RNA Polymerase II Carboxyl-terminal Domain Phosphorylation Regulates Protein Stability of the Set2 Methyltransferase and Histone H3 Di- and Trimethylation at Lysine 36*§

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Background: Set2 catalyzes mono-, di-, and trimethylation of lysine 36 on histone H3 (H3K36).

Results: Phosphorylation of the C-terminal domain of RNA polymerase II (RNAPII CTD) regulates Set2 protein levels and H3K36 methylation.

Conclusion: Set2 protein is regulated to maintain balanced levels of H3K36 methylation.

Significance: These results suggest that different methylation states perform distinct biological functions.

Methylation of lysine 36 on histone H3 (H3K36) is catalyzed by the Set2 methyltransferase and is linked to transcriptional regulation. Previous studies have shown that trimethylation of H3K36 by Set2 is directed through its association with the phosphorylated repeats of the RNA polymerase C-terminal domain (RNAPII CTD). Here, we show that disruption of this interaction through the use of yeast mutants defective in CTD phosphorylation at serine 2 results in a destabilization of Set2 protein levels and H3K36 methylation. Consistent with this, we find that Set2 has a short half-life and is co-regulated, with RNAPII CTD phosphorylation levels, during logarithmic growth in yeast. To probe the functional consequence of uncoupling Set2-RNAPII regulation, we expressed a truncated and more stable form of Set2 protein levels and H3K36 methylation. Phosphorylation at serine 2 results in a destabilization of Set2 protein levels and H3K36 methylation. Consistent with this, we find that Set2 has a short half-life and is co-regulated, with RNAPII CTD phosphorylation levels, during logarithmic growth in yeast. To probe the functional consequence of uncoupling Set2-RNAPII regulation, we expressed a truncated and more stable form of Set2 that is capable of dimethylation but not trimethylation in vivo. Results of high throughput synthetic genetic analyses show that this Set2 variant has distinct genetics from either SET2 or set2Δ and is synthetically sick or lethal with a number of transcription elongation mutants. Collectively, these results provide molecular insight into the regulation of Set2 protein levels that influence H3K36 methylation states.

The N-terminal tails of histones are richly decorated with post-translational modifications that regulate a variety of DNA-templated processes through the recruitment of additional factors to chromatin. These factors interact with modifications such as acetylation, methylation, and phosphorylation through specific protein domains such as bromo-, chromo-, PHD, and TUDOR domains (1). Furthermore, these domains are usually found in the context of multiprotein complexes that perform diverse functions in chromatin remodeling, replication, transcription, and DNA repair, among others (2–4). Thus, there is great interest in both identifying biological important post-translational modifications on histones and understanding how these modifications function within chromatin.

One such post-translational modification is methylation of lysine 36 on histone H3. In yeast, H3K363 is modified by the methyltransferase Set2, which can add one, two, or three methyl groups to the e-amino group of the lysine side chain using S-adenosylmethionine as the methyl donor (5). Genome-wide chromatin immunoprecipitation studies have demonstrated that H3K36 methylation is generally associated with actively transcribed genes; however, dimethylation (H3K36me2) and trimethylation (H3K36me3) show different patterns of localization (6–8). H3K36me2 appears uniformly distributed across transcribed genes and is not correlated with transcription frequency, whereas H3K36me3 seems to be most prevalent at the 3’ end of coding regions and is positively correlated with the rate of transcription. These data suggest that the deposition of different levels of H3K36 methylation is controlled within the cell and that the different degrees of this modification likely direct distinct biological functions.

H3K36 methylation has been reported to be recognized by the Rpd3(S) deacetylase complex in Saccharomyces cerevisiae through the chromodomains of Eaf3 (a member of both the Rpd3(S) deacetylase and NuA4 histone acetyltransferase complexes) (9–11). Subsequent studies revealed that H3K36me2 is the preferred substrate for Rpd3(S) binding (12–14) and that a primary function for H3K36 methylation is to recruit the repressive activity of Rpd3(S) to genes to “reset” or reestablish

