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H3K4me3 Interactions with TAF3 Regulate Preinitiation Complex Assembly and Selective Gene Activation

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SUMMARY

Histone modifications regulate chromatin-dependent processes, yet the mechanisms by which they contribute to specific outcomes remain unclear. H3K4me3 is a prominent histone mark that is associated with active genes and promotes transcription through interactions with effector proteins that include initiation factor TFIID. We demonstrate that H3K4me3-TAF3 interactions direct global TFIID recruitment to active genes, some of which are p53 targets. Further analyses show that (1) H3K4me3 enhances p53-dependent transcription by stimulating preinitiation complex (PIC) formation; (2) H3K4me3, through TAF3 interactions, can act either independently or cooperatively with the TATA box to direct PIC formation and transcription; and (3) H3K4me3-TAF3/TFIID interactions regulate gene-selective functions of p53 in response to genotoxic stress. Our findings indicate a mechanism by which H3K4me3 directs PIC assembly for the rapid induction of specific p53 target genes.

INTRODUCTION

Histone modifications contribute to gene regulation through direct effects on chromatin structure and through the recruitment of effector proteins (Bannister and Kouzarides, 2011; Campos and Reinberg, 2009; Taverna et al., 2007). H3K4me3 is a histone mark associated with promoters and early-transcribed regions of active genes (Bernstein et al., 2005; Heintzman et al., 2007). Whereas studies in yeast indicate that H3K4me3 is a consequence of transcription, studies in metazoans favor more causal roles in transcription for this mark (reviewed in Vermeulen and Timmers, 2010). Nonetheless, the functions of H3K4me3 in initiation versus early elongation are not well established. However, in support of a function at initiation, a seminal study has demonstrated an interaction of H3K4me3 with the TAF3 subunit of TFIID (Vermeulen et al., 2007). TFIID is a general initiation factor (GTF) that, through core promoter recognition, plays a key role in preinitiation complex (PIC) assembly. TFIID is comprised of the TATA-binding polypeptide (TBP) and ~14 TBP-associated factors (TAFs) that interact with transcriptional activators, core promoter elements, and other GTFs (Burley and Roeder, 1996; Juven-Gershon et al., 2008; Thomas and Chiang, 2006). TAF3, while showing tissue-specific functions through interactions with the TBP-related factor TRF3 (Deato et al., 2008; Deato and Tjian, 2007; Hart et al., 2009), is nonetheless an integral component of TFIID (Gangloff et al., 2001). H3K4me3 interactions with TAF3 (Vermeulen et al., 2007) offer a mechanism for TFIID recruitment/stabilization that is complementary to mechanisms involving interactions of TBP and TAF with core promoter elements (Juven-Gershon et al., 2008), and these interactions have been suggested to be important specifically for TATA-less promoters (Vermeulen et al., 2007). However, this interesting possibility has not been tested and, importantly, there is no corresponding information on either global or gene-specific functions of H3K4me3-TAF3 interactions or on their underlying mechanism of action.

Following a global analysis of H3K4me3-TAF3 interactions, we focus on possible gene-specific effects of these interactions in the regulation of gene expression by the tumor suppressor p53. As a transcriptional activator, p53 strikes a balance between cell-cycle arrest and programmed cell death through the differential activation of cognate genes (reviewed in Vousden and Prives, 2009). The kinetics of p53 target gene activation vary, ranging from a rapid induction of cell-cycle control genes to a delayed induction of proapoptotic genes (Zhao et al., 2000). However, the mechanisms that regulate the distinct transcription programs remain to be fully elucidated. A potential role for H3K4me3 in p53-mediated gene activation is supported by a DNA damage-enhanced accumulation of H3K4me3 at the GADD45A promoter (An and Roeder, 2004). Also, ubiquitylated H2B, which directly stimulates hSET1-dependent H3K4 di- and trimethylation (Shilatifard, 2006), accumulates at the p21 promoter in response to DNA damage (Kim et al., 2009). However, the downstream mechanisms underlying H3K4me3 function in p53-dependent activation remain to be determined.
Figure 1. H3K4me3 Facilitates Global TAF3/TFIID Recruitment

ChiP-seq (A and B) and HIT-seq (C–E) analyses.

(A) Heat maps showing the distributions of RNAPII, TAF3, and H3K4me3 within 2 kb of the TSSs of 17,347 Refseq genes, rank-ordered by the size of RNAPII peaks.

(B) Venn diagram showing the overlap of genes occupied by TAF3 and enriched for H3K4me3 (p < 1.3 × 10^-5, false discovery rate <5%).

(C) HIT-seq signals for full-length WT and M880A mutant TAF3 and the corresponding isolated PHD TAF3 proteins at the indicated loci as compared to the distributions of H3K4me3, RNAPII, and expressed RNAs from published data sets (Shen et al., 2012).

(D) HIT-seq binding sites with the total integrations including: TAF3 WT (22,844), TAF3 M880A mutant (39,695), TAF3 PHD WT (26,087), and TAF3 PHD M880A mutant (35,363). The number of TAF3 peaks that overlap with H3K4me3 or the TSS (+/− 2 kb) are shown.

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Here, we show that H3K4me3, through interactions with TAF3, directs global TFIIID recruitment. In subsequent biochemical and cell-based analyses of p53 function, we show that H3K4me3-TAF3/TFIID interactions both enhance p53-dependent transcription by directly stimulating PIC formation and facilitate selective gene activation by p53 in response to DNA damage.

