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Phosphatidylserine colocalizes with epichromatin in interphase nuclei and mitotic chromosomes

Igor Prudovsky,1 Calvin P.H. Vary,1 Yolanda Markaki,2 Ada L. Olins3 and Donald E. Olins3,*

1 Maine Medical Center Research Institute; Scarborough, ME USA; 2 Department of Biology II; Ludwig Maximilians University; Munich, Germany; 3 Department of Pharmaceutical Sciences; College of Pharmacy; University of New England; Portland, ME USA

Keywords: nuclear architecture, glycerophospholipid, chromatin, confocal and deconvolution microscopy

Abbreviations: PS, phosphatidylserine; PC, phosphatidylcholine; PFA, formaldehyde; NE, nuclear envelope; INM, inner nuclear membrane; 3D-SIM, structured illumination microscopy; PLC(D), phospholipase C(D)

Cycling eukaryotic cells rapidly re-establish the nuclear envelope and internal architecture following mitosis. Studies with a specific anti-nucleosome antibody recently demonstrated that the surface (‘epichromatin’) of interphase and mitotic chromatin possesses a unique and conserved conformation, suggesting a role in postmitotic nuclear reformation. Here we present evidence showing that the anionic glycerophospholipid phosphatidylserine is specifically located in epichromatin throughout the cell cycle and is associated with nucleosome core histones. This suggests that chromatin bound phosphatidylserine may function as a nucleation site for the binding of ER and re-establishment of the nuclear envelope.

Introduction

Peripheral chromatin, located adjacent to the inner nuclear membrane (INM) of the interphase nucleus, is immunologically distinct from internal chromatin when examined with a specific autoimmunere mouse monoclonal antibody (PL2-6) directed against a conformational epitope within the nucleosome.3,4 Furthermore, mitotic chromosomes exhibit surface staining with PL2-6, even though the nuclear envelope (NE) is no longer present during “open” mitosis. We have named this distinctive surface region “epichromatin” and hypothesized that it represents a chromatin conformation which persists through mitosis, facilitating postmitotic nuclear envelope reformation and the re-establishment of interphase nuclear architecture.5 The structure and mechanism of formation of the epichromatin epitope remain largely unknown, although histones H2A, H2B and DNA are likely involved.3 Localization of epichromatin in proximity to the INM suggests that it may contain bound phospholipids. Phosphatidylserine (PS) is the major anionic glycerophospholipid within eukaryotic membranes, being distributed asymmetrically within lipid bilayers and involved in binding to membrane associated polycationic proteins.6 During apoptosis, PS is externalized within lipid bilayers and involved in binding to membrane-associated phosphatidylserine during the division cycle, we performed an immunostaining comparison of anti-PS antibody (1H6) with anti-epichromatin antibody (PL2-6) on rapidly growing tissue culture cells (i.e., human U2OS and mouse NIH 3T3). With the use of 1H6 on formaldehyde-fixed (PFA) and Triton X-100 permeabilized U2OS cells, the deconvolution microscopy images (Fig. 1A-F) bore a remarkable resemblance to U2OS cells immunostained with anti-epichromatin PL2-6 (ref. 2, Fig. 2A). For both antibodies, the interphase nuclei showed strong staining adjacent to the INM. During prophase, 1H6 staining was apparent on the most peripheral chromosomal regions, while at metaphase it surrounded the condensed chromosomes. During early anaphase, 1H6 reactivity remained confined to the periphery of chromosomes, whereas by late anaphase it was detectable on trailing chromosome arms. During telophase and G1 phase, the decondensing chromosomes showed only peripheral surface staining, similar to results for interphase epichromatin. We have obtained identical images using 1H6 on U2OS cells fixed with methanol (-20°C, 10 min), indicating that the fixation reagent is not a factor. In addition, we have recently identified a second independently derived purified mouse monoclonal anti-phosphatidylserine (Abcam clone 4B6), which yields identical immunofluorescent staining, compared with 1H6, with either PFA or ethanol fixation (Fig. S1). A more direct demonstration of colocalization of PS within the epichromatin region is presented in Figure 1G–I. Interphase and mitotic chromosomes

