The histone variant H2A.Bbd is enriched at sites of DNA synthesis

Viola Sansoni1, Corella S. Casas-Delucchi2, Malini Rajan2, Andreas Schmidt1, Clemens Bönisch3, Andreas W. Thomae1, Martin S. Staeg4, Sandra B. Hake1,3, M. Cristina Cardoso2 and Axel Imhof1,3,*

1Munich Center of Integrated Protein Science, Ludwig Maximilians University of Munich, 80336 Munich, Germany, 2Technische Universität Darmstadt Schnittspahnstr. 10, 64287 Darmstadt, Germany, 3Adolf-Butenandt Institute, Ludwig Maximilians University of Munich, 80336 Munich, Germany and 4Department of Pediatrics, Martin Luther University Halle-Wittenberg, Ernst-Grube-Str. 40, 06097 Halle, Germany

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ABSTRACT

Histone variants play an important role in shaping the mammalian epigenome and their aberrant expression is frequently observed in several types of cancer. However, the mechanisms that mediate their function and the composition of the variant-containing chromatin are still largely unknown. A proteomic interrogation of chromatin containing the different H2A variants macroH2A.1.2, H2A.Bbd and H2A revealed a strikingly different protein composition. Gene ontology analysis reveals a strong enrichment of splicing factors as well as components of the mammalian replisome in H2A.Bbd-containing chromatin. We find H2A.Bbd localizing transiently to sites of DNA synthesis during S-phase and during DNA repair. Cells that express H2A.Bbd have a shortened S-phase and are more susceptible to DNA damage, two phenotypes that are also observed in human Hodgkin’s lymphoma cells that aberrantly express this variant. Based on our experiments we conclude that H2A.Bbd is targeted to newly synthesized DNA during replication and DNA repair. The transient incorporation of H2A.Bbd may be due to the intrinsic instability of nucleosomes carrying this variant or a faster chromatin loading. This potentially leads to a disturbance of the existing chromatin structure, which may have effects on cell cycle regulation and DNA damage sensitivity.

INTRODUCTION

Chromatin consists of an equal mass of DNA and histone proteins and is bound by a plethora of associated factors (1–6). The fundamental unit of chromatin is the nucleosome, a structure that contains 147 bp of DNA wrapped around a histone octamer made up by the four core histones H2A, H2B, H3 and H4. Individual nucleosomes can differ from each other by a multitude of post-translational modifications and by the incorporation of distinct histone variants. Whereas relatively few variants for H4 and H2B are known, most eukaryotes have several isoforms of H3 and H2A (7). Canonical histones, which constitute the majority of the histones in a cell, are incorporated during S-phase whereas most histone variants are assembled independently of DNA synthesis. Genome-wide association studies in various organisms demonstrate a non-random association of histone variants across the genome (8) suggesting a possible function in setting up defined chromatin structures.

The structure of chromatin can undergo massive and global rearrangements during cellular differentiation (9–12), which are often accompanied by a substantial difference in the chromatin composition. Changes in chromatin structure and nuclear morphology are frequently observed during oncogenic transformation and are often used as diagnostic markers. In this light it is noteworthy that the over-expression of specific histone variants has recently been shown to be a hallmark of several cancer types. The gene encoding the replication-independent H3.3 variant for example has been found to carry a driving mutation in paediatric glioblastoma (13,14). The two prevalent mutations result in a disturbed modification of H3K27 and/or H3K36 presumably causing a mis-regulation of epigenetic marks. The centromeric H3 variant CENP-A on the other hand...
is overexpressed in primary colorectal cancers, which leads to a mis-targeting of this variant (15) resulting in mitotic defects and an increase in aneuploidy, a hallmark of cancerous cells (16). Besides H3, several H2A variants have also been shown to be mis-regulated in cancer (17–20). The two most divergent H2A variants H2A.Bbd and macroH2A show a striking dichotomy when it comes to their involvement in cancer: whereas H2A.Bbd RNA is overexpressed in strongly proliferating Hodgkin’s lymphoma (HL) cells (17), the expression of macroH2A significantly suppresses melanoma progression (18). These findings suggest that the two variants may have opposing properties in regulating cellular physiology.

MacroH2A has been named after its large C-terminal macro domain (21). Initially it was shown to be enriched on the inactive female X-chromosome in mammals (22,23) but has later been shown to also localize to large autosomal domains (1–3,8,24–26). MacroH2A stabilizes nucleosome core particles (27) and interferes with the remodeling activity of the SW1/SNF complex (22,28). Consistent with a rather repressive function, macroH2A mostly localizes to heterochromatic regions and correlates with H3K27me3-rich regions (24,29,30). H2A.Bbd preferentially binds H4 hyperacetylated regions and was named Barr body-deficient according to its striking exclusion from the inactive female X-chromosome when expressed ectopically (17,31). H2A.Bbd-containing nucleosomes organize less DNA and are less stable than canonical nucleosomes (16,32). The mouse genome contains four genes encoding putative homologues of H2A.Bbd termed H2A.Lap1–4 (33,34). The H2A.Lap1 isoform is expressed in testis and brain and localizes to the transcription start sites of active genes in round spermatids where it can even localize to the inactive female X-chromosome (27,29,30). When ectopically expressed in HeLa cells, H2A.Bbd is enriched on active genes where it is involved in transcriptional activation and mRNA processing (29,30,36). This potential role of H2A.Bbd in transcriptional activation and splicing regulation may have important implications in HL cells that express increased levels of H2A.Bbd RNA is overexpressed in primary colorectal cancers, which leads to a mis-targeting of this variant (15) resulting in mitotic defects and an increase in aneuploidy, a hallmark of cancerous cells (16). Besides H3, several H2A variants have also been shown to be mis-regulated in cancer (17–20). The two most divergent H2A variants H2A.Bbd and macroH2A show a striking dichotomy when it comes to their involvement in cancer: whereas H2A.Bbd RNA is overexpressed in strongly proliferating Hodgkin’s lymphoma (HL) cells (17), the expression of macroH2A significantly suppresses melanoma progression (18). These findings suggest that the two variants may have opposing properties in regulating cellular physiology.

