recovered over time ($t_{1/2} = 9.63$ min) (Figure 4). In control experiments using the H3 product peptide (H3\textsuperscript{1-21}), the progress curve appears linear, as there is no distinguishable delay in the recovery of activity (Figure S8). Additionally, $t_{1/2}$ was unable to be defined by the data, suggesting that this interaction is rapidly reversible (Table 2). We also used 2-PCPA as a mechanism-based irreversible inhibitor control,\textsuperscript{36} in the presence of which KDM1A did not recover activity as expected (Figure 4). The jump dilution experiment confirms the long residence time and reversibility of the H3/KDM1A interaction.

**Surface Plasmon Resonance (SPR) Measurements.** In order to confirm the tight-binding interaction between KDM1A and full-length H3 independent of catalysis,
we turned to SPR measurements. Here, an antibody specific to the histone H3 C-terminus was covalently immobilized on the chip surface as the capturing antibody to which H3 was immobilized. This experimental set-up allowed subsequent analysis of the H3/KDM1A binding interaction (Figure 5A). As is shown in Figure 5B, the increase in response units over time represents the amount of KDM1A bound to the antibody-immobilized histone H3. This response is proportional to the association rate constant \( (k_{on}) \) of \( 9.26 \times 10^4 \pm 1.5 \times 10^4 \text{ M}^{-1}\text{s}^{-1} \). After the association of KDM1A and immobilized H3, additional buffer was injected over the flow cell to dissociate the bound enzyme. This analysis yielded a dissociation rate constant \( (k_{off}) \) of \( 8.35 \times 10^{-4} \pm 3.4 \times 10^{-5} \text{ s}^{-1} \). These calculated rate constants result in an equilibrium dissociation constant \( (K_d) \) of \( 9.02 \pm 2.27 \text{ nM} \) that is within approximately 2-fold of that calculated in the continuous assay. This technique independently confirmed that the interaction between KDM1A and H3 has a \( K_d \) in the low nanomolar, tight-binding range and is suggestive of a 1:1 stoichiometry. Additionally, the off-rate determined from this experiment is in good agreement with the results of the rapid dilution experiment. We also suggest that the presence of DTT in the running buffers strongly precludes the formation of disulfide bonds as a contributing factor of the tight-binding nature of the H3/KDM1A interaction.

**Additional Core Histone Proteins have no Effect on KDM1A.** Several groups have illustrated that KDM1A catalyzes the oxidative demethylation of histone H3K4me1/2 and H3K9me1/2 as well as non-histone substrates.\(^6-8, 21\) Despite a broad substrate specificity profile, the demethylase has never been reported to recognize PTMs on the additional core histone proteins. In order to confirm this observation in terms of
inhibition of demethylation activity, dose-response titrations were completed for each of
the other core histone proteins, H2A, H2B and H4, against KDM1A. Consistent with
these reports, none of the other core histone proteins inhibited KDM1A activity (Figure
S9). We can safely conclude that the demethylase specifically recognizes histone H3 and
is not negatively regulated by the other core histones. However, this analysis does not
prove that KDM1A does not contain surface recognition elements that bind these other
core histones.

DISCUSSION

Many groups have suggested that the substrates and products of KDM1A exit the
expanded yet restricted active site pocket and interact with a secondary binding site on
the enzyme surface.\textsuperscript{11, 15, 27, 28} This secondary surface element is expected to that facilitate
substrate/product binding and recognition. To assess this hypothesis, we investigated the
ability of unmodified, full-length histone proteins to inhibit KDM1A dependent
demethylation activity. We chose longer, “native-like” products in order to determine
whether or not the enzyme could potentially recognize a larger portion of these species,
thereby increasing the apparent binding affinity. Before we began this study we chose to
assess whether or not differential CoREST constructs (CoREST1/RCOR1) altered
KDM1A catalytic efficiency toward peptide substrates.

Using various truncated CoREST constructs, we observed no statistical difference
in catalytic efficiency suggesting that this binding partner doesn’t control activity toward
the minimal substrate footprint. This result supports the idea that KDM1A catalytic
activity is modular in nature, as it often exists as a member of multiprotein complexes
that dictate function.\textsuperscript{19} We should note that KDM1A catalytic efficiency and substrate specificity might be further altered if chromatin or nucleosomes are utilized as substrates.\textsuperscript{22} In fact, Tan and coworkers recently illustrated that the KDM1A/CoREST-C complex has greater than a 25-fold increase in catalytic efficiency when using site-specifically methylated nucleosomes with extranucleosomal DNA as substrates; a phenomenon driven by $K_{M}^{app}$.\textsuperscript{19} Despite this evidence, there appears to be further confounding aspects to the regulation of KDM1A activity. For example, CoREST paralogs (CoREST2/RCOR2 and CoREST3/RCOR3) were recently shown to alter KDM1A activity toward nucleosomal substrates as assessed by their DNA binding ability and also affected catalytic efficiency toward peptide substrates.\textsuperscript{17, 47, 48} Since we are utilizing the minimal peptide substrate (H3K4me2\textsuperscript{17-21}) in these studies, and inhibitor potency seems to be unaffected by the presence of CoREST \textit{in vitro},\textsuperscript{23, 49} we moved forward with our investigation using isolated KDM1A.

