Liquid droplet formation by HP1α suggests a role for phase separation in heterochromatin

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Gene silencing by heterochromatin is proposed to occur in part as a result of the ability of heterochromatin protein 1 (HP1) proteins to spread across large regions of the genome, compact the underlying chromatin and recruit diverse ligands1–3. Here we identify a new property of the human HP1α protein: the ability to form phase-separated droplets. While unmodified HP1α is soluble, either phosphorylation of its N-terminal extension or DNA binding promotes the formation of phase-separated droplets. Phosphorylation-driven phase separation can be promoted or reversed by specific HP1α ligands. Known components of heterochromatin such as nucleosomes and DNA preferentially partition into the HP1α droplets, but molecules such as the transcription factor TFIIB show no preference. Using a single-droplets, but molecules such as the partition into the HP1α heterochromatin such as nucleosomes and DNA preferentially generated two different phosphorylated versions of HP1α in vivo (Fig. 1a). Phosphorylation was validated on HP1α in vitro (Fig. 1a, Phos-HP1α) and enhanced specificity for the H3K9me3 mark within nucleosomes5. To investigate if NTE phosphorylation has additional effects on peptidase and enhances specificity for the H3K9me3 mark within nucleosomes6,7. A plausible hypothesis is that the less conserved and less structured regions (hinge, N and C termini) are responsible for the unique properties of the different HP1 proteins8–11. Recent work has shown that phosphorylation of the N-terminal extension (NTE) of human HP1α (Fig. 1a) is important for formation of heterochromatin foci in cells12. The corresponding phosphorylation sites are absent in HP1β and HP1γ, which do not show detectable gene-silencing and gene-activating roles7. NTE phosphorylation increases affinity for an H3K9me3 tail peptide and enhances specificity for the H3K9me3 mark within nucleosomes5. To investigate if NTE phosphorylation has additional effects on HP1α, we generated different types of phosphorylated HP1α proteins as described previously12 (Fig. 1a). Phosphorylation was validated by mass spectrometry and H3K9me3 peptide binding5,12 (Extended Data Figs 1 and 7d). To separate the effects of phosphorylation on either the NTE or the hinge, which is also phosphorylated in vivo, we generated two different phosphorylated versions of HP1α (Fig. 1a, nPhos-HP1α and hPhos-HP1α, which are generated by respectively phosphorylating HP1α proteins that have their hinge or NTE serine residues mutated to alanine).

While working with nPhos-HP1α we observed the formation of a turbid solution upon cooling the protein on ice (Fig. 1b, left panel). The turbid solution became clear upon raising the temperature or upon treatment with alkaline phosphatase (Supplementary Videos 1 and 4). Investigation of the turbid material under a microscope revealed liquid droplets (Fig. 1b, right panel, Supplementary Video 2). Together, these observations are indicative of phase separation, a characteristic of proteins with intrinsically disordered regions and the capacity for multivalency14,15. Indeed the NTE, hinge, and C-terminal extension (CTE) regions contain sequences with a high propensity for intrinsic disorder16. Unlike nPhos-HP1α, wild-type HP1α did not phase-separate upon cooling. We quantified the saturation concentration for phase separation using two independent methods at room temperature (around 22–24°C) (Fig. 1c, e and Methods). This is the concentration at which the HP1α solution will appear as two separate phases. The saturation concentrations for the nPhos-HP1α, Phos-HP1α and hPhos-HP1α proteins increased in that order while wild-type HP1α, HP1β and HP1γ did not show detectable phase separation at the highest concentration tested (Fig. 1e and Extended Data Fig. 2). Replacing the NTE serine residues in HP1α with glutamate (nE-HP1α) did not cause any phase separation (Extended Data Fig. 2).

Given that phase separation is associated with multivalent interactions, we investigated whether nPhos-HP1α forms higher-order oligomers. We found that, in contrast to wild-type HP1α and HP1β, which do not show detectable higher-order oligomerization, nPhos-HP1α does form higher-order oligomers beyond a dimer (Fig. 2a and Extended Data Figs 3a, b, 4). Overall, HP1α proteins that were competent for phase-separation displayed higher-order oligomerization, while those that did not phase-separate appeared incapable of higher-order oligomerization (Fig. 2a and Extended Data Figs 3a, b). These data suggest that phase separation depends on inter-dimer contacts. Pairwise distance measurements using small angle X-ray scattering (SAXS) indicated that nPhos-HP1α is a substantially more elongated molecule than wild-type HP1α (Dmax ≈ 220 Å versus 130 Å, respectively, Fig. 2b). This extended conformation upon phosphorylation was further supported by size-exclusion chromatography–multi-angle light-scattering studies (Extended Data Fig. 6a).

