A Common Docking Domain in Progesterone Receptor-B links DUSP6 and CK2 signaling to proliferative transcriptional programs in breast cancer cells

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ABSTRACT

Progesterone receptors (PR) are transcription factors relevant to breast cancer biology. Herein, we describe an N-terminal common docking (CD) domain in PR-B, a motif first described in mitogen-activated protein kinases. Binding studies revealed PR-B interacts with dual-specificity phosphatase 6 (DUSP6) via the CD domain. Mutation of the PR-B CD domain (mCD) attenuated cell cycle progression and expression of PR-B target genes (including STAT5A and Wnt1); mCD PR-B failed to undergo phosphorylation on Ser81, a ck2-dependent site required for expression of these genes. PR-B Ser81 phosphorylation was dependent on binding with DUSP6 and required for recruitment of a transcriptional complex consisting of PR-B, DUSP6 and ck2 to an enhancer region upstream of the Wnt1 promoter. STAT5 was present at this site in the absence or presence of progestin. Furthermore, phospho-Ser81 PR-B was recruited to the STAT5A gene upon progestin treatment, suggestive of a feed-forward mechanism. Inhibition of JAK/STAT-signaling blocked progestin-induced STAT5A and Wnt1 expression. Our studies show that DUSP6 serves as a scaffold for ck2-dependent PR-B Ser81 phosphorylation and subsequent PR-B-specific gene selection in coordination with STAT5. Coregulation of select target genes by PR-B and STAT5 is likely a global mechanism required for growth promoting programs relevant to mammary stem cell biology and cancer.

INTRODUCTION

Progesterone is an ovarian steroid hormone essential for breast development and implicated in breast cancer progression. Progesterone receptors (PR) exist primarily as two coexpressed isoforms, PR-A and PR-B (1,2), encoded by the same gene downstream of distinct promoters (3). PR-B, the full-length receptor, contains 164 amino acids at the N-terminus, not present in PR-A, termed the B-upstream segment (BUS) (4). Both receptors contain the same DNA-binding domain (DBD), a hinge region (H) and two activator function (AF) domains; PR-B contains a third AF domain in the BUS (5). Unliganded PR rapidly shuttles between the cytoplasm and the nucleus. After ligand binding, however, PR undergoes dimerization and is retained in the nucleus. Nuclear PR, together with coactivators and corepressors, activates or represses transcription of PR target genes, either directly through DNA binding to progesterone response elements (PREs) or indirectly through tethering interactions with other transcription factors (AP1, SP1, STATs) (6–9).

PR-mediated regulation of gene expression is controlled by many posttranslational modifications to the receptor, primarily on N-terminal serine (phosphorylation) and lysine (ubiquitination, sumoylation and acetylation) residues (10–15). These modifications significantly alter receptor stability, localization, transcriptional activity and target gene selectivity. PR is phosphorylated on serines (Sers) 294, 345 and 400 by mitogen-activated protein kinase (MAPK) and cyclin-dependent kinase 2 (cdk2). PR-B is also phosphorylated on Ser81 by ck2 (formerly casein kinase II; (16–19]), a ubiquitously expressed, constitutively active kinase that is overexpressed in every cancer examined thus far, including breast cancer (16,20). ck2-dependent PR-B phosphorylation of Ser81 regulates a specific subset of PR-B target genes involved in breast...
cancer cell growth and pro-survival, including BIRC3, HSD11B2 and HbEGF (19). In addition, ck2 is recruited along with Ser81-phosphorylated PR-B to enhancer sites of a subset of progesterone-responsive target genes (19). Notably, these studies have shown that ck2-dependent phosphorylation of PR-B Ser81 is unique to PR-B and thereby a primary determinant of PR isoform-specific activity. However, the molecular interactions necessary to support PR-B Ser81 phosphorylation have yet to be understood.

Posttranslational modifications to PR regulate the receptor’s interactions with other proteins (21). PR protein interaction domains include the estrogen receptor (ER) interaction domains [ERIDs (22)] and a poly-proline-rich (p-Pro) domain that is required for interaction between PR and the SH3-domain of c-Src (23). PR interacts with many other proteins via unknown domains (i.e. MEK1, FGFR2, and the SH3-domain of c-Src (23). PR interacts with many other proteins via unknown domains (i.e. MEK1, FGFR2, STAT5) (21,24–26). In a recent in silico analysis of the PR amino acid sequence aimed at identifying protein interaction domains, we identified a putative common docking (CD) domain in the N-terminal BUS region of full-length PR-B (21), a region that is not present in other PR isoforms. CD domains are commonly found in members of the MAPK family, where they mediate interactions between MAPKs and their upstream activators (MEKS), negative regulators (MAPK phosphatases [MKPs] or dual-specificity phosphatases [DUSPs]) and downstream targets (27,28). CD domains are characterized by clusters of negatively charged amino acids (aspartic or glutamic acid) that form electrostatic interactions with a positively charged ‘D domain’ in their respective binding partner [MKKs, DUSPs, MAPK substrates; reviewed in (29)]. The PR-B CD domain is an identical match to the CD domain of the MAPK family member, Erk2. This identical match suggests that PR-B interacts with the same D domain-containing proteins as Erk2. However, the function of this domain, unique to the PR-B isoform, has not yet been determined.

We predict that both PR-B Ser81 phosphorylation and PR-B CD domain interactions may be involved in breast cancer progression. Indeed, PR has been implicated in breast cancer progression in recent clinical studies of hormone-replacement therapy. These studies found that women taking estrogen and progesterone had more and larger breast tumors than women taking estrogen alone (30,31). In addition, we recently identified a phosphorylated PR-B gene signature associated with decreased survival in women with luminal (ER+ and/or PR+) breast cancer whose disease stopped responding to anti-ER therapy with tamoxifen (13). Mitogenic protein kinases—such as MAPK, c-Src, cdk2 and ck2—are commonly upregulated in cancer (17,32–34) and represent likely pathway components in PR-mediated gene expression in breast cancer. Understanding how PR interacts with these protein kinases and their regulatory protein partners is critical to understanding breast tumor etiology and developing better treatments. Herein, we sought to identify proteins that interact with PR-B via a novel N-terminal CD domain and how protein–protein interactions through this domain alter PR-B phosphorylation and transcription factor function in breast cancer models.