RESULTS

H3K4me3 Facilitates Global TAF3/TFIID Recruitment

To investigate a role for H3K4me3 in the genome-wide recruitment of TAF3, we employed chromatin immunoprecipitation sequencing (ChIP-seq) to examine the distribution of TAF3, RNAPII, and H3K4me3 in human colorectal carcinoma HCT116 cells (Bunz et al., 1998). Stringent TAF3 binding peaks (n = 11,765, p < 1 × 10⁻⁵, false discovery rate <5%) were identified with TAF3 bound at 46% of protein-coding gene promoters. Heat maps (Figure 1A) and composite profiles (Figure S1A available online) demonstrate a significant colocalization of the peak binding sites for TAF3 and RNAPII with the peak sites of H3K4me3 enrichment on active promoters. The vast majority (98%) of the TAF3-occupied regions were marked with significant H3K4me3 levels, and conversely, TAF3 was bound at most (81%) H3K4me3-enriched regions (Figure 1B). There also exists a genome-wide overlap in the length and height of the H3K4me3 and TAF3 peaks (Figure S1B). Thus, our ChIP-seq results establish a strong correlation between TAF3 binding sites with sites of H3K4me3 enrichment (Figures 1C and 1D), and a strong correlation in the sizes of the TAF3 binding sites across the genome. Further, H3K4me3 HIT-seq data to compare the binding of WT and mutant TAF3 near the TSSs of genes that were sorted into bins of 100 based on expression level (Figure 1E). The largest HIT-seq peaks for both intact TAF3 and the isolated PHD finger, as well as the greatest differences between the binding of the WT and M880A mutant TAF3 proteins and the corresponding WT and mutant PHD fingers, were identified at highly expressed genes (Figure 1E). Gene ontology enrichment (GO) analysis revealed that the TAF3 binding sites identified by HIT-seq were significantly enriched at genes involved in cell-cycle control and transcriptional regulation (data not shown). These findings suggest that direct interaction of TAF3 with H3K4me3 facilitates global TFIIID recruitment/stabilization at the promoters of actively transcribed genes. Given the propensity for PHD-dependent binding of TAF3/TFIID to cell-cycle control genes, we focused on p53-dependent transcription.

H3K4me3 Enhances PIC Formation and p53/p300-Dependent Activation

We employed a cell-free transcription system (Figure 2A; Figure S2B; An and Roeder, 2004) to determine how H3K4me3 and H3K4me3-TAF3/TFIID interactions affect transcription initiation. Chromatin was assembled by recombinant factors (Figure S2A) using a pG5ML template (Figure S2C) and reconstituted histone octamers containing either unmodified H3 or the H3K4me3 analog H3K4me3 (Figure S2D). Micrococcal nuclease (MNase) digestion showed that both templates are chromatinized equivalently (Figure 2B). Robust transcription from the unmodified template was dependent upon p53 and p300 (Figure 2C, lanes 1–4). In contrast, the H3K4me3-modified template showed a significant level of p53-dependent transcription in the absence of p300 (lane 6), likely due to endogenous cofactors in the nuclear extract (NE). This activity was further enhanced (~7-fold) by ectopic p300 to yield an absolute level of transcription ~4-fold higher than observed with the unmodified template (lane 8 versus lane 4), and this increase was observed over a range of chromatin assembly conditions (Figure S2E). Importantly, H3K4me3 contributed more significantly than H3Kac4me1 and H3Kac4me2 to stimulation of p53-dependent transcription (Figure S2F). These results demonstrate a direct effect of H3K4me3 on p53-dependent transcription.

To determine whether H3K4me3 enhances transcription by stimulating PIC formation, in vitro ChIP was performed using a purified system (Figure 2D; Figures S2G and S2H). Chromatin was incubated with or without p53, acetylated by p300, digested with MNase, and immunoprecipitated with antibodies against
Figure 2. H3K4me3 Enhances PIC Formation and p53-Dependent Transcription

(A) Schematic of the in vitro transcription assay.

(B) MNase digestion showing similar nucleosome spacing for chromatin assembled with unmodified H3 and H3Kc4me3-modified octamers.

(C) NE-based transcription assay with deletions and additions as indicated. Relative transcription levels were quantitated by a phosphoimager. After background subtraction, all values were normalized to lane 4. The bar graph represents the average of two independent experiments and the error bars denote the standard deviation. ND, nondetectable.

(D) Schematic of the in vitro ChIP assays.

(E) Schematic of the pG5ML template indicating the amplicons used for quantitative real-time PCR.

(F) ChIP on chromatin assembled with unmodified H3 or H3Kc4me3 was performed with IgG and the indicated antibodies. H3K4me3 levels are relative to H3 levels. An average of two independent ChIP experiments is shown with error bars representing the standard deviation.

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p53, H3, H3K4me3, TAF3, RNAPII, and TBP. The precipitated DNA was analyzed by quantitative real-time PCR using primer sets (Figure 2E) that scan both a promoter region (A), containing p53 binding sites, and a distal vector region (B). On the unmodified template, there was selective binding of p53 and selective p53-enhanced binding of TBP, TAF3 (TFIID), and RNAPII to the promoter region (Figure 2F). On the H3K4me3 template, we observed (1) comparable p53 binding to the promoter region, (2) a broad distribution of H3K4me3 across the template (A and B), and (3) a p53-dependent binding of TFIID and RNAPII to the promoter region. In the presence of p53, the H3K4me3 template showed a significant increase in promoter-assembled TFIID and RNAPII relative to the unmodified template (Figure 2F). This H3K4me3-dependent increase was detected exclusively at the promoter, despite the uniform distribution of H3K4me3 indicative of the importance of the TATA box and promoter-bound p53 in contributing to TFIID stabilization at the promoter. The lack of an effect of H3K4me3 on p53 binding also indicates that enhanced TFIID/RNAPII recruitment is not due to an H3K4me3-mediated stabilization of p53. Notably, TFIID showed comparable binding to H3 peptides with natural K4me3 versus the K4me3 analog (Figure S2I). Together, these results establish a role for H3K4me3 in directing enhanced PIC formation/stabilization.