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1 These abbreviations used are: H3K36, lysine 36 on histone H3; CTD, C-terminal domain; 5-FOA, 5-fluoro-orotic acid; H3K36me2, H3K36 dimethylation; H3K36me3, H3K36 trimethylation; RNAPII, RNA polymerase II; SRI, Set2-Rpb1 interacting; PAF, RNA polymerase-associated factor.
chromatin structure between multiple rounds of transcription (15–17). Such resetting, or at least maintaining a more compact chromatin structure within gene bodies, also protects genes from spurious transcription from improper initiation sites (cryptic transcription) (15–18). To date, no separate function has been reported for H3K36me3. However, H3K36me3 can be recognized by both the chromodomain of Eaf3 and in vitro by the PHD domain of Nto1, a member of the NuA3 histone acetyltransferase complex, suggesting that H3K36me3 might have a positive role in transcription through the recruitment of histone acetyltransferase complexes (19). Consistent with this, NuA3 association to chromatin has been shown to be partially dependent on Set2 methylation (20).

Set2 interacts with the repeating C-terminal domain (CTD) of RNA polymerase II (RNAPII) (12, 21–25). In yeast, the CTD of RNAPII consists of 26 repeats of a heptapeptide repeat sequence (YSPTSPS) in which all three serine residues can be phosphorylated (26–28). Set2 preferentially recognizes the Ser2/Ser5-modified form of the CTD through a region at the C terminus of Set2 termed the SRI (Set2-Rpb1 interacting) domain (21, 29). Consistent with this, H3K36me3 is well correlated with Ser2- and Ser2/Ser5-modified forms of the CTD that occur within the bodies and 3′ ends of genes (6, 30–32).

We recently reported that deletion of the kinase responsible for Ser2 phosphorylation of the CTD, Ctk1, results not only in loss of H3K36me3 but also decreased Set2 protein levels (14). This finding suggested a novel mechanism for regulating histone methyltransferase activity. Here, we present a molecular basis for the regulation of Set2 protein levels. We find that events that alter or perturb Ser2 CTD phosphorylation levels result in reduced levels of Set2 protein and global loss of H3K36me3 without much change in H3K36me2. Furthermore, we demonstrate that expression of a stable truncation of Set2, incapable of H3K36me3, greatly compromises the growth of several transcription elongation mutants. Lastly, high throughput epistatic (E-MAP) analysis reveals different interactions for strains capable of catalyzing different levels of H3K36 methylation in vivo. Collectively, our data delineate a novel mechanism for Set2 regulation and support the model that different degrees of H3K36 methylation likely harbor distinct functions.

EXPERIMENTAL PROCEDURES

Yeast Strains and Plasmids—Strains used in this study are detailed in supplemental Table S1. Strains created for this study were constructed using heterologous gene replacement (33, 34) or through classical yeast mating in combination with tetrads dissection or high throughput strain construction techniques (35). Set2 expression plasmids (14, 21) were transformed using the lithium chloride technique (36), and shuffle strains (where important genes were genomically deleted but carried on a ura3-selectable plasmid) were maintained on selective media until the time of the appropriate experiment, and plasmid loss was instigated by plating on media containing 5-fluoroorotic acid (5-FOA) (37).

Western Blot Analysis—Yeast cultures were grown in either rich medium (YPD) or synthetic defined medium from a start- ing A600 of between 0.1 and 0.2 and grown to mid-log phase (A600 0.6–1.0) with the exception of cultures started from stationary phase. In this instance, 200-ml cultures were started from saturated cultures of yeast grown in synthetic defined medium for >7 days and inoculated at an initial A600 of 0.5. In all cases, five optical units of cells were harvested by centrifugation, and extracts were prepared using a protocol adapted from Ref. 38. Briefly, cell pellets were resuspended in 75 μl of SUMEB buffer (1% SDS, 8 m urea, 10 mM MOPS, pH 6.8, 10 mM EDTA, 0.01% bromphenol blue), mixed with 200 μl of glass beads, and vortexed for 3 min. To the beads was added 175 μl of additional buffer, and the total supernatant was collected and clarified to give a total volume of 250 μl of cell extract. Extracts were heated at 95 °C for 5 min and frozen at −80 °C until use. For Western blot analysis of histones, 5 μl of extract was separated by 17% SDS-PAGE. For Western analysis of Set2 and RNAPII, proteins were separated by 8% SDS-PAGE. Gels were transferred to PVDF for 90 min at 45 mA and dried in methanol to block according to the manufacturer’s instructions (Millipore-Immobilon-P). Dried membranes were rehydrated briefly in methanol and incubated with primary antibodies overnight at 4 °C in PBST (phosphate-buffered saline including 0.05% Tween 20) containing 5% blocking (Pierce). Antibodies were used at the following dilutions: anti-H3K36me2 (Active Motif), 1:1,000; anti-H3K36me3 (Abcam ab9050), 1:2,500; anti-H3 (Abcam ab9050), 1:2,500; anti-H3 1:5,000; anti-Set2 (1:8,000), anti-G6PDH (1:100,000), anti-Ser(P)2 CTD (gift of D. Eick.), 1:200. Western blots were visualized using HRP-conjugated secondary antibodies and ECL Plus chemiluminescence (GE Healthcare).

