A novel histone deacetylase pathway regulates mitosis by modulating Aurora B kinase activity

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Histone deacetylase (HDAC) inhibitors perturb the cell cycle and have great potential as anti-cancer agents, but their mechanism of action is not well established. HDACs classically function as repressors of gene expression, tethered to sequence-specific transcription factors. Here we report that HDAC3 is a critical, transcription-independent regulator of mitosis. HDAC3 forms a complex with A-Kinase-Anchoring Proteins AKAP95 and HA95, which are targeted to mitotic chromosomes. Deacetylation of H3 in mitosis requires AKAP95/HA95 and HDAC3 and provides a hypoacetylated H3 tail that is the preferred substrate for Aurora B kinase. Phosphorylation of H3S10 by Aurora B leads to dissociation of HP1 proteins from methylated H3K9 residues on mitotic heterochromatin. This transcription-independent pathway, involving interdependent changes in histone modification and protein association, is required for normal progression through mitosis and is an unexpected target of HDAC inhibitors, a class of drugs currently in clinical trials for treating cancer.

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well studied. Acetylation of core histone tails by histone acetyltransferases [HATs] is a paradigm for transcriptional activation and is reversed by histone deacetylases [HDACs] [Grunstein 1997; Wade et al. 1997; Cheung et al. 2000b]. Notably, hypoaecytlation of histones has also been found during mitosis. The acetylation of histones begins to decrease during the early stages of mitosis, is absent in metaphase and anaphase, then gradually increases in the late mitotic phase [Marian and Wintersberger 1982; Jeppesen et al. 1992; Kruhlak et al. 2001; Chen et al. 2005; Valls et al. 2005; Nishiyama et al. 2006].

A role for histone deacetylation during mitosis has been further suggested by the cell cycle effects of HDAC inhibitors [Marks et al. 2001; Wong et al. 2005]. HDAC inhibitors have attracted considerable interest in the clinic, with early studies suggesting anti-tumor efficacy for a range of malignancies including T-cell lymphoma [Marks et al. 2001; Zhang et al. 2005a]. However, the molecular mechanisms of their anti-tumor effects are not well established. Most investigators have focused on transcriptional mechanisms, and in particular on the induction of the cell cycle inhibitor p21, but while HDAC inhibitors globally increase histone acetylation, the level of transcription is altered in only 2%–5% of all expressed genes [Della Ragione et al. 2001; Johnstone and Licht 2003]. Moreover, transcriptional mechanisms do not explain the dramatic mitotic effects of HDAC inhibition, which impairs mitotic progression with associated defects in chromosome condensation, segregation, and kinetochore assembly [Qiu et al. 2000; Sandor et al. 2000; Shin et al. 2003; Warrener et al. 2003; Mikhailov et al. 2004; Robbins et al. 2005].

HDAC3 is widely expressed, highly regulated, and mediates its function through protein–protein interactions [Emiliiani et al. 1998; Guenther et al. 2002; Yang et al. 2002]. Immature HDAC3 is folded by the TCP-1 ring complex (TRIC) before interacting with the transcriptional corepressor protein N-CoR [nuclear receptor corepressor] and SMRT [silencing mediator for retinoid and thyroid hormone receptors] [Li et al. 2000; Guenther et al. 2002]. N-CoR and SMRT serve as activating cofactors for HDAC3 enzymatic activity [Guenther et al. 2001; Zhang et al. 2002]. HDAC3 activity is also regulated through phosphorylation by casein kinase II and dephosphorylation by protein phosphatase 4 [Zhang et al. 2005b]. HDAC3 is required for transcriptional repression by nuclear hormone receptors [Ishizuka and Lazar 2003, 2005] and also suppresses the transcriptional potential of other transcription factors including YY1, GATA-2, and TFII-I [Ozawa et al. 2001; Yao et al. 2001; Wen et al. 2003]. HDAC3 knockdown was recently shown to increase cells in G2/M phase in human colon cancer cells [Wilson et al. 2006], but the molecular mechanism of this effect is unknown.

Here we report that HDAC3 forms a complex with A-Kinase Anchor Protein 95 (AKAP95) and its relative Homologous to AKAP95 (HA95) in mitotic cells. AKAP95 and HA95 have been previously shown to be recruited to mitotic chromosomes and involved in mitotic progression, but their mechanism of action is obscure [Collas et al. 1999; Orstavik et al. 2000; Steen et al. 2000]. We find that HDAC3, along with AKAP95/HA95, is required for normal mitotic progression, and that both HDAC3 and AKAP95/HA95 are essential for global histone deacetylation that occurs during mitosis. Unexpectedly, HDAC3 and AKAP95/HA95 are also required for maximal phosphorylation of H3S10 in mitosis. Reduction in HDAC3 level and pharmacological inhibition of HDAC activity leads to increased histone acetylation, decreased H3S10 phosphorylation, and accumulation of cells in mitosis, suggesting that inhibition of HDAC3 is sufficient to explain the cell cycle effects of HDAC inhibitors. The reduced phosphorylation H3S10 can be explained by the observation that the H3S10 kinase activity of Aurora B is dependent on the histone acetylation state, with hypoaecytlated histone tails being the preferred substrate. HP1 dissociation from mitotic chromosomes is prevented by knockdown of AKAP95/HA95, selective loss of HDAC3, or pharmacological HDAC inhibitors, demonstrating that the AKAP–HDAC3–Aurora B axis is upstream of the Aurora B–HP1 “meth-phos switch.” This novel pathway explains the changes in histone modifications that occur during mitosis, and suggests that HDAC inhibitors used in cancer treatment target HDAC3 via this nontranscriptional mechanism.