Results

Immunofluorescent analysis of nuclear phosphatidylserine localization. In order to examine the distribution of chromatin-associated phosphatidylserine during the division cycle, we performed an immunostaining comparison of anti-PS antibody (1H6) with anti-epichromatin antibody (PL2-6) on rapidly growing tissue culture cells (i.e., human U2OS and mouse NIH 3T3). With the use of 1H6 on formaldehyde-fixed (PFA) and Triton X-100 permeabilized U2OS cells, the deconvolution microscopy images (Fig. 1A-F) bore a remarkable resemblance to U2OS cells immunostained with anti-epichromatin PL2-6 (ref. 2, Fig. 2A). For both antibodies, the interphase nuclei showed strong staining adjacent to the INM. During prophase, 1H6 staining was apparent on the most peripheral chromosomal regions, while at metaphase it surrounded the condensed chromosomes. During early anaphase, 1H6 reactivity remained confined to the periphery of chromosomes, whereas by late anaphase it was detectable on trailing chromosome arms. During telophase and G1 phase, the decondensing chromosomes showed only peripheral surface staining, similar to results for interphase epichromatin. We have obtained identical images using 1H6 on U2OS cells fixed with methanol (-20°C, 10 min), indicating that the fixation reagent is not a factor. In addition, we have recently identified a second independently derived purified mouse monoclonal anti-phosphatidylserine (Abcam clone 4B6), which yields identical immunofluorescent staining, compared with 1H6, with either PFA or ethanol fixation (Fig. S1). A more direct demonstration of colocalization of PS within the epichromatin region is presented in Figure 1G–I. Interphase and mitotic chromosomes

*Correspondence to: Donald E. Olins; Email: dolins@une.edu
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of methanol-fixed NIH 3T3 cells were reacted first with PL2-6, washed with PBS and reacted with FITC-anti-mouse IgG. Subsequent PBS washes were followed by incubation with an excess of normal mouse IgG, in order to saturate all binding sites on the anti-mouse IgG. Lastly, the slides were incubated with TRITC-1H6, washed, incubated with TOPRO-3 and visualized by confocal microscopy. Supporting images document that the peripheral staining by 1H6 is not a consequence of an antibody-established barrier against diffusion (Fig. S2A). For immunostaining, we normally employ 1H6 at 1:200 dilution; identical, but weaker, images are obtained up to a 1:5400 dilution, making it unlikely that a concentration-dependent “layer” of bound antibody impedes diffusion of 1H6 deeper into the nucleus. A related control experiment (Fig. S2B) demonstrates that there is neither a 1H6 barrier blocking the nuclear penetration of an unrelated mouse mAb (BM28), nor does the strong peripheral FITC staining cause any obvious attenuation of the exciting and emitted light.

Employing 3D Structured Illumination Microscopy (3D-SIM), we have obtained high resolution immunostaining images (Fig. 1J–L) of interphase U2OS cells reacted with 1H6, PL2-6 and PL2-7 (an independent monoclonal anti-nucleosome antibody obtained from the same mouse as PL2-6). Both 1H6 and PL2-6 stained a thin rim at the periphery of the DAPI-stained DNA, while PL2-7 reacted throughout the chromatin. The similarity of staining in the epichromatin region by both 1H6 and PL2-6 (but not PL2-7) throughout the cell cycle is dramatically demonstrated with single 3D-SIM slices (Fig. 2).
Co-immunostaining comparisons of 1H6 with other nuclear antibodies are presented in Figure 3. A single U2OS nucleus co-stained with 1H6, anti-lamin B and DAPI and imaged by 3D-SIM is shown (Fig. 3A–C). Examination of the nuclear envelope at higher magnification (Fig. 3C, insert), demonstrated that lamin B is exterior to 1H6 staining (see also, Figure 4). The images also show that the nucleoplasmic reticulum is stained by anti-lamin B (arrows), but not by 1H6 (also not stained with PL2-6). Employing deconvolution microscopy to examine co-immunostained mitotic chromosomes, compared with 1H6, we found that: (1) the “mitotic marker” anti-H3S10p is present at the surface of metaphase chromosomes, penetrating more deeply than 1H6 (Fig. 3D–F); (2) the INM protein lamin B receptor (LBR) is retained within the cytoplasm, while 1H6 binds at mitotic chromosome outer surfaces (Fig. 3G–I) and (3) the INM protein emerin is distributed in a similar manner to LBR (Fig. 3J–L). Identical colocalization images comparing PL2-6 with H3S10p, LBR and emerin in U2OS cells have been published earlier (Figs. 3 and 4). Association of the phosphatidylserine epitope with compacted interphase peripheral chromatin. It has long been known that exposure of live tissue culture cells to hypertonic (hyperosmotic) buffer conditions rapidly leads to interphase chromatin condensation resembling the beginning of prophase. Presently, this induction of compacted interphase chromatin is regarded as an example of macromolecular crowding (volume exclusion), rather than the induction of mitosis. A recent study clearly demonstrated, by fluorescence microscopy and thin section electron microscopy, that treatment of MCF7 cells with a hypertonic buffer (320 mM sucrose added to tissue culture medium) for about 20 min induced compaction of peripheral chromatin, which moved away (retracted) from the still intact interphase nuclear envelope, generating “lacunas of fine fibrillar material.”

