Succinylation of H3K122 destabilizes nucleosomes and enhances transcription

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Abstract

Histone post-translational modifications (PTMs) are key players in chromatin regulation. The identification of novel histone acylations raises important questions regarding their role in transcription. In this study, we characterize the role of an acylation on the lateral surface of the histone octamer, H3K122 succinylation (H3K122succ), in chromatin function and transcription. Using chromatin succinylated at H3K122 in in vitro transcription assays, we show that the presence of H3K122succ is sufficient to stimulate transcription. In line with this, we found in our ChiP assays H3K122succ enriched on promoters of active genes and H3K122succ enrichment scaling with gene expression levels. Furthermore, we show that the co-activators p300/CBP can succinylate H3K122 and identify sirtuin 5 (SIRT5) as a new desuccinylase. By applying single molecule FRET assays, we demonstrate a direct effect of H3K122succ on nucleosome stability, indicating an important role for histone succinylation in modulating chromatin dynamics. Together, these data provide the first insights into the mechanisms underlying transcriptional regulation by H3K122succ.

Keywords acetylation; histones; succinylation

Subject Category Chromatin, Transcription & Genomics

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Introduction

The organization of DNA into chromatin allows the DNA to be stored within the nucleus of a cell, while at the same time safeguarding, when necessary, its accessibility. Histones are the main protein component of chromatin (Luger et al., 1997). Multiple studies have described how different post-translational modifications (PTMs) on the histone N-terminal tails, which extend from the nucleosomal core, indirectly regulate chromatin function, e.g., by recruiting or excluding binding proteins (Jenuwein & Allis, 2001). However, more direct effects of histone PTMs are still poorly understood (Kouzarides, 2007; Lawrence et al., 2016).

Histone post-translational modifications can occur, apart from the N-terminal tails, also within the globular domains of histones (Mersfelder & Parthun, 2006; Tropberger & Schneider, 2013). In addition to acetylation, multiple different histone acylations have been identified within the nucleosomal core, including butyrylation, crotonylation, malonylation, and succinylation (Tan et al., 2011; Tessarz & Kouzarides, 2014; Kebede et al., 2015). All these modifications change the charge of the side chain of the modified lysine residue and thus can, depending on their position, potentially alter the interactions with other histones, neighboring nucleosomes or with the nucleosomal DNA (Mersfelder & Parthun, 2006).

Several studies have shown that lysine acetylations within the globular domain of histones can affect nucleosome stability (Neumann et al., 2009; Tropberger et al., 2013; Di Cerbo et al., 2014) and regulate chromatin-dependent processes (Lawrence et al., 2016). Namely, the acetylation of histone H3K122 (H3K122ac), located on the so-called lateral surface of the histone octamer, can directly increase transcriptional output (Tropberger et al., 2013). Recently, Bao et al. (2019) described that a new type of acylation, glutarylation of lysine 91 on histone H4 (H4K91glut), impairs the interaction between the H3-H4 tetramer and the H2A-H2B dimers, resulting in the destabilization of the nucleosome.

Lysine succinylation was first described in Zhang et al. (2011), and a year later, the succinylation of H3K122 (H3K122succ) was identified by mass spectrometry (Xie et al., 2012). It converts, in contrast to the majority of other acylations, the positive charge of the lysine residue to a negative charge and increases the steric hindrance due to its larger volume. Lysine succinylation could therefore affect interactions involving the lysine side chain. Interestingly, H3K122 is positioned on the dyad axis of the nucleosome,
where its side chain is in contact with the nucleosomal DNA. Any perturbation of the dyad axis conformation has the potential to affect nucleosome dynamics, as this is the region where the nucleosomal DNA and the histone octamer are in closest proximity and their interactions are the strongest (Cutter & Hayes, 2016). Previously, the desuccinylation of H3K122succ by sirtuin 7 (SIRT7) has been shown to be associated with DNA double strand break repair and genome stability (Li et al, 2016); however, the role of H3K122succ in transcription is still unclear.

The important location of H3K122 and the reversal of the charge of lysine 122 by succinylation prompted us to study the possible role of H3K122 succinylation in transcription. Here, we find by using our new antibodies H3K122succ enriched on the promoter regions of actively transcribed genes and we identify novel enzymes responsible for the establishment and removal of this mark. By comparing site-specific succinylation chromatin with unmodified chromatin, we demonstrate that H3K122succ is sufficient to increase transcriptional output. We reveal that H3K122succ destabilizes nucleosomes and, thus, gain insights into potential mechanisms on how H3K122succ acts.

Results and Discussion

In order to specifically study H3K122succ, we raised and affinity purified three rabbit polyclonal antibodies (ABs) against H3K122succ. All three ABs recognized in peptide dot blot assays the immunizing H3K122succ peptide with no cross-reactivity observed to any other peptide tested (Figs 1A and EV1A and B). In immunoblot, all ABs recognized H3 specifically and this signal was competed away by the H3K122succ peptide (see Fig EV1C for an example). Succinylated H3, but not equal amounts of unmodified H3 nor H3 mutated on K122 to an arginine (Fig 1B) were detected. Additionally, also in native chromatin (ChIP input) H3 was specifically recognized (Fig EV1D) and this reactivity did not depend on the presence of the H3 N-terminal tail (Fig EV1E). Taken together, these data demonstrate the specificity of our ABs for H3K122succ in native chromatin as well as on histone peptides.

H3K122succ is enriched at transcription start sites

We next investigated the genomic distribution of this modification by carrying out native chromatin immunoprecipitation followed by deep sequencing (ChIP-seq) on the human MCF7 breast cancer cell line using all three independent antibodies against H3K122succ. We identified in our ChIP 4745, 4788, and 4346 peaks with AB #1, AB #2, and AB #3, respectively. Reads enrichment in all H3K122succ peaks were highly correlated (Spearman’s ρ ≥ 0.9, Fig 1C). We found that H3K122succ peaks obtained in our ChIP experiments with all three antibodies were preferentially located in gene promoter regions (≥ 50%) and specifically close to and at transcription starting sites (TSS), with 46–49% of H3K122succ peaks at TSS regions, compared to only 5% in a random peak set (Fig 1D). This localization of H3K122succ peaks to TSS regions suggests that H3K122succ may play a role in transcriptional regulation.

H3K122succ marks active genes

Since our three antibodies gave consistent results, we focused on AB #2 for all subsequent analysis. A first visual inspection of the genome browser tracks revealed an enrichment of H3K122succ around the TSS of active genes together with other histone modifications (Fig 2A). Approximately three-quarters (76.6%) of the genes with H3K122succ ChIP-seq peaks detected at their TSS have CpG island (CGI) containing promoters and more than one-third (35.2%) are housekeeping genes (HKGs; Fig 2B). Furthermore, genome wide our H3K122succ ChIP peaks co-localized with peaks for two hallmark of active promoters (Santos-Rosa et al, 2002)—H3K4me3 and H3K9ac (Fig 2C). More specifically, at TSS 70–80% of the H3K122succ peaks we detected were also marked with “active” histone tail modifications such as H3K4me3, H3K9ac, and H3K27ac (Fig 2D). When comparing enrichment profiles at the positions of our H3K122succ ChIP peaks with datasets for the histone acetyltransferase p300 and the acetylated histone variant H2A.Z, previously shown to occur on active regulatory regions (Giaimo et al, 2019), we found enrichment of p300 and H2A.Zac at the majority of H3K122succ peaks (Fig EV2A). Additionally, the H3K122succ peaks overlapped with open chromatin regions identified both by FAIRE-seq and DNase-seq, at TSS (Fig 2E) and overall (Fig EV2B). Taken together, this suggests that H3K122succ marks promoters of genes in an open and “active” chromatin environment. Indeed, comparison with RNA-seq data revealed that the H3K122succ ChIP-seq read density at TSS correlated with the mRNA levels (Fig 2F). Thus, local H3K122succ enrichment can be indicative of the steady-state mRNA level of the respective gene.