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Despite a large body of work with regard to the structure and localization of nucleosomes carrying these two histone variants, surprisingly little is known about their function under physiological or pathophysiological conditions. Mice lacking both macroH2A genes are viable and fertile (39–41) but their somatic cells are more susceptible to epigenetic remodeling compared to wild type suggesting that this histone variant acts as a barrier to induced pluripotency (42). Mice lacking the H2A.Bbd homologues H2A.lap1–4 have not been reported so far but the testsis-specific expression pattern suggests a role of this histone variant during male gametogenesis (35,43). In order to better understand the role of H2A.Bbd upon expression in somatic cells and its role during tumor progression, we ectopically expressed the H2A.Bbd variant in mouse embryonic fibroblasts (MEFs) and analyzed the composition of chromatin domains containing H2A.Bbd by comparing it to domains that either contained canonical H2A or the differentially localizing macroH2A.1.2 variant.

**MATERIALS AND METHODS**

**Cell culture and cell transfection**

Female MEF cells were grown under standard conditions in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum (Sigma) and penicillin/streptomycin (PAA). Human HL cell lines L-1236 (44), L-540 (45), L-428 (27,46), HDLM-2 (28,46) and KM-H2 (24,30,47) were obtained from the Deutsche Sammlung für Mikroorganismen und Zellkulturen (Braunschweig, Germany) HL cell lines and control Epstein-Barr virus-immortalised cell lines (LCL) were cultured in RPMI 1640/GlutaMAX (Gibco) supplemented with 10% fetal bovine serum (Sigma), betamercaptoethanol and penicillin/streptomycin (PAA). Peripheral blood mononuclear cells (PBMCs) from healthy donors and patients with HL were isolated as described (31,48) with informed consent and approval by the ethics committee of the Medical Faculty of the Martin Luther University Halle-Wittenberg. MEF cells were transfected with GFP-H2A.Bbd and GFP-H2A expression plasmids using FuGENE® HD transfection reagent (Roche) according to the manufacturer’s instructions. Stable cell lines were selected using G418 (PAA) at a concentration of 600 µg/ml. The GFP-macroH2A expression plasmid was transfected using the TransIT LT1 reagent (Mirus) according to the manufacturer’s instructions. For live cell imaging, MEF cells stably expressing H2A.Bbd were transfected with mRFP-PCNA (49) using Amaxa nucleofection as previously described (50).

**FACS sorting and FACS analysis**

GFP single cell sorting was carried out using a MoFlo cell sorter (Beckman Coulter). Single cell clone suspensions of GFP-H2A.Bbd, GFP-macroH2A and GFP-H2A MEFs were analyzed using a FACS Canto flow cytometer (Beckman Coulter). The GFP signal was detected in the Fluorescein Isothiocyanate (FITC) channel, and the data analysis to quantify the GFP positive cells was performed using FlowJo software.

**Chromatin purification**

Adherently growing MEFs were detached by trypsinization followed by centrifugation at 200 × g. Cells were washed twice with phosphate buffered saline (PBS) and nuclei were isolated by the addition of ice-cold PBS/0.3% Triton-X. After a 10 min incubation step at 4°C, nuclei were pelleted for 10 min at 1000 × g, washed once with PBS and resuspended in EX-100 (10 mM Hepes pH 7.6, 100 mM NaCl, 1.5 mM MgCl2, 0.5 mM EGTA, 10% v/v glycerol) freshly supplemented with protease inhibitors. Chromatin was solubilized by 20 min micrococcal nuclease (MNase) digestion at 26°C in the presence of 2 mM CaCl2. The reaction was stopped by adding 1/50th volume of 0.5 M EGTA. All further steps were carried out at 4°C. The first supernatant (S1) was recovered after centrifugation at 1000 × g and the pellet was resuspended in resuspension buffer (1× PBS, 150 mM NaCl, 2 mM EDTA, 0.1% Triton) and over-head rotated over night (o/n) to further extract solubilized chromatin fragments. The second supernatant (S2) was recov-
erated by a 30 min centrifugation, pooled with the first supernatant to yield the purified chromatin. A chromatin aliquot was used to check the MNase digestion by DNA purification and agarose gel electrophoresis to measure the size of chromatin fragments.

GFP pull-down

MNase-digested chromatin was precleared with 20 μl of a 1:1 (v/v) slurry of agarose beads in EX-100 (10 mM HEPES pH 7.6, 100 mM NaCl, 1.5 mM MgCl2, 0.5 mM EGTA, 10% v/v glycerol) for 1 h at 4°C. The supernatant was collected after 2 min of centrifugation at 2000 × g, 4°C. Twenty microliters of a 1:1 slurry (v/v) of GFP-binder (Chromotek) were equilibrated three times with EX-100 and then added to the supernatant. Immunoaffinity purification was performed for 2 h at 4°C. The beads were washed three times with resuspension buffer (1× PBS, 150 mM NaCl, 2 mM EDTA, 0.1% Triton) plus protease inhibitors and centrifuged at 4°C at 2000 × g. Fifteen microliters of 2× Lamelli buffer were added to the beads and boiled for 15 min at 95°C. Samples were run on a 15% SDS gel for 2 cm and stained with Coomassie Brilliant Blue G250 for 20 min. After destaining with 10% acetic acid, every lane was cut into eight slices and every slice was collected into a tube.