We decided to expose U2OS cells to the same hypertonic conditions, followed by formaldehyde fixation and immunostaining with 1H6 and PL2-6, to see whether the PS and epichromatin epitopes remained at the nuclear envelope or retracted with the compacted peripheral chromatin (Fig. 4). Using deconvolution microscopy, the resulting images convincingly demonstrated that both epitopes remained associated with the retracted chromatin in the epichromatin region. With respect to PS localization, this experiment strongly argues that its interaction with peripheral chromatin is much stronger than its interaction with other lipids of the INM.

Figure 2. 3-D SIM images of immunostained U2OS cells throughout the cell cycle. (A) 1H6 (red) and DAPI (blue). (B) PL2-6 (red) and DAPI (blue). (C) PL2-7 (red) and DAPI (blue). Each row is a gallery of single optical slices with the same antibody. Columns (left to right): interphase; prophase; metaphase; late anaphase-telophase. Note the peripheral staining by 1H6 and PL2-6, and the deep nuclear staining of PL2-7. Scale bar: 10 μm.
Conserved location of chromatin-associated phosphatidylserine. *Drosophila melanogaster,* like most eukaryotes, possesses PS in its cellular membranes. The evolutionary conservation of PL2-6 staining has been demonstrated in diverse animal and plant species. To examine whether the 1H6 epitope is also conserved, we immunostained *Drosophila* Kc cells with 1H6 or PL2-6 combined with the mitotic marker, anti-H3S10p, using deconvolution microscopy to view interphase and mitotic cells. Figure 5 demonstrates that PL2-6 and 1H6 yield identical and highly conserved staining patterns, similar to immunostained mammalian cells.

Characterizing the differences between 1H6 and PL2-6. Given the apparent identity of immunostaining by 1H6 and PL2-6, we attempted to distinguish the epitopes recognized by these two purified monoclonal antibodies of independent origins; the immunogen of 1H6 being liposomes containing PS, whereas PL2-6 is derived from an autoimmune mouse. Toward this goal, 1H6 and PL2-6 were absorbed with three types of liposome suspensions in PBS, separately prepared from PS, PC (phosphatidylcholine) or a combination of PS + PC (50:50 mixture), prior to immunostaining ethanol-fixed NIH 3T3 cells (Fig. 6A). The results demonstrated that absorption by PS significantly reduces
the staining with 1H6, but not with PL2-6; absorption by PC had a negligible effect. The liposome suspension containing both phospholipids was also effective in absorbing 1H6, but not PL2-6. This strongly argues that PL2-6 does not interact with PS-containing liposomes. We also examined whether PL2-6 can bind to PS which has been coated onto an ELISA plate (Fig. 6B). The experiment clearly indicated that whereas 1H6 reacted with PS-coated wells neither normal mouse IgG nor PL2-6 showed more than negligible reactivity.

In order to further define the PS epitope present within the fixed cells, we performed two types of extractions on coverslip-attached NIH 3T3 cells. Ethanol-fixed cells were incubated with chloroform overnight at -20°C. Following chloroform extraction, the coverslips were washed in PBS and blocked with BSA/PBS prior to immunostaining. 1H6 gave negligible staining, PL2-6 was moderately reduced and BM28 (an unrelated MCM nuclear protein) was unaffected by exposure to chloroform (Fig. 7A). The reduction of PL2-6 staining may reflect perturbation of the epichromatin conformational epitope, but was clearly not as profound as the reduction of 1H6 staining. For the second type of extraction experiment, we attempted selective removal of histones from methanol-fixed NIH 3T3 cells by incubation of the coverslips with various NaCl concentrations in 20 mM Tris, pH 7.4 (0.6 M NaCl has been shown to extract H1 histone from chromatin; 1.2 M NaCl removes H2A and H2B). Immuno-staining demonstrated that PL2-6 staining was unaffected by 0.6 M NaCl extraction, whereas 1H6 staining was strikingly reduced (Fig. 7B). After incubation with 1.2 M NaCl, both
strips with phospholipase C or D strongly reduced subsequent reactivity with 1H6, arguing that PS in milk associates with histones bound to PVDF, resulting in significant 1H6 binding. In conclusion, when "properly" blocked (e.g., casein or BSA in TBST), 1H6 displays negligible reactivity with acid-extracted histones separated by SDS-PAGE and blotted onto PVDF membranes, whereas, PL2-6 reacts with inner histones (probably H2A and H2B), but exhibits negligible reaction with H1.

