THEMATIC REVIEW

90 YEARS OF PROGESTERONE
Molecular mechanisms of progesterone receptor action on the breast cancer genome

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Abstract

Gene regulation by steroid hormones has been at the forefront in elucidating the intricacies of transcriptional regulation in eukaryotes ever since the discovery by Karlson and Clever that the insect steroid hormone ecdysone induces chromatin puffs in giant chromosomes. After the successful cloning of the hormone receptors toward the end of the past century, detailed mechanistic insight emerged in some model systems, in particular the MMTV provirus. With the arrival of next generation DNA sequencing and the omics techniques, we have gained even further insight into the global cellular response to steroid hormones that in the past decades also extended to the function of the 3D genome topology. More recently, advances in high resolution microscopy, single cell genomics and the new vision of liquid-liquid phase transitions in the context of nuclear space bring us closer than ever to unravelling the logic of gene regulation and its complex integration of global cellular signaling networks. Using the function of progesterone and its cellular receptor in breast cancer cells, we will briefly summarize the history and describe the present extent of our knowledge on how regulatory proteins deal with the chromatin structure to gain access to DNA sequences and interpret the genomic instructions that enable cells to respond selectively to external signals by reshaping their gene regulatory networks.

Introduction

The steroid hormone progesterone was initially considered to be involved mainly in menstrual cycle, pregnancy, and mammary gland function. Meanwhile, it was also implicated in multiple other functions outside the sexual organs. In this review we will concentrate on the role of progestins (Pg) on the regulation of gene expression via their specific progesterone receptor (PR), focusing on breast cancer cells. All the steroid hormone receptors share a similar organization of domains – a central short DNA-binding domain composed of two zinc fingers coordinated each by four cysteines; a C-terminal domain that is responsible for hormone binding and interaction with co-regulators; and a N-terminal domain of variable length, which is mainly unstructured and includes multiple residues that can be post-translationally modified and fine tune the receptors’ functional activity.
(see (Gronemeyer 1992; Huang et al. 2010), for review on steroid hormone receptor structure). In the case of PR, alternative initiation of the same gene leads to the generation of two PR isoforms, the complete isoform named PRB and the PRA isoform, which lacks the first 164 amino acids (Kastner et al. 1990). The difference in the properties of these two isoforms has been a matter of debate as the results observed depended on the nature of the assays used, which mainly were in vitro assays or transient transfection experiments (Jacobsen and Horwitz 2012; Khan et al. 2012). However, the importance of the N-terminal region of PR for hormone action and some recent experiments point to a significant functional difference between the two isoforms that will be addressed in the last section of this review.

This review will mainly focus on the mechanisms that enable the PR to access DNA sequences on chromatin and to modulate the transcriptional rate of subsets of hormone responsive genes. Before going into the more recent results obtained with genome-wide approaches, we will summarize as a historical introduction the concepts elaborated at the turn of the last century based on the findings related to PR regulation of the MMTV expression.

PR before omics: the MMTV promoter as a model system.

The role of progestins in breast cancer cells has been mostly neglected in favor of the more obvious effect of estrogens as drivers of cell proliferation (Carroll et al. 2017). This is particularly the case when we consider the most popular cell-culture model of luminal breast cancer, namely the MCF-7 cell line, which exhibits higher levels of estrogen receptor alpha (ERα) compared to PR. However, more recently, the interplay between estrogens and progestins has received considerable attention, predominantly based not only on results from cell-culture models but also on whole animal studies (Mohammed et al. 2015; Need et al. 2015; Singhal et al. 2016). Most studies on progestin action in cell culture have used the T47D cell line that, in contrast to MCF7 cells, expresses much higher constitutive levels of PR compared to ERα.

Prior to the establishment of whole genome approaches, most of the studies on PR action were based on the MMTV provirus model, which was initially used to study regulation by the glucocorticoid receptor (GR) (Ringold et al. 1975). Later, MMTV expression was shown to be regulated by Pg via the PR (Ahe et al. 1985). Both GR and PR were shown to bind cooperatively to DNA sequences of the MMTV LTR, upstream of the TATA box of the main MMTV promoter (Payvar et al. 1983; Scheidereit et al. 1983) (Fig. 1A). Similar sequences were found in many other Pg-regulated genes, although at different locations relative to the gene promoter. Comparison of these sequences led to the identification of the 15-mer palindromic hormone response element (HRE) with the consensus GGTACAnnnTGTTCT, which is recognized by a homodimer of PR in a head-to-head orientation (Beato 1989). The same 15-mer was also bound by GR, androgen receptor (AR) and the mineralocorticoid receptor. In the region bound by PR on the long terminal repeat (LTR) of the MMTV provirus, one canonical palindromic HRE and four incomplete HREs containing only one half of the palindromic sequence were identified. Downstream of the cluster of receptor-binding sequences is a palindromic binding site for the transcription factor nuclear factor 1 (NF1), which is important for efficient hormonal regulation. Intriguingly, PR and NF1 synergize in cell assays (Truss et al. 1995) but compete for binding to free DNA (Bruggemeier et al. 1990), suggesting that free DNA is not a sufficient template to completely explain the mechanism of hormonal gene regulation. This observation on free DNA led us to investigate the manner in which the MMTV promoter is organized in the context of chromatin.

When assembled in chromatin in vitro and in cells, the MMTV promoter positions precisely one nucleosome that includes the five HREs and the NF1 binding site (Richard-Foy and Hager 1987; Piña et al. 1990). The orientation of the DNA major groove in the surface of the histone octamer enables binding of PR to the palindromic HRE-I and to the half palindromic HRE-IV. This is possible because PR only contacts a narrow sector of the DNA double helix (Fig. 1B and C). In contrast, NF1 cannot bind its target sequence when assembled in nucleosomes as it contacts the whole circumference of the DNA helix (Fig. 1B and D) (Scheidereit and Beato 1984). However, in cells carrying an integrated copy of the MMTV promoter (Truss et al. 1995) or when assembled on the surface of MMTV minichromosomes (Di Croce et al. 1999), PR and NF1 synergize in an ATP-dependent process upon exposure to hormone. Thus, efficient MMTV induction depends on the proper nucleosome organization of its promoter, as demonstrated by the lack of synergism in histone-depleted yeast (Chávez and Beato 1997). This seems paradoxical, as the DNA binder with the weaker DNA affinity (PR) recognizes its target sequence on nucleosomes and upon ATP-dependent remodeling enables the occupancy of the stronger DNA binding factor (NF1) (Bruggemeier et al. 1990). However, something similar also occurs with the
classical pioneer factors, such as FOXA1, which exhibit an even weaker affinity for DNA, compared to PR, but are capable of enabling binding of other transcription factors even in the absence of ATP (Zaret and Carroll 2011; Zaret et al. 2016).