Protein Turnover Experiments—Turnover experiments were adapted from a protocol from Ref. 39. Briefly, 100-ml cultures of yeast in YPD were grown to an A600 of ~0.6, and cycloheximide was added to a final concentration of 200 μg/ml. Five optical units of cells were taken at various time points, and 1 ml of 10 m sodium azide and 1 ml of dimethyl sulfoxide were added. Cells were then immediately frozen in liquid nitrogen. Frozen samples were thawed simultaneously, and extracts were prepared using the SUMEB method described above.

Yeast Growth Assays—For phenotypic growth assays, yeast were grown overnight in appropriate media (YPD or synthetic complete medium with necessary additives). Saturated cultures were used to start fresh cultures in the same medium at an A600 of ~0.2. Cells were allowed to double at least two times, and cells were harvested and resuspended to an A600 of 1.0 in sterile water in a 96-well plate. Cells were 5-fold serially diluted and then spotted onto plates containing necessary selections using a 48-pin replicating tool. Plates were incubated at indicated temperatures and imaged daily.

Epistatic Miniarray Analysis—E-MAP analysis was carried out as described (40, 41). Growth rates were assessed by automated image analysis of colony size (42).

RESULTS

Set2 and H3K36 Methylation Are Regulated Differentially during Cell Growth—It has been well established that methylation by Set2 is correlated with hyperphosphorylation of the CTD of RNAPII. More than a decade ago, West and Corden showed that CTD phosphorylation was highly correlated with log phase growth and the diauxic shift in yeast (43). We therefore measured Set2 protein and histone modifications in differ-
ent stages of yeast growth. Starting from stationary phase cells, a 200-ml culture of yeast in YPD was inoculated at an initial \(A_{600}\) of 0.5, and time points/\(A_{600}\) measurements were taken at regular intervals (Fig. 1A). Extracts from each time point were separated by SDS-PAGE and probed for Set2 as well as RNAPII phosphorylation and histone modifications by Western blot analysis. As shown in Fig. 1B, and consistent with previous findings (43), RNAPII and CTD phosphorylation levels peaked at late log phase and during diauxic shift (\(t/\text{H}11005\) 6–8 h). Significantly, we found that Set2 protein levels also changed under these same conditions and correlated strongly with the occurrence of RNAPII CTD phosphorylation (Fig. 1B). In examination of H3K36 methylation levels, we observed that H3K36me2 was most abundant at \(t/\text{H}11005\) 0 and 2 h but decreased during mid- to late log phase (Fig. 1C). In contrast, H3K36me3 levels were lowest at \(t/\text{H}11005\) 0–4 h but increased during mid- to late log phase at the same time where Set2 levels and CTD phosphorylation levels increased (Fig. 1C). Although Set2 and CTD phosphorylation levels decreased during late log phase, H3K36me3 levels remained constant. These data uncover that Set2 and H3K36me2 and H3K36me3 are dynamically regulated and that Set2 proteins levels and the occurrence of H3K36me3 are tightly coupled to CTD phosphorylation.

**Set2 Regulation by RNA Polymerase CTD Phosphorylation**

**FIGURE 1.** Set2, H3K36 methylation, and RNAPII phosphorylation levels are dynamically regulated during cell growth. A, growth curve of wild-type BY4741 cells in complete synthetic defined medium with approximate growth stages. s.p., stationary phase; lag, lag phase; log, anaerobic fermentation of glucose; d.s., diauxic shift; and res., aerobic respiration of ethanol. B, Western blot analyses of extracts from the time course shown in A, indicating the presence of RNAPII as well as phosphorylated forms of the CTD and Set2. C, Western blot analyses from samples taken in A probing for the presence of various post-translational modifications (H3K36me2, H3K36me3, H3K4me3, and H3K27ac) on histone H3.