Trypsin digestion and mass spectrometry

Gel pieces were washed once with milliQ water and two times with 20 mM NH4HCO3. Destain of the gel pieces was preformed in 50% ACN, 50% NH4HCO3 at 37°C for 1 h. Samples were washed two times with 20 mM NH4HCO3 and dehydrated with 100% ACN (supplier and quality grade ultrapure). Reduction was performed in DTT 10 mM diethylthithiocarbamate (supplier and quality grade ultrapure). Reaction was stopped by adding 50 μl of 20 mM IAA. The samples were incubated overnight at 37°C. Before trypptic digestion, the samples were washed once with 20 mM NH4HCO3 and dehydrated with 100% ACN. 30–50 μl of trypsin (25 ng/μl) was added to the gel pieces in ice till the gel pieces hydrated completely. Not absorbed trypsin was removed and 50 μl of 20 mM NH4HCO3 was added. The samples were incubated overnight at 37°C. The supernatant containing peptides was collected and placed in a fresh vial. Gel pieces were incubated twice in 50% ACN, 0.25% TFA, 10 min at room temperature followed by two times 100% ACN, 10 min at room temperature. All supernatants were pooled, dried and then resuspended in 15 μl 0.1% FA. Fifty percent of the sample was injected in an Ultimate 3000 HPLC system (LC Packings Dionex). Samples were desalted on-line by a C18 trapping column (300 μm i.d. 5 mm, packed with C18 PepMapTM, 5 μm, 100 Å by Thermo-Fisher Scientific), and subsequently peptides were separated with a gradient from 5% to 60% acetonitrile in 0.1% formic acid over 40 min at 300 nl/min on an analytical column (75 μm i.d.×10 cm, packed with C18-PepMap, 3 μm, 100 Å). The eluent from the HPLC was directly electro-sprayed into the LTQ Orbitrap mass spectrometer (Thermo Fisher Scientific). The mass spectrometry (MS) instrument was operated in the data-dependent mode to automatically switch between full scan MS and MS/MS acquisition. Survey MS spectra (m/z 350–2000) were acquired in the Orbitrap with resolution 60 000 at m/z 400. Fragment ion spectra were recorded in the linear trap of the instrument. For all measurements with the Orbitrap detector, three lock-mass ions from ambient air (m/z = 371.10123, 445.12002, 519.13882) were used for internal calibration as described. The six most intense peptide ions with defined charge states of 2+ and higher were sequentially isolated in a 2 Da window around the precursor signal (window = 350–2000 m/z) to a target value of 10 000 and fragmented in the linear ion trap by collision-induced dissociation (CID). The ion selection threshold for triggering acquisition of an MSMS spectrum was 10 000 counts. Previously fragmented precursor signals were excluded from repeated fragmentation analysis for 30 s. Typical mass spectrometric conditions were: spray voltage, 1.4 kV; no sheath and auxiliary gas flow; heated capillary temperature, 200°C; normalized collision energy, 35% for CID in linear ion trap.

Mass spectrometry data analysis

For protein identification, the raw data were analyzed with the Andromeda algorithm of the MaxQuant protein analysis package (version 1.2.2.5) against the IPI mouse v.3.68. Fastai including reverse sequences and contaminants. The trypsin/P enzyme was selected, allowing for maximally two missed cleavages. Carbamidomethylation of C was set as fixed modification whereas methionine oxidation and protein N-acetylation were included as variable modifications. The mass tolerance of the initial search was 20 ppm; after recalibration, a 6 ppm mass error was applied for the main search. Fragment ions were searched with a mass offset of 0.5 Da using the six most intense signals per 100 Da. Searching for secondary peptide hits within already assigned MSMS spectra was enabled. The search results were filtered with a peptide and protein false discovery rate of 0.01 with a minimum peptide length of six amino acids. Protein identifications with at least one unique or razor peptide were accepted. For the quantification, the Intensity Based Absolute Quantification (iBAQ) values were calculated from peptide intensities and the protein sequence information (51) of unmodified, M/oXidated and acetylated peptide species with a minimum of two peptides per protein.

Statistical data analysis

Statistical analysis of protein enrichment was performed from the three affinity purification-MS replicates of the three GFP-H2A variants. First, all replicates were normalized using the median of abundant histone H4 peptides, including only peptides without modifications, less than two missed cleavages and excluding trypptic cleavage after Pro, as normalization factor. No-GFP data were not normalized, since detected H4 levels were too low. Second, normalized iBAQ values were log2 transformed and missing values were imputed from a Gaussian distribution that was shifted by a difference of about 2.5 and had a width of 0.3 using...
Figure 1. Female MEF single clones stably express the GFP-tagged histone variants. (A) FACS profile of female MEF cell populations derived from sorted single cell clones that express H2A.Bbd, macroH2A.1.2 and H2A tagged with GFP (solid line) in comparison to untransfected control female MEF cells (dotted and filled histogram). (B) Western blot analysis of GFP-tagged histone variant expression levels with antibody against GFP. Detection of histone H3 served as loading control. (C) Nuclear localization of the histone variants GFP-H2A.Bbd and macroH2A.1.2 in female MEFs stained with H3K27me3 (red). Arrows indicate the Barr body. Scale bar: 5 μm.

the Perseus software tool (53). The average difference of the GFP-H2A variants over cells not expressing GFP (no-GFP) was calculated and only proteins having a difference of more than 3.5 and quantitative information in at least two experiments were kept. To determine enrichment factors and p-values for enrichment, ANOVA was performed in DanteR and resulting p-values were subsequently corrected for multiple hypothesis testing using the Benjamini–Hochberg method in DanteR (54). Data were plotted with Perseus software. The gene ontology was performed using the Database for Annotation, Visualization and Integrated Discovery (DAVID) version 6.7. (55,56) Functional Annotation Charts were created using default settings.