We hypothesized that the extended conformation exposes positively charged hinge residues, allowing the phosphorylated NTE in one dimer to interact with hinge residues in another dimer (Fig. 2c). To test this possibility, we mutated a conserved basic patch in the hinge to alanines (residues 89–91, basic patch mutant) (Fig. 1a, Phos-HP1α(BPM)). Phos-HP1α(BPM) was defective for both phosphorylation-driven oligomerization and phase separation (Extended Data Figs 2a, 3a, c). Furthermore, amongst various chimeraes of HP1β and HP1α that we

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generated, only the chimera that has both the hinge and phosphorylated NTE of HP1α swapped into HP1β formed higher-order oligomers (Extended Data Fig. 3d). These results suggest that sequence features of both the hinge and NTE that are specific to HP1α are required for oligomerization.

To identify HP1α regions that help stabilize the compact conformation of wild-type HP1α we performed cross-linking by BS3 (bis(sulfo-succinimidyl)suberate) followed by mass spectrometry (Extended Data Fig. 1d). We found several putative inter-HP1α cross-links between the CTE and the hinge (21 out of 74, Supplementary Table). Deleting the 14-amino-acid CTE in the context of NTE phosphorylation (nPhos-HP1αΔCTE) lowers the saturation concentration by approximately tenfold compared to nPhos-HP1α (Fig. 1d, e). These results suggest that interactions between the CTE and the hinge stabilize the HP1α dimer in a compact auto-inhibited state that cannot make multivalent interactions (Fig. 2c). We therefore hypothesized that ligands which bind the CSD–CTE interface might alter the equilibrium between the closed and open states of nPhos-HP1α, regulating the ability of the CTE to stabilize the compact state. The shugoshin 1 (Sgo1) and lamin B receptor (LBR) proteins have been shown to interact directly with the CSD dimer of HP1α via a specific PXXVXL-like sequence in shugoshin, and a different sequence in LBR13. We therefore investigated the effects of these sequences as peptides on the phase-separation properties of nPhos-HP1α.

Consistent with previous work, both peptides bind specifically to the CSD–CTE interface12–15 (Extended Data Fig. 7a, b). Notably, the Sgo peptide promoted phase separation, lowering the saturation concentration by approximately 100-fold compared to nPhos-HP1α (Fig. 1d, e). The Sgo peptide also promoted nPhos-HP1α oligomerization (Extended Data Fig. 7c). H3K9 methylated and unmethylated tail peptides and equivalent concentrations of spermine also promoted phase separation (Extended Data Fig. 8a). As the H3 tail peptide is rich in lysine and arginine, these results suggest that in addition to specific ligands such as the Sgo peptide, which directly regulate HP1α oligomerization, other molecules can contribute to phase separation though general electrostatic effects (Extended Data Fig. 8).

In the model in Fig. 2c, the phosphates on the NTE make bridging contacts with the hinge region of a neighbouring dimer. The hinge also binds DNA in the context of HP1α and HP1β. We therefore wondered if DNA could bridge adjacent HP1 molecules and through its inherent multivalency promote phase separation. Saturating concentrations of DNA resulted in droplet formation by wild-type HP1α but did not cause droplet formation with HP1β (Fig. 3a). Mutating the basic patch in the HP1α hinge that is proposed to interact with DNA eliminated droplet formation (Fig. 3a). To understand this phenomenon better we used a DNA curtain assay and total internal reflection fluorescence microscopy to visualize the effects of HP1 on λ bacteriophage DNA molecules via the fluorescent dye YOYO-1, which intercalates into DNA4 (Fig. 3b, c).

The action of wild-type HP1α on DNA appears to be cooperative, as suggested by the emergence of fluorescent puncta, a result of local compaction leading to higher local concentrations of YOYO-1-labelled DNA (Fig. 3c–e). A non-cooperative mechanism would manifest as a global increase in fluorescence during compaction. Typically, for wild-type HP1α, a single puncta appears, followed by rapid (vav = 2.25 ± 0.026 s (s.e.m.); vav is average speed of DNA compaction) compaction of the rest of the 48.5 kb (approximately 12 μm) λ-DNA molecule without increased fluorescence outside the puncta (Fig. 3d, f). This compaction appears to be driven largely by electrostatic interactions, as increasing the level of monovalent salts reverses compaction (Fig. 3f, g). By contrast, YOYO-1 intensity was less localized and DNA strands often exhibited multiple fluorescent puncta during initial DNA binding and compaction.
by nPhos-HP1α (Fig. 3e, g, h), suggesting that cooperative binding to the DNA is disturbed by phosphorylation of HP1α. Furthermore, wild-type HP1α was able to compact DNA at much lower concentrations (Extended Data Fig. 9e). Finally, while nPhos-HP1α was able to completely compact λ-DNA, it compacted the DNA at a slower rate (1.1 μm s⁻¹ ± 0.15 (s.e.m.)) with much more variation than wild-type HP1α (Fig. 3f–h).