We next examined the relationship between gene expression level and the presence of H3K122succ alone and/or in combination with H3 tail modifications (K4me3, H9ac, and K27ac; Fig 2G). In general, we found that genes with H3K122succ peaks at their TSS (peaks ± 500 bp around TSS) were more expressed than genes without H3K122succ peaks. Genes where we detect H3K122succ in combination with the three other active marks (H3K9ac, H3K27ac, and H3K4me3) showed the highest expression level of all the groups tested, higher than ones without H3K122succ peaks. To investigate

Figure 1. H3K122succ peaks localize to gene promoters.

A Peptide dot blot array probed with anti-H3K122succ AB #2. Peptides with indicated modifications were spotted on membranes. Peptide sequences can be found in Table 1. For other two antibodies, see Fig EV2A and B.

B Immunoblot analysis with anti-H3K122succ AB #2 on indicated histone octamers. Octamers with H3K122succ were specifically recognized.

C Pairwise scatterplots of H3K122succ read counts within peaks from ChIP experiments using all three H3K122succ antibodies (AB #1, AB #2, and AB #3). Note high degree of correlation as indicated by the Spearman correlation coefficient (ρ ≥ 0.9).

D Genomic feature distribution of H3K122succ peaks obtained with the three H3K122succ antibodies. The percentage of peaks in the 1st exon, remaining exons, intergenic regions, 5′ intron, other introns, promoter/ TSS (≤ 1 kb from TSS), promoter (1–2 kb), and 3′ UTR is displayed. A random distribution of peaks is shown for comparison (see Materials and Methods for additional information on the random distribution).
Figure 1.
whether the higher transcription levels simply reflected higher acetylation levels on the histone tails, we used H3K27ac as an example and compared the number of H3K27ac read in these different gene groups (Fig EV2C). These results show that the gene group associated with the highest transcription levels (H3K122succ + 3 histone tails) did not have the most H3K27ac reads, indicating that the higher expression of genes enriched in H3K122succ cannot simply be explained by, e.g., higher H3K27 acetylation levels.

H3K122 can not only be succinylated but also acetylated (Tropberger et al., 2013). We therefore compared the genes we associated with H3K122succ at their TSS with the ones with H3K122ac peaks (Fig EV2D) and found an overlap of 61%. Overall, the number of H3K122succ peaks we detected was lower than the ones for H3K122ac what is in line with the lower overall abundance of histone succinylation compared to histone acetylation (Xie et al., 2012) and could suggest a more specific function for histone succinylation compared to acetylation. However, we cannot rule out that different antibodies affinities or linear ranges of antibodies contribute to the differences in the peak numbers. Interestingly, we found that genes at which we detect H3K122succ peaks at their TSS were in general higher expressed than those with H3K122ac peaks (Fig EV2E). Further studies will be needed to determine whether and how these marks are functionally connected.

Taken together, these results suggest that H3K122succ may promote transcriptional activation and could play a role in creating an optimal chromatin environment allowing for maximal transcriptional output together with other histone modifications.

**p300/ CBP can establish H3K122succ**

To address the role of H3K122succ in transcriptional regulation, we first sought to identify the enzyme(s) catalyzing the succinylation of H3K122. It has been shown that histone acetyltransferases (HAT) can succinylate proteins (Hirschey & Zhao, 2015; Wang et al., 2017). Therefore, we depleted selected HATs by small interfering RNA (siRNA; Fig EV3A) and assayed the levels of H3K122succ by immunoblot (Figs 3A and EV3B). As shown in Fig 3A, upon double knock down (KD) of p300 and its parologue CREB binding protein (CBP), we observed decreased levels of H3K122succ. Additionally, we treated MCF7 cells with curcumin, a p300/ CBP inhibitor (Morimoto et al., 2008) that promotes degradation of p300 and CBP. Indeed, upon curcumin treatment we detected decreased levels of H3K122succ (Fig EV3C). To demonstrate that p300 can catalyze H3K122succ in vitro, we performed next succinyltransferase assays with recombinant p300 on recombinant histone octamers (rOctamers; Figs 3B and EV3D), as well as on peptides spanning the region around H3K122 (Fig 3C) as substrates. In these assays, recombinant p300 can indeed succinylate H3 at position K122. Next, we compared how adding unlabeled acetyl-coA (ac-CoA) or succinyl-CoA (succ-CoA) affects radioactive in vitro histone acylation assays (Fig EV3E). Although succ-CoA can compete, ac-CoA has a stronger effect on signal intensity. Thus, we suggest that the local availability of CoAs, e.g., in metabolic microniches (Katada et al., 2012), could determine if p300 acts as acetyl or succinyltransferase.

In summary based on our in vitro and in vivo assays, we conclude that p300 can succinylate a specific histone residue. This is in line with previous findings that p300 can act as a lysine succinyltransferase in vitro (Hu et al., 2014), as well as with the co-localization between H3K122succ and p300 enrichment that we observed (Fig EV2A). Whether (and which) other enzymes contribute to H3K122 succinylation and whether p300 can succinylate additional residues in histones is still an open question.

**Sirtuin 5 can desuccinylate H3K122**

Sirtuins have been identified as protein desuccinylases (Maurer et al., 2012). Specifically sirtuin 5 (SIRT5) has been reported as a protein desuccinylase (Du et al., 2011) and, more recently, SIRT7 as an enzyme that can desuccinylate H3K122succ during DNA damage response (Li et al., 2016). Although SIRT5 is mainly a mitochondrial enzyme, it can also be found in the cytosol and in the nucleus (Hirschey & Zhao, 2015).

To explore the role of SIRT5 in the removal of H3K122succ, we performed in vitro desuccinylase assays on H3K122succ peptides with recombinant SIRT5 and SIRT7 (Figs 3D and E, and EV3F and G). As expected, SIRT7 can desuccinylate H3K122; however, SIRT5 can also desuccinylate H3K122. To confirm the activity of SIRT5 in vivo, we compared the levels of H3K122succ on histones isolated

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**Figure 2. H3K122succ is enriched at active TSS.**

A IGV browser snapshots showing the distribution of reads around the TSS of the actively transcribed PPM1D gene (chr17:60,587,313-60,643,064; top panel) and of the inactive CD40 gene (chr20:46,114,180-46,129,721; bottom panel) for the indicated histone modifications, input, and RNA-seq.

B Venn diagrams showing the overlap between genes with H3K122succ peaks detected at their TSS regions and genes with CpG island associated to their promoters (CpG) (upper panel), as well as housekeeping genes (HKG) (bottom panel). P-values were calculated through hypergeometric tests.

C Venn diagram showing the number of H3K122succ peaks detected overlapping with H3K4me3 peaks and H3K9ac peaks.

D Correlation matrix displaying the degree of overlap between H3K122succ, H3K4me3, H3K9ac, and H3K27ac peaks from our ChIP experiments at TSS (peaks within ± 500 bp of TSS). Total number and fraction of overlapping peaks are displayed.