With MMTV minichromosomes assembled in Drosophila embryo extracts, the synergism between PR and NF1 is improved in the presence of histone H1, which represses transcription by each of the factors added separately (Koop et al. 2003). The reason being that the presence of histone H1 increases the proportion of chromatin templates that adopt the correct nucleosome organization. During the transcription process, H1 becomes phosphorylated in a PR-dependent manner and is partially displaced from the chromatin template by the NURF complex (Koop et al. 2003). Thus, histone H1 fulfills a complex and dynamic role in the regulation of the MMTV promoter, keeping the promoter silent in the absence of hormone and enabling efficient induction in response to hormone. Later we found that MSK1 phosphorylation at S10 of histone H3 and the displacement of H2A/H2B dimers from the MMTV promoter nucleosome by the SWI/SNF complex are part of the chromatin remodeling required for MMTV activation (Vicent et al. 2004; Vicent et al. 2006). Thus, PR-induced transcription of the MMTV promoter is initiated on the surface of H3/H4 tetramer that is more sensitive to DNAse I at its dyad (Vicent et al. 2010).

The PR cistrome in breast cancer cells

The arrival of whole genome approaches using next generation DNA sequencing made it possible for the first time to explore the global landscape of PR occupancy over the entire genome and to correlate this with the changes in the transcriptome upon hormone exposure. In T47D cells only a few hundred weak PR-binding sites were detected by chromatin immunoprecipitation and sequencing (ChIP-seq) with a PR-specific antibody prior to hormone exposure. In contrast, upon 30–60 min of exposure to 10 nM promegestone (R5020, abbreviated as Pg), we identified around 25,000 PR-binding sites and

Figure 1
The hormone responsive region of the MMTV promoter. (A) Of the five hormone response elements (HREs) in the MMTV promoter, only HRE-I is a perfect palindromic sequence. The other four HREs are half palindromic, although in in vitro DNA binding experiments homodimers of the PR bind cooperatively to each HRE. The nucleotides upstream of the transcription start sites and the position of the TATA box are indicated (Scheidereit et al. 1983; Chávez & Beato 1997). PR and NF1 compete for binding to the MMTV promoter DNA in vitro. The presence of an excess of PR or preincubation with PR precludes NF1 binding (top). In contrast, the presence of an excess of NF1 or preincubation with NF1 diminishes PR binding (bottom). This is due to the proximity of the HRE-IV and the NF1 binding sequence and to the cooperative binding of PR to all HREs (Bruggemeier et al. 1990). PR can bind its target sequence in nucleosomes while NF1 cannot. Panel (B) shows the contacts of PR with the HRE IV and of NF1 with its target sequence in a side view and panels (C) and (D) in a view along the DNA helix axis taking PR and NF1 as view points, respectively. Contacts with guanines (G), phosphates (P) and thymines (T) are highlighted. The non-accessible contacts are those facing the core histone (Scheidereit & Beato 1984).
observed significant changes in expression of around 2000 genes using RNA-seq (Ballare et al. 2013). Sequence analysis revealed that most of these PR binding sites do not exhibit a complete palindromic HRE, but just one or several half HREs. As measured by MNase-seq, the large majority of PR binding sites were located over regions of chromatin enriched in nucleosomes prior to hormone exposure (Ballare et al. 2013) (Fig. 2, left panel, − hormone). However, after 30–60 min of hormone exposure, we found a significant decrease in the number of MNase reads over the PR binding sites, suggesting that the nucleosomes have been remodeled and become sensitive to MNase digestion (Ballare et al. 2013) (Fig. 2, left panel, + hormone). It is worth noting that other ligands binding to PR induced different binding dynamics and gene regulation. While Pg-occupied PR binds rapidly and exchanges rapidly as well, PR occupied by the partial agonist RU486 binds and exchanges more slowly, while PR occupied by the full antagonist ZK98299 does not bind to the MMTV promoter (Rayasam et al. 2005).

The extent of nucleosome remodeling was estimated by the ratio of MNase reads before and after hormone exposure, which we named Nucleosome Remodeling Index (NRI). Classifying the PR-binding sites according to their NRI (Fig. 2, left panel) showed that sites with the highest NRI corresponded to the strongest PR peaks that were associated with hormone responsive genes and were heavily remodeled upon hormone exposure (Fig. 2, right top panel). On the contrary, PR binding sites with the lowest NRI showed weaker peaks not associated with hormone responsive genes that were not remodeled upon hormone exposure (Fig. 2, right lower panel) (Ballare et al. 2013). Therefore, the genome-wide analysis of PR binding confirmed the original observation made with the MMTV model system years earlier, supporting that optimal PR binding, as observed in around 2500 high NRI sites, requires the PREs to be organized in an accessible way on nucleosomes that are remodeled upon hormone exposure. These findings are in conflict with a very general assumption that for transcription factor access to their DNA target sequences requires the previous displacement of nucleosomes. This idea does hold true for factors like NF1 that interact with DNA embracing the double helix, but does not apply to PR that only contacts a narrow sector of the DNA helix, as the distance between the two halves of the palindrome is 10 bp, both halves are similarly exposed on the surface of a perfect HRE (Piña et al. 1990) (Fig. 1B).

In the case of estrogens and glucocorticoids, it was reported that binding of their ligand-activated receptors to DNA occurs mainly at chromatin regions that are already accessible prior to hormone exposure, judged by the hypersensitivity of these regions to nucleases (John et al. 2011; Zaret and Carroll 2011). This is mainly attributed to the action of the pioneer factor FOXA1, which is postulated to prepare the chromatin for subsequent hormone receptor binding (Zaret and Carroll 2011). GATA3 could also act upstream of FOXA1 to enable ERα binding to its target sequences (Theodorou et al. 2013). Pioneer factors may remain attached to chromatin even during mitosis, enabling transcriptional memory (Palozola et al. 2019). In line with this, we have also uncovered that a fraction of the PR binding sites are already highly occupied by FOXA1 prior to hormone exposure and that the amount of FOXA1 does not increase after adding hormone, while a larger proportion of sites is marked by FOXA1 and ligand-activated PR favors further recruitment of FOXA1 (Nacht et al. 2019). However, we interpret the requirement of FOXA1 for PR binding as a consequence of FOXA1-mediated displacement of histone H1 (Iwafuchi-Doi et al. 2016) rather than to the removal of nucleosome core particles. In addition, we cannot exclude that these sites may reflect loops formed by distant interactions (Glont et al. 2019). Interestingly, we found a subset of PR binding sites that are assisted by C/EBPα binding prior to hormone exposure (Nacht et al. 2019). These regions exhibit epigenetic marks of active enhancers and C/EBPα favors contact of these enhancers with their target promoters.