In some instances, inter-strand DNA interactions were clearly visible, indicating bridging across micrometre scales (Fig. 3i). Such linking would require, at a minimum, mesoscale protein–DNA networks consisting of around 100 individual wild-type HP1α dimers. Thus the energetics of phase separation could play a crucial role in the dynamics of DNA binding, compaction and organization, without the formation of macroscopic droplets. Notably, despite containing several positively charged residues in its hinge, HP1β was unable to compact DNA or produce puncta (Fig. 3j and Extended Data Fig. 9a, c), consistent with the inability of HP1β to form droplets with DNA (Fig. 3a). The DNA compaction behaviour of HP1α (BPM) was severely attenuated (Fig. 3j and Extended Data 9b, d), indicating that, like the ability to phase-separate and bind DNA, DNA compaction by HP1α involves this basic patch in the hinge.

On the basis of these data, we propose that phosphorylation and DNA binding have related roles. In this model, the phosphorylated residues of the NTE in one dimer make electrostatic interactions with basic residues in the hinge of another dimer to generate higher-order oligomers. These NTE–hinge interactions could help relieve CTE-mediated auto-inhibition and generally allow exposure of additional residues for making higher-order interactions (Fig. 2c and Extended Data Fig. 8c). Binding of DNA to the hinge could analogously displace the CTE in wild-type HP1α, thereby exposing new interaction surfaces. In addition, binding of multiple wild-type HP1α molecules to DNA could increase the local concentration of wild-type HP1α, possibly further promoting higher-order HP1α interactions and droplet formation. Alternatively, bridging of two regions of DNA by one wild-type HP1α dimer could locally alter DNA conformation in a manner that promotes the binding of additional wild-type HP1α dimers, without higher-order oligomerization. Our model also implies that NTE phosphorylation should compete with DNA binding and helps explain why DNA compaction by nPhos-HP1α is less cooperative and slower compared to that observed with wild-type HP1α (Extended Data Fig. 8c). Consistent with such competition, previous work has shown that NTE phosphorylation of HP1α reduces DNA binding⁵.

We next investigated how known nuclear components interact with phase-separated HP1α. We used Cy3-labelled components to visualize and measure their partitioning into nPhos-HP1α droplets (Fig. 4a). Core nucleosomes, 147 bp double-stranded DNA and aurora B kinase, a known interaction partner of HP1α, all localized within the
Figure 4 | Partitioning of specific macromolecules into HP1α phase and behaviour of HP1α molecules in cells. a. Micrographs of phase separated nPhos-HP1α droplets with either Cy3 labelled or YOYO-1 labelled macromolecules visualized using Cy3 fluorescence or YOYO-1 fluorescence, respectively. Scale bar, 50 μm. b, NIH3T3 cells transduced with Cy3-labelled HP1 proteins and classification of puncta distribution. Top right, average number of distinct puncta per cell. Bottom right, percentage of cells that have at least one large puncta. A large puncta is defined as having a diameter >5 μm in any direction within a z projection. Scale bars, 10 μm. Error bars represent s.e.m. nPhos-HP1α, n = 38; HP1α(CSDm), n = 26. c, Model for the role of regulated phase separation in chromatin organization. Swi6 has demonstrated such higher-order oligomerization and its importance for silencing in vivo. To date however, higher-order oligomerization of human HP1 proteins has not, to our knowledge, been reported. Instead, previous work suggests that dimers of HP1α and HP1β bridge nucleosomes and compact chromatin. Here we show that higher-order oligomerization of HP1α can be promoted by NTE phosphorylation, could provide qualitatively different means of regulating heterochromatin. For example, phase separation coupled to DNA binding may be used when rapid chromatin compaction is needed. Notably, wild-type HP1α compacts λ-DNA at rates around 15 times faster than its own capsid packaging motor, although against smaller forces. Our findings provide experimental evidence for such a possibility. A role for phase separation in HP1 heterochromatin is also suggested by work in Drosophila. Most simply, phase-separated HP1α droplets allow the means to physically sequester and compact chromatin while enabling recruitment of repressive factors.

The results in Fig. 4a raised the possibility that phase separation helps compartmentalize heterochromatin components in cells. We therefore investigated how the phase-separation behaviour of HP1α correlates with its localization within the nucleus. We directly delivered chemically labelled HP1α proteins into NIH3T3 cells using the Chariot delivery system. We used direct protein delivery to ensure high levels of NTE phosphorylation and because genetically encoded GFP tags inhibit phosphorylation, could provide qualitatively different means of regulating heterochromatin. For example, phase separation coupled to DNA binding may be used when rapid chromatin compaction is needed. Notably, wild-type HP1α compacts λ-DNA at rates around 15 times faster than its own capsid packaging motor, although against smaller forces. Our findings provide experimental evidence for such a possibility. A role for phase separation in HP1 heterochromatin is also suggested by work in Drosophila. Most simply, phase-separated HP1α droplets allow the means to physically sequester and compact chromatin while enabling recruitment of repressive factors.