We considered the possibility that SDS-PAGE separation of the acid-extracted histones might destroy conformational epitopes recognized by either 1H6 or PL2-6. Therefore, we prepared histone subfractions from NIH 3T3 cells by a method which better preserves conformational epitopes within histone complexes.\textsuperscript{17} ELISA assays were employed to compare the binding of 1H6 and PL2-6 to plates coated with the histone subfractions (H1, H2A + H2B and H3 + H4) extracted using an Active Motif kit, based upon.\textsuperscript{17} The subfractions were also examined for "purity" on a 15% SDS-PAGE and stained with Coomassie blue (Fig. S5). Three different concentrations (1.0, 0.1 and 0.01 µg/ml) of monoclonal antibodies and normal mouse IgG were tested on the histone-coated ELISA plates after blocking with 0.5% casein in TBST (Fig. 8D). The results demonstrated that PL2-6 yielded stronger reactions with H2A + H2B and H3 + H4. 1H6 exhibited clear reactions with the same histone subfractions (although the reaction with H3 + H4 was obviously more antibody concentration-dependent than with PL2-6). Thus, 1H6 recognizes H2A + H2B by ELISA, but exhibits only weak reactivity on immunoblots of SDS-PAGE resolved histones (Fig. 8A).

Comparing the binding of 1H6 and PL2-6 to histones. In a series of experiments, we compared 1H6 and PL2-6 binding to histones using electrophoretic immunoblotting. Previously, we demonstrated that PL2-6 reacts with the "inner histones" (especially with H2A and H2B) by immunoblotting or "dot blotting" onto PVDF membranes.\textsuperscript{6} When both 1H6 and PL2-6 were tested on PVDF membranes containing immunoblotted total acid-extracted histones\textsuperscript{14} resolved by SDS-PAGE, they unexpectedly yielded similar patterns of interaction with histones (Fig. 8A and B, lanes M). These membrane strips had been blocked with 5% low fat milk powder in TBST. Subsequently, we learned that milk contains histones and DNA\textsuperscript{15} and PS.\textsuperscript{16} Therefore, we prepared histone subfractions from NIH 3T3 cells by a method which better preserves conformational epitopes within histone complexes. The ELISA experiments implied that the epitopes, as present on fixed NIH 3T3 cells, can be distinguished by their different sensitivities to extraction; the PS epitope is more readily extracted by chloroform extraction; the epichromatin epitope is more readily extracted by sodium dodecyl sulfate (SDS) and NaCl, than is the epichromatin epitope. Therefore, we prepared histone subfractions from NIH 3T3 cells by a method which better preserves conformational epitopes within histone complexes.\textsuperscript{17} ELISA assays were employed to compare the binding of 1H6 and PL2-6 to plates coated with the histone subfractions (H1, H2A + H2B and H3 + H4) extracted using an Active Motif kit, based upon.\textsuperscript{17} The subfractions were also examined for "purity" on a 15% SDS-PAGE and stained with Coomassie blue (Fig. S5). Three different concentrations (1.0, 0.1 and 0.01 µg/ml) of monoclonal antibodies and normal mouse IgG were tested on the histone-coated ELISA plates after blocking with 0.5% casein in TBST (Fig. 8D). The results demonstrated that PL2-6 yielded stronger reactions with H2A + H2B and H3 + H4. 1H6 exhibited clear reactions with the same histone subfractions (although the reaction with H3 + H4 was obviously more antibody concentration-dependent than with PL2-6). Thus, 1H6 recognizes H2A + H2B by ELISA, but exhibits only weak reactivity on immunoblots of SDS-PAGE resolved histones (Fig. 8A).

Figure 6. (A) Comparison of anti-phosphatidylserine (1H6) and anti-epichromatin (PL2-6) after absorption with PS, followed by immunostaining. Confocal images of NIH 3T3 cells immunostained with unabsorbed antibodies or after absorption with phosphatidylserine (PS), phosphatidylinositol (PI) or a 50/50 mixture (PS/PI) are shown (top: 1H6; bottom: PL2-6). Color scheme: antibodies, red; TOPRO-3, blue. Scale bar: 20 µm. (B) ELISA comparison of PL2-6, 1H6 and normal mouse IgG (M IgG) binding to PS-coated wells. Y-axis: A490 from bound HRP anti-mouse IgG. 0.25, 0.06 and 0.01 mAb (ng) binding to PS-coated wells. X-axis: A490 from bound HRP anti-mouse IgG.