E Correlation matrix displaying the overlap between our H3K122succ ChIP-seq peaks and DNase-seq and FAIRE-seq enriched regions at TSS (peaks within ± 500 bp of TSS). Total number and percentage of overlapping peak regions are displayed.

F H3K122succ occupancy profile over a meta gene as a function of expression level (based on RNA-seq quartiles: Q1—least expressed, Q4—most expressed). All genes were normalized to the same length.

G Boxplots visualizing expression values of genes (reads per kb; RPK) with peaks for indicated different combinations of histone PTMs at their TSS. Histone modifications are (from left to right): none (no PTM, black); only H3K122succ (green); H3K122succ plus one tail modification (H3K4me3, or H3K9ac, or H3K27ac, green and light orange); H3K122succ plus two tail modifications (green, yellow, and light orange); H3K4me3 and H3K9ac and H3K27ac without H3K122succ (yellow, light orange, and dark orange); H3K122succ and all three tail modifications (green, yellow, light orange, and dark orange). The number of genes within the groups is indicated below each boxplot. Boxes indicate the range between the first and third quartile, the central line depicts the median, and the whiskers span the range of the data while extending no further than 1.5 times the interquartile range. Each distribution was compared to the “no PTM” distribution. P-values were calculated by Wilcoxon signed-rank test.
Figure 2.

A

B

C

D

E

F

G

Expression quartile

Mean density (tag/50bp)

-5Kb TSS TTS +5Kb

Figure 2. [Image]
Figure 3. p300/ CBP can succinylate H3K122, SIRT5 and SIRT7 desuccinylate it.

A H3K122succ levels (detected by immunoblot), relative to the scramble siRNA control, upon depletion of indicated HAT enzyme(s) (GCN5, pCAF, p300, and/or CBP) from MCF7 cells by siRNA. The bar graph shows a representative experiment (for biological replicate see Fig EV3B).

B In vitro succinyltransferase assay on recombinant histone octamers as substrate. After incubation with p300 and suc-CoA, the H3K122succ levels were assessed by immunoblot. Ponceau staining of membrane is shown as a loading control. For additional controls, see Fig EV3D.

C In vitro succinyltransferase assay on unmodified peptides spanning H3K122. Samples were incubated with increasing amounts of p300 (0–160 ng) in the presence of radiolabeled suc-CoA. The radioactivity incorporated was quantified by liquid scintillation counting. The plot displays average counts per minute (CPMs), ± SD and individual datapoints of two independent replicates. Note that increasing enzyme concentrations resulted in increased succinylation.

D, E In vitro desuccinylation assay on H3K122succ peptides with recombinant SIRT7 (D) and SIRT5 (E). Unmodified H3K122 peptide was used as control (left dot). Peptides were probed with H3K122succ AB #2 after incubation with sirtuins. For additional controls, see Fig EV3F and G.

F H3K122succ levels in Sirt5-KO cells. Histones were acid extracted from wild type (WT) and Sirt5-KO MEFs, and the levels of H3K122succ determined by immunoblot. The level of H3K122succ was normalized to histones extracted from WT MEFs. Average, ± SD, and individual datapoints of three biological replicates are shown. P-value was calculated by two-tailed paired t-test.
from WT and Sirt5-KO mouse embryonic fibroblasts (MEFs (Park et al., 2013); Fig 3F). In accordance with our in vitro desuccinylase assay, the levels of H3K122succ were higher in the Sirt5-KO MEF cell line compared to the WT, identifying SIRT5 as a novel enzyme that can desuccinylate H3K122 both in vitro and in vivo.

**H3K122succ stimulates transcription in vitro**

The location of H3K122 at the dyad axis prompted us to investigate the effect of H3K122succ on transcription in a well-controlled system. For this purpose, we performed in vitro transcription (IVT) assays (Orphanides et al, 1998) on unmodified chromatin templates or chromatin site-specifically succinylated on H3K122. Briefly, we expressed WT H2A, H2B, and H4 in E. coli and purified the histones. Site-specifically modified H3K122succ, as well as unmodified H3, were generated by protein synthesis. We assembled chromatin on a pG5-MLP plasmid (Dignam et al, 1983), with the help of the nucleosome assembly protein (NAP1) histone chaperone and the ATP-dependent chromatin assembly factor (ACF) remodeling complex (Ito et al, 1997). To control for chromatin assembly efficiency, we performed MNase digestion and sucrose gradient fractionation on the assembled chromatin (Fig EV4A and B). The scheme of the general in vitro transcription reaction is shown in Fig 4A.

First, we investigated whether adding suc-CoA to the reaction can stimulate transcription and how its effect compares to adding ac-CoA, which is required for efficient transcription on chromatinized templates. For this, we performed in vitro transcription on unmodified chromatin in the presence of ac-CoA or suc-CoA or with no CoA. Our results show that not only ac-CoA but also suc-CoA can stimulate transcription (Fig 4B and C). Next, in order to study the impact of site-specific H3K122succ on transcription, we performed in vitro transcription on unmodified and H3K122succ chromatin. Remarkably, we observed in our assays that the presence of H3K122succ stimulated transcription (compared to unmodified chromatin) by approximately 1.6-fold (Fig 4D and E). Together, these results add H3K122succ to the so far very few examples of site-specific histone modifications whose presence is sufficient to stimulate transcription.

It has been previously shown that most individual acetylations of the H3 tail residues do not affect transcription in vitro; however, the acetylation on the lateral surface of the histone octamer at H3K122 can stimulate transcription by approximately 1.7-fold (Tropberger et al, 2013). Thus, the two H3K122 modifications—acetylation and succinylation—stimulate transcription in vitro to a similar extent.

**Succinylation of H3K122 decreases nucleosome stability**

We hypothesized that the stimulation of transcription by H3K122succ could be due to a weakening of DNA histone interactions and, thus, destabilization of nucleosomes. To investigate such a mechanism, we explored the effects of H3K122succ on nucleosome stability by single molecule Förster resonance energy transfer (smFRET) assays. We inserted FRET labels in the nucleosomal DNA (at positions F-48 and R-28, Fig EV5A), assembled mononucleosomes with synthesized H3, either unmodified (Fig 5A top panel) or H3K122succ (Fig 5A bottom panel), and performed smFRET measurements under different NaCl concentrations. The merged smFRET efficiency histograms (Fig 5A) showed a peak at a FRET efficiency of around E = 0.8 representing the intact nucleosomes and a no-FRET peak representing free DNA or a very open nucleosome conformation. As expected, increased salt concentrations led to a reduction of the nucleosome peak. Interestingly, at intermediate salt concentrations the nucleosome population is reduced for H3K122succ-containing nucleosomes compared to unmodified H3-containing nucleosomes. Calculating the normalized fraction of FRET events in the nucleosome population (E ≥ 0.45; Fig 5B) revealed that H3K122succ nucleosomes are indeed more sensitive to destabilization by increased NaCl concentrations. Determining the NaCl concentration at which half of the nucleosomes have opened (x0) showed a Δx0 of ~0.12 M NaCl for H3K122succ compared to unmodified nucleosomes (see Materials and Methods for details). Together, these results reveal that H3K122succ nucleosomes are less stable than unmodified nucleosomes, possibly rendering the underlying DNA more accessible and, therefore, facilitating transcription. This is in line with reports of SIRT7-mediated chromatin condensation in cells upon DNA damage (Li et al, 2016).