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However, many fundamental questions remain: what is the nature of the physico-chemical environment within phase-separated heterochromatin; how do other heterochromatin components alter this environment; and is a predominant HP1α conformation maintained in the phase-separated state? Building on the approaches used to study other phase-separated cellular bodies will provide appropriate methods to address these questions.15,23

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions A.G.L. developed the overall experimental plan with guidance from G.J.N. and carried out the majority of the experiments and their interpretation. D.E. developed and implemented new software for analysing the SAXS data and helped conceive further experiments, M.J.T. performed and analysed the cross-linking and phosphate mapping mass spectrometry experiments, J.B.J. performed the native mass spectrometry experiments with guidance from A.L.B. D.A.A. provided guidance on SAXS experiments. M.K. performed and analysed the DNA curtains experiments and S.R. oversaw their design and interpretation. G.J.N. and A.G.L. wrote the bulk of the manuscript with major contributions from S.R. G.J.N. oversaw the overall project.

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METHODS

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment.

**Protein purification.** Full-length human HP1α was cloned into a pBH4 expression vector, mutants were made using site-directed mutagenesis following the fastochoning protocol and proteins were purified from *E. coli* 

Phosphorylated HP1α was obtained by co-expression with the catalytic subunits of CDK1 in a pRSF-Duet vector. HP1α proteins were purified from *E. coli* Rosetta (DE3) strains as follows. Cells were grown to OD 0.4 at 30 °C in 2XLB with 50 μg ml⁻¹ carbenicillin and 25 μg ml⁻¹ chloramphenicol. For co-expression with the pRSFduet CKII plasmid, 25 μg ml⁻¹ kanamycin was added. Cells were then grown to OD 0.8 at 18 °C and induced with 0.3 mM isopropyl-β-D-thiogalactopyranoside for 14 h. Harvested cells were resuspended in lysis buffer (1× PBS pH 7.2, 300 mM NaCl, 10% glycerol, 7.5 mM Imidazole, and protease inhibitors phenylmethanesulfonyl fluoride, pepstatin A, aprobin, and leupeptin). Following lysis in an Emsuliflex (Avestin), cell debris was removed by centrifugation at 25,000g for 35 min. Clarified lysate was incubated with Cobalt-NTA affinity resin (Clontech) for 40 min at 4 °C. Resin was then washed with approximately 50 ml of lysis buffer and eluted with 20 mM HEPES pH 7.2, 150 mM KCl, and 400 mM imidazole. Proteins were cleaved overnight with 3 mg ml⁻¹ TEV protease while dialyzing into 20 mM HEPES pH 7.2, 150 mM KCl, 3 mM DTT to remove imidazole. Protein was then injected on a Mono-Q 4.6/100 PE anion exchange column (GE), was washed, and eluted with a 120–800 mM KCl gradient over 15 column volumes. Protein was then concentrated in an Amicon Ultracel-10k spin concentration before injection on a Superdex 75 16/60 size-exclusion column run with storage buffer (20 mM HEPES pH 7.2, 200 mM KCl, 1 mM DTT, 10% glycerol). Proteins were concentrated to about 1 mM in Amicon concentrators before flash freezing in LN₂. Concentrations were measured by UV absorbance at 280 nm and using the calculated extinction coefficient, ε = 29,405. Removal of the 14-amino-acid CTE in the context of unphosphorylated HP1α protein (HP1αΔCTE) leads to rapid degradation and an unstable protein. However co-expression of HP1α protein with CKII stabilizes the protein (nPhos-HP1αCTE). The nPhos-HP1αΔCTE protein was concentrated by dialysis against 35,000 Mₗ lutelyene glycol at 25 °C owing to its propensity to form hydrogels during spin concentration. Wild-type and phosphorylated proteins were subsequently concentrated in this manner to ensure no artefacts were observed by analytical ultracentrifugation (AUC).

Human TFIIb and aurora B DNA sequences were ordered in gBlocks from IDT and cloned into the pBHH vector. Proteins were purified similar as described above, though a Mono-S cation exchange column was used in place of the Mono-Q.

The nPhos-HP1αΔ mutant has serine residues in the hinge mutated to alanines, such that it can only be phosphorylated in the NTE. By contrast, the nPhos-HP1αΔ mutant has serine residues of the NTE mutated to alanines such that it can only be phosphorylated in the hinge.

