Mitogenic Signals Stimulate the CREB Coactivator CRTC3 through PP2A Recruitment

**HIGHLIGHTS**

The mammalian CRTC family is distinctly regulated by protein phosphatases

CRTC1 and CRTC2 bind to calcineurin, and PP2A is recruited to CRTC3

CRTC3-PP2A complex formation is mediated by phosphorylation at S391 of CRTC3

Mitogenic signals induce S391 phosphorylation via MAPKs and CDKs

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Mitogenic Signals Stimulate the CREB Coactivator CRTC3 through PP2A Recruitment

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SUMMARY

The second messenger 3',5'-cyclic adenosine monophosphate (cAMP) stimulates gene expression via the cAMP-regulated transcriptional coactivator (CRTC) family of cAMP response element-binding protein coactivators. In the basal state, CRTCs are phosphorylated by salt-inducible kinases (SIKs) and sequestered in the cytoplasm by 14-3-3 proteins. cAMP signaling inhibits the SIKs, leading to CRTC dephosphorylation and nuclear translocation. Here we show that although all CRTCs are regulated by SIKs, their interactions with Ser/Thr-specific protein phosphatases are distinct. CRTC1 and CRTC2 associate selectively with the calcium-dependent phosphatase calcineurin, whereas CRTC3 interacts with B55 PP2A holoenzymes via a conserved PP2A-binding region (amino acids 380–401). CRTC3-PP2A complex formation was induced by phosphorylation of CRTC3 at S391, facilitating the subsequent activation of CRTC3 by dephosphorylation at 14-3-3 binding sites. As stimulation of mitogenic pathways promoted S391 phosphorylation via the activation of ERKs and CDKs, our results demonstrate how a ubiquitous phosphatase enables cross talk between growth factor and cAMP signaling pathways at the level of a transcriptional coactivator.

INTRODUCTION

The second messenger 3',5'-cyclic adenosine monophosphate (cAMP) is a potent driver of cellular responses to hormonal and environmental cues (Montminy, 1997). cAMP mediates the activation of the protein kinase A (PKA) holoenzyme (Taylor et al., 2012), which subsequently phosphorylates cellular substrates (Shabb, 2001). The transcriptional response to cAMP proceeds via the PKA-mediated phosphorylation of cAMP response element (CRE)-binding protein (CREB) family members (CREB1, ATF1, and CREM) (Mayr and Montminy, 2001). Phosphorylation of CREB at S133 induces a conformational change that enables the association of CREB with the histone acetyltransferases CREB-binding protein (CBP) and p300 (Parker et al., 1996). In parallel, cAMP also stimulates the association of CREB with the family of cAMP-regulated transcriptional coactivators (CRTC, CRTC1–3) over relevant genomic CREB-binding sites (CRE sites) (Altarejos and Montminy, 2011).

Under basal conditions, AMP-activated protein kinase family members, most notably the salt-inducible kinases (SIKs, SIK1–3), sequester the three CRTCs in the cytoplasm by phosphorylation at conserved sites, which promotes phosphorylation-dependent 14-3-3 protein binding (Sonntag et al., 2017; Wein et al., 2018). cAMP stimulation triggers the PKA-mediated phosphorylation and inhibition of SIK1–3 (Sonntag et al., 2018), leading to the dephosphorylation of CRTCs, which shuttle to the nucleus and drive cAMP/CREB target gene expression (Wein et al., 2018).

In addition to their regulation by SIKs, CRTCs are also controlled by calcium signaling via the Ca²⁺/calmodulin-dependent protein phosphatase calcineurin (CaN) (Perrino et al., 1995), which binds to and dephosphorylates the CRTCs at 14-3-3 binding sites (Altarejos and Montminy, 2011). The most abundant cellular phosphatases protein phosphatase 1 (PP1) and PP2A also appear to stimulate the dephosphorylation of CRTC2 (Shi, 2009; Uebi et al., 2010), although the context in which these ubiquitous enzymes regulate CRTC activities is unclear.

Multiple CRTC family members are co-expressed in mammalian tissues: CRTC1 and CRTC2 are produced in the brain (Altarejos et al., 2008; Jeanneteau et al., 2012) and pancreatic islets (Eberhard et al., 2013; Malm Clayton Foundation Laboratories for Peptide Biology, The Salk Institute for Biological Studies, La Jolla, CA 92037, USA
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et al., 2016), for example, whereas CRTC2 and CRTC3 are found in fat (Henriksson et al., 2015; Song et al., 2010), liver (Koo et al., 2005; Patel et al., 2014), and bone marrow-derived immune cells (Kim et al., 2017; MacKenzie et al., 2013). Nevertheless, individual CRTCs appear to execute dominant roles in certain tissues, with CRTC1 controlling memory (Nonaka et al., 2014) and behavior (Breuillaud et al., 2012), entraining the circadian clock (Jagannath et al., 2013), and promoting energy expenditure (Altarejos et al., 2008). By contrast, CRTC2 regulates liver gluconeogenesis (Koo et al., 2005) and beta-cell viability (Blanchet et al., 2015; Eberhard et al., 2013), whereas CRTC3 promotes adipose tissue (Song et al., 2010) and anti-inflammatory macrophage function (MacKenzie et al., 2013). However, it is unknown whether these differences reflect the relative abundance or distinct activation properties of CRTC family members in relevant tissues.

Based on the importance of CRTC dephosphorylation for transcriptional activation, we systematically examined the relative association of individual CRTC family members across the mammalian complement of Ser/Thr-specific protein phosphatases by mass spectrometry. We identified selective interactions of CRTC1/2 with CaN and uncovered an unexpected role for PP2A in binding to and activating CRTC3 in response to extracellular signals. These results point to a novel role for PP2A in mediating effects of growth factor cues on the selective activation of a distinct CRTC family member.

