Abstract

The SET and MYND Domain (SMYD) proteins comprise a unique family of multi-domain SET histone methyltransferases that are implicated in human cancer progression. Here we report an analysis of the crystal structure of the full length human SMYD3 in a complex with an analog of the S-adenosyl methionine (SAM) methyl donor cofactor. The structure revealed an overall compact architecture in which the “split-SET” domain adopts a canonical SET domain fold and closely assembles with a Zn-binding MYND domain and a C-terminal superhelical 9 α-helical bundle similar to that observed for the mouse SMYD1 structure. Together, these structurally interlocked domains impose a highly confined binding pocket for histone substrates, suggesting a regulated mechanism for its enzymatic activity. Our mutational and biochemical analyses confirm regulatory roles of the unique structural elements both inside and outside the core SET domain and establish a previously undetected preference for trimethylation of H4K20.

Introduction

SET domain histone methyltransferases (HMTases) mediate epigenetic post-translational histone modifications that govern transcriptional activity, in part by modulating chromatin structure and accessibility of transcription factors and RNA polymerase II (pol II) to promoters [1]. Pol II promoters are typically repressed by histone H3 lysine 9 trimethylation (H3K9me3), H3 lysine 27 trimethylation (H3K27me3), and/or H4 lysine 20 di- and tri-methylation (H4K20me2, H4K20me3). In contrast, active pol II promoters are generally unmethylated [2,3] and are associated with a permissive chromatin state enriched in histone H3 and H4 acetylation and H3 lysine 4 di- and tri-methylation (H3K4me2, H3K4me3) [4]. Human cancer leads to both global and gene-specific modifications of the “epigenome” [5,6,7,8,9]. Tumorigenesis is often accompanied by a general loss of repressive marks from bulk chromatin, which leads to disruption of heterochromatin structure and transcriptional repression. Alternatively, promoter-associated CpG islands can become heavily methylated during oncogenesis, resulting in local changes in chromatin structure (e.g., nucleosome repositioning) and replacement of active histone marks by repressive ones [10,11,12,13,14,15].

SMYD3 and its 4 vertebrate paralogs (Fig. 1A) derive from an ancient family of SET HMTases with orthologs present in plants, animals, fungi, and some (typically parasitic) protozoa [16]. All SMYDs have the N-terminal terminal portion of SET (N-SET), sequentially, and includes critical catalytic folds. The SMYD structure “core” ends in a cysteine-rich zinc binding fold (post-SET). The N-terminal terminal portion of SET (N-SET) is followed by a Myeloid translocation protein 8, Nervy, and DEAF-1 (MYND) domain, N-terminal to an intermediate or linker sequence (I-SET) of variable length and configuration [17,18]. The remainder of the SET domain (C-SET) comes next, sequentially, and includes critical catalytic folds. The SMYD SET “core” ends in a cysteine-rich zinc binding fold (post-SET). SMYDs 1–4 have an additional, previously uncharacterized 150 residue C-terminal domain (CTD), whereas SMYD3 has primarily insertions in its MYND and I-SET sequences. Most prototypic SET active site residues are conserved in SMYDs [19,20], but there are notable exceptions (discussed further below). SMYD1 and SMYD3 were identified as H3K4me3-specific HMTases [5,21], whereas SMYD2 catalyzes H3K36me2 [19]. The only previous characterization of SMYD3 HMTase was performed by...
Silva et al. [22], who reported that substrate release is facilitated by tumor-specific proteolysis of the SMYD3 N-terminal 34 residues. Aside from this, little has been done to establish the functional interface of SMYD3 with its substrates or its structural underpinnings.

Conversely, numerous studies have strongly implemented SMYD3 as a protooncogene in hepatocellular, colon and breast carcinoma, based on its high levels of endogenous expression, cancer-associated promoter polymorphisms, and cell proliferative effects produced by enforced SMYD3 over-expression in normal cells or SMYD silencing in tumors [5,22,23,24,25]. Approximately 80 genes have been identified as targets of SMYD3 HMTase, including Nkx2.8, a homeobox transcriptional regulator upregulated in hepatocellular malignancies as well as cell cycle mediators, oncogenes, and developmental fate determinants [5,22,23,24,25]. The considerable if not unprecedented interest in SMYD structure and its implications for putative anti-cancer drug development is evidenced by publication of three structures which appeared just prior to [26,27] and during [28] the submission phase of this manuscript (addressed in sections below). We present here, in addition to the independent high resolution co-crystal structure of the full length human SMYD3 with the S-adenosyl methionine (SAM) analog Sinefungin, a detailed mutational and biochemical assessment of SMYD3 function. We provide a structural basis for the proposed [29,30] differential regulation of SMYD HMTase activities via their MYND domain binding partners. We demonstrate that SMYD3 can function as a transcriptional repressor via MYND interactions as well as through hitherto undetected H4K20 HMTase activity. We show that in addition to the MYND domain, the aromatic cage structure throughout the methyltransferase active site and the unique carboxy terminal domain have the potential to regulate SMYD HMTase methylation state and substrate specificity.

**Results and Discussion**

**Preferential H4K20 activity of SMYD3**

Human his-tagged SMYD3 was purified following baculoviral or bacterial expression (Fig. S1). In addition to the expected H3K4me3 activity, SMYD3 methylated all histones to various degrees with highest activity for histone H4 when measured on mixed calf thymus histone acid extracts or on individual recombinant histones (Fig. 2). Western blotting with anti-H4 antibodies indicated that the maximal activity was for H4K20me3 (Fig. 3A), which was unanticipated given that this has generally been associated with establishment of heterochromatin. Using a series of synthetic H4 peptides bearing mono-, di-, and tri-methylation states at K20, we confirmed this specificity and also observed significant activity toward H4K20me2 (Fig. 3B). It is generally thought that the majority of H4K20 methylation occurs in a stepwise process in which monomethylation by the SET HMTase PR-SET7/SET8 serves as a substrate for di- and trimethylation by SUV420H [12,14]. That H4K20me2 served as a far better substrate than unmethylated or monomethylated species (Fig. 3B) indicated that SMYD3 alone, at least *in vitro*, is capable of progressive methylation at this lysine mark. H4K20 methylation is not a general property of SMYDs, as evidenced by the near baseline activity of SMYD1 (Fig.3A).