In line with our hypothesis, initial dose-response analysis of full-length H3 versus KDM1A demethylation activity suggested that the interaction was in the theoretical tight-binding limit of the assay (i.e $IC_{50} \leq 10$-fold $[E]_{T}$). If we consider the basic definition of the equilibrium dissociation constant for competitive inhibitors, where $K_i = k_{off}/k_{on}$, it is easy to see that affinity is driven by either rapid association of the binary $EI$ complex or a slow rate of release of the ligand from the target enzyme.\textsuperscript{50} In order to assess the onset of inhibition we examined the initial rate data upon assay initiation. These initial rates appeared linear suggesting that H3 and KDM1A reach rapid equilibrium under the specified assay conditions. Additional support for this observation came from preincubation of H3 with KDM1A. Even with extended incubation times of up to 1 h the
average IC$_{50}$ did not change significantly. In order to next assign the modality of full-length H3, we next determined the IC$_{50}$ of H3 at a wide range of substrate concentrations.$^{50}$

Our data support that the binding of full-length histone H3 to KDM1A is competitive in nature and displayed nearly a 100-fold increase in binding affinity ($K_i$ of 18.9 ± 1.2 nM) as compared to the 21-mer product peptide. This value of magnitude increase strongly suggests some form of additional product recognition with the full-length histone and time-dependence of the binding interaction. We therefore looked to investigate the source of this increase in observed binding affinity as low $K_i$ values typical of tight-binding inhibitors are driven mainly by very slow rates of complex dissociation (i.e. low values of $k_{off}$) and by relation large target residence times.$^{50}$

We note that the increase in binding affinity of the H3/KDM1A interaction may be at least be partially attributed to the long residence time, as the $t_{1/2}$ of the interaction was determined to be 9.63 min with a $k_{off}$ of $1.2 \times 10^{-3}$ s$^{-1}$. Additionally, we independently confirmed the tight-binding nature of the interaction and slow rate of dissociation of the binary complex with SPR experiments in which histone H3 was immobilized to the chip surface via interaction with a C-terminal specific antibody. Importantly, our assessment that increasing ionic strength results in an increase in relative IC$_{50}$ suggests that the interaction of full-length histone H3 with both the flavin-containing active site and secondary exo-site is electrostatic in nature. Notably, there are several regions on the surface of KDM1A that are electronegative in nature and could thereby provide the secondary binding site for histone H3.
Consistent with our proposal of a secondary binding site on the enzyme surface of KDM1A, the paralog KDM1B contains such a site that was identified through crystallographic efforts. Additionally, we envision that this secondary binding site may play a role in the “stick-and-catch” model recently proposed by Mattevi and coworkers. Here, KDM1A can probe a nucleosomal substrate using both its active site and secondary recognition element and lock the complex into a competent binding mode. On the other hand, after demethylation the secondary binding site is expected to contribute to the stability of the nucleosome/demethylase complex. Importantly, KDM1A can remain bound to genetic loci on time scales that are in line with large-scale chromatin rearrangement. Upon formation of this stable product complex, KDM1A can ‘lie in wait’ and serve as a docking element for protein, transcription factor, or repressor recruitment as first suggested by Forneris and coworkers. These proteins can then cause concomitant gene expression or repression or facilitate the dissociation of KDM1A from the product complex. The role of KDM1A as a docking module is also consistent with the observation that demethylation activity at specific estrogen responsive elements (EREs) is required for ERα recruitment and concomitant target gene expression. Our observations of an increased binding affinity for full-length products and target residence time not only have implications in the role of KDM1A as a docking element, but also the kinetic mechanism of the enzyme.