RESULTS

Selective Binding of Ser/Thr-Specific Protein Phosphatases to CRTC Family Members

Although they are all regulated by cAMP and the SIKs, individual CRTC family members can perform unique roles (Figures 1A and 1B). We hypothesized that these distinct phenotypes reflect in part differences in protein-protein interactions, rather than differences in CRTC protein abundance. In previous immunoprecipitation (IP)-mass spectrometry (MS) studies, we identified two conserved SIK/14-3-3-binding sites that inhibit CRTC activity (Sonntag et al., 2017) (S171 and S275 in CRTC2; Figure 1C). Using the corresponding HEK 293T cell dataset to identify Ser/Thr phosphatases that interact with overexpressed CRTC1–3 (Figures 2A and 2B), we detected the regulatory and catalytic subunits of CaN in IPs of CRTC1 and CRTC2, but not CRTC3 (Figures 2C and 2D). Conversely, CRTC3 but not CRTC1/2 associated very strongly with PP2A holoenzymes; spectral counts for catalytic (PPP2CA/B), scaffold (PPP2R1A/B), and 55-kDa regulatory B (B55, PPP2R2A–D) subunits were more than 10-fold enriched in IPs of CRTC3 relative to CRTC1/2. Other phosphatases (PP1, PP4, and PP6) appeared to associate comparably, albeit weakly with all CRTC family members (Figure S1).
We confirmed the mutually exclusive interactions of CRTC2 with CaN and CRTC3 with PP2A by co-immunoprecipitation (co-IP) (Figure 2F). Exposure to the adenylyl cyclase activator forskolin (Fsk) triggered the dephosphorylation of both CRTC2 and CRTC3 at SIK/14-3-3 sites (CRTC2/3: S171/S162 and S275/S273), thereby decreasing their association with 14-3-3 proteins. Co-treatment with the CaN inhibitor cyclosporin (CsA) (Jin and Harrison, 2002) selectively impaired the Fsk-mediated dephosphorylation of CRTC2 at S171, leading to increases in 14-3-3 binding. Fsk stimulation decreased the association of CRTC2 with CaN, but it had no effect on the association of CRTC3 with PP2A. We considered that CRTC3 phosphorylation at 14-3-3 binding sites may modulate PP2A interaction (Figure 2G). Although it eliminated 14-3-3 binding, mutation of all five SIK/14-3-3 sites (Sonntag et al., 2017) in CRTC3 had no effect on its association with PP2A. Collectively, these results indicate that the interaction of CRTC3 with the B55 PP2A holoenzyme is independent of SIK-mediated phosphorylation.
Identification of a Conserved PP2A-Binding Region in CRTC3

To characterize the PP2A-binding region (PBR) in CRTC3, we generated serial N- and C-terminal truncations of CRTC3 and analyzed these for PP2A association by co-IP (Figure S2). This analysis uncovered a 20-amino acid core PBR (amino acids 380–401) located within the central regulatory domain of CRTC3 (Figures 3A and 3B). Pointing to an important function for this region, the PBR amino acid sequence is identical between humans and mice (Figure 3C).

**Figure 3. Characterization of the PP2A-Binding Region (PBR) in CRTC3**

(A) Western blot analysis showing recovery of endogenous PP2A from IPs of FLAG-tagged CRTC3 mutants. The only fragment that abolished PP2A binding is highlighted in red (201–383).
(B) To-scale representation of CRTC3 protein fragments assayed for interaction with PP2A. In orange the PBR of CRTC3 (Mus musculus amino acids 380–401).
(C) CRTC3 sequence alignment of the PBR from multiple vertebrate species (CRTC3 sequences were obtained via UniProt; The UniProt Consortium, 2017).
(D) Sequence alignment of the PBR from vertebrate CRTC3 proteins compared with Mus musculus CRTC2 425–444 (in purple).
(E and F) (E) To-scale representation of the CRTC2 (light green) and CRTC3 (dark green) hybrid proteins assayed for PP2A binding. Hybrid proteins: CRTC2 H1 [mCRTC2(328–421) replaced with mCRTC3(326–379), in blue], CRTC2 H2 [mCRTC2(328–449) replaced with mCRTC3(326–402)], and CRTC3 H1 [mCRTC3(322–407) replaced with mCRTC2(324–454)]. (F) Western blot showing recovery of endogenous PP2A from IPs of FLAG-tagged CRTC2/3 mutants.

Identification of a Conserved PP2A-Binding Region in CRTC3

To characterize the PP2A-binding region (PBR) in CRTC3, we generated serial N- and C-terminal truncations of CRTC3 and analyzed these for PP2A association by co-IP (Figure S2). This analysis uncovered a 20-amino acid core PBR (amino acids 380–401) located within the central regulatory domain of CRTC3 (Figures 3A and 3B). Pointing to an important function for this region, the PBR amino acid sequence is identical between humans and mice (Figure 3C).
Although CRTC2 does not bind PP2A, it contains a PBR-related sequence at 425–444 (Figure 3D). We generated CRTC2/3 hybrids to test whether sequences flanking this core region modulate the PP2A association (Figures 3E and 3F). Similar to the PP2A interaction patterns of truncated CRTC3 protein fragments, insertion of mCRTC3 326–402 promoted binding of CRTC2 to PP2A (CRTC2 H2), whereas insertion of the corresponding region of CRTC2 (mCRTC2 324–454) into CRTC3 (CRTC3 H1) abolished PP2A binding. However, insertion of CRTC3 sequences flanking the PBR into CRTC2 (mCRTC3 326–379; CRTC2 H1) increased PP2A binding to a level that was intermediate between CRTC2 and CRTC3. These findings indicate that although mCRTC3 380–401 is absolutely required for the strong interaction with PP2A, regions outside the PBR also contribute to complex formation.

Phosphorylation of the PBR Increases PP2A Binding
The CRTC3 PBR contains three conserved Ser/Thr residues that appear to undergo phosphorylation (Hornbeck et al., 2015) (Figure 4A). We confirmed the phosphorylation of CRTC3 at S391 and S396 by MS analysis (Figure S3). To assess the potential effects of these modifications within the PBR on the CRTC3-PP2A interaction, we performed mutagenesis studies (Figure 4B). Although alanine substitutions at T394 and S396 had no effect on PP2A binding, mutation of either S391 or the flanking P392 to alanine abolished this interaction. To monitor phosphorylation at the critical S391 site, we generated a phospho-specific antiserum (P-CRTC3 S391). Western blot analysis revealed robust S391 phosphorylation following overexpression of CRTC3 wild-type (WT) and S391A mutant proteins in HEK 293T cells.