Results

AKAP95 and HA95 are present in high-molecular-weight HDAC3 complexes and interact directly with HDAC3

To explore the biological functions of HDAC3, we purified endogenous proteins that coimmunoprecipitated with Flag-HDAC3 stably expressed in 293T cells. Proteins identified through silver staining and mass spectrometry included expected components of HDAC3 complexes including N-CoR, SMRT, TBL1, and GPS2 [Fig. 1A; Guenther et al. 2000; Li et al. 2000; Zhang et al. 2002]. An additional band, corresponding to a molecular mass of ~95 kDa, was found to contain both AKAP95 and HA95, and further verified by Western analysis [Fig. 1A]. To further test the intracellular interaction of AKAP95 with HDAC3, HA-epitope tagged-AKAP95 was expressed in 293T cells, followed by immunoprecipitation with anti-HA antibody. Overexpressed AKAP95 was associated with endogenous HDAC3, as well as endogenous N-CoR [Fig. 1B] and SMRT [data not shown]. Endogenous HDAC3 was also reliably detected in immunoprecipitates of endogenous HA95, establishing that the in vivo association of AKAP95/HA95 with HDAC3 was not a result of protein overexpression [Fig. 1C]. Moreover, recombinant HDAC3 [Fig. 1D], but not other components of HDAC3 complexes [data not shown], interacted with recombinant AKAP95 and HA95 in vitro, suggesting that AKAP95 and HA95 are recruited to the complex through HDAC3 [Fig. 1D]. Activation of HDAC3 by the SMRT deacetylase-activating domain

AKAP–HDAC3–Aurora B–HP1 pathway regulates mitosis
Interaction of HDAC3 with AKAP95 and HA95 increases during mitosis

AKAP 95 and HA95 have been previously found to be associated with mitotic chromosomes (Collas et al. 1999, Martins et al. 2000, Steen et al. 2000, Eide et al. 2002), and so we next investigated the AKAP-HDAC3 interaction during mitosis. Immunoprecipitation of endogenous HDAC3 with endogenous AKAP95 or HA95 was markedly increased in nocodazole-synchronized cells blocked in G2/M (Fig. 2A). Conversely, coimmunoprecipitation of AKAP95 and HA95 with HDAC3 was also increased in nocodazole-treated cells (Fig. 2B). In contrast, the interaction of HDAC3 with SMRT or N-CoR was essentially unaltered in nocodazole-synchronized cells (Fig. 2B). To exclude the possibility that the increased HDAC3-AKAP95/HA95 interactions were due to drug toxicity, 20 h of aphidicolin treatment was used to synchronize 293T cells at the G1/S transition, then cells were released from aphidicolin treatment for 2 or 12 h to obtain S-phase- and G2/M-enriched cell populations, respectively. The AKAP-HDAC3 interactions were not increased either in G1/S- or S-phase-synchronized cells (Fig. 2C). However, the interaction between endogenous AKAP95 and HDAC3 was robustly increased in G2/M cells, which have the highest level of H3S10 phosphorylation (Fig. 2C). Using immunofluorescence staining and confocal microscopy, AKAP95 and HDAC3 colocalization was also evident in early mitotic cells compared with cells at interphase (Fig. 2D). To better characterize the mitotic AKAP/HDAC3 complex, HDAC3 complexes prepared from nocodazole-synchronized Flag-HDAC3 stable cells were eluted from immunoprecipitates using Flag peptide then size separated by gel filtration (Fig. 2E). HDAC3 cochromatographed with SMRT in two catalytically active high-molecular-weight complexes (∼1.5–2 mDa and ∼550 kDa) (Guenther et al. 2000; Li et al. 2000; Zhang et al. 2002), and was also detected in lower-molecular-weight complexes containing the TCP1β subunit of the TRiC chaperone machinery (Guenther et al. 2002). AKAP95 and HA95 were present in the higher-molecular-weight complexes that also contained SMRT (and N-CoR, TBL1, and GPS) [data not shown].

