Estrogen receptor beta repurposes EZH2 to suppress oncogenic NFκB/p65 signaling in triple negative breast cancer

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Triple Negative Breast Cancer (TNBC) accounts for 15–20% of all breast cancer cases, yet is responsible for a disproportionately high percentage of breast cancer mortalities. Thus, there is an urgent need to identify novel biomarkers and therapeutic targets based on the molecular events driving TNBC pathobiology. Estrogen receptor beta (ERβ) is known to elicit anti-cancer effects in TNBC, however its mechanisms of action remain elusive. Here, we report the expression profiles of ERβ and its association with clinicopathological features and patient outcomes in the largest cohort of TNBC to date. In this cohort, ERβ was expressed in approximately 18% of TNBCs, and expression of ERβ was associated with favorable clinicopathological features, but correlated with different overall survival outcomes according to menopausal status. Mechanistically, ERβ formed a co-repressor complex involving enhancer of zeste homologue 2/polycomb repressive complex 2 (EZH2/PRC2) that functioned to suppress oncogenic NFκB/RELA (p65) activity. Importantly, p65 was shown to be required for formation of this complex and for ERβ-mediated suppression of TNBC. Our findings indicate that ERβ+ tumors exhibit different characteristics compared to ERβ− tumors and demonstrate that ERβ functions as a molecular switch for EZH2, repurposing it for tumor suppressive activities and repression of oncogenic p65 signaling.

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INTRODUCTION

Breast cancer is the most prevalent cancer among women worldwide1. Triple negative breast cancer (TNBC) accounts for 15–20% of breast cancer cases and is defined by the lack of estrogen receptor alpha (ERα) and progesterone receptor (PR) expression and the absence of Her2 overexpression/amplification. TNBC is a heterogeneous disease and represents the most aggressive subtype of breast cancer, accounting for a disproportionately high fraction of breast cancer mortalities2. Current standard of care for newly diagnosed TNBC patients involves a combination of chemotherapy, radiation, and surgery3. Unfortunately, approximately 46% of TNBC patients will develop distant metastasis, and survival time after diagnosis of metastatic disease is grim, with a median of only 13.3 months3. Further complicating the care of TNBC patients is the lack of treatment strategies that can be used in the adjuvant setting to prevent disease recurrence3. For these reasons, it is imperative to better understand the biology of TNBC and to discover and develop novel therapeutic strategies to combat this deadly disease.

Endocrine-based therapies such as tamoxifen and aromatase inhibitors remain the most utilized and effective class of drugs for the treatment of ERα-positive breast cancer but are not used to treat TNBC due to lack of ERα expression. Approaches to target other hormone receptors in TNBC such as the glucocorticoid receptor (GR) and androgen receptor (AR) are being explored with some early signs of success in a proportion of patients4,5. A second form of the estrogen receptor, ERβ, is reported to be expressed in approximately 20-30% of TN breast tumors6,7. ERβ is largely thought to function as a tumor suppressor in the breast given its high expression in normal mammary epithelium and loss of expression in most tumors8,9. Ligand-mediated activation of ERβ is known to elicit anti-cancer activity in TNBC cell lines in vitro and in vivo, effects that are reversed by anti-estrogens. Intriguingly, estrogen remains the only known drug to not only prevent breast cancer development, but to also reduce breast cancer mortality10,11,12. It is conceivable that ERβ plays a central role in mediating these clinical benefits of estrogen, a possibility that has not yet been explored. Therefore, a better understanding of ERβ in breast cancer is required. Further, elucidation of ERβ’s expression profiles, role in pathophysiology, and molecular mechanisms of action are likely to uncover novel biological processes that could be exploited for therapeutic purposes.

Here we report the incidence of ERβ protein expression in the largest cohort of centrally reviewed TN breast tumors to date13, as well as its association with clinicopathological features, TNBC subtypes and patient outcomes. Using multiple model systems and unbiased global screens, we have discovered a molecular mechanism by which ERβ suppresses TNBC proliferation and migration. Specifically, ERβ is shown to potentely inhibit the oncogenic transcriptional activities of nuclear factor kappa b (NFκB/RELA/p65) by repressing enhancer of zeste homologue 2 (EZH2) from a co-activator to a co-repressor of p65. The functions of this co-repressor complex rely on expression of p65 and the methyltransferase activity of EZH2 for ERβ to elicit anti-cancer...
effects. These results describe new biology of ERβ in TNBC and further our understanding of its mechanisms of action in this subtype of the disease.

RESULTS

ERβ expression in TNBC and its association with clinicopathological features

ERβ protein expression was assessed in 567 triple negative breast tumors from the Mayo Clinic TNBC Cohort using the well-validated PPGS/10 ERβ monoclonal antibody. ERβ positivity (ERβ+) was defined by moderate or strong nuclear staining in at least 25% of tumor cells. Based on this definition, ERβ was expressed in 102 of the 567 tumors (18%) (Table 1). Representative images of ERβ+ and ERβ- tumors have been previously published. None of the patients included in this cohort received neo-adjuvant therapy. Patients with ERβ+ tumors were less likely to present with lymph node involvement than those with ERβ-negative (ERβ-) tumors (26.0% in ERβ+ versus 37.1% in ERβ-; \(p = 0.04\)). The proportion of patients with 15% or more stromal tumor infiltrating lymphocytes (TILs) was greater in ERβ- tumors (43.1% in ERβ+ versus 55.6% in ERβ-; \(p = 0.03\)) (Table 1).