To determine the importance of S391 phosphorylation for the CRTC3-PP2A interaction, we tested whether substitution of this phosphoacceptor with alanine or a phosphomimetic (Glu) alters complex formation...
In contrast to WT CRTC3, neither S391A nor S391E mutant CRTC3 protein associated detectably with PP2A. Correspondingly, phospho-S391-defective CRTC3 proteins were phosphorylated to a greater extent at 14-3-3 binding sites (P-CRTC3 S273), and as a result, the association of S391A/E mutants with 14-3-3 proteins was stronger compared with WT.

In a transient CRE luciferase-based assay for the quantification of cAMP-dependent gene expression (Conkright et al., 2003), the phosphorylation-defective S391A mutant CRTC3 exhibited ~2-fold lower basal transcriptional activity compared with WT CRTC3 (Figure 4D). WT and S391A CRTC3 CRE activities were more similar following exposure to Fsk, however, suggesting that S391 modulates CRTC3 activity in a cAMP-independent manner. We performed immunofluorescence studies to determine the subcellular localization of the S391A mutant (Figures 4E and 4F). In contrast to the predominantly nuclear WT CRTC3, S391A mutant protein was primarily localized in the cytoplasm under basal conditions. In keeping with its effects on CRE reporter activity, Fsk promoted the nuclear translocation of CRTC3 WT and S391A comparably (Figure S4).

PP2A has multifaceted roles in the cell cycle and is thought to function as a tumor suppressor (Haesen et al., 2014; Wlodarchak and Xing, 2016). Various cellular proteins can inhibit PP2A activity, including the phosphorylation-dependent regulators of mitosis ENSA, ARPP19, and Bod1 (Gharbi-Ayachi et al., 2010; Porter et al., 2013). In contrast to CRTC3 (Song et al., 2010), these proteins are small (12–20 kDa) (The UniProt Consortium, 2017) and knockdown or overexpression of any one regulator typically leads to growth arrest and embryonic lethality (Dupre et al., 2014; Matthews and Evans, 2014). Arguing against a role for CRTC3 as a PP2A inhibitor, cellular PP2A activity and proliferation appeared comparable between cells expressing WT or phosphorylation-defective (S391A) CRTC3 in HEK 293T cells (Figure S5). These results support the notion that S391 phosphorylation selectively increases CRTC3 activity by promoting its association with the PP2A holoenzyme, which in turn dephosphorylates CRTC3 at remote SIK/14-3-3 sites, leading to CRTC3 nuclear translocation and CREB target gene activation.

**MAPKs and CDKs Promote PBR Phosphorylation at S391**

Realizing that S391 phosphorylation of CRTC3 increases its interaction with PP2A, we set out to identify relevant protein kinases that mediate the phosphorylation of this site. Based on the importance of a flanking proline at the +1 position relative to S391, we addressed the potential involvement of proline-directed kinases in this process by pharmacological inhibition (Figure 5A). Exposure to the cyclin-dependent kinase (CDK1/2/5) inhibitor roscovitine (IC50: CDK1/2/5 = 0.2–0.8 µM, ERK1/2 = 15–30 µM) (Meijer et al., 1997) and the ERK1/2 inhibitor SCH772984 (IC50: ERK1/2 = 0.004/0.001 µM) (Morris et al., 2013) reduced both CRTC3 S391 phosphorylation and PP2A binding following CRTC3 overexpression in HEK 293T cells. CDKs (Holst et al., 2009) and mitogen-activated protein kinases (MAPKs) such as ERK1/2 (Carlson et al., 2011) are proline-directed Ser/Thr kinases that appear capable of phosphorylating CRTC3 at S391. The CK1α/δ inhibitor longdaysin (IC50: CK1α/δ = 6–9 µM, ERK2 = 52 µM) (Hirota et al., 2010) also inhibited S391 phosphorylation, with the PBR (S391PLT394) containing a casein kinase 1 (CK1) consensus motif pS/Txx(x)T (underlined = CK1 phosphorylated residue) (Knippschild et al., 2014). In contrast to its effects on WT CRTC3, longdaysin did not impair PP2A binding in the context of the T394A mutant (Figure S6), indicating that S391 and T394 phosphorylation are required for full binding of PP2A in cells overexpressing WT CRTC3.

To evaluate the potential role of CDKs in modulating endogenous CRTC3 S391 phosphorylation, we arrested HEK 293T cells in the G2/M phase with nocodazole (Figure 5B). Although undetectable in the basal state, phosphorylations of CRTC3 at S391 and nucleophosmin (NPM) at T199, a known CDK target (Tokuyama et al., 2001), were both upregulated following nocodazole arrest. Consistent with these effects, short-term CDK inhibition with roscovitine and dinaciclib (IC50: CDK1/2/5 = 0.001–0.003 µM) (Parry et al., 2010) blocked the phosphorylation of both proteins.

To test the effect of ERK and cAMP signaling on endogenous S391 phosphorylation of CRTC3, we stimulated HEK 293T cells with either 12-O-tetradecanoylphorbol-13-acetate (TPA) (Schonwasser et al., 1998) or Fsk (Figure 5C). In contrast to Fsk, which promoted the dephosphorylation of CRTC2 and CRTC3 at 14-3-3 sites (P-CRTC3 S273: top CRTC2 and bottom CRTC3), exposure to TPA upregulated ERK activity (P-ERK T202 Y204) and induced CRTC3 phosphorylation at S391. These effects were suppressed by pretreatment with the ERK1/2 inhibitor SCH772984.
Following exposure to TPA or nocodazole, we detected two immunoreactive bands using the S391 phospho-specific antiserum. Both bands correspond to endogenous CRTC3, as revealed in knockdown studies using CRTC3-specific short hairpin RNA (shRNAs) (Figure S7). In previous studies with HEK 293T cells, we found that TPA does not activate the predominant CRTC family member CRTC2 (Ravnskjaer et al., 2007). By contrast, exposure to TPA decreased 14-3-3 protein binding to CRTC3, thereby enhancing its nuclear localization (Figures 5D and 5E). Taken together, these results indicate that MAPKs and CDKs both regulate the S391 phosphorylation and activation of endogenous CRTC3.