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We obtained clear evidence that both antibodies 1H6 and PL2-6 can interact with inner histones in solution; whereas, only PL2-6 significantly precipitates H1 (Fig. 8F). Examples of identified representative histone peptide sequences are presented in Figure S4. This experiment identifies specific histones that were detected in the immunoprecipitates, but does not yield information on the quantities of the histones, whether they were part of a protein complex or whether (in the case of 1H6) there was bound phosphatidylserine. Collectively, the immunoblotting, ELISA and immunoprecipitation plus mass spectroscopy argue that 1H6 does interact with H2A + H2B. But it is likely that this interaction is mediated through PS bound to H2A or to the H2A + H2B dimer. However, the interaction of PL2-6 with the inner histones does not depend upon the presence of PS (see Figs. 3B and 8B).

Discussion

The present study provides evidence that PS colocalizes with epichromatin. We have documented that two independently derived and purified mouse monoclonal antibodies (1H6 and 4B6), developed by immunizing with liposomes containing PS, and used to detect apoptosis (comparable to PS-specific Annexin V), stain the surfaces of chromatin in an interphase nucleus and on mitotic chromosomes, a region also recognized by the mouse antinucleosome monoclonal antibody PL2-6.8 Despite the similar immunostaining pattern of these mouse antibodies, they exhibit significant differences. Comparing 1H6 and PL2-6: (1) 1H6, but not PL2-6, can be absorbed by PS-containing liposomes, prior to immunostaining of ethanol-fixed NIH 3T3 cells. (2) By ELISA assay, 1H6 binds to PS, while PL2-6 does not. (3) Extraction of ethanol-fixed NIH 3T3 cells with chloroform depletes 1H6 immunostaining more effectively than PL2-6 staining. (4) NaCl extracts the 1H6 epitope from methanol-fixed NIH 3T3 cells at 0.6 M, with no reduction in PL2-6 staining. These observations strongly argue that 1H6 recognizes PS complexed with chromatin, while PL2-6 does not bind to PS.

There are very few studies on the consequences of glycerophospholipid interactions for chromatin structure and function. In a series of studies on isolated rat liver nuclei,19-21 addition of PS but not PC, vesicles was shown to enhance DNase I sensitivity, facilitate production of mononucleosomes by micrococcal nuclease, extract histone H1 (but not inner histones) and stimulate endogenous RNA polymerase activity. Two other studies22,23 focused on the strong electrostatic interactions of anionic phospholipids [PS and cardiolipin (CL)] with histones. By ELISA, the interaction of histones with CL was shown to be ionic strength dependent with strong binding at 0.3 and 0.6 M NaCl but severely reduced binding at 1.2 and 2.4 M NaCl.22 With the use of surface plasmon resonance, histone H2A has the highest association constant with PS-containing vesicles compared with the other individual histones.23 The authors suggest that the binding of histones to phospholipids “may contribute to the binding of histones to surfaces and blebs of apoptotic cells.” This conclusion is supported by recent evidence that H2B is tethered at the surface of macrophage and apoptotic cells via electrostatic interactions with PS.24 From the present and prior18 studies, there is little doubt that PL2-6 recognizes histones, principally the inner histones H2A and H2B. It is still unclear whether 1H6 will bind histones independently of the presence of PS. The present immunoblotting and ELISA experiments strongly indicate that reactions of 1H6 with histones or histone complexes are significantly reduced by treatment with phospholipases C and D prior to the immunoblot assay. It remains possible that 1H6 may be directed against an epitope consisting of PS plus histone amino acid residues. Such multireactive antibodies are not rare in autoimmunity and are...
currently viewed as a consequence of exposed nucleosome determinants at the cell surface of apoptotic cells. 25