**Sedimentation velocity analytical ultracentrifugation.** Proteins were dialyzed into 20 mM HEPES pH 7.2, 75 mM KCl, and 1 mM DTT overnight at 4 °C. Concentrations were checked by UV absorbance at 280 nm. The samples were incubated at the appropriate temperature for 50 min in a pre-equilibrated rotor under vacuum. Runs were performed at 50,000 r.p.m. for 8–10 h in a Beckman XL-A analytical ultracentrifuge. Scans were collected at 250 or 280 nm with a radial step size of 0.003 cm at approximately 60-s intervals. Runs were completed in triplicate to ensure no experimental artefacts were incorporated into the analysis. SV analysis was done with SEDFIT/SEDPHAT (NIH) software and plots were generated using GUSSS20. Experimental Parameters were calculated using the Sednterp software and were as follows: HP1α, 0.72820 HP1β, 0.72794; buffer density, 1.002; buffer viscosity, 0.0089. Attempts to assess the oligomerization properties of HP1α protein (HP1αΔCTE) were inconclusive as the lowest concentrations of nPhos-HP1αΔCTE that are detectable by AUC displayed substantial phase-separation.

**SEC-MALs.** Proteins were injected on a Shodex KW-803 column at 25 °C. Buffer was 20 mM HEPES pH 7.2 and 75 mM KCl with 1 mM DTT. Samples were run again at pH 6.8 to exclude possible influence of column interaction on elution volume. BSA standard was run previous to injections for internal calibration.

**SAXS measurement and analysis.** All samples were dialyzed before measurement to obtain a matching buffer. SAXS experiments were done using an in-house instrument (Anton Paar SAXSSEXS MC2) with line-collimated illumination. Buffer-subtracted data was then corrected for aggregation using averaging programs. Further data processing was done using custom software written in Python and Fortran90 using the NumPy, SciPy, Matplotlib and PyQtGraph libraries (UCSF SAXS, code available at https://github.com/delnatan/UCSFsaxs). The software implements a Bayesian algorithm for determining the optimal maximum particle diameter and smoothing factor to fit the scattering data to a pairwise interatomic distance distribution, \( P(r) \). This was done by solving a regularized least squares equation (in matrix form, matrices are bolded):

\[
J(\alpha) = \| A x - b \|^2 + \alpha \| L_{xx} x + L_{zb} z \|^2
\]

Where \( J(\alpha) \) is the least-squares objective function, and \( \alpha \) is the smoothing factor that balances data overfitting and sum of squared enrichment. For aggregation, \( P(r) \) consists of zeros everywhere except on the first and last element, which is set to 1. Matrix \( P \) penalizes zero values for the end-point of the \( P(r) \) with an arbitrarily large value, \( \beta \) (which is set to be 100%). This optimization was done using the non-negative least squares (NLLS) routine from SciPy.Optimize.

Models in Fig. 2b (inset) were generated using the ensemble optimization method from SAXS envelopes and two crystallized HP1α domains CSD(3Q6S) and CD(3FDT), and the structures represent two possible conformations for the HP1α dimer.

**Cross-linking mass spectrometry and phosphopeptide analysis.** Wild-type HP1α was cross-linked with 1 mM BS3 for 5 min, at room temperature and quenched by addition of 10 mM Tris base. Samples were resolved by SDS–PAGE using precast 4–20% Bis-Tris gels and stained by Coomassie. Gel bands corresponding to monomeric and dimeric HP1α were excised and trypsin digested. Extracted peptides were desalted and analysed on an Orbitrap Velos (Thermo Scientific) mass spectrometer coupled with a nanoelectrospray ion source and NanoAcuity UPLC system (Waters). Peptides were separated on a 15 cm x 75 μm ID PepMap C18 column (Thermo) using a 60-min gradient from 3–28% acetonitrile containing 0.1% formic acid. Precursor MS5 scans were measured in the Orbitrap analyser scanning from 350–1,800 m/z (mass resolution: 30,000). The six most intense triply charged or higher precursors were isolated in the linear ion trap (isolation window: 4 m/z), dissociated by HCD (normalized collision energy: 30), and the product ion spectra were measured in the Orbitrap (mass resolution: 7,500). A 30 s dynamic exclusion window was applied. Three technical replicates were analysed per condition.