PBR Phosphorylation Increases CRTC3 Activity
Based on the importance of CRTC3 for adipose function (Song et al., 2010; Yoon et al., 2018), we examined CRTC3 S391 phosphorylation in brown-adipose-tissue-derived stromal vascular fraction (bSVF) cells, which are enriched in preadipocytes (Han et al., 2015) (Figure 6A). In the basal state, S391-phosphorylated CRTC3...
was detected in WT, but not in CRTC3 knockout (KO) bSVFs. Having seen their effects on CRTC3 S391 phosphorylation in HEK 293T cells, we tested CDK and ERK inhibitors in bSVFs (Figure 6B). Although both roscovitine and dinaciclib inhibited CDK activity as revealed by the corresponding reduction in NPM T199 phosphorylation, the ERK1/2 inhibitor SCH772984 was most effective in reducing CRTC3 S391 phosphorylation. We evaluated the importance of ERK signaling for CREB target gene expression by exposing bSVFs to SCH772984 (Figure 6C). SCH772984 treatment partially blocked the Fsk-mediated induction of canonical CREB target genes NR4A1 and regulator of G-protein signaling-2 (RGS2) (Ravnskjaer et al., 2007; Song et al., 2010). The inhibitory effects of SCH772984 appear to be CRTC3 dependent, because they were absent in CRTC3 KO bSVFs. Taken together, these results demonstrate that in bSVFs ERK activity functions cooperatively with cAMP/PKA signals to promote CREB target gene expression through its effects on CRTC3.

**DISCUSSION**

The second messenger cAMP promotes cellular gene expression via the PKA-mediated phosphorylation and inhibition of the SIKs, leading to CRTC nuclear translocation upon dephosphorylation and release from 14-3-3 proteins (Figure 7A). Here we show that the association of CRTC members with different protein phosphatases provides distinct stimulus-dependent mechanisms for the induction of CREB target genes (Figures 7B and 7C).

The calcium-responsive phosphatase CaN interacts with substrates that contain PxIxIT motifs (Li et al., 2011), including vertebrate CRTC1 and CRTC2, but not CRTC3 (mCRTC1: P219G/NIF, P555NL/IL and mCRTC2: P249G/NIF, P614NL/IL). Although PxlIT motifs regulate the substrate affinity of CaN (Li et al., 2007; Roy et al., 2007), its phosphatase activity is dependent upon Ca\(^{2+}\)/calmodulin (Perrino et al., 1995). As a result, the Ca\(^{2+}\)/CaN-dependent regulation is limited to CRTC1 and CRTC2 (Altarejos et al., 2008; Kovacs et al., 2007; Screaton et al., 2004; Bittinger et al., 2004; Ch'ng et al., 2012).

By contrast, CRTC3 binds to the B55 PP2A holoenzyme following phosphorylation of CRTC3 at S391 by MAPK/CDKs. Within the PBR (mCRTC3 PBR: 380–401) the S391 phosphorylation site (PVS391P) forms part

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**Figure 6. ERK-Mediated Activation of CRTC3 in Undifferentiated bSVFs**

(A) Western blot analysis of primary brown adipose stromal vascular fraction (bSVF) cells generated from wild-type C57BL/6 (WT) and CRTC3 knockout mice (KO).

(B) Western blot analysis showing the effects of 4-hr treatment with roscovitine (20 μM), dinaciclib (200 nM), and SCH772984 (200 nM) on phospho-S391 CRTC3 levels in bSVF WT and KO cells.

(C) NR4A1 and RGS2 mRNA levels of bSVF CRTC3 WT and KO cells following (co-)treatment with SCH772984 (SCH, 200 nM, 4 hr) and Fsk (10 μM, 1 hr). Values represent data from three independent experiments normalized to RPL32 and relative to CRTC3 WT DMSO. (n = 3, ± SD; ns, not statistically significant, **p < 0.01)
of an optimal ERK1/2 motif (Px[S/T]P) (Carlson et al., 2011). The PBR also contains an MAPK docking motif (D domain), with characteristic basic and alternating hydrophobic residues (RRRQPPV S391PL TL) (Remenyi et al., 2005). Stimuli that upregulate ERK activity (epidermal growth factor and angiotensin II), also promote the phosphorylation of CRTC3 at S391/S396 (Olsen et al., 2006; Christensen et al., 2010). Superimposed on effects of ERKs, CDKs also appear to phosphorylate CRTC3 at S391 in growth-arrested cells (Olsen et al., 2010; Franz-Wachtel et al., 2012). Indeed, CRTC3 was also identified in a study of mitotic phosphorylation substrates within the chromosomal passenger complex (Yang et al., 2007), which controls cell division together with the CDKs (Carmena et al., 2012). Consistent with the importance of PP2A-mediated activation of CRTC3, in primary calvarial osteoblasts the parathyroid-hormone-induced nuclear translocation of CRTCs was blocked by the PP2A/PP1 inhibitor okadaic acid in case of CRTC3, but not for CRTC2 (Ricarte et al., 2018).

Although the regulatory subunits of PP2A appear important in distinguishing between different substrates, relevant substrate recognition motifs for the PP2A holoenzyme have not been reported until recently (Hae-sen et al., 2014; Wlodarchak and Xing, 2016). The B55 PP2A holoenzyme has been found to recognize substrates containing a CDK1 phosphorylation site flanked on both sides by a polybasic region (Cundell et al., 2016); a short LxxIxE motif appears to mediate interaction with the B56 PP2A holoenzyme (Hertz et al., 2016). The PBR of CRTC3 includes an N-terminal Arg triplet but lacks any basic C-terminal sequences (mCRTC3: RRRQPPVS391PLTLSPGPE). Only B55 PP2A subunits were recovered from IPs of all CRTCs, which contain a common sequence inside their PBR-related region consisting of two hydrophobic residues flanking an SP motif (mCRTC1/2/3: PL S332PI TQ/PL S434PL S434/PL S391PL TL). Although it lacks an essential glutamate at position 6, this common PBR-like motif in CRT family members bears more similarity with the LxxIxE motif described for the B56 PP2A holoenzyme. Similar to our findings, phosphorylation of residues within the LxxIxE motif also increases B56 PP2A affinity, leading to the dephosphorylation of distal residues in relevant substrates such as Repo-Man (LS PIPE), BubR1 (LSPILE), and RacGAP1 (LSTIDE; underlined = phosphorylated residues) (Hertz et al., 2016; Qian et al., 2013; Kruse et al., 2013). Future biochemical studies should reveal the extent to which the PBR we identified represents a distinct binding motif for the B55 PP2A holoenzyme.