**FIGURE 2.** Set2 is a highly unstable protein. A, Western blot analysis of a time course experiment probing for Set2 protein levels in wild-type and proteasome-deficient (cim3-1) cells following addition of the translation inhibitor cycloheximide (200 \(\mu\)g/ml). B, quantification of Set2 protein levels as a function of time following cycloheximide treatment. Protein abundance was calculated by densitometry of Western blot analysis and plotted as the natural log of the intensity. C, schematic of Set2 domain structure and truncation mutants used in D. D, Western blot analyses showing the relative abundance of Set2 and the corresponding H3K36 methylation levels of several Set2 truncation mutants.

**Set2 Protein Levels Are Regulated in Vivo**—We previously noted that Set2 protein was lost in CTK1 delete cells (ctk1\(\Delta\)), and this was not the result of changes in SET2 gene expression (14, 25). These data, combined with our observation that Set2 protein levels correlate with RNAPII CTD phosphorylation levels, prompted us to explore the regulation of the Set2 protein further. Accordingly, we examined the abundance of Set2 protein upon treatment with the translational inhibitor cycloheximide (Fig. 2A). Results showed in a time course experiment that Set2 exhibited a rapid turnover with a half-life of \(\sim9\) min (Fig. 2, A and B). In contrast, performing the same experiment in a temperature-sensitive proteasome-deficient strain (cim3-3) at
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the nonpermissive temperature showed increased stability of Set2 (Fig. 2, A and B).

We next sought to identify the region responsible for regulating Set2 stability. Using strains harboring truncation mutants of Set2 (Fig. 2C), we were able to map the region responsible for Set2 instability to several hundred amino acids of unknown function between the Set2 domain and the SRI domain responsible for interaction with RNAPII (Fig. 2D). Interestingly, analysis of Set2 and histone methylation levels in these strains revealed unexpected behavior of the different Set2 truncations. Deleting solely the SRI domain of Set2 (Set2(1–618)) resulted in no significant change in the abundance of Set2 protein but also no observable H3K36me2 or H3K36me3 (Fig. 2D). However, expression of just the SET domain (1–261) of Set2 resulted in a marked increase in Set2 protein and H3K36me2 but no detectable H3K36me3 (Fig. 2D).

Set2 Protein Levels Are Decreased in Mutants That Alter RNAPII CTD Phosphorylation—Set2 is known to bind to the Ser2/Ser5-phosphorylated form of RNAPII. Because Ctk1 is the major kinase responsible for Ser2 phosphorylation, we next wanted to determine whether the lower Set2 protein levels found in ctk1Δ strains (Fig. 3A) were related to the ability of Set2 to bind to the CTD of RNAPII. We therefore examined the levels of Set2 protein in a number of yeast mutants known to alter the phosphorylation state of the CTD. Set2 protein levels were noticeably decreased in mutants of the PAF complex, which is known to play an important role in transcription elongation and affect CTD Ser2 phosphorylation (Fig. 3B) (44). Significantly, Set2 protein levels were only decreased in PAF mutants that also exhibited changes in CTD Ser2 phosphorylation, cdc73Δ, paf1Δ, and ctr9Δ. Decreased levels of Set2 correlated with decreased levels of H3K36me3 but not H3K36me2, consistent with the model that CTD association is necessary for H3K36me3 by Set2. We also note that overall histone H3 levels are reduced in both the ctr9Δ and paf1Δ strains, likely also resulting in the lower levels of even the cross-reacting band in these two strains (Fig. 3B). Furthermore, Set2 levels were decreased in a temperature-sensitive allele of Kin28, as well as mutant alleles of the phosphatases Ssu72 (ssu72Δ-2), Fcp1 (fcp1Δ-1), and the cyclin-like protein Bur2 (supplemental Fig. S1). Because many of the mutants we tested were enzymes themselves, we wanted to confirm that the phenotype of decreased Set2 protein was directly due to a loss changes in its ability to recognize the CTD rather than other potential activities of these enzymes. Accordingly, we looked at Set2 protein levels in cells expressing mutant forms of the RNAPII CTD itself (Fig. 3C). Each strain carries an RNAPII gene with a CTD comprising wild-type repeats (WT) or mutations at either Ser2 or Ser5 within the given number of repeats. Again, we observed a decrease in Set2 protein levels when a number of Ser2 or Ser5 residues within the CTD repeat were mutated to alanine. However, we saw a strong reduction in H3K36me3, which was accompanied by either no change or a slight increase in the levels of H3K36me2 in these cells.