Peaklists were generated using Proteome Discoverer (Thermo) and searched using Protein Prospector 5.14.20 (ref. 33). An initial unbiased search of the data against SwissProt (535,248 entries from March 21, 2012) showed the sample to consist of predominantly human HP1α by spectral counts (>$90%). Subsequent searches for cross-linked peptides were restricted to the sequence of human HP1α and the next 15 most abundant proteins which consisted of minor contaminants from *E. coli* as well as residual TEV protease. Additionally, randomized versions of these proteins were included in the search database for false discovery rate (FDR) analysis. Cross-linking searches were performed against the 85 most intense product ion peaks with the following parameters: enzyme specificity: tryptic, 3-missed cleavages; mass tolerance: 8 p.p.m. (precursor), 25 p.p.m. (product); cross-linking reagent: DSS/BS3; variable modifications: phosphorylation at Ser/Thr/Tyr, oxidation at Met, N-terminal Glu to pyroGlu conversion, loss of Met and/or acetylation at the protein N terminus, dead-end modification of lys by semi-hydrized BS3; constant modification: carbamidomethylation of Cys. Cross-linked peptides were reported with a Prosporctor score greater than 20 and score difference greater than 6.5 corresponding to an FDR below 1%. Inter-molecular HP1α cross-links were putatively assigned by taking the set of cross-linked residue pairs uniquely identified in the dimer gel bands when compared to the corresponding monomer bands.

Phosphopeptide analysis of N-terminal phosphorylated HP1α was performed on a Q-Exactive Plus Orbitrap instrument (Thermo Scientific) from in-solution trypsinized sample. Peptides were analysed directly (without phospho-enrichment). Three technical replicates were run, and phosphopeptides were searched for as variable modifications on Ser and Thr residues using Protein Prospector.

**Native MS.** Native mass spectrometry was carried out using the Exactive Plus EMR instrument (Thermo Scientific) that was externally calibrated using a 5 μg ml⁻¹ CsI solution prepared in water. Prior to analysis, the protein samples were buffer-exchanged into 150 mM ammonium acetate, pH 7.5 using MicroRiospin-6 columns (Bio-Rad) that had been pre-equilibrated in the same buffer. Protein samples were introduced into the mass spectrometer using offline Au/Pd-coated borosilicate emitters (NanoES Spray Capillaries, Medium, E3880, Thermo Scientific) at a flow rate of 10–40 nl per min. Spectra were acquired over the range of 500–20,000 m/z in positive ion mode, were averaged, and then exported for deconvolution and subsequent generation of the zero-charge mass values using PeakSeeker and Unidec. Samples were analysed with the following experimental parameters: spray voltage (0.8–1.5 kV), injection flattople = 5;
inter flatop lens = 5; bent flatop = 5; transfer multipole = 2; C-trap entrance lens = 2, source DC offset (25 V), fragmentation energies (CE = 25 and CID = 65), injection time (200 μs), trapping gas pressure (7.5, 37,1000, 250 °C), 5-ε RF levels (200 V), microscans (10), and AGC (1 × 10^6).

Saturation concentration measurements. These experiments were carried out at room temperature (approximately 22–24 °C) in a buffer containing 75 mM KCl, 20 mM HEPES pH 7.2, and 1 mM DTT.

Centrifugal spin-down assay. In this method we spun the phase separated material at 10,000 × g in a two-phase solution, with the low concentration phase existing in the top layer and the high concentration HP1β phase existing at the bottom of the tube (Fig. 1e). The concentration of HP1β in the top layer was measured to obtain the saturation concentration of phase separation. 10-μl samples were incubated at the appropriate temperature for 5 min then spun at 10,000 g for 5 min in a tabletop centrifuge. 4 μl of supernatant was removed for A280 measurements in triplicate on a nanodrop instrument. Pipetting or vortexing returned the sample to a turbid solution (Supplementary Video 6).

Turbidity method. This assay measures the turbidity of the phase-separated solution by absorbance at 340 nm (Fig. 1c). In this method, the saturation concentration was defined by the concentration at which the turbidity was at a half maximal value. Serial dilutions were performed in 12 tube PCR strips. 20 μl of sample was then added to a clear bottom 384-well plate (Corning) and absorbance was read at 340 nm in a Spectramax M5 plate reader. For peptide addition, 1 μl of peptide at the appropriate concentration was added, mixed, and incubated for 5 min before reading.

Microscopy. Microscopy of the droplets was done on a Leica Axiovert 200 M microscope using a 10 × or 40 × air objective. For Cy3 detection samples were excited with a 520 nm laser and a 560 × 20 nm emission filter. A custom nitrogen curtain microscope was used for cooling experiments to eliminate condensation. Imaging of YOYO-1-labeled arrays and NIH/3T3 cells was performed on a Nikon Eclipse confocal microscope through a 10 × air objective. For Cy3 detection samples were excited with a 520 nm laser and a 560 × 20 nm emission filter. A custom nitrogen curtain microscope was used for cooling experiments to eliminate condensation. Imaging of YOYO-1-labeled arrays and NIH/3T3 cells was performed on a Nikon Eclipse microscope using a 10 × air objective. For Cy3 detection samples were excited with a 520 nm laser and a 560 × 20 nm emission filter.

