A Mutation in Histone H2B Represents a New Class of Oncogenic Driver

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ABSTRACT

By examination of the cancer genomics database, we identified a new set of mutations in core histones that frequently recur in cancer patient samples and are predicted to disrupt nucleosome stability. In support of this idea, we characterized a glutamate to lysine mutation of histone H2B at amino acid 76 (H2B-E76K), found particularly in bladder and head and neck cancers, that disrupts the interaction between H2B and H4. Although H2B-E76K forms dimers with H2A, it does not form stable histone octamers with H3 and H4 in vitro, and when reconstituted with DNA forms unstable nucleosomes with increased sensitivity to nucleases. Expression of the equivalent H2B mutant in yeast restricted growth at high temperature and led to defective nucleosome-mediated gene repression. Significantly, H2B-E76K expression in the normal mammary epithelial cell line MCF10A increased cellular proliferation, cooperated with mutant PIK3CA to promote colony formation, and caused a significant drift in gene expression and fundamental changes in chromatin accessibility, particularly at gene regulatory elements. Taken together, these data demonstrate that mutations in the globular domains of core histones may give rise to an oncogenic program due to nucleosome dysfunction and deregulation of gene expression.

SIGNIFICANCE: Mutations in the core histones frequently occur in cancer and represent a new mechanism of epigenetic dysfunction that involves destabilization of the nucleosome, deregulation of chromatin accessibility, and alteration of gene expression to drive cellular transformation.

See related commentary by Sarthy and Henikoff, p. 1346.

INTRODUCTION

With the introduction of next-generation sequencing and the elucidation of now tens of thousands of cancer genomes, it has become evident that among the commonest class of genes recurrently mutated in human cancers are those encoding components of the gene-expression machinery, including transcription factors, DNA methyltransferases, histone methyltransferases and demethylases, histone acetyltransferases, components of the SWI/SNF chromatin remodeling system required for nucleosome movement, and components of the cohesin complex necessary for promoter/enhancer looping interactions (1). Collectively, these mutations may affect chromatin modification states and accessibility of DNA to transcription factors, DNA replication, and DNA-repair machinery. Epigenetic regulation of gene expression occurs by modifying the accessibility of DNA wrapped around the histone octamer consisting of two molecules each of histone H2A, H2B, H3, and H4 to sequence-specific transcription factors and RNA polymerase (2, 3). Formation of a nucleosomal array inhibits gene activity, and effective transcriptional activation requires the movement of nucleosomes at key sites in response to physiologic stimuli (4, 5). Histones are among the most highly conserved proteins across species, and the genes encoding the canonical histones are located in four clusters encompassing a total of 72 histone genes that include 16 genes for histone H2A, 22 genes for histone H2B, 14 genes for histone H3, 14 genes for histone H4, and 6 genes for histone H1 (6). The canonical histones are synthesized during S phase to package newly synthesized DNA. In addition, each core histone has specialized replication-independent variant genes dispersed throughout the genome that can differ significantly in sequence from their canonical counterparts and substitute for canonical histones in context-dependent instances to form alternative nucleosome conformations (7). These alternative nucleosome configurations provide functional heterogeneity to chromatin dynamic processes such as DNA repair or regulation of gene expression.

Chromatin packaging and gene expression are controlled in part by the nature of chemical modifications of histones, particularly on tail residues which are unstructured and do not directly interact with DNA, but rather serve as recognition sites for epigenetic writers of chromatin marks, erasers of such marks, and readers that bind to such marks to further modify chromatin. For example, the histone H3 lysine 27 trimethyl (H3K27me3) modification created by the PRC2/EZH2 complex is recognized by the PRC1 complex,
which contains chromatin-remodeling ATPases that actively condense and close chromatin (8). Mutation of H3K27 to methionine inhibits the activity of PRC2 and leads to global decrease of H3K27me3, aberrantly open chromatin, and dysregulation of gene expression that contributes to the pathogenesis of diffuse intrinsic pontine glioma in children (9–11). Missense mutations in H3 at amino-acid residues 34 and 36 are associated with gliomas, sarcomas, and head and neck cancers and lead to inhibition of methyltransferases that create H3K36 methyl marks, which are associated with open and transcribed chromatin (12–15). Hence, loss of the H3K36me2 or H3K36me3 may lead to aberrantly closed chromatin.

To gain a more complete understanding of histone mutations in cancer, we used the cBioPortal of cancer patient genomics data to identify mutations in the four core histones that recur with high frequency in cancer (16). In agreement with a recent report (17), we observed that the most common histone mutations in cancer were often found in the globular histone fold domains of canonical histones rather than the tail regions, and many of these mutations were located in discrete structural regions important for histone octamer and nucleosome integrity. To analyze the functional impact of histone mutations, we characterized the most frequent canonical histone mutation observed in cancer, a glutamate to lysine mutation of histone H3. This mutation occurs frequently in human cancers.

RESULTS

Missense Mutations at Specific Residues of Core Histones Frequently Occur in Cancer Patient Samples

We analyzed a curated set of 159 nonredundant cancer studies with a total of 41,738 sequenced patient samples using cBioPortal to identify mutations that frequently occur in genes encoding H2A, H2B, H3, and H4, in patient samples of all cancer types (16). Because there are multiple gene paralogs for each histone, we determined the cumulative number of patients with a mutation at each conserved amino acid residue across all canonical gene paralogs for each histone (Fig. 1A). Across the four core histones, ~90% of the mutations reported were missense due to a one base substitution with frameshift or indels being much less common. Generally, most residues of histones were reported to be mutated at least once, and on average each residue was reported to be mutated in 6 to 10 patients. Across all four histones, there were clear hotspots where a recurrent mutation was noted at a much higher frequency. A Z score was calculated for the number of patients with mutations at each amino acid, and a score of $Z > 2$ was considered significant.