**Figure 2**
Classification of the genome PR-binding sites according to their Nucleosome Remodeling Index (NRI). Top panels: Around 25,000 PR-binding sites detected upon 60 min of hormone exposure were classified according to their Nucleosome Remodeling Index (NRI) and are shown as a heat map of nucleosome reads before and after hormone exposure ordered from the highest NRI to the lowest NRI. Bottom panels: Right: Nucleosome occupancy plot around the 2500 PR-binding sites with the highest NRI, before hormone exposure (red) and after 60 min of hormone exposure (lilac). Left: A similar nucleosome occupancy plot with the 2500 PR-binding sites with the lowest NRI (Ballare et al. 2013).
by facilitating binding of RAD21, YY1 and the mediator complex (Nacht et al. 2019). In these sites, C/EBPα acts as a modulator of progestin action, enabling only a single round of cell division and slowing down the growth of xenografted cells (Nacht et al. 2019). A similar facilitating role of C/EBPα had been described for GR recruitment to the genome (Grontved et al. 2013). In endometrial cancer cells, binding of PR occurs in regions that are pre-marked by PAX2 binding (La Grecia et al. 2019). These findings support the notion that gene regulation in differentiated cells depends on the combined action of various cell and sequence-specific transcription factors acting sequentially or simultaneously to fine-tune the gene expression program required.

Chromatin remodeling and gene regulation in response to hormone

How does hormone binding to PR promote nucleosome remodeling? In that respect, it is important to note that hormone-induced chromatin remodeling and gene expression is also dependent on the activation of PR attached to the cell membrane. This small fraction of the PR is indeed attached to the cell membrane via cysteine palmitoylation (Pedram et al. 2007) and forms a complex with ERα and Src (Migliaccio et al. 1998). Upon binding of progesterin, the membrane-attached PR activates Src, either directly (Boonyaratankornkit et al. 2001) or via ERα (Ballare et al. 2003), leading to the activation of the SRC/ERK/MSK1 kinase pathway. Activated ERK1–2 phosphorylates PR that translocates to the cell nucleus in the form of a ternary complex PR/extracellular signal-regulated kinases 1 and 2 (ERK1–2)/Mitogen- and stress-activated protein kinase-1 (MSK1), which once located at Pg target genes phosphorylates histone H3 at S10 (Vicent et al. 2006). This step is crucial for the regulation of gene expression. Later we discovered that many more enzymes and steps are involved in hormone-induced nucleosome remodeling and gene activation, a process that can be divided into two consecutive cycles (Vicent et al. 2009; Vicent et al. 2011; Wright et al. 2012). The first cycle takes place within the first 1–5 min after hormone exposure, involves CDK2/CyclinA-mediated phosphorylation and activation of poly (ADP-ribose) polymerase 1 (PARP1), the MLL complex and the NURF complex. Ultimately these enzymes lead to the phosphorylation and PARylation of histone H1 (H1), resulting in its displacement from chromatin (Fig. 3, left panel). The second cycle takes place between 5 and 30 min after hormone exposure and requires MSK1, the histone acetyl transferase PCAF and the BAF ATP-dependent chromatin remodeling complex, leading to additional nucleosome remodeling and the
displacement of histone H2A/H2B dimers (Fig. 3, right panel) (Vincent et al. 2006; Yang et al. 2007). Therefore, the actual activation of transcription takes place on a histone H3/H4 tetramer (Vincent et al. 2010), which can be cleaved by MNase, explaining the decrease in the density of nucleosome reads upon hormone exposure (Fig. 2). Later, we discovered that hormone-activated chromatin remodeling also requires nuclear synthesis of ATP from ADP-Ribose and PPi by NUDIX5 (Wright et al. 2016) (see subsequent section).

Most of what has been said previously was discovered by studying hormonal gene activation, but the addition of hormone to breast cancer cells also represses a substantial number of genes (Ballare et al. 2013). Curiously, around 20% of the genes that will be activated by progestins are initially silenced by a repressive complex recruited by unliganded PR and contain heterochromatin protein 1-γ (HP1γ), lysine-specific histone demethylase 1 (LSD1), histone deacetylase 1 and 2 (HDAC1/2), REST corepressor 1 (CoRest), lysine demethylase 5B (KDM5B) and the steroid receptor RNA activator (SRA). Upon hormone exposure and as a result of H3S10 phosphorylation by MSK1, the HP1γ-LSD1 repressive complex is displaced, enabling the recruitment of additional co-regulators needed for full de-repression (Vincent et al. 2013). In a subset of other genes, hormone exposure leads to the recruitment of HP1γ and Brahma-related gene-1 (BRG1) resulting in hormone-induced gene down-regulation (Vincent et al. 2013). In these genes, repression is mediated by BRG1-dependent deposition of linker histone H1.2 (Nacht et al. 2016). Likely, other mechanisms of hormone mediated gene repressions still remain to be discovered.

Role of 3D genome structure

Beyond the first level of genome packaging in nucleosomes, higher levels of genome folding have entered the field of epigenetics due substantially to the development of chromosome conformation capture techniques (Dekker et al. 2002); particularly, the genome-wide detection of chromatin proximity by ligation using Hi-C (Lieberman-Aiden et al. 2009). Hi-C and the techniques derived thereof continue to provide information about the different levels of genome folding, from chromosome territories to chromosome compartments, topologically associated domains (TADs), sub-TADs and loops (Bonev and Cavalli 2016). Formation of TADs is accepted to be generated by the mechanism of loop extrusion mediated by Cohesins and controlled by architectural proteins, such as the CCCTC-binding factor (CTCF) (Fudenberg et al. 2016). The RNA-binding region of CTCF participates in self-association and loop formation (Hansen et al. 2019; Saldana-Meyer et al. 2019).

We have used Hi-C approaches to explore the role of genome topology in gene regulation by progestins. Curiously, we found that the division of the genome in TADs of around 1 Mb size is not affected by exposure of breast cancer cells to progestins, but that many of the hormone regulated genes are clustered in a subset of hormone responsive TADs. Moreover, all the coding and non-coding genes within a regulated TAD tend to respond in the same direction, suggesting that these TADs represent units of hormone response (Le Dily et al. 2014) (Fig. 4A). Hormone activated TADs expand in size and lose histones when exposed to hormone, while hormone repressed TADs become more compacted and enriched in histones (Le Dily et al. 2014). A more detailed analysis of a few interesting TADs revealed that all the gene promoters within a TAD interact with one single 20–90 kb long hormone control region (HCR) encompassing several PR- and ERα-binding sites. An example is the TAD containing the ERS1 gene encoding Erα, where the genes are activated by estrogens and repressed by progestins (Le Dily et al. 2019) (Fig. 4A and B). In response to estrogens (+E2), the HCR interactions with the promoters within the TAD are enhanced, whereas in response to progestins (+Pg), the HCR-promoters interactions are destabilized (Fig. 4C). We identified around 200 HCRs in T47D breast cancer cells and found that, prior to hormone exposure, the HCRs interact with each other at long distances at higher frequency than expected in cells expressing ERα and PR but not in receptor negative cells of the same epithelial luminal origin (Le Dily et al. 2019) (Fig. 4D). Thus, via the HCRs, the hormone receptors contribute to the higher order structure of the genome in a cell-type-specific manner and preform the conditions for an optimal hormone response. Similar structure has been proposed using Tri-C as regulatory hubs for mouse globin loci in primary mouse erythroid cells (Oudelaar et al. 2018).