MATERIALS AND METHODS

Cell lines and constructs

The estrogen-independent ER/PR-positive T47Dco (T47D) variant cell line has been previously described (35), and is the parent cell line from which all T47D-variants used herein were created. T47D-Y (PR negative), T47D-YB (stably expressing wt PR-B) and T47D-YA (stably expressing wt PR-A) cells were characterized by Sartorius et al. (36). T47D-S79/81A PR-B cells have been previously described (19). T47D-mCD PR-B cells were created by stable expression of pSG5-mCD PR-B and pSV-neo in T47D-Y cells using FuGene-HD (Roche). Individual colonies were selected in 500 µg/ml G418 and maintained in 200 µg/ml G418 after initial selection in cMEM (as described in the Supplementary Materials and Methods). All cells were maintained as previously described or described in the Supplementary Materials and Methods (19). Creation of the pSG5-mCD PR-B, CD-PR-A and mCD-PR-A plasmids was performed as described in the Supplemental Materials and Methods. The DUSP6 construct [pcDNA3.1(-)Mye-His] was a gift from Stefanie Dimmel, University of Frankfurt.

Gene expression profiling

T47D cells stably expressing pSG5 empty vector, wt PR-B or mCD PR-B were serum starved in modified iMEM (Gibco) for 1 day and treated with R5020 (10 nM) or vehicle control for 6 h. Total RNA was extracted using a RNeasy kit (QIAgen) with on-column DNase I treatment (QIAgen). Triplicate RNA samples were labeled and hybridized to the Illumina HT-12v4 bead chip platform according to the manufacturer’s protocols. Chip scanning within Genome Studio software produced raw expression values that were analyzed within R software using the Bioconductor (37) package called lumi, where raw intensities were log2 transformed and quantile normalized. Differentially expressed genes were analyzed using the limma package, where empirical Bayes was used to better estimate the variance of the genes. Reported gene expression data contain log2 normalized intensities and biological comparisons (e.g. R5020/vehicle) contain log2 fold change with the Benjamini and Hochberg (BH)-adjusted P-value (38). Heat maps were generated through unsupervised hierarchical clustering of probes by the heatmap.2 function in the gplots R software package. Clustering was performed using Euclidean distance and complete linkage. Rows were scaled to have mean zero and standard deviation (SD) equal to one. Gene expression data are available in the NCBI Gene Expression Omnibus database (accession number: GSE46850).

Pathway and gene set analysis

Ingenuity Pathway Analysis (IPA) software (Ingenuity Systems, www.ingenuity.com) was used to compare biological functions or network pathways in cells expressing wt or mCD PR-B after progesterin treatment (wt + R5020/ wt-R5020 or mCD + R5020/mCD – R5020). Default IPA
settings for Core analyses were used to evaluate upregulated genes (fold change >1.5, P < 0.05). IPA Comparison analyses were used to reveal whether cells expressing wt or mCD PR-B regulated functionally distinct pathways. Analyses were scored based on significance (the BH-adjusted P-value, corrected for multiple hypothesis testing) and the threshold for a gene list to be significantly involved in a particular biological function was P < 0.05 [i.e. −log10(BH-adjusted P-value) > 1.30].

Gene set enrichment (GSEA) and Leading Edge analysis (39,40) was performed using the javaGSEA desktop software; all five gene set collections (c1–c5) from the Molecular Signatures Database (MSigDB) version 3.1 were queried. Data set files were developed based on normalized Illumina expression intensities from cells that constitutively express PR. Specifically, the log2 fold change values were compared for two phenotypes in our ligand-dependent analysis: (wt +R5020/−R5020) versus (mCD +R5020/−R5020). GSEA was executed using the default settings, except the permutation type was set to Gene set with 1000 permutations, and the metric for ranking genes was set to Diff_of_Classes because normalized expression data were log2 transformed. Each MSigDB collection was analyzed individually in multiple GSEA runs.

**Immunoblotting**

Immunoblotting was performed as previously described (19) and in the Supplementary Materials and Methods.

**Coimmunoprecipitation experiments**

For coimmunoprecipitation (CoIP) experiments, cell lysates were collected in RIPA (supplemented as described in the Supplementary Materials and Methods) and incubated on ice for 60 min. Cell lysates containing equivalent protein concentrations (1000 μg) were incubated overnight at 4°C with 2 μg appropriate antibody or control IgG. Protein G agarose (Roche Diagnostics) was added for the final 1 h of incubation time. Immune complexes were washed three times with supplemented RIPA buffer, resuspended in Laemmli sample buffer containing dithiothreitol and β-mercaptoethanol, boiled for 5 min and subjected to western blotting analysis.

**Transient transfections**

Transient transfections were performed as previously described (19) and in the Supplementary Materials and Methods.

**H2O2 assays**

After T47D-wt PR-B cells were starved overnight in serum-free iMEM, they were pretreated with 1 mM H2O2 for 20 min, followed by 30 min with 10 nM R5020. Protein lysates were isolated and analyzed as described above.

**Luciferase transcription assays**

Luciferase assays were performed as previously described (41) using the Dual Luciferase Reporter Assay (Promega). Relative luciferase units were normalized to *Renilla* ± SD.

siRNA

ON-TARGETplus SMARTpool designed to target human DUSP6 (DUSP6) and nonsilencing siRNA controls were purchased from Dharmacon. For siRNA experiments, T47D-YB cells were plated, and 24 h later they were transfected with 50 nM nonsilencing or DUSP6 siRNA. After 72 h, cells were treated with EtOH or 10 nM R5020 for 60 min. Protein lysates were isolated and analyzed as described in the Supplementary Materials and Methods.

**Reagents**

Cells were treated with the following reagents (when applicable): R5020 (10 nM; Sigma), AG490 (50 μM; CalBioChem) and H2O2 (1 mM; from Cell Biolabs OxiSelect™ Intracellular ROS Assay Kit).

**Cell cycle analysis/flow cytometry**

Flow cytometry for cell cycle analysis was performed as previously described (42).

**Real-Time Quantitative polymerase chain reaction**

Real-time quantitative polymerase chain reaction (RT-qPCR) was performed as described previously (19) and in the Supplementary Materials and Methods. Primer sets used for qPCR are listed in Supplementary Table S2. Relevant genomic sequence information and enhancer positioning for Wnt1 genomic sequence is based on GR37 Release 57 (March 2010).

**ChIP assays**

ChIP was performed using the ChIP-IT Express Kit (Active Motif) according to the manufacturer’s instructions using sonication as the method for chromatin shearing. Lysates were immunoprecipitated (IP) overnight (18 h) with the following antibodies: PR (ThermoScientific #MS-298-P), STAT5 (Santa Cruz Biotechnology sc-1081 and sc-836), ck2α (Santa Cruz sc-12738), DUSP6 (Santa Cruz Biotechnology sc-100374) or an equal amount of mouse or rabbit IgG. Resulting DNA was analyzed using qPCR as described above, and data are represented as a percentage of input DNA. In silico analysis using MatInspector (Genomatix) identified potential PRE-binding sites using the following consensus sequence: RGNACNRNNTGTNCY. Primer sets used for ChIP-qPCR are listed in Supplementary Table S2.