Depletion of AKAP95 and HA95 induces G2/M arrest and mitotic defects

AKAP95 and HA95 have been proposed to have a role in chromosome condensation during mitosis (Collas et al. 1999, Martins et al. 2000, Steen et al. 2000, Eide et al. 2002). We used short-hairpin RNA [shRNA] to knock down AKAP95 and HA95 [Fig. 3A] in order to study their function in the cell cycle. Knockdown of AKAP95 or HA95 individually had modest effects on the cell cycle (data not shown). However, simultaneous depletion of both AKAP95 and HA95 substantially increased the percentage of G2/M cells as determined by flow-assisted cytometric analysis [FACS] (Fig. 3B). This was, in large part, due to a significantly increased proportion of cells in mitosis (Fig. 3C). Of note, many of these mitotic cells were of quite aberrant morphology, which included incomplete chromosomal condensation or “lagging” chromosomes [Fig. 3D, indicated by the arrowhead in the middle panel, and in higher magnification inset], chromosome segregation defects like a chromosome bridge [Fig. 3D, the right panel], as well as tri- or multipolar mitoses [Fig. 6E [below], bottom right panel]. In contrast,
few mitoses were found after control shRNA, and these were largely unperturbed (Fig. 3C,D).

Depletion of HDAC3 induces G2/M arrest and mitotic defects similar to those caused by AKAP95/HA95 knockdown

We next tested the effects of HDAC3 knockdown, again via shRNA [Fig. 4A]. Knockdown of HDAC3 also substantially increased the percentage of G2/M cells [Fig. 4B], and mitotic counts again revealed an increased proportion of mitotic cells [Supplementary Fig. 2A]. A large proportion of the cells exhibited aberrant mitoses in the absence of HDAC3 [Supplementary Fig. 2B], similar to that seen with knockdown of AKAP95/HA95 [cf. Fig. 3D]. Thus, knockdown of HDAC3 or AKAP95/HA95 led to similar impairment of mitotic progression and increased aberrancy of mitotic figures.

HDAC inhibition blocks the cell cycle at G2/M phase in a transcriptional-independent manner

Nonspecific inhibition of HDACs by pharmacological inhibitors has been shown to induce cell cycle arrest beyond prometaphase, in association with mitotic defects reminiscent of the effects of selective loss of HDAC3 demonstrated above [Qiu et al. 2000; Sandor et al. 2000; Shin et al. 2003; Mikhailov et al. 2004; Robbins et al. 2005]. Indeed, the nonselective HDAC inhibitor trichostatin A (TSA) caused G2/M arrest both in human MCF7 breast carcinoma cells [Fig. 5A] as well as in human 293T embryonic kidney cells [Fig. 5B], similar to the effect of selective depletion of HDAC3. In contrast, shRNA depletion of the related HDAC1 did not cause mitotic arrest [Supplementary Fig. 3a], suggesting that HDAC3 is a major target by which HDAC inhibitors induce G2/M arrest. The G2/M arrest due to TSA was reversible [Supplementary Fig. 4], and two additional
nonspecific HDAC inhibitors (sodium butyrate and valproic acid), also blocked the cell cycle in G2/M (Fig. 5C). Importantly, the G2/M arrest due to each of the three HDAC inhibitors was maintained in the presence of actinomycin D (Act D) (Fig. 5C), at a concentration that blocked global transcription by >90% (Supplementary Fig. 5). Act D alone has been reported to induce a partial G1 arrest (Kim et al. 2005), and this was observed in our studies as well (Fig. 5C). These data implied that the major effect of HDAC inhibition on the cell cycle—i.e., G2/M arrest—was not due to effects of these drugs on transcription. This, in turn, suggested that failure to deacetylate histones during mitosis directly impairs mitosis progression.

Histone deacetylation during mitosis requires HDAC3 and AKAP95/HA95

Our finding of a role for HDAC3 and AKAP95/HA95 in mitosis led us to consider a potential regulatory role for histone deacetylation. Consistent with previous observations (Marian and Wintersberger 1982; Jeppesen et al. 1992; Kruhlak et al. 2001; Valls et al. 2005), we observed that H3 and H4 are globally hypoacetylated during mitosis (Fig. 6A). This result was confirmed by quantitative mass spectrometric analysis of H3 isolated from nonsynchronized or nocodazole-synchronized HeLa cells (Fig. 6B). Abundance levels of unmodified and trimethylated K9 (Fig. 6B), as well as mono- and dimethylated K9 (data not shown), remain consistent in nonsynchronized and mitotic HeLa cells. However, single acetylation of K9 or K14 and K9/K14 double acetylation dramatically decreased [twofold and sevenfold, respectively] in mitotic cells (Fig. 6B). Similarly, we also noted the decreases in single acetylation of H3K18 or K23 as well as K18/K23 dual acetylation in nocodazole-synchronized HeLa cells (Supplementary Fig. 6). These results further confirm that H3 is globally deacetylated on multiple lysines in mitotic cells.

