Mitogen- and stress-activated protein kinase 1 is required for gonadotropin-releasing hormone–mediated activation of gonadotropin α-subunit expression

Majd Haj1, Andrea Wijeweera1, Sergei Rudnizky1, Jack Taunton6, Lilach Pnueli1, and Philippa Melamed11

From the 1Faculty of Biology, Technion-Israel Institute of Technology, Haifa 32000, Israel and the 6Department of Cellular and Molecular Pharmacology, University of California, San Francisco, California 94158

Edited by Joel Gottesfeld

Pituitary gonadotropin hormones are regulated by gonadotropin-releasing hormone (GnRH) via MAPK signaling pathways that stimulate gene transcription through activating mitogen-and stress-activated protein kinase 1 (MSK1) and the hormone-specific β-subunits of gonadotropin. We have reported previously that GnRH-induced activities at these gene promoters are mediated via histone modifications, but we did not examine histone phosphorylation. This modification adds a negative charge to residues of the histone tails that interact with the negatively charged DNA, is associated with closed chromatin during mitosis, but is increased at certain genes for transcriptional activation. Thus, the functions of this modification are unclear. We initially hypothesized that GnRH might induce phosphorylation of Ser-10 in histone 3 (H3S10p) as part of its regulation of gonadotropin gene expression, possibly involving cross-talk with H3K9 acetylation. We found that GnRH increases the levels of both modifications around the Cga gene transcriptional start site and that JNK inhibition dramatically reduces H3S10p levels. However, this modification had only a minor effect on Cga expression and no effect on H3K9ac. GnRH also increased H3S28p and H3K27ac levels and those of activated mitogen- and stress-activated protein kinase 1 (MSK1). MSK1 inhibition dramatically reduced H3S28p levels in untreated and GnRH-treated cells and also affected H3K27ac levels. Although not affecting basal Cga expression, MSK1/2 inhibition repressed GnRH activation of Cga expression. Moreover, ChIP analysis revealed that GnRH-activated MSK1 targets the first nucleosome just downstream from the TSS. Given that the elongating RNA polymerase II (RNAPII) stalls at this well-positioned nucleosome, GnRH-induced H3S28p, possibly in association with H3K27ac, would facilitate the progression of RNAPII.

Chromatin structure often responds to external cues to determine gene expression through changing the accessibility of regulatory DNA. We have previously reported that the hypothalamic gonadotropin releasing hormone (GnRH)2 up-regulates expression of the gonadotropin hormones via various actions on the chromatin. In partially differentiated gonadotropes, GnRH was seen to induce the removal of histone deacetylases from the repressed hormone β-subunit gene promoters (1, 2), while also activating expression of these genes by targeting trimethylation of histone H3 lysine 4 (H3K4me3) to their promoters (3). There are several reports that GnRH induces recruitment of transcriptional coactivators with histone acetyltransferase (HAT) activity to these gene promoters (e.g. Refs. 4–6) indicating that the histones are also acetylated as part of the induction of expression of these genes, although the specific residues affected and their precise roles in transcription have not been reported.

In contrast to histone acetylation and methylation, whose functions in transcriptional regulation are relatively well understood, the role of histone phosphorylation is much less clear (7, 8). The better studied phosphorylation of H3 at serine 10 (H3S10p) has been implicated in transcriptional regulation of various induced genes (7, 9, 10), but is also involved in the DNA damage response, apoptosis, and mitosis/meiosis (11–13), and is associated both with histone acetylation and deacetylation during mitosis and chromatin relaxation during gene expression (11, 12). Such opposing biological effects of H3 phosphorylation of the same residue indicate that the interpretation of this modification is highly context-dependent and that the individual residues cannot be considered in isolation.

Besides altering the charge of the histone that leads to a loosening of its interaction with the DNA, phosphorylation may serve as a recognition signal for other regulatory proteins, although the only proteins found to bind specifically to phosphorylated histones in association with gene transcription are the 14–3–3 family members that can recruit modifying enzymes such as HATs for subsequent histone acetylation (14–19). Indeed H3 phosphorylation and acetylation are often seen together at active promoters and both are crucial for the activation of certain genes, such as the immediate early genes following stimulation of MAPK pathways (10, 20, 21). Furthermore, HAT enzymes such as GCN5, which targets H3K9ac, were shown to acetylate preferentially H3S10p peptides over protein; qPCR, quantitative PCR; HAT, histone acetyltransferase; MSK1, mitogen- and stress-activated protein kinase 1; TSS, transcriptional start site.
non-phosphorylated ones (22), providing a possible mechanism for such cross-talk between these modifications.

A number of kinases are reported to phosphorylate H3, and the mitogen-activated protein kinases (MAPKs) MSK and RSK have been implicated to mediate the response to various extracellular stimuli (12, 23–25). These kinases are downstream of the ERK and p38 MAPK that mediate many signaling pathways, and MSK1/2 knock-out mice were seen to have vastly reduced phospho-H3 levels (26). However, MSK1/2 also target various transcription factors (27, 28). Another MAPK, JNK which mediates part of the stress response, translocates to the nucleus following its activation and is also sometimes associated with chromatin (23), was shown to phosphorylate specifically H3S10 at its target genes during neuronal differentiation. This was shown to involve JNK recruitment to the chromatin via NF-Y, which was essential for gene expression (30). Other kinases are also capable of phosphorylating H3S10, although the specific and diverse roles and contributions of each of these events to the transcriptional regulation are not understood (12).