An unresolved puzzle is why 1H6 can recognize PS in the epichromatin region and when exposed in apoptosis but does not detect PS in other cellular membranes. From analytical cell fractionation studies,3,26 PS is present in measurable quantities in the ER, Golgi and the plasma membrane. When GFP-lactadherin C2 domain was expressed in various tissue culture cell lines (A431, BHK-21, HeLa, COS7) PS was detected at the cytoplasmic side of the plasma membrane, the Golgi and endocytic vesicles and at the luminal side in ER and Golgi.27 GFP-Annexin V was transfected into Neuro 2A cells and internal calcium levels elevated by treatment with ionomycin, leading to detection of PS “in the cytoplasmic leaflet of the plasma membrane and possibly in the nucleoplasmic face of the nuclear envelope.” The lack of detection by 1H6 of PS in these various cell membranes might be due to, at least, three causes. The fixation procedures may lead to a selective loss of PS from most cellular membranes,28 sparing the PS associated with epichromatin. Second, the PS epitope recognized by 1H6 may be "masked" in most membranes. Third, the PS epitope recognized by 1H6 may be different in its presentation when bound to histones or chromatin, as in apoptotic bodies and in the nucleus, than when present in most membranes. When interphase chromatin is compacted by hypertonic sucrose (320 mM), the epitopes for both 1H6 and PL2-6 remain associated with the chromatin surface, clearly showing that they can be separated from the inner nuclear envelope membrane. This experiment also indicates that induced chromatin retraction does not result in a random redistribution of chromatin-bound PS; i.e., PS is tightly associated with the epichromatin region. However, we cannot rule out the possibility of PS redistribution from membranes specifically to the epichromatin region during the fixation and permeabilization protocols needed for immunofluorescent staining.

Nuclear envelope breakdown and reformation during mitosis must reproduce a faithful rendition of the parental nucleus in the derived daughter cells.29,30 An important question is how epichromatin might play a role in directing postmitotic reformation. Persistence of the epichromatin epitope throughout mitosis elicited the hypothesis2 that the surface of interphase and mitotic chromatin possesses a conformation which facilitates attachment of NE components. The present study suggests that phosphatidylserine from the interphase inner nuclear membrane may remain attached to epichromatin during mitosis, functioning as “nucleation sites” for the binding of ER and the re-establishment of the NE during postmitotic nuclear reformation.
Materials and Methods

Cell lines. Human U2OS cells were cultivated in DMEM medium, plus 20% fetal calf serum and 2 mM L-glutamine. Human HL-60/S4 suspension cells were maintained in RPMI 1640 medium, plus 10% heated fetal calf serum and 2 mM L-glutamine. Mouse NIH 3T3 cells (ATCC, Manassas VA) were maintained in DMEM, plus 10% calf serum. Drosophila melanogaster Kc 167 cells were maintained at 25°C, in Schneider’s Drosophila medium, plus 20% fetal calf serum and 2 mM L-glutamine. Mouse NIH 3T3 cells (ATCC, Manassas VA) were maintained in DMEM, plus 10% calf serum. Drosophila Kc cells were used prior to confluence. For confocal imaging, NIH 3T3 cells were plated on fibronectin-coated glass coverslips.

Antibodies. The sources and use of mouse monoclonal PL2-6, PL2-7, rabbit anti-H4S10ph, guinea pig anti-LBR and guinea pig anti-emerin have been described earlier.8 PL2-6 and PL2-7 were purified from tissue culture supernatants by affinity chromatography on protein G-Sepharose columns.18 1H6 and PL2-6 were purified from tissue culture supernatants by affinity chromatography on protein G-Sepharose columns. For colocalization experiments, 1H6 was labeled with TRITC using fluoropporter protein labelling kit (Life Technologies). A second purified mouse monoclonal anti-phosphatidylserine (Abcam clone #4B6) was examined, based upon a published use of this antibody for immuno-staining. For immunostaining experiments the mouse monoclonal anti-phosphatidylserine (Abcam clone #4B6) was examined, based upon a published use of this antibody for immuno-staining. Mouse monoclonal anti-BM28 a from Transduction Laboratories. Goat anti-lamin B is from Santa Cruz Biotechnology. Normal mouse IgG was obtained from Sigma Aldrich. For immunostaining experiments mouse monoclonal antibodies were used at ~5–10 µg/ml; immunoblots were performed ~0.5–1.0 µg/ml.

Immunostaining. A number of different protocols were employed, following slightly different methods in the different laboratories using different microscopes. When employing the DeltaVision deconvolution microscope at the German Cancer Research Center, PFA (formaldehyde) fixation, permeabilization and visualization were as described earlier for the study of U2OS, HL60/S4 and Drosophila Kc cells.6 Confocal data collected on NIH 3T3 cells using a Leica SP1 microscope (Maine Medical Center Research Institute) employed methanol or ethanol fixation (−20°C, 10 min), with DNA stained by TOPRO-3 and mounted in Vectashield. For the colocalization experiment (confocal imaging), PL2-6 was reacted first, followed by FITC-anti-mouse IgG, an excess of normal mouse IgG and, finally, TRITC-1H6. Besides demonstrating colocalization of the two mAbs, this experiment demonstrated that prior binding by PL2-6 did not significantly inhibit 1H6.