While it would be ideal to have a clear structural rationale for the substrate selectivity demonstrated here, crystal structures available at the time of writing do not provide enough detail to make a clear and definitive statement. Alignment of the SMYD3 (or SMYD1) structures with other structures featuring an H4 peptide fragment bound to an MTase, such as in the SET8 structure [31], shows considerable clashes between the H4 peptide and the SMYD protein. Close inspection of the overlay indicates that the H4 peptide forms part of the support for the SAM binding
Figure 2. SMYD3 preferentially catalyzes histone 4 lysine methylation in vitro. (A, B) SMYD3 purified from baculovirus methylates all histones (H4->H2A->H3->H2B) in vitro. Histone methyltransferase (HMTase) assays employed mixed histones from HeLa cells as substrate. Upper panel, fluography showing H- incorporation into H3 (17 kD) and into species smaller bands (H2A/ H2B and H4). Lower panel, Coomassie-stained SDS-PAGE gel used to verify equal loading. (C) SMYD3 purified from bacteria methylates histones H4->H3->H2A in vitro. Recombinant histones or mixed histones were used, as indicated, for substrate. Fluorography is shown and the bands corresponding to each histone are indicated. doi:10.1371/journal.pone.0022290.g002

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Conventional SET and novel features of the SMYD3-Sinefungin complex

Baculoviral SMYD3 was co-crystallized with the SAM analog Sinefungin, and the structure was solved to 1.8 Å resolution (Fig. 1B; Table 1) [37]. SMYD3-Sinefungin crystallized as 2 symmetry-related molecules/unit cell (P21). However, no convincing dimer interface exists, and the mass of the purified protein following gel filtration was 50,187d (Fig. S1A), consistent with a monomeric form. Thus, we confined our analysis to the monomer of Fig. 1B.

The domains comprising the “core” of the split-SET (N-SET, C-SET, post-SET) of SMYD3 and the MYND domains (Fig. 1B) overlay with those of corresponding conventional domains (Figs. S2A-D) [17,18,38,39,40,41,42,43,44]. Modification of the strictly conserved and catalytically essential Y239 results in the expected loss of function (Fig. 4A). Mutation of several residues conserved within many conventional N-SEts (e.g., G15, G17) and C-SEts [17,44] (e.g., C186, E192 and H206) abrogated SMYD3 HMTase activity (Fig. 4B), confirming the functional conservation of the split SET domain. About one third of the SMYD3 substrate binding site is formed by the Intermediate SET spacer (I-SET) region located C-terminal to MYND (Fig. 1). The significance of this variable linker region in SET substrate selectivity has already been noted [39,41,44,45]. However, the SMYD3 I-SET is unusually long and exhibits extraordinary structural conservation, in lieu of primary sequence similarity, with the I-SET of the Rubisco Large Subunit Methyltransferase (RLSMT) (Fig. S2E).

The close structural similarity to other SET domains allowed us to superimpose onto SMYD3 the H3K4 peptide coordinates from the SET7/9 ternary complex (Fig. 5A) [41]. The peptide is bound in the conventional manner; i.e., the methyl-lysine is oriented on the opposite surface of SMYD3 from the SAM/Sinefungin methyl donor, with a narrow channel connecting the two surfaces of the SET domain. The orientation is similar to that modeled in mouse (m)SMYD1 [26] (Fig. S5B), with selectivity opportunities on either flank of the target lysine. The relatively conservative mutation T184A, which contacts the N-terminal side of the peptide, confers not only increased activity toward H4, but a striking gain of activity toward H3. The C-terminal of the modeled peptide clashes with the CTD, suggesting that the CTD also regulates the specificity of substrate binding (more below).

The SMYD3 aromatic cage

The SMYD3 post-SET provides another commonly shared feature—an essential aromatic residue, Y257 (see Fig. 4B), that anchors against the conserved SET core to form the hydrophobic channel interface with substrate (Fig. 5B). A notable difference in SMYDs is that a critical SAM-contacting tyrosine, which occupies
this position in other N-SETs (e.g., Y335 in SET9) [17], is not conserved. To apparently compensate for the lost N-SET contact, SMYD3 and its paralogs contain a structurally homologous and essential (Fig. 4B) aromatic contact (F183 in SMYD3) within the β-1 of the C-SET (Figs. 1B and 5). The bracketing pi-cation interactions of the aromatic cage [46] are likely essential for efficient MTase

Figure 3. SMYD3 trimethylates H4-K20 preferentially. (A) SMYD3 trimethylates H4-K20. Right panel: unmethylated [H4(0)], mono-[H4(1)], di-[H4(2)], and, as a negative control, tri-methylated [H4(3)] peptides were employed in an in vitro HMTase proximity bead assay with baculoviral SMYD3 and SMYD1 (negative control). Degree of methylation was measured by scintillation counting in CPM. Left panel: Western analysis using anti-mono- and trimethyl-specific antibodies (Upstate) confirm in vitro specificity of SMYD3 for H4-K20me3. (B) SMYD3 preferentially trimethylates H4-K20 in reconstituted chromatin. Recombinant oocyte nucleosomes were assembled into chromatin, followed by in vitro HMTase assays and SDS-PAGE resolution of reaction products. SMYD3 inputs were increased from 0.5 μg to 2.4 μg, (triangle above lanes), and western analyses were performed with the indicated histone H4 methylation state-specific antibodies (middle panels), with a pan-anti-H4 (lower panel) providing a loading control for chromatin input.

doi:10.1371/journal.pone.0022290.g003
activity. Extended pi cloud interactions between aromatic side-chains extending from the aromatic cage appear common in MTases. For example, residue F259 interacts with the adenine ring of Sinefungin and the Y239 of the aromatic cage, which itself packs against Y257. F216 packs against Y198 which packs against the F183 of the aromatic cage. A similar network may be seen in the SMYD1 structure, with preservation of the aromatic network around Y252, the equivalent of Y239 in SMYD3.