**CEAS**

Web-based CEAS analysis (http://ceas.cbi.pku.edu.cn/index.html) was performed on a publicly available PR ChIP-Chip data set (http://cistrome.org/PR_Cistrome/Cistrome.html). TRANSFAC and JASPAR motifs were used to determine putative transcription factor binding sites.

**Statistics**

Statistical significance for all experiments was determined using an unpaired Student’s *t*-test, unless otherwise specified.
RESULTS

PR-B CD domain is required for progestin-induced S-phase entry

We previously identified a putative CD domain (Figure 1A) (21) located in the N-terminal BUS region of full-length PR-B, a region that is absent from other PR isoforms (Figure 1B). To study the importance of this newly identified CD domain in modulating PR-B-specific functions, we mutated the critical negatively charged amino acids (D-aspartic acid, E-glutamic acid) to alanines (A), creating an mCD PR-B mutant (Figure 1B). Similar mutational strategies have been used to study CD domains present in Erks and other MAPKs (27,43). We then determined whether mCD PR-B was able to bind DNA and activate PRE-dependent transcription in luciferase reporter gene assays. PRE-luciferase expression levels were increased at similar levels in HeLa cells transiently expressing wt or mCD PR-B following treatment with vehicle (ethanol) or 10 nM R5020 (synthetic progesterone) (Figure 1C). These data suggest that the PR-B CD domain is not required for intrinsic PR-B transcriptional activity.

Progesterone or synthetic progestins (R5020) induce S-phase entry in breast cancer cells expressing PR-B (44–46), but not PR-A (47). Experimental isolation of PR isoform-specific actions is complicated by the fact that estrogen itself is a potent mitogen, but it tightly controls PR isoform expression (48). To overcome this barrier, we used the ER-positive T47Dco cell line, which expresses abundant PR-A and PR-B in the absence of added estrogen (35). A naturally occurring PR-negative variant of T47Dco cells, termed T47D-Y, was used to create stable cell lines constitutively expressing either wt PR-B (T47D-YB) or wt PR-A (T47D-YA) (49). We then tested the contribution of the PR-B CD domain to cell cycle progression by stably expressing mCD PR-B (T47D-mCD PR-B) in T47D-Y cells and analyzed progestin-induced S-phase entry. As predicted, there was an increase in progestin-induced S-phase entry in T47D-YB cells but not in T47D-YA cells (Figure 1D). Interestingly, cells stably expressing mCD PR-B also failed to enter S-phase upon treatment with R5020; these cells resembled PR-A expressing cells (Figure 1D). These data suggest that the PR-B CD domain is essential for proliferative signaling in breast cancer cells, as measured by progestin-induced S-phase entry.

PR-B CD domain regulates select PR-B target genes

Although mutation of the PR-B CD domain did not appreciably alter the absolute levels of PR-B transcriptional activity, posttranslational modifications can dramatically alter PR target gene selectivity, directing PR to specific enhancer or promoter regions in chromatin (50–52). This is an important feature of steroid hormone receptor action that is missed in reporter assay systems such as luciferase.

To determine whether the PR-B CD domain functions in the regulation of endogenous PR target genes, we...
performed global gene expression analyses using Illumina HT-12v4 whole genome bead arrays. Triplicate gene expression analyses were performed on T47D-Y (PR-null), T47D-YB and T47D-mCD PR-B cells after 6 h of treatment with R5020 or vehicle (ethanol). Transcriptional differences between cells expressing wt and mCD PR-B are evident in the heat map of significantly upregulated or downregulated (>1.5-fold) genes (Figure 2A). Differential regulation of numerous genes that require an intact CD domain for ligand-induced expression was validated by RT-qPCR (Figure 2B). Using IPA software, we compared the wt and mCD PR-B gene sets to a large database of genes that have been manually assigned to molecularly defined pathways, biological functions or disease states. Interestingly, genes that were specifically upregulated (>1.5-fold) in cells expressing wt but not mCD PR-B (P < 0.05, BH adjusted) were identified as significantly involved in pathways regulating cell proliferation, survival and cancer (Figure 2C; line represents P = 0.05 significance). Importantly, the individual genes validated by RT-qPCR (Figure 2B) were included in the CD-regulated gene set assigned to these IPA-defined pathways. These data suggest that the CD domain in PR-B is critical for PR’s ligand-dependent contributions to cell growth and survival pathways, and verify that the CD domain regulates a biologically significant subset of PR-B target genes.

**PR-B CD domain is required for PR-B Ser81 phosphorylation in response to ligand**

PR phosphorylation occurs on multiple sites and is a key determinant of receptor localization, ubiquitin-dependent turnover, tethering interactions and hormone responsiveness at selected PR target genes [reviewed in (53)]. We therefore screened for differences in basal and regulated phosphorylation of wt and mCD PR-B using phospho-specific PR antibodies. Notably, Ser81, a basally phosphorylated site that is further upregulated by ck2 in response to ligand-binding (19), failed to undergo basal (absence of ligand) phosphorylation in HeLa cells transiently expressing mCD PR-B (Figure 3A) or T47D cells stably expressing mCD PR-B (Figure 3B). Similarly, ligand-induced PR-B Ser81 phosphorylation was greatly diminished in cells expressing mCD PR-B relative to wt PR-B (Figure 3A and B). A time course of PR-B Ser81 phosphorylation in response to R5020 treatment verified that mCD PR-B is persistently weakly phosphorylated relative to wt PR-B (Figure 3C). Additionally, we analyzed PR phosphorylation on proline-directed sites (Sers 294, 345 and 400) in HeLa cells transiently transfected with either wt or mCD PR-B and then treated with R5020. Despite relatively equal levels of total wt or mCD PR-B expression, PR phosphorylation occurred with faster kinetics in cells expressing mCD PR-B relative to cells expressing wt PR-B. After 60 min, however, similar levels of phosphorylation were achieved in both groups (Supplementary Figure S1). These data suggest that mutation of PR-B’s CD domain dramatically alters PR phosphorylation in response to ligand. Namely, Ser81 fails to be persistently phosphorylated, while a number of proline-directed sites (Sers 294, 345...
and 400) appear to exhibit transient or temporary hyper-phosphorylation that is not persistently maintained.