One question that has not been sufficiently explored is the role of the underlying DNA sequence information on the higher order folding of chromatin. Ultimately, if chromatin folding is subjected to evolution, one would expect that the DNA sequence carries at least part of the information. However, except for the significance of repeated elements, very little has been established on the role of DNA sequence. In the filamentous fungus Epichloë Festucae, blocks of repeated elements act as boundaries that organize the 3D genome structure and separate gene-rich genomic domains (Winter et al. 2018). In Drosophila,
SINE elements have been shown to maintain and reshape 3D chromosome structure (Courmac et al. 2016). In the human genome, MIR and L2 transposable elements share long-range interactions, acting as enhancers that shape gene regulatory networks (Cao et al. 2019). Active Alu elements are very abundant in the human genome and have binding sequence for very relevant transcription factors (Bennett et al. 2008). In breast cancer cells, we have identified a class of Alu elements bound by the CHAP complex that, in response to serum starvation, reshape the 3D genome structure by binding of general transcription factor 3C (TFIIIC). TFIIIC acetylates H3 at K18 and forms loops that contact CTCF sites contained within cell cycle genes, maintaining them ready to respond to serum addition (Ferrari et al. 2019).

In addition to repeated elements, the biophysical properties of DNA not only influence the positioning of nucleosomes (Segal et al. 2006), but can also be used to identify special regulatory elements. Very recently, it has been shown that the increased DNA flexibility and accessibility of sequences with a high propeller twist are an optimal predictor of regulatory enhancer regions (Pataskar et al. 2019). Many years ago, Giorgio Bernardi proposed that the structure of chromosomes is determined by compositional DNA structure that determines what he now calls the genomic code. He first discovered that calf thymus DNA exhibits a reversible decrease in viscosity at sub-melting temperatures that affects ~35% of the GC-poor DNA molecules (Freund and Bernardi 1963). Using chromatography of human DNA on hydroxyapatite...
columns and Cs$_2$SO$_4$/Ag$^+$ density gradients, he established the concept of isochores, large DNA sequences, ~0.9 Mb average size, that can be separated into five families of increasing GC content, decreasing size and increasing gene content, L1, L2, H1, H2, H3 (Macaya et al. 1976; Thierry et al. 1976). The isochore H3 with the highest GC content accounts for only ~3% of the human genome and is very gene rich, constituting the ‘genome core’ (Bernardi et al. 1985). FISH experiments showed that the isochores with high-GC content are located in the center of the nucleus. Their size and location correspond to the TADs as defined in Hi-C experiments (Dixon et al. 2012; Nora et al. 2012). In contrast, the low-GC content isochores represent gene deserts and are associated to the nuclear lamina, corresponding to the lamina associated domains (LADs) (Guelen et al. 2008). The isochores behave as domains of DNA replication timing, with the short gene-rich isochores replicating early in S phase and the large gene-poor isochores replicating later (Costantini and Bernardi 2008). This bimodal distribution of the compositional genome structure is reminiscent of the two A and B chromosome compartments identified in Hi-C experiments (Lieberman-Aiden et al. 2009). Moreover, in the mouse genome interspersed GC-poor LINEs and GC-rich SINEs repeats constrain the base composition of isochores. These results suggest that there is a DNA-guided folding of chromatin, a kind of genome code, which could be the genetic basis for genome topology. For further reading on this interesting proposal, see Bernardi’s recent review (Bernardi 2019).

Role of nuclear ATP synthesis

As mentioned previously, one of the key steps involved in progestin signaling is the rapid activation of PARP1. PARP1 is activated in response to hormone via CDK2 mediated phosphorylation within its NAD$^+$ binding active site. Activation in this manner increases subsequent nuclear poly(ADP-ribose) (PAR) (Wright et al. 2012). Blocking this step via inhibition of PARP1 prevents the regulation of 80% of the hormone-controlled genes and inhibits chromatin remodeling and histone H1 displacement. Initially, we assumed that the increase in PAR synthesis and the PARylation of histones was only needed for chromatin decompaction that would facilitate access for transcription factors, other histone modifying enzymes and cofactors to ensure correct transcriptional regulation (Wright et al. 2012). Unexpectedly, however, we found that the hydrolysis of PAR to its ADPR subunits by poly ADP-ribose glycohydrolase (PARG) was also required for hormonal gene regulation, a finding that led us to think about other possible functions of ADPR in hormone regulation (Wright et al. 2016), a concept that was initially formulated in the context of DNA repair (Maruta et al. 1997; Oei and Ziegler 2000; Maruta et al. 2007). Exploring the interactome of PAR in breast cancer cells we identified NUDIX5 (also known as NUDT5) as a hormone-induced interactor. This member of the large NUDIX family of enzymes is known to hydrolyze ADPR to AMP and ribose-5-P and, in principle, it could use pyrophosphate (PPI) instead of H$_2$O to convert ADPR to ATP and ribose-5-P. Therefore, we tested whether NUDIX5 was required for hormonal gene regulation. We found that depleting NUDIX5 by specific siRNAs resulted in a dramatic inhibition of hormonal gene regulation as well as of cell proliferation in response to either estrogens or progestins (Wright et al. 2016). Moreover, recombinant NUDIX5 can generate ATP when incubated with ADPR and PPI. Finally, using FRET or bioluminescence sensors for ATP visualization in living cells, we could detect a transient increase in nuclear ATP around 30 min after hormone exposure that returned to basal levels after 60 min (Wright et al. 2016) (Fig. 5A). Depleting NUDIX5 also resulted in an inhibition of hormone-induced chromatin remodeling and, therefore, we assumed that the nuclear ATP was required for the ATPases of the chromatin remodeling enzymes.

NUDIX5 is the only member of the NUDIX family that forms a homodimer (Zha et al. 2006). The NUDIX5 homodimer has two active clefts formed at the interface of the antiparallel oriented monomers. In the inactive state prior to hormone exposure, the monomers are tightly connected by ionic interaction between phosphorylated Threonine45 (T45) of one monomer and Lysine27 (K27) of the other (Wright et al. 2016). In this conformation, there is no space for PPI to enter the active cleft and the enzyme hydrolyzes ADPR (Fig. 5B, left panel). We wondered how NUDIX5 is activated upon hormone exposure and found that 1 min after adding hormone T45 is rapidly dephosphorylated (Wright et al. 2016) (Fig. 5B, right panel). Dephosphorylation of T45 weakens the interaction between the two monomers, allowing them to flip and form a hexamer that exhibits more open substrate clefts able to accept PPI (Wright et al. 2016). The phosphomimetic NUDIX5 mutant T45D cannot synthesize ATP and acts as dominant negative mutation on gene regulation (Wright et al. 2016). Thus, we have identified a novel nuclear pathway in which nicotinamide nucleotide adenyllytransferase 1 (NMNAT1) uses ATP and nicotinamide mononucleotide (NMN) to synthesize...
NAD\(^+\), that is used by PARP1 to attach PAR chains to itself and to chromatin proteins; PARG hydrolyses PAR to ADPR, which NUDIX5 can either hydrolyze to AMP or use ADPR along with PPI to synthesize ATP, depending on its phosphorylation state (Fig. 5C).