Overexpression of More Stable Form of Set2(1–261) Is Lethal in Combination with Various Transcription Mutants—A well studied phenotype for set2Δ is the ability to bypass deletion of essential transcription factors such as members of the Bur1 complex and Spt16 (11, 45, 46). We therefore hypothesized that Set2 degradation might alter the levels of H3K36me2/me3 to facilitate transcription elongation during periods of transcriptional stress, when, for example, chromatin compaction by Rpd3(S) would inhibit rapid gene activation of stress response genes (47, 48). To examine this possibility, by way of using mutant forms of Set2 that have different behaviors in degradation and H3K36me, we expressed the different truncated forms of Set2 (see Fig. 2C) on plasmids in a bur2Δ sets2Δ double deletion strain harboring a wild-type copy of BLUR2 and then analyzed the growth of these mutants after selection for the loss of the BLUR2 plasmid on 5-FOA (Fig. 4A). As expected, deletion of SET2 bypasses the poor growth of bur2Δ strain. However, reintroduction of Set2(1–261) resulted in worse growth than either the SET2 control or vector-only controls. These results suggest that the inability to degrade Set2 and/or reduce H3K36me2 levels results in impaired growth due to inappropriate transcriptional repression through maintained recruitment of Rpd3(S). Accordingly, we next deleted RCO1 (rcolΔ), a unique subunit of the Rpd3(S) complex, to see whether Rpd3(S)
activity was necessary for this lethality. In fact, rco1Δ was able to rescue the lethality caused by expression of Set2(1–261) in the bur2 background (Fig. 4A).

High Throughput Genetic Analysis of Set2 Truncation Mutants—The truncated form of Set2(1–261) offered a unique tool to assess the biological consequence of having a stable form of catalytically active Set2, which is not able to catalyze H3K36me3. We therefore performed epistatic miniarray analysis against 1,534 yeast mutant strains (Fig. 4B and supplemental Table S1). Our analysis revealed several classes of interactions. For example Set2(1–261) and set2Δ exhibit similar negative (synthetically sick) interactions with members of the SWR1 and PAF complexes, whereas only Set2(1–261) results in synthetic sickness with members of the COMPASS complex (Fig. 4B). We were able to confirm several interactions by classical genetic analysis, most notably those with SWR1 (Fig. 4C). Five representative tetrads from crosses between set2 mutant constructs and a swr1Δ strain are shown in Fig. 4C. We found that Set2(1–261), but neither Set2-FLAG nor set2Δ was lethal in combination with swr1Δ (Fig. 4C and Table 1). Because the
SWR1 complex is primarily responsible for depositing the histone variant HTZ1 into nucleosomes, we also tested genetic interactions between our set2 mutant constructs and HTZ1 (Table 1). In this instance, both deletion of SET2 and expression of the Set2(1–261) construct resulted in synthetic lethality (supplemental Table S2).

DISCUSSION

Using a combination of genetics and biochemistry, we have uncovered a significant mechanism for the regulation of Set2 methyltransferase activity in yeast. The SRI domain is known to bind to the CTD of RNAPII when it is phosphorylated at both the Ser2 and Ser5 positions within the CTD repeats. It was shown previously that disruption of this interaction, by deleting the CTD kinase Ctk1, resulted in a loss of H3K36 methylation without affecting Set2 mRNA levels (25). Here, we show that loss of H3K36me3 results from changes in the stability of the Set2 enzyme itself, which we demonstrate is mediated via its interaction with the phosphorylated RNAPII CTD.

In support of a role for Set2-RNAPII interaction in governing Set2 stability, we found that mutations in enzymes that either act on the CTD (Ctk1, Kin28, Bur1, Ssu72, Fcp1), or mutations to the CTD sequence itself resulted in lower levels of Set2 protein and H3K36me3 (Fig. 3 and supplemental Fig. S1). Interestingly, we also found that deletion of CTK1 altered the protein levels of other factors that associate with the CTD, suggesting this mechanism may not be restricted to just Set2 (supplemental Fig. S2).