We next wanted to investigate the effects of H3K122ac on nucleosome stability in a similar setup. We, therefore, performed smFRET assays with nucleosomes containing recombinantly expressed WT H3, H3K122ac, and H3K122E, where the latter was previously used to mimic H3K122succ (Li et al, 2016). As expected, we observed a reduced stability of the H3K122ac nucleosomes (Δx0 of ~0.05 M NaCl; Fig EV5B and C). The stability of H3K122E nucleosomes was similar to the corresponding WT H3-containing nucleosomes. Although glutamate inverts the charge of the side chain similarly to lysine succinylation, our results suggest that the shorter amino acid side chain of glutamate (compared to lysine) could prevent the H3K122E mutant from being an effective mimic of a succinylated lysine in nucleosome stability assays. Indeed, this is in line with previously reported H3 lysine acetylation mimics that did not fully recapitulate the effects of acetylation (Suganuma & Workman, 2008). When comparing our results between H3K122ac and H3K122succ nucleosomes, it is notable that although H3K122succ nucleosomes are destabilized more than H3K122ac nucleosomes (Fig EV5D and E), the increased instability, compared to H3K122ac, does not translate into increased transcriptional output. This suggests that there is a degree of nucleosomal destabilization that provides maximal transcription output and that further destabilization might have no additional effect on transcription, at least in vitro, possibly because transcription is an active process with multiple energy barriers, not simply depending on nucleosome opening.

Our results show that succinylation of H3K122 can be mediated by p300/ CBP, leading to nucleosome destabilization and possible displacement of the histone octamer from the DNA, resulting in increased DNA accessibility. This process is particularly important at regulatory regions such as TSS, suggesting that H3K122succ can facilitate the binding of transcriptional factors resulting in transcriptional activation and maintenance of active transcription. Conversely, the desuccinylation of H3K122succ by SIRT5 (or SIRT7 (Li et al, 2016)) leads to chromatin stabilization and compaction, decreasing, thereby DNA accessibility (Fig 5C). Thus, we gained for the first-time insights into the role of a site-specific histone
**Figure 4.** H3K122succ is sufficient to stimulate transcription.

A. Schematic representation of the IVT assay: chromatin assembly, density selection by sucrose gradient, and in vitro transcription per se in the presence of \( ^{32}P\)-rUTP. For details, see Materials and Methods.

B. In vitro transcription in the presence of GAL4-VP16 and p300, in the presence of no CoA, ac-CoA or suc-CoA. Shown is a representative autoradiogram of the IVT product.

C. Quantification of IVT reactions. Expression relative to no CoA is plotted. Average, ± SD, and individual datapoints of four experiments are shown. P-values relative to no CoA were calculated by two-tailed paired t-test.

D. In vitro transcription on unmodified chromatin or chromatin site-specifically succinylated at H3K122. Shown is a representative autoradiogram.

E. Quantification of IVT reactions. Expression relative to transcription on unmodified chromatin is plotted. Average, ± SD and individual datapoints of four experiments are shown. P-value was calculated by two-tailed paired t-test.
succinylation in transcription, expanding the role of histone acylations in transcriptional regulation.

We are currently only beginning to understand the links between cellular metabolism and chromatin via histone acylations (Trefely et al., 2020). Recently, production of succ-CoA in the nucleus through nuclear oxoglutarate dehydrogenase complex (OGDH) has been reported, allowing α-ketoglutarate conversion to succinyl-CoA, which can be used by the acetyltransferases to succinylate histones (Wang et al., 2017). However, it is still unclear whether nuclear, local production of CoAs results in metabolic microniches with increased concentrations of CoAs (Katada et al., 2012) that could determine the type of histone acylations to be set on chromatin. Upon changes in CoA availability (e.g. as a consequence of an altered metabolic cell state), specific acylations at the lateral surface of the histone octamer, such as H3K122succ, at distinct genomic regions, could directly modulate nucleosome stability and thus transcriptional outcome. Further studies will be required to better understand metabolic regulation of H3K122succ as well as the interplay between different H3K122 acylations.

**Materials and Methods**

**Antibody purification and characterization**

For the generation of anti-H3K122succ ABs, rabbits were immunized with the H3K122succ peptide (peptide sequences can be found in Table 1) according to the immunization protocol from BioGenes Gmbh (Germany). Specific ABs were affinity purified from rabbit serum in a two-step purification protocol using SulfoLink™ Coupling Resin (Thermo Fisher Scientific, USA): initially enriching for ABs specific for H3K122succ and later eliminating ABs recognizing H3K122un.
For AB characterization by peptide dot blot, serial dilutions of differently modified peptides (Table 1) were directly spotted onto 0.1 μm-pore nitrocellulose membranes. After completely air-dried, the membranes were blocked with 4% BSA-TBST (0.5% Tween® 20) for 1 h at room temperature and probed with Abs, at 1:1,000 dilution, O/N at 4°C. Regarding the characterization by immunoblot, histone acidic extracts were used (Di Cerbo et al., 2014). The primary AB was diluted 1:1,000 in 4% BSA-TBST (0.25% Tween® 20), and for competition experiments, the diluted AB was pre-incubated on ice with 200 pmol/ml of competitor peptide for 30 min prior to adding to the membrane.

### Cell culture

All cells were maintained at 37°C in a humidified atmosphere containing 5% CO₂. MCF7 and HEK293 were grown in low-glucose DMEM, supplemented with 10% fetal bovine serum (FBS), 2 mM l-glutamine, 100 U/ml penicillin, and 0.1 mg/ml streptomycin. MEFs were cultured in high-glucose DMEM, supplemented with 15% FBS, 1x pyruvate, 20 mM HEPES, 100 μM non-essential amino acids, 2 mM l-glutamine, 100 U/ml penicillin, and 0.1 mg/ml streptomycin.

For p300/ CBP inhibition, MCF7 cells were treated with 40 μM curcumin (previously shown to inhibit p300 significantly (Zhang et al., 2015)) O/N, followed by histone extraction and quantification of H3K122 succinylation levels by immunoblot. H3K122suc signal was quantified by Image Lab™ and normalized to the corresponding loading control (ponceau staining of membrane).

### Native chromatin immunoprecipitation

ChIP on native chromatin was performed as described previously with minor modifications (Daujat et al., 2009). For H3K122 succ, chromatin immunoprecipitation was performed independently with three different antibodies from a batch of native chromatin; the H3K9ac, H3K27ac, and H3K4me3 ChIP were performed in one replicate since comparable data have already been reported for MCF7 cells. MCF7 (5 × 10⁹–5 × 10¹⁰ cells) grown with 10% FBS was collected and washed with PBS. Cell pellets were resuspended in 4 ml of buffer 1 (0.3 M sucrose; 60 mM KCl 15 mM NaCl; 5 mM MgCl₂; 0.1 mM EDTA; 15 mM Tris–HCl, pH 7.5; 0.5 mM DTT; 0.1 mM AEBSP; 5 mM sodium butyrate; 5 mM nicotinamide; 1x Complete™ protease inhibitors) and divided into two tubes. Next, 2 ml of buffer II (buffer I supplemented with 0.4% IGEPAL CA-630® (Sigma-Aldrich, USA)) was added to each tube. Samples were gently mixed and incubated on ice for exactly 10 min. The samples were carefully layered on top of 8 ml of sucrose cushion (buffer 1 with 1.2 M sucrose) and centrifuged in a swing-out rotor at 10,000 g for 20 min at 4°C. The supernatant was carefully removed, and the pellets were resuspended in 1 ml of MNase digestion buffer (0.32 M sucrose; 50 mM Tris–HCl, pH 7.5; 4 mM MgCl₂; 1 mM CaCl₂; 0.1 mM AEBSP). Chromatin was fractionated by MNase digestion (at a final concentration of 2 U/ml) for 7–12 min at 37°C, which is known to release accessible chromatin first. The digestion was stopped with the addition of 5 mM EDTA.