Regarding treatment, patients with ERβ+ tumors were less likely to receive aggressive regimens of chemotherapy (Anthracycline + Taxane) (8.8% in ERβ+ versus 22.2% in ERβ-; \(p = 0.009\)) (Table 1). ERβ expression was not found to differ with respect to age \((p = 0.48)\), BMI \((p = 0.84)\), histologic subtype \((p = 0.23)\), grade \((p = 0.39)\), tumor size \((p = 0.31)\), Ki67 \((p = 0.80)\), or AR expression \((p = 0.87)\) (Table 1). Regarding overall survival (OS), there were 58 deaths among the 215 pre-menopausal women. After accounting for age, tumor size, nodal positivity and receipt of adjuvant chemotherapy, pre-menopausal women with ERβ+ disease were more likely to have shorter OS than pre-menopausal women with ERβ- disease (HRadj = 2.06; 95% CI: 1.14–3.71; \(p = 0.017\)). Among 342 post-menopausal women, there were 138 deaths. After accounting for

| Characteristics               | Range        | ERβ positive \((n = 102)\) | ERβ negative \((n = 465)\) | \(p\) value |
|--------------------------------|--------------|-----------------------------|-----------------------------|-------------|
| Age (years)                    | <50          | 29 (28.4%)                  | 161 (34.6%)                 | 0.48        |
|                                | 50–69        | 54 (52.9%)                  | 221 (47.5%)                 |             |
|                                | >70          | 19 (18.6%)                  | 83 (17.8%)                  |             |
| BMI (kg/m²)                    | <25          | 40 (39.6%)                  | 171 (37.3%)                 | 0.84        |
|                                | 25–29.9      | 32 (31.7%)                  | 159 (34.7%)                 |             |
|                                | ≥30          | 29 (28.7%)                  | 128 (27.9%)                 |             |
|                                | Missing      | 1                           | 7                           |             |
| Histology                      | Apocrine differentiation | 6 (5.9%)                   | 29 (6.2%)                   | 0.23        |
|                                | Medullary features | 10 (9.8%)                  | 83 (17.8%)                  |             |
|                                | Metaplastic carcinoma | 9 (8.8%)                   | 35 (7.5%)                   |             |
|                                | Other        | 77 (75.5%)                  | 318 (68.4%)                 |             |
| Nottingham grade               | 1–2          | 14 (13.7%)                  | 50 (10.8%)                  | 0.39        |
|                                | 3            | 88 (86.3%)                  | 415 (89.2%)                 |             |
| Maximum tumor dimension (cm)   | at most 2.0  | 59 (57.8%)                  | 230 (49.6%)                 | 0.31        |
|                                | 2.1–5.0      | 38 (37.3%)                  | 210 (45.3%)                 |             |
|                                | 5.1 or larger | 5 (4.9%)                   | 24 (5.2%)                   |             |
|                                | Not stated   | 0                           | 1                           |             |
| Lymph node involvement         | positive     | 26 (26.0%)                  | 170 (37.1%)                 | 0.04        |
|                                | negative     | 74 (74.0%)                  | 288 (62.9%)                 |             |
|                                | not evaluated | 2                           | 7                           |             |
| Ki67                           | ≤15%         | 22 (22.0%)                  | 106 (23.5%)                 | 0.80        |
|                                | >15%         | 78 (78.0%)                  | 346 (76.5%)                 |             |
|                                | not obtained | 2                           | 13                          |             |
| Androgen Receptor              | 0            | 55 (67.1%)                  | 273 (67.4%)                 | 0.87        |
|                                | 1–15%        | 5 (6.1%)                    | 20 (4.9%)                   |             |
|                                | ≥15%         | 22 (26.8%)                  | 112 (27.7%)                 |             |
|                                | not obtained | 20                          | 60                          |             |
| Stromal TILs                   | 1–15%        | 58 (56.9%)                  | 204 (44.4%)                 | 0.03        |
|                                | ≥15%         | 44 (43.1%)                  | 255 (55.6%)                 |             |
|                                | not obtained | 0                           | 6                           |             |
| Chemotherapy type              | Anthracycline-based | 26 (25.5%)              | 96 (20.7%)                  | 0.009       |
|                                | Anthracyline/Taxane | 9 (8.8%)                  | 103 (22.2%)                 |             |
|                                | Taxane-based    | 4 (3.9%)                    | 6 (1.3%)                    |             |
|                                | Non-Anthracycline/non-Taxane | 14 (13.7%)              | 58 (12.5%)                  |             |
|                                | None/unknown  | 49 (48.0%)                  | 202 (43.4%)                 |             |

BMI: Body Mass Index, TILs: Tumor Infiltrating Lymphocytes.
the same variables, OS was not found to differ significantly with respect to ERβ status (HRadj = 0.96; 95% CI: 0.63–1.48; p = 0.867).

**ERβ expression among distinct subtypes of TNBC**

TNBC is a highly heterogeneous disease that has been subtyped by various groups. We therefore assessed the distribution of ERβ+ and ERβ− tumors across these different subtypes. In the four subtypes described using the non-negative matrix factorization (NMF) method, the majority of ERβ+ tumors were classified as basal-like immune activated (BLIA) (34.4% of all ERβ+ tumors) or basal-like immunosuppressed (BLIS) (43.8% of all ERβ− tumors), and no differences were observed between the distribution of ERβ+ versus ERβ− tumors across these four subtypes (p = 0.891) (Table 2). In the subtypes described by Jezequel et al. (Fuzzy), most ERβ+ tumors were classified as basal TNBC (bTNBC) (53.1%), but again no differences in the distribution of ERβ+ versus ERβ− tumors across subtypes were detected (p = 0.596) (Table 2). Last, using a classification system being developed by the Kalari group (CALAR), the vast majority of all tumors were classified as basal (bTNBC) (80.9%), and no significant differences were detected for the distribution of ERβ+ versus ERβ− cancers (p = 0.596) (Table 2). These data demonstrate that ERβ positivity is not significantly enriched within a particular subtype of TNBC.