Mutations in 15 canonical histone paralog genes encoding H2A were observed in 864 of 40,317 sequenced patients (2.1%) of all cancer types (Fig. 1A). The overall somatic mutation frequency in each individual H2A paralog ranged from 0.1% to 0.2%. When the H2A mutations were sorted by cancer type using studies with at least 50 cases, H2A mutations were commonly found in bladder cancer (10.7%), endometrial carcinomas (10.1%), head and neck carcinomas (8.2%), esophagogastric adenocarcinomas (7.8%), and cervical squamous cell carcinomas (7.2%). The most frequently found missense mutation in H2A was a glutamate to lysine mutation at amino-acid residue 121 (Fig. 1A; Supplementary Table S1). Mutation at E121 was the only mutation observed across all H2A histone cluster 1 paralog genes.

Mutations in 18 canonical H2B paralogs were observed in 1,224 of 40,317 sequenced patient samples (Fig. 1B; 2.8%). The mutation frequency in individual H2B paralog genes ranged from 0.2% to 0.3%. When sorted by cancer type using studies with at least 50 cases, the five cancers with the most frequently observed H2B mutations were endometrial carcinomas (13.8%), bladder urothelial carcinomas (13.4%), cervical squamous cell carcinomas (10.8%), head and neck squamous cell carcinomas (10.3%), and esophageal squamous cell carcinomas (9.5%). The most common missense mutation in H2B was a glutamate to lysine mutation at amino-acid residue 76, H2B-E76K, and this was also the most frequently observed canonical histone gene mutation across all cancer types (Fig. 1B; Supplementary Table S2).

Histone H3 mutations in 12 canonical paralog genes were observed in 1,533 of 40,317 cancer patient samples (3.8%), and the frequency of mutation in individual paralog H3 genes ranged from 0.1% to 0.5% (Fig. 1C; Supplementary Table S3). When sorted by cancer type using studies with at least 50 cases, the five cancers with the most frequently observed H3 mutations were high-grade glioma (19.3%), bladder cancer, not otherwise specified (13.7%), esophageal squamous cell carcinoma (13.7%), endometrial carcinoma (11.4%), and nonmelanoma skin cancers (9.2%). The most frequent amino-acid residue found mutated in the canonical H3 histone genes was the glutamate residue at position 105 (Fig. 1C; Supplementary Fig. S1). Mutations at these latter residues were well characterized to be significant in cancer, suggesting that other frequently occurring histone mutations identified by this study may be of similar importance. Along these lines, we also uncovered a recurrent change, A114G, in H3.3. The H3F3A gene that encodes H3.3 uses the sequence AAG to encode lysine at position 27 and is found frequently mutated to ATG (K→M) in cancers. Although more than half of the genes encoding canonical histone H3.1 and H3.2 use AAG to encode K27, the K27M missense mutation is not observed frequently among these proteins, suggesting that the significance of the H3K27M mutation may be specific to histone H3.3 function. Also of note, although few mutations have been cataloged for the centrosome-specific H3 histone CENPA, two mutations, R69C/L and R130Q/W/L, were among the most frequent (Supplementary Fig. S1; Supplementary Table S3).

Mutations in 14 canonical histone H4 paralogs were observed in 687 of 40,317 queried patients with cancer (1.7%), and the frequency of mutation in individual paralog H4 genes ranged from 0.1% to 0.2% (Fig. 1D; Supplementary Table S4). When sorted by cancer type using studies with at
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Figure 1. Survey of the most frequently found canonical histone mutations in cancer patient samples. A cross-cancer mutation summary was performed using the cBioPortal to search a total of 41,738 nonredundant patient samples across all cancer types. The number of patients reported to have a missense mutation at each amino-acid residue across histone paralogs was graphed. A red line denotes the average number of mutations for each histone across all paralogs, and blue shaded regions indicate the first two standard deviations from the average. Location of the cumulative missense mutations missense mutation at each amino-acid residue across histone paralogs was graphed. A red line denotes the average number of mutations for each histone using the cBioPortal to search a total of 41,738 nonredundant patient samples across all cancer types. The number of patients reported to have a missense mutation at each amino-acid residue across histone paralogs was graphed. A red line denotes the average number of mutations for each histone across all paralogs, and blue shaded regions indicate the first two standard deviations from the average. Location of the cumulative missense mutations across all cancer patient samples. The scores indicate the most frequently found Z,C,14 canonical H3 genes (E/F/G/H/I/J/K, HIST2H2BE/F, HIST3H2BB, D, HIST1H3A/B/C/D/E/F/G/H/I/J/K/L, HIST2H3D, HIST3H3HIST1H2AA/B/C/D/E/G/H/I/J/K/L/M, HIST2H2AB/C, HIST3H2A); 15 canonical H2A genes (B, across all paralogs, and blue shaded regions indicate the first two standard deviations from the average. Location of the cumulative missense mutations found in: A, 15 canonical H2A genes (HIST1H2AA/B/C/D/E/G/H/I/J/K/L/M/N/O, HIST2H2AB/C, HIST3H2BB); B, 18 canonical H2B genes (HIST1H2BA/B/C/D/E/F/G/H/I/J/K/L/M/N/O, HIST2H2AB/C, HIST3H2BB); C, 12 canonical H3 genes (HIST1H3A/B/C/D/E/F/G/H/I/J/K/L/M/N/O, HIST2H3D, HIST3H3); D, 14 canonical H4 genes (HIST1H4A/B/C/D/E/F/G/H/I/J/K/L/M/N/O, HIST2H4A, HIST4H4) across all cancer patient samples. The Z scores indicate the most frequently found mutations (Z > 2) in each histone. Amino-acid residues displayed in red were significantly more frequent in cBioPortal than in the general SNP database.

least 50 cases, the five cancer types with the most frequently observed H4 mutations were endometrial carcinomas (9%), bladder urothelial carcinomas (7.3%), head and neck squamous cell carcinomas (6.3%), esophageal adenocarcinomas (5.6%), and colorectal adenocarcinomas (5.3%). The most frequently observed mutation across H4 paralogs was the arginine at position 3 (Fig. 1D; Supplementary Table S4).