Soluble chromatin fractions were collected by centrifugation of digested chromatin at 12,000 g for 10 min. The supernatant (S1) was collected, while the pellet was resuspended in the same volume of dialysis buffer (1 mM Tris–HCl, pH 7.5; 0.2 mM EDTA; 0.2 mM AEBSP; 5 mM sodium butyrate; 5 mM nicotinamide) and dialyzed O/N against dialysis buffer. On the next day, the samples were centrifuged as before and the supernatant was collected (S2). Both fractions underwent quality control on an agarose gel electrophoresis.

Per IP, 40 μg of digested chromatin (20 μg of S1 + 20 μg of S2) was diluted in 500 μl of ChIP incubation buffer (50 mM NaCl; 50 mM Tris–HCl, pH 7.5; 0.1 mM AEBSP; 5 mM sodium butyrate; 5 mM nicotinamide) and dialyzed O/N against dialysis buffer. On the next day, the samples were then incubated O/N at 4°C with different amounts of AB. To each sample, 20 μl of 50% protein A 50% protein G Dynabeads™ was added and incubated on a rotating wheel for 2–4 h. The beads were washed 3 times with ChIP incubation buffer, before being eluted with 2 × 125 μl of elution buffer (50 mM NaCl; 50 mM Tris–HCl, pH 7.5; 0.1 mM PMSF; 5 mM EDTA; 1% SDS (w/v)). The samples were then digested with RNase A (Fermentas, USA) for 30–45 min at 37°C and then purified using the QIAquick PCR Purification columns (Qiagen, Germany) according to the manufacturer’s instructions.

### ChIP-seq library preparation

ChIP samples were purified using Agencourt AMPure XP beads (Beckman Coulter) and quantified with the Qubit (Invitrogen). For the H3K122 succ samples and respective input, the ChIP-seq libraries were prepared from 5 to 10 ng of double-stranded purified DNA using the MicroPlex Library Preparation kit v2 (C05010014, Diagenode s.a., Seraing, Belgium), according to manufacturer’s instructions. While for H3K4me3, H3K9ac, H3K27ac, and respective input, the ChIP-seq libraries were prepared from 5 ng of double-stranded purified DNA using the NEBNext® Ultra II (E7645S, New England Biolabs). In the first step, the DNA was repaired and yielded molecules with blunt ends. In the next step, stem-loop adaptors with

### Table 1. List of peptides used in this study.

| Peptide abbreviation | Peptide name | Peptide sequence |
|----------------------|--------------|------------------|
| H3K9ac               | H3K9 acetylation | TKQTAR-Kac-STGGKAGC |
| H3K18ac              | H3K18 acetylation | GCGKAPR-Kac-QLATKAGC |
| H3K27ac              | H3K27 acetylation | ATKAAR-Kac-SAPATGC |
| H3K56ac              | H3K56 acetylation | IRRYQ-Kac-STEIIGGC |
| H3K64ac              | H3K64 acetylation | STELLIR-Kac-LPFQRLVGC |
| H3K115ac             | H3K115 acetylation | CAIHA-Kac-IVTIPMK |
| H3K122un             | H3K122 unmodified | CCGVTIMP-K-DIQLA |
| H3K122succ           | H3K122 succinylation | CCGVTIMP-Ksuc-DIQLA |
| H3K122ac             | H3K122 acetylation | CCGVTIMP-KAc-DIQLA |
| H3K122glut            | H3K122 glutarylation | CCGVTIMP-Kglut-DIQLAR |
| H3K122crot            | H3K122 crotonylation | CCGVTIMP-Kcrot-DIQLAR |
blocked 5’ ends were ligated to the 5’ end of the genomic DNA, leaving a nick at the 3’ end. The adaptors cannot ligate to each other and do not have single-strand tails, avoiding non-specific background. In the final step, the 3’ ends of the genomic DNA were extended to complete library synthesis and Illumina compatible indexes were added through a PCR amplification (8 cycles). Amplified libraries were purified and size-selected using Agencourt AMPure XP beads (Beckman Coulter) to remove unincorporated primers and other reagents. Libraries were sequenced on Illumina Hiseq 4000 sequencer as single-end 50 bp reads (H3K122succ samples and respective input) or 2 × 100 bp reads (H3K4me3, H3K9ac, H3K27ac, and respective input) following Illumina’s instructions. Image analysis and base calling were performed using RTA 2.7.3 and bcl2fastq 2.17.1.14.

**ChIP-seq analysis**

In this study, the generated ChIP-seq data and published data (see Table 2) were mapped to the human genome (assembly hg38) using Bowtie v1.0.0 (Langmead et al, 2009) with default parameters except for “-p 3 -m 1” (Cook et al, 2007). For H3K27ac data, MACS2 was run using the following parameters «-g hs --broad --broad-cutoff 0.1». Peaks falling into ENCODE blacklisted regions were removed. Peaks were annotated relative to genomic features using Homer v4.11.1 (Sven et al, 2010) (annotations were extracted from gtf file downloaded from Ensembl v94) and using the Bioconductor package ChIPseeker v1.20 (Yu et al, 2015). Random distribution of genomic features was obtained by randomly selecting genomic regions of the same amount and width as in the H3K122succ AB #2 peak set with Bedtools shuffle v2.26.0 Quinlan and Hall (2010). Picked regions were covered by at least one read in the input sample to avoid selecting regions in closed chromatin. Heatmaps at genomic loci were generated with seqMINER v1.3.3g (Ye et al, 2011). Other heatmaps and boxplots were generated using the R package ggplot2 v3.3 (Wickham, 2016). Venn diagrams were generated using the Bioconductor package DiffBind v2.12 (Ross-Innes et al, 2012) with R package VennDiagram v3.6.20.

For the comparison of our ChIP-seq dataset with the H3K122ac, H2A.Z, H2A.Zac, and p300 datasets, as well as for comparing the expression between H3K122succ and H3K122ac marked genes, we used only genes that did not show growth condition dependent changes in their expression (MCF7 grown in 10% FBS versus starved conditions with 5% charcoal-stripped FBS). For this, we used RNA-seq data and read counts per gene normalized across samples using the median-of-ratios method proposed by Anders and Huber (Anders & Huber, 2010) to make the read counts comparable.