**Molecular profiles of ERβ+ TNBC**

Given the association of ERβ with tumor suppressive phenotypes in TNBC, we sought to elucidate the molecular consequences of ERβ expression in this disease. Towards this goal, we screened a panel of TNBC cell lines to identify ERβ+ models. Of the 11 cell lines in our panel, none expressed appreciable levels of ERβ mRNA or protein (Fig. 1a, b) and none exhibited alterations in cell proliferation rates when treated with estradiol (E2), the ERβ-specific agonist 17βestradiol (E2), the ERβ-specific antagonist ICI 427277, or the ER antagonist fulvestrant (ICI), with the exception of HCC1937 and BT20 cells, which exhibited increased proliferation in response to ICI (Fig. 1c). Given the lack of TNBC models with endogenous ERβ expression, we utilized cell lines that stably express full length ERβ in a doxycycline (dox)-inducible manner (Fig. 1a, b). Expression of ERβ was shown to suppress TNBC cell proliferation following E2 and LY treatment, effects that were completely blocked by ICI in all three models (Fig. 1d).

To elucidate the gene expression changes elicited by ERβ, RNA-sequencing (RNA-seq) was performed on MDA-MB-231-ERβ cells following vehicle and E2 treatment for 5 days (GSE155685).

**Relevance of ERβ-mediated suppression of NFκB/p65 signaling**

RNA-seq data was available for a sub-set of ERβ+ (n = 32) and ERβ− (n = 225) patient tumors from the Mayo Clinic TNBC Cohort. Interrogation of these data revealed that the expression of multiple NFκB/p65 target genes were diminished in the ERβ+ tumors relative to ERβ− tumors, including BCL2A1, CXCL1, IL8, CXCL8, and VCAM1 (Fig. 2a). GSEA of differentially expressed genes in ERβ+ TN breast tumors revealed significant negative correlations with publically available gene sets related to NFκB signaling (Fig. 2b). To assess
whether ERβ-mediated suppression of NFκB/p65 target genes was reflected at the protein level, we performed a Cytokine/Chemokine array (Eve Technologies) using conditioned medium from MDA-MB-231-ERβ cells treated with vehicle or E2. Indeed, the protein levels of multiple NFκB/p65 target genes were significantly decreased in E2 conditioned medium (Fig. 2c). We next performed a co-culture experiment using various ratios of parental MDA-MB-231 cells expressing nuclear red fluorescent protein (RFP) and MDA-MB-231-
ERβ cells expressing nuclear green fluorescent protein (GFP) to determine the potential impact of changes in the ERβ regulated secretome on TNBC cell proliferation. As expected, ERβ- parental MDA-MB-231 cells were unaffected by E2 treatment in the absence of ERβ+ cells (Fig. 2d, left). However, the presence of ERβ+ cells resulted in inhibition of ERβ- cell proliferation rates following E2 treatment (Fig. 2d, middle panels), indicating that the alterations in the ERβ regulated secretome are of relevance to the anti-proliferative effects of ERβ-targeted therapies (Fig. 2d).

To further assess the importance of changes in canonical NFκB signaling with regard to the anti-cancer effects of ERβ, we determined the impact of p65 knockdown (Fig. 2e, f) and constitutive activation (Fig. 2g, h) on ERβ+ cell proliferation. siRNA-mediated knockdown of p65 in MDA-MB-231-ERβ cells significantly inhibited proliferation to an extent that was identical to E2 treatment (Fig. 2f). Further, the inhibitory effects of E2 were lost in the setting of p65 knockdown (Fig. 2f). Expression of a constitutively active form of NFκB/p65 (CA p65) in MDA-MB-231-ERβ cells (Fig. 2g) resulted in increased cell proliferation (Fig. 2h). Intriguingly, E2 was a more potent inhibitor of proliferation in cells with constitutively active p65 (Fig. 2h). Taken together, these data suggest that inhibition of p65 signaling is a primary mechanism by which ERβ elicits anti-cancer effects in TNBC cells.