Missense mutations were found to occur beyond two standard deviations of the average frequency at only 18 amino-acid residues across all four core canonical histones, and patients usually displayed only one histone gene mutation (Fig. 1; Supplementary Tables S1–S4). We examined the Single Nucleotide Polymorphism database (dbSNP), the database of short genomic variation (18), to determine if some mutations could actually represent rare polymorphisms. This revealed that in general specific histone protein changes were much more common in the cancer genomics database than in the general population (Supplementary Fig. S2; Supplementary Tables S1–S4). Only H2A amino acids S1 and R35 had similar polymorphism frequencies between the general population and cancer patient databases, indicating they may not be pathogenic. By contrast, polymorphisms at H2B-E76 were found to be approximately 100-fold more common in patients with cancer than in the general population (Supplementary Fig. S2). Furthermore, the average variant allele frequency of these 18 mutations, including those at H2A S1 and R35, was in the 20% to 30% range in diploid tumor samples, suggesting that histone mutations are subclonal and acquired rather than germline mutation or polymorphisms (Supplementary Tables S1–S4).

A mutation in 1 of the 18 recurrently mutated histone residues was found in up to 8% of some solid tumors (Fig. 2A). Most of these mutations lie within the histone fold of each protein rather than the tail region and several map to locations important for nucleosome integrity, suggesting that nucleosome destabilization may play a role in cancer. For instance, E76 of H2B and R92 of H4 (the seventh most frequently mutated amino-acid residue in H4) are predicted to form salt bridges bolstered by hydrogen bonds with each
other at the H2B–H4 interface, and H4-D68 is also in this vicinity in the nucleosome crystal structure (Supplementary Fig. S3A). In addition, E105 of H3 interacts with G42 of H4 in the nucleosome (Supplementary Fig. S3B). Molecular dynamics simulations and a recently solved structure indicate that mutation of H2B at E76 disrupts the association of H2B with H4 (Supplementary Fig. S3C and S3D; ref. 19).

Missense mutations at H2B-E76, H4-D68, or H4-R92 are found in about 1% to 2% of cervical, bladder, and uterine cancer patient samples (Fig. 2B). Furthermore, these mutations are often found to occur together with common oncogenes in cancer patient samples. For instance, the most frequent histone mutation was at E76 of H2B, and mutations at this residue tend to co-occur with common oncogenes such as RAS, KDM6A, KMT2C, and TP53 at a rate greater than chance (Fig. 2C). Taken together, these findings suggest that mutations at any of these 18 residues may contribute to oncogenesis potentially by destabilizing the histone octamer and nucleosome structure in such a way as to favor the cancer phenotype without being lethal to the cell.

**Mutation of H2B at E76 Disrupts Histone Octamer Formation and Nucleosome Structure**

To test the effect of histone mutations on nucleosome structure, gene expression, and cellular phenotype, we chose to fully characterize the effect of H2B mutation at glutamate 76 (E76) because this mutation occurred at the highest frequency among cancer patient samples, and previous reports indicated that it destabilized nucleosome structure (19, 20). Alteration at H2B at residue E76 was rarely reported as a...
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The MNase susceptibility of E76K nucleosomes is distinct from nucleosomes formed only with tetrasomes. A time course by gel (left) and densitometry quantification (right) of intact nucleosomes following MNase nuclease (MNase) sensitivity assay performed on nucleosomes made with WT, E76Q, and E76K H2B mutants shows more rapid digestion of E76K nucleosomes containing H2B-E76K and E76Q have an altered migration pattern, intermediate between a tetrasome and a WT nucleosome.

C Micrococcal nuclease (MNase) sensitivity assay performed on nucleosomes made with WT, E76Q, and E76K H2B mutants shows more rapid digestion of E76K containing nucleosomes than those with WT-H2B. A time course by gel (left) and densitometry quantification (right) of intact nucleosomes following MNase treatment. D The MNase susceptibility of E76K nucleosomes is distinct from nucleosomes formed only with tetrasomes.

The E76K mutation in H2B destabilizes the histone octamer and fails to protect the nucleosome from nuclease treatment in vitro. A The H2B-E76K mutant was unable to form stable octamers in vitro. Recombinant human histones (H2A, H2B, H3, and H4) were mixed and histone octamers resolved from [H3–H4], tetramers, H2A–H2B dimers, and free histones by gel filtration chromatography. B Nucleosomes were reconstituted by mixing equimolar amounts of DNA (147 bp) and octamers or, in the case of E76K, of tetramers and dimers (1:2 molar ratio) and resolved by native PAGE. Nucleosomes containing H2B-E76K and E76Q have an altered migration pattern, intermediate between a tetrasome and a WT nucleosome. C Micrococcal nuclease (MNase) sensitivity assay performed on nucleosomes made with WT, E76Q, and E76K H2B mutants shows more rapid digestion of E76K containing nucleosomes than those with WT-H2B. A time course by gel (left) and densitometry quantification (right) of intact nucleosomes following MNase treatment. D The MNase susceptibility of E76K nucleosomes is distinct from nucleosomes formed only with tetrasomes.

polymorphism in dbSNP (18), and the average variant allele frequency of H2B-E76K in tumors was 0.23 ± 0.13, suggesting it is not a founding “driver” lesion but rather an acquired subclonal mutation that may increase cancer development (Supplementary Fig. S2).