Table 1. Crystallization, data collection, and refinement statistics for the SMYD3/Sinefungin structure.

| Space group | P2₁ |
|-------------|-----|
| Unit Cell   | a = 58.175Å, b = 118.073Å, c = 82.901Å, α = γ = 90°, β = 91.579° |
| Wavelength (Å) | 1.2815 |
| Resolution (Å) | 30.0–1.85 |
| Redundancy | 7.3 |
| Unique Reflections | 94,957 |
| Completeness (%) | 99.8 (99.9) |
| Molecules/Assym. Unit | 2 |
| R/Rwork/ Rfree (%) | 20.1/20.0/21.6 |

*Numbers in parentheses refer to the highest resolution shell.

R = 2|Fo–Fc|/|Fo|, where Fo and Fc are the observed and calculated structure factors, respectively. Rwork is calculated using the formula for R, but employing only the 95% of reflections used in the refinement and Rfree is calculated using a randomly-selected 5% subset of reflections not used in the refinement.

doi:10.1371/journal.pone.0022290.t001

Figure 4. Mutational analysis of residues critical to SMYD3 structure and function. (A) Wild-type SMYD3, but not catalytic point mutant Y239F, methylates recombinant H3 and H4 in an in vitro HMTase assay. Upper panels: Fluorographs with bands corresponding to H3 (left) and H4 (right) indicated. Lower panels: Coomassie-stained PVDF membranes used to verify equal loading. (B) Substitution and (C) truncation mutants, constructed in E coli as described in Methods, were compared in in vitro HMTase assays to wildtype SMYD3 and to SET7/9. Inputs (upper panels, ~500 ng) were assayed for ³H-SAM incorporation (middle panels) either on recombinant histone 4 or mixed histones, as indicated (lower panels). Alanine substitutions of most SMYD3 residues predicted to be catalytically essential eliminate HMTase activities. An exception is T184/A which, as described in the text, appears to affect H3-H4 substrate specificity (note change in relative ratios of H3/H4). N-terminal truncation through position 44 removes the entire N-SET domain, while truncation through position 74 eliminates both the N-SET domain and half the MYND domain.

doi:10.1371/journal.pone.0022290.g004
Despite the sequence identity and near identical backbone placement of the aromatic residues around F182 (the equivalent to F183 in SMYD3), the SMYD1 network assumes a very different set of side chain conformations, driven by the insertion of the adjacent leucine side-chain into the arrangement observed for SMYD3 (Fig. 5B). In fact, F182 in SMYD1 is rotated away from the catalytically competent conformation [26]. As suggested by this geometry, SMYD1 should be, and is, a less efficient MTase than SMYD3 in catalyzing higher methylation states of the common H3 substrate (Fig. 6). Fig S3 shows the similarities of the aromatic network among other lysine MTases, with the extent of the aromatic networking trending with the amount of methylation preferred. In general, stabilization of the biologically active conformation of the aromatic residues forming the cage surrounding the target lysine of the substrate should lead to more efficient transfer rates and therefore, indirectly, to the MTase’s proclivity toward mono-, di-, or tri-methylation.

An intact MYND domain is required for catalysis and transcriptional specificity

The MYND domain is the principal distinguishing element separating the SMYDs from other SET domain-containing proteins. MYND consists of two interlocking zinc binding folds and is present in several transcriptional regulators where it facilitates interactions with partner proteins through PXLXP motifs [47,48,49]. Though unfettered by SET domain constraints, the integrity of MYND is essential to SMYD3 basal function, as substitution of its Zn2+ ligating residues (C49 or C87) eliminated HMTase activity (Fig. 4B). This observation is consistent with previous analyses of the AML1/ETO MYND domain which indicated that coordination of zinc atoms is essential to maintain the intact conformation of that MYND domain, with loss of zinc coordination leading to a disordered domain [49]. Loss of coordination here also likely leads to a disordered domain, but more importantly, the lack of order affects the catalytic fidelity of SMYD3, indicating that some constraints on the linking sequence between the N- and C-SET domains exist.

The intact MYNDs of AML-1/ETO and SMYD3 bind a common PXLXP-containing protein, the N-CoR transcriptional co-repressor (Fig. 7A) [50]. That N-CoR can bind to SMYD3 and ETO similarly is consistent with our finding that SMYD3 can act as a MYND-dependent transcriptional repressor (Fig. 7B,C). These data confirm that the nature of the MYND-bound ligand influences SMYD3 transcriptional outcomes [49].

Potential contribution of SMYD3 and SMYD1 CTDs to catalysis

SMYD1-4 have an additional ~150 residue C-terminal domain (CTD) whose function was recently proposed [26] to regulate MTase activity of SMYDs. The SMYD3 CTD is a superhelical 9 α-helical bundle which constrains the floor of the substrate binding site opposite to the I-SET domain, preventing the trivial insertion of substrates (Fig. 8). In fact, the CTD clamps further down on the peptide binding space of SMYD3 than of SMYD1, featuring a greater superhelical pitch, such that it contacts the MYND domain (circled region of Fig. 8). The difference in pitch is likely driven by the larger turn in the C-SET domain of SMYD1 which significantly displaces the entire CTD relative to its location in SMYD3. There is still a relatively large space near the C-terminus of the modeled peptide where the inner wall of the pocket is decorated by polar residues from the CTD (mainly helix 4). We suggest that these polar residues would cooperate with the post-SET residues to select for specific sequences N- and C-terminal to the methyl-lysine, even in the absence of a significant displacement. In this context, the CTD could function as a cap necessary to bind substrates effectively and selectively.
Complexities of SMYD3 substrate entry/release

Constrictions imposed by the I-SET, post-SET and CTD domains onto the peptide C-terminus suggest that substrate release is a complicated process for SMYD3. Silva et al. [22] reported that substrate release is facilitated by tumor-specific proteolysis of the SMYD3 N-terminal 34 residues; that is, the N-SET is "auto-inhibitory" to catalysis. To the contrary and consistent with our structure, we found that elimination of the N-SET by truncation at position 44 or 74 or by destabilizing its conserved first β-turn, eliminated HMTase activity (Fig. 4B, C). We suggest, instead, that ...
substrate release will require a significant conformational change in the CTD, which should be readily detected by differential shifts in the geometry/contacts of unmethylated and methylated peptides.