To further investigate the impact of H3K4me3 on the rate of PIC formation and initiation, kinetic transcription analyses were performed with addition of sarkosyl at various times during PIC formation (Figure 2G). Whereas chromatin assembled with H3K4me3 supported the assembly of sarkosyl-resistant PICs as early as 30 min (Figure 2H, lane 9), unmodified chromatin was significantly less efficient in assembling a functional PIC, even after 2 hr (lanes 1–5) and despite the fact that this template shows the same sensitivity to sarkosyl (Figure S2J) and directs efficient transcription in the absence of sarkosyl (Figure 2H, lane 6). Because sarkosyl restricts transcription to a single round, a comparison of transcription with and without sarkosyl reveals that once formed, PICs bound to chromatin assembled with H3K4me3 or unmodified H3 undergo almost the same number of rounds of transcription (~7, lane 12 versus lane 11, versus ~5, lane 6 versus lane 5, respectively). These data are consistent with the in vitro ChIP results that show enhanced TFIID/RNAPII recruitment at the promoter of chromatin templates assembled with H3K4me3 versus unmodified H3 (Figure 2F). Also consistent with the sarkosyl results, B2 RNAs (Espinoza et al., 2004) had little to no effect on transcription from chromatin assembled with H3K4me3 as early as 30 min following NTP addition (Figure S2K, lane 9). By comparison, transcription from unmodified chromatin was inhibited in the presence of B2 RNA at all time points (Figure S2K, lanes 1–5). Taken together, these results establish a direct link between an increase in the rate of PIC formation and the stimulation of transcription by H3K4me3.

### H3K4me3 Facilitates Primary TFIID Recruitment and Cooperates with the TATA Box to Enhance PIC Assembly and Initiation

To elucidate the significance of H3K4me3 with respect to TATA-TBP/TFIID interactions, we assessed p53-dependent transcription on chromatin assembled with unmodified H3 or H3K4me3 and containing a wild-type (TATAA) or a mutated TATA box (TATGG). Transcription from the unmodified template was reduced 2.5-fold by the TATA mutation (Figure 3A, lane 4 versus lane 2). The enhanced transcription from H3K4me3-modified templates also was reduced 2.5-fold by the mutant TATA (Figure 3A, lane 8 versus lane 6), although the residual signal was nonetheless comparable to the signal from chromatin assembled with unmodified H3 and a wild-type TATA (Figure 3A, lane 8 versus lane 2). This indicates that H3K4me3 can partially compensate for the mutant TATA in mediating transcription. Chromatin containing both H3K4me3 and a wild-type TATA supported a significantly higher transcription level (Figure 3A, lane 6) than did chromatin that contained only H3K4me3 (Figure 3A, lane 8) or a wild-type TATA (Figure 3A, lane 2), which strongly suggests that the TATA box and H3K4me3 cooperate to enhance transcription.

To further investigate the role of H3K4me3 with respect to TFIID/TBP-TATA interactions, in vitro ChIP was performed. Mutation of the TATA box caused a significant decrease in TAF3/TFIID and RNAPII binding to both unmodified and H3K4me3 templates (Figure 3B). However, the levels of TAF3/TFIID and RNAPII bound to H3K4me3-modified chromatin containing a mutant TATA were equal to, or greater than, the levels bound to unmodified chromatin with a wild-type TATA (Figure 3B), showing that H3K4me3 can compensate for the mutant TATA in facilitating GTF/RNAPII recruitment. In addition, the amount of TFIID/RNAPII recruited to chromatin containing both H3K4me3 and a wild-type TATA was higher than that recruited to chromatin containing either H3K4me3 or a wild-type TATA (Figure 3B). These results parallel the transcription data (Figure 3A) and show that H3K4me3 can support a significant level of PIC assembly and transcription initiation in the presence of a mutant TATA box and that H3K4me3 can cooperate with TATA to enhance PIC assembly and transcription.

### TAF3/TFIID Binding to H3K4me3 Facilitates PIC Formation and Function on Chromatin Containing a Mutant TATA

To determine whether TAF3/TFIID-H3K4me3 interactions mediate the effects of H3K4me3 on p53-dependent transcription, we purified TAF3/TFIID complexes (Figure 4A; Figures S3A and S3B) containing either wild-type or M880A mutant TAF3. As predicted, the WT, but not the mutant, TAF3/TFIID complex binds selectively to K4 trimethylated H3 peptides relative to unmodified H3 peptides (Figure 4B). To assess the transcription activities of the TAF3/TFIID complexes, we

(G and H) Kinetic analyses of PIC formation were performed ±sarkosyl. Relative transcription levels were quantitated by a phosphoimager. After background subtraction, all values were normalized to lane 1. The bar graph represents the average of two independent experiments. Error bars denote the standard deviation. See also Figure S2.
immunodepleted endogenous TFIID from HeLa NE (Figure S3C, ΔIID). ΔIID NE was unable to support transcription from naked DNA (Figures S3E and S3F, lanes 3 and 4) or chromatin (Figures 4C and 4D, lanes 3 and 4), whereas mock-depleted NE supported transcription from naked DNA (data not shown) and chromatin (Figure S3D) to levels comparable to those observed with control NE. To examine the rescue potential of the TAF3/TFIID complexes, equivalent amounts of each complex were examined on chromatin containing a wild-type TATA. Both the WT and mutant TAF3/TFIID complexes are equally efficient in restoring transcription to levels comparable to control NE, both on naked DNA (Figure S3E, lanes 6–9 and 11–14 versus lane 2) and on chromatin assembled with unmodified H3 or H3K4me3 (Figure 4C, lanes 6–9 and 11–14 versus lane 2).

Next, we examined the rescue potential of the TAF3/TFIID complexes from chromatin that contains a mutant TATA. Both the WT and mutant TAF3/TFIID complexes are equally efficient in restoring transcription to levels comparable to control NE, both on naked DNA (Figure S3E, lanes 6–9 and 11–14 versus lane 2) and on chromatin assembled with unmodified H3 or H3K4me3 (Figure 4C, lanes 6–9 and 11–14 versus lane 2).