GnRH exerts its actions on the gonadotropes to induce gonadotropin gene expression through the activation of various MAPKs including JNK (31, 32), while the Cga gene promoter contains an NF-Y binding motif, suggesting a possible pathway for GnRH induction of HS10p, which might up-regulate Cga expression. We thus considered it highly likely that gonadotrope exposure to GnRH induces histone phosphorylation to activate this gene, possibly involving also cross-talk with H3 acetylation. We thus aimed to examine the GnRH-induced changes in histone acetylation and phosphorylation of the Cga gene and their role in Cga gene expression. This led us to understand that phosphorylation at neither H3S10 nor H3S28 is required for the on-going transcription of this gene, but H3S28p is elevated by GnRH particularly at the +1 nucleosome, where it likely helps transition through the first nucleosome, and is required for the full Cga transcriptional response to GnRH.

Results

The open chromatin at the Cga gene transcriptional start site (TSS) is enriched with phosphorylated H3S10 compared with the more closed chromatin at the β-subunit genes

The αT3-1 cell line represents an immature partially-differentiated gonadotrope, and expresses high levels of the Cga gene, which encodes the common α subunit, but only low levels of the hormone-specific β-subunit genes, Lhb and Fshb (Fig. 1A). To examine whether the phosphorylation status of the histones around the TSSs of these genes correlates with their diverse expression levels, we first examined the levels of phosphorylation at H3S10 (H3S10p). Chromatin immunoprecipitation (ChIP) using antisera to H3S10p showed higher levels of this modification at the Lhb and Fshb genes as compared with the Cga gene (Fig. 1B). However H3 is found at much lower levels on this region of the Cga gene than on the β-subunit genes (Fig. 1C), and when the levels of this modification were normalized to those of total H3, the nucleosomes were clearly enriched with this modification at the Cga gene, correlating with the higher gene expression levels (Fig. 1D).

Basal levels of H3 histone acetylation at Lys-9, -18, and -27 also largely correlate with gene expression

Histone acetylation is characteristically found at actively expressed genes, and levels of H3K9ac at these genes often correlate with those of H3S10p. Although we saw previously that the β-subunit genes are associated with histone deacetylases in these cells, the actual acetylation status of these genes, and how this compares with the acetylation levels at the more active Cga gene, are not known. We therefore measured, also by ChIP, the levels of acetylation of lysines 9, 18, and 27 of histone H3 at the same upstream regions of these three genes. The levels of acetylation at each of these lysine residues normalized to the level of total H3 in the same samples correlated with levels of expression, although K18ac and K27ac were at similarly low levels at the two β-subunit genes (Fig. 2, A–C).

Exposure of these cells to GnRH elevates levels of H3S10p and H3 acetylation globally and at the Cga gene, correlating with an increase in its expression

GnRH is a major regulator of gene expression in the gonadotropes as the activation of MAPK pathways target many genes, not just the gonadotropins, so we first assessed whether it had any global effects on histone acetylation and/or phosphorylation. Western analysis of histone extracts from cells exposed to 0–6 h GnRH revealed that although there was an initial drop in H3S10p levels, global levels of this modification and K27ac then increased, whereas the H3K9ac and K18ac appeared unaffected on a global scale (Fig. 3A).

Although expression of all three gonadotropin genes is induced by GnRH, these cells are considered the most appropriate model for study of the GnRH stimulation of the Cga gene, due to its higher expression levels and more predictable response. After confirming that GnRH elevates Cga mRNA levels in these cells (Fig. 3B), we went on to examine whether there was change in levels of H3S10p and H3ac at the Cga gene. ChIP was carried out with antisera to the particular modified histone and also total H3 for normalization. Acetylation of all three lysines was increased in the GnRH-treated cells but the response of K27ac was the most notable (Fig. 3, C–E). Phosphorylation of H3S10 was also significantly, albeit marginally, elevated after GnRH treatment (Fig. 3F).

JNK plays a crucial role in determining H3S10p levels, but this histone modification barely affects basal Cga gene expression and its absence does not influence H3K9ac

GnRH signaling to gonadotropin gene expression is through the activation of various MAPK phosphorylation cascades including JNK (31). JNK was reported to target H3S10p in neuronal cells, being recruited by the NF-Y transcription factor (30), which also has a binding motif on the Cga gene promoter (33). Therefore we examined whether the levels of H3S10p at the Cga gene might be regulated by JNK, first through exposure of the cells to a chemical JNK inhibitor, SP600125. This inhibitor dramatically reduced global levels H3S10p, which were not restored by exposure of the cells to GnRH (Fig. 4, A and B). This effect of the JNK inhibitor on H3S10p was apparent also specifically at the Cga gene in ChIP analysis, which revealed that the
inhibitor reduced H3S10p levels to below 20% those in the control cells (Fig. 4C).

To determine the effect of this drop in H3S10p on Cga gene expression, we carried out quantitative PCR (qPCR) in serum-starved cells exposed to the inhibitor for 30 min before and during treatment with GnRH for 8 h. Surprisingly, the inhibitor had no apparent effect on the basal Cga mRNA levels, although the GnRH response was reduced somewhat (Fig. 4D), in line with previous reporter gene assays on the activity of this promoter (34). To determine whether an effect on basal gene expression might be seen during a longer time frame, we exposed cells to the inhibitor for up to 24 h; still there was only a minor effect on Cga mRNA levels that dropped, at the most, to around 70% that of the untreated controls (Fig. 4E). To confirm this result, we also transfected the cells with a construct to express a dominant-negative CDC42, which is upstream of JNK, and a similar effect was seen (Fig. 4F).

To investigate whether this drop in H3S10p is accompanied by a drop in the neighboring acetylation of H3K9, we carried out ChIP for this modification in the SP600125-treated cells. The levels of H3K9ac at the Cga gene promoter/TSS were unaffected by the JNK inhibition (Fig. 4G) and a similar lack of effect on H3K9ac was seen at a global level in Western analysis (Fig. 4H). Together, these results show clearly that JNK is responsible for H3S10p at the Cga gene in these cells, but this modification plays little role in determining the levels of Cga expression, and does not influence the levels of H3K9ac.