A crystal structure of the human SMYD3-sinefungin complex was reported by Sirinupong et al. [27] during the final drafting of this manuscript. Despite space group/crystal packing differences (please compare Table 1 of both manuscripts), the two HMTase-substrate inhibitor complexes could be virtually superimposed. Indeed, a number of active site and MYND domain residues, predicted in that paper as important for basal catalysis or PXLXP-binding interactions, were confirmed by our mutational (Fig. 4) and biochemical (Fig. 7) analyses. Based on the differential geometries adopted by the CTDs of SMYD1 and SMYD3 (code supra), Sirinupong et al. [26,27] speculated that the CTD must undergo a hinge-like movement to relieve its inherent auto-inhibition of substrate entry and/or release. However, neither of the structural analyses rules out the possibility that, at least for basal catalysis, the CTD performs a positive enzymatic function by stabilizing the active site. As shown previously [5], SMYD3 HMTase is stimulated by HSP90, a chaperone whose deregulation is also strongly implicated in a broad array of malignancies [51,52]. It will be critical to determine if HSP90 binds directly to SMYD3, and if so, whether this interaction generates a CTD conformational change of the nature they proposed.

Another structural analysis of SMYD3 was published by Xu et al. [28] during the review process. Notwithstanding their considerably lower resolution (none better than 2.8A), their structure overlays very closely with ours. Much as in the work of Sirinupong et al. [26,27], Xu et al. [28] speculate on the previously observed [5,21] association of SMYD3 with HSP90. While they do not establish a causal link, they do help establish some of the residues necessary for basal activity against an uncharacterized admixture of histones. The two residues lowering activity (D241 and D392) have a structural role, making apparently key intramolecular hydrogen bonds, while the one that does not make any intramolecular hydrogen bonds (E192) fails to alter basal activity. Interestingly, E192 is proximal to T184 in space, suggesting the trajectory of the N-termini of histones lie less towards the CTD and more towards the MYND domain, which may explain why an intact MYND domain is essential for activity. Given that Xu et al. [28] find weak but dose dependent SMYD3 HMTase activation with DNA binding to the MYND domain, one might speculate that the influence of MYND domain conformation changes may lie not only with its interactions with the C-SET residues adjacent to the catalytic binding site but also with the histone on the exterior surface.

Conclusions

SMYD MTases share many key features in their SAM binding and lysine side-chain binding sites. A key beta-turn motif in the N-SET is essential for activity, with deletion of the motif or mutation of the superfAMILY signature residues G15 and G17 leading to a complete loss of activity. This motif serves as a flap that partially encloses the active site and provides residues that can interact with SAM. Targeting the disruption of this loop therefore becomes a logical objective for oncology research, as it should be sufficient to eliminate SMYD3 activity. The residues which comprise the motif are typically quite diverse and only modestly conserved, suggesting that selectivity may be achieved as well. The main drawback to targeting the loop is that the current motif features a relatively shallow groove and inhibitors would have to induce a conformational change that cannot be visualized from the current structures. Nevertheless, simulation methods could be used to explore this region of the protein.

A more likely approach to targeting SMYD3 activity is to design inhibitors that bind either the SAM- or substrate-binding pockets. Our examination of the active site suggests that disruption of the aromatic cage structure is likely to succeed, even if the site of catalysis is not occupied by an inhibitor. Differences in intramolecular aromatic-aromatic contacts lead to different stabilizations of the catalytically competent protein conformation. These differences in stability likely influence the MTase activity and hence the preference for the extent of methylation conferred on their substrates. The difference in MTase activity between SMYD1 and SMYD3 highlights this disparity; even though the sequences are identical, subtle changes in the packing influence the aromatic cages, with the more active SMYD3 retaining a stronger aromatic network than the less active SMYD1.

The MYND domain inserts into an otherwise structurally conserved SET motif that extends back to bacteria and viruses. We established that SMYD3 function is dependent on a properly folded MYND domain, suggesting that its role is not only in attracting partner binding partners but also in influencing the conformation of the N- and C-SET domains. Consistent with this hypothesis, T184 is on the far end of a beta sheet connected to the MYND domain. We establish that the rather conservative mutation of that residue to alanine leads to increased activity and promiscuity. This result suggests that small changes in the chemistry and position of the threonine side chain can lead to significant changes in catalytic activity and preference. Such changes may be possible through propagated changes in MYND domain conformation on the substrate binding pocket or may arise from changes in its direct association with a portion of the histone. More research is needed to refine these possibilities and to clarify which other residues confer the substrate preference for H4K20.

Although the MYND domain helps provide functional selectivity toward SMYD substrates, the CTD may also regulate the level of HMTase activity, serving as a cap necessary to bind substrates effectively and selectively. More experimentation is necessary to clarify the roles played by the CTD of SMYD3. New opportunities to design potent and selective agents may arise from the further characterization of these two domains and their interrelatedness to the SET domains.

Nevertheless, how do we explain the apparent biologic paradox that the oncogenic SMYD3 catalyzes histone lysine marks that promote both localized promoter activation (H3K4me3) and, even more aggressively, the repressive stabilization of heterochromatin (H4K20me3) [10,11,13,53]? While global reduction of H4K20 trimethylation has been suggested to be a hallmark of human cancer [11,15], stable and heritable H4K20-mediated repression of selected pol II genes, including tumor suppressors, has recently become appreciated as an epigenetic feature of cancer [4,11]. For example, the tumor suppressor target of methylation-induced silencing (TMS1/ASC) becomes methylated and silenced in human breast and other cancers [54,55,56,57]. Silencing is accompanied by a local shift from a histone activating mark, H4K16 acetylation (Ac), to H4K20 trimethylation [58]. Selective promoter-proximal "pausing" results, such that initiated Pol II accumulates just downstream of the transcription start site [59]. Taken together, SMYD3 may serve both as a repressor of tumor suppressor expression and a promoter of oncogene expression. These studies illustrate the complexities of gene-specific regulatory mechanisms in the epigenetic program and underscore the critical importance of tightly regulating the targeting of SMYD3 for regional deposition of H3K4me3 and H4K20me3.
Materials and Methods