**Table 2. List of publicly available datasets used in this study.**

| Dataset name       | GEO data  | Year | Authors                                                                 | Link                                                                 |
|--------------------|-----------|------|-------------------------------------------------------------------------|----------------------------------------------------------------------|
| MCF7_RNA-seq       | GSM3110713| 2019 | Handa T, Katayama A, Yokobori T, Yamané A, Fujii T, Obayashi S, Kurozumi S, Kawabata-Iwakawa R, Gombodorj N, Nishiyama M, Asao T, shirabe K, Kuwano H, and Oyama T | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM3110713          |
| MCF7_NS_FAIRE_rep1 | GSM1825697| 2016 | Hardy K, Wu F, Tu W, Zafar A, Boulding T, McCuaig R, Sutton C R, Theodoratos A, Rao S | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1825697          |
| MCF7_Dnase-seq     | GSM2476264| 2017 | Liu Y, Chen S, Wang S, Soares F, Fisvher M, Meng F, Du Z, Lin C, Meyer C, DeCaprio J A, Brown M, Liu X S, He H H | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM2476264          |
| p300_ChIP          | GSM1059392| 2013 | Tropberger P, Sott S, Keller C, Kamieniarz-Gdula K, Caron M, Richter F, Li G, Mittler G, Liu E T, Bühler M, Margueron R, Schneider R | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1059392          |
| H3K122Ac_ChIP      | GSM1059385| 2013 | Tropberger P, Sott S, Keller C, Kamieniarz-Gdula K, Caron M, Richter F, Li G, Mittler G, Liu E T, Bühler M, Margueron R, Schneider R | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1059385          |
| MCF7_H2A.Z_ChIP    | GSM1059387| 2013 | Tropberger P, Sott S, Keller C, Kamieniarz-Gdula K, Caron M, Richter F, Li G, Mittler G, Liu E T, Bühler M, Margueron R, Schneider R | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1059387          |
| MCF7_H2A.Zac_ChIP  | GSM1059388| 2013 | Tropberger P, Sott S, Keller C, Kamieniarz-Gdula K, Caron M, Richter F, Li G, Mittler G, Liu E T, Bühler M, Margueron R, Schneider R | https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1059388          |
between samples. The comparison between the conditions was performed with the Bioconductor package DESeq2 v1.6.3 with R v3.1.1. Genes having a log2FC ≥ 1 and a P-value < 0.05 were considered as differentially expressed and thus excluded from further analysis.

Expressed genes (at least one read in RNA-seq) with H3K122succ and H3K122ac peaks (± 500 bp from their TSS) were compared based on their Ensembl Gene IDs, and results are shown as a Venn diagram created by using R scripts.

CGI positions were extracted as a BED file using table cgIsland-Ext of the UCSC table browser. CGIs were annotated relative to Ensembl v94 genomic features using Homer v4.11.1 (Sven et al., 2010). Genes having H3K122succ peaks and CGIs at their TSS regions were compared based on their Ensembl Gene IDs, and results are shown as a Venn diagram created using R scripts.

Gene symbols of genes associated to H3K122succ peaks were compared to the human housekeeping gene list from: https://www.tau.ac.il/~elieis/HKG/HK_genes.txt (Eisenberg & Levanon, 2013).

**RNA-seq analysis**

Reads were mapped onto the hg38 assembly of *Homo sapiens* genome using STAR version 2.5.3a (Dobin et al., 2013). Gene expression quantification was performed from uniquely aligned reads using htsqucoount version 0.6.1p1 (Anders et al., 2015), with annotations from Ensembl v94 and “union” mode. Read counts have been normalized across samples with the median-of-ratios method proposed by Anders and Huber (Anders & Huber, 2010), to make these counts comparable between samples.

**siRNA KD**

Depletion of HAT enzymes was achieved by transient transfection using Lipofectamine™ RNAiMAX (Thermo Fisher Scientific, USA) combined with a reverse transcription protocol following the supplier’s instructions. The following ON-TARGETplus SMARTpools from Dharmacon (Horizon Discovery, UK) were used: p300, L-003486-00-0010; CBP, L-003477-00-0010; pCAF, L-005055-00; GCN5, L-009722-00; scramble negative control, D-001810-10. In brief, a mixture of 10–50 nM of siRNA and lipofectamine was preincubated for 20 min at room temperature and mixed with MCF7 cells before plating them. Cells were grown for 72 h.

**p300 purification**

His-tagged p300 was expressed on baculovirus-infected S9 insect cells. After expression, the cells were harvested by centrifugation for 5 min at 400 g at 4°C. Unless stated otherwise, all steps were carried out on ice. The cell pellet was resuspended in 2.5 ml of buffer B (20 mM Tris–HCl, pH 7.5; 250 mM NaCl; 0.1% IGEPAL CA-630; 30 mM imidazole; 1× cComplete™ EDTA-free protease inhibitor cocktail) and homogenized with 10 strokes using an “A” pestle. The sample was incubated on ice for 15 min before being aliquoted into 700–800 µl aliquots and sonicated on a Branson sonicator (3 series of 5 min, with 90% amplitude, 30 s ON and 30 s OFF). The aliquots were pooled together and centrifuged for 15 min at 16,000 g at 4°C. The clarified supernatant was loaded on a pre-equilibrated 1 ml HisTrap™ HP column (GE Healthcare, USA) connected to an Äkta™ pure. After loading, the column was washed with buffer B and eluted with 10 column volumes (CV) of elution buffer (20 mM Tris–HCl, pH 7.5; 100 mM NaCl; 0.1% IGEPAL CA-630; 250 mM imidazole; 10% glycerol; 1× cComplete™ EDTA-free protease inhibitor cocktail). 500 µl fractions were collected during the elution. The different fractions were analyzed on a 4–12% SDS–PAGE gel followed by coomassie staining. The fractions containing the purified p300 were pooled together and diluted with 50% glycerol before being aliquoted and snap-frozen in liquid nitrogen. The aliquots were stored at −80°C.

**Succinyltransferase assays**

For the in vitro succinyltransferase assays on octamers, 1 µg of recombinant octamers was incubated with 40 ng of p300 in the presence of suc-CoA O/N at 30°C. The reaction products were separated by SDS–PAGE and analyzed by immunoblotting with anti-H3K122succ peptides, the samples were incubated with 0–160 ng of p300 in the presence of approximately 17 µCi/ml of radiolabeled suc-CoA (American Radiolabeled Chemicals, USA) O/N at 30°C. The reactions were spotted on cellulose chromatography paper P81. Chromatography papers were washed three times with 10% TFA before the incorporated radioactivity was measured by liquid scintillation counting. For “competition assays” on WT H3, 0.5 µg of H3 was incubated with 40 ng of p300 in the presence of approximately 17 µCi/ml of [3H]ac-CoA (Hartmann Analytic, Germany). Unlabeled ac-CoA or suc-CoA (0-70.4 mmol) was added as “competitor” to the reaction. After 4 h of incubation, the reactions were stopped, and the samples ran on an 18.7% SDS–PAGE gel and transferred to a 0.45 µm nitrocellulose membrane. The membrane was air-dried before being exposed to a detection screen. The results were read on an Amersham™ Typhoon™ Biomolecular Imager and signal intensities quantified. In all of these assays, the final reaction buffer consisted of 5% glycerol; 50 mM Tris, pH 8; 0.1 mM EDTA; 7.5 mM sodium butyrate; 7.5 mM nicotinamide; 1 mM DTT; and 1× cComplete™ protease inhibitors (Roche, Switzerland).

**Desuccinylation assays**

For the in vitro desuccinylation assays, 1 µg of H3K122succ peptides was incubated with 0.5–1.8 µg of SIRT5 or SIRT7 in reaction buffer (20 mM Tris–HCl, pH 7.5; 1 mM DTT) supplemented, or not, with NAD⁺. After 3 h at 37°C, 2 µl of each reaction was spotted on 0.1 µM-pore nitrocellulose membrane and probed with AB anti-H2K122succ #2.