PR-B CD domain interacts with DUSP6

The fact that mCD PR-B lacks Ser81 phosphorylation suggests that the CD domain may facilitate a specific interaction(s) between PR-B and one or more factors that are required for this phosphorylation event. An interaction between ck2 and DUSP6 [also called MAPK phosphatase 3 (MKP3)] has previously been reported (54). To test whether PR-B also interacts with DUSP6, COS cells were transiently cotransfected with constructs encoding wt or mCD PR-B and DUSP6 (myc-tagged) or vector-only controls. In cells transfected with wt PR-B and DUSP6, myc-tagged DUSP6 clearly communoprecipitated with wt PR-B in a largely ligand-independent manner (Figure 4A; lanes 3–4). In contrast, myc-tagged DUSP6 weakly copurified with mCD PR-B under the same experimental conditions (Figure 4A; lanes 7–8), indicating that the CD domain is mediating an interaction between PR-B and DUSP6.

To determine the specificity of PR-B’s interaction with DUSP6, we engineered two additional PR-specific mutants (Figure 4B) in which a wt or mutant PR CD domain was fused to the N-terminus of PR-A (termed CD-PR-A and mCD-PR-A, respectively). COS cells were transiently transfected with control (wt PR-B) and mutant PR constructs, as well as myc-tagged DUSP6. Reproducibly, wt PR-B and DUSP6 exhibited robust interaction as measured by CoIP, and mCD PR-B again displayed greatly reduced interaction with DUSP6 (Figure 4C; lanes 2 and 4). Wt PR-A coimmunoprecipitated with low levels of myc-tagged DUSP6, similar to levels observed for mCD PR-B (Figure 4C; lanes 5 and 4). Transient expression of CD-PR-A and mCD-PR-A fusion proteins remained poor relative to wt PR-A (Figure 4C; lanes 6–7 of Input lysates). However, wt PR-A and both PR-A fusion proteins were visible in western blots of immunoprecipitates (Figure 4C; lanes 5–7 of IP:PR blot). Despite relatively poor expression, the CD-PR-A fusion receptor communoprecipitated with myc-tagged DUSP6 at greater levels than wt PR-A (Figure 4C; lanes 6 and 5). However, the mCD PR-A fusion receptor completely reversed this effect (Figure 4C; lane 7), returning DUSP6 CoIP levels to approximately those seen with wt PR-A alone (no CD domain) and mCD PR-B. An additional region(s) common to both PR-B and PR-A (i.e. outside the BUS) is likely capable of weak interaction with DUSP6 (lanes 5 and 7). These data indicate that the PR-B CD domain is primarily responsible for mediating the interaction between wt PR-B and DUSP6.

DUSP6 is required for ck2-dependent PR-B Ser81 phosphorylation

We next sought to determine how PR-B’s interaction with DUSP6 is related to PR-B Ser81 phosphorylation. We previously identified PR-B Ser81 as a ck2-dependent site regulated in response to treatment of breast cancer cells with progestin, and during the S-phase of the cell cycle in the absence of progestin (19). If DUSP6 primarily functions to recruit ck2 for PR-B Ser81 phosphorylation, then loss of DUSP6 should block this phosphorylation event. To test this hypothesis, a DUSP6-specific siRNA was used to knock down DUSP6 CoIP levels to approximately those seen with wt PR-A alone (no CD domain) and mCD PR-B. An additional region(s) common to both PR-B and PR-A (i.e. outside the BUS) is likely capable of weak interaction with DUSP6 (lanes 5 and 7). These data indicate that the PR-B CD domain is primarily responsible for mediating the interaction between wt PR-B and DUSP6.
consistently exhibited decreased PR-B Ser81 phosphorylation relative to cells transfected with nonsilencing control siRNA (Figure 5A, left panel); a 50% decrease in DUSP6 protein levels resulted in at least 75% reduction of PR-B Ser81 phosphorylation (Figure 5A, right panel). As a control for functional DUSP6 knockdown, we measured Erk1/2 phosphorylation under similar conditions because DUSP6 phosphatase activity is a negative regulator of Erk1/2 phosphorylation. As expected, T47D-YB cells transfected with DUSP6 siRNA contained elevated levels of Erk1/2 phosphorylation relative to controls, indicating effective DUSP6 knockdown (Figure 5B). To confirm these results using independent methods, we chemically modulated DUSP6 phosphatase activity. Reactive oxygen species (ROS), produced as a result of treatment of cells with agents such as H2O2, block MKP enzyme activity (55,56), thereby resulting in high levels of Erk1/2 phosphorylation (Figure 5C, left panel). T47D-YB cells treated with either 1 mM H2O2 or vehicle alone, followed by R5020, exhibited similar levels of PR-B Ser81 phosphorylation (Figure 5C, right panel) despite effective DUSP6 enzyme (i.e. phosphatase) inhibition, as measured by increased phospho-Erk1/2. DUSP6 protein levels remained unchanged in the presence of high ROS (Figure 5C, right panel). Phosphorylation on other selected PR sites (Ser345 and Ser400) was relatively insensitive to H2O2 treatment (Supplementary Figure S2). These data suggest that DUSP6 enzyme activity is not required for PR-B Ser81 phosphorylation, as phospho-Ser81 levels remained unchanged even under conditions where DUSP6 phosphatase activity was greatly diminished. Cumulatively, these data suggest that the DUSP6 protein, but not its phosphatase activity, is required for efficient PR-B Ser81 phosphorylation, indicating that DUSP6 serves as a scaffolding protein that supports ck2-dependent PR-B Ser81 phosphorylation.