To measure the extent to which the H2B-E76K mutation could affect nucleosome structure or stability, we reconstituted nucleosomes using recombinant histones. Gel filtration chromatography showed that wild-type (WT) H2B yielded mostly a peak of histone octamer and a small peak of H2A–H2B dimer when the dimer is added in 10% excess (Fig. 3A, left). By contrast, the H2B-E76K mutant failed to form histone octamers and two peaks were resolved, one consisting of H3–H4 tetramers and a second of H2A–H2B dimers (Fig. 3A, right). In addition, the H2B-E76Q mutant, which was less frequent than H2B-E76K but was recurrent in cancer, formed histone octamers less efficiently than WT, as a greater fraction of H2A–H2B dimers was observed than when WT-H2B was used (Fig. 3A, middle). Despite octamer instability, H2B-E76K and H2B-E76Q histones can be reconstituted into nucleosomes from H2A–H2B dimers and (H3–H4), tetramers with a strong positioning DNA sequence. However, nucleosomes containing H2B-E76K or H2B-E76Q displayed faster electrophoretic mobility in a native gel, suggesting a qualitative change in the complex. The DNA–protein complex formed by nucleosomes containing H2B-E76K was significantly more sensitive to micrococcal nuclease (MNase) digestion than nucleosomes reconstituted with WT-H2B or H2B-E76Q (Fig. 3B and C). However, DNA–protein complexes containing H2B-E76K nucleosomes were more resistant to MNase than H3–H4 tetrasomes (Fig. 3D). Collectively, these data suggest that H2B-E76K forms unstable nucleosomes with the full complement of histones, whereas H2B-E76Q more closely resembles WT histone H2B.

H2B Mutation Deregulates Growth and Gene Expression in Yeast

To begin to characterize the biological effects of the mutant histones, we expressed them in S. cerevisiae, which have only one gene for each of the core histones. In yeast, one of the two endogenous WT H2B alleles was replaced with mutant H2B. Conversion to H2B-E79K (equivalent to human E76K; ref. 21) or H2B-E79Q caused temperature sensitivity, whereas conversion to an alanine residue did not have an effect (Fig. 4A and B), suggesting that E76K or E76Q changes protein folding/octamer stability at higher temperatures. Chromatin...
from yeast heterozygous for E79K had a global change in accessibility as exhibited by increased sensitivity to MNase (Fig. 4C). In addition, H4 was more readily extracted from chromatin by salt from yeast expressing H2B-E79K (Fig. 4D), indicating that the nucleosomes in yeast with H2B-E79K have decreased stability.

To test if nucleosome instability may disrupt nucleosome-mediated gene repression, we examined the PHO5 gene, which is inactive when yeast are grown in complete media and induced when grown in phosphate-free media. Compared with WT yeast, cells expressing the E79K allele displayed a 4-fold increase in PHO5 expression under nonpermissive conditions, and yeast with the E79Q allele demonstrated a 2-fold increase (Fig. 4E). The expression of PHOS in yeast harboring E79K in nutrient-replete media was comparable with the level of expression of this gene in WT yeast in phosphate-free media, and when E79K or E79Q yeast was grown without phosphate, there was a further increase in gene expression (Fig. 4E). Using a nucleosome scanning assay, we mapped the presence of a well-positioned nucleosome present at the “−2” position relative to the transcription start site (TSS), which occludes critical transcription factor binding sites (22). The qPCR readout of this assay showed the presence of positioned nucleosomes at the −1, −2, and −3 positions in yeast with WT H2B but the loss of the −2 nucleosome in yeast expressing H2B-E79K (Fig. 4F). Thus, nucleosome instability and loss of nucleosome-mediated gene repression may cause aberrant gene expression and growth phenotype of cells harboring mutant H2B.

Expression of H2B-E76K in Mammary Epithelial Cells Alters Cellular Growth and Gene Expression

Because H2B-E76K was observed in a number of cancers, we sought to characterize its ability to confer oncogenic properties to a human cell line model. To measure the effects of H2B mutation in the absence of other oncogenic mutations,
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WT-H2B or H2B-E76K was expressed as either a FLAG epitope–tagged version or as an inducible GFP fusion in the immortalized but nontransformed mammary epithelial cell line MCF10A. Exogenously expressed H2B accounted for 5% to 10% of total H2B and was incorporated into chromatin as evidenced by coprecipitation with H4, presence in salt-extracted chromatin, or GFP signal in the chromosomes of dividing cells (Supplementary Fig. S4A–S4D and data not shown). To determine how H2B mutation might contribute to malignancy, we measured the growth of MCF10A cells stably transduced with WT-H2B or H2B-E76K. MCF10A cells were grown continuously, and cumulative cell counts were determined. After several weeks, cells expressing the mutant H2B displayed significant increased growth (Fig. 5A).

In order to characterize the pathways that may be altered by mutant histone, we performed expression profiling of MCF10A cells expressing either WT-H2B or H2B-E76K by RNA sequencing (RNA-seq). Relative to MCF10A cells expressing WT-H2B, expression of 247 genes was increased, whereas expression of 149 genes was decreased in cells expressing H2B-E76K (Fig. 5B). Gene ontology analysis (Enrichr, ref. 23; Revigo, ref. 24) revealed that genes upregulated by H2B-E76K tended to regulate differentiation, apoptosis, proliferation, migration, and cellular signaling (Fig. 5C). Genes decreased by expression of H2B-E76K were primarily involved in the regulation of cellular biosynthetic processes, response to growth factors and adhesion, mitochondrial membrane transport, and glucose homeostasis (Fig. 5D).