In interpreting these results, we were faced with a problem since chromatin remodeling is virtually finished 30 min after hormone exposure (Vicent et al. 2009), the time when we start to see accumulation of ATP. We reasoned that this was an excess of ATP not used by the chromatin remodeling ATPases and were wondering about its possible function. At that time a paper from Tony Hyman’s laboratory appeared in Science showing that ATP at millimolar concentrations could act as an hydrotrope that would facilitate high concentrations of macromolecules as required for macromolecular condensates and phase transitions (Patel et al. 2017). This appeared as a very plausible role for the large amount of nuclear ATP that we detected and that could serve to transiently facilitate chromatin transcription and RNA processing (Wright et al. 2019). Another possible function of mM ATP concentrations could be to bind free Mg\(^{2+}\) ions reducing their free concentration and thereby promoting chromatin decompaction (Maeshima et al. 2018). In fact, Mg\(^{2+}\) ions at mM concentrations neutralize the charge of the histone tails and have been used to precipitate chromatin in the past (Widom 1986). The energy expensive synthesis of NAD\(^+\) by MNAT1 in the nucleus using mitochondrial ATP and the conversion of NAD\(^+\) to PAR by PARP1 leads to the accumulation of large amount of chemical energy in ADPR. Part of this chemical energy is used for nuclear ATP synthesis by activated NUDIX5 to fulfill various consecutive nuclear functions, including ATP-dependent chromatin remodeling, likely also histone chaperones function, followed by chromatin decompaction as an hydrotrope and as a chelator of free Mg\(^{2+}\) ions (Wright et al. 2019) (Fig. 6). The availability of NUDIX5 specific inhibitors (Page et al. 2018) will help to clarify these processes in more sophisticated biological models, including 3D cell cultures and mouse intraductal xenografts (Sflemos et al. 2016). We have recently found that nuclear synthesis of ATP by NUDIX5 is essential for the formation of oncospheres by breast cancer cells grown in 3D cultures, due to the dependence of the cancer stem cells on nuclear ATP (Pickup et al. 2019). As NUDIX5 is overexpressed in breast tumors and correlates with poor prognosis (Wright et al. 2016), it represents a novel possible target for the pharmacological control of cancer growth. In addition, we know that nuclear ATP synthesis is needed for DNA damage repair (Wright et al. 2016). As cancer cells are addicted to DNA repair mechanisms, blocking NUDIX5 alone or in combination with PARP...
inhibitors could be another possible strategy for the pharmacological management of breast cancer.

The next frontier: liquid-liquid phase transitions of macromolecular condensates

One aspect that is becoming increasingly relevant for the complete understanding of nuclear function is the formation of macromolecular condensates and liquid-liquid phase transitions (LLPT) (Peng and Weber 2019). Given that the cell nucleus has many different structures that are not separated by membranes, including the nucleolus, Cajal bodies, paraspeckles, transcription factories and the two phases of chromatin – euchromatin and heterochromatin – LLPT is a plausible mechanism for regulating the functions of these macromolecular condensates. In particular, chromatin as a long heavily charged polymer has intrinsic properties encoded mainly in the non-structured charged tails of linker and core histones. This will facilitate phase transitions in response to changes in ionic strength or to post-translational modifications of the histone tails, in particular acetylation of lysines (Gibson et al. 2019), but also methylation of lysines and arginines, citrullination of arginines, phosphorylation of serines and threonines and ADP-ribosylation of serines (Altmeyer et al. 2015; Larsen et al. 2018), or in response to binding of regulatory proteins (for example: transcription factors, readers of epigenetic marks or members of the basic transcriptional machinery including the CTD of RNA polymerase II) (Gibson et al. 2016). The discovery that the state of phosphorylation of the CTD can drive the transition from a transcriptional condensate to a splicing condensate (Guo et al. 2019) is relevant in view of the fact that hormonal regulation influences the state of the CTD not only in terms of its phosphorylation state, but also via PADI2-mediated citrullination of arginine 1810 that favors promoter release and elongation by RNA polymerase II (Sharma et al. 2019).

In relation to the action of PR, it is important to note that the two isoforms A and B may differ in the type of transcription factor and kinases with which they interact (Bellance et al. 2013; Kaya et al. 2015) and that the PRA isoform lacks one transactivation function (Tung et al. 2006). Methylation of K464 in the activation function 1 (AF1) activates PR (Chung et al. 2014) and the triple mutant K464, K481 and R492 to QQQ makes PR hyperactive in the absence of ligand (Woo et al. 2019), indicating relevant functional interactions of the AF1. This may be physiologically relevant since very recently it has been found that transgenic mice expressing only the PRB isoform develop ovarian neoplasms, while those expressing only PRA displayed a reduced frequency of tumor development (Wetendorf et al. 2019). We know that, as in the case of androgen receptor (De Mol et al. 2015), the N-terminal unstructured region of PR can form droplets in vitro. Therefore, it could be involved in cluster formation detected in cell nuclei in response to hormone that was different for the A and B isoforms of PR (Lim et al. 1999). We are now investigating how this physical behavior of PR may be combined with the modifications of the CTD of the RNA polymerase II and with changes in chromatin PARylation or other chromatin epigenetic marks, along with changes in nuclear ATP and Mg$^{2+}$ ions, to facilitate the formation of macromolecular condensates that mediate hormonal gene regulation. In experiments with electron microscopy and immunogold, it was shown that PR binds preferentially to the condensed heterochromatin near the lamina and that, in response to hormone, there is disaggregation of the condensed chromatin to smaller...
fragments and the immunogold particles accumulate in the border between this dispersed chromatin and the nucleolus (Perrot-Applanat et al. 1986). The recent advances in high resolution microscopy and single molecule tracking will be key for exploring the dynamics of nuclear phase transitions in living cells. In particular, the transitions between the active and the inactive states of chromatin can be studied using soft X-ray tomography (Le Gros et al. 2016; Strom et al. 2017) and may be correlated with visualization of specific chromatin regions via oligo-painting and oligoSTORM (Beliveau et al. 2015; Beliveau et al. 2017) and localization of PR tagged with an appropriate fluorophore. Combined with single-cell advanced studies (Jin et al. 2015), approaches like combinatorial cellular indexing (Cusanovich et al. 2015), RNA seqFISH (Eng et al. 2019), single cell Hi-C (Nagano et al. 2017; Ramani et al. 2017; Stevens et al. 2017), single cell MNase-seq (Lai et al. 2018), ihChIP-seq (Ai et al. 2019), Dip-C (Tan et al. 2019) or CHIA-DROP (Zheng et al. 2019) will permit insight into the mechanism that uses dynamic variability in structure (Finn and Misteli 2019) and the stochastic nature of gene transcription (Raser and O’Shea 2004) to permit adaptation of the cell to changes in the environment, such as the exposure to variable levels of steroid hormones.