ERβ blocks ligand-mediated activation of canonical NFκB/ p65 signaling

Based on our findings that ERβ repressed basal p65 signaling in TNBC, we next investigated whether ERβ could block ligand-mediated activation of the canonical NFκB/p65 signaling pathway. We chose to activate canonical NFκB/p65 signaling with tumor necrosis factor alpha (TNFα), a potent ligand for canonical NFκB/p65 signaling known to promote proliferation, migration, and primary tumor growth of TNBC [24-25]. TNFα was also chosen since it was identified as a top-upstream regulator in our IPA analysis of the ERβ transcriptome (Fig. 1d). RNA-seq analysis identified 915 genes regulated by TNFα in MDA-MB-231-ERβ cells (Fig. 3a, Supplemental Table 2). TNFα failed to significantly regulate 458 of these genes (50%) when E2 was present (E2+ TNFα treatment) (Fig. 3a, Supplemental Table 3). Indeed, E2 treatment largely blocked or attenuated the ability of TNFα to induce or repress NFκB/p65 target genes (Fig. 3b). Raw sequencing data is available in GEO under accession number GSE155685. These findings were confirmed via RT-qPCR in MDA-MB-231-ERβ cells (Fig. 3c) and in HS578T-ERβ cells (Supplemental Fig. 2a). As with the basal gene expression levels, treatment with doxycycline (dox) alone (i.e., expression of ERβ alone) also diminished the ability of TNFα to activate p65 target genes (Supplemental Fig. 2b). In addition, ERβ also prevented TNFα-mediated activation of a NFκB/p65 luciferase reporter construct in ERβ+ TNBC cells (Fig. 3d, e). Finally, TNFα induced migration of MDA-MB-231-ERβ cells similar to that of cells expressing a constitutively active form of NFκB (Fig. 3f). E2 blocked TNFα-induced migration and potently suppressed migration of the CA p65 cell line (Fig. 3f).

To gain insight into the mechanisms by which ERβ suppresses p65 signaling, we determined the impact of ERβ on phosphorylation of p65 (RELA) and its upstream inhibitor, IκBα. As expected, TNFα induced phosphorylation of both p65 and IκBα and induced loss of total IκBα in TNBC cells (Fig. 3g). However, E2 treatment had no impact on phosphorylation of p65 or IκBα alone or in the presence of TNFα (Fig. 3g), nor did E2 treatment of ERβ expressing cells affect p65 nuclear localization following TNFα stimulation (Fig. 3h).

ERβ interacts with p65 and alters its genomic localization

Given that ERβ did not alter phosphorylation or nuclear localization of p65, we hypothesized that ERβ suppresses p65 transcriptional activity by altering its chromatin binding profiles. To address this possibility, ChIP-seq was used to elucidate p65 chromatin binding sites following TNFα stimulation in the absence and presence of E2 (GSE155684). A total of 4618 and 2703 peaks were identified for p65 in MDA-MB-231-ERβ cells following TNFα treatment alone or in combination with E2, respectively (Fig. 4a). Approximately 60% of p65 binding sites identified following TNFα stimulation (2794 out of 4618) were diminished in the presence of E2, while 40% (1824 of 4,618) were maintained (Fig. 4a). Expression of a constitutive activation of NFκB/p65 (CA p65) in TNBC cells (Fig. 3g) resulted in increased cell proliferation (Fig. 2h). Intriguingly, E2 was a more potent inhibitor of proliferation in cells with constitutively active p65 (Fig. 2h). Taken together, these data suggest that inhibition of p65 signaling is a primary mechanism by which ERβ elicits anti-cancer effects in TNBC cells.
demonstrating that E2 treatment suppresses basal transcription of these genes and blocks their induction by TNFα (Fig. 4h).

ERβ assembles a co-repressor complex with EZH2 to inhibit p65 transcriptional activity

Since ERβ did not completely block p65 association with chromatin and was found to be co-localized at many NFκB/p65 binding sites throughout the genome, we next assessed the impact of ERβ on chromatin architecture at these genomic loci. ChIP-seq tracks for H3K27me3 and H3K27ac (GSE155684) at representative NFκB/p65 target genes are shown in Supplemental Fig. 4a. At NFκB/p65 binding sites, TNFα treatment resulted in a substantial decrease in the transcriptionally repressive H3K27me3 mark with minimal changes to the transcriptionally active H3K27ac mark (Fig. 5a, b). However, E2 treatment in combination with TNFα significantly increased H3K27me3 relative to TNFα alone (Fig. 5a, b). Given that EZH2 is the catalytic component of the PRC2 responsible for trimethylation of H3K27, we determined if ERβ interacts with EZH2 and other members of the complex. Co-immunoprecipitation studies using MDA-MB-231-ERβ nuclear extracts revealed that ERβ associates with EZH2, EED, and SUZ12 (Fig. 5c). p65 was also shown to be a component of this repressive complex (Fig. 5c). ChIP-qPCR at known ERβ binding sites near specific NFκB/p65 target genes demonstrated increased ERβ binding in response to E2 treatment, which also resulted in recruitment of p65 and EZH2 and was associated with increased H3K27me3 (me3) (Fig. 5d). Considering enrichment of H3K27me3 and recruitment of EZH2 to ERβ binding sites in close proximity to NFκB/p65 target genes, we examined the impact of drug-
mediated inhibition of EZH2 on NFκB/p65 target gene expression (Fig. 5e, Supplemental Fig. 4b). Treatment of MDA-MB-231-ERβ cells with GSK126, an inhibitor of EZH2 catalytic activity, resulted in significant up-regulation of five out of the 6 p65 target genes examined (Fig. 5e). In the presence of GSK126, the effects of E2 were attenuated and the ability of ERβ to suppress the expression of these genes below basal expression levels was lost with the exception of IL1B (Fig. 5e). Similar results were obtained in the Hs578T-ERβ cell line (Supplemental Fig. 4b).

p65 is required for function of the ERβ/EZH2 co-repressor complex

Since p65 was shown to be a component of the ERβ/EZH2 repressive complex and was recruited to ERβ-bound sites near NFκB/p65 target genes, we determined the necessity of p65 for ERβ-mediated gene silencing. As expected, siRNA-mediated knockdown of p65 reduced basal expression of NFκB/p65 target genes relative to control siRNA transfected cells (Fig. 5f). However, the inhibitory effects of E2 on the expression levels of these genes were diminished or completely lost in the setting of p65 knockdown (Fig. 5f). Conversely, NFκB/p65 target gene expression was induced in cells expressing constitutively active p65 (Fig. 5g), and E2 potently suppressed their expression (Fig. 5g). When p65 was silenced, recruitment of ERβ, p65, and EZH2 to known ERβ binding sites near NFκB/p65 target genes in response to E2 was lost (Fig. 5h). This coincided with a loss of H3K27me3 at these loci following E2 treatment (Fig. 5h).