In vitro ChIP was performed to examine the requirement for H3K4me3-TAF3/TFIID interactions in PIC formation. On chromatin with a wild-type TATA, the WT and mutant TAF3/TFIID complexes were equivalent with respect to TAF3/TFIID and RNAPII recruitment to promoters (A) both on unmodified H3 chromatin and on H3K4me3 chromatin, although the overall levels of recruitment were higher on the H3K4me3 template (Figure 4E). On chromatin templates with a mutant TATA, the WT TAF3/TFIID complex supported low levels of TAF3 and RNAPII recruitment on the unmodified H3 template and higher levels of recruitment on the H3K4me3 template (Figure 4F). In contrast, mutant TAF3/TFIID supported significantly reduced levels of TAF3 and RNAPII recruitment to the H3K4me3 template compared to the WT TAF3/TFIID complex (Figure 4F). These findings are consistent with the transcription data (Figures 4C and 4D) and show that H3K4me3-TAF3 interactions are important for GTF/RNAPII recruitment and in promoting transcription from templates containing a mutant TATA.

**TAF3 Is an Essential p53 Coactivator**

We examined the significance of TAF3/TFIID and H3K4me3-TAF3/TFIID interactions in regulating the expression of endogenous p53 target genes using RNAi-mediated knockdown of TAF3 in HCT116 cells. Relative to a nontargeting control siRNA (control) (Figure 5A, left), two pairs of oligonucleotides directed against TAF3 mRNA (TAF3-1 and TAF3-2) significantly reduced both the uninduced and doxorubicin-induced level of TAF3.
(The doxorubicin-induced increase in TAF3 RNA levels is specific because the RNA levels of other TAF subunits were not affected [data not shown]. Whereas doxorubicin induced TAF3 RNA levels, no change was apparent at the protein level.) Both siRNAs also significantly decreased TAF3 protein levels (Figure 5A, right) without affecting the levels of other TFIIID subunits (Figure S4A) or the integrity of the TFIIID complex (Figure S4B). TAF3-1 (hereafter TAF3) siRNA was selected for further studies. Following siRNA transfection, cells were treated with doxorubicin and induction of p53 target gene expression was validated by qRT-PCR (Figure 6A) and immunoblot (Figure S4C).

Microarray analysis comparing the transcriptome of control and TAF3-depleted cells (+/− doxorubicin) showed that TAF3 depletion significantly alters (≥2-fold, p < 0.05) the expression of a small subset (1,234 of 22,000) of genes, which suggests that TAF3 selectively regulates transcription. As shown in Figure 5B (red), the vast majority of the genes (n = 1,115, 90%) affected by TAF3 depletion are genes that are significantly induced (≥2-fold) by doxorubicin in the control cells. Conversely, only a small percentage (n = 119, 9%) of genes were affected by TAF3 depletion in the absence of doxorubicin (Figure 5B, blue). Analysis of the doxorubicin-induced genes revealed 271 (22%) TAF3-repressed genes and 844 (68%) TAF3-activated genes (Figure 5B, red). Several key regulators of the DNA damage response were specifically identified as TAF3-activated genes, which is consistent with a role for TAF3 in mediating p53 transactivation. Overall our microarray data indicate that TAF3 is involved primarily in gene induction and, to a much lesser extent, in basal gene expression.

GO analysis of the TAF3-regulated genes did not reveal any significant functional clusters for the small group of genes whose expression changed in the absence of doxorubicin; however, several significant (p < 0.01) categories were identified in the larger set of doxorubicin-induced genes. Of the 271 TAF3-repressed genes, only 36 (13%) showed significant enrichment for GO categories, which included ubiquitin ligase activity, protein biosynthesis, and steroid biosynthesis (Figure 5C, left). GO annotations related to transcription and DNA damage response pathways were not significantly identified in the TAF3-repressed gene profile; however, these annotations were among the most prominent in the TAF3-activated data set. Specifically, 817 (97%) of the 844 TAF3-activated genes were classified into ten significant categories, the most prominent of which are related to cell-cycle control and transcription (including DNA metabolic processes, nucleotide binding, and transcriptional regulation) (Figure 5C, right). Although several proapoptotic genes were identified in our microarray data for TAF3-activated genes, these genes were not recognized as a functional cluster and are significantly underrepresented relative to the highly significant cell-cycle category. Together, the microarray and GO data suggest that the repression and activation mechanisms that require TAF3, under doxorubicin-induced conditions, are functionally unrelated, as reflected by the lack of overlap in the functional categories between the two data sets. By comparison, the high level of significance and functional coherence of GO terms associated with the TAF3-activated genes suggests that the regulation of p53 target gene expression by TAF3 is largely mediated through DNA damage-induced activation events and not through negative gene regulation.

Consistent with our microarray results, qRT-PCR analysis of representative genes revealed that TAF3 depletion had little to no effect on uninduced gene expression but caused 2.2- to 6-fold decreases in the doxorubicin-induced mRNA levels of the cell-cycle-regulated genes p21, BTG2, CDK5RAP1, and IGFI R, the autoregulatory gene MDM2, and the proapoptotic gene PUMA (Figure 5D)—all of which display early activation kinetics in response to p53 (Yu et al., 2001; Zhao et al., 2000). In contrast, delayed expression of the proapoptotic genes DR5, FAS, Apaf-1, sertad1, and Noxa, was not significantly affected by TAF3 depletion (Figure 5D). Consistent with these data, an immunoblot confirmed a decrease in the protein levels of p21 and BTG2, but not DR5 and FAS, as a result of TAF3 depletion (Figure S4D). The gene-selective effects of TAF3 are reproducible in osteosarcoma U2OS cells (Figure S4E), are not due to altered protein levels of p53 (Figure S4F), and are dependent on p53 because these effects were not identified in the p53-knockout HCT116 cell line (data not shown). Moreover, core promoter sequence analysis of the TAF3-dependent genes showed that TAF3 is required for the regulation of both TATA-less and TATA-containing genes (Figure 5D). Together, our global expression profiles parallel our gene-specific analysis, demonstrating that TAF3 selectively regulates rapidly induced p53 target genes.