**Figure 5.** Expression of H2B-E76K alters growth properties and gene expression in human mammary epithelial cells. A, Proliferation of MCF10A transduced with lentivirus expressing either WT-H2B or H2B-E76K was measured every 3 to 4 days. The results from three different infections are shown. Linear regression analysis was used to calculate statistical significance. B, Heat maps depicting differential gene expression between MCF10A cells expressing either WT-H2B or H2B-E76K. C and D, Bubble plots of gene ontology analysis identify biological processes significantly different in cells expressing H2B-E76K compared with cells expressing WT-H2B. Bubble area is relative to the number of genes identified in each classification. Ontology of genes with increased (C) or decreased (D) expression in cells expressing H2B-E76K. E, MCF10A cells expressing WT-H2B or H2B-E76K and PIK3CA(H1047R) were grown in soft agar for 14 days, and colonies were counted.
To determine the relevance of the H2B mutation to cancer, we tested its ability to cooperate with mutant PIK3CA, which tends to co-occur with H2B-E76K in breast cancer (Fig. 2C). Expression of oncogenic PIK3CA-H1047R along with mutant but not WT H2B led to increased colony formation in soft agar (Fig. 5E; Supplementary Fig. S4E). Taken together, these results demonstrate that even a small fraction (<10%) of mutant histone causes altered cellular phenotypes that may collaborate with other oncogenes during tumorigenesis.

Expression of H2B-E76K Relaxes Chromatin Structure and Alters H2A–H2B Dimer Dynamics

We next tested the effect of H2B-E76K on the chromatin structure. Mass spectrometry analysis demonstrated that expression of mutant histone did not alter global histone methylation patterns, suggesting that the effect of H2B-E76K on gene expression was not due to altered histone post-translation modification states (Supplementary Fig. S5). However, expression of H2B-E76K in MCF10A cells significantly increased the sensitivity of genomic DNA to MNase, indicative of a relaxed chromatin structure (Fig. 6A). FRAP analysis of histone H2A-GFP dynamics in MCF10A cells expressing only H2A-GFP (n = 20 cells) or both H2A-GFP and a mutant H2B-E76K-mCherry fusion (n = 25 cells) with standard deviation envelopes. The inset demonstrates coexpression of H2A-GFP and E76K-mCherry. E, Representative prebleach, bleach, and postbleach images of nuclei expressing H2A-GFP in WT MCF10A cells or in cells coexpressing mCherry-tagged H2B-E76K. F, 100-nm particles injected into the nucleus had significantly increased mean square displacement (MSD) over time in MCF10A cells stably expressing exogenous H2B-E76K (red) compared with cells expressing WT-H2B (blue). N = 68 (WT) and N = 67 (E76K) cells were analyzed. Error bars, SEM.
particle tracking experiments. In MCF10A cells expressing H2B-E76K, nuclear nanoparticle travel was significantly increased compared with WT cells, indicating either an increase in active forces on the particles or a decrease in resistance to particle motion in the mutant nucleus (Fig. 6F; Supplementary Fig. S6C).

**The H2B-E76K Mutant Fundamentally Alters Chromatin Accessibility**

We performed the assay for transposase-accessible chromatin using sequencing (ATAC-seq) on MCF10A cells that had expressed WT or mutant H2B to directly measure the extent to which mutant H2B altered chromatin accessibility in MCF10A cells. ATAC-seq reads were categorized into common peaks present in both cell lines or new peaks that displayed significantly increased chromatin accessibility in H2B-E76K-expressing compared with WT-H2B-expressing cells (Fig. 7A). In addition, a small subset of peaks were observed to have increased signal in cells expressing WT-H2B compared with H2B-E76K (Supplementary Fig. S7A). In new open chromatin regions of H2B-E76K cells, the frequency of guanine-cytosine (GC)–rich (>60% GC content) peaks was decreased, and the frequency of adenine-thymine (AT)–rich (>60% AT content) peaks increased compared with the frequency of GC- or AT-rich peaks observed in the common peaks shared by WT-H2B– and H2B-E76K–expressing cells, suggesting nucleosome formation is preferentially decreased by H2B-E76K in AT-rich genomic regions (Supplementary Fig. S7B). When the distribution of ATAC-seq peaks in various chromatin states was analyzed (25), open chromatin regions in common to WT-H2B– and H2B-E76K–expressing MCF10A cells were predominantly found in promoter and enhancer regions, whereas more than 20% of new peaks caused by expression of mutant histone appeared in normally transcriptionally inactive chromatin regions such as heterochromatin (Fig. 7B). In addition, for those ATAC-seq peaks in common between WT and E76K cells, the peaks in E76K cells tended to have an average increased width of about 50 bp compared with the same peak in WT cells (Supplementary Fig. S7C). Expression of mutant H2B caused significantly increased chromatin accessibility in the promoter regions of more than 3,200 genes (Fig. 7C), and these genes also displayed increased gene expression, whereas genes without an increased ATAC-seq signal around the TSS had no significant difference in expression.
expression (Fig. 7D). For example, expression of H2B-E76K promoted both a significant increase in chromatin accessibility at the normally repressed promoter region of the HOXB6 gene in MCF10A cells and a corresponding significant increase in gene expression (Fig. 7E). Thus, histone mutations that affect nucleosome stability may significantly alter chromatin structure and cause aberrant gene expression.