DISCUSSION

Treatment options for patients with TNBC are limited, and identifying new biomarkers and therapeutic targets for this disease remains critical for improving outcomes. We assessed ERβ protein expression in the largest reported TNBC cohort using a well-validated monoclonal antibody\(^{16-18}\), and found that ERβ was expressed in 18% of TN breast tumors. Patients with ERβ+ tumors were more likely to present with decreased clinical stage (smaller tumors and no nodal involvement). Like ERα expression\(^{17}\), ERβ positivity was associated with significantly decreased stromal TILs, although the composition and frequency of specific types of lymphocytes in ERβ+ versus ERβ− tumors remains to be elucidated. Interestingly, NFκB/p65 is an essential player in the recruitment of TILs to the tumor stroma\(^ {28-29}\). Thus, ERβ-mediated suppression of p65 signaling may be in part responsible for the decrease in TILs in ERβ+ tumors. The proportion of AR positivity among the ERβ+ and ERβ− tumors was nearly identical in this TNBC cohort, demonstrating that these two important hormone receptors are not mutually inclusive or exclusive in this form of the disease. ERβ+ and ERβ− tumors were also similarly distributed.
across the known subtypes of TNBC. Interestingly, in pre-menopausal women with ERβ+ tumors, OS was found to be significantly worse than pre-menopausal women with ERβ− tumors. This association was not seen in post-menopausal patients. There could be multiple reasons for these findings including the fact that aggressive chemotherapy regimens were less frequently administered to women with ERβ+ disease. It is also possible that chemotherapy responsiveness may be diminished in ERβ+ tumors as is known to be the case for ERα+ disease. Further, the fact that ERβ+ tumors were less likely to have high levels of stromal TILs may have contributed to these findings given that high numbers of TILs are associated with improved outcomes in TNBC patients as well as increased chemotherapy responsiveness. Finally, it is possible that ERβ+ tumors that develop in young pre-menopausal women represent a different entity of disease compared to ERβ+ tumors that

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**Fig. 4** ERβ interacts with p65 and alters its genomic localization. a) Venn diagram depicting overlap of p65 binding sites following TNFα or E2+ TNFα stimulation in dox treated MDA-MB-231-ERβ cells as determined by ChIP-seq. b) Aggregate plots and (c) heat maps of p65 binding intensity in the presence of TNFα alone (Unique T), E2+ TNFα (Unique E + T), or both (Common) following indicated treatments. d) GIGGLE analysis of identified p65 binding sites assessing their similarity with other known protein binding sites in publicly available datasets. e) Venn diagram depicting overlap of ERβ and p65 binding sites identified via ChIP-seq. f) Bar graph of all p65 binding sites and their distribution relative to ERβ binding sites. Note the 936 overlapping sites from (e) are at the exact same genomic location. g) Co-immunoprecipitation experiments using nuclear lysates from dox treated MDA-MB-231-ERβ cells demonstrating protein interactions between ERβ and p65, but not RELB. h) ChIP-seq tracks from IGV for ERβ, p65, and RNA Polymerase II phospho-Ser2 ChIP-seq following dox plus veh, E2, T, and E2+ T treatment at NFκB/p65 target gene loci. See also Figure S3.
Develop in older post-menopausal patients. These possibilities warrant further study in additional patient cohorts and highlight the need to include menopausal status as a stratification factor when assessing ERβ associations with tumor characteristics and patient outcomes.

The uses of tamoxifen and aromatase inhibitors for chemoprevention of breast cancer are known to successfully reduce the incidence of ERα+ breast cancer but not TNBC31–34. However, these treatment interventions fail to reduce breast cancer mortality31–34. In contrast, long-term follow-up results from the randomized, placebo-controlled Women’s Health Initiative (WHI) trials in post-menopausal women showed that the use of estrogen alone as a menopausal hormone therapy decreased breast cancer incidence and death for patients with both ERα+ disease and...
In ERβ+ TNBC cells (left), EZH2 associates with p65 in a PRC2-independent manner and functions to enhance p65 transcriptional activity and promote aggressive cancer phenotypes. In ERβ− TNBC cells (right), ERβ induces formation of a co-repressor complex involving p65, EZH2, and the PRC2 complex (including SUZ12, EED, and EZH2). This repressive complex results in trimethylation of H3K27, chromatin compaction, and subsequent blockade of NFκB/p65 target gene expression, ultimately resulting in anti-cancer effects and less aggressive cancer phenotypes.

Additionally, there were reductions in lymph node positive disease for estrogen treated women, which parallel our findings for ERβ. Finally, E2 has been shown to improve OS of breast cancer patients when compared to tamoxifen. While the basis for these observations has yet to be elucidated, further studies evaluating the role of ERβ in these settings is warranted.