Binding of TFII D/TAF3 and RNAPII Correlates with H3K4me3 Enrichment at the Promoters of Rapidly Induced p53 Target Genes

To assess a direct role for TAF3 in the regulation of selective p53 target genes, ChIP-quantitative-real-time-PCR was performed in HCT116 cells (+/− doxorubicin). Immunoprecipitated DNA was analyzed with primer sets (Figure 6B) specific to the promoter region (A) and a distal control region (B) of the early-response cell-cycle genes p21 and BTG2 and of the delayed-response proapoptotic genes DR5 and FAS. ChIP-quantitative-real-time-PCR revealed preassociated TFII D and RNAPII before doxorubicin treatment, and substantial doxorubicin-induced increases in these factors at the promoters, but not at the downstream regions, of all four genes (Figure 6B; Figure S5). During DNA-damage-induced activation, the levels of TAF3 recruited to the promoters of p21 and BTG2 increased rapidly and peaked at 4 hr, which correlates with the TBP and RNAPII binding profiles. By comparison, TAF3, TBP, and RNAPII recruitment at the DR5 and FAS promoters was significantly more delayed, with peak binding at 8 and 24 hr (Figure 6B; Figure S5). In addition, the peak levels of TFII D and RNAPII were comparable at the p21 and BTG2 promoters but significantly (3- to 7-fold; note the different scales) higher than the peak levels at the DR5 and FAS promoters. Thus, the temporal binding profiles of TFII D and RNAPII correlate strongly with the rapid versus delayed activation kinetics of the cell-cycle and proapoptotic genes, respectively (Figure 6A).

To determine whether TAF3 contributes to the DNA damage-induced recruitment of RNAPII, ChIP was performed in doxorubicin-treated HCT116 cells that had been transfected with control or TAF3 siRNA. TAF3 binding in the TAF3-depleted cells...
Figure 4. Functional Analysis of the TAF3/TFIID Complexes

(A) SDS/PAGE-silver stain analysis of the purified TAF3/TFIID complexes.

(B) Wild-type and mutant TAF3/TFIID complexes were analyzed for binding to H3K4-methylated peptides, and bound proteins were analyzed by immunoblots with indicated antibodies.

(C and D) In vitro transcription assays performed with ΔIID NE on chromatin templates assembled with H3 or H3K4me3-modified octamers and containing a wild-type (WT) TATA (C) or a mutant (M) TATA box (D). These assays were performed in the presence or absence of p53 and the wild-type or mutant TAF3/TFIID complexes. The amount of TFIID added in lanes 8 and 13 is comparable to the amount of endogenous TFIID in the NE in lane 2. Relative transcription levels were quantitated by a phosphoimager. After background subtraction, all values in (C) and (D) were normalized to lane 2 in the upper part of (C). Note that the assays in (C) and (D) were performed and analyzed in parallel such that the absolute levels of transcription are directly comparable. The bar graphs represent the average of two independent experiments. Error bars denote the standard deviation. ND, nondetectable.

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is reduced significantly and comparably (~65%–76%) at the promoter regions of *p21*, *BTG2*, *DR5*, and *FAS* (Figure 6C). RNAPII binding to the *p21* and *BTG2* promoters was significantly reduced (by 80% and 50%, respectively) in TAF3-depleted versus control cells; however, the levels of RNAPII remained largely unaffected at the promoters of *DR5* and *FAS* (Figure 6C). TBP recruitment to the promoters of all four genes was largely unaffected by TAF3 knockdown (Figure 6C). These results validate the specificity of our TAF3 antibody in ChIP and demonstrate that TAF3 directly impacts RNAPII recruitment to effect the activation of rapidly induced p53 target genes.

Having demonstrated that H3K4me3 facilitates global TFIID recruitment to promoters, particularly of cell-cycle control genes, and that H3K4me3-TAF3/TFIID interactions regulate p53-dependent activation in vitro, we next investigated whether H3K4me3 enrichment overlaps with TAF3/TFIID recruitment at the promoters of select p53 target genes. ChIP revealed pre-associated H3K4me3 prior to doxorubicin treatment and a substantial enhancement in H3K4me3 in response to DNA damage (Figure 6D), which parallels the temporal changes in TAF3, TBP, and RNAPII binding at the promoter regions of *p21* and *BTG2* (Figure 6B; Figure S5). In contrast, there was no parallel enrichment of H3K4me3 and TAF3/TBP (TFIID) at the promoters of *DR5* and *FAS*. Specifically, H3K4me3 rapidly increased at the *DR5* promoter in response to DNA damage (Figure 6D), whereas there was little to no change in H3K4me3 before or after doxorubicin treatment at the *FAS* promoter (Figure 6D). The doxorubicin-induced levels of H3K4me3 are significantly higher at the *p21* and *BTG2* compared to the *DR5* and *FAS* promoters (Figure 6D), which is consistent with our TAF3, TBP, and RNAPII ChIP results (Figure 6B; Figure S5). Our ChIP studies thus demonstrate a role for TAF3 in the regulation of selective p53 target genes. Furthermore, the strong overlap in TAF3 binding and H3K4me3 enrichment at the TAF3-dependent genes suggests that H3K4me3-TAF3/TFIID interactions are important in regulating the expression of select p53 target genes.

The Binding of TAF3 to H3K4me3 Regulates the Expression of Select p53 Targets

To investigate a direct role for H3K4me3-TAF3/TFIID interactions in the expression of select p53 target genes, we examined (1) the effects of siRNA-mediated knockdown of WDR5, which reduces global levels of H3K4me3 (Dou et al., 2006; Steward et al., 2006), and (2) the relative ability of siRNA-resistant wild-type TAF3 and the M880A TAF3 mutant to rescue p53 target gene expression in TAF3 siRNA-transfected cells. qRT-PCR revealed that WDR5 depletion significantly reduced H3K4me3 (Figure 7A) and the doxorubicin-induced levels of *p21* and *BTG2* (3- and 8-fold, respectively) but had minimal (2- and 1.4-fold, respectively) effects on the doxorubicin-induced levels of *DR5* and *FAS* (Figure 7B). H3K4me3 also appears to repress uninduced p53 target gene expression, as revealed by the 2- to 3-fold increase in basal levels of *p21*, *BTG2*, *DR5*, and *FAS* (Figure 7B, Doxo 0 hr) following WDR5 knockdown. However, this effect is TAF3 independent because TAF3 depletion had little to no effect on uninduced p53 target gene expression (Figure 5D).