**DISCUSSION**

Whole-genome sequencing of human patient samples has revealed that dysfunction of almost every mechanism governing chromatin structure and regulation of gene expression occurs in cancer (1). Mutations found in tumors include those that affect the function of transcription factors, chromatin-modifying enzymes, chromatin-remodeling factors, and mutations that disrupt the covalent modification of histone tail residues. Now, by examination of the eBiobrain patient genomics database, we have identified a new class of mutations in the core histones (H2A, H2B, H3, and H4) that frequently occur in cancer and may function to destabilize the histone octamer and nucleosome. Although some amino acid–residue changes in histones we identified in our survey of cancer genomes can be found as polymorphisms in the general population, for most sites the frequency of specific amino acid–residue changes was 10 to 100 times more common in the cancer cases than in normal populations, and many of the recurrent histone amino acid–residue changes we identified were not observed in databases of >200,000 genomes. This, plus the fact that histone mutations were found to have a variant allele frequency of 20% to 30%, suggested that these variations are acquired somatic mutations that may contribute to carcinogenesis and not passenger polymorphisms. Such oncohistone mutations may by themselves or in combination with other oncogenic drivers change the dynamics of chromatin access in a way that alters gene expression to a state that favors tumorigenesis. Thus, our data suggest that histones are directly affected in malignancy, and chromatin may assume a global pathogenic state.

Support for histone alterations as cancer drivers is growing, and the prevalence of histone mutations in cancer was recently noted by others (17). Mutations have also been reported in the H1 linker histone, which couples nucleosomes to facilitate chromatin compaction (26). H1 has diverse roles in gene expression, and loss of H1 has pronounced effects on the heterochromatin structure that alters gene expression, sensitivity to DNA damage, and embryonic development (27). Downregulation of histone H1.0 was proposed to alter gene expression in glioblastoma and cause more heterogeneity of gene expression in cell populations that could contribute to oncogenesis (28). In addition, aberrant histone gene expression has been reported to have an oncogenic role in cancer (29, 30).

Mutations in the core histones were previously reported among a compilation of other cancer mutations in tumors but not in the systematic manner presented here and not using such a large set of patient data (17, 19, 20, 30–32). Histones H2A and H2B were reported to be mutated in carcinosarcoma of the ovary and predicted to affect histone structure (32). Expression of these mutant histones in cells yielded biological effects, but no molecular correlates in chromatin configuration or gene expression were assayed. Notably, these mutations were not among the 18 recurrent mutations we found to be significantly elevated in this survey of cancer genomes.

Our analysis and that of Nacev and colleagues reveal that the most common histone mutations in human tumor samples tend to occur in the globular, histone fold domains of the core histone proteins rather than in N-terminal tail regions (17). Furthermore, among the most frequent mutations in the four core histones, two specific structural regions of the nucleosome were significantly affected. Missense mutations altering H2B-E76, H4-R92, or H4-D68 affect a region of the nucleosome important for H2B–H4 interaction, whereas missense mutations affecting H3-E105 and H4-G42 affect a region important for H3–H4 interaction (Supplementary Fig. S3A and S3B). Thus, we propose that core histone mutations that disrupt nucleosome integrity may represent a new class of driver mutations in cancer. To test this concept, we evaluated the H2B-E76K mutation both in vitro and in cell models. This mutation was the most common cancer-associated missense mutation in core histones and often coincides with gain-of-function mutations of oncoproteins, including RAS and PIK3CA (Fig. 2C and ref. 31). The subclonal nature of the H2B-E76K and other histone mutations suggests that the mutations may not be initial drivers of the cancer phenotype but instead contribute to cancer progression. In accordance with this idea, expression of mutant H2B in MCF10A cells caused a modest increase in cell growth and a shift of gene expression. However, when coexpressed with well-characterized driver mutations of PIK3CA, enhanced colony formation was noted, consistent with the idea that histone mutations may contribute to the progression of cancer.

Our results and those of others who crystallized the histone octamer in the presence of the H2B-E76K mutation indicate that H2B-E76K inhibits histone octamer formation by destabilizing the interaction of H2B with H4 (19). Furthermore, our results demonstrate that expression of even a limited amount of mutant histone led to widespread change in chromatin accessibility, nucleosome function, gene expression, and cell proliferation. However, the impact of histone core mutations such as H2B-E76K may be more far reaching; for instance, future studies must now include determining the impact of these mutations on histone deposition and chromatin structure during DNA replication, DNA damage response, and DNA repair. Importantly, our results indicate that H2B-E76K can cooperate with oncogenes such as PIK3CA that tend to co-occur in cancer to alter cellular growth, suggesting that loss of nucleosome integrity synergizes with activated oncogenic signaling to propel the cancer program. What chromatin structures and configurations and specific genes and pathways are affected by such collaboration in the pathogenesis or progression of cancer remains to be uncovered.

**METHODS**

**Cells Lines and Reagents**

MCF10A cells were purchased from ATCC and maintained in mammary epithelial cell growth basal medium (Lonza) supplemented with bovine pituitary extract, hydrocortisone, hEGF, insulin, and 100 ng/mL cholera toxin. The HIST1H2BD open reading frame was cloned into pCDH-EF1-MCS-Puro (SBI) and the E76K mutant
made using a QuikChange Site-Directed Mutagenesis Kit (Agilent). Lentivirus was packaged in 293T cells, and supernatant was used to generate MCF10A cells expressing either WT-H2B or H2B-E76K. In addition, WT-H2B or H2B-E76K were cloned into a custom pLenti-cumate-switch vector to create cumate-inducible GFP-tagged constructs. To induce expression, cells were grown in media containing 1x cumate (SB8). For generating dual expression H2A-GFP and E76K-H2B-mCherry cell lines, we custom generated TRE3G-H2A-mGFP-rTATA3-T2A-hygromycin and TRE3G-H2B-E76K-mCherry-rTATA3-T2A-elasticidin vectors by modifying the commercially available cumate-switch vector. To induce dual expression, the cells were induced with doxycycline. Western blotting was performed to evaluate protein expression using antibodies to H2B (cat. #12364, Cell Signaling Technology), H4 (cat. #D2 4X, Cell Signaling Technology), FLAG (cat. #F3165, Sigma), and alpha-tubulin (cat. #7291, Abcam). All cell lines generated were screened for Mycoplasma monthly and authenticated to match MCF10A cells once a year by short tandem repeat analysis (Genetica DNA Laboratories). Cells were used in experiments within 10 to 20 passages after thawing.