The second messenger cAMP is traditionally thought to inhibit cell division, but in a subset of tissues, cAMP appears to cooperate with MAPK/ERK signals in promoting proliferation and differentiation (Dumaz and Marais, 2005). This cross talk is noteworthy in adipose tissue wherein β-adrenergic signaling triggers the activation of both cAMP and MAPK signaling pathways (Collins, 2011), often in combination with growth factors such as insulin/insulin growth factor-1 (Kajimura et al., 2015; Tang and Lane, 2012). As loss of CRTC3 expression increases brown adipocyte proliferation and differentiation and protects against high-fat-diet-induced adipose expansion (Song et al., 2010; Yoon et al., 2018), future studies should
address the specific roles of S391 phosphorylation in this context. This cooperative effect of cAMP and MAPK signals on CRTC3 activity has also been reported in regulatory macrophages where lipopolysaccharides (LPS) and prostaglandin E2 (PGE2) promote expression of the anti-inflammatory interleukin 10 (Clark et al., 2012; Mackenzie et al., 2013). Based on the ability of PGE2 to stimulate the cAMP pathway and LPS to activate multiple MAPK family members (Symons et al., 2006), our results suggest that these CRTC3-specific effects proceed via the recruitment of B55 PP2A.

Limitations of the Study
We used HEK 293T cells to characterize CRTC3-PP2A complex formation and its regulation by MAPK- and CDK-mediated phosphorylation. Although this approach was valuable in characterizing these molecular components and their downstream effects on CRTC3 activation, the relevant extracellular signals that act upstream to trigger the interaction of CRTC3 with PP2A remain relatively unknown. Our studies with brown pre-adipocytes support a potential role for ERK in mediating effects of β-adrenergic signals on recruitment of PP2A and on the subsequent activation of CRTC3. Future studies with phospho (S391)-specific antiserum should reveal the physiological contexts in which this site is phosphorylated, and mutant mice expressing phospho-S391 defective CRTC3 should provide insight into the physiological importance of this modification.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Transparent Methods and seven figures and can be found with this article online at https://doi.org/10.1016/j.isci.2018.12.012.

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AUTHOR CONTRIBUTIONS
T.S. conceived the study, designed and performed experiments, analyzed data, and wrote the manuscript. J.O. performed SVF mRNA experiments, analyzed data, provided reagents (CRTC3 shRNA plasmids), and edited the manuscript. J.M.V. generated antisera for P-CRTC3 S391 (PBL #7408), hCRTC3 (PBL #7019), and mCRTC2 (PBL #6,896). J.J.M. and J.R.Y. performed MS experiments and data analysis. Y.-S.Y. provided SVF cells. M.M. conceived the study and reviewed/editing the manuscript. All authors reviewed results and approved the final version of the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Mitogenic Signals Stimulate
the CREB Coactivator CRTC3
through PP2A Recruitment

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Supplemental Figures

Fig. S1

Tables show the IP-MS recovery of protein phosphatases (PP) 1, 4, and 6 subunits (comparing N-terminally tagged CRTC1-3; SPCs = spectral counts).

Fig. S1. Interaction of CRTC1-3 with protein phosphatases 1, 4, and 6. Related to Fig. 2.
Fig. S2. Characterization of the PP2A-binding region (PBR) in CRTC3. Related to Fig. 3.

Western blot analysis of Co-IPs of FLAG-tagged CRTC3 truncation mutants with endogenous PP2A and 14-3-3 proteins. Mutants that abolished PP2A interaction are highlighted in red (201-421 AAA = mCRTC3 201-421 S273A S329A S370A).
Fig. S3. Phosphorylation of S391 and S396 within the PP2A-binding region (PBR) of CRTC3. Related to Fig. 4.

A) Detection of MS spectra (387-407) derived from the PBR with corresponding peaks labeled (in B)). Phosphorylation was detected on amino acid s391 (Ascore = 14.89) and s396 (Ascore = 7.53; for Ascore see (Beausoleil et al., 2006)).
Fig. S4

**A** Immunofluorescence of HEK 293T cells transfected with FLAG-tagged CRTC3 WT and S391A. Cells were stained for FLAG epitope and counterstained with DAPI. (Fsk treatment for 30 min; scale bar indicates 20 µm)

**B** Graph shows relative subcellular localization of CRTC3 WT and S391A mutant.

**Fig. S4. Localization of CRTC3 WT and S391A mutant upon Forskolin treatment.** Related to Fig. 4.

A) Immunofluorescence of HEK 293T cells transfected with FLAG-tagged CRTC3 WT and S391A. Cells were stained for FLAG epitope and counterstained with DAPI. (Fsk treatment for 30 min; scale bar indicates 20 µm) B) Graph shows relative subcellular localization of CRTC3 WT and S391A mutant.
Fig. S5. Effect of CRTC3 overexpression on cellular PP2A activity and cell proliferation. Related to Fig. 4.