Phosphorylation at H3S28 also increases following GnRH, correlating with K27ac, and this plays a role in GnRH-induced gene expression

As H3S10p did not appear to play much of a role in determining Cga gene expression levels, we next examined phosphorylation at H3S28 (H3S28p), which is thought to be associated with transcriptional activation, but is much less studied than H3S10p. Initial global Western analysis in αT3-1 cells revealed that, like H3S10p, levels of this modification also dropped during the first hour after GnRH exposure, but it increased by 3 h of GnRH treatment (Fig. 5A). A similar effect was seen with another gonadotrope cell line, LβT2, although both the initial drop and increase above basal levels were somewhat delayed, the maximum response (3.12 ± 0.7; n = 5) was also reached
after 4–6 h (Fig. 5B). H3S28p reportedly can be phosphorylated by MSK (9, 11, 13, 35), which is downstream of the ERK and p38 MAPK signaling pathways both of which are activated by GnRH, with ERK being particularly crucial to up-regulation of the Cga gene (31). We therefore tested the effects of a broad chemical inhibitor of MSK, H89, which was applied to the cells 30 min before and during exposure to GnRH. The inhibitor virtually abolished the GnRH stimulation of Cga gene expression, but did not affect the basal levels (Fig. 5C), suggesting a possible role for this pathway in the GnRH up-regulation of this gene.

We went on to examine whether H3S28p is induced by GnRH specifically at the Cga gene. For these experiments we utilized high resolution ChIP (36) which, through extensive sonication and multiple primer sets, is able to distinguish between the +1 nucleosome, which is the first one downstream of the TSS, and the −1 nucleosome, which is the closest of the promoter nucleosomes to the TSS, located 400 bp upstream (36). ChIP for H3 after this protocol is able to differentiate clearly between these two nucleosomes separated by a nucleosome-free region, and both nucleosomes are affected by GnRH (Fig. 5D). Subsequently, ChIP analysis for H3S28p revealed that the levels of phosphorylation at both nucleosomes are enriched following GnRH treatment, which was mimicked by the levels of H3K27ac (Fig. 5, E and F). Clearly actions on either one of these nucleosomes has distinct implications for the initiation and/or elongation mechanisms of regulating transcription.

**GnRH activates MSK, which is required for S28p and also K27ac**

We next examined whether GnRH-activated MSK might be responsible for H3S28p at either of these nucleosomes. West-
GnRH targets H3S28p via MSK to activate Cga transcription

Figure 3. Exposure to GnRH elevates levels of H3S10p and H3ac globally and at the Cga gene, correlating with its increase in expression. A, cells treated with GnRH for 0–6 h were harvested at the same time, and Western blot analysis was carried out on the histone fraction using antibody to H3S10p, H3K9ac, H3K18ac, and H3K27ac, following stripping of the membrane after each reaction. Total histone H3 was used as the loading control. (The experiment was performed several times and blotting was performed in various sequences, but for the results shown here, the sequence was H3K27ac, H3, K18ac, S10p, and K9ac.) Relative levels of each modified protein signal are shown below the blot as fold over the level in untreated cells, after normalization to H3. B, serum-starved αT3-1 cells were treated with GnRH for 8 h after which total RNA was extracted for qPCR. Cga mRNA levels are shown after normalization with those of Rplp0, and relative to the mean in untreated control cells, as described in the legend to Fig. 1. Statistical analysis was Student’s t test: **, p < 0.01. C–F, ChIP analysis at the Cga gene promoter/5′ end was carried out in untreated or GnRH-treated αT3-1 cells using antibodies for total H3 and: C, H3K9ac; D, H3K18ac; E, H3K27ac; or F, H3S10p. Levels of precipitated DNA were quantified by qPCR and analyzed as described in the legend to Fig. 1, presented relative to the mean levels in the untreated control; *, p < 0.05; **, p < 0.01; ***, p < 0.001.

ern blotting of the active phosphorylated MSK1 (MSK1p) showed that its levels are indeed increased in αT3-1 cells within 30 min of GnRH treatment (Fig. 6A), and a similar effect was seen after exposure to PMA for 1 h (Fig. 6B). The LβT2 gonadotropes responded similarly to both GnRH (Fig. 6C) and PMA (Fig. 6D), with 4–6.5-fold increases over control levels. Moreover, we were able to reduce H3S28p levels dramatically through the use of the highly specific MSK-1 inhibitor, RMM-64 (Fig. 6E). The levels of phosphorylated H3S28, after exposure to 3–10 μM of the inhibitor, dropped even more than those of other known MSK1 targets, CREB and ATF1, which are both phosphorylated also by other kinases.

Subsequently we tested whether this inhibitor also affects the levels of H3K27ac, which might suggest cross-talk between these modifications. There was a clear drop in the levels of H3K27ac in cells exposed to the inhibitor, and neither this modification, nor the H3S28p could be rescued by GnRH, whereas there was no apparent effect on levels of H3K18ac (Fig. 6F). Confirmation of the role of MSK1 in H3S28p was then provided through shRNA-mediated knockdown (KD) of MSK1, which significantly reduced S28p levels, although it had less effect on K27ac. It is not clear whether this discrepancy is due to the different efficacy in reducing H3S28p levels, and/or might be due to nonspecific effects of the inhibitor on the K27ac. Notably, however, no effect was apparent on basal Cga mRNA levels (Fig. 6G).