Structural Modeling and Molecular Dynamics Simulations

The atomic structure of the simulated system was taken from the crystal structure of a nucleosome complex (Protein Data Bank ID: 1AOI), and only the chains B and D, corresponding to the H4 and H2B units, were used in the following MD simulations. The E76K mutation was generated using a method implemented in VMD (33). The missing H atoms were added using the H++ package. The H4 and H2B chains were then solvated in a water box filled with TIP3P water molecules. Periodic boundary conditions were applied in the MD simulations, and the dimensions of the water box were set at 110, 98, and 78 Å in the x, y, and z directions, respectively. In addition, Na+ and Cl− ions were added to neutralize the system and make the concentration of Na+ 0.15 mol/L. The force-field parameters for the protein were obtained from the ff4SB force field (34). Molecular dynamics simulations were carried out using NAMD (35). A 10-ns molecular dynamics simulation at 310 K with an NPT ensemble was performed to equilibrate the system. In the production period, the system was simulated for 20 ns using the NPT ensemble. Pressure was maintained at 1 atm using the Langevin piston method with a piston time constant of 50 fs, and a piston temperature of 310 K (36). No atomic coordinates were performed to equilibrate the system. In the production period, the molecular dynamics simulation at 310 K with an NPT ensemble was simulated for 20 ns using the NPT ensemble. Pressure was maintained at 1 atm using the Langevin piston method with a piston period of 100 fs, a damping time constant of 50 fs, and a piston temperature of 310 K (36). No atomic coordinates were performed to equilibrate the system. In the production period, the molecular dynamics simulation at 310 K with an NPT ensemble was simulated for 20 ns using the NPT ensemble. Pressure was maintained at 1 atm using the Langevin piston method with a piston period of 100 fs, a damping time constant of 50 fs, and a piston temperature of 310 K (36). No atomic coordinates were

FRAP Assay

Five single imaging scans were acquired followed by an iterative bleaching (5 pulses of 200 ms each) using a spot of 1 μm in diameter without scanning. Z-stack images were then collected at 20-second intervals. For imaging, the laser power was attenuated to 0.6% of the bleach intensity (25 mW, 488 nm laser excitation). FRAP recovery curves were generated from background-subtracted images. Spontaneous photobleaching was corrected by tracking changes in background fluorescence at every time point. The level of autofluorescence in cells not expressing the fusion proteins was 1% or less than in expressing cells. All quantitative values represent averages from at least 10 cells from three independent experiments.

Micrococcal Nuclease Digestion Assay

Micrococcal nuclease (MNase) was purchased from Worthington. To assay for MNase accessibility, 2 × 106 cells in 1 well of a 6-well plate were washed with PBS and incubated in 1 mL of NP-40 buffer (10 mMol/mL Tris Cl pH 7.4, 10 mMol/L NaCl, 5 mMol/L MgCl2, 0.5% NP-40, 0.15 mMol/L spermine, 0.5 mMol/L spermidine) on ice for 5 minutes. Cells were washed with MNase buffer (10 mMol/L Tris Cl pH 7.4, 15 mMol/L NaCl, 60 mMol/L KCl, 0.15 mMol/L spermine, 0.5 mMol/L spermidine, 1 mMol/L CaCl2) twice, and 0.75 mL of MNase buffer containing 0.015 U Mnase was added for the indicated times. Reactions were stopped by adding equal volume 10% SDS, 0.5 mol/L EDTA. Samples were harvested by scraping cells and transferring to a 15-μL tube. Cells were incubated with proteinase K at 65°C overnight. Next, NaCl was added, and a phenol–chloroform extraction was performed. The nucleic acid fraction was incubated with RNase A at 37°C for 1 hour, and DNA was isolated by chloroform extraction and ethanol precipitation. The final DNA product was resuspended in 0.1X TE buffer, and Mnase sensitivity was measured by resolving 100 ng of each sample on a 2.5% agarose gel in TAE buffer.

Recombinant Nucleosome Assembly and Stability Assay

Recombinant human core histones (H2A, H2B, H3, and H4) were prepared and mixed at a 1:2:1:2:1:1 ratio in denaturing buffer and subsequently dialyzed into high-salt native buffer to promote protein refolding and histone octamer formation (39, 40). Histone octamers were resolved from (H3–H4), tetramers, H2A/H2B dimers, and free histones by gel filtration chromatography (Superdex 200 10/30 GL, GE Healthcare), and selected fractions were visualized by SDS-PAGE. Equimolar amounts of 601 Sequence DNA (147 bp) and octamers or crs, in the case of E76K, of tetramers and dimers (1:2 molar ratio) were mixed in high salt buffer, and then dialyzed against a salt gradient to reconstitute nucleosomes (41). Nucleosomes were then resolved by native PAGE. Mnase susceptibility assays were performed on nucleosomes made with H2B-WT, H2B-E76Q, and H2B-E76K H2B by mixing each nucleosome (0.4 pmol/mL) with 9.6 U of Mnase (Worthington) and incubating for indicated amount of time. Reactions were stopped with a 20 mMol/L EGTA + 20 mMol/L EDTA solution and resolved by native PAGE. Relative amounts of nucleosomes at each time point were measured by densitometry and plotted.