Due to the time scale of dissociation of the H3/KDM1A binary complex, we also suspect that the overall demethylation reaction in vivo and in vitro may be either partially or wholly rate-limited by product release with full-length histones or nucleosomes (Scheme 1). In this model, the rate of product release, $k_3$, is much smaller than the rate of
catalysis and product rebinding \( (k_2 \text{ and } k_{-3}, \text{ respectively}) \) resulting in an \( EP \) binary complex with an observable half-life. Well in line with this hypothesis, there was more than a 5-fold reduction in \( k_{\text{cat}}^{\text{app}} \) values observed when nucleosomal substrates were utilized.\(^ {19} \) Additionally, the observation that KDM1A completely demethylates p53 (K370) \textit{in vitro} but only removes a single methyl mark when this activity is probed in cell culture further supports our claim. Together, our and others’ results suggest that the product dissociation rate of KDM1A may be “tuned” by several factors including substrate, binding partners, or splice variations to the enzyme itself.

In summary, our kinetic study has verified the tight-binding interaction between full-length histone H3 and KDM1A suggesting the existence of secondary binding site on the demethylase surface. The contact between the H3/KDM1A binary complex likely occurs through an extensive interaction interface with broad implications for enzyme function and control. As the complex also has a measurable residence time, the presence of KDM1A on specific genetic loci, may be further investigated in order to provide insights into the order of partner protein recruitment, complex nucleation, or dissociation.

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Alam. We also thank Jennifer E. Link and members of the McCafferty laboratory for their thoughtful insight during the preparation of the manuscript.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website. A detailed description of Materials and Methods and Figures S1–S9.

REFERENCES


40. Cheng, Y., and Prusoff, W. H. (1973) Relationship between the inhibition constant ($K_I$) and the concentration of inhibitor which causes 50 per cent inhibition ($I_{50}$) of an enzymatic reaction, *Biochemical pharmacology* 22, 3099-3108.


**Tables**

**Table 1.** Representative steady-state kinetic parameters of KDM1A and differential KDM1A/CoREST complexes using an H3K4me2\(^{1-21}\) peptide substrate.

<table>
<thead>
<tr>
<th></th>
<th>(K_M^{app}) (µM)</th>
<th>(k_{cat}^{app}) (min(^{-1}))</th>
<th>(k_{cat}^{app}/K_M^{app}) (µM(^{-1}) min(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>nΔ150 KDM1A(^a)</td>
<td>4.46 ± 0.27</td>
<td>6.00 ± 0.07</td>
<td>1.35 ± 0.25</td>
</tr>
<tr>
<td>nΔ150 KDM1A/CoREST-linker(^a)</td>
<td>8.89 ± 0.65</td>
<td>9.42 ± 0.10</td>
<td>1.06 ± 0.37</td>
</tr>
<tr>
<td>nΔ150 KDM1A/CoREST-C(^a)</td>
<td>8.14 ± 0.52</td>
<td>9.54 ± 0.09</td>
<td>1.17 ± 0.30</td>
</tr>
</tbody>
</table>

\(^a\)In 50 mM HEPES-\(\text{Na}\) (pH 7.5), 125 µM 4-AAP, 1.0 mM 3,5 DCHBS, 300 nM \([E]\), & 1 U/mL HRP at 25°C in air saturated buffer; peptide titrated from 50 µM to 0 in 2-fold dilution series (n=3).
Table 2. Inhibition constants for full-length histones and histone peptides against KDM1A.

<table>
<thead>
<tr>
<th>Entry</th>
<th>IC&lt;sub&gt;50&lt;/sub&gt; (nM)</th>
<th>K&lt;sub&gt;i&lt;/sub&gt; (nM)</th>
<th>k&lt;sub&gt;off&lt;/sub&gt; (s&lt;sup&gt;-1&lt;/sup&gt;)</th>
<th>residence time (τ) (min)</th>
<th>half-life (t&lt;sub&gt;1/2&lt;/sub&gt;) (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H3&lt;sup&gt;1-21&lt;/sup&gt;</td>
<td>4840</td>
<td>1770</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>H3</td>
<td>153.3</td>
<td>18.9</td>
<td>1.2 x 10&lt;sup&gt;-3&lt;/sup&gt;</td>
<td>13.9</td>
<td>9.63</td>
</tr>
<tr>
<td>H4</td>
<td>N/A</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>H2A</td>
<td>N/A</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>H2B</td>
<td>N/A</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

N/A = not applicable; entry did not effectively inhibit KDM1A. IC<sub>50</sub> values were determined at 5.0 µM (~K<sub>app</sub>) and 175 nM KDM1A. ND = not determined, inhibition was rapidly reversible. Residence time was calculated with the formula; τ = 1/k<sub>off</sub>, assuming that the dilution is large enough that the rate of compound rebounding is assumed insignificant. Half-life was calculated with the formula; t<sub>1/2</sub> = 0.693/k<sub>off</sub>.
SCHEMES