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Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

References
Von Der Ahe D, Janich S, Scheidereit C, Renkawitz R, Schutz G & Beato M 1985 Glucocorticoid and progesterone receptors bind to the same sites in two hormonally regulated promoters. Nature 313 706–709. (https://doi.org/10.1038/313706a0)
Ai S, Xiong H, Li CC, Luo Y, Shi Q, Liu Y, Xu X, Li C & He A 2019 Profiling chromatin states using single-cell ihChIP-seq. Nature Cell Biology 21 1164–1172. (https://doi.org/10.1038/s41556-019-0383-5)

Altmeyer M, Neelsen KJ, Teloni F, Poznyakova I, Pellegrino S, Groffe M, Rask MB, Streicher W, Jungmichel S, Nielsen ML, et al. 2015 Liquid demixing of intrinsically disordered proteins is seeded by poly(ADP-ribose). Nature Communications 6 8088. (https://doi.org/10.1038/ncomms9088)
Ballare C, Castellano G, Gaveglia L, Althammer S, Gonzalez-Vallinas J, Eyras E, Le Dily F, Zaurin R, Soronellas D, Vicent GP, et al. 2013 Nucleosome-driven transcription factor binding and gene regulation. Molecular cell 49 67–79. (https://doi.org/10.1016/j.molcel.2012.10.019)
Ballare C, Uhrig M, Bechtold T, Sancho E, Di Domenico M, Migliaccio A, Auricchio F & Beato M 2003. Two domains of the progesterone receptor interact with the estrogen receptor and are required for progesterone activation of the c-Src/Erk pathway in mammalian cells. Molecular and Cellular Biology 23 1994–2008. (https://doi.org/10.1128/MCB.23.6.1994-2008.2003)
Beato M 1989 Gene regulation by steroid hormones. Cell 56 335–344. (https://doi.org/10.1016/0092-8674(89)90237-7)
Beliveau BJ, Boettiger AN, Avendano MS, Jurgenmann M, McCole RB, Joyce EJ, Kim-Kiselak C, Bantigieves F, Fonseka CY, Erreg J, et al. 2015 Single-molecule super-resolution imaging of chromosomes and in situ haplotype visualization using Oligopaint FISH probes. Nature Communications 6 7147. (https://doi.org/10.1038/ncomms8147)
Beliveau BJ, Boettiger AN, Nir G, Bintu B, Yin P, Zhuang X & Wu CT 2017 In situ super-resolution imaging of genomic DNA with OligoSTORM and OligoDNA-PAINT. Methods in Molecular Biology 1663 231–252. (https://doi.org/10.1007/978-1-4939-7265-4_19)
Bellance C, Khan JA, Meduri G, Guiochon-Mantel A, Lombs B & Loosfelt H 2013 Progesterone receptor isoforms PRA and PRB differentially contribute to breast cancer cell migration through interaction with focal adhesion kinase complexes. Molecular biology of the cell 24 1363–1374. (https://doi.org/10.1091/mbc.e12-11-0807)
Bennett EA, Keller H, Mills RE, Schmidt S, Moran JV, Weichenreider O & Devine SE 2008 Active Alu retrotansposons in the human genome. Genome research 18 1875–1883. (https://doi.org/10.1101/gr.081737.108)
Bernardi G 2019 The genomic code: A pervasive encoding/molding of chromatin structures and a solution of the “non-coding DNA” mystery. Bioessays e1900106. (https://doi.org/10.1002/bies.201900106)
Bernardi G, Olofsson B, Filipski J, Zerial M, Salinas J, Cuny G, Meunier-Rottival M & Rodier F 1985 The mosaic genome of warm-blooded vertebrates. Science 228 953–958. (https://doi.org/10.1126/science.4001930)
Bonev B & Cavalli G 2016 Organization and function of the 3D genome. Nature reviews Genetics 17 772. (https://doi.org/10.1038/nrg.2016.147)
Boonyaratavornkitt V, Scott MP, Ribon V, Sherman L, Anderson SM, Maller JL, Miller WT & Edwards DP 2001 Progesterone receptor contains a proline-rich motif that directly interacts with SH3 domains and activates c-Src family tyrosine kinases. Molecular Cell 8 269–280. (https://doi.org/10.1016/S1097-2765(01)00304-5)
Bruggemeier U, Rogge L, Winnacker EL & Beato M 1990 Nuclear factor A acts as a transcription factor on the MMTV promoter but competes with steroid hormone receptors for DNA binding. The EMBO journal 9 2233–2239. (https://doi.org/10.1002/j.1460-2075.1990.tb0791.x)
Cao Y, Chen G, Wu G, Zhang X, McDermott J, Chen X, Xu C, Jiang Q, Chen Z, Zeng Y, et al. 2019 Widespread roles of enhancer-like transposable elements in cell identity and long-range genomic interactions. Genome research 29 40–52. (https://doi.org/10.1101/gr.235747.118)
Carroll JS, Hickey TE, Tarulli GA, Williams M & Tilley WD 2017 Deciphering the divergent roles of progestogens in breast cancer. Nature Reviews Cancer 17 54–64. (https://doi.org/10.1038/nrc.2016.116)
Costantini M & Bernardi G 2008 Replication timing, chromosomal bands, and isochores. PNAS 105 3433–3437. (https://doi.org/10.1073/pnas.0710587105)