Under uninduced conditions, WDR5 knockdown resulted in an ~80%–90% loss of H3K4me3 at the *p21*, *BTG2*, and *DR5* promoters and a slightly lower, but still significant (~55%), decrease at the *FAS* promoter (Figure 7C). This decrease in H3K4me3 resulted in a partial (40% and 55%, respectively) reduction of TAF3 binding at the *BTG2* and *p21* promoters but had little to no effect on TAF3 binding at the *FAS* promoter and led to a partial (~50%) increase of TAF3 binding at the *DR5* promoter (Figure 7D). These modest effects of H3K4me3 depletion on TAF3 recruitment are consistent with a role for H3K4me3 in the regulation of uninduced gene expression that is independent of TAF3 (Figure 5D). In doxorubicin-treated cells, WDR5 knockdown caused substantial losses of the doxorubicin-induced increase in TAF3 at the *p21* and *BTG2* promoters (70% and 80%, respectively) (Figure 7D) but had a much smaller effect on TAF3 binding at the *DR5* and *FAS* promoters (20% and little to no change, respectively) (Figure 7D). The significant effect of H3K4me3 on TAF3 recruitment in response to DNA damage (Figures 7C and 7D) is in agreement with the expression data implicating both TAF3 (Figure 5D) and H3K4me3 (Figure 7B) in the doxorubicin-induced activation of both *p21* and *BTG2*, but not *DR5* and *FAS*. These results demonstrate that H3K4me3 is required for TFIID recruitment to cell-cycle promoters in response to DNA damage.

To examine the importance of TAF3-H3K4me3 interactions, TAF3 siRNA-resistant constructs expressing wild-type or mutant M880A TAF3 were cotransfected with TAF3 siRNA into HCT116 cells. Both TAF3 expression constructs equivalently reconstituted TAF3 mRNA (Figure 7E, left) and protein (Figure 7E, right) levels. Notably, as shown in Figure 7F, wild-type TAF3 fully rescued the expression of the TAF3-dependent genes (*p21*, *MDM2*, *BTG2*, *PUMA*, *IGF1R*, and *CDK5RAP1*), whereas the TAF3 PHD point mutant M880A was significantly (1.8- to 3.3-fold) less efficient. These rescue experiments confirm the specificity of our TAF3 siRNA experiments (Figure 5D) and, taken together with our in vitro analyses, unequivocally demonstrate a direct requirement for the H3K4me3-TAF3/TFIID effector pathway in the regulation of rapidly induced p53 target genes.

DISCUSSION

H3K4me3 Modulates Global TFIID Recruitment through Direct Interactions with TAF3

We demonstrated that H3K4me3-TAF3 interactions direct global TAF3 recruitment by showing a strong genome-wide overlap of TAF3 and H3K4me3 and by a significant decrease in the recruitment of the TAF3 M880A PHD mutant relative to WT TAF3. However, whereas recruitment of TAF3, through H3K4me3 interactions, to the promoters of constitutively expressed genes (E and F) in vitro ChIP analysis on chromatin templates assembled with H3 or H3K4me3-modified octamers and containing a wild-type (E) or mutant (F) TATA box. The amplicons used for real-time PCR are shown in the schematic (Figures 2E and 3B). An average of two independent ChIP experiments is shown with error bars denoting the standard deviation. See also Figure S3.
Figure 5. TAF3 Functions as an Essential Coactivator for p53

(A) HCT116 cells were transfected with nontargeting control (control) or TAF3 siRNAs (TAF3-1 and TAF3-2) and cultured with doxorubicin for 0 or 24 hr. qRT-PCR (left) and immunoblot (right) revealed significant reduction of TAF3 compared to transfection with the control siRNA. The expression levels determined after doxorubicin treatment are relative to the levels before doxorubicin. Error bars denote the standard deviation from duplicate reactions by real-time PCR.

(B) Microarray analysis of genes affected by TAF3 depletion in HCT116 cells. Genes were sorted before (left, blue, No Doxo) or after (right, red, Doxo 24 hr) doxorubicin treatment. Genes induced more than 2-fold after doxorubicin treatment in the control cells are represented (right, red). The numbers of affected gene probes (out of a total of 22,000) are shown with the indicated fold of down- or upregulation due to TAF3 siRNA-mediated depletion.

(C) GO analysis of the genes significantly affected by TAF3 depletion following 24 hr of doxorubicin treatment. The most highly represented categories are shown with ontology terms and the percentage of genes in each category.

(D) qRT-PCR analysis of p53 target genes in control and TAF3-depleted HCT116 cells both before (Doxo 0 hr) and after (24 hr) doxorubicin treatment. The expression levels determined after doxorubicin are relative to the levels before doxorubicin. Error bars denote the standard deviation from duplicate reactions by real-time PCR.

See also Figure S4.
Figure 6. TAF3 Binding Is Required for RNAPII Recruitment and Correlates with H3K4me3 Accumulation at the p21 and BTG2, but Not the DR5 and FAS, Promoters

(A) HCT116 (p53+/+ and p53−/−) cells were treated with doxorubicin (for 0, 2, 4, 8, 12, and 24 hr), and qRT-PCR was performed to measure the rapid p53-dependent induction of the cell-cycle genes p21 and BTG2 and delayed induction of the proapoptotic genes DR5 and FAS. The expression levels after doxorubicin treatment are provided relative to the levels before doxorubicin. Error bars denote the standard deviation from duplicate reactions by real-time PCR.
was widespread, TAF3 depletion had little effect on constitutive gene expression. Thus, despite the specific association of TAF3 with constitutive gene promoters, there was little to no change in constitutive gene expression following TAF3 knockdown. This finding is consistent with the results of a recent study of gene expression following TAF3 knockdown in embryonic stem (ES) cells (Liu et al., 2011). The results, in both cases, could reflect sufficiency of the low levels of TAF3 that remain after siRNA-mediated knockdown for steady-state gene expression levels. Another possibility is that the high levels of preassociated TAF3 prime corresponding genes and influence their expression only under certain conditions. In this regard, TAF3 could function by poising PICs for a rapid response to stress. This possibility is consistent with a report of PICs at p53 target genes prior to DNA damage (Espinosa et al., 2003) and is also supported by our finding that H3K4me3-TAF3/TFIID interactions contribute to TAF3 recruitment at the promoters of constitutive genes whose GO categories parallel the categories of TAF3-dependent genes following doxorubicin treatment. Thus, in addition to the significant genome-wide impact of H3K4me3-TAF3 interactions in mediating TFIID recruitment, our genome-wide and gene ontology analyses underscore the significance of H3K4me3-TAF3/TFIID interactions in the regulation of TFIID recruitment to p53-regulated cell-cycle control genes.