PR-B Ser81 phosphorylation is required for STAT5A and Wnt1 expression

To link CD domain-dependent regulation of PR-B Ser81 to gene expression, we examined known PR isoform-specific target genes for sensitivity to disruption of the CD domain. STAT5A (signal transducer and activator of transcription 5A) (25,57) and Wnt1 (wingless-type MMTV integration site family, member 1) (41,58–60) are primarily regulated by PR-B in response to progestin (25,41). To study the effects of PR-B Ser81 on STAT5A and Wnt1 gene expression, we used a previously well-characterized PR-B phospho-mutant that cannot be phosphorylated on Ser81, S79/81A PR-B (Ser81 and nearby Ser79 were mutated to ensure that this PR mutant remained fully resistant to phosphorylation on Ser81) (19). T47D cells expressing empty vector control (PR-null) or wt, mCD or S79/81A PR-B were treated with progestin (0, 1 and 6 h), and mRNA was harvested for RT-qPCR analyses. After progestin treatment, cells stably expressing wt PR-B robustly activated STAT5A and Wnt1 relative to PR-null cells, whereas cells expressing S79/81A PR-B exhibited greatly diminished STAT5A and Wnt1 mRNA relative to cells expressing wt PR-B
Interestingly, cells expressing mCD PR-B displayed an intermediate (reduced expression) phenotype, consistent with our finding that this mutant is only weakly phosphorylated on Ser81 over time (Figure 3B). STAT5B expression remained unaffected in cells expressing wt or mutant receptors (Supplementary Figure S3A). Notably, cells expressing the S79/81A phospho-mutant PR-B were phenotypically identical to cells expressing wt PR-A with regard to STAT5A and Wnt1 gene expression (Supplementary Figure S3B), confirming that these are PR-B–specific target genes. Additionally, tissue factor (TF) mRNA expression, which is independent of PR-B Ser81 (19), was similar in cells expressing either wt or S79/81A PR-B (Supplementary Figure S4A). Taken together, these data suggest that PR-B Ser81 phosphorylation—which is dependent on a CD domain-mediated scaffolding interaction between PR-B, DUSP6 and cdk2—is required for expression of select PR-B target genes (STAT5A and Wnt1) known to be critical mediators of mammary gland development, stem cell self-renewal and breast cancer cell proliferation.

**Figure 5.** PR-B’s interaction with DUSP6 is required for PR-B Ser81 phosphorylation. (A) DUSP6 is necessary for ligand-induced PR-B Ser81 phosphorylation. Left: T47D cells stably expressing wt PR-B (T47D-YB) were transfected with 50 nM nonsilencing (NS) or DUSP6 siRNA. Seventy-two hours after transfection, cells were treated with 10 nM R5020 or EtOH for 60 min. Lysates were analyzed via western blotting using p-S81, PR-B and DUSP6 antibodies. Right: Densitometry was used to determine the ratio of Ser81 phosphorylated PR-B in R5020-treated cells to total PR-B in cells transfected with NS or DUSP6 siRNA. (B) Knockdown of DUSP6 increased MAPK activation. T47D-YB cells were transfected and treated as in (A), and lysates were analyzed via western blotting using p-Erk1/2 and Erk1/2 antibodies. (C) DUSP6 phosphatase activity is not needed to mediate ligand-induced PR-B Ser81 phosphorylation. Left: Schematic showing how ROS production (increased through treatment with H2O2) decreases DUSP6 phosphatase activity, which subsequently leads to an increase in Erk1/2 activity and phosphorylation. Right: T47D-YB cells were pretreated with 1 mM H2O2 for 20 min, followed by 10 nM R5020 (or EtOH) for 30 min. Lysates were analyzed via western blotting using p-S81, PR, p-Erk1/2, Erk1/2 and DUSP6 antibodies.

GSEA is a powerful computational tool that can be used to determine whether publicly available gene sets (via the MSigDB) are significantly enriched in gene expression data sets (39,40). GSEA can identify gene sets that are significantly regulated in a particular microarray sample group (e.g. wt PR-B or mCD PR-B). Using GSEA, we compared ligand-regulated wt and mCD PR-B expression data sets (wt PR-B R5020/EtOH versus mCD PR-B R5020/EtOH) and identified enriched gene sets from the e5 Gene Ontology collection. Interestingly, genes from the
Figure 6. Endogenous phospho-Ser81 PR-B-target genes are JAK/STAT-dependent. (A) Select genes regulated by PR-B Ser81 phosphorylation. T47D-Y cells stably expressing either wt PR-B, mCD PR-B, S79/81A PR-B or unmodified (PR-null) cells were starved for 18 h in serum-free media, followed by treatment with 10 nM R5020 or EtOH for 0–6 h. mRNA levels were analyzed by qPCR. Error bars represent ± SD. (B) GSEA comparing gene expression data sets obtained using the Illumina Microarray described in Figure 3. Left: wt PR-B R5020/EtOH compared with mCD PR-B R5020/EtOH is shown as compared with the c5 Gene Ontology database, JAK_STAT_Cascade gene set. Right: Leading Edge analysis within GSEA (c2 Curated Gene Set) included 229 gene sets significantly enriched (FDR < 0.05) in wt PR-B cells, as compared with mCD PR-B. A subset of 38 gene sets (boxed region) contained substantial overlap (green shading) and were significantly involved in interferon signaling pathways. See Supplementary Table S1 for gene set names and statistics. (C) Ser81-dependent PR-B target genes are JAK/STAT-dependent. T47D-Y cells stably expressing wt PR-B (T47D-YB) were starved for 18 h in serum-free media. Cells were then pretreated (60 min) with 50 μM AG490, followed by 10 nM R5020 for 6 h. mRNA levels were analyzed by qPCR. Error bars represent ± SD. (D) PR-B Ser81 phosphorylation is unaffected by JAK/STAT-inhibition. T47D-YB cells were starved for 18 h in serum-free media. Cells were then pretreated (60 min) with 50 μM AG490, followed by 10 nM R5020 for 30 min. Lysates were analyzed via western blotting using p-S81 and total PR antibodies.
JAK/STAT signaling pathway (JAK_STAT_Cascade gene set) were significantly enriched in cells expressing wt PR-B relative to cells expressing mCD PR-B (Figure 6B, left panel), suggesting that mCD PR-B loses the ability to regulate genes in the JAK/STAT pathway. Leading edge analysis is a deeper examination used to compare only the significant gene sets (identified through GSEA) to each other to identify the following: (i) the individual genes that are highly associated in a particular sample group and (ii) the core gene sets that contain the majority of those highly associated genes. These methods help identify the pathways (and the core subset of genes within these pathways) that are significantly associated with a sample group. We performed leading edge analysis on gene sets that were significantly enriched (229 c2 Curated gene sets with FDR < 0.05) in cells expressing wt PR-B relative to mCD PR-B. Our analyses identified 38 gene sets that contained substantial overlap (Figure 6B, right panel; Supplementary Table S1) and revealed a substantial overlap of gene sets involved in interferon signaling, which is primarily mediated via the actions of STAT proteins. Cumulatively, these data suggest that the CD domain in PR-B is critical for progestin-dependent regulation of interferon/JAK/STAT-related signaling pathways.