GnRH-induction of Cga expression requires MSK induced H3S28p at the +1 nucleosome

Having established that the H3S28p is regulated by MSK1, that MSK1 is activated following GnRH treatment, and that this can be blocked most effectively by the specific inhibitor, we sought to determine how this modification affects Cga transcription. The dramatic reduction in H3S28p levels by incubation of the cells with the MSK inhibitor, as after the MSK1 KD (Fig. 6G), did not appear to affect the basal levels of Cga expression, but clearly repressed GnRH-induced up-regulation of the gene in both gonadotrope cell lines (Fig. 7, A and B). Moreover, in confirmation of the specificity of the effect of the inhibitor, this repressive effect was significantly abated in cells expressing a RMM-64-insensitive MSK1 mutant (Fig. 7C).
High resolution ChIP analysis of H3S28p revealed that the inhibition of MSK affected primarily the GnRH-induced levels of this modification at the +1 nucleosome, although the basal levels at the −1 nucleosome were also affected (Fig. 7, D and E). This suggests that phosphorylation of H3S28 at the upstream nucleosome is quite easily removed when MSK is not active, but that H3S28p at this nucleosome plays little role in basal rates of on-going transcription, as seen by the lack of effect on basal Cga expression (Fig. 7, A and B). In contrast, the GnRH-induced increase in H3S28p at the +1 nucleosome implies a role in promoter escape and/or the early steps in transcriptional elongation.

**Discussion**

GnRH is the major central regulator of the reproductive axis due to its role in stimulating production of the gonadotropin hormones, primarily via several MAPK pathways that activate gene-specific transcription factors (31, 32). Despite a crucial role for these kinases in increasing gonadotropin gene expression, there have not yet been any reports of a role for histone phosphorylation in the activation the GnRH target genes. In fact, although phosphorylation is a crucial and central player in intracellular signaling pathways in general, the role of histone phosphorylation as an end point in these pathways to activate gene transcription has remained understudied. We report here that, although H3S10p is found at all three gonadotropin genes in unstimulated gonadotropes, and is relatively enriched on the more highly expressed Cga gene, neither H3S10p nor H3S28p play much of a role in determining basal levels of Cga expression. Nevertheless, GnRH increases phosphorylation at both serines at the 5’ end of this gene, as well as increasing H3 acety
GnRH targets H3S28p via MSK to activate Cga transcription

The GnRH-induced increase in H3S28p at the first nucleosome downstream of the TSS appears crucial for the GnRH stimulation of Cga expression, presumably facilitating RNAPII transition of this nucleosome and promoter escape (Fig. 7F).

Although JNK is activated by GnRH (31) and is clearly essential for the basal phosphorylation of H3S10, this modification surprisingly has very little effect on the on-going transcription of the Cga gene. This is in strong contrast with the situation in differentiating neuronal cells in which JNK-mediated phosphorylation of H3S10 is essential for expression of the target genes (30). The fact that the dramatic reduction in H3S10p levels following JNK inhibition also did not affect acetylation of the neighboring H3K9 indicates that this acetylation is JNK-independent and supports a previously suggested model that H3K9ac and H3S10p may be independent through spatially linked processes, and that phosphorylation is not necessarily a prerequisite for this acetylation (13, 35).

Reduction in the basal levels of H3S28p at the upstream nucleosome also had no apparent effect on the basal levels of Cga transcription, and was a transient response globally after GnRH exposure, presumably as a result of rapid phosphatase activation (32). However, after 3–6 h of GnRH, when increases in Cga expression are first noted, this modification was increased at both nucleosomes, as was H3K27ac. The link between H3S28p and H3 acetylation in stimulus-induced transcription is less well studied than for H3S10p, but cross-talk has been noted in several contexts, including at stress-responsive promoters where H3S28p is accompanied by increased histone acetylation (13), while targeting MSK1 to the /H9251-globin promoter induced both H3K27ac and H3S28p (37). More recently it was demonstrated that H3S28p increases directly p300/CBP-dependent transcription in vitro, and MSK1 was seen to facilitate p300 recruitment to a chromatin template, leading to the proposition that H3S28p helps stabilize p300 association with the chromatin (25).

Our findings suggest a crucial role for MSK1 in the phosphorylation of H3S28 and in vitro kinase assays have previously shown that it is a direct target of this kinase (26, 38, 39). Moreover, MSK1 was shown to bind directly to the MLL1 complex (40), which we previously reported is found at the Cga gene TSS (3). We cannot exclude the possibility that the inhibitor also has other nonspecific effects; generally negligible effects on other kinases were reported (25), but the increase in S28p and CREBp...
at the lowest doses suggest a possible inhibition of the phosphatases that target these proteins although not ATF-1. In this study, however, we utilized the doses that were clearly inhibitory to the kinase activity. The MSK1/2 might also phosphorylate transcription factors that are responsible for recruiting the HAT enzymes to this gene, although CREB does not mediate the GnRH effect on this gene (41). It is also possible that MSK1/2 itself recruits HAT enzymes to the nucleosome; MSK1 was shown previously to co-immunoprecipitate with multiple HATs including CBP and p300 both of which acetylate H3K27 (25, 42). However, in the context of these other findings, our results suggest that H3S28p facilitates H3K27ac at the 5′ end of the Cga gene as part of the GnRH-induced transcriptional response.

The precise roles of promoter H3S10p and S28p in basal Cga expression are not yet clear, but MSK was previously reported to cause displacement of repressive polycomb group proteins in the activation of a subset of genes via various signaling-response and neuronal differentiation pathways involving H3S28p (43). It is thus possible that these modifications are required primarily for the initial activation of closed chromatin, whereas in gonadotrope cells in which the Cga gene is already expressed, histone phosphorylation has little effect on the already open chromatin.