We next used shRNA to reduce cellular HDAC3 levels and determine whether deacetylation of histones during mitosis is dependent on HDAC3. Indeed, HDAC3 knockdown abolished the global histone deacetylation characteristic of mitotic, nocodazole-treated cells, but had little effect on global histone acetylation in nonsynchronized cells (Fig. 6C). This result indicated that HDAC3 is required for the deacetylation of mitotic histones. Knockdown of N-CoR or SMRT individually did not significantly change the cell cycle progression or histone acetylation [data not shown], suggesting that they redundantly activate HDAC3 in this context. We next used shRNA to knock down AKAP95 and HA95 to address whether the HDAC3-dependent histone deacetylation during mitosis requires these AKAPs. Individual knockdown of AKAP95 or HA95 modestly reduced the deacetylation of H3 and H4 during mitosis (data not shown). More strikingly, simultaneous knockdown of AKAP95 and HA95 completely prevented the mitotic deacetylation of H3 and H4 (Fig. 6D). Note that phosphorylation of H3S10, which occurs in mitosis, did not alter our ability to detect H3 acetylation level (Supplementary Fig. 7).
HDAC3 and AKAP95/HA95 are required for normal phosphorylation of H3 during mitosis

H3S10 phosphorylation is a mitotic marker, and therefore should increase in HDAC3- or AKAP95/HA95-depleted cells that contain numbers of mitotic cells. Paradoxically, however, phosphorylation of H3S10 was actually reduced when HDAC3 or AKAP95 levels were depleted (Fig. 6C,D). Immunofluorescence analysis of AKAP95/HA95 double-knockdown HeLa cells also revealed marked reduction in H3 S10 phosphorylation within mitotic nuclei (Fig. 6E). These data are consistent with recent reports that TSA treatment decreased phosphorylation of H3S10 in cancer cells (Dowling et al. 2005; Robbins et al. 2005). Indeed, we confirmed that TSA treatment reduced H3S10 phosphorylation in MCF7 breast cancer cells (Fig. 6F) despite increased the number of cells in G2/M (see Fig. 5A). These results suggest an unanticipated relationship between AKAP95/HA95–HDAC3-dependent histone deacetylation and the mitosis-associated phosphorylation of H3S10.

Histone acetylation is inversely correlated with H3S10 phosphorylation in cells

The hypothesis that phosphorylation of H3S10 occurs mainly on hypoacetylated histone during mitosis was tested in 293T cells treated with nocodazole for 6 h to partially enrich the mitotic cells. Chromosome-bound histones were isolated by acid extraction, and the histones were incubated with anti-acetylated H3 (H3ac) anti-
tibodies to immunoprecipitate the hyperacetylated histones; hypoacetylated histones remained in the supernatant [Fig. 7A]. H3S10 phosphorylation was detected in the hypoacetylated H3 fraction [supernatant], but not in the hyperacetylated H3 fraction [immunoprecipitate] [Fig. 7A].

We also tested this hypothesis by quantitative mass spectrometry [Fig. 7B]. We reasoned that if H3S10 phosphorylation is affected by a given modification, that modification should be disproportionately represented in the H3S10-phosphorylated tails [cf. the H3 tails as a whole]. Indeed, acetylation of H3 at K9 or K14 was underrepresented by ~10-fold in H3S10-phosphorylated tails [0.22% as compared with 2.6% in the total H3 pool]. In contrast, we observed a similar level of H3K9Me in total H3 tail peptides [22.6%] as in H3S10-phosphorylated peptides [24.8%], suggesting that H3S10 phosphorylation is not affected by methylation of K9. Moreover, we could not detect any S10 phosphorylated, doubly acetylated H3 tail peptide with current MS detection limits [Supplementary Fig. 8c; the calculated mass is 561.2708 m/z], although it is possible that the double acetyl/phospho-peptide exists at extremely low levels that may be detected with higher sensitivity immunoassay methods [Kruhlak et al. 2001]. These data, while correlative, are consistent with the conclusion that acetylation of H3 tails interferes with phosphorylation of the H3 tails, and, thus, hypoacetylation of H3 tails appears to be a prerequisite for mitotic H3S10 phosphorylation.

**The H3S10 kinase Aurora B is present in HDAC3 complexes during mitosis**

Thus far we have demonstrated that AKAP95/HA95–HDAC3-dependent histone deacetylation correlates with the mitotic phosphorylation of H3S10. We hypothesized that the mechanism underlying this phenomenon involved Aurora B, the kinase that phosphorylates H3S10 during mitosis [Crosio et al. 2002; Meraldi et al. 2004]. Indeed, we noted that Aurora B is associated with the HDAC3 complex during mitosis [Fig. 2B], and that after gel filtration of purified HDAC3 complexes, Aurora B is present in the high-molecular-weight fractions that also contain AKAP95/HA95 and SMRT [Fig. 2E]. Thus, Aurora B is a present in high-molecular-weight HDAC3 complexes.