This work is licensed under a Creative Commons Attribution 4.0 International License.
endometrial cancer cell gene expression. bioRxiv 739466. (https://doi.org/10.1101/739466)
Lai B, Gao W, Cui K, Xie W, Tang Q, Jin W, Hu G, Ni B & Zhao K 2018 Principles of nucleosome organization revealed by single-cell micrococcal nuclease sequencing. Nature 562 281–285. (https://doi.org/10.1038/s41586-018-0567-3)
Larsen SC, Hendriks IA, Lyon D, Jensen L & Nielsen ML 2018 Systems-wide analysis of serine ADP-ribosylation reveals widespread occurrence and site-specific overlap with phosphorylation. Cell Reports 24 2493–2505 e2494. (https://doi.org/10.1016/j.celrep.2018.07.083)
Le Dily E, Bau D, Pohl A, Vicent GP, Serra F, Soronellas D, Castellano G, Wright RH, Ballare C, Filion G, et al. 2014 Distinct structural transitions of chromatin topological domains correlate with coordinated hormone-induced gene regulation. Genes and Development 28 2151–2162. (https://doi.org/10.1101/gad.241422.114)
Le Dily E, Vidal E, Cuartero Y, Quilez J, Nacht AS, Vicent GP, Carbonell-Caballero J, Sharma P, Villanueva-Canas JL, Ferrari R, et al. 2019 Hormone-control regions mediate steroid receptor-dependent genome organization. Genome research 29 29–39. (https://doi.org/10.1101/gr.248324.118)
Le Gros MA, Clowney EJ, Magklara A, Yen A, Markenscoff-Papadimitriou E, Colquitt B, Myllys M, Kellis M, Lomvardas S & Larabell CA 2016 Soft X-ray tomography reveals gradual chromatin compaction and reorganization during neurogenesis in vivo. Cell Reports 17 2125–2136. (https://doi.org/10.1016/j.celrep.2016.10.060)
Lieberman-Aiden E, van Berkum NL, Williams L, Imakaev M, Ragaçzy T, Telling A, Amit I, Lajoie BR, Sabo PJ, Dorschner MO, et al. 2009 Comprehensive mapping of long-range interactions reveals folding principles of the human genome. Science 326 289–293. (https://doi.org/10.1126/science.1181369)
Lim CS, Baumann CT, Htun H, Xian W, Irie M, Smith CL & Hager GL 1999 Differential localization and activity of the A- and B-forms of the human progesterone receptor using green fluorescent protein chimeras. Molecular Endocrinology 13 366–375. (https://doi.org/10.1210/mend.13.3.0247)
Macaya G, Thiery JP & Bernardi G 1976 An approach to the involvement of ATP produced via (ADP-Ribose)n in the transcriptional memory propagation through mitosis. iScience 29 444–451 e446. (https://doi.org/10.1016/j.iScience.2017.12.035)
Maruta H, Matsumura N & Tanuma S 1997 Role of (ADP-ribose)n catabolism in DNA repair. Biochemical and Biophysical Research Communications 236 265–269. (https://doi.org/10.1006/bbrc.1997.6910)
Maruta H, Okita N, Takasawa R, Uchiumi F, Hatanaka T & Tanuma S 2007 The involvement of ATP produced via (ADP-ribose)n in the maintenance of DNA replication apparatus during DNA repair. Biological and pharmaceutical bulletin 30 447–450. (https://doi.org/10.1248/bpb.30.447)
Migliaccio A, Piccolo D, Castoria G, Di Domenico M, Bilancio A, Lombardi M, Gong W, Beato M & Auricchio F 1998 Activation of the src/p21+raf/erk pathway by progesterone receptor via a crosstalk with estrogen receptor. The EMBO Journal 17 2008–2018. (https://doi.org/10.1093/emboj/17.7.2008)
Mohammed H, Russell IA, Stark R, Rueda OM, Hickey TE, Tarulli GA, Serandour AA, Birrell SN, Bruna A, Saadi A, et al. 2015 Progesterone receptor modulates Elkalpha action in breast cancer. Nature 523 313–317. (https://doi.org/10.1038/nature14583)
Nacht AS, Ferrari R, Zaurin R, Scabia V, Carbonell-Caballero J, Le Dily E, Quilez J, Leopoldi A, Brikken C, Beato M, et al. 2019 C/EBPa mediates the growth inhibitory effect of progestins on breast cancer cells. The EMBO Journal 38 e101426. (https://doi.org/10.15252/embj.2018101426)
Nacht AS, Pohl A, Zaurin R, Soronellas D, Quilez J, Sharma P, Wright RH, Beato M & Vicent GP 2016 Hormone-induced repression of genes requires BRG1-mediated H1.2 deposition at target promoters. The EMBO Journal 35 1822–1843. (https://doi.org/10.15252/embj.201593260)
Nagano T, Luling B, Varnai C, Dudley C, Leung W, Baran Y, Mendelson Cohen N, Wisgett S, Fraser P & Tanay A 2017 Cell-cycle dynamics of chromosomal organization at single-cell resolution. Nature 547 61–67. (https://doi.org/10.1038/nature23001)
Nee EF, Seltl LA, Trotta AP, Leach DA, Giorgio L, O’Loughlin MA, Smith E, Gill PG, Ingman WV, Graham JD, et al. 2015 The unique transcriptional response produced by concurrent estrogen and progesterone treatment in breast cancer cells results in upregulation of growth factor pathways and switching from a Luminal A to a Basal-like subtype. BMC Cancer 15 791. (https://doi.org/10.1186/1475-2820-15-1819-3)
Nora EP, Lajoie BR, Schulz EG, Giorgetti L, Okamoto I, Servant N, Piiolet T, van Berkum NL, Meisj J, Sedat J, et al. 2012 Spatial partitioning of the regulatory landscape of the X-inactivation centre. Nature 485 381–385. (https://doi.org/10.1038/nature11049)
Oei SL & Ziegler M 2000 ATP for the DNA ligation step in base excision repair is generated from poly(ADP-ribose). The Journal of Biological Chemistry 275 23234–23239. (https://doi.org/10.1074/jbc.M002429200)
Oudelaar AM, Davies JOJ, Hanssen LLP, Telenius JM, Schwessinger R, Liu Y, Brown JM, Downes DJ, Chiariello AM, Bianco S, et al. 2018 Single-allele chromatin interactions identify regulatory hubs in dynamic compartmentalized domains. Nature Genetics 50 1744–1751. (https://doi.org/10.1038/s41588-018-0253-2)
Page BDG, Valerice NCK, Wright RHG, Walliser O, Isaksossn R, Carter M, Rudd SG, Loseva O, Jemth AS, Almlof I, et al. 2018 Targeted NUDT5 inhibitors block hormone signaling in breast cancer cells. Nature Communications 9 250. (https://doi.org/10.1038/s41467-017-02293-7)
Palozola KC, Lerner J & Zaret KS 2019 A changing paradigm of transcriptional memory propagation through mitosis. Nature Reviews Molecular Cell Biology 20 55–64. (https://doi.org/10.1038/s41580-018-0077-2)
Pataskar A, Vanderlinden W, Emminger J, Singh A, Liptert J & Tiwari VK 2019 Deciphering the gene regulatory landscape encoded in DNA by biophysical features. iScience 21 638–649.
Patel A, Malinovska L, Saha S, Wang J, Alberti S, Krishnan Y & Hyman AA 2017 ATP as a biological hydrotrop. Science 356 753–756. (https://doi.org/10.1126/science.aaf846)
Payvar F, DeFrando D, Firestone GL, Edgar B, Wrangle O, Okret S, Gustafsson JA & Yamamoto KR 1983 Sequence-specific binding of glucocorticoid receptor to MTV DNA at sites within and upstream of the transcribed region. Cell 35 381–392. (https://doi.org/10.1016/S0092-8674(83)90171-X)
Pedram A, Razandi M, Sainson RC, Kim JK, Hughes CC & Levin ER 2007 A conserved mechanism for steroid receptor translocation to the plasma membrane. Journal of Biological Chemistry [in press].
Peng A & Weber SC 2019 Evidence for and against liquid-liquid phase separation in the nucleus. Noncoding RNA 5 50. (https://doi.org/10.3399/nrcnrn050050)
Perrot-Applanat M, Groyer-Picard MT, Logeat F & Milgrom E 1986. Ultrastructural localization of the progesterone receptor by an immunogold method: Effect of hormone administration. Journal of Cell Biology 102 1191–1199. (https://doi.org/10.1083/jcb.102.4.1191)
Pickup KE, Pardow F, Carbonell-Caballero J, Lioutas A, Villanueva-Canas JL, Wright RHG & Beato M 2019 Expression of oncogenic drivers in 3D cell culture depends on nuclear ATP synthesis by NUDT5. Cancers (Basel) 11 1337. (https://doi.org/10.3390/cancers11091337)
Progesterone action in breast cancer