Uncovering the mechanisms through which ERβ functions is critical towards the goal of developing it as a useful prognostic biomarker and drug target. Here, we provide evidence that ERβ+ TN breast tumors have decreased p65 pathway activity and demonstrate that ERβ directly suppresses p65 transcriptional activity in TNBC cells. We also provide evidence that suppression of p65 signaling is a primary mechanism by which ERβ elicits its anti-cancer effects. Importantly, we found through co-culture studies that ligand mediated activation of ERβ resulted in alterations in cell-to-cell communication that led to decreased proliferation of non-ERβ expressing TNBC cells, potentially as a result of decreased levels of p65 regulated cytokines. These findings indicate that ERβ targeted therapies may still elicit anti-cancer effects in tumors with heterogenous expression of ERβ, a possibility that will need to be confirmed in cell line and PDX tumor models as well as clinical trials.

Canonical NFκB signaling via p65/RelA is known to promote carcinogenesis, enhance tumor cell survival and tumor progression, promote the development of metastatic disease, and induce resistance to standard of care chemotherapy regimens in TNBC. In addition, decreased expression of NFκB/p65 target genes is associated with improved outcomes for TNBC patients and decreased invasiveness of TNBC cell lines. Finally, suppression of IL6 and IL8, genes shown to be repressed by ERβ in the present study, have also been shown to potently inhibit proliferation, migration, and tumor formation of TNBC cells. Although drugs specifically targeting the canonical NFκB/p65 pathway elicit potent anti-cancer effects in vitro, they have thus far failed in the clinic due to off-target effects, immune suppression-related issues, and substantial toxicity. Our data suggest that ligand-mediated activation of ERβ is an effective way to block canonical NFκB/p65 signaling and invoke potent anti-cancer effects in TNBC.

Previous studies have shown that ERβ can suppress canonical NFκB/p65 signaling in various cell and tissue types, however the mechanism(s) remains unclear and these effects have not been studied in TNBC. Mechanistically, we provide evidence that ERβ does not alter the upstream signaling pathways required for p65 activation, nor does it impact p65 phosphorylation or nuclear localization. Instead, we show that ERβ modifies the genomic distribution of p65 in TNBC cells. At some genomic loci, ERβ partially or completely, displaced p65 association with chromatin in the setting of E2 treatment. Indeed, a substantial proportion (20%) of all p65 binding sites directly overlapped with an identified ERβ binding site. However, an equal number of p65 binding sites were identified that were either maintained or gained in the presence of E2, suggesting that ERβ alters p65 transcriptional activity via multiple mechanisms that extend beyond simply displacing p65 from chromatin. Interestingly, in the setting of p65 and ERβ activation (i.e. TNFα+E2 treatment), p53 motifs represented the most enriched binding site for p65. p53 and p65 are known to interact with one another and oppose each other’s activity. Since ERβ has also been reported to interact with p53, it is possible that estrogen treatment of ERβ+ TNBC cells drives the formation of a complex consisting of p65 and p53 to further dampen NFκB/p65 signaling. Indeed, we confirmed an interaction between ERβ and p65 using co-immunoprecipitation and proximity-based ligation assays. Further, we showed that E2 is a more potent inhibitor of proliferation and migration in TNBC cells expressing a constitutively active form of p65, suggesting that p65 plays an important role in mediating the anti-cancer effects of ERβ and indicating that ERβ-targeted therapies may be most effective in tumors with high NFκB/p65 activity.

To further understand the mechanisms by which ERβ suppresses p65 transcriptional activity, we assessed histone modifications and chromatin architecture at p65 binding sites. Our studies revealed that E2 treatment, both in the presence and absence of TNFα stimulation, significantly increased H3K27me3, a histone mark associated with gene silencing. Tri-methylation of H3K27 primarily occurs through the catalytic activity of EZH2 as a part of the PRC2 complex. Previous studies have shown that ERβ interacts with the PRC2 complex. We demonstrate that the methyltransferase activity of EZH2 is critical for ERβ-mediated silencing of NFκB/p65 target genes. Intriguingly, knockdown of p65 prevented ERβ-mediated growth suppression of TNBC cells and diminished ERβ’s ability to repress NFκB/p65 target gene expression. Knockdown of p65 also disrupted association of ERβ, EZH2, and H3K27me3 at ERβ binding sites near NFκB/p65 target genes, further confirming that these three proteins participate in the formation of a co-repressor complex that has not been previously described.

In contrast with our findings, previous studies have found that EZH2 mediates gene expression in breast and prostate tumors and functions as an oncogene by promoting canonical NFκB/p65 signaling in TNBC. However, it is important to note that these previous described “co-activator” properties of EZH2 occur through methyltransferase- and PRC2-independent mechanisms.
EZH2 overexpression in TNBC is known to enhance tumor progression and metastasis and is associated with advanced tumor stage and increased mortality.63–66 Paradoxically, H3K27me3 is associated with improved breast cancer patient outcomes.64,67 These findings suggest that in TNBC, the histone methyltransferase activity of EZH2 is usurped for non-canonical oncogenic functions, including the activation of NFκB signaling.62 Here, we demonstrate that in the absence of ERβ expression and activity, these functions of EZH2 occur, given that canonical NFκB/p65 signaling is high and H3K27me3 near p65 binding sites is low. However, upon treatment with inducible cells were maintained in the same medium supplemented with doxycycline (dox) to induce ERβ expression and, if necessary, treated with ethanol vehicle or 1 nM E2 for 5 days, with a media change on day 3. After 5 days, 20 ng/mL of TNFα was added for 24 h. RNA was isolated using TRIzol Reagent (Thermo Fisher Scientific, Waltham, MA, USA) and cDNA was generated from 1 µg of total RNA using the iScript cDNA Synthesis Kit (Bio-Rad). Real-time quantitative PCR (RT-qPCR) was conducted using PerfeCTa SYBR Green Fast Mix™ (Quanta Bioscience, Gaithersburg, MD, USA) and a Bio-Rad CFX Real-Time PCR detection system (Bio-Rad). RT-qPCR primer sequences are listed in Supplemental Table 4.