Interestingly, female STAT5 and PR-B knockout mice exhibit similar developmental blocks in mammary gland alveologenesis (61). Furthermore, numerous studies have implicated the JAK/STAT pathway in PR target gene expression (11). Progestin-induced expression of selected PR-B target genes, including HSD11b2 and STAT5A, was significantly blocked by the JAK/STAT inhibitor, AG490 (62). Similar to STAT5A (Figure 6A), we previously showed that progestin-induced expression of HSD11b2 requires phospho-Ser81 PR-B (19). To test the requirement for JAK/STAT signaling in PR-B Ser81-dependent regulation of Wnt1 expression, T47D cells stably expressing wt PR-B were pretreated (1 h) with AG490 (50 μM) followed by progesterin (10 nM R5020 for 6 h) or respective vehicle controls. AG490 significantly diminished progestin-induced Wnt1 mRNA expression (Figure 6C); HSD11b2 and STAT5A were included as additional Ser81-dependent PR-B target genes and positive controls (62). In contrast, c-myc, a well-characterized PR target gene whose regulation is independent of both PR-B Ser81 (19) and JAK/STAT (62) remained unaffected by AG490 inhibition of JAK/STAT signaling (Figure 6C). Importantly, progestin-induced PR-B Ser81 phosphorylation was unaffected by pretreatment of cells with AG490 (Figure 6D). Thus, in addition to PR-B Ser81 phosphorylation, JAK/STAT signaling is also required for progestin-induced expression of a subset of PR-B target genes (STAT5A, Wnt1, HSD11b2).

To determine the role of STAT5 in coregulating phospho-Ser81 PR-B-dependent transcription, we analyzed a publicly available (http://cistrome.org/Cistrome/Cistrome_Project.html) PR chromatin immunoprecipitation ChIP-chip data set for the presence of STAT5 binding sites within or nearby PR binding sites. These ChIP-chip data were created using T47D breast cancer cells treated with vehicle or estrogen (to increase endogenous PR expression) followed by progesterone treatment for 45 min. Interestingly, CEAS [cis-regulatory element annotation system; (63)] analysis revealed a 1.8-fold enrichment of STAT5 DNA sequence binding motifs within PR binding sites as compared with random genome sampling (P = 8.22 × 10^{-11}). These data suggest that STAT5 may function as a pioneer factor by ‘opening’ sites in chromatin for subsequent transcriptional activation by PR-B, DUSP6 and ck2.

### PR-B Ser81 phosphorylation is required for binding to an enhancer region upstream of the Wnt1 promoter

Wnt1 is a potent breast oncogene and stem cell regulatory factor/morphogen. We previously demonstrated the existence of a PR-driven autocrine loop in which frizzled receptors, activated by progestin-induced Wnt1, transactivated epidermal growth factor receptor (EGFR), leading to increased cyclin D1 expression and breast cancer cell proliferation (41). Importantly, knockdown of Wnt1 blocked progestin-induced breast cancer cell growth in soft agar. To further understand how phospho-Ser81 PR-B regulates Wnt1 expression in collaboration with STATs, we performed an in silico analysis of Wnt1 promoter and enhancer regions. We identified four putative full-length PRE binding regions (Figure 7A; triangles), including a site located in the proximal enhancer region (~1 kb upstream from the transcriptional start site [TSS]; PRE1) and three sites located downstream of the TSS (+44, 65 and 66 kb from the TSS; PREs 2–4, respectively). To determine whether PR-B directly regulates Wnt1, we performed ChIP assays to detect relative PR-B recruitment to sites within the Wnt1 enhancer region. After cross-linking and sonication was performed, lysates from vehicle- (EtOH) or R5020-treated T47D-YB cells were subjected to ChIP using PR-specific antibodies. PR-null (T47D-Y) cells served as a negative control. In the presence of ligand, we detected robust recruitment (~16-fold) of wt PR-B to PRE1, while moderate levels of PR-B recruitment (~2–4-fold with ligand) were detected at the other PREs located downstream of the Wnt1 TSS (PREs 2, 3 and 4; Figure 7B). We next tested whether DUSP6 and ck2 were present in the transcription complexes at PRE1, the same region where wt PR-B was detected. Both DUSP6 and ck2 proteins were IP’d from similarly prepared lysates of ligand-treated T47D-YB cells; PR-B was also IP’d as a positive control (Figure 7C). In addition, STAT5-specific antibodies revealed that STAT5 was present at this Wnt1 regulatory site, although protein recruitment was not ligand-dependent (Figure 7C, bottom right). Cumulatively, these data demonstrate direct regulation of the Wnt1 enhancer at region PRE1 by PR-B, DUSP6 and ck2. Although these molecules were recruited on progestin treatment, STAT5 was pre-associated with this site.

### PR-B Ser81 phosphorylation mediates recruitment of PR-B to Wnt1 and STAT5A enhancer regions

Finally, we examined PR-B recruitment to Wnt PRE1 in cells expressing either wt or S79/81A PR-B. In contrast to efficient recruitment of wt PR-B to the Wnt1 enhancer region, S79/81A PR-B failed to be recruited to PRE1 at

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by guest on 29 July 2018
Figure 7. PR-B regulates Wnt1 transcription through binding to Wnt1 enhancer regions. (A) Schematic of the Wnt1 promoter/enhancer regions. PRE sites (triangles) are located upstream and downstream of the Wnt1 transcription start site (TSS; denoted with an arrow). Distance of the regulatory regions from the TSS is listed in kilobases (kb). (B) PR-B is recruited to Wnt1 PREs. T47D-Y cells stably expressing wt PR-B or unmodified cells (PR-null) were serum-starved for 18 h. Cells were then treated with 10 nM R5020 or EtOH for 60 min. Fixed lysates were subjected to ChIP with antibodies against PR-B or species-specific IgG (control), and qPCR was performed on the isolated DNA using primers designed to amplify the respective regulatory region (PREs 1–4). Recruitment of PR-B to PREs is shown as fold change between R5020- or EtOH-treated cells (R5020/EtOH). Error bars represent ±SD of triplicate experiments. (C) PR-B Ser81 phosphorylation-scaffolding complex is recruited to Wnt1 enhancer. T47D-Y cells stably expressing wt PR-B were serum-starved for 18 h. Cells were then treated with 10 nM R5020 or EtOH for 60 min. Fixed lysates were subjected to ChIP with antibodies against PR-B, ck2, DUSP6, STAT5 or species-specific IgG (controls), and qPCR was performed on the isolated DNA using primers designed to amplify Wnt1 PRE1. ChIP experiments were performed in triplicate. A representative experiment is shown here; error bars represent ±SD of technical replicates. (D) PR-B Ser81 phosphorylation is necessary for PR-B recruitment to Ser81-regulated genes. T47D-Y cells stably expressing wt PR-B or S79/81A PR-B were serum-starved for 18 h. Cells were then treated with 10 nM R5020 or EtOH for 60 min. Fixed lysates were subjected to ChIP with antibodies against PR or species-specific IgG (controls), and qPCR was performed on the isolated DNA using primers designed to amplify Wnt1 PRE1 or a STAT5A-enhancer site. ChIP experiments were performed in triplicate. A representative experiment is shown here; error bars represent ±SD of technical replicates.
the Wnt1 enhancer (Figure 7D, left panel). Similarly, ChIP assays revealed that wt but not S79/81A PR-B was recruited to a regulatory region within the STAT5A gene (Figure 7D, right panel). Importantly, both wt and S79/81A PR-B showed similar levels of recruitment to PRE regions associated with expression of TF [Supplementary Figure S4B and (19)], indicating that the lack of S79/81A PR-B recruitment to Wnt1 and STAT5A enhancer regions is not due to an inherent nuclear localization or DNA-binding defect of phospho-mutant S79/81A PR-B. These data suggest a possible feed-forward mechanism involving phospho-Ser81 PR and STAT5. Cumulatively, these results provide a basis for understanding mechanisms of PR-B isoform-specific target gene selection and suggest that phosphorylation of Ser81 is required for PR-B recruitment to selected PREs located within JAK/STAT-regulated genes.