Immunofluorescence

Cells grown on glass coverslips were fixed in PBS/3.7% FA (Roth:7398.1) for 10 min at room temperature. After fixation, the cells were washed in PBS and then permeabilized in 3.3 mM sodium citrate and 0.1% Triton X-100 for 5 min at room temperature. The cells were then washed two times with PBS and two times with washing solution I (PBS-Tween (0.1%) plus 0.25% BSA). Blocking was performed with blocking solution I (5 ml washing solution I/2.5%
Figure 2. Chromatin containing GFP-tagged histone variants is released after MNase digestion. (A) Chromatin purification protocol (left) and DNA analysis of the first and second supernatants containing chromatin (S1 and S2, respectively). (B) Western blot analysis of the GFP-tagged histone variant release compared to H3 in presence or absence of chromatin MNase digestion.

BSA) for 30 min at room temperature. Primary antibodies were diluted in blocking solution I. Incubation with the primary antibody was performed overnight at 4°C. The next day after washing with washing solution I the cells were incubated with the secondary antibodies diluted in blocking solution II (blocking solution I plus 10% normal goat serum) for 1 h at room temperature. Slides were mounted with Vectashield containing 4',6-diamidino-2-phenylindole (DAPI; Vectorex Laboratories) and sealed with nail polish. Images were acquired using a Zeiss LSM 510 META confocal microscope equipped with a 63× oil immersion objective NA = 1.4 or a Leica TCS SP5 confocal microscope with a 63× glycerol immersion objective NA = 1.3. Digital image processing, light linear adjustment of brightness and contrast, was done using ImageJ.

The primary antibodies used were anti-H3K27me3 (kind gift of Gunnar Schotta) and anti-GFP (mixture of mouse monoclonal antibodies from Roche). For detection, the cells were stained with donkey anti-mouse Alexa 488 (Jackson Immuno Research), goat anti-rabbit Cy3 (Millipore), TO-PRO-3 (Invitrogen). For live cell imaging, MEF transfected cells were plated on a glass bottom p35 dish and grown under standard conditions. 3D stacks were acquired at 20 min intervals overnight. Time lapse imaging was carried out on a UltraVIEW VoX spinning disk confocal system (PerkinElmer, UK) in a closed live cell microscopy chamber (ACU control, Olympus, Japan) heated to 37°C, with 5% CO2 and 60% air humidity control, mounted on a Nikon Ti microscope (Nikon, Japan), using a 60×/1.45 NA Planapochromat oil immersion objective lens. Images were obtained with a cooled 14 bit EMCCD camera (C9100-50, CamLink). Maximum intensity Z-projections were assembled into videos and annotated using ImageJ (http://rsb.info.nih.gov/ij/). For cyclobutane-pyrimidine dimer (CPD) analysis, cells were grown on glass coverslips. Fixation of the cells was done with 3.7% formaldehyde for 10 min at room temperature followed by a combined permeabilization and DNA denaturation step with ice-cold 0.7% Triton X-100/0.4 N HCL/PBS for 10 min. For blocking step, the cells were incubated with 4% BSA for 30 min. Immunostaining for CPD was performed by incubating the cells with monoclonal anti-CPD (Kamiya Biomedical Company, Cat no.: MC-062) at a dilution of 1:200 for 1 h at room temperature. For detection, cells were incubated with polyclonal anti-mouse IgG Cy3 (Jackson Laboratories, Cat no.: 715–165–151) at a dilution of 1:330 for 1 h at room temperature. The DNA content was stained with DAPI and finally mounted in Mowiol 4-88 (Sigma-Aldrich Chemie).
In situ replication labeling

To visualize newly synthesized DNA, cells grown on coverslips were pulse labeled with either 10 μM EdU (Figure 4) or 100 μM 5-bromo-2′-deoxyuridine (BrdU, Sigma-Aldrich) (Figure 5) and either fixed immediately or chased for the indicated time in pre-warmed medium supplemented with 200 μM thymidine. Cells were fixed with 3.7% formaldehyde for 10 min and permeabilized with 0.5% Triton X-100 for 20 min. BrdU detection was performed with rat-anti-BrdU (Gentaur, Cat no.: OBT0030CX, dilution 1:100) in the presence of DNase I as described (50). Click reaction was performed using the click-it kit (Invitrogen). Alexa Fluor 647 azide was added in a final concentration of (2 μM). The click reaction was performed for 30 min at room temperature. The cells were then stained with an anti-GFP antibody as described above.

Nuclear extract for western blot analysis

GFP-histone variant transfected MEFs and HL cell lines were incubated with 0.1% Triton in PBS supplemented with
Figure 4. H2A.Bbd localizes to DNA synthesis sites and is replaced within 1 h. (A) Single confocal slice of female MEFs expressing the GFP-tagged histone variants pulsed labeled with EdU. (B) Single confocal slice of female MEFs expressing the GFP-tagged histone variants pulse labeled with EdU and chased for 1 h before fixation. DAPI: DNA (blue); GFP: GFP-tagged histone variant (green); EdU: replication foci (red). Scale bar: 5 μm. The channel intensity recorded along the white line is plotted on a diagram. Blue line: DNA; green line: GFP-tagged histone variants; red line: replication foci.
proteinase inhibitor for 10 min at 4°C. After centrifugation at 3000 rpm the nuclear pellet was resuspended in 1× PBS supplemented with proteinase inhibitors and the DNA was digested with Benzonase (VWR) for 15 min at 37°C. 2× Laemmli buffer was added to the suspension of nuclei and boiled for 15 min at 95°C.