Crystallography

X-ray diffraction data are summarized in Table 1. Details of protein purification and crystallization are provided below. The data were indexed and integrated using the program MOSFLM [60] and then merged using the program SCALA [61]. The subsequent conversion of intensity data to structure factor amplitudes was carried out using the program TRUNCATE [60]. The program SnB [61] was used to determine the location of Zn sites in the protein using the Bijvoet differences in data collected at the Zn peak wavelength. The refinement of the Zn sites and the calculation of the initial set of phases were carried out using the program MLPHARE [60]. The electron density map resulting from this phase set was improved by density modification using the program DM [60]. The initial protein model was built into the resulting map using the program ARP/WARP [62] and XTALVIEW/XFIT [63]; (available on request from San Diego Super Computer Center). This model was refined using the program REFMAC [60] with interactive refitting carried out using the program XTALVIEW/XFIT [63]; (available on request from San Diego Super Computer Center).

Molecular biology

Immunoprecipitations, histone methyltransferase assays, and mutagenesis were performed as previously described [19]. Details of each of these experiments and a list of the templates and mutagenic primers employed are provided below. Dual luciferase assays using GALA-DBD-SMYD3 wildtype and GALA-DBD-SMYD3 mutants (C49G and C87G) were performed and normalized following transient transfection into 293T cells as previously described [19] and are detailed below.

Cloning and baculoviral expression

The full length human SMYD3 protein (Genbank Accession No. AAH31010; SEQ. ID NO:1) was engineered to contain a C-terminal hexa-histidine tag. Sequence verified clones were each transformed into DH10 BAC chemically competent cells (Invitrogen Corporation, Cat.#10361012). The transformation was then plated on selective media. 1-2 colonies were picked into minipreps and total DNA isolated. The bacmids were transfected and plated on selective media. 1-2 colonies were picked into minipreps and plasmid DNA isolated. The bacmids were transfected into 293T cells using the standard Bac to Bac protocol (Invitrogen Corporation, Cat. #10359-016) to generate viruses for protein expression. SF9 cells were used for 48 hr expressions in SF-900 II media.

The full length cDNA of HSP90 was cloned from Hep G2 cells [ATCC HB-8065]. The chaperone HSP90 was co-expressed with SMYD3 by co-infection with virus for each. Cells were collected into the resulting map using the program ARP/WARP [62] and XTALVIEW/XFIT [63]; (available on request from San Diego Super Computer Center).

Mutagenesis, cloning, and bacterial expression

Point mutants were generated using the GeneEditor in vitro Site-Directed Mutagenesis System (Promega) according to the instructions of the manufacturer For PCR, samples were heated to 94°C for 5 min, subjected to amplification for 16 cycles of 0.5 min at 94°C, 0.5 min at 55°C, and 0.5 min at 68°C and extended after the last cycle at 72°C for 7 min. Polyhistidine (6xHis)-tagged SMYD3 wildtype, truncation and substitution mutants were cloned into Gateway (Invitrogen) pET™-DEST42. High level expression was induced by IPTG in E. coli strains MG232 (Scarab LTM) or Hsp90Plus™ (Expression Technologies Inc). Primers and mutagenic oligos were:

Substitution mutants

G1519A

cgcgaacag ggg aaa ggg ctggegggcctg forward + strand
cgcgaacag ggg aaa ggg ctggegggcctg reverse + strand
CGAGGC GGGCAGC GGGTTTGGCCCTGTTGGCG reverse C49S
cgtggcccctg ccagcccctg forward + strand
cgtggcccctg ccagcccctg reverse + strand
AAGGGCAGGTCGGCGACGACGCCACAG reverse C87A
cacaaggggaa tgg aaatgcttact forward + strand
cacaaggggaa tgg aaatgcttact reverse + strand
TTAAAGGGCAGTCGCTCCGCCTTGGTTCG reverse S182A
cattgctgacact gcacatcctgt forward + strand
cattgctgacact gcacatcctgt reverse + strand
ACAGATGTTGAAAGCGTATGACAGATCATC reverse T184A
tgacattgacttacct gcacatcctgt forward + strand
tgacattgacttacct gcacatcctgt reverse + strand
CGCATATCGATGGCGAAAGGGTTGCA reverse G186A
ctcttcatgggtct gctgctggct forward + strand
tctcttcatgggtct gctgctggct reverse + strand
CATCTCCCGATCGAGGATGTTGGAAGA reverse E192A
gggtgatctcggaa gccggtgacgc forward + strand
gggtgatctcggaa gccggtgacgc reverse + strand
GCCAAGCCAACTGCTCGCATCTCCCGC reverse H206A
cattgctaatc tcacgttgcttct forward + strand
cattgctaatc tcacgttgcttct reverse + strand
GGGGTACAAGCCTGGCATGGAGCAAGA reverse Y257A
cattgctaatc tcacgttgcttct forward + strand
cattgctaatc tcacgttgcttct reverse + strand
CATCTCCCGATCGAGGATGTTGGAAGA reverse E192A
gggtgatctcggaa gccggtgacgc forward + strand
gggtgatctcggaa gccggtgacgc reverse + strand
GCCAAGCCAACTGCGCATCTCCCGC reverse H206A
cattgctaatc tcacgttgcttct forward + strand
cattgctaatc tcacgttgcttct reverse + strand
GGGGTACAAGCCTGGCATGGAGCAAGA reverse Y257A
cattgctaatc tcacgttgcttct forward + strand
cattgctaatc tcacgttgcttct reverse + strand