Set2 levels correlated strongly with changes in CTD phosphorylation as wild-type cells pass through log phase and into diauxic shift (t ~ 8–10 h) (Fig. 1, A and B). Interestingly, we observed H3K36me2 in whole cell extracts at time points where Set2 was barely detectable but H3K36me3 was not observed until log phase where Set2 and CTD Ser(P)2 levels peaked (Fig. 1). These data indicate that the low levels of transcription found in stationary phase are accompanied by low levels of Set2 and CTD phosphorylation, yet H3K36me2 levels are maintained. Not only are these results consistent with previous findings that the interaction of Set2 with the CTD is required for H3K36me3 (14), but they provide important insight into the dynamic regulation of H3K36me2 versus H3K36me3 and suggest that these two methyl marks have distinct functions in the cell.

Consistent with SRI regulating H3K36me3, several other transcription complexes or kinases have been shown to regulate H3K36 trimethylation, notably the PAF and BUR kinase complexes (46, 49). Prelich and co-workers demonstrated several years ago that deletion of several PAF complex members, cdc73Δ, ctr9Δ, or paf1Δ, resulted in a decrease in H3K36me3 (49). Similarly, Jaehning and co-workers showed that CTD Ser2 phosphorylation is reduced in these same three PAF mutants (44). Our data show that Set2 protein levels are also decreased in cdc73Δ, ctr9Δ, and paf1Δ strains (Fig. 3B). We take these data to conclude that decreased CTD phosphorylation in PAF mutants results in reduced Set2 protein levels which ultimately changes the balance of H3K36 methylation in the cell. Consistent with PAF effects of Ser2 phosphorylation, we also see a decrease of Set2 levels in BUR2 deletions (data not shown). Given that Bur1/Bur2 has recently been demonstrated to function on both Spt5 (DSIF) and the CTD (50), it is plausible that Bur1 also affects H3K36me3/Set2

### Table 1

| Mutant/Tetrad analysis of set2 mutant strains with htz1Δ and swr1Δ | | |
|--------------------------------------------------|------------------|------------------|
| Mutant | Tetrads | Viable double mutant spores |
|--------|---------|-----------------------------|
| swr1Δx | | |
| SET2 | 12 | 100 (12/12) |
| SET2(1–261) | 14 | 0 (0/15) |
| set2Δ | 13 | 100 (15/15) |
| htz1Δx | | |
| SET2 | 20 | 100 (15/15) |
| SET2(1–261) | 14 | 0 (0/13) |
| set2Δ | 18 | 11 (2/19) |

**FIGURE 5. Proposed model of Set2 regulation.** Set2 is in equilibrium between a free state and association with the hyperphosphorylated form of RNAPII. The free form of Set2 is degraded in a proteasome-dependent manner with a half-life in yeast of ~9 min. In its free form, Set2 is capable of dimethylation of H3K36, which is sufficient for the recruitment of the repressive Rpd3(S) complex. Binding to RNAPII is required for H3K36me3, which may perform a positive function in transcription elongation.
levels by directly influencing Ser\textsuperscript{2} phosphorylation and/or through DSIF recruitment of the PAF complex.

In our analysis of Set2 regulation, we noted that Set2 protein turnover was reduced in a temperature-sensitive allele of a proteasome subunit. We take these data to suggest that Set2 is degraded via the proteasome (Fig. 2). Consistent with this notion, our data are in agreement with a high throughput study of protein stability where it was predicted that Set2 is an unstable protein (51). However, perhaps due to the rapid turnover of Set2 or the relatively small amounts of enzyme normally found in cells, we have been unable to identify a ubiquitin ligase, ubiquitin protease, or a site of ubiquitin attachment to Set2 (data not shown).