In these cells, the chromatin structure of the Cga, although it is a highly inducible gene, is more characteristic of a readily-expressed gene. In its basal state, the active proximal promoter is relatively free of nucleosomes, rendering it highly accessible for binding and activation by transcription factors. In contrast, the first nucleosome downstream of the TSS is well positioned, and provides an additional level of control by forming a strong barrier for early elongation (36). Such a barrier causes RNAPII stalling and backtracking, from which the RNAPII can proceed only after a recovery process involving spontaneous wrapping and unwrapping of the nucleosome (44–46). We have shown previously that incorporation of the H2A.Z histone variant...
allows greater mobility of the nucleosome to help overcome this barrier (36), whereas here we demonstrate that histone H3 phosphorylation, which reduces the histone-DNA charge interactions, likely also plays a role in the recovery from this backtracked state. Likewise, this barrier to elongation would be facilitated by the elevation in histone acetylation at H3K27, which removes a positive charge on the histone tail, and is reportedly facilitated by H3S10p-mediated HAT recruitment (25). Interestingly, a similar role has been shown in Drosophila for H3S10p, which is phosphorylated by JIL-1, a kinase essential for early transcriptional elongation; the 14-3-3 proteins recruited by H3S10p interact with the HAT Elongator protein 3, which acetylates H3K9 and this is also required for elongation to proceed (19).

The present study has thus demonstrated a specific role for H3S28 phosphorylation and reveals that it is regulated by a MAPK-activated pathway though MSK1. Furthermore, this study emphasizes the importance of context in understanding the role of histone phosphorylation at specific nucleosomes. GnRH modulates the chromatin of the Cga gene to activate its expression primarily at the level of transcription elongation by modulating the +1 nucleosome barrier, so facilitating RNA pol II progression. The GnRH induction of H3S28 phosphorylation by MSK1 thus plays a major role in regulating the Cga expression, by targeting this nucleosome.

These findings corroborate the recent report that H3S28p is a key event in stimulus-activated transcription in macrophages (25) and support the idea that this may be a widespread event in the induction of gene expression in response to specific stimuli, as a consequence of the activation of the commonly utilized MAPK signaling pathways. However, our study also places this in the context of the first nucleosome downstream of the TSS, with implications for promoter escape, linking the known activation of MAPK cascades as part of common signal transduction cascades to diverse downstream epigenetic effects at the target genes.

Experimental procedures

Cell culture

The murine gonadotrope αT3-1 and LβT2 cells (gifts from P. L. Mellon, University of California, San Diego) were cultured as reported (41), with serum removed (LβT2) or reduced to 0.5% (αT3-1) for 16 h before exposure to GnRH or PMA (both 100 nM), H89 (13.5 μM), SP600125 (all Sigma) or RMM-64 (both 10 μM; 25), with the inhibitors being added 30–120 min

![Figure 7. GnRH induction of Cga expression requires MSK induced S28p at the +1 nucleosome.](image)
prior to the GnRH. Length of exposure to GnRH was characteristically 6–8 h for assessment of effects on gene expression, and 2–5 h when measuring histone modifications, according to the initial empirical evaluation of optimal response. After replacement of serum and antibiotics (30–60 min), cells at 70–80% confluence were transfected with plasmids expressing a dominant-negative CDC42 (a gift from BC Low, National University of Singapore) or MSK1 C440V (25), using PolyJet In Vitro DNA transfection reagent (Signagen) according to the manufacturer's instructions (3 μl reagent; 1 μg of DNA). MSK1 was knocked down using the targeting sequence: AAAGT-GAGCTTCCGTATCC in pSUPER, as previously described (41).

Quantitative PCR analysis
RNA was extracted using TRIzol (Invitrogen), total RNA was reverse transcribed using a High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) and real-time quantitative PCR (qPCR) carried out using PerfeCTa SYBR Green FastMix (Quanta Biosciences, Gaithersburg, MD) all as previously described (3). The primers for detection of Cga mRNA levels were, forward, ATGGATTACTACAGAAAATATGCAG and reverse, CCTGAATAATAAAGTTCTCAGG, and for Msk1, forward, TCTTACTGTCAGGCACG and reverse, AGGTGTTCCAGGGCAACACA. The normalizing gene was RpLp0, with forward primer, GGCACCTGGAAGTCCAACTA and reverse primer, ATCTGCTTGGAGCCCACAT. A standard curve, generated by 1:2 serial dilutions, was used to determine the levels in each sample. All reactions were repeated three times and averaged.

Protein isolation and purification
Cells were placed on ice and washed with ice-cold PBS before harvest in fresh PBS, transferred to a pre-cooled microcentrifuge tube, and centrifugation at 3000 rpm for 5 min at 4°C. The supernatant was removed leaving the cell pellet on ice. For whole cell extract, cells were lysed in radioimmunoprecipitation assay buffer (RIPA: 0.1% SDS, 0.5% sodium deoxycholate, 1% NP-40, 2 mM EDTA, 150 mM NaCl, 50 mM Tris-HCl, pH 8) containing protease and phosphatase inhibitors (2 μM/ml of aprotinin; 10 μg/ml of leupeptin; 1 μg/ml of pepstatin A, 1 mM Na3VO4, 1 mM Na2VO4) for 30 min. After centrifugation at 12,000 rpm for 20 min at 4°C, the supernatant was removed leaving the cell pellet on ice. For the isolation of cytoplasmic and nuclear proteins, the NE-PER nuclear and cytoplasmic extraction kit (Pierce) was used, as previously described in Ref. 3, centering at 4°C, with 1% NP-40, 2 mM EDTA, 150 mM NaCl, 50 mM Tris-HCl, pH 8) containing protease and phosphatase inhibitors (2 μM/ml of aprotinin; 10 μg/ml of leupeptin; 1 μg/ml of pepstatin A, 1 mM Na3VO4, 1 mM Na2VO4) for 30 min. After centrifugation at 12,000 rpm for 20 min at 4°C, the supernatant was collected and transferred into a fresh tube. For the isolation of cytoplasmic and nuclear proteins, the NE-PER nuclear and cytoplasmic extraction kit (Pierce) was used, as previously described (47). The isolation of histones, cells were lysed in Triton extraction buffer (TEB: PBS containing 0.5% Triton X-100 (v/v)) containing the protease and phosphatase inhibitors (as above). After centrifugation at 2000 rpm for 10 min at 4°C, the pellet was washed in one-half volume of TEB and centrifuged as before. The supernatant was removed, the pellet was resuspended in 0.2 M HCl, and the samples were incubated overnight at 4°C on ice. The samples were then centrifuged at 2000 rpm for 10 min at 4°C and the supernatant was collected and transferred into a fresh tube. The protein concentrations were determined via Bradford Protein Assay reagent (Bio-Rad) according to the manufacturer's instructions, following which the samples were used immediately or stored at −80°C.