Protein purification

Frozen cells were lysed in buffer [50 mM Tris-HCl pH 7.7, 250 mM NaCl with protease inhibitor cocktail (Roche Applied Science, Cat. #11-873-580-001)] and centrifuged to remove cell debris. The soluble fraction was purified over an IMAC column charged with nickel (GE Healthcare, NJ), and eluted under native conditions with a step gradient of 10 mM, then 500 mM imidazole. Proteins were then further purified by gel filtration using a Superdex 200 column (GE Healthcare, NJ), into 25 mM Tris-HCl pH 7.6, 150 mM NaCl, and 1 mM TCEP. Protein was pooled based on SDS-PAGE and concentrated to 1-10 mg/ml.
Crystal preparation

Diffraction quality crystals were obtained by hanging or sitting drop containing 0.73 µl of protein 10 mg/ml and 1 mM Sinefungin in 25 mM Tris-HCl pH 7.6, 150 mM NaCl, 1 mM TCEP and 0.75 µl reservoir solution: 100 mM Tris-HCl pH 8.5, 17% PEG 20 K, 100 mM Magnesium Chloride hexahydrate in a sealed container containing 500 µl reservoir solution, incubated overnight at 21°C. Crystals were also grown with a reservoir solution of 100 mM HEPES pH 7.5, 16% PEG 3350, 200 mM Magnesium Chloride.

The crystals were individually harvested from their trays and transferred to a cryoprotectant consisting of 75–90% reservoir solution plus 20–25% glycerol or PEG400. After ~2 min, crystals were collected and transferred into liquid nitrogen and then transferred to the Advanced Photon Source (Argonne National Laboratory), where a two wavelength MAD experiment was collected, using a Zn peak wavelength and a high energy remote wavelength.

Immunoprecipitation (IP) and Western blotting

293T cells were transiently transfected, harvested 48 hours later, and then lysed in RIPA buffer (150 mM NaCl, 1% NP-40, 0.5% DOC, 50 mM Tris pH 8, 0.1% SDS) containing protease inhibitors (Roche Molecular Biochemicals, Indianapolis, IN). Cell supernatants were incubated with primary anti-tag mAb or polyclonal anti-HI3 Ab (0.5–2 µg/ml) centrifuged at 4°C, and then incubated with protein A-Sepharose/protein G PLUS-agarose (Santa Cruz Biotechnology) at 4°C with rotation for 1 hour. Resulting immune complexes were washed 6 times and immunoprecipitated proteins were resolved on 8–15% SDS-PAGE. Separated proteins were transferred to nitrocellulose (Protran BA, Schleicher and Schuell, NH), blocked using 5% nonfat milk (10 g nonfat milk, 150 mM NaCl, 10 mM Tris pH 8, 0.05% Tween-20) overnight at 4°C. Membranes were incubated with 1° antibody for 1 hour at room temperature, extensively washed, then incubated with 2° antibodies for 1 hour at room temperature. Blots were exposed and developed using the ECL blot detection reagent (Amersham Pharmacia Biotech) according to the instructions of the manufacturer.

Histone methyltransferase assays

For in vitro HMTase assays, SMYD3 proteins (0.1–1 µg) /– equivalent amt. of human HSP90x (Assay Designs, Ann Arbor, MI, USA, cat. no SPP-776D) were incubated with 1 µg of mixed histones from calf thymus (Sigma) or recombinant core histones (Upstate), or with 1 µg reconstituted chromatin generated from oocyte nucleosomes (graciously provided by Dr. Yali Dou, Univ. Michigan Med School) prepared as described previously [64,65]. Recombinant oocyte histones were assembled onto a 201 bp ‘60I’ DNA template [66] by mixing ~1.5 µg octamers with 1 µg DNA template in a volume of 10 µl containing 2 M NaCl, 10 mM Tris (pH 8.0), 0.1 mM EDTA, and 10 mM β-mercaptoethanol, followed by stepwise, 10-fold reduction of salt by addition of Tris–EDTA to a final concentration of ~0.2 pmol nucleosome/µl 200 mM NaCl/Tris–EDTA. For radioactive based assays, 2 µCi S-adenosyl-L-[methyl-3H] methionine (SAM; Amershams Biosciences) was included as a methyl donor. All reactions were carried out in 40 µl HMT reaction buffer (10 mM dithiothreitol, 100 mM NaCl, 4 mM MgCl2, and 50 mM Tris-HCl at pH 8.8) at 30°C for 3 hours. An 18% SDS-PAGE gel was used to resolve the samples and fluorography was used visualize positive methylation.

Substrate loading was visualized by Coomassie blue staining.

Specificity of SMYD3 activity was determined, following transient transfection into 293T cells, by incubating immunoprecipitated proteins with recombinant histones and 20 µM unlabeled SAM (Sigma) in 40 µl HMT reaction buffer at 30°C for 1 hour. Western blot analysis was conducted using antibodies against H3K4me2, H3K4me3, H3K9me2, H3K9me3, H3K27me3, H3K36me2, H3K79me2, H4K20me2, and H4K20me3 (all from Upstate, Charlotteville, VA).

Differential H4 methylation state catalyzed by SMYD3 was confirmed by proximity bead HMT assays as follows: 2 µCi of 3H-SAM (Amersham Biosciences) were incubated with 0.1 µg of SMYD3 and 0.1 µg of histone H4peptide, non-, mono-, di-, or trimethylated at K20 (sequence of the peptide: acetyl-GGKGLKGGAKRHRKVL-biotin). The assay was carried out for three hours at 30°C in 20 µl HMT reaction buffer. At the end of the incubation time, 100 µl of binding buffer (1x PBS containing 1% NP-40 and 0.1% SDS) was added. The substrate was then precipitated using 10 µl of Streptavidin PVT SPA Scintillation Beads (Amersham Biosciences; used as 50% slurry in binding buffer) for one hour at room temperature on a rocking platform, followed by five washes in binding buffer and scintillation counting.

Transcription assays

The SV40-luciferase reporter, containing five copies of the GAL4-UAS, was obtained from J. Milbradutt [19]. pRL-TK was purchased from Promega. The GAL4-SMYD3 WT and MYND mutant mammalian expression vectors were constructed by PCR amplification (5’ ATG CGC GCC GAG GCC CGC; 3’ TCA GTG GCT CCT AAC CTC CTG) and restriction digestion (Not I; Xba I) followed by subcloning into the GAL4-DBD plasmid [20]. Dual luciferase assays were performed and normalized following transient transfection into 293T cells as previously described [19].