Western blot antibodies

The anti-GFP antibody (a mixture of two mouse anti-GFP monoclonal antibodies (Roche)) was used at a dilution of 1:1000, the anti-H3 antibody (Abcam ab1719) was used at a concentration of 1:5000 and the H2A.Bbd antibody (Millipore polyclonal) at a concentration of 1:1000. HRP-coupled secondary antibodies were used at a concentration of 1:5000 and detection was performed using ECL chemiluminescent reagents (GE Healthcare Life Sciences).

Microirradiation experiments

Microirradiation experiments were carried out in Nikon Ti spinning disk microscope (see live cell imaging) with a 405 nm diode laser set to 100% transmission. Approximately 1 μm spots were selected inside the nucleus and microirradiated with 1 mJ. Pre- and post-bleach images were acquired as indicated.

UV-C irradiation experiments

For UV-C irradiation experiments, the cells were irradiated with a UV-C source containing 5.8 W short wave lamps (Amersham Biosciences) at a wavelength of 254 nm. For damage induction, the media were removed from the cells and were globally irradiated for 10 s with a final dose of 1 mJ/cm². Unirradiated cells were used as control.

CPD quantification

The levels of DNA damage were calculated from confocal 3D images of CPD stainings and DNA counterstained with DAPI using the image analysis platform Priithon (http://code.google.com/p/priithon/). The DAPI images were used to create 3D nuclear masks by applying a median filter (sigma = 500 nm) and segmenting the filtered images using automatic thresholding. The integrated CPD signal within the nuclear mask was divided by the mask volume to obtain the mean CPD intensity in each nucleus.

Quantitative reverse transcription-polymerase chain reaction

RNA from HL cell lines and PBMCs was isolated using Trizol reagent (Invitrogen, Karlsruhe, Germany) following the manufacturer’s protocol. After reverse transcription of 2 μg of RNA, quantitative reverse transcription-polymerase chain reaction (qRT-PCR) was performed essentially as described (57). The following primer combinations have been used: actin beta (ACTB): 5’-ggc atc tgt gac tcc g-3’ and 5’-gtc gga agg tgt acg gcg a-3’; H2A.Bbd: 5’-tgc ttt tca gtg atg gac tcc g-3’ and 5’-cag agg gac tga agc cag gt-3’. Quantification of gene expression was performed using the 2ΔΔCt method [H], and for comparative analysis the median Dct value from PBMC (21 independent donors) was set as one.

RESULTS

To investigate the composition of H2A.Bbd-containing chromatin we generated female MEF clonal cell lines expressing GFP-H2A.Bbd, macroH2A.1.2 and -H2A. These cell lines express the corresponding histones at approximately similar levels (Figure 1A). After several weeks in culture, we noticed that whereas the levels of GFP-H2A.Bbd and GFP-H2A remained the same, the amount of GFP-macroH2A.1.2 dropped, as seen by immunoblot analysis of nuclear extracts (Figure 1B). This phenomenon was observed in several attempts and may be caused by a negative effect of high levels of macroH2A.1.2 on the cell cycle (18) and a subsequent negative selection of the highly expressing cell clones. In agreement with previously published results, we detect GFP-macroH2A.1.2 co-localizing with the inactive X-chromosome labeled with an antibody recognizing H3K27me3 (22) whereas GFP-H2A.Bbd is largely excluded from the Barr Body (31) (Figure 1C). The canonical GFP-H2A in contrast does not show enrichment on a particular chromosome or nuclear domain (data not shown).

Upon establishment of stable lines expressing the different GFP-tagged histones, we isolated chromatin fragments from MEFs using micrococcal nuclease (MNase) digestion. As chromatin is organized in large domains in which not all nucleosomes contain the same histone isoform (36), we pooled chromatin fragments of different lengths that were released from the nuclei using a combination of different extraction protocols (Figure 2A). The tagged histone variants (Figure 2B) were released with similar efficiency from...
the nuclei, suggesting that the physicochemical properties of the nucleosomal arrays carrying specific variants in MEFs are roughly the same. MNase treatment released more than 80% of the tagged chromatin, which is comparable to the amount isolated by Tolstorukov et al. (29) and significantly higher than the levels released in experiments performed by Ishibashi et al. (34). This difference may be due to experimental differences or differences in cell culture conditions as we also observe a small fraction of GFP-H2A.Bbd in the nuclear matrix fraction. However, our isolation procedure was selective for chromatin-bound histones, as we could not release histones from nuclei without MNase digestion (Figure 2B).

The nucleosomal arrays containing GFP-H2A.Bbd, GFP-macroH2A.1.2 or GFP-H2A were purified by GFP-affinity chromatography (Figure 3A). The analysis of the protein composition of these fibres showed that, not surprisingly, the oligonucleosomal arrays contain not only the tagged versions of H2A used for purification but also all four canonical histones. This is in good agreement with our previous findings that chromatin domains are not homogeneous with regard to their histone isoform/variant composition (36).