H3K4me3 Directly Enhances TFIID and RNAPII Recruitment and Initiation

Given the diverse factors that interact with H3K4me3 (reviewed in Chi et al., 2010; Vermeulen and Timmers, 2010), a number of functions for H3K4me3 in transcription or transcription-coupled processes may be envisioned. Roles for H3K4me3 in transcription initiation are supported by the direct interactions between H3K4me3 methyltransferases and transcriptional activators (reviewed in Ruthenburg et al., 2007a; Vermeulen and Timmers, 2010), the contribution of H3K4me3 to the recruitment of the chromatin remodeler CHD1 to the PIC (Lin et al., 2011), and the direct interaction of H3K4me3 with TAF3 (Vermeulen et al., 2007). Our studies provide additional insights into a role for H3K4me3-TAF3 interactions in initiation. First, genome-level ChIP experiments revealed the significance of H3K4me3-TAF3/TFIID interactions in global TFIID recruitment. Second, complementation assays in doxorubicin-induced cells expressing WT and mutant TAF3 highlighted the importance of H3K4me3-TAF3 interactions for the activation of a select group of p53 target genes. Third, in vitro ChIP assays demonstrated that TAF3-H3K4me3 interactions enhance TFIID recruitment to a model promoter, and kinetic analyses showed that H3K4me3 increased the rate of PIC formation rather than the rounds of transcription. In addition, H3K4me3-TAF3/TFIID interactions were sufficient to nucleate PIC assembly and potentiate transcriptional activation from a TATA-mutated template that was deficient in PIC assembly and activation in the absence of H3K4me3. Overall, our results provide strong evidence for the functional significance of H3K4me3 and for H3K4me3-TAF3/TFIID interactions in stimulating initiation through an effect on TFIID recruitment and PIC formation. However, our results do not rule out the possibility that H3K4me3, through interactions with other proteins, may also act at later steps of the transcription cycle.

H3K4me3 Regulation of TFIID Recruitment to TATA-less and TATA-Containing Promoters

The demonstration that only about 10% of mammalian promoters contain a classical TATA box suggested that other core promoter elements may direct transcription initiation (Carinci et al., 2006) and, indeed, initiator (INR) and downstream core promoter elements (e.g., DPE) were shown to facilitate TFIID recruitment to TATA-less promoters (reviewed in Juven-Gershon et al., 2008). We extend these findings by revealing that H3K4me3 can act in the absence of a functional TATA box to facilitate PIC formation and activation in vitro. The specificity of this H3K4me3 function, through an interaction with TAF3/TFIID, was demonstrated by showing that the wild-type, but not the M880A mutant, TAF3/TFIID complex facilitates TFIID and RNAPII recruitment and rescues transcription initiation from a TATA-mutated, H3Kc4me3-containing chromatin template.

Consistent with these in vitro results, TAF3 and H3K4me3 were shown to be required for the DNA damage-induced activation of endogenous p53 target genes BTG2, PUMA, and CDK5RAP1, all of which have TATA-less promoters. In addition, this finding supports our observation that H3K4me3 is required for TAF3/TFIID, and subsequently RNAPII recruitment to these TATA-less p53 target genes. siRNA-based TAF3 knockdown and complementation experiments further revealed that the DNA damage-induced expression of these TATA-less genes is rescued by wild-type, but not M880A mutant, TAF3. Collectively, our in vitro and in vivo findings define a mechanism by which H3K4me3-TAF3/TFIID interactions, perhaps acting in concert with other TFIID core promoter interactions, are sufficient to direct TFIID recruitment and subsequent transcription of both a TATA-mutated in vitro chromatin template and endogenous p53 target genes containing TATA-less promoters.

Our cell-based assays of doxorubicin-induced genes also showed cooperativity between H3K4me3 and the TATA box that enhances TFIID and RNAPII recruitment/stabilization and transcription. However, whereas siRNA rescue experiments demonstrated a direct contribution of H3K4me3-TAF3/TFIID interactions to the activation of TATA-containing p53 target genes in cells, we were unable to detect a significant contribution to p53-dependent transcription from a model TATA- and
Figure 7. TAF3 Interacts with H3K4me3 to Facilitate the Rapid Induction of Selective p53 Target Genes

(A) HCT116 cells were transfected with nontargeting control (control) or WDR5 siRNA and treated with doxorubicin for 0 or 24 hr. Immunoblot analysis of cell lysates with the indicated antibodies.

(B) qRT-PCR analysis of p53 target genes in HCT116 cells that were treated as described in (A). The expression levels determined after doxorubicin are relative to the levels before doxorubicin. Error bars denote the standard deviation from duplicate reactions by real-time PCR.

(C and D) ChIP analysis of HCT116 cells transfected with control or WDR5 siRNA and treated with doxorubicin for 0 or 8 hr. The levels of H3K4me3 are relative to H3 levels. Schematic showing the amplicons used for real-time PCR. An average of two independent ChIP experiments that are representative of at least three is shown with error bars denoting the standard deviation.