Samples prepared for 3-D SIM analysis (University of Munich) followed a protocol similar to the earlier publication,2 with the following differences: 2% PFA/PBS, 10 min, RT; gradient exchange from fixative to 0.02% Tween 20/PBS, 20 mM glycine/PBS, 10 min; 0.5% Triton X-100/PBS, 10 min; blocking with 2% BSA, 0.5% fish skin gelatin, 0.1% Tween 20/PBS; primary and secondary antibodies dissolved in blocking buffer; 4 washes in blocking buffer; post-fixation in 4% PFA/PBS; gradient buffer exchange (as above); DAPI staining and washing in PBS; mounting in Vectashield. Regardless of the specific fluorophor conjugated to the secondary antibody, images of 1H6 staining are artificially colored red in all figures. The other antibodies are presented in red or green, for convenience. DNA (chromatin) is artificially colored blue, regardless of whether the staining was with DAPI or TOPRO-3. All adjustments of brightness, contrast or color balance were linear adjustments and applied to the whole image.

Immunoblotting. Total acid extracted histones were prepared from undifferentiated HL-60/S4 cells following a published procedure.19 Gradient (10–20%) SDS-PAGE (BioRad Criterion) was run at 200 V for 1 hr. Each lane had acid extract from ~3 × 106 cells. Electrophoretic transfer to PVDF membrane was performed with a BioRad semi-dry apparatus in 250 mM glycine, 25 mM Tris, 0.05% SDS and 10% methanol, at 130 mA for 45 min. PVDF membranes were placed on both sides of the gels; each was stained with Ponceau S to confirm that histone migration was toward the anode. After the histone containing membrane was dry, it was cut into strips. For the blocking experiment, all strips were wetted with methanol, washed with TBST (Tris buffered saline + Tween 20) for 50 min, blocked with 5% milk in TBST, 1% casein (Sigma-Aldrich) in TBST, 5% BSA in TBST or TBST alone for 30 min at RT. Primary antibody dilutions (1H6 or PL2-6 in TBST) were incubated for 1 h at RT under fresh parafilm strips.

Following 6 × 5 min TBST washes of each strip in separate Petri dishes, each strip was incubated in HRP-antimouse IgG (1:5000 dilution in TBST) 1 h at RT under fresh parafilm strips and washed 6 × 5 min in separate fresh Petri dishes with TBST, ECL and film exposure were as described earlier.2 For testing the reactivity of 1H6 on the PVDF strips after milk blocking followed by phospholipase digestion, strips were incubated with 5% milk/TBST (1 h, 42°C), washed in TBST, followed by 25 mM Tris/HCl (pH 7.4), 140 mM NaCl, 2 mM CaCl2, then phospholipase C (PLC, 2.2 units/ml; Sigma P7633) or 50 mM Tris/HCl (pH 8.0), 140 mM NaCl, 5 mM CaCl2, containing phospholipase D (PLD, 3.8 units/ml; Sigma P0515) and incubated 1 h at 37°C. The strips were washed with TBST and incubated with 1H6. After washing with TBST, the strips were incubated in 5% milk (10 min, 37°C), then HRP conjugated 2° Ab (30 min), washed and developed with ECL for 1 and 5 min. This experiment argues that glycerophospholipids in milk (presumably PS) bind to blotted acid-extracted histones, leading to reactions with 1H6. Milk does not lead to nonspecific binding of the secondary antibody, only of 1H6.

ELISA. For analysis of the binding of 1H6, PL2-6 and normal mouse IgG to NIH 3T3 histone subfractions (H1, H2A/H2B and H3/H4), these fractions were diluted to 10 µg/ml in PBS. 50 µl of each sample was loaded per well on a PVC 96 well ELISA plate, covered with a sealing membrane and incubated ON at 4°C. Wells were washed three times with 100 µl TBST and blocked with 0.5% casein in TBST, 1 h at RT. 100 µl 1H6, PL2-6 or mouse IgG at 1.0, 0.1 and 0.01 µg/ml in TBST, or TBST alone, was added to each well and incubated 2 h at RT. Following
three washes with 200 µl TBST, 150 µl of HRP-antimouse IgG (1:5000 in TBST) was added for 1 h at RT. Following 4 × 200 µl TBST washes, OPD (o-phenyldiamine dihydrochloride) plus H2O2 in citrate buffer (pH 5.5) was added. Absorbance was read at 490 nm.