Protein labelling. A GSCKK peptide tag inserting a labile cysteine was cloned in C-terminal to the sequence of HP1α. Cys133 was mutated to serine to favour C-terminal labelling. Proteins were dialysed into labelling buffer (20 mM HEPES pH 7.2, 350 mM KCl, 0.5 mM TCEP) and concentration adjusted to 200 μM of protein. Each Cys residue was labelled with YOYO-1 dye and visualized using a 488 nm laser. In this method we spun the phase separated material at 10,000 × g in a two-phase solution, with the low concentration phase existing in the top layer and the high concentration HP1β phase existing at the bottom of the tube (Fig. 1e). The concentration of HP1β in the top layer was measured to obtain the saturation concentration of phase separation. 10-μl samples were incubated at the appropriate temperature for 5 min then spun at 10,000 g for 5 min in a tabletop centrifuge. 4 μl of supernatant was removed for A280 measurements in triplicate on a nanodrop instrument. Pipetting or vortexing returned the sample to a turbid solution (Supplementary Video 6).

The appropriate concentration was added, mixed, and incubated for 5 min before reading.

Data availability. All relevant data are included in the main manuscript, Extended Data figures and Supplementary Information. Any additional data are available from the corresponding author upon reasonable request.
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37. Gallardo, I. F. et al. High-throughput universal DNA curtain arrays for single-molecule fluorescence imaging. Langmuir 31, 10310–10317 (2015).
Extended Data Figure 1 | Mass-spectrometric analysis of HP1α proteins. Cross-linking mass spectrometry of HP1α identifies extensive interactions between the CSD and the hinge region. a, Phosphorylation of HP1α occurs almost exclusively at the N terminus. Left, annotated HCD (higher energy collision dissociation) product ion spectra of a quadruply phosphorylated, doubly charged HP1α peptide at Ser11, Ser12, Ser13, and Ser14. Neutral loss of phosphoric acid from b-ions is indicated by b*. Right, relative occupancy of observed HP1α phosphorylation sites as estimated by spectral counting. 41.7% of product ion spectra from peptides containing serine at residues 11–14 were observed quadruply phosphorylated (393 of 943 spectra). An additional 32.9% (310 spectra), 12.8% (121 spectra), and 8.5% (80 spectra) were identified triply, doubly, and singly phosphorylated, respectively, while only 4.1% (39 spectra) were observed with no phosphorylation. By contrast, phosphorylation was observed at other positions (Ser45, Thr132 and/or Ser135, Thr145, and Thr188) with 1–2.5% occupancy (1,059, 2,243, 1,586, 1,042 total spectra observed for peptides containing these residues). b, Native MS charge state envelopes for wild-type, Phos- and nPhos-HP1α. c, Table with predicted and observed masses is also shown. The deconvoluted masses fit best to dimeric HP1α modified by eight phosphates in Phos-HP1α and nPhos-HP1α samples. d, Cross-links were identified by separating cross-linked HP1α by SDS–PAGE and excising bands corresponding to monomeric and dimeric HP1α. Putative inter-protein cross-links, diagrammed here, were identified by taking the set of cross-links that are unique to the dimer band (from three replicates). Only cross-links identified by four or more product ion spectra are shown for clarity.
Extended Data Figure 2 | Phase separation is an isoform-specific capability of phosphorylated HP1α that is perturbed by GFP fusion.