**DISCUSSION**

Our studies identify a novel protein interaction domain (CD domain) in the PR-B N-terminus that mediates interaction between PR-B, DUSP6 and ck2, and is required for progesterin-induced S-phase cell cycle entry of breast cancer cells. DUSP6 binding to PR-B is required for ck2-dependent PR-B Ser81 phosphorylation and subsequent regulation of Wnt1 and STAT5A, two PR-B-specific target genes known to be critically involved in mammary stem cell maintenance and breast cancer cell proliferation. DUSP6 appears to play a novel scaffolding role in this complex, as DUSP6 knockdown but not inhibition of phosphatase activity blocked PR-B Ser81 phosphorylation. Cumulatively, these data support a model where DUSP6 binds the PR-B CD domain and scaffolds activated ck2 to mediate PR-B Ser81 phosphorylation (54). In addition, we defined Wnt1 and STAT5A as members of the same gene set regulated by phospho-Ser81 PR-B and JAK/STAT signaling. This finding complements previous studies showing that PR-B Ser81 phosphorylation is required for progesterin-induced expression of HSD11B2 and BIRC3 (19) and that both genes are sensitive to JAK/STAT pathway inhibition (62). Notably, we found that PR-B, STAT5, DUSP6 and ck2 are recruited to the same hormone-responsive PRE-containing region of the Wnt1 enhancer. Taken together, our data demonstrate a role for PR-B Ser81 phosphorylation in target gene selection of JAK/STAT-dependent genes that are seminal to mammary gland biology (Figure 8).

Furthermore, our data demonstrate a functional linkage between PR-B and STAT5 that provides a paradigm for coordinate activation of proliferative gene programs in mammary gland development and, pathologically, in ER+/PR+ breast cancer.

**CD domains and nuclear receptors**

The CD domain of PR-B is located in the N-terminal BUS region of full-length PR-B, a region that is absent from other PR isoforms (21). An *in silico* analysis of the most closely related, steroid hormone-activated nuclear receptors (i.e. ER, glucocorticoid receptor and androgen receptor) revealed the presence of weak potential candidate CD domain-like regions. However, none contained the conserved Erk2 amino acid sequence similarity found in the PR-B CD domain. A similar domain was identified in PPARγ, a more distant member of the nuclear receptor family, which mediates an interaction between PPARγ and MEK1 (64). Interestingly, deletion of the CD-like domain in PPARγ did not completely abrogate MEK1 interaction, suggesting that other areas of PPARγ also make contact with MEK1 or proteins that bind MEK1 (64). Similarly, in our study, mutation of PR-B’s CD domain did not completely disrupt DUSP6 binding. Additionally, mutation of PR-B’s CD domain did not alter MEK1 binding (data now shown), indicating that the CD domain in PR-B is not required for PR-MEK1 complex formation (21). Stable PR binding to components of MAPK pathways (MEKs or DUSPs) occurs via multiple domains (CD domain, p-Pro, ERIDs) in PR that are required for robust progesterin-induced MAPK activity. Moreover, both c-Src and ER are also found in protein complexes with components of the MAPK pathway (22,23,65). PR interaction with multiple signaling molecules illustrates that these pathways are fully integrated, and this integrated circuitry provides a basis for the highly selective context-dependent regulation of PR target genes. Notably, both the CD domain and p-Pro region occur in human but not mouse PR-B. This fact may explain why mouse models have not strongly implicated PR-B as a breast oncogene; PR-B is not well expressed in virgin mice relative to PR-A, and mouse PR-B lacks key domains required for linkage of mitogenic signaling pathways to PR-B–specific gene expression.

**Regulation of ck2-dependent PR-B Ser81 phosphorylation**

ck2 is a ubiquitously expressed kinase with >300 substrates (16). Unlike traditional growth factor-activated
protein kinases that require upstream inputs for full activation, ck2 is constitutively active. Although ck2 regulation is poorly understood, evidence suggests that ck2 activation is modulated primarily through subcellular localization, substrate distribution/complex formation, ck2 holoenzyme formation, small molecule interactions or autophosphorylation. In a previous report, we show that PR-B Ser81 is phosphorylated by ck2 in the presence of progesterin, but in the absence of progesterin PR-B Ser81 is mainly phosphorylated by ck2 during the S-phase of the cell cycle, when ck2 is nuclear and exposed to PR-B (19). Herein, we discovered that DUSP6 binding via the PR-B CD domain provides a mechanism for robust ck2-dependent phosphorylation of PR-B Ser81. Perhaps PR-B must be bound to DUSP6 to ‘accept’ Ser81 phosphorylation by ck2, either because of proximity restrictions whereby DUSP6 recruits ck2 into close proximity with PR-B Ser81 or because of substrate conformation changes in which DUSP6 binding to PR-B induces conformational changes that permit ck2-dependent Ser81 phosphorylation. Importantly, we observed constitutive PR-B binding with DUSP6 in the absence or presence of progesterins. However, transcriptional complexes containing PR-B, DUSP6 and ck2 are clearly recruited to the Wnt1 enhancer (PRE1) in a progesterin-dependent manner. Whether these protein–protein and protein–DNA interactions are regulated by additional factors (e.g. growth factors or STAT5) or circumstances (e.g. cell cycle) are questions requiring further study. Notably, ck2 is upregulated in many human cancers, including breast cancer (16,20). Preliminary data obtained from a small subset of PR-positive breast tumors [described in (13)] demonstrated that roughly half contained phospho-Ser81 PR-B (A. Daniel and C. Lange, unpublished observations). These findings suggest that PR-B Ser81 phosphorylation is clinically relevant, and underscore the importance of further study of PR-B phosphorylation and associated isoform-specific target gene expression in human breast tumors.