Scheme 1. Overview of simplified model of KDM1A substrate turnover and product release. Here $k_3 \ll k_{-3}$ and $k_2$ and the $EP$ binary complex would therefore have an observable half-life. This model suggests that the rate-limiting step of catalysis may in some cases be the rate of product release.
FIGURE LEGENDS

**Figure 1.** Domain and structural overview of KDM1A and CoREST. (A) Domain maps of KDM1A and CoREST1. KDM1A: SWIRM domain is shown in green, AOD is shown in blue, and tower domain is shown in magenta. CoREST: Linker is shown in orange and SANT2 is shown in cyan. The ELM1 and SANT1 domains are shown in gray due to the lack structural information. (B) Structural overview of KDM1A in complex with CoREST-C (residues 286-482). Coloring scheme follows that of the domain maps above, FAD is denoted in yellow, and N and C-termini are labeled (PDB 2IW5).

**Figure 2.** Investigation of time-dependence of inhibition of full-length histone H3 against KDM1A as monitored in fluorescence-based HRP coupled assay. (A) Linear fit of the initial velocity of KDM1A in the presence of increasing concentrations of histone H3 with no preincubation prior to assay initiation and H3K4me2\textsuperscript{1-21} substrate held at apparent $K_M$. (B) Investigation of $IC_{50}$ values with preincubation of KDM1A in the presence of full-length H3 prior to assay initiation into substrate solution show no apparent time-dependence with an average of $153.3 \pm 26.0$ nM ($[E]_T = 175$ nM).

**Figure 3.** Inhibition profiling of full-length histone H3 against KDM1A as monitored in fluorescence-based HRP coupled assay reveals the tight-binding nature of the interaction. (A) Concentration-response plot of H3 (1.5-fold dilution series from 30-fold $[E]_T$) against KDM1A fit to Morrison’s equation revealing a $K_{i}^{\text{app}}$ of $59.1 \pm 5.95$ nM. (B) Plot of $IC_{50}$ as a function of $[E]_T$ measured over a range of concentrations from 88.9 to 400 nM.
KDM1A generating a $K_i^{\text{app}}$ of 61.2 ± 5.78 nM. (C) Plot of $IC_{50}$ as a function of $[S]/K_M$ over a range of concentrations from 0.156 to 10.0 x $K_M$ is best fit to a competitive model generating a $K_i$ of 18.9 ± 1.2 nM. (D) Plot of $IC_{50}$ as a function of NaCl concentration from 0 to 150 mM suggests that the H3/KDM1A interaction is ionic in nature.

**Figure 4.** The H3/KDM1A binary complex has an observable residence time. 2.5 µM of KDM1A and 3.0 µM H3 (50× $K_i^{\text{app}}$) or 200 µM 2-PCPA (10× $IC_{50}$) were incubated for 1 h followed by 100-fold dilution into assay buffered containing 25 µM peptide substrate, 50 µM Amplex Red, 1 U/mL HRP. The recovery of enzyme activity was monitored over time by monitoring resorufin-derived absorbance at 572 nm. Histone H3’s inhibition of KDM1A is slowly reversible with an off rate of $1.2 \times 10^{-3}$ s$^{-1}$ as compared to 2-PCPA in which the activity was not recovered.

**Figure 5.** Surface plasmon resonance experiments independently confirm the tight-binding nature of the H3/KDM1A interaction. (A) Diagram of the experimental set-up in which anti-H3 antibody was covalently immobilized to chip surface in order to capture histone H3. KDM1A was subsequently injected over chip surface to monitor binding. (B) Response curves obtained during and after injection of KDM1A over anti-body immobilized H3 chip surface. Global curve fitting of sensograms reveal an association rate constant ($k_{\text{on}}$) of $9.26 \times 10^4 \pm 1.5 \times 10^4$ M$^{-1}$s$^{-1}$ and dissociation rate constant ($k_{\text{off}}$) of $8.35 \times 10^4 \pm 3.4 \times 10^5$ s$^{-1}$ and were used to calculate a dissociation constant ($K_d$) of 9.02 ± 2.27 nM.
FIGURES

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Figure 2
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\[
E + S \xrightleftharpoons[k_{-1}]{k_1} ES \xrightarrow[k_2]{t_1/2} EP \xrightleftharpoons[k_{-3}]{k_3} E + P
\]
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J.M.B. and D.G.M. conceived this study. J.M.B. designed and performed experiments, analyzed data, and wrote the manuscript. J.J.G. performed experiments. K.R.M designed experiments. J.J.G and K.R.M. helped analyze the data and revised the manuscript. D.G.M. designed the research, analyzed the data, revised the manuscript, and supervised all aspects of this study.

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Notes

The authors declare no competing financial interest.