To examine the PS binding capability of 1H6, PL2–6 and normal mouse IgG, wells of a solid plastic immunoplate (Costar, #3798) were coated with 25 µl/well of PS (Avanti Polar Lipids) dissolved in hexane (20 µg/ml), which was allowed to evaporate ON at RT. Duplicate wells were blocked with 1% BSA in PBS for 2 h at RT, washed with 500 µl PBS, incubated with 25 µl of serial 2-fold dilutions of 1H6, PL2–6 or normal mouse IgG (starting at 1 µg protein/ml) in 1% BSA/PBS and incubated 1 h at RT. Wells were washed with 5 × 300 µl PBS, followed by 25 µl HRP-anti-mouse IgG (1:7500) in 1% BSA/PBS for 1 h at RT, washed again with 8 × 300 µl PBS and treated with OPD plus H2O2 in citrate buffer, followed by reading at 490 nm. To examine the effect of PLC and PLD upon the H2A + H2B subfraction, wells were coated with the histone subfraction (as above), washed in PBS, then incubated for 3 h at 37°C with PLC in PLC buffer (2 units/ml), PLD in PLD buffer (8 units/ml), with control wells containing either PLD buffer or PBS. Following the digestion, all wells were washed in TBST, incubated in 0.5% casein/TBST for 2 h at RT, washed in TBST, incubated in 0.1% casein/TBST for 2 h at RT. After extensive washing with TBST, reaction with HRP-antimouse IgG and further washing with TBST, the wells were assayed with OPD, as described above.

Immunoprecipitation and mass spectrometric analysis. Three full 15 cm dishes of NIH 3T3 cells were fractionated using the Qproteome Cell Compartment Kit (QIAGEN). Fraction 3 (Nuclear Proteins) was divided into two equal aliquots, precleared by addition of normal mouse IgG, anti-mouse IgG, protein A/G sepharose beads and incubated ON at 4°C with continuous rotation. After centrifugation, the supernatants were incubated with either specific mAb or with normal mouse IgG and protein A/G sepharose, followed by rotation for 6 h at 4°C. The experiment was performed twice; once comparing 1H6 with IgG, and once comparing PL2–6 with IgG. After extensive washing, the immunoprecipitated pellets were reduced using trichloroethyl phosphine, alkylated (iodoacetamide), and digested with trypsin. 8 The reaction with HRP-anti-mouse IgG and further washing with TBST, the wells were assayed with OPD, as described above.

Immuno-precipitated proteins were digested with trypsin and analyzed by mass spectrometry using a QSTAR Elite Triple Quadrupole/Time-of-flight mass spectrometer (Applied Biosystems). Following desalting (PepClean C18 column, Pierce), tryptic peptides were separated using a linear water/acetonitrile gradient (0.1% formic acid) on a Acclaim PepMap reversed-phase capillary column (3 µm, C18, 100 Angstrom pore size, 75 µm ID × 15 cm, 15 µm tip, LC Packings), with an inline PepMap 100 precolumn (C18, 300 µm ID × 5 mm, LC Packings) as a loading column. Mass spectrometry was conducted as previously described. 32,33 Peptides were eluted inline with a hybrid quadrupole/time-of-flight mass spectrometer (QSTAR, API Sciex), using a solvent gradient of 2–30% solvent B in 120 min and then 30–80% B in 10 min, in which solvent A is 0.1% formic acid and solvent B is 80% acetonitrile containing 0.1% formic acid. Data acquisition on the mass spectrometer was performed in information-dependent mode, whereby precursor peptides were automatically selected based on an MS-only scan and subjected to collision activated dissociation (MS/MS) in subsequent scans using a mass-dependent profile to set the collision energy for each MS/MS-interrogated peptide. Peak list generation, peptide mass fingerprint peak picking, and relative quantification and protein identification were performed with the ProteinPilotTM software (version 4.0; ABSciex). Default parameters were used for all analyses.

Protein searches were conducted using the UniProtKB/Swiss-Prot protein knowledgebase release 2011-03. MS/MS spectrum was searched against a database of human protein sequences using a mass tolerance of 0.6 Da and a detected protein threshold confidence of greater than 90%, set to detect contaminants (e.g., serum albumin). The number of unique proteins searched was 76,053. Threshold score for accepting individual MS/MS spectra was 95% confidence, based on confirmation by western blot analysis. Trypsin autolysis peaks and keratin tryptic peptides were known contaminants that were excluded from database searching. Peptide prevalence was estimated using the count of peptides for each protein that achieved > 95% significance. The detected protein threshold was set to 2.0 to achieve 95% confidence as determined by the Paragon algorithm (Applied Biosystems) to minimize redundancy, as recently described 34,35 and using automated bias correction.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Supplemental Material
Supplemental materials may be found here:
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