**DUSP6 may function as a scaffolding protein to promote cancer growth**

DUSP6 is a potent phosphatase responsible for reversing Erk1/2 phosphorylation and thus is a negative regulator of MAPK activity. Because of its role as a negative regulator of MAPK signaling, the central dogma has been that DUSP6 functions as a tumor suppressor in cancer, even though DUSP6 overexpression was often predictive of poor clinical outcomes [reviewed in (67,68)]. Recent data, however, have implicated DUSP6 overexpression as a negative prognostic marker in cancer development, progression and survival in many different types of primary cancers and cancer cell lines, including thyroid (69), lung (70), myeloma (71), melanoma (72,73), breast (74,75), colon (76), cervical (77), pancreatic (78) and glioblastoma (79). These paradoxical data suggest that DUSP6 may have multiple biological functions, independent of its long-studied role in attenuating MAPK activation. Herein, our data support a novel mechanism in which DUSP6 functions as a scaffold for assembly of transcriptional coactivators that drive tumor growth. We found that DUSP6 phosphatase activity was not required for PR-B Ser81 phosphorylation, but rather, DUSP6 acted as a scaffold protein to bridge the interaction between PR-B and ck2, thereby bringing ck2 into close proximity with its substrate (i.e. PR-B Ser81) (19,54). This scaffolding action represents a unique role for DUSP6. Indeed, DUSP6’s scaffolding function, rather than its phosphatase activity, may provide a molecular explanation for the growing body of data linking DUSP6 overexpression to poor clinical outcomes in many different types of cancer, including breast (74,75). In addition, classically defined roles of kinases and phosphatases clearly have broader scopes of action, such as bridging pathways previously thought to be unrelated (i.e. PR-B regulation by ck2) and direct participation in gene regulation as part of transcription complexes (19,21,80).

**Functional linkage of STAT5 and PR-B signaling**

Whole genome (cistrome) analyses have identified ‘pioneer factors’ for nuclear receptors. Pioneer factors are specialized subsets of transcriptional coregulators that bind to transcriptional enhancers, making them competent for subsequent transcriptional activation by transcription factors, such as steroid hormone receptors [reviewed in (81,82)]. For example, multiple pioneer factors (FOXA1, PBX-1, GREB1, AP2-γ) have been identified for modulation of ER binding, and similar factors have been identified for other nuclear receptors (81,83–86). Importantly, pioneer factors bind DNA before activation of transcription and function to ‘open’ sites in chromatin for subsequent transcriptional activation. We found that JAK/STAT pathway inhibition blocked the expression of multiple phospho-Ser81 PR-B target genes. We also identified STAT5 DNA sequence binding motifs within PR binding sites and STAT5 protein associated with PR-B target genes. Thus, it is tempting to speculate that STAT5 may act as a pioneer factor for phospho-Ser81 PR-B binding by ‘opening’ the enhancer regions of phospho-PR-B target genes. Interestingly, we also observed that phospho-Ser81 PR-B was required for STAT5A mRNA expression, suggestive of a feed-forward regulatory loop wherein many PR-B and STAT5 genes are coordinately regulated.

Recent reports provide some insight into how PR-B/DUSP6/ck2-containing protein complexes and STAT5A coordinate gene expression during breast cancer development and early breast cancer progression. STAT5A and Wnt1 have recently been implicated in PR control of mammmary stem cell maintenance and mammmary gland biology (60,87). Like PR-B, STAT5A is required for mammmary gland development, and both STAT5A and PR-B knockout mice have similar defects in mammmary gland development (87). Progesterone is a known activator of STAT5A mRNA and protein expression (25,57); however, the mechanism by which progesterone induces STAT5A expression is not well understood. Similarly, Wnts are important mediators of progesterone action in the normal (58) and pregnant mammmary gland (59).
Previous work published from our lab showed that phospho-PR-B-dependent upregulation of Wnt1 is required for breast cancer cell soft-agar growth in response to progestins (41). Wnts have recently been shown to be critical paracrine mediators of progesterone-induced expansion of mammary stem cells (60), and Wnt loss as a result of early parity has been linked to protection shown to be critical paracrine mediators of progesterone-response to progestins (41). Wnts have recently been required for breast cancer cell soft-agar growth in phospho-PR-B-dependent upregulation of Wnt1 is deregulation of the Wnt/β-catenin signaling pathway has been found in many human cancers, including breast cancer (89). Interestingly, unlike most other cancers, direct mutations of positive and negative regulators of the Wnt/β-catenin signaling pathway are rarely seen in breast cancer, despite the clear upregulation of downstream pathway endpoints, such as β-catenin stabilization and nuclear accumulation (90,91). Notably, progesterone/PR-B is a direct activator of this pathway (58,60). Potential involvement of these key mediators (Wnts, STAT5A) of mammary gland biology in progestin-induced breast cancer development or early tumor progression underscores the need to understand precisely how PR-B regulates these genes.

PR-B Ser81 phosphorylation is a major determinant of PR isoform specificity

We showed previously that phosphorylation of Ser81 is a significant determinant of PR isoform specificity; mutation of this residue in PR-B (or mutation of CD) confers PR-A–specific behavior with regard to target gene expression (19) and cell cycle entry (Figure 2C). Coordinate regulation of PR-B–specific target genes by phospho-Ser81 PR-B and STAT5 may explain the requirement for both factors during the same stage of mammary gland development. Notably, paracrine factors (i.e. Wnts, RANKL) derived from PR-B-positive progenitors or luminal precursor cells are believed to induce self-renewal of PR-null stem cells (60,92). PR-A and PR-B are usually coexpressed in the same tissues; cells that express only a single PR isoform are rare (93–95). A 1:1 ratio of PR-A to PR-B seen in normal tissues is often altered in malignant breast tissues, suggesting that balanced isoform action is crucial to normal adult mammary gland biology (93,96). PR-A, but not PR-B, gene silencing via promoter methylation was significantly associated with tamoxifen-resistant breast cancer (97).

Understanding the critical differences between PR-A and PR-B-dependent gene regulation as linked to DUSP6-dependent PR-B Ser81 phosphorylation by ck2 and JAK/STAT signaling may allow for highly selective isoform-specific therapies. Restoration of the balance between PR isoform actions may provide an innovative and complementary approach to existing endocrine therapies.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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