A-C) The activity of immunoprecipitated catalytic PP2A subunit (PPP2CA) was quantified by malachite green and KRptIRR phosphopeptide (PP2A Immunoprecipitation Phosphatase Assay Kit, Upstate). Prior to PP2A activity measurement FLAG-tagged GFP, CRTC3 and CRTC3 S391A S396A were overexpressed in HEK 293T cells for 48 h. A) Picture of the 96 well plate after the colorimetric PP2A assay. (A/B indicates independent transfections). B) Corresponding PP2A activity upon normalization to the standard curve. (n = 4, ± SD). C) Corresponding Western blot analysis of the cell lysate used for the PP2A activity assay. D) BrdU Cell Proliferation ELISA Kit (Abcam) measuring the effects of transient-overexpression of GFP, CRTC3, and CRTC3 S391A S396A S396E.
mutants on HEK 293T proliferation. BrdU treatment occurred either for 22 h (5 h post-transfection) or for 4 h (23 h post-transfection) prior to the colorimetric assay. ($n = 5, \pm$ SEM).
**Fig. S6**

Western blot analysis of the Co-IP of FLAG-tagged CRTC3 WT and T394A mutant with endogenous PP2A. 4 h prior to the IP HEK 293T cells were treated with Longdaysin (20 µM; IC₅₀ µM: CK1α/δ = 6-9, ERK2 = 52 (Hirota et al., 2010)) and SCH772984 (100 nM; IC₅₀ µM: ERK1/2 = 0.004/0.001 (Morris et al., 2013)).

**Fig. S6.** The effects of Longdaysin on the CRTC3-PP2A complex formation requires the intact CK1 motif inside the PBR. Related to Fig. 5.
**Fig. S7.** 

**P-CRTC3 S391 antibody evaluation.** Related to Fig. 5.

A) Western blot analysis showing the effects of transient CRTC3 knockdown and TPA treatment (200 nM, 1 h) on S391 phosphorylation of endogenous CRTC3. HEK 293T cells were transfected with three different shRNA constructs against *H. sapiens* CRTC3 (C3 no 1-3; neg. = shRNA negative control). After 48 h cells were treated for 1 h with TPA (200 nM),
Transparent Methods

Sections of the methods were previously described and are reprinted here, partly verbatim, for reference (Sonntag et al., 2017).

Small molecules

Small molecules were solubilized in DMSO (ACS, Sigma-Aldrich) at the indicated concentrations and stored until usage at -80 °C (long term storage) or -20 °C (working dilution): 2 mM Carfilzomib (PR-171; Selleck Chemicals), 200 µM Cyclosporin A (cyclosporin; Sigma-Aldrich), 100 µM Dinaciclib (SCH727965; Selleck Chemicals), 20 mM Forskolin (Sigma-Aldrich), 20 mM Longdaysin (Sigma-Aldrich), 200 µg/ml nocodazole (Cell Signaling Technology), 1 mM Palbociclib (PD-0332991; freshly prepared in ddH2O; Selleck Chemicals), 20 mM Roscovitine (Seliciclib, CYC202; Selleck Chemicals), 100 µM SCH772984 (Selleck Chemicals), 400 µM TPA (Cell Signaling Technology).

Antibodies

The antibodies used in this study were purchased from Abcam (PPP2R1A), Covance (GFP), EMD Millipore (α-tubulin), Santa Cruz Biotechnology (14-3-3 ε, 14-3-3 ζ), Sigma-Aldrich (FLAG M2), and Cell Signaling Technology (14-3-3 [pan], Calcineurin A [CaN; pan], P-Cdc2 Y15, P-CREB S133, CRTC3 H. sapiens AA139 [#2720], P-CRTC2 S171, P-NPM S4, P-NPM T199, P-p44/42 MAPK [Erk1/2] T202/Y204, PPP2CA, PPP2R1A, PPP2R2A). The P-CRTC3 S273 antiserum was previously described (Sonntag et al., 2017).

See antiserum production for the rabbit P-CRTC3 S391 (PBL #7408), the rabbit hCRTC3(414-432) (PBL #7019), and the rabbit mCRTC2(454-477) (PBL #6896) antisera.

Antiserum production

Animal Care

All animal procedures were approved by the Institutional Animal Care and Use Committee of the Salk Institute...
and were conducted in accordance with the PHS Policy on Humane Care and Use of Laboratory Animals (PHS Policy, 2015), the U.S. Government Principles for Utilization and Care of Vertebrate Animals Used in Testing, Research and Training, the NRC Guide for Care and Use of Laboratory Animals (8th edition) and the USDA Animal Welfare Act and Regulations. All animals were housed in an AAALAC accredited facility in a climate controlled environment (65-72 degrees Fahrenheit, 30-70% humidity) under 12-hour light/12-hour dark cycles. Upon arrival, animals were physically examined by veterinary staff for good health and acclimated for at least two weeks prior to initiation of antiserum production. Each animal was monitored daily by the veterinary staff for signs of complications and weighed every two weeks. Routine physical exams were also performed by the veterinarian quarterly on all rabbits.

For production of each antiserum (P-CRTC3 S391, mCRTC2(454-477) and hCRTC3(414-432)), three 10 to 12-week old, female New Zealand white rabbits, weighing 3.0 to 3.2 kg at beginning of the study, were procured from Irish Farms (I.F.P.S. Inc., Norco, California, USA). Rabbits were provided with ad libitum feed (5326 Lab Diet High Fiber), micro-filtered water and weekly fruits, vegetables and alfalfa hay for enrichment.

**Preparation of Antigens**

Synthetic peptides were synthesized in house or by RS Synthesis (Louisville, KY), HPLC purified to >95%, and amino acid sequenced verified by mass spectrometry. Peptides were covalently attached to large carrier proteins for use as immunogens. Cys^{383}.pSer^{391} mCRTC3(383-395)-NH_{2}, and Cys^{414} hCRTC3(414-432) were conjugated to maleimide activated Keyhole Limpet Hemocyanin (KLH) per manufacturer’s instructions (Thermo Fisher Scientific). mCRTC2(454-477) was conjugated to bovine thyroglobulins via glutaraldehyde, 1% final, using reagents purchased from Sigma-Aldrich.