**Hypoacetylated H3 is the preferred substrate for Aurora B kinase**

Consistent with a previous report [Shindo et al. 1998], Aurora B level is increased in cells arrested in mitosis [Supplementary Fig. 9]. However, depletion of AKAP95/HA95 or HDAC3 had little effect on Aurora B expression [Supplementary Fig. 9]. We therefore hypothesized that the HDAC3-dependence of H3S10 phosphorylation was due to preferential phosphorylation of hypoacetylated H3 tails by Aurora B. This hypothesis was tested using an in vitro kinase assay to compare the ability of purified recombinant Aurora B to phosphorylate H3S10 in hyperacetylated versus hypoacetylated histone substrates. We first confirmed that H3S10 phosphorylation level was low and similar in the preparations of hypoacetylated and hyperacetylated histones [Supplementary Fig. 10]. Consistent with our hypothesis, Aurora B more efficiently phosphorylates the hypoacetylated histone substrate, although its autophosphorylation was the same under both conditions [Fig. 8A].

We also examined whether the presence of the HDAC3/SMRT complex would facilitate S10 phosphorylation by native Aurora B complex purified from 293T cells. The result indicated that incubation of the purified HDAC3/SMRT complex with purified endogenous Aurora B kinase complex dramatically increased the phosphorylation of H3S10 in an in vitro kinase assay [Fig. 8B]. This result was not an artifact of the Western detection of H3S10 phosphorylation interfered by acetylation of H3, because in vitro deacetylation of the hyperacetylated histones using recombinant HDAC3/SMRT complex after the kinase assay did not increase the phosphorylation signal [Supplementary Fig. 11]. Taken together, these data suggest that Aurora B kinase preferentially phosphorylates hypoacetylated H3. Based on these observations, we reason that hypoacetylated H3 is the preferred substrate for Aurora B kinase during mitosis.

**Figure 7.** Phosphorylation of H3S10 occurs mainly on hypoacetylated H3 tails. (A) 293T cells were synchronized with nocodazole for 6 h to produce ~40% G2/M cells. The cells were acid-extracted, and 10 μg of extract was incubated with anti-H3ac antibody. The supernatant [Sup] and immunoprecipitate [IP] fractions were analyzed for phosphorylation and acetylation levels by Western blot. (B) Relative abundance of total unmodified H3K9, trimethylated H3K9 [H3K9me3], as well as acetylated H3K9 or H3K14 [H3K9ac or K3K14ac] peptides [open bars], with the percentage of each modified peptide in combination with H3S10 phosphorylation in the phosphorylated peptide population [black bars] isolated from nocodazole-synchronized mitotic HeLa cells as determined by mass spectrometry [Supplementary Fig. 8c].
which explains why phosphorylation of H3S10 decreased in HDAC3 and AKAP95/HA95 complex-depleted mitotic cells.

**The AKAP95/HA95–HDAC3–Aurora B pathway is required for HP1β dissociation during mitosis**

The physiological function of mitotic H3S10 phosphorylation has been recently clarified by the recent finding that Aurora B phosphorylation of H3S10 results in the dissociation of HP1 proteins, notably HP1β/H9252, despite persistent H3K9 trimethylation (Fischle et al. 2005; Hirota et al. 2005). We hypothesized that the AKAP95/HA95–HDAC3–Aurora B pathway we have uncovered is upstream of this “meth-phos switch.” Indeed, the overall decrease in H3S10 phosphorylation after knockdown of either HDAC3 or AKAP95/HA95 also occurs on H3 tails in which H3K9 is trimethylated, although the overall trimethylation of H3K9 was unchanged in HDAC3 or AKAP95/HA95 knockdown cells (Fig. 9A). Knockdown of HDAC1, which did not change acetylation levels of histones from mitotic-arrested cells, also did not reduce this double modification [Supplementary Fig. 3b].

If the AKAP95/HA95–HDAC3–Aurora B pathway is, indeed, a prerequisite for normal mitotic H3S10 phosphorylation leading to HP1β dissociation, then knockdown of the individual components of this pathway should result in retention of HP1β on mitotic chromatin. To test this hypothesis, HeLa cells were transfected with shHDAC3 or shAKAP95/shHA95 constructs, then stained with anti-HP1β after extraction of the soluble proteins to reduce the interference from soluble non-chromatin-bound proteins. As expected, in cells treated with control shRNA, HP1β was not extracted from interphase cells, but was completely extracted from mitotic cells [Fig. 9B, arrow]. In contrast, HP1β was retained by mitotic chromatin in cells depleted of HDAC3 or AKAP95/HA95 [Fig. 9B]. This was very similar to the effect of pharmacological inhibition of Aurora B using hesperadin, the hallmark of the “meth-phos switch” [Fischle et al. 2005; Hirota et al. 2005], which we also observed [Fig. 9C]. These results demonstrate that normal HP1 dissociation from mitotic chromosomes requires the HDAC3–AKAP95/HA95–Aurora B pathway, acting upstream of the H3S10-phosphorylation-dependent displacement of HP1 from heterochromatin. They further predict that HDAC inhibitors acting on this pathway to induce mitotic arrest should cause HP1β retention. Indeed, the dissociation of HP1β from mitotic...
heterochromatin was prevented by treatment with TSA (Fig. 9C).