Nuclear Nanoparticle Tracking

MCF10A nuclei were microinjected with polyethylene glycol (PEG)–modified, 100-nm, red fluorescent, carboxylate-modified nanoparticles (Invitrogen). We followed established protocols to coat carboxylate-modified nanoparticles (Invitrogen) with amine-terminated PEG (42, 43). PEGylated particles (10 μL; 3.6 × 1010 particles/mL) were sonicated for 30 minutes before being loaded to Femtotop II (0.5 μm; Eppendorf). The particles were then microinjected into the nuclei using an Eppendorf InjectMan microjection system. Cells were incubated at least 6 hours before imaging. Time-lapse images of circulating particles embedded in the nucleus were acquired on a Nikon TE2000 inverted epifluorescent microscope with 60×, 1.49 NA oil immersion objective (Nikon) and CCD camera (CoolSNAP HQ; Photometrics). Cells were maintained at 37°C in 5% CO2. Tracking of particles was achieved using the TrackMate plugin of ImageJ (44). The trajectories of the particles were then used to calculate time-averaged mean-squared displacements using msdanaalyzer Matlab tool (45).

Next-Generation Sequencing and Analysis

For gene-expression analysis, cells were collected at approximately 50% to 70% confluence and stored in RNALater (Thermo Fisher Scientific) until RNA could be prepared using the Aurum total RNA isolation kit (Bio-Rad). RNA-seq libraries were prepared from 0.5 μg of total RNA and 1:100 ERCC spike in control using a KAPA RNA-seq kit with Ribozera (Roche). Sequencing was performed on a NextSeq 500 with a 2 × 150 bp read length kit to achieve approximately 30 to 40 million reads per sample (Illumina). Sequences were trimmed using Trimmomatic with quality control performed before and after trimming using FastQC. Input sequences were aligned to the
transcriptome using STAR, 2.5.2a, and differential gene expression was analyzed using RSEM v1.2.31. Differentially expressed genes in each contrast were defined as abs(log2(FC)) ≥ 1.0 and FDR-corrected P value ≤ 0.01. Gene ontology analysis of biological processes was performed using the Enrichr and REVIGO web tools (24, 46).

ATAC-seq was performed as described (47). Libraries were size selected by SPRIsel, and library QC was performed by measuring the ratio of GAPDH promoter to a genomic desert region by qPCR prior to sequencing using a NextSeq 500 (2 × 150 bp) to achieve approximately 60 million reads per sample. Short reads were trimmed using Trimmomatic (v 0.36), and QC on the original and trimmed reads was performed using FastQC (v 0.11.4) and MultiQC (v 1.1). The reads were aligned to the human genome (version GRCh38) using Bowtie 2 (v. 2.3.3). ATAC peak calling and differential peak analysis were performed using the MACCS2 (v. 2.0.10) “callpeak” command. Insert size distribution was analyzed with the Collect-InsertSizeMetrics tool (Picard v 2.9.2). Custom scripts were used to collect heat maps and enrichment plots. The MANorm script with custom modifications was used to compare differences in ATAC-seq peak size. The ChromHMM database was used to assess enrichment of open chromatin at broad classes of chromatin states (25). Sequencing data can be accessed through the sequence read archive.

**Proliferation and Cell Growth Assays**

A cumulative cell count method was used to measure cell proliferation differences between cultures expressing either WT or mutant histone. Briefly, 1.5 × 10^6 cells were plated into a 6-well dish. After 3 days of growth cells were harvested and counted, and 1.5 × 10^5 cells were replated. Experiments were done in triplicate, and population doubling was calculated using the formula n = 3.32*(log CC − log I) + X, where n is the final population doubling number at the end of a given subculture, CC is the cell count at that point, I is the number of cells used as inoculum to begin that subculture, and X is the doubling level of the inoculum used to initiate the subculture being quantitated (48). A linear regression analysis with GraphPad Prism 7.03 was used to measure the significance of proliferation differences. Colony formation assay was performed using a soft-agar assay. Briefly, 10,000 cells were embedded in growth medium containing 0.3% agar in each well of a 6-well dish. After 3 weeks, colonies were fixed and stained with crystal violet.

**Measurement of Growth and Nucleosome Occupancy in Yeast**

The yeast H2B gene was PCR amplified from BY4741 yeast and cloned into pRS316. Mutations were introduced using the QuickChange mutagenesis kit (Stratagene). The WT or mutant H2B was incorporated into BY4741 by integrative transformation of the SacI-, Xhol-digested plasmid. For rich-media growth, cells were grown in YPD. For phosphate-starvation growth, the cells were washed with PBS and transferred to complete synthetic dropout media without potassium phosphate for 3 hours. A nucleosome scanning assay was used to examine the nucleosome architecture of the yeast PHOS promoter (22). Briefly, yeast grown in rich media was converted to spheroplasts, chromatin was digested with MNase, mononucleosomal DNA was purified, and MNase protection of the PHOS promoter was determined by quantitative PCR as previously described (22).

**Disclosure of Potential Conflicts of Interest**

J.A. Oyer is principal scientist at Pfizer Inc. and senior scientist II at AbbVie Inc., and has ownership interest (including stock, patents, etc.) in Pfizer Inc. and AbbVie Inc. A.J. Ruthenberg is a consultant/advisory board member for Epicypher Inc. J.D. Licht is chief scientific officer at the Samuel Waxman Cancer Research Foundation and reports receiving a commercial research grant from Celgene. No potential conflicts of interest were disclosed by the other authors.

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**REFERENCES**


Histone Mutations in Cancer


A Mutation in Histone H2B Represents a New Class of Oncogenic Driver