Specific peptides used to generate antisera are as follows:

Cys^{383}.pSer^{391} mouse CRTC3(383-395)-NH_{2}, CRRRQQPV(pS)PLTL-NH_{2};

mouse CRTC2(454-477), KQFSPTMSPTLSSITQGVPLDTSK; and

Cys^{414} human CRTC3(414-432), CLAPYPTSQMVSSDRLS.
Injection and Bleeding of Animals

The antigen was delivered to host animals using multiple intradermal injections of peptide-KLH conjugate in Complete Freund's Adjuvant (initial inoculation) or incomplete Freund's adjuvant (booster inoculations) every three weeks. Rabbits were bled, <10% total blood volume, one week following booster injections and bleeds screened for titer and specificity. Rabbits were administered 1-2 mg/kg Acepromazine IM prior to injections of antigen or blood withdrawal. At the termination of study, rabbits were exsanguinated under anesthesia (ketamine 50 mg/kg and aceprozamine 1 mg/kg, IM) and euthanized with an overdose of pentobarbital sodium and phenytoin sodium (1 ml/4.5 kg of body weight IC to effect). After blood was collected death of animals was confirmed. All animal procedures were conducted by experienced veterinary technicians, under the supervision of Salk Institute veterinarians.

Characterization and purification of antisera

Each bleed from each animal was tested at multiple doses for the ability to recognize the synthetic peptide antigen; bleeds with highest titers were further analyzed by Western blot for the ability to recognize the full-length endogenous protein and to check for cross-reactivity with other proteins. Antisera with the best characteristics of titer against the synthetic peptide antigen, ability to recognize the endogenous protein, and specificity were affinity purified and used for all studies. Rabbit PBL #7408 anti P-CRTC3 S391, rabbit PBL #6896 anti-CRTC2, rabbit PBL #7019 anti-hCRTC3 were purified using Cys383.pSer391.mCRTC3(383-395)-NH$_2$, or mCRTC2(454-477), or Cys414.hCRTC3(414-432), respectively, covalently attached to Sulfolink agarose (Thermo Fisher Scientific) for cysteine containing CRTC3 peptides or to Affi-Gel 10 (BioRad) for the CRTC2 peptide with N-terminal lysine. Coupling of peptides to resins was per manufacturer’s instructions. To ensure that the same batch of purified antibodies could be used for this and future studies, large volumes, 20 ml rabbit sera, from bleeds with similar profiles were purified.

Plasmids
For overexpression studies plasmids were used that contained the *H. sapiens* Ubiquitin C promoter (pUbC), whose activity is unaffected by cAMP signaling. The plasmids - pUbC-3xFLAG-TEVsite-His6-MCS-IRESeGFP and pUbC-MCS-His6-TEVsite-3xFLAG-IRESeGFP - have been described previously (*MCS* = multiple cloning site) (Sonntag et al., 2017).

CRTC1-3 and EGFP (Clontech) overexpression constructs were previously generated (Sonntag et al., 2017, Sonntag et al., 2018). The plasmids code for the following proteins (UniProt identifier): mCRTC1 (Q68ED7-1), mCRTC2 (Q3U182-1), and mCRTC3 (Q91X84-1) (*m* = *Mus musculus*).

The CRTC3 mutants were cloned by restriction enzyme digest, site-directed mutagenesis, fusion PCR (CRTC3 Δ369-379), inverse PCR (CRTC3 Δ330-364), and restriction-free (RF) cloning (CRTC2/3 hybrid proteins). Fusion PCR and RF cloning were performed as previously described (Sonntag and Mootz, 2011, Sonntag, 2017).

The CRTC3 shRNA constructs were generated by cloning 5’ phosphorylated double stranded hCRTC3 oligonucleotides into the *Bam*HI and *Hind*III cut pSilencer 2.1-U6 puro plasmid (Applied Biosystems):

no. 1: 5’ GATCCGCTTCAGCAACTGCGCCTTTTCAAGAGAAAGGCGCAGTTGCTGAAGTTTTTTGGAAA

no. 2: 5’ GATCCGAAGCTCCTCTGGTCTCCATTCAAGAGATGGAGACCAGAGGAGCTTCTTTTTTGGAAA

no. 3: 5’ GATCCGCACATCAAGGTTTCAGCATTCAAGAGATGCTGAAACCTTGATGTGCTTTTTTGGAAA

The empty plasmid served as negative control.

**Cell culture**

HEK 293T cells were purchased from ATCC (CRL-11268) and propagated in DMEM (Gibco®, high glucose) supplemented with 10% Fetal Bovine Serum (FBS; Gemini Bio-Products) and 100 U/ml penicillin-streptomycin (Corning Inc.). Primary stromal vascular fraction (SVF) cells of brown adipose tissue (BAT) were generated from C57BL/6 wild type mice and CRTC3 knockout mice (Yoon et al., 2018):

Interscapular BAT was collected and digested for 30 min in collagenase buffer (100 mM HEPES pH 7.4, 1.5 mg/ml collagenase I [Sigma C6885], 125 mM NaCl, 5 mM KCl, 1.3 mM CaCl₂, 5 mM D-glucose, and 2% BSA) with gentle shaking at 37°C. After digestion and centrifugation, SVF cells were separated from floating mature adipocytes. SVF cells were filtered and incubated in RBC lysis buffer (0.017 M Tris pH 7.4; 0.16 M NH₄Cl, and
0.01 M EDTA) for 10 min at room temperature. SVF cells were centrifuged, washed, plated, and ultimately propagated in DMEM supplemented with 10% FBS and 100 U/ml penicillin-streptomycin.

**Overexpression & immunoprecipitation (IP)**

Experiments were performed in 6 well plates by reverse transfecting HEK 293T cells (2.5 × 10⁶ cells) with 2 µg plasmid DNA using Lipofectamine® 2000 (Invitrogen). 48 h post transfection cells were collected in PBS and resuspended in lysis buffer (50 mM Tris, 150 mM NaCl, 10% glycerol, 1% Igepal [Sigma-Aldrich], 1 mM DTT, EDTA-free cOmplete™ Protease Inhibitor Cocktail [Roche], Phosphatase Inhibitor Cocktail 2 and 3 [Sigma-Aldrich], 1 µM Carfilzomib; pH 8.0). The supernatant (= cell lysate) was either used in IP experiments or directly mixed with SDS-PAGE loading buffer. In all IP experiments, cells were pre-treated for 1 h with 1 µM Carfilzomib prior to cell lysis. Cell lysates were incubated with anti-FLAG® M2 magnetic beads and 3xFLAG peptide (100 µg/ml final) was used to elute bound proteins (both Sigma-Aldrich).