Materials and reagents
17β-estradiol (E2), doxycycline (dox), and Tumor Necrosis Factor alpha (TNFα) were purchased from Sigma-Aldrich (St. Louis, MO, USA). LY500307 (LY) was provided by Eli Lilly (Indianapolis, IN, USA). IC182,780 (ICI) was purchased from Tocris Bioscience (Bristol, United Kingdom). GS126 was purchased from Selleckchem (Houston, TX, USA).

Cell culture
HCC1937, HCC1143, MDA-MB-435, MDA-MB-436, BT549, MDA-MB-231, BT20, MDA-MB-468, and Hs578T cells were obtained from ATCC. SUM185 and SUM159 cells were purchased from BioVIT (Westbury, NY, USA). All parental cell lines were maintained in phenol red-free DMEM/F12 medium supplemented with 10% fetal bovine serum (FBS) and 1% antibiotic/antimycotic (AA). MDA-MB-231, MDA-MB-431, and Hs578T cells were engineered to express full-length human ERβ in a doxycycline (dox)-inducible manner (MDA-MB-231-ERβ) as previously described.12,22 Dox-inducible cells were maintained in the same medium supplemented with 5 µg/mL blasticidin S and 500 µg/mL zeocin. Parental MDA-MB-231 and MDA-MB-231-ERβ cells were developed to stably express nuclear red and green fluorescent protein, respectively, following infection with NucLight Red Lentivirus (#4625, Sartorius Stedim Biotech, Göttingen, Germany) and NucLight Green Lentivirus (#4624, Sartorius, Germany) at a multiplicity of infection (MOI) of 5. Following 72 h of infection, virus-containing medium was removed and replaced with fresh medium containing 500 µg/mL puromycin for selection. MDA-MB-231-ERβ cells stably expressing a constitutively active form of NFκB (S276G) were generated by transfection of the HA-tagged pCMV3 NFκB p65 expression vector (Sino Biological, Beijing, China) after point mutation generation via Quikchange PCR. A clonal cell line was generated following selection with 50 µg/mL hygromycin B. Experiments utilizing ERβ ligands were performed in growth medium containing 10% troglitazone (dexamethasone containing) and cDNA (Shanghai HyClone, GE Healthcare Life Sciences, Pittsburg, PA, USA). All cell lines were routinely checked for mycoplasma contamination and were authenticated via IDEXX BioAnalytics (Columbia, MO).

Real-time qPCR
Cells were seeded in 12-well plates in medium containing 100 ng/mL doxycycline (dox) to induce ERβ expression and, if necessary, treated with ethanol vehicle or 1 nM E2 for 5 days, with a media change on day 3. After 5 days, 20 ng/mL of TNFα was added for 24 h. RNA was isolated using TRIzol Reagent (Thermo Fisher Scientific, Waltham, MA, USA) and cDNA was generated from 1 µg of total RNA using the iScript cDNA Synthesis Kit (Bio-Rad). Real-time quantitative PCR (RT-qPCR) was conducted using PerfeCTa SYBR Green Fast Mix™ (Quanta Biosciences, Gaithersburg, MD, USA) and a Bio-Rad CFX Real-Time PCR detection system (Bio-Rad). RT-qPCR primer sequences are listed in Supplemental Table 4.

Western blotting
Cells were seeded in 6-well plates and treated as described above with the exception that TNFα treatments persisted for 10 min. Cell lysates were harvested using NETN buffer (150 mM NaCl, 1 mM EDTA, 20 mM pH 8.0 Tris, 0.5% NP-40 containing 1X Complete™ Protease Inhibitor Cocktail without EDTA (PI) (Sigma-Aldrich) and 1X PhosSTOP™ phosphatase inhibitor (Sigma-Aldrich). Protein concentrations were determined using the Pierce BCA Protein Assay Kit (Thermo Fisher Scientific) and a SpectraMax Microplate Reader (Molecular Devices, San Jose, CA, USA). Equal amounts of protein were separated on 7.5% SDS polyacrylamide gels, transferred to PVDF membranes which were subsequently blocked for 1 h at room temperature using 5% milk or 5% BSA in 1X TBST depending on primary antibody. Membranes were incubated with primary antibody overnight at 4°C, washed with 1X TBST, incubated with secondary antibody for 1 h at room temperature, and washed again. Blots were imaged on the Odyssey FC (LI-COR, Lincoln, NE, USA) using the ChemiImager (Alpha Innotech, San Leandro, CA). Intensities were calculated using NIH Image J software.