(E) qRT-PCR (left) and immunoblot (right) analysis of TAF3 in HCT116 cells that were transfected with control or TAF3 siRNA together with empty vector control or siRNA-resistant plasmids expressing wild-type or M880A mutant TAF3. The expression levels determined after doxorubicin are relative to the levels before doxorubicin, and error bars denote the standard deviation from duplicate reactions by real-time PCR.

(F) qRT-PCR analysis of p53 target gene expression in HCT116 cells treated as described in (E). Error bars denote the standard deviation from duplicate reactions by real-time PCR.

See also Figure S6.
INR-containing promoter in vitro. This inconsistency between the in vivo and in vitro results regarding the requirement for H3K4me3-TAF3/TFIID interactions likely reflects the use, in vitro, of a model p53-responsive promoter (with five promoter-proximal p53 sites and a very strong core promoter) and the lack of other natural constraints (limiting factors, alternative chromatin structures, etc.).

Our current studies provide insight into the mechanisms of TFIID recruitment by showing that the stability and/or specific recruitment of TFIID is enhanced by bivalent interactions involving H3K4me3 and the TATA box. Based on previously described TFIID core promoter interactions (reviewed in Juven-Gershon et al., 2008) and studies reported here, we support a model (Figure S6) for the functional interplay between histone modifications and core DNA sequences that emphasizes a link between the total number of TFIID interactions with the core promoter and the corresponding level of transcription initiation. This model is consistent with previous reports that have shown that multiple core promoter elements are involved in mediating cooperative GTF recruitment (Hsu et al., 2008; O’Shea-Greenfield and Smale, 1992; Smale and Kadonaga, 2003) and that multiple histone modifications facilitate the multivalent recruitment of transcription factors (Ruthenburg et al., 2007b; Vermeulen and Timmers, 2010).

**H3K4me3-TAF3/TFIID Interactions Enhance the Recruitment of TAF3 for the Expression of Select p53 Target Genes**

In response to stress, p53-dependent signaling pathways direct damaged cells to undergo growth arrest or apoptosis (Levine, 1997; Prives and Hall, 1999). Although several mechanisms have been implicated in gene-selective functions of p53 (reviewed in Espinosa et al., 2003; Oda et al., 2000; Sykes et al., 2006; Tang et al., 2006), the full spectrum of mechanisms that control gene activation by p53 and determine the appropriate response remain to be fully elucidated. Our studies reveal a transcriptional mechanism in which H3K4me3-TAF3 interactions are required to regulate the expression of select p53 target genes. Importantly, our microarray data identified a specific subset of genes affected by TAF3 knockdown, which supports a gene-specific, rather than a global, effect of TAF3 on gene expression. The vast majority of the DNA damage-induced genes that were affected by TAF3 depletion are key regulators of the cell-cycle pathway and are rapidly induced in response to DNA damage (Zhao et al., 2000). In contrast, proapoptotic genes that display delayed induction (Zhao et al., 2000), were significantly under-represented in our microarray data. Consistent with our global studies, qRT-PCR analysis of a subset of p53 target genes confirmed this gene-selective coactivator role for TAF3. Our cell-based findings are corroborated by our in vitro mechanistic studies showing the importance of H3K4me3 in coordinating rapid PIC formation and enhanced transcriptional activation.

Our studies also reveal that, in response to DNA damage, a significant increase in TAF3/TFIID recruitment correlates with a significant increase in H3K4me3 accumulation at the promoters of several cell-cycle genes. In contrast, this overlap was not identified at several proapoptotic gene promoters. Strikingly, the significant increase in H3K4me3 accumulation and TAF3 recruitment that was observed at the cell-cycle promoters also correlates strongly with the rapid increase (from a low basal level) in the mRNA levels of these genes. More specifically, either H3K4me3 depletion or the loss of H3K4me3-TAF3/TFIID interactions was found to disrupt TFIID and RNAPII recruitment to the cell-cycle promoters while having no effect on their recruitment to proapoptotic promoters. The recovery of cell-cycle gene expression by siRNA-resistant wild-type, but not M880A mutant, TAF3 in TAF3-depleted cells underscores the importance of H3K4me3-TAF3 interactions in regulating cell-cycle gene responses. Taken together, our studies reveal that TAF3, and associated H3K4me3 interactions, have a direct role in gene-specific regulation within the p53 network.

**EXPERIMENTAL PROCEDURES**

**RNA Interference Experiments**

HCT116 and U2OS cells were transfected with a nontargeting siRNA control, a human WDR5 siRNA SMART pool, or TAF3 siRNA duplexes (listed in Table S1) (all siRNAs are from Thermo Scientific). At 48 hr after transfection, 0.5 μM doxorubicin (Sigma) was added. siRNA rescue experiments were performed by cotransfecting TAF3 siRNA with pcDNA5/FRT/TO-TAF3 siRNA resistant plasmids using Dharmafect Duo.

**In Vitro ChIP**

ChIP assays were performed as described (Kundu et al., 2000). For details and primer sequences, see Extended Experimental Procedures and Table S2.

**RNA Purification and Quantitative Real-Time PCR**

Total RNA was prepared using the RNeasy kit (QIAGEN), reverse-transcribed, and amplified with SYBR Green PCR master mix (Applied Biosystems). Details and primer sequences can be found in Extended Experimental Procedures and Table S3.

**Gene Expression Microarray Analysis**

Standard methods for labeling and hybridization to the illumina HumanHT-12 v3 Beadchip were performed. For details, see Extended Experimental Procedures.

**In Vivo ChIP and ChIP-Seq**

For experimental details and primer sequences, see Extended Experimental Procedures and Table S4.

**Chromatin Assembly and Transcription Assays**

Methyl-lysine analogs were installed as described (Simon et al., 2007). Chromatin was assembled and transcription assays were performed as described (An and Roeder, 2004). See Extended Experimental Procedures for details.

**HIT-Seq**

For experimental details and primer sequences, see Extended Experimental Procedures and Table S6.

**ACCESSION NUMBERS**

The GEO accession number for the microarray, ChIP-seq, and HIT-seq data sets reported in this paper is GSE43542.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, six figures, and six tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.01.052.
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