**NAP1 purification**

NAP1 was purified from cell pellets of baculovirus-infected S9 insect cells expressing N-terminally His-tagged NAP1 as previously described (Peterson, 2009). Briefly, after harvesting the cells by centrifugation, the cell pellet was washed with ice-cold PBS. All subsequent steps were performed on ice unless stated otherwise. The cell pellet was resuspended in lysis buffer (50 mM sodium phosphate, pH 7.6; 500 mM NaCl; 20 mM imidazole; 15% glycerol; 0.01% NP40; 10 mM β-glycerophosphate; 0.2 mM PMSF; 0.5 mM EDTA-free protease inhibitor cocktail). The sample was incubated in ice for 15 min before being aliquoted into 700–800 µl aliquots and sonicated on a Branson sonicator (3 series of 5 min, with 90% amplitude, 30 s ON and 30 s OFF). The aliquots were pooled together and centrifuged for 15 min at 16,000 g at 4°C. The clarified supernatant was loaded on a pre-equilibrated 1 ml HisTrap™ HP column (GE Healthcare, USA) connected to an Äkta™ pure. After loading, the column was washed with buffer B and eluted with 10 column volumes (CV) of elution buffer (20 mM Tris–HCl, pH 7.5; 100 mM NaCl; 0.1% IGEPAL CA-630; 250 mM imidazole; 10% glycerol; 1× cComplete™ EDTA-free protease inhibitor cocktail). 500 µl fractions were collected during the elution. The different fractions were analyzed on a 4–12% SDS–PAGE gel followed by coomassie staining. The fractions containing the purified p300 were pooled together and diluted with 50% glycerol before being aliquoted and snap-frozen in liquid nitrogen. The aliquots were stored at −80°C.

**Succinyltransferase assays**

For the in vitro succinyltransferase assays on octamers, 1 µg of recombinant octamers was incubated with 40 ng of p300 in the presence of suc-CoA O/N at 30°C. The reaction products were separated by SDS–PAGE and analyzed by immunoblotting with anti-H3K122succ peptides, the samples were incubated with 0–160 ng of p300 in the presence of approximately 17 µCi/ml of radiolabeled suc-CoA (American Radiolabeled Chemicals, USA) O/N at 30°C. The reactions were spotted on cellulose chromatography paper P81. Chromatography papers were washed three times with 10% TFA before the incorporated radioactivity was measured by liquid scintillation counting. For “competition assays” on WT H3, 0.5 µg of H3 was incubated with 40 ng of p300 in the presence of approximately 17 µCi/ml of [3H]ac-CoA (Hartmann Analytic, Germany). Unlabeled ac-CoA or suc-CoA (0-70.4 mmol) was added as “competitor” to the reaction. After 4 h of incubation, the reactions were stopped, and the samples ran on an 18.7% SDS–PAGE gel and transferred to a 0.45 µm nitrocellulose membrane. The membrane was air-dried before being exposed to a detection screen. The results were read on an Amersham™ Typhoon™ Biomolecular Imager and signal intensities quantified. In all of these assays, the final reaction buffer consisted of 5% glycerol; 50 mM Tris, pH 8; 0.1 mM EDTA; 7.5 mM sodium butyrate; 7.5 mM nicotinamide; 1 mM DTT; and 1× cComplete™ protease inhibitors (Roche, Switzerland).
benzamidine-HCl) in 1/40 of the initial cell culture volume and homogenized on a Branson sonicator (3 series of 1 min, with 50% amplitude, 0.5 s ON, and 0.5 s OFF). The sample was cleared by centrifugation for 10 min at 15,000 g at 4°C, and the supernatant was loaded on a pre-equilibrated 5 ml HisTrap™ HP column (GE Healthcare, USA) connected to an Äkta™ pure. After loading the sample, the column was washed with lysis buffer and with wash buffer (50 mM sodium phosphate, pH 7.6; 100 mM NaCl; 20 mM imidazole; 15% glycerol; 0.01% NP40; 10 mM β-glycerophosphate; 0.2 mM PMSF; 0.5 mM benzamidine-HCl). NAP1 was eluted with 42 ml of elution buffer, and fractions of 500 µl were collected and analyzed on an SDS–PAGE gel. The fractions containing NAP1 were pooled together and dialyzed against two changes of 4 l of dialysis buffer (25 mM HEPES, pH 7.6; 1 mM EDTA; 10% glycerol; 100 mM NaCl; 0.01% NP40; 10 mM β-glycerophosphate; 1 mM DTT; 0.2 mM PMSF), where the first dialysis was carried out O/N and the second for 2 h and against one change of 4 l of NAP1 buffer (10 mM HEPES, pH 7.6; 1 mM KCl; 1.5 mM MgCl2; 0.5 mM EDTA; 10% glycerol; 0.01% NP40; 10 mM β-glycerophosphate; 1 mM DTT; 0.2 mM PMSF) plus 100 mM NaCl for 2 h. After dialysis, any precipitate was removed by centrifugation for 20 min at 20,000 g. The soluble protein was quantified on an 8% SDS–PAGE gel followed by coomassie staining, using BSA as a mass standard. The sample was loaded on a pre-equilibrated 6 ml Resource™ Q (GE Healthcare, USA) column connected to an Äkta™ pure. The column was washed with NAP1 buffer plus 200 mM NaCl and eluted with a 20 CV of a linear gradient from NAP1 buffer supplemented with 200 mM NaCl to NAP1 buffer plus 500 mM NaCl. Fractions of 0.5 ml were collected during elution and analyzed on an 18% SDS–PAGE gel. The fractions containing NAP1 were pooled together and dialyzed against two changes of 4 l of dialysis buffer (25 mM Tris–HCl, pH 7.9; 500 mM NaCl; 4 mM MgCl2; 0.4 mM EDTA; 2 mM DTT; 20 mM β-glycerophosphate; 20% glycerol; 0.4 mM PMSF; 1 mM benzamidine-HCl; 4 µg/ml leupeptin; 2 µg/ml aprotinin) and dounce homogenized using an “A” pestle (three series of ten strokes over 30 min, on ice). Homogenate was centrifuged for 10 min at 14,500 g at 4°C. The supernatant was transferred to a new tube and mixed with 250 µl of FLAG-M2 resin and 7 ml of dilution buffer (20 mM Tris–HCl, pH 7.9; 150 mM NaCl; 2 mM MgCl2; 0.2 mM EDTA; 1 mM DTT; 10 mM β-glycerophosphate; 15% glycerol; 0.01% NP40; 0.2 mM PMSF; 0.5 mM benzamidine-HCl; 2 µg/ml leupeptin; 1 µg/ml aprotinin). The slurry mix was incubated for 2–4 h at 4°C on a rotation wheel. After incubation, the samples were centrifuged for 3 min at 775 g at 4°C. The supernatant was removed and the pellet resuspended in 12 ml of wash buffer F (20 mM Tris–HCl, pH 7.9; 150 mM NaCl; 2 mM MgCl2; 0.2 mM EDTA; 1 mM DTT; 10 mM β-glycerophosphate; 15% glycerol; 0.01% NP40; 0.2 mM PMSF; 0.5 mM benzamidine-HCl; 2 µg/ml leupeptin; 1 µg/ml aprotinin). The slurry mix was incubated for 20 min at 20,000 g and the supernatant was centrifuged for 20 min at 80°C. The samples were centrifuged for 30 min at 30°C. The reaction was chased with the addition of a 6 µl rUTP-mix (600 µM cold rUTP and 6 U/µl RNase T1) and incubated for 20 more minutes. The transcribed RNA was phenol-chloroform extracted, precipitated with ethanol, and analyzed on 6% acrylamide denaturing gels (Margueron et al., 2008). A phosphor screen was used to capture the signal O/N and was then measured on an Amersham™ Typhoon 5 Biomolecular Imager (GE Healthcare, USA). The signal quantifications were done by ImageJ (Java).