The non-histone proteins that were isolated together with the tagged short nucleosomal arrays were identified and quantified by separation of the proteins via SDS-PAGE, and a subsequent identification by liquid chromatography-mass spectrometry (LC-MS). Proteins were identified from these runs using the Andromeda algorithm of the MaxQuant protein analysis package and quantified using their iBAQ values (51,58). We identified a total number of 1428 proteins in three independent biological replicates for each variant and control. More than 50% of the proteins are identified in at least two independent biological replicates. The overlap varies between samples: 53.4% for H2A.Bbd, 51.6% for H2A and 53.1% for macroH2A.1.2 (Supplementary Figure S1). A protein was considered to be specifically chromatin-bound when its average log2 transformed iBAQ value had a difference of at least 3.5 between the GFP-H2A variant sample and the background control and it was detected in at least two samples. The application of these filters resulted in the identification of 607 bona fide chromatin proteins. For every chromatin-bound protein, we determined its variant specificity value as the log2 value of the ratio of the corresponding enrichment factors (Figure 3B). As expected based on the different chromosomal distributions (22,31) and Figure 1C, the GFP-H2A.Bbd- and GFP-macroH2A.1.2-containing short nucleosomal arrays were the most divergent (Figure 3B). For the functional analysis, we focused our further analysis towards the proteins enriched in the GFP-H2A.Bbd-containing chromatin. Very little is known about the effect of H2A.Bbd on chromatin structure and its molecular function. We therefore wondered whether a global analysis of the H2A.Bbd bound proteome would provide a deeper insight. To this end, we performed a gene ontology analysis for factors enriched in the GFP-H2A.Bbd compared to the GFP-macroH2A.1.2 and the GFP-H2A samples (upper right panel, Figure 3B). Consistent with the recent finding that H2A.Bbd might play a role in RNA splicing (29,30), we observe a strong overrepresentation of gene ontology categories related to RNA maturation (Tables 1 and 2). In addition, when we compared the H2A.Bbd- with macroH2A.1.2-containing chromatin, we found a strong enrichment of factors involved in DNA replication such as PCNA, the MCM proteins, the FACT complex and the CAF1 complex (Tables 1 and 3). These proteins are also strongly enriched when comparing GFP-H2A.Bbd chromatin with chromatin isolated from cells expressing the GFP-tagged H2A isoform (Tables 2 and 3), suggesting that they are specific for GFP-H2A.Bbd rather than excluded from chromatin that contains high levels of GFP-macroH2A.1.2.

When characterizing the nuclear distribution of GFP-H2A.Bbd, we noticed a subpopulation of cells that had a particular GFP-H2A.Bbd localization pattern. In contrast with the hypothesis that H2A.Bbd is preferentially found in transcriptionally active chromatin, these cells showed a strong association with DAPI dense regions within the nucleus, which are usually considered to be transcriptionally inert (Supplemental Figure S2). Based on our observation that GFP-H2A.Bbd is associated with the replication machinery, we wondered whether the cells that showed this particular localization pattern are in fact engaged in replicating heterochromatic DNA.

We therefore labeled asynchronously growing MEFs ectopically expressing GFP-H2A.Bbd with EdU to identify cells that are in S-phase. Consistent with our hypothesis we observed a striking co-localization of the EdU signal with GFP-H2A.Bbd in all S-phase cell lines (Figure 4A and Supplementary Figure S3). However, after a 1 h chase following the 30 min EdU labelling, the co-localization is lost suggesting a transient interaction (Figure 4B). To get a better understanding of the dynamic localization of H2A.Bbd to sites of DNA synthesis, we performed live cell imaging of cell lines expressing GFP-tagged H2A.Bbd (Figure 5 and Supplementary movie). High-resolution microscopy revealed that H2A.Bbd is incorporated slightly after PCNA is recruited to sites of DNA replication, which confirms our finding that H2A.Bbd interacts with PCNA-containing chromatin (Figure 5). Using this technique we could also determine a residence time of H2A.Bbd at replication foci of less than 10 min at early replicating foci and 30 min at sites of late replication (Supplementary Figure S4). As we see a close association with PCNA, we wondered whether GFP-H2A.Bbd is also incorporated at sites of DNA damage. For this we microirradiated MEFs expressing GFP-tagged H2A.Bbd at 450 nm for 1200 ms. After irradiation, PCNA quickly localizes to sites of DNA damage followed by GFP-H2A.Bbd, which is similar to their timing of localization to DNA replication sites. However, in contrast to replication sites, GFP-H2A.Bbd can be detected at repair foci for at least 2 h (Figure 6).

In order to investigate whether the expression of H2A.Bbd has an effect on replication and DNA damage, we compared the S-phase length of cells expressing GFP-H2A.Bbd with the ones expressing the canonical GFP-H2A. In agreement with the hypothesis that H2A.Bbd facilitates chromatin remodeling or assembly during DNA replication, we detect a 20% shorter S-phase in the GFP-H2A.Bbd expressing MEFs compared to the GFP-H2A expressing cells (Figure 7A). Moreover, the expression of H2A.Bbd also increases the sensitivity to UV based DNA damage (Figure 6B).
damage as measured by the increased formation of CPDs (Figure 7B). The shortening of S-phase and increased sensitivity toward DNA damage are hallmarks of cancer cells. Indeed, the analysis of Gene Expression Omnibus microarray data sets from HL cell lines and primary HL samples in comparison with normal tissue revealed increased levels of H2A.Bbd RNA in a large number of HL lines ((17); Figure 7C). Consistent with this finding, chromatin isolated from HL cells contains much higher levels of H2A.Bbd than control cells (Figure 7D) where H2A.Bbd is not detectable. Interestingly, the total levels of H2A.Bbd vary substantially in different HL lines with the ones containing a lot of H2A.Bbd having a shorter doubling time than the ones with lower expression levels ((32); Figure 7E), suggesting that H2A.Bbd overexpression might have a similar effect in MEFs and tumor cells.