a, 1 μl of a 400 μM solution of each protein was spotted on a plastic coverslip and imaged at 10 ×. Scale bars, 50 μm. Buffer was 75 mM KCl, 20 mM HEPES pH 7.2, 1 mM DTT. Phos-HP1α is phosphorylated in the N terminus and hinge, nE-HP1α has the N-terminal serines replaced with glutamate, Phos-GFP–HP1α is a N-terminal GFP fusion phosphorylated in the N terminus and hinge, Phos-HP1α–GFP is a C-terminal GFP fusion phosphorylated in the N terminus and hinge, Phos-HP1α(BPM) has the KRK hinge sequence mutated to alanines and is phosphorylated in the N terminus and hinge, Phos-HP1α–KCK has a C-terminal GSKCK tag added and is phosphorylated in the N terminus and hinge. b, Complete comparison of saturation concentration measurements between spin-down assay (left) and 340 nm turbidity-based measurement (right), some data are repeated from Fig. 1.
Extended Data Figure 3 | Estimation of oligomeric potential by sedimentation velocity analytical ultracentrifugation. a, Representative sedimentation velocity runs from high-concentration HP1 samples. Percentage of the loaded sample higher than 6 S was quantified to estimate oligomeric species higher than a dimer. b, Table showing the comparison of high-concentration AUC runs. Average sedimentation coefficient was quantified by integrating from 1–20 S and higher order oligomers were estimated by integrating signal from 6–20 S. c, Analytical ultracentrifugation c(S) analysis of fully phosphorylated HP1α and the fully phosphorylated basic patch mutant. d, Analytical centrifugation c(S) analysis of fully phosphorylated HP1α and the fully phosphorylated HP1α/β chimaera (PhosNH-α/β chimaera). Representative traces from three independent experiments are shown in a–d (n = 3).
Extended Data Figure 4 | Estimation of HP1α dimerization affinity by isothermal calorimetry and analytical ultracentrifugation. a. Isothermal calorimetry data showing the measured dimerization $K_d$ for the HP1α CSD. The calculated $K_d$ is $1.1 \mu$M. b. An analytical ultracentrifugation isotherm used to estimate the dimerization $K_d$ for wild-type HP1α. Estimated $K_d$ for dimerization using an isodesmic association model is $1.12 \mu$M.
Extended Data Figure 5 | Scattering and Guinier fits of SAXS on wild-type and nPhos-HP1α show homogeneous populations. a, Raw X-ray scattering intensity of wild-type (blue points) and nPhos-HP1 (green points) at 3.5 mg ml$^{-1}$ (150 μM) concentration. Black lines are Fourier transforms of the fitted interatomic distance distribution, $P(r)$, with $\chi^2$ values of 1.186 and 1.199 for wild type and nPhos, respectively. b, Guinier plots of wild-type (blue points) and nPhos-HP1 (green points) at 150 μM. Black lines are linear fits to the data plotted as log intensity versus $q^2$. The range of data used in the linear fits extend up to $q \times R_g$ of 1.3. $R_g$ is radius of gyration and $q$ is scattering vector. The corresponding residuals for each fit are shown below as vertically shifted horizontal lines for clarity.
Extended Data Figure 6 | Phosphorylated HP1α elutes as an extended dimer when examined by SEC-MALS. a, Elution profiles of wild-type HP1 and nPhos-HP1 examined by SEC-MALS. The horizontal green, and blue lines correspond to the calculated masses for nPhos-HP1 and wild-type HP1, respectively. b, MALS trace of fully phosphorylated HP1α run under identical conditions to those in a.
Extended Data Figure 7 | Measuring shugoshin 1, lamin B receptor, H3K9me3 peptide affinity, and the effect of shugoshin peptide binding on oligomerization. a, b, Fluorescence anisotropy plots showing the $K_d$ measurements (values in $\mu$M next to symbols for wild-type versus HP1$\alpha$(CSDm)) for LBR and Sgo1 peptide binding to wild-type HP1$\alpha$ and the I163A CSD mutant (CSDm), which can no longer form dimers. c, Comparative analytical ultracentrifugation runs of approximately 50$\mu$M nPhos HP1$\alpha$ with and without 100$\mu$M shogushin or LBR. d, Fluorescence anisotropy plots with a 15-mer trimethylated H3K9 peptide showing the relevant HP1 isoforms can bind the nucleosome tail.
Extended Data Figure 8 | Effects of additional ligands on saturation concentrations. a, Bar graphs displaying the effects of 100 μM of the polyamine spermine along with the H3K9 and H3K9me3 peptides on phase-separation behaviour. b, Schematic of the assay used to quantify the partitioning of Cy3-labelled substrates into the two phases. The blue bars represent the total concentration of the labelled species before spin down; the orange bars represent the concentration of Cy3-labelled species remaining in the upper phase after spin down. The lower phase contains HP1α at a higher concentration than in the upper phase. Error bars represent standard error of the mean from three independent measurements. c, Model for phosphorylation or DNA-driven HP1α phase separation. Phosphorylation or DNA binding relieves intra-HP1 contacts and opens up the dimer. The location(s) of the intra- and inter-dimer contacts that change during this transition are not fully understood, but are predicted to involve interactions between the CTE, hinge and NTE.
Consequences of the interaction between HP1 and DNA. a, b, Wide-field TIRF microscopy images of DNA compaction by HP1β (a) and HP1α(BPM) (b) at different time points. Scale bars, 5 μm. c, d, Average kymograms for HP1β (c; n = 368) and HP1α(BPM) (d; n = 318) overlayed with fits for average compaction speed (dashed line) and standard deviation (solid lines). e, f, Individual kymograms showing compaction by wild-type HP1α (e) and nPhos-HP1α (f) at different protein concentrations.
Extended Data Figure 10 | Additional micrographs of NIH3T3 cells transduced with HP1. NIH3T3 cells transduced with 0.3 μg of HP1 proteins and imaged under identical conditions. a, nPhos-HP1α; b, HP1α (CSDm); c, wild-type HP1α.