**Preparation of double labeled nucleosomes and smFRET assay**

Into the 147 bp DNA sequence, the dye molecules were incorporated via PCR with labeled primers (IBA, Table 3). The double labeled DNA was then purified by ethanol precipitation and size exclusion chromatography with a
Superose™ 6 Increase 3.2/300 column (GE Healthcare USA). Assembly of double labeled nucleosomes was performed as described earlier (Schwarz et al., 2018). In brief, a DNA mixture of the 147 bp labeled DNA (1/10) and unlabeled 200 bp 601 DNA (9/10) in a total concentration of 400 nM was used for the assembly. The unlabeled DNA was used in order to increase the overall nucleosome concentration during single molecule experiments. Additionally, an equal amount of low affinity competitor DNA was used for buffering nucleosome assembly. Purified octamer solution, with specific modifications or unmodified, was titrated against the DNA solution in order to maximize assembly. The nucleosomes were assembled via an O/N salt-gradient dialysis in a TE buffer with DTT (0.25 mM) from the DNA mixture and the histone octamers. For this the Slide-A-Lyzer™ MINI Dialysis Device (7 kDa cutoff, Thermo Fisher Scientific) with the sample was placed in 300 ml of buffer with 2 M NaCl and the salt concentration was reduced stepwise by adding buffer with 50 mM NaCl until about 3 l. Then, a final dialysis step was performed in buffer with 50 mM NaCl. The assembled nucleosomes were analyzed on a 0.4x TBE, 6% PAA gel.

For performing the smFRET measurements, the nucleosomes were diluted to picomolar concentrations of the labeled nucleosomes in measurement buffer containing 10 mM Tris–HCl pH 7.8, 1 mM EDTA, and 0.1 mg/ml BSA with varying NaCl concentrations. The sample was incubated in the buffer for 1.5–2 h on ice, and then, 30 µl was pipetted on a PEG-coated coverslip with a 0.25 to 1.1 NaCl of the respective construct in order to account for a small fraction of free DNA present after assembly.

At least three measurements per sample and salt concentration were performed. After taking the mean, the Boltzmann sigmoid function was fitted to the data \( x = \frac{1}{1 + \exp(-dx)} \) where \( x_0 \) is the NaCl concentration where 50% of the nucleosomes have disassembled and \( dx \) is a parameter describing the slope of the decay.

### Data availability

The ChIP-seq data generated for this study were deposited in the Gene Expression Omnibus (GEO) with accession number GSE152019 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE152019).

### Acknowledgements

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**Table 3. MMTV-A sequence and primer sequences used in the stability assays.**

| 147 bp DNA sequence based on the MMTV-A sequence and primers | Comments |
|-------------------------------------------------------------|----------|
| **5'- ACTTGC AACAG GCC TCG TCG GCC CGC GAC CTC TG -3'**      | T Cy3B attached via C6 linker |
| **5'- CAA AAA ACT GTG CCG CAG TCG GCC GAC CTG**              | T Alexa647 attached via C2 linker |
| **TTTTT TG -3'**                                             | T Position 0 (dyad) |
| **T Donor Cy3B attached to T on forward strand (F-48)**      | A Acceptor Alexa647 attached to T on reverse strand (R-28) |

The data were analyzed with the software PAM (PIE analysis with MATLAB v1.2) (Schrömpfl et al., 2018). A burst search (Nir et al., 2006) was performed with a minimum threshold of 50 photons per burst, and additionally, at least 5 photons had to be in a time window of 500 µs. To filter for the FRET population, a stoichiometry (raw) cut (0.2–0.85) and an ALEX-2CDE filter (max. 11) (Tomov et al., 2012) were applied to those data. After correcting the data for background (between about 0.2 and 0.5 kHz), crosstalk (0.10), direct excitation (0.06), the detection correction factor (0.39), and the excitation correction factor (2.07), a FRET efficiency cut (~0.25 to 1.1) was applied. Due to the design of the dye molecule attachment sites, intact nucleosomes show a peak at a high FRET efficiency and unwrapped nucleosomes show a peak at a FRET efficiency of (close to) zero. The fraction of intact nucleosomes was calculated by dividing the number of events in the nucleosome population (FRET population with a FRET efficiency equal or higher than 0.45) by the overall events in the FRET population. Those values were normalized to the mean value at 0 M NaCl of the respective construct in order to account for a small fraction of free DNA present after assembly.

The measurements were performed as described before (Bönsch et al., 2012). The measurement was performed for 1 h at room temperature on a custom-built confocal microscope which was previously described in detail (Schwarz et al., 2018). It uses pulsed interleaved excitation (PIE) in combination with multiparameter fluorescence detection (MFD) (Kudryavtsev et al., 2012). The dyes were excited with a green laser (532 nm) and a red laser (640 nm) with an excitation power of 90 mW for both lasers measured before the objective. For each laser, a repetition rate of 20 MHz was used.

The data were analyzed with the software PAM (PIE analysis with MATLAB v1.2) (Schrömpfl et al., 2018). A burst search (Nir et al., 2006) was performed with a minimum threshold of 50 photons per burst, and additionally, at least 5 photons had to be in a time window of 500 µs. To filter for the FRET population, a stoichiometry (raw) cut (0.2–0.85) and an ALEX-2CDE filter (max. 11) (Tomov et al., 2012) were applied to those data. After correcting the data for background (between about 0.2 and 0.5 kHz), crosstalk (0.10), direct excitation (0.06), the detection correction factor (0.39), and the excitation correction factor (2.07), a FRET efficiency cut (~0.25 to 1.1) was applied. Due to the design of the dye molecule attachment sites, intact nucleosomes show a peak at a high FRET efficiency and unwrapped nucleosomes show a peak at a FRET efficiency of (close to) zero. The fraction of intact nucleosomes was calculated by dividing the number of events in the nucleosome population (FRET population with a FRET efficiency equal or higher than 0.45) by the overall events in the FRET population. Those values were normalized to the mean value at 0 M NaCl of the respective construct in order to account for a small fraction of free DNA present after assembly.

At least three measurements per sample and salt concentration were performed. After taking the mean, the Boltzmann sigmoid function was fitted to the data \( x = \frac{1}{1 + \exp(-dx)} \) where \( x_0 \) is the NaCl concentration where 50% of the nucleosomes have disassembled and \( dx \) is a parameter describing the slope of the decay.

### Data availability

The ChIP-seq data generated for this study were deposited in the Gene Expression Omnibus (GEO) with accession number GSE152019 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE152019).

### Acknowledgements

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Author contributions
LZS, MH, and SN performed and designed experiments. SLG analyzed the ChiP-seq data. LZS, JM, SD, and RS conceived and designed the project. AM and AG expressed p300, ACf, and NAP1. LZS and RM setup and performed the IVT experiments. LZS, JM, SLG, SD, and RS wrote the manuscript with input from the other authors.

Conflict of interest
The authors declare that they have no conflict of interest.

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