Discussion

The classic role of HDAC3 has been that of a transcriptional repressor of gene expression, as part of a complex tethered to sequence-specific transcription factors. Here we have reported the unexpected finding that HDAC3 has a critical, transcription-independent function in mitosis. In interphase cells, AKAP95/HA95 binds to the nuclear matrix (Collas et al. 1999; Martins et al. 2000; Akileswaran et al. 2001) and is less associated with HDAC3. HP1 proteins are recruited to methylated H3K9 in heterochromatin. When cells enter into mitosis, AKAP95/HA95 may target the HDAC3 complex to deacetylate H3, in a reaction that is blocked by HDAC inhibitors, and thereby provides a hypoacetylated H3 tail as substrate for Aurora B to phosphorylate on S10. Phosphorylation of S10 by Aurora B then dissociates HP1 proteins from methylated H3K9 residues on mitotic heterochromatin, which has been referred to as the “meth-phos switch” (Fischle et al. 2005; Hirota et al. 2005). These interdependent changes in histone modification and protein association are required for normal progression through mitosis, perhaps by facilitating chromosome condensation, or by serving as the indicator for the mitotic checkpoint to control proper cell division (Fig. 10).

While the transcriptional effect of HDAC inhibitors on specific genes, such as p21 and other cell cycle-regulated genes, has been reported to contribute to their anti-tumor actions, especially in G1-phase arrest (Archer et al. 1998; Sambucetti et al. 1999; Xiao et al. 1999; Richon et al. 2000; Johnstone 2002), their direct effects on histone acetylation levels may be equally important for the anti-tumor activity because of the important functions of histones in different cellular processes, including mitosis [Johnstone and Licht 2003]. It is increasingly clear that HDAC inhibition induces G2/M arrest in many human cell lines and causes mitotic defects in different cancer cell lines (Qiu et al. 2000; Taddei et al. 2001; Bali et al. 2005; Dowling et al. 2005; Nome et al. 2005; Wong et al. 2005). We have confirmed this and, furthermore, find that this effect of HDAC inhibition is independent of ongoing gene transcription, suggesting direct effects of histone hyperacetylation on mitosis. Our results indicate that the hyperacetylation of histones induced by HDAC inhibitors directly interfere with mitotic progression.

Global histone acetylation is reduced during mitosis (Marian and Wintersberger 1982; Jeppesen et al. 1992; Kruhlak et al. 2001; Chen et al. 2005; Li et al. 2005; Valls et al. 2005; Nishiyama et al. 2006). Our studies reveal that HDAC3 and its partner proteins AKAP95 and HA95 are required for global histone deacetylation during mitosis. Of note, the most dramatic change in acetylation that occurs during mitosis is hypoacetylation of Lys 5 of H4 (Kruhlak et al. 2001), which matches the substrate specificity of HDAC3 [Hartman et al. 2005]. Moreover, our results clearly show that HDAC3 is required for normal mitotic progression. This is consistent with a recent study in which knockdown of HDAC3, but not HDAC1 or HDAC2, increased cells in G2/M phase in human colon cancer cells [Wilson et al. 2006]. Furthermore, knockdown of HDAC3 or AKAP95/HA95 also mimicked the effects of nonselective HDAC inhibition on phosphorylation of H3S10 and retention of HP1β proteins on mitotic chromosomes. Inhibition of HDAC3 is therefore likely to be the mechanism by which HDAC inhibitors induce the G2/M block in the cell cycle. The transcription independence of this effect, while unexpected, is completely consistent with a direct mitotic function of HDAC3 in the context of the novel pathway that we report here.

Specific patterns of histone modification at gene promoters regulate transcription via a “histone code” [Jenuwein and Allis 2001]. Notably, the transient phosphorylation of H3S10 has been reported in the promoter region of many mammalian immediate-early genes, which are
HP1 pathway regulates mitosis