**Immunoprecipitation & mass spectrometry (IP-MS)**

The IP-MS protocol and data analysis of N- and C-terminally FLAG-tagged CRTC1-3 has been previously described (Sonntag et al., 2017).

**Immunoprecipitation of endogenous CRTC3**

Immunoprecipitation was performed from 4 x 100 mm dishes of HEK 293T cells (1.5 × 10⁷ cells). 48 h post seeding cells were treated for 2 h with 200 nM SCH772984 as well as 1 h with 1 µM Carfilzomib and 200 nM TPA. Cell lysis was performed as described in the IP protocol. HEK 293T lysate was incubated with CRTC3 rabbit hCRTC3(414-432) (PBL #7019) antiserum immobilized on Protein A magnetic beads (Dynabeads®, Life Technologies) and washed three times with lysis buffer. Bound CRTC3 protein was eluted using a CRTC3 peptide (Tyr⁴¹⁴ 415-432: YLAPYPTSQMVSSDRQLS) resuspended in lysis buffer (100 µg/ml final).

**Immunofluorescence**
HEK 293T cells (0.75 × 10⁶ cells) were plated in Poly-D-Lysine coated glass bottom dishes (MatTek Corporation) and under certain conditions reverse transfected with Lipofectamine® 2000 (Invitrogen) using 1 µg of mCRTC3 plasmid DNA (pUbC-3xFLAG backbone). 24 h post seeding/transfection, cells were either directly fixed with 4% paraformaldehyde or treated for 30 min with DMSO (0.05 % final), 10 µM Forskolin, 200 nM TPA prior to fixation. After incubation with primary antibodies (FLAG M2, PPP2R1A #2041, hCRTC3(414-432) PBL #7019), microscopy samples were incubated with secondary antibodies conjugated with Alexa Fluor® - 568 (goat anti-mouse / goat anti-rabbit) or in case of FLAG/PPP2R1A co-staining Alexa Fluor® - 568 and - 647 (568 - goat anti-rabbit & 647 - goat anti-mouse) (Life Technologies). Counterstaining with DAPI (Cayman Chemical Company) was performed before image acquisition (LSM 710; Carl Zeiss).

**Luciferase reporter assays**

Assays were performed in 96 well plates by reverse transfecting HEK 293T cells (100,000 cells). For each well 80 ng of DNA was used: 10 ng of EVX-Luc reporter plasmid (2x CRE half-sites, firefly luciferase) (Sonntag et al., 2017), 10 ng of FLAG-tagged CRTC1-3 plasmids, 60 ng of empty pUbC plasmid. 24 h post transfection 10 µM Fsk or 200 nM TPA were added and cells further incubated for 4 h (Fsk) or 5 h (TPA). Under all circumstances, DMSO served as the control treatment (each well 1% DMSO final). Next, 10 µl of Bright-Glo™ (Promega) was added per well and luciferase activity measured in a GloMax® multi microplate reader (Promega).

**Gene expression analysis**

CRTC3 WT and KO SVF cells were propagated to 60-80% confluency before (co-)treatment with 200 nM SCH772984 and DMSO (0.1 % final) for 4h and/or 10 µM Forskolin for 1h. Subsequently, cells were lysed in TRIzol® (Thermo Fisher Scientific) and RNAs extracted. cDNA was synthesized from 500 ng input RNA using First Strand cDNA synthesis kit (Roche) and quantified on a LightCycler® 480 II (Roche).

Primers used in quantitative real-time PCR (qPCR) experiments:

*RPL32*: 

14
FP 5’ TCTGGTGAGCCCAAGATCG 3’; RP 5’ CTCTGGTTTCCGAGGCTT 3’

NR4A1:
FP 5’ CTCTGGTCCCTGGACGTTA 3’; RP 5’ AGTACCAGGCTGAGCAGAA 3’

RGS2:
FP 5’ AACGGCCCCAGGTCGAGGA 3’; RP 5’ CGCTTCCTCAGGAGAAGGCTT 3’

**PP2A activity assay**
Experiments were performed in 6 well plates by reverse transfecting HEK 293T cells (2.5 × 10⁶ cells) with 2 µg plasmid DNA using Lipofectamine® 2000 (Invitrogen). 48 h post transfection cells were treated for 1 h with 1 µM Carfilzomib, collected in PBS, and resuspended in lysis buffer (50 mM Tris, 150 mM NaCl, 10% glycerol, 1% Igepal [Sigma-Aldrich], 1 mM DTT, EDTA-free cOmplete™ Protease Inhibitor Cocktail [Roche], 1 µM Carfilzomib; pH 8.0). The supernatant (= cell lysate) was either used in the PP2A activity assay or directly mixed with SDS-PAGE loading buffer. PP2A catalytic subunit (PPP2CA) was immunoprecipitated from cell lysates according to manufacturer’s instructions (PP2A Immunoprecipitation Phosphatase Assay Kit, Upstate). Phosphate release from 0.75 mM threonine phosphopeptide (KRPpTIRR) was detected by Malachite Green and measured in a Synergy™ H1 microplate reader (BioTek Instruments).

**Cell proliferation assay**
Experiments were performed in 96 well plates by reverse transfecting HEK 293T cells (100,000 cells). In each well 80 ng of GFP or CRTC3 plasmid DNA was used. Effects on HEK 293T proliferation were measured using the BrdU Cell Proliferation ELISA Kit (Abcam) according to manufacturer’s instructions in a Synergy™ H1 microplate reader (BioTek Instruments). BrdU treatment occurred either for 22 h (5 h post transfection) or for 4 h (23 h post transfection) prior to the colorimetric assay 27 h post transfection.
Sequence alignment

Amino acid sequences were aligned using MegAlign and Clustal W method (DNASTAR v7).

Statistical analysis

Data are either presented as the mean ± SEM or ± SD. One-way ANOVA was used for qPCR and CRE reporter data analysis. Statistical analyses were performed using either Microsoft Excel (Microsoft Corporation) or PRISM (GraphPad). Graphical presentations were generated using PRISM (GraphPad) and SigmaPlot (Systat Software Inc.).

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