Western blot analysis was carried out as reported previously (47), after loading the same amounts of protein in each lane of a 10 or 12.5% SDS-polyacrylamide electrophoresis gel and with nitrocellulose membranes. The membrane was incubated with a primary antibody diluted to 1,000–15,000 overnight at 4°C or for 2 h at room temperature, and after washing with 1× PBST, the membrane was incubated with horseradish peroxidase-conjugated anti-mouse, -rabbit, or -rat diluted to 1:10,000 in 1× PBST for 1 h at room temperature. The immunoreactive proteins were detected using the Super Signal Pico West chemiluminescent system (Pierce Chemical Co.) according to the manufacturer's instructions, followed by exposure in the Las-4000 image analyzer (Fujiﬁlm) and analysis using the Imagel software (NIH). The magnitude of each signal peak was measured as a percent of the total size of the measured peaks and this value was normalized against that of the loading control in the same sample. Loading controls for histone modifications were the total H3 levels, which even after stripping give a strong signal. Modified MSK1 and CREB were normalized first to the RNAPII loading control, which is not affected by the treatments in this study; the total/unmodified proteins in the same samples were assessed similarly in parallel due to their lower levels and weaker signals, after which these relative values were normalized.

Chromatin immunoprecipitation (ChIP)
Cells were grown to ~70–80% confluence before treatment with GnRH and/or the inhibitors. Cross-linking was by addition of 1% formaldehyde in ice-cold PBS for 10 min at room temperature on a shaking platform and terminated by 0.125 M glycine for 5 min at room temperature. Cells were washed twice and harvested in ice-cold 1× PBS, and ChIP was carried out essentially as previously reported (41). The sonication to obtain fragments averaging 500–700 bp was carried out using 10 pulses of 33% amplitude for 15 s, with 10-s intervals between each pulse. Alternatively, to obtain DNA fragments of 100–150 bp (high resolution ChIP), which is able to distinguish between the nucleosomes, 60 pulses of 33% amplitude for 15 s, with 10-s intervals between each pulse were used (as demonstrated in Ref. 36). The immunoprecipitated DNA was collected and purified using the PCR purification kit (Qiagen) and quantified by qPCR as above using primers to the proximal promoters described in Ref. 3, centering at −60 to −160 bp, or as follows for Cga: forward +47, GTGGTCACAATTTATTACTCTTT; forward −429, GGAGCAATTTTTTTTTCTTG; forward −525: CACACCTGGACATATCTACTGT; reverse +147, ATCACCTGCCAGAAACA; R −139: TTAGCATATACACATCAACCCA; reverse −329, ATTAGCTAAGTACCTGATATTTCA; reverse −425, CAATTGCTCCAATTTCCTTAAATTAC.

Antibodies
Antibodies used were: anti-H3 (Abcam; ab1791), anti-H3K27ac (Cell Signaling; number 4729), anti-H3K18ac (Abcam; ab9675), anti-H3S10ph (Abcam; ab14955; all with various lot numbers over several years during which experiments...
Acknowledgment—We acknowledge the gift from Pamela Mellon.

For Western blot analysis, antibodies for total H3 and histone modifications were diluted 1:5000, except for that against H3S28p, which was diluted 1:2000. The antibodies for MSK1 and phospho-MSK1 were diluted 1:1000 and 1:1500, respectively, for CREB and phospho-CREB at 1:2000, RNAPII at 1:5000, and IgG was diluted 1:10,000. For ChIP, they were diluted 1:280, except for the antibodies against H3 and H3S28p, which were used at 1:350 and 1:140 dilutions, respectively.

Statistical analysis

All experiments were repeated on at least two occasions, and representative individual experiments or compiled results are presented. One-way analysis of variance, followed by Bonferroni’s t test or Student’s t test were employed to determine statistically different means. Differences were considered significant when $p < 0.05$.

Author contributions—M. H. conception, design, and performing the research, analysis of results, writing of draft manuscript. A. W. conception, design, and performing the research and analysis of results. S. R. conception and experimental design. J. T. supplied reagents that he validated. L. P. contributed to experimental design, performed the research, and analysis of results. P. M. conceived the project, oversaw the work, analysis, and interpretation of data, wrote the final version of the paper.

Acknowledgment—We acknowledge the gift from Pamela Mellon (University of California, San Diego) of the gonadotrope cell lines.