Tan L, Xing D, Daley N & Xie XS 2019 Three-dimensional genome structures of single sensory neurons in mouse visual and olfactory systems. *nature structural and molecular biology* 26 297–307. (https://doi.org/10.1038/s41594-019-0205-2)

Theodorou V, Stark R, Menon S & Carroll JS 2013 GATA3 acts upstream of FOXA1 in mediating ESR1 binding by shaping enhancer accessibility. *Genome research* 23 12–22. (https://doi.org/10.1101/gr.139469.112)

Thiery JP, Macaya G & Bernardi G 1976 An analysis of eukaryotic genomes by density gradient centrifugation. *Journal of Molecular Biology* 108 219–235. (https://doi.org/10.1016/0022-2836(76)90104-0)

Truss M, Bartsch J, Schellert A, Haché RJG & Beato M 1995 Hormone induces binding of receptors and transcription factors to a rearranged nucleosome on the MMTV promoter in vivo. *The EMBO journal* 14 1737–1751. (https://doi.org/10.1002/j.1460-2075.1995.tb07163.x)

Tung L, Abdel-Hafiz H, Shen T, Harvell DM, Nitao LK, Richer JK, Sartorius CA, Takimoto GS & Horwitz KB 2006 Progesterone receptors (PR)-B and -A regulate transcription by different mechanisms: AF-3 exerts regulatory control over coactivator binding to PR-B. *Molecular Endocrinology* 20 2656–2670. (https://doi.org/10.1210/me.2006-0105)

Vicent GP, Ballaré C, Nacht AS, Clausell J, Subtil-Rodríguez A, Jordan A & Beato M 2006 Induction of progesterone target genes requires activation of Erk and Msk kinases and phosphorylation of histone H3. *Molecular Cell* 24 367–381. (https://doi.org/10.1016/j.molcel.2006.01.011)

Vicent GP, Nacht AS, Font-Mateu J, Castellano G, Gavaglia L, Ballare C & Beato M 2011 Four enzymes cooperate to displace histone H1 during the first minute of hormonal gene activation. *Genes and Development* 25 845–862. (https://doi.org/10.1101/gad.215993.111)

Vicent GP, Nacht AS, Smith CL, Peterson CL, Dimitrov S & Beato M 2004 DNA instructed displacement of H2A and H2B at an inducible promoter. *Molecular Cell* 16 439–452. (https://doi.org/10.1016/j.molcel.2004.10.025)

Vicent GP, Nacht AS, Zaurin R, Font-Mateu J, Soronellas D, Le Dilly F, Reyes D & Beato M 2013. Unliganded progesterone receptor-mediated targeting of an RNA-containing repressive complex silences a subset of hormone-inducible genes. *Genes and Development* 27 1179–1197. (https://doi.org/10.1101/gad.215993.113)

Vicent GP, Zaurin R, Nacht AS, Font-Mateu J, Le Dilly F & Beato M 2010 Nuclear factor 1 synergizes with progesterone receptor on the mouse mammary tumor virus promoter wrapped around a histone H3/H4 tetramer by facilitating access to the central hormone-responsive elements. *Journal of Biological Chemistry* 285 2622–2631. (https://doi.org/10.1074/jbc.M109.060848)

Vicent GP, Zaurin R, Nacht AS, Li A, Font-Mateu J, Le Dilly F, Vermeulen M, Mann M & Beato M 2009 Two chromatin remodeling activities cooperate during activation of hormone responsive promoters. *pLoS Genetics* 5 e1000567. (https://doi.org/10.1371/journal.pgen.1000567)

Wetendorf M, Li R, Wu S-P, Liu J, Creighton CJ, Wang T, Janardan K, Wilson CJ, Lanz RB, Murphy BD, et al. 2019. Constitutive expression of the progesterone receptor isoforms promotes hormone-dependent development of ovarian neoplasms. *bioRxiv* 816934. (https://doi.org/10.1101/816934)

Widom J 1986 Physicochemical studies of the folding of the 100 A nucleosome filament into the 300 A filament. Cation dependence. *Journal of Molecular Biology* 190 411–424. (https://doi.org/10.1002/0022-2836(86)90012-4)

Winter DJ, Ganley ARD, Young CA, Liachko I, Schardl CL, Dupont PY, Berry D, Ram A, Scott B & Cox MP 2018 Repeat elements organise 3D genome structure and mediate transcription in the filamentous fungus *Epichloë festucae*. *PLoS Genetics* 14 e1007467. (https://doi.org/10.1371/journal.pgen.1007467)
Woo ARE, Sze SK, Chung HH & Lin VC 2019 Delineation of critical amino acids in activation function 1 of progesterone receptor for recruitment of transcription coregulators. Biochimica et Biophysica Acta-Gene Regulatory Mechanisms 1862 522–533. (https://doi.org/10.1016/j.bbagrm.2019.01.004)

Wright RH, Lioutas A, Le Dily F, Soronellas D, Pohl A, Bonet J, Nacht AS, Samino S, Font-Mateu J, Vicent GP, et al. 2016 ADP-ribose-derived nuclear ATP synthesis by NUDIX5 is required for chromatin remodeling. Science 352 1221–1225. (https://doi.org/10.1126/science.aad9335)

Wright RHG, Castellano G, Bonet J, Le Dily F, Font-Mateu J, Ballare C, Nacht AS, Soronellas D, Oliva V & Beato M 2012 CDK2-dependent activation of PARP-1 is required for hormonal gene regulation in breast cancer cells. Genes and Development 26 1972–1983. (https://doi.org/10.1101/gad.193193.112)

Wright RHG, Le Dily F & Beato M 2019 ATP, Mg(2+), nuclear phase separation, and genome accessibility. Trends in Biochemical Sciences 44 565–574. (https://doi.org/10.1016/j.tibs.2019.03.001)

Yang X, Zaurin R, Beato M & Peterson CL 2007 Swi3p controls SWI/SNF assembly and ATP-dependent H2A-H2B displacement. Nature Structural and Molecular Biology 14 540–547. (https://doi.org/10.1038/nsmb1238)

Zaret KS and Carroll JS 2011 Pioneer transcription factors: establishing competence for gene expression. Genes and Development 25 2227–2241. (https://doi.org/10.1101/gad.176826.111)

Zaret KS, Lerner J & Iwafuchi-Doi M 2016 Chromatin scanning by dynamic binding of pioneer factors. Molecular cell 62 665–667. (https://doi.org/10.1016/j.molcel.2016.05.024)

Zha M, Zhong C, Peng Y, Hu H & Ding J 2006 Crystal structures of human NUDT5 reveal insights into the structural basis of the substrate specificity. Journal of Molecular Biology 364 1021–1033. (https://doi.org/10.1016/j.jmb.2006.09.078)

Zheng M, Tian SZ, Capurso D, Kim M, Maurya R, Lee B, Piecuch E, Gong I, Zhu JJ, Li Z, et al. 2019 Multiplex chromatin interactions with single-molecule precision. Nature 566 558–562. (https://doi.org/10.1038/s41586-019-0949-1)

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