Supporting Information

Figure S1 Expression and purification of recombinant human SMYD3. (A) Baculoviral SMYD3. 6X-his-SMYD3 was expressed in sf9 cells as detailed in Methods, purified by Ni-NTA, Hitrap-Q, and Superdex-75 column chromatography (left) and confirmed for purity by mass spectrometry (right). SMYD3 purified as a monomer of predicted (50189) mass. These fractions were suitable for crystallization and further biochemical analyses (described in text). (B) Bacterial SMYD3. 6X-his-SMYD3 (wild-type, catalytic mutant H206/A, and other mutants analyzed in Suppl. Fig. 4) were cloned into Invitrogen Gateway plasmids as described in Methods. Following IPTG-induction in Scarab MG232 (left), proteins were purified by Ni-NTA (center) and confirmed with polyclonal anti-SMYD3 (right). (TIF)

Figure S2 Conventional SET and MYND domain architectures are unaltered in SMYD3. (A-C) Ribbon representations in cross-eye stereo of the N-SET(A), C-SET (B), and POST-SET (C) domains of SMYD3 (colored by domain as in Fig. 1) superimposed on the corresponding regions of SMYD1 (magenta), SET1 (red brown), SET9 (black), Rubisco LSMT (white), Dim5 (green), CIf4 (blue-green), and the viral SET of the Chlorella virus (gold). Zn locations are indicated by spheres. Sinefungin is represented in green wireframe. (D) Structure of the MYND domain in cross-eye stereo of SMYD3 (yellow) superimposed on the MYND domains of ZMYND10 (green), ETO (white), CBF2ATI (black), and SMYD1 (magenta). Zn locations are indicated by spheres. (E) Overlay of the complete SET domain of SMYD3 (colored by domain as in Fig. 1) with those of SMYD1 (magenta) and Rubisco LSMT (white) in cross-eye stereo. Sinefungin is represented in green wireframe. Of all MTase structures currently available, only these three almost completely
overlays, including the commonly conformationally and sequentially variable I-SET region.

**Figure S3** Aromatic residues in the catalytic site in cross-eye stereo. Comparison of the SMYD3 catalytic site (colored by domain) with the corresponding site in (A) SET8 (red box, B) SET7/9 (black), (C) Rubisco LMST (white), (D) DIM5 (green), (E) CLR4 (blue green), and (F) the SET domain from the Chlorella virus (gold). The modeled lysine from the SET7/9 structure (black carbons) and Sinelfungin (green carbons) are displayed for reference.