References

1. Lim, S., Luo, M., Koh, M., Yang, M., bin Abdul Kadir, M. N., Tan, J. H., Ye, Z., Wang, W., and Melamed, P. (2007) Distinct mechanisms involving diverse histone deacetylases repress expression of the two gonadotropin $\beta$-subunit genes in immature gonadotropes, and their actions are overcome by gonadotropin-releasing hormone. Mol. Cell. Biol. 27, 4105–4120

2. Melamed, P. (2008) Histone deacetylases and repression of the gonadotropin genes. Trends Endocrinol. Metab. 19, 25–31

3. Wijeweera, A., Hajj, M., Feldman, A., Pnueli, L., Luo, Z., and Melamed, P. (2015) Gonadotropin gene transcription is activated by medin-mediated effects on the chromatin. Biochim. Biophys. Acta 1849, 328–341

4. Melamed, P., Kadir, M. N., Wijeweera, A., and Seash, S. (2006) Transcription of gonadotropin $\beta$ subunit genes involves cross-talk between the transcription factors and co- regulators that mediate actions of the regulatory hormones. Mol. Cell. Endocrinol. 252, 167–183

5. Mouillot, J. F., Sonnenberg-Hirche, C., Yan, X., and Sadovsky, Y. (2004) p300 regulates the synergy of steroidogenic factor-1 and early growth response-1 in activating luteinizing hormone-$\beta$ subunit gene. J. Biol. Chem. 279, 7832–7839

6. Miller, R. S., Wolfe, A., He, L., Radovick, S., and Wondisford, F. E. (2012) CREB binding protein (CBP) activation is required for luteinizing hormone-$\beta$ expression and normal fertility in mice. Mol. Cell. Biol. 32, 2349–2358

7. Bannister, A. J., and Kouzarides, T. (2011) Regulation of chromatin by histone modifications. Cell Res. 21, 381–395

8. Baek, S. H. (2011) When signaling kinases meet histones and histone modifiers in the nucleus. Mol. Cell. 42, 274–284

9. Banerjee, T., and Chakravarti, D. (2011) A peek into the complex realm of histone phosphorylation. Mol. Cell. Biol. 31, 4858–4873

10. Dunn, K. L., and Davie, J. R. (2005) Stimulation of the Ras-MAPK pathway leads to independent phosphorylation of histone H3 on serine 10 and 28. Oncogene 24, 3492–3502

11. Rossetto, D., Arvakumov, N., and Côté, J. (2012) Histone phosphorylation: a chromatin modification involved in diverse nuclear events. Epigenetics 7, 1108

12. Sawicka, A., and Seiser, C. (2012) Histone H3 phosphorylation: a versatile chromatin modification for different occasions. Biochemie 94, 2193–2201

13. Sawicka, A., and Seiser, C. (2014) Sensing core histone phosphorylation: a matter of perfect timing. Biochim. Biophys. Acta 1839, 711–718

14. Walter, W., Clynes, D., Tang, Y., Marmorstien, R., Mellor, I., and Berger, S. L. (2008) 14–3–3 interaction with histone H3 involves a dual modification pattern of phosphorycetyl. Mol. Cell. Biol. 28, 2840–2849

15. Winter, S., Simboeck, E., Fischle, W., Zupkovitz, G., Dohnal, I., Mechtler, K., Ammerer, G., and Seiser, C. (2008) 14–3–3 proteins recognize a histone code at histone H3 and are required for transcriptional activation. EMBO J. 27, 88–99

16. Zippo, A., Serafini, R., Rocchigiani, M., Pennacchini, S., Krepleva, A., and Oliviero, S. (2009) Histone crosstalk between H3S10ph and H4K16ac generates a histone code that mediates transcription elongation. Cell. 138, 1122–1136

17. Drobic, B., Pérez-Cadahía, B., Yu, J., Kung, S. K., and Davie, J. R. (2010) Promoter chromatin remodeling of immediate–early genes is mediated through histone phosphorylation at either serine 28 or 10 by the MSK1 multi-protein complex. Nucleic Acids Res. 38, 3196–3208

18. Simboeck, E., Sawicka, A., Zupkovitz, G., Senese, S., Winter, S., Dequidt, F., Ogris, E., Di Croce, L., Chiocca, S., and Seiser, C. (2010) A phosphorylation switch regulates the transcriptional activation of cell cycle regulator p21 by histone deacetylase inhibitors. J. Biol. Chem. 285, 41062–41073

19. Karam, C. S., Kellner, W. A., Takenaka, N., Clemmons, A. W., and Corces, V. G. (2010) 14-3-3 mediates histone cross-talk during transcription elongation in Drosophila. PLoS Genet. 6, e1000975

20. Cheung, P., Tanner, K. G., Cheung, W. L., Sassone-Corsi, P., Denu, J. M., and Allis, C. D. (2000) Synergistic coupling of histone H3 phosphorylation and acetylation in response to epidermal growth factor stimulation. Mol. Cell. 5, 905–915

21. Clayton, A. L., Rose, S., Barratt, M. J., and Mahadevan, L. C. (2000) Phosphoacetylation of histone H3 on c-fos- and c-jun-associated nucleosomes upon gene activation. EMBO J. 19, 3714–3726

22. Lo, W. S., Trievel, R. C., Rojas, R. J., Duggan, L., Hsu, J. Y., Allis, C. D., Marmorstien, R., and Berger, S. L. (2000) Phosphorylation of serine 10 in histone H3 is functionally linked in vitro and in vivo to Gcn5-mediated acetylation at lysine 14. Mol. Cell. 5, 917–926

23. Klein, A. M., Zaganjor, E., and Cobb, M. H. (2013) Chromatin- tethered MAPKs. Curr. Opin. Cell Biol. 25, 272–277

24. Yang, S. H., Sharrocks, A. D., and Whitmarsh, A. J. (2013) MAP kinase signalling cascades and transcriptional regulation. Gene 513, 1–13

25. Josephowicz, S. Z., Shimada, M., Armache, A., Li, C. H., Miller, R. M., Lin, S., Yang, A., Dill, B. D., Molina, H., Park, H. S., Garcia, B. A., Taunton, J., Roeder, R. G., and Allis, C. D. (2016) Chromatin kinases act on transcription factors and histone tails in regulation of inducible transcription. Mol. Cell. 64, 347–361