**DISCUSSION**

So far most studies addressing the function of the histone variant H2A.Bbd focused either on the structural characteristics of nucleosomes containing these variants ((33)) or on the DNA sequences that are associated with it ((27,29,30)). However, the physicochemical properties of chromatin and its function are mainly defined by associated proteins. As the current knowledge about the proteome bound to the H2A.Bbd-containing chromatin is limited, we performed a detailed proteomic analysis on chromatin isolated from cells expressing a GFP-tagged version of H2A.Bbd and used the purification of nucleosomal arrays containing GFP-macroH2A.1.2 and GFP-H2A.Bbd as highly specific filters for a differential proteomics approach. As the cells we use to analyze H2A.Bbd bound chromatin do not show significant expression levels of this variant, the system most likely reflects the situation in tumor cells that have a high expression level for H2A.Bbd.

In agreement with earlier findings investigating chromatin fibres containing H3 variants H3.1 and H3.3 ((36)), we observed that arrays of nucleosomes containing either one of the two H2A variants also contain a sizeable amount of canonical histones. Despite this apparent heterogeneity,

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**Table 1. DNA replication factors are significantly over-represented in the GFP-H2A.Bbd compared to the GFP-macroH2A.1.2 pull-down**

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<tr>
<th>Category Term</th>
<th>Count</th>
<th>%</th>
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<th>Bonferroni</th>
<th>Benjamini</th>
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Top enrichment terms for the factors enriched in the H2A.Bbd/macroH2A.1.2 sample. The terms and their related p-values for Biological Processes (BP) are displayed. Category: original database/resource where the terms orient; Term: enriched term associated with the gene list; Count: gene in the list involved in the term; Percentage: involved genes/total genes; p-value: modified Fisher exact p-value; EASE score.
Table 2. Replication factors are significantly over-represented in the GFP-H2A.Bbd compared to the GFP-H2A pull-down

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Top enrichment terms for the factors enriched in the H2A.Bbd/H2A sample. The terms and their related p-values for Biological Processes (BP) are displayed. Category: original database/resource where the terms orient; Term: enriched term associated with the gene list; Count: gene in the list involved in the term; Percentage: involved genes/total genes; p-value: modified Fisher exact p-value, EASE score.

Table 3. Replication associated factors are enriched in chromatin containing GFP-H2A.Bbd

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<tr>
<th>Protein names</th>
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Enriched replication factors in the GFP-H2A.Bbd-containing chromatin are displayed according to their enrichment factors and p-values calculated in the GFP-H2A.Bbd/GFP-macroH2A.1.2 and in the GFP-H2A.Bbd/GFP-H2A samples.

of the purified fibres, these histone variants have a major impact on the proteomic composition of chromatin. Note-worthy, the number of proteins detected in GFP-H2A.Bbd chromatin is higher compared to chromatin isolated from GFP-H2A or GFP-macroH2A 1.2. This observation could be due to an intrinsic feature of H2A.Bbd containing chromatin or a result of slight expression differences of the histone variants within the cells. In order not to bias our quan-titative analysis, we normalized the resulting quantitative information of all variant specific chromatin proteins to the total amount of isolated chromatin using the histone H4 iBAQ values.

We observed a striking enrichment of factors involved in DNA replication within the GFP-H2A.Bbd-containing nucleosomal arrays in an asynchronous population of cells. Although most proteins reported to reside in the man-
malian replisome are enriched in the GFP-H2A.Bbd associated chromatin, not all enrichment values are significantly different among the variant containing chromatin when analyzed by rigorous statistical testing. This may well be due to the fact that only a fraction of cells within the population is actively replicating and incorporating H2A.Bbd at sites of DNA synthesis.

Nevertheless, the presence of factors associated with DNA replication in the chromatin containing GFP-H2A.Bbd prompted us to analyze its involvement in this process using alternative methods. Indeed, when we analyzed DNA replication by immunohistochemistry and live cell imaging, we observed a strong co-localization pattern of GFP-H2A.Bbd with sites of DNA replication. Interestingly, a previous report had reported a dotted staining pattern in a small fraction of H2A.Bbd transfected MEFs (34), which in light of our findings most likely reflects replicating chromatin. Histone variants have been shown to be recruited to DNA repair foci (37), however, they have not been found co-localizing with replicating DNA. H2A.Bbd is a striking exception to this rule. Interestingly, this enrichment is only transient as the co-localization of H2A.Bbd with newly synthesized DNA is only detected for less than 60 min, which may be the reason why it has gone unnoticed so far. This apparent instability of the newly formed nucleosome is in agreement with the observed low intrinsic stability of the H2A.Bbd-containing particles and its inability to bind the linker histone H1 (39,41).

H2A.Bbd is incorporated at both early and late replicating chromatin during S-phase, which points to a general role of the DNA synthesis machinery in the incorporation of this histone variant. Alternatively, the localization of H2A.Bbd to sites of DNA synthesis might reflect its ability to bind to naked DNA at replication sites. However, we do not think this is the sole mechanism of incorporation as we do not observe it for the canonical H2A variant that is expressed to comparable levels and has a similar affinity for naked DNA. The differential residence time of H2A.Bbd in early or late replicating foci may be related to the different efficiency of replication fork progression in eu- and heterochromatin or to a different binding dynamics of an H2A.Bbd interacting factor. What is intriguing, however, is the substantial increase in residence time on DNA repair foci, which may be due to an extended presence of PCNA at those regions.