Materials and methods

Plasmids and reagents

Full-length AKAP95 and HA95 were obtained from HeLa cells by RT–PCR and cloned into pcDNA3.1A vector (Invitrogen). HA-epitope tags were added to the N terminus of HA95 and the C terminus of AKAP95. The pTRE-HDAC3-Flag expression vector was constructed by inserting HDAC3-Flag [Guenther et al. 2001] into the pTRE vector [BD Clontech]. The SMRT DAD domain or SMRT 1–763, fused to the Gal4 DNA-binding domain, were described previously [Yu et al. 2003]. Anti-HA95 and AKAP95 antibodies were raised against KLH-conjugated peptides HA95 [amino acids 65–85] and AKAP95 [amino acids 672–688] and affinity-purified. Anti-GPS2 antibody was raised against the peptide from 307 to 327 [Covance]. N-CoR and TBL1 antibodies were described previously [Huang et al. 2000; Guenther et al. 2001]. Other antibodies were purchased as follows: H4, H3ac, H4ac, H3S10ph, and H3Kme3S10ph from Upstate Biotechnology, H3, H3K9acS10ph, anti-HDAC3 [rabbit], and anti-Aurora B from Abcam; anti-HDAC3 (mouse), anti-SMRT, and anti-Ran from BD PharMingen; mouse anti-HDAC3 for immunoprecipitation from Upstate Biotechnology [clone 3G6]; anti-SMRT/N-CoR from Affinity Bioreagents; anti-GAPDH from Santa Cruz Biotechnology; anti-Tcp1β from Stressgen Biotechnologies; anti-Flag from Sigma; and anti-HA tag from Roche Applied Science. TSA, sodium butyrate, VPA, nocardazole, paclitaxel, and aphidicolin were purchased from Sigma.

Cell line establishment and culture conditions

293T, HeLa, and MCF7 cells were kept in DMEM with 10% FBS. The 293T tet-off cell line [BD Clontech] was transfected with pTRE-HDAC3-Flag or pTRE empty vector and selected under 50 µg/mL hygromycin in DMEM with 5% FBS. Positive clones were confirmed by Western blot. The HDAC3-Flag expression level of the selected clone was about five times higher than endogenous HDAC3 [data not shown]. The HDAC3 stable cell lines were kept in DMEM with either 5% tet-approved FBS [BD Clontech] to express HDAC3-Flag or 5% FBS with 2 µg/mL Doxycycline [Sigma] to turn off HDAC3 expression.

Affinity purification and gel filtration

Flag-HDAC3 stable cell pellets were resuspended in BC100 buffer [20 mM HEPES, 100 mM KCl, 0.2% NP-40, 0.2 mM EDTA, 10% glycerol, 0.2 mM DTT; protease inhibitor cocktail [Roche]] followed by brief sonication to lyse the cells. Whole-cell lysates were incubated with anti-Flag M2 agarose (Sigma) in BC100 buffer overnight at 4°C. After extensive washing with BC100 and BC300 [20 mM HEPES, 300 mM KCl, 0.2% NP-40, 0.2 mM EDTA, 10% glycerol, 0.2 mM DTT], associated complexes were eluted by 0.5 mg/mL Flag peptide (Sigma) or 0.2 mg/mL 3xFlag peptide (Sigma) in BC100 buffer or BC100 without NP-40 (for gel filtration). For gel filtration, eluents were concentrated to 0.15 mL on an UltrAFree cellulose concentration device [Millipore]. The samples were loaded on a Superose 6 10/30 gel filtration column [Amersham Bioscience] in BC150 [150 mM KCl] without NP-40. Eluents were collected at 0.5 mL per fraction. Fractions from the Superose 6 column were subjected to immunoblot analysis.

Protein identification

Protein identification by mass spectrometry was performed as described [Jung et al. 2005].
Immunoprecipitation

Cells were lysed with BC100 for 10 min on ice followed by brief sonication. After centrifugation, the supernatants were pre-cleared with protein A agarose, and then allowed to bind either anti-HDAC3 or anti-HA95 and anti-AKAP95 overnight at 4°C. Protein A agarose beads were added and incubated for another 2 h. The beads were washed five times in BC100 and once in BC300 and subjected to immunoblot analysis. For in vitro binding assays, Flag-HDAC3 and Gal-SMRT[1–763] were in vitro translated with the T7 TNT Quick Coupled Transcription/Translation System (Promega) and incubated with anti-Flag M2 agarose (Sigma) in BC100 buffer overnight at 4°C. HDAC3-coupled beads were collected and incubated with in vitro translated, 35S-labeled HA95 or AKAP95, which were pre-cleared with anti-Flag M2 agarose in BC100 buffer. After extensive washing, bound proteins were subjected to autoradiography and immunoblot analysis.

shRNA transfection

Vectors expressing shRNA under the control of the human H1 promoter were constructed by inserting pairs of annealed DNA oligonucleotides into the pSuper vector (Oligoengine) between BglII and HindIII restriction sites or pSilencer vector (for HDAC1 and control for HDAC1; Ambion). The target sequences were as follows: nontargeted control: 5′-AGACACACGCCACTCGTC-3′; HA95-1: 5′-GCCAGGCATCTTGACAGCAA-3′; AKAP95-1: 5′-GCCAGGACCTCTTCTCACA-3′; HDAC1, HDAC3 control for HDAC1: see Ishizuka and Lazar (2003).