26. Solbaga, A., Thomson, S., Wiggins, G. R., Rampersaud, N., Dyson, M. H., Hazzalin, C. A., Mahadevan, L. C., and Arthur, J. S. (2003) MSK2 and MSK1 mediate the mitogen- and stress-induced phosphorylation of histone H3 and HMG-14. EMBO J. 22, 2788–2797

GnRH targets H3S28p via MSK to activate Cga transcription
GnRH targets H3S28p via MSK to activate Cga transcription

27. Vermeulen, L., Vanden Berghe, W., Beck, I. M., De Bosscher, K., and Haegeman, G. (2009) The versatile role of MSKs in transcriptional regulation. Trends Biochem. Sci. 34, 311–318
28. Reyskens, K. M., and Arthur, J. S. (2016) Emerging roles of the mitogen and stress-activated kinases MSK1 and MSK2. Front. Cell Dev. Biol. 4, 56
29. Tiwari, V. K., Studier, M. B., Wirbelauer, C., Paro, R., Schübeler, D., and Beisel, C. (2011) A chromatin-modifying function of JNK during stem cell differentiation. Nat. Genet. 44, 94–100
30. Lim, S., Pnueli, L., Tan, J. H., Naor, Z., Rajagopal, G., and Melamed, P. (2009) Negative feedback governs gonadotrope frequency-decoding of gonadotropin releasing hormone pulse-frequency. Proc. Natl. Acad. Sci. U.S.A. 106, 10131–10136
31. Jorgensen, J. S., Quirk, C. C., and Nilson, J. H. (2004) Multiple and overlapping combinatorial codes orchestrate hormonal responsiveness and dictate cell-specific expression of the genes encoding luteinizing hormone. Endocr. Rev. 25, 521–542
32. Hodges, C., Bintu, L., Lubkowska, L., Kashlev, M., and Bustamante, C. (2005) Transcript profiling of immediate early genes reveals a unique role for activating transcription factor 3 in mediating activation of the glycoprotein hormone alpha-subunit promoter by gonadotropin-releasing hormone. Mol. Endocrinol. 19, 2624–2638
33. Sawicka, A., Hartl, D., Goiser, M., Pusch, O., Stocsits, R. R., Tamir, I. M., Mechtler, K., and Seiser, C. (2014) H3S28 phosphorylation is a hallmark of the transcriptional response to cellular stress. Genome Res. 24, 1808–1820
34. Rudnizky, S., Bavly, A., Malik, O., Pnueli, L., Melamed, P., and Kaplan, A. (2016) H2A.Z controls the stability and mobility of nucleosomes to regulate expression of the LH genes. Nat. Commun. 7, 12958
35. Lau, P. N., and Cheung, P. (2011) Histone code pathway involving H3 S28 phosphorylation and K27 acetylation activates transcription and antagonizes polycomb silencing. Proc. Natl. Acad. Sci. U.S.A. 108, 2801–2806
36. Dyson, M. H., Thomson, S., Inagaki, M., Goto, H., Arthur, S. J., Nightingale, K., Iborra, F. J., and Mahadevan, L. C. (2005) MAP kinase-mediated phosphorylation of distinct pools of histone H3 at S10 or S28 via mitogen- and stress-activated kinase 1/2. J. Cell Sci. 118, 2247–2259
37. Zhong, S., Jansen, C., She, Q. B., Goto, H., Inagaki, M., Bode, A. M., Ma, W. Y., and Dong, Z. (2001) Ultraviolet B-induced phosphorylation of histone H3 at serine 28 is mediated by MSK1. J. Biol. Chem. 276, 33213–33219
38. Wiersma, M., Bussiere, M., Halsall, J. A., Turan, N., Slany, R., Turner, B. M., and Nightingale, K. P. (2016) Protein kinase MSK1 physically and functionally interacts with the KMT2A/MLL1 methyltransferase complex and contributes to the regulation of multiple target genes. Epigenetics Chromatin 9, 52
39. Janknecht, R. (2003) Regulation of the ER81 transcription factor and its coactivators by mitogen- and stress-activated protein kinase 1 (MSK1). Oncogene 22, 746–755
40. Gehani, S. S., Agrawal-Singh, S., Dietrich, N., Christophersen, N. S., Helin, K., and Hansen, K. (2010) Polycomb group protein displacement and gene activation through MSK-dependent H3K27me3 S28 phosphorylation. Mol. Cell. Biol. 30, 886–900
41. Hodges, C., Bintu, L., Lubkowska, L., Kashlev, M., and Bustamante, C. (2009) Nucleosomal fluctuations govern the transcription dynamics of RNA polymerase II. Science 325, 626–628
42. Kireeva, M. L., Hancock, B., Cremona, G. H., Walter, W., Studitsky, V. M., and Kashlev, M. (2005) Nature of the nucleosomal barrier to RNA polymerase II. Mol. Cell. 18, 97–108
43. Yosefzon, Y., David, C., Tsukerman, A., Pnueli, L., Qiao, S., Boehm, U., and Melamed, P. (2015) RNA transcription Elongation factors and stress activated kinases MSK1 and MSK2. Front. Cell Dev. Biol. 5, 71
44. Petesch, S. J., and Lis, J. T. (2012) Overcoming the nucleosome barrier to RNA polymerase II. J. Biol. Chem. 287, 20720–20731
45. Feng, J., Lawson, M. A., and Melamed, P. (2008) A proteomic comparison of immature and mature mouse gonadotrophs reveals novel differentially expressed nuclear proteins that regulate gonadotropin gene transcription and RNA splicing. Biol. Reprod. 79, 546–561
46. Pnueli, L., Rudnizky, S., Yosefzon, Y., and Melamed, P. (2015) RNA transcription from a distal enhancer is required for activating the chromatin at the promoter of the gonadotropin α-subunit gene. Proc. Natl. Acad. Sci. U.S.A. 112, 4369–4374