Like most other studies performed before, we do not investigate H2A.Bbd containing chromatin isolated from cells that naturally express this variant. It is therefore difficult to make a conclusion about its physiological function. However, it is tempting to speculate that the incorporation of H2A.Bbd at sites of DNA synthesis could physiologically contribute to the maturation of sperm and the replacement of the canonical histone with protamines. The putative mouse homologue H2A.Lap1 is expressed in all stages of spermatogenesis with the highest level between the late pachytene and round spermatid stage when it is exported to the cytosol and another H2A.Bbd-like histone becomes incorporated into the chromatin (17–20,35). The presence of H2A.Bbd-like histones in mouse coincides with episodes of increased DNA breakage in sperm (17,18,21,35,59). Moreover, the DNA repair machinery might facilitate the incorporation of H2A.Bbd, thereby facilitating DNA recombination and protamine exchange. As spermatocytes expressing H2A.Bbd are postmitotic, it is unlikely that H2A.Bbd plays a role in DNA replication.

As pointed out above, our system more closely resembles the elevated expression of H2A.Bbd in HL cells (Figure 7C.

Figure 6. H2A.Bbd accumulates at DNA damage sites following PCNA. Time-lapse imaging of microirradiated MEF cells stably expressing GFP-H2A.Bbd (green) and transfected with mRFP-PCNA (red) at 405 nm for 1200 ms. Images were collected before microirradiation and 5 s, 10/50 and 110 min after microirradiation. Scale bars: 5 μm.
Figure 7. MEF cells and HL lines expressing H2A.Bbd replicate faster and are more sensitive to DNA damage. (A) Doubling time and S-phase length in MEF cells stably expressing GFP-H2A.Bbd and GFP-H2A. (B) Mean intensity of nuclear CPD signal 10 and 30 min post 1 mJ UV-C irradiation. Error bars represent the 95% confidence interval. N > 30 (C) qRT-PCR expression analysis of H2A.Bbd in HL cells and control cells. cDNA from HL cell lines (two independent experiments) and normal PBMC (21 donors) was used as template for quantitative PCR. For calculation of relative expression values, actin beta was used as housekeeping control and the median expression in normal PBMC was set as 1. (D) Western blot analysis of H2A.Bbd in Hodgkin’s lymphoma cells (HL) and control cell lines. Ponceau staining of histone H3 is used as loading control. (E) Dot plot showing the correlation in Hodgkin’s lymphoma cell lines between the doubling time derived from (32) (y-axis) and the expression levels of H2A.Bbd (x-axis) measured by qRT-PCR. Every dot represents a specific Hodgkin’s lymphoma cell line.
and D). Such an overexpression might result in a faster S-phase progression similar to what we observe in MEFs expressing GFP-H2A.Bbd (Figure 7). Indeed, HL cell lines with lower expression of H2A.Bbd (cell lines L-540 and HDLM-2) have longer doubling times than cell lines with higher expression of H2A.Bbd (22,23,32). HL high sensitivity to radio- and chemotherapy is known for a long time and has led to the development of highly effective treatment regimens (reviewed in (60)). H2A.Bbd is specifically overexpressed in HL cell lines and in HL cells in vitro (17) and might be involved in the high success of radio- and chemotherapy for treatment of these patients.

CONCLUSION

Our data support a functional compartmentalization of chromatin by histone variants and their associated factors. The finding that H2A.Bbd transiently localizes to sites of DNA synthesis in cells thereby increasing S-phase progression and sensitizing the cells for DNA damaging UV irradiation opens up new avenues for the investigation of the function of this exceptional histone variant in healthy and diseased tissue.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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Conflict of interest statement.

REFERENCES


Supplementary Material

Supplementary Figure S1: Identified proteins overlap among the replicates. Protein factors identified after the GFP-H2A variants chromatin purification using the GFP nanotrap beads are displayed in Venn diagrams showing the identified proteins overlap in three independent biological replicates (e.g. H2A.Bbd exp.#1, #2 and #3).

Supplementary Figure S2: GFP-H2A.Bbd co-localizes with DNA-dense regions in a subpopulation of cells. Single confocal slice of asynchronous population of female MEF cells that express GFP-H2A.Bbd. To-PRO: DNA (blue); GFP: GFP-H2A.Bbd (green). The arrow indicates GFP-H2A.Bbd co-localizing with DNA dense regions. The two panels represent different areas of acquisition. Scale bar upper panel: 5 µm; lower panel: 10 µm.

Supplementary Figure S3: GFP-H2A.Bbd localizes to sites of replication during all stages of S-phase. Single confocal slice of female MEFs expressing H2A.Bbd labeled with EdU and stained with anti-GFP at different time of replication. EdU: replication foci (red); GFP-H2A.Bbd: GFP (green); DAPI: DNA (blue). Early S-phase: eS; late S-phase: IS. Scale bar: 5 µm.

Supplementary Figure S4: GFP-H2A.Bbd has different resident time in early- and late- replicating chromatin. Optical sections of confocal images of replicating MEF cells stably expressing GFP-H2A.Bbd and pulsed labeled with BrdU (red) for 5 min and fixed 0 / 5 / 10 min thereafter as indicated. Scale bars: 5 µm.

Supplementary Movie. Localization of H2A.Bbd to sites of DNA synthesis in vivo. Live cell imaging of MEF cells stably expressing GFP-H2A.Bbd (green) transfected with mRFP-PCNA (red). 3D-stacks were acquired at 20-minute intervals.
over night using a spinning disk microscope equipped with temperature, humidity and CO$_2$ control. Shown are maximum intensity projections of each time point. Scale bar: 5 µm.
Supplementary figure S1, Sansoni et al.
Supplementary figure S2, Sansoni et al.
Supplementary figure S4, Sansoni et al.
Supplementary figure S4, Sansoni et al.