293T or HeLa cells were transfected with shRNAs using Lipofectamine 2000 (Invitrogen Gibco) according to the manufacturer’s instructions. Cells were divided 24 h after transfection, followed by a second transfection 24 h later with the same shRNA constructs. Cells were split after an additional 24 h incubation, and specific drugs were added at the indicated concentration and collected at various time points.

In vitro kinase assay and HDAC assay

One microgram of sodium-butyrate-treated [hyperacetylated] or untreated histones from HeLa cells [Upstate Biotechnology] was incubated with 2.5 µM of active Aurora B (Upstate Biotechnology) in TAB kinase buffer (Upstate Biotechnology) for 10 min at 30°C. The reaction included either Mg2+ ATP cocktail (Upstate Biotechnology) or 0.2 µCi of [γ-32P]ATP plus Mg2+ ATP cocktail. The reactions were stopped by addition of 3× SDS loading buffer and boiling for 5 min at 95°C. Phosphorylation of histones was detected by Western blot or autoradiography. After kinase assay, the samples were diluted with HDAC buffer together with beads alone or anti-Flag agarose-immobilized HDAC3 and SMRT DAD, which are overexpressed in 293T cells. HDAC assay was performed as described previously [Yu et al. 2003]. In vitro kinase assays were also performed using affinity-purified Aurora B complex from 293T cells. Five micrograms of hyperacetylated histones were incubated with immobilized HDAC3-SMRT DAD complex or beads alone at 37°C in TAB kinase buffer. After 20 min of incubation, affinity-purified Aurora B complex from 293T cells was added to the reactions with Mg2+ ATP cocktail. The reactions were incubated for 15 min more and stopped by addition of SDS loading buffer and boiling.

FACS

Cells were trypsinized, washed with PBS, and fixed with ice-cold 70% ethanol overnight at 4°C. Nucleus DNA was stained using a solution with 50 µg/mL propidium iodide [Sigma] and 1 mg/mL RNase A in PBS (Jin et al. 2002). The cells were analyzed on FACS Calibur (BD Biosciences) using CellQuest and ModFit data analysis software.

Immunofluorescence

For colocalization staining of AKAP95 and HDAC3, HeLa cells cultured on Lab-TekII Chamber slides were washed three times in PBS, fixed for 15 min in 3% paraformaldehyde [PFA] at room temperature, washed, and fixed with cold methanol for 5 min. After two additional washes, cells were incubated with blocking buffer [5% donkey serum in PBS] for 1 h at room temperature. Primary antibodies were diluted in blocking buffer and incubated with cells at 4°C overnight. After washing with PBS three times, cells were incubated with secondary antibodies conjugated with either Alexa 594 or Alexa 488 (Molecular Probes) for 1 h at room temperature. Then DNA was stained with 0.1 mg/mL 4′,6-diamidino-2-phenylindole [DAPI, Sigma] for 15 min at room temperature. After washing, slides were mounted with Prolong Gold anti-fade reagent. Slides were viewed with a Zeiss LSM-510 META confocal microscope, using LSM 510 version 3.2 software to acquire images.

Alternatively, cells were cultured on chamber slides and incubated with 0.5% Triton X-100 in PBS for 5 min at room temperature to extract non-chromatin-bound proteins. After fixing and staining with indicated antibodies or DAPI, additional imaging was performed with a 100× PlanNeofluor objective mounted on a Nikon TE-200 microscope equipped with epifluorescence optics. With the latter, images were captured with a Hamamatsu CCD camera that was controlled with IP LabSpectrum version 2.0.1 (Scanalytics, Inc.).

Mass spectrometry (MS)

The mass spectrometry data shown in Supplementary Figures 6 and 8 are summarized in Figures 6B and 7B. H3 from asynchronously grown and mitotically arrested HeLa cells were first chemically derivatized with propionylation reagent [Aldrich] as previously described [Syka et al. 2004]. H3 was dried to near dryness and redissolved in 25 µL of deionized water, diluted with 50 µL of 100 mM ammonium bicarbonate buffer solution [pH 8], and digested with trypsin [Promega] at a substrate:enzyme ratio of 20:1 for 5 h at 37°C. All reactions were quenched by the addition of concentrated acetic acid and freezing. Propionylated histone digest mixtures were loaded onto capillary precolumns packed with 5 cm of irregular C18 resin, washed with 0.1% acetic acid, and connected to analytical columns packed with 8 cm of regular C18 resin [5 µm, YMC, Inc.] constructed with integrated electrospray emitters as previously published [Martin et al. 2000]. All samples were analyzed by nanoflow HPLC-microelectrospray ionization on a Finnigan linear quadrupole ion trap-Fourier Transform Ion Cyclotron Resonance (LTQ-FT-ICR) mass spectrometer [Thermo Electron] as previously described [Hake et al. 2006].

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