A significant question to then ask is why would cells need to dynamically regulate Set2 levels? In addition to providing a rapid means of regulating H3K36 di- and trimethylation during logarithmic cell growth, the control of Set2 levels might have a role in the response to cellular stress as mentioned above, where the transient changes in H3K36 methylation may alter recruitment of the repressor Rpd3(S) to sites of transcription to alleviate chromatin repression during transcription as proposed in Fig. 5. This would explain why deletion of SET2 is able to bypass a number of transcription elongation mutations including mutations in Spt6, Spt16, and Bur1/2 (11, 14, 45). In this scenario, removal of Set2 and hence both H3K36me2/me3, would result in more open chromatin structure and improved transcription of stress-response genes. In support of this idea, studies have shown that a wide number of stress-response genes are up-regulated in a SET2 deletion (47).

As mentioned above, the ability to rapidly alter Set2 levels to control the balance of H3K36me2 and H3K36me3 supports the idea that H3K36me2 and H3K36me3 have different biological functions. To explore this possibility further, we examined the genetic interactions of a Set2 fragment that shows an inability to catalyze H3K36me3 (Set2(1–261)). Importantly, this Set2 fragment is properly recruited to gene coding regions in a transcription-dependent manner (14). Our study compared the Set2(1–261) fragment against wild-type Set2, where all three methylation states at H3K36 are present, and a SET2 deletion, where no H3K36 methylation is observed. We found that Set2(1–261) shows distinct genetics from either deletion of SET2 or the wild-type case when crossed with mutations in other factors important for chromatin structure and/or transcription elongation. This is particularly significant because Set2(1–261) exhibits partial function. This could be attributed to an imbalance in the levels of H3K36 methylation and supports that Set2 has both positive and negative roles in regulating transcription. For example, a lack of H3K36me3 could lead to increased recruitment of repressive factors such as Rpd3(S) to H3K36me2, or it may result in the inability to recruit effector proteins that recognize H3K36me3. Interestingly, Eaf3, the factor that recognizes H3K36me in the Rpd3(S) complex, is also an integral member of the NuA4 histone acetyltransferase complex. Furthermore, evidence shows that a member of the NuA3 histone acetyltransferase complex also interacts with H3K36me3 (and this complex is known to associate with chromatin in a Set2-dependent manner) (19, 20). Perhaps then, H3K36me3 recruits activating complexes such as NuA3 and NuA4 to certain genes to facilitate transcription, and methylation at H3K36 can act as both an activating and repressive mark, as is seen with lysine methylation at H3K4 (52, 53). If this were the case, it is possible that, in response to changes in CTD phosphorylation caused by an environmental cue or transcriptional stress, degradation of Set2, rather than just disruption of the interaction between Set2 and the CTD is necessary to maintain some proper balance of H3K36me2/me3 in the cell. As Keogh et al. proposed several years ago, such a mechanism might also be used to attenuate regulation of certain genes or control transcription from normally silent cryptic promoters (11).

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As mentioned above, the ability to rapidly alter Set2 levels to control the balance of H3K36me2 and H3K36me3 supports the idea that H3K36me2 and H3K36me3 have different biological functions. To explore this possibility further, we examined the genetic interactions of a Set2 fragment that shows an inability to catalyze H3K36me3 (Set2(1–261)). Importantly, this Set2 fragment is properly recruited to gene coding regions in a transcription-dependent manner (14). Our study compared the Set2(1–261) fragment against wild-type Set2, where all three methylation states at H3K36 are present, and a SET2 deletion, where no H3K36 methylation is observed. We found that Set2(1–261) shows distinct genetics from either deletion of SET2 or the wild-type case when crossed with mutations in other factors important for chromatin structure and/or transcription elongation. This is particularly significant because Set2(1–261) exhibits partial function. This could be attributed to an imbalance in the levels of H3K36 methylation and supports that Set2 has both positive and negative roles in regulating transcription. For example, a lack of H3K36me3 could lead to increased recruitment of repressive factors such as Rpd3(S) to H3K36me2, or it may result in the inability to recruit effector proteins that recognize H3K36me3. Interestingly, Eaf3, the factor that recognizes H3K36me in the Rpd3(S) complex, is also an integral member of the NuA4 histone acetyltransferase complex. Furthermore, evidence shows that a member of the NuA3 histone acetyltransferase complex also interacts with H3K36me3 (and this complex is known to associate with chromatin in a Set2-dependent manner) (19, 20). Perhaps then, H3K36me3 recruits activating complexes such as NuA3 and NuA4 to certain genes to facilitate transcription, and methylation at H3K36 can act as both an activating and repressive mark, as is
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