**References**

1. Venters B, Pugh BF (2007) Chromatin meets RNA polymerase II. Genome Biology 8: 319.
2. Bird AP (1989) CpG-rich islands and the function of DNA methylation. Nature 321: 209–213.
3. Cheng B, Price DH (2007) Properties of RNA polymerase II elongation complexes before and after the P-TEFB-mediated transition into productive elongation. J Biol Chem 282: 21901–21912.
4. McCabe MT, Brandes JC, Veronos PM (2009) Cancer DNA methylation: molecular mechanisms and clinical implications.Clin Cancer Res 15: 3927–3937.
5. Hamamoto R, Furukawa Y, Morita M, Imura Y, Silva FP, et al. (2004) SMYD3 encodes a histone methyltransferase involved in the proliferation of cancer cells. Nat Cell Biol 6: 731–740.
6. Muta IS, Figgitt M, Clements CS, Kill IR, Bridger JM (2007) Alterations to nuclear architecture and genome behavior in senescent cells. Ann N Y Acad Sci 1106: 250–263.
7. Jones PA, Baylin SB (2007) The epigenomics of cancer. Cell 128: 683–692.
8. Jones PA, Baylin SB (2007) The fundamental role of epigenetic events in cancer. Nat Rev Genet 3: 415–428.
9. Ducasse M, Brown MA (2006) Epigenetic aberrations and cancer. Mol Cancer 5: 60.
10. Beneti R, Gonzalo S, Jiao I, Schotta G, Klatt P, et al. (2007) Suv4-20 h deficiency results in telomere elongation and derepression of telomere recombination. J Cell Biol 178: 925–936.
11. Fraga MF, Ballestar E, Villar-Garea A, Boix-Chornet M, Espada, J et al. (2005) Loss of acetylation at Lys16 and trimethylation at Lys20 of histone H4 is a common hallmark of human cancer. Nat Genet 37: 391–400.
12. Pesavento JJ, Yang H, Pesavento JJ, Liu Z, O’Malley S, Harris JV, et al. (2002) m-Bop, a repressor protein essential for cardiogenesis, interacts with 4K20, a heart- and muscle-specific transcription factor. J Biol Chem 277: 26324–26329.
13. Srivastava D, Gottlieb PD, Olson EN (2002) Mechanisms of nuclear architecture and genome behavior in senescent cells. Ann N Y Acad Sci 1106: 250–263.
14. Couture JF, Dirk L, C. Brunzelle JS, Houtz RL, Trivel RC (2006) Structural origins for the product specificity of SET domain protein methylesterases. Proceedings of the National Academy of Sciences 105: 20659–20664.
15. Beisel C, Infuso A, Greene J, Kremmer E, Sauer F (2002) Histone methylation by the Drosophila epigenetic transcriptional regulator Ash1. Nature 419: 857–862.
16. Rayamas GV, Wendling O, Angrand PO, Mark M, Niederreither K, et al. (2003) NSD1 is essential for early post-implantation development and has a catalytically active SET domain. EMBO J 22: 3153–3163.
17. Byrd KN, Sheara A (2003) ASH1, a Drosophila trithorax group protein, is required for methylation of histone 4 residues on histone H3. Proc Natl Acad Sci U S A 100: 11535–11540.
18. Tanaka Y, Katagiri Z, Kawaihashi K, Kousu D, Kitajima S (2007) Trithorax-group protein ASH1 methylates histone H3 lysine 36. Gene 397: 161–168.
19. Yang H, Pesavento JJ, Stumnes TW, Cryderman DE, Walraat LL, et al. (2008) Preferential demethylation of histone H4 lysine 20 by Suv4-20. J Biol Chem 283: 12045-12052.
20. Protein Databank (PDB) accession ID 3R1U.
21. Zhang X, Tamaru H, Khan SI, Horton JR, Kelleci L, et al. (2003) Structure of the Neurospora SET domain protein DIM-5, a histone H3 lysine methyltransferase. Cell 111: 117–127.
22. Trivel RC, Flynn EM, Houtz RL, Hurley JH (2003) Mechanism of histone methylation by the SET domain enzyme Rubisco LSmT. Nat Struct Biol 10: 545–552.
23. Min J, Zhang X, Cheng X, Grewal SI, Xu RM (2002) Structure of the SET domain histone lysine methyltransferase CBX4. Nat Struct Biol 9: 828–832.
24. Kwon T, Chang JH, Kook E, Lee CW, Joachimiak A, et al. (2003) Mechanism of histone lysine methyl transfer revealed by the structure of SET7/9-AdoMet. EMBO J 22: 292–303.
25. Jacobs SA, Harp JM, Devarakonda S, Kim Y, Rastinejad F, et al. (2002) The active site of the SET domain is constructed on a knot. Nat Struct Biol 9: 833–836.
26. Couture JF, Collazo E, Brunzelle JS, Trivel RC (2005) Structural and functional analysis of SET1, a histone H4 Lys-20 methyltransferase. Genes Dev 19: 1455–1460.
27. Marmostein R (2003) Structure of SET domain proteins: a new twist on histone methylation. Trends Biochem Sci 28: 59–62.
28. Zhou CQa-M (2006) SET domain protein lysine methyltransferases: Structure, specificity and catalysis. Cellular and Molecular Life Sciences 65: 9.
46. Taverna SD, Li H, Ruthenburg AJ, Allin CD, Patel DJ (2007) How chromatin-binding modules interpret histone modifications: lessons from professional pocket pickers. Nat Struct Mol Biol 14: 1025–1040.
47. Veraksa A, Kermison J, McGinnis W (2002) DEAF-1 function is essential for the early embryonic development of Drosophila. Genesis 33: 67–76.
48. Spadaccini R, Perrin H, Bottomley MJ, Anseau C, Sattler M (2006) Structure and functional analysis of the MYND domain. J Mol Biol 358: 498–508.
49. Liu Y, Chen W, Gaudet J, Cheney MD, Routhia L, et al. (2007) Structural basis for recognition of SMRT/N-CoR by the MYND domain and its contribution to AML1/ETO's activity. Cancer Cell 11: 483–497.
50. Erickson P, Gao J, Chang KS, Look T, Whisenant E, et al. (1992) Identification of breakpoints in t(8;21) acute myelogenous leukemia and isolation of a fusion transcript, AML1/ETO, with similarity to Drosophila segmentation gene, runt. Blood 80: 1825–1831.
51. Brown MA, Zhu L, Schmidt C, Tucker PW (2007) Hsp90--from signal transduction to cell transformation. Biochem Biophys Res Commun 363: 241–246.
52. Whitesell L, Lindquist SL (2005) HSP90 and the chaperoning of cancer. Nat Rev Cancer 5: 761–772.
53. Congdon LM, Houston SI, Veerappan CS, Spektor TM, Rice JC (2010) PR-Srt7-mediated monomethylation of histone H4 lysine 20 at specific genomic regions induces transcriptional repression. J Cell Biochem 110: 609–619.
54. Collard RL, Harya NS, Monzon FA, Maier CE, O'Keefe DS (2006) Methylation of the ASC gene promoter is associated with aggressive prostate cancer. Prostate 66: 607–610.
55. Conway KE, McConnell BB, Bowering CE, Donald CD, Warren ST, et al. (2000) TMS1, a novel proapoptotic caspase recruitment domain protein, is a target of methylation-induced gene silencing in human breast cancers. Cancer Res 60: 6236–6242.
56. McConnell BB, Vertino PM (2004) TMS1/ASC: the cancer connection. Apoptosis 9: 5–18.
57. Stone AR, Bobo W, Brat DJ, Devi NS, Van Meir EG, et al. (2004) Aberrant methylation and down-regulation of TMS1/ASC in human glioblastoma. Am J Pathol 165: 1151–1161.
58. Kapoor-Vazirani P, Kagey JD, Powell DR, Vertino PM (2008) Role of hMOF-dependent histone H4 lysine 16 acetylation in the maintenance of TMS1/ASC gene activity. Cancer Res 68: 6810–6821.
59. Kapoor-Vazirani P, Kagey JD, Vertino PM (2011) SUV420H2-Mediated H4K20 Trimethylation Enforces RNA Polymerase II Promoter-Proximal Pausing by Blocking hMOF-Dependent HKK16 Acetylation. Mol Cell Biol 31: 1594–1609.
60. Powell HR (1999) The Rossmann Fourier autoindexing algorithm in MOSFLM. Acta Crystallogr D Biol Crystallogr 55: 1690–1695.
61. Weeks C, Miller R (1999) The design and implementation of vSs v2.0. J Appl Cryst 32: 120–124.
62. Perrakis A, Morris R, Lamzin VS (1999) Automated protein model building combined with iterative structure refinement. Nat Struct Biol 6: 450–463.
63. McRee D (1993) Struct Biol 125: 156–165.
64. Luger K, Rechsteiner TJ, Richmond TJ (1999) Preparation of nucleosome core particle from recombinant histones. Methods Enzymol 304: 3–19.
65. Luger K, Rechsteiner TJ, Flaus AJ, Waye MM, Richmond TJ (1997) Characterization of nucleosome core particles containing histone proteins made in bacteria. J Mol Biol 272: 301–311.
66. Thastrom A, Lowary PT, Widlund HR, Cao H, Kubista M, et al. (1999) Sequence motifs and free energies of selected natural and non-natural nucleosome positioning DNA sequences. J Mol Biol 288: 213–229.