Proliferation assays
Cells were seeded in 96-well plates in replicates of six and treated with or without dox for 24 h prior to indicated treatments. Following treatment, cells were allowed to proliferate for seven days, at which point they were fixed with 25% (v/v) glutaraldehyde (Sigma-Aldrich) for 10 min, washed four times with water, stained with crystal violet, and washed again. Crystal violet was solubilized with 100 µl of a solution containing 100 mM sodium citrate and 50% ethanol and quantitated using a plate reader at 550 nm excitation. Co-culture proliferation assays were performed in the same manner with various proportions of ERβ– and ERβ+ cells seeded in the same wells for a total of 1000 cells/well at the start of the assay. Proliferation rates were monitored in real-time for the two populations by tracking the number of red and green nuclei using an IncuCyte® S3 system (Sartorius).
RNA was isolated using TRIzol® Reagent and a miRNeasy Mini kit (Qiagen, Hilden, Germany) following the manufacturer’s protocol. Total RNA was submitted to the Mayo Clinic Genome Analysis Core for sequencing (Rochester, MN). RNA libraries were prepared using 200 ng of total RNA and the TrueSeq Stranded mRNA Sample Prep Kit (Illumina, San Diego, CA, USA) according to the manufacturer’s instructions. Fifty base-pair paired-end reads were generated using an Illumina HiSeq 4000 sequencer and software (HD 3.4.0.38) with approximately 50 million fragment reads per sample. Base-calling was performed using Illumina’s RTA version 2.7.7. Library preparation and primary analysis was performed by the Mayo Clinic Medical Genome Facility Genome Analysis Core. Mapped reads were assigned using featureCounts and batch effect correction was performed with R/P lineage using a curated set of housekeeping genes to normalize the batch effect. An RPMK cutoff was used to remove lowly expressed genes (RPMK < 1) from further analyses. Comparison tests were performed using edgeR and significance was measured using $|\log_2(\text{fold change})| \geq 1.5, p < 0.05$, and FDR $\leq 0.1$.

RNA-seq of TN breast tumors

RNA-Sequencing data was generated for 301 formalin-fixed paraffin-embedded (FFPE) tissues from the Mayo Clinic TNBC Cohort. The raw sequencing files from RNA-seq were processed through the Mayo Analysis Pipeline for RNA-Seq (MAP-RSeq). After applying quality control (FDR) normalization (CQN) to account for gene length and library size. Differential expression results were estimated in the CQN method. Differential expression analysis was performed using the edgeR package by modeling the raw gene counts predicted by ER$^+$.p values reported for multiple testing adjustments.

Biological pathway and gene set enrichment analysis

Differentially expressed genes were utilized for canonical pathway and upstream regulator identification with Ingenuity Pathway Analysis software (IPA, Ingenuity Systems, Inc., Redwood City, CA, USA; http://www.ingenuity.com). Associations of gene signatures derived from RNA-seq data using MDA-MB-231-ERβ cell lines and human tumors with publically available databases were performed using Gene Set Enrichment Analysis (GSEA).

Cytokine/chemokine array

MDA-MB-231-ERβ cells were plated in triplicate in six-well plates in dox-containing medium and were treated with ethanol vehicle or 1 nM E2 for five days with a media change on day 2. On day 4, cells were washed with 1X PBS and media was changed to serum-free while maintaining indicated treatments. Conditioned medium was collected on day 5 and centrifuged at 4,000 rpm for 5 min at 4°C to remove any debris. 100 µl conditioned medium was flash frozen and sent to Eve Technologies (Calgary, Canada) for analysis using the Discovery Assay® 65-Plex Human Cytokine Array/Chemokine Array Panel (HD65). Results are presented as log2 fold-change between groups with false discovery rate (FDR) p values reported for multiple testing adjustments.

Migration assays

Cells were seeded in 10 cm dishes and treated for five days with indicated treatments with a media change and treatment refresh on day 3. At the time of the media change, cells were lifted and re-seeded at 30,000 cells/well in IncuCyte® imageLock 96-well plates while continuing pretreatment. After 5 days of treatment, wounds were created using the IncuCyte® WoundMaker Tool (#4563, Sartorius), cells were washed once, and medium with treatment was re-added. Plates were then placed in the IncuCyte® S3 machine and imaged once every two hours using the Scratch Wound Protocol. Wound closure was assessed using the Cell Migration Analysis software module (#9600-0012, Sartorius).

siRNA-mediated knockdown of NFκB

Pooled siRNAs designed to specifically target human p65 were purchased from Dharmacon (Lafayette, CO, USA). Cells were transfected with 5 nM ON-TARGETplus SMARTpool siRNA using DharmaFECT 1 transfection reagent (T-2005-01, Dharmacon) according to the manufacturer’s protocol. Non-Targeting siRNA Pool 1 (D-001206-13, Dharmacon) was used as a negative control. Cells were transfected with siRNAs 24 h prior to performing indicated treatments.

Luciferase assays

Cells were seeded in 24-well plates and treated with dox for 24 h prior to transfection with 100 ng of a pGL3 luciferase reporter construct containing NFκB response elements (NRE) using FuGENE 6 (Roche, Basel, Switzerland). Twenty-four hours post-transfection, treatments were treated with vehicle control, 1 nM E2, 20 ng/ml TNFa, or E2 + TNFa for an additional 24 h. Cells were washed once with 1X PBS and lysed using 1X Passive Lysis Buffer (Promega). Equal amounts of protein lysate were assessed for luciferase activity using Luciferase Assay Reagent and a GloMax-Dual Luminometer (Promega). Treatments were conducted in replicates of 6.

p65 nuclear localization assay

MDA-MB-231-ERβ cells were seeded in four-well Nunc™ Lab-Tek™ II Chamber Slides™ (Thermo Fisher) and treated in duplicate the following day with dox and vehicle or 1 nM E2. After 24 h, 20 ng/ml TNFa was added for 30 min. Cells were washed with 1X PBS, fixed for 15 min at room temperature with 4% formalin freshly diluted in 1X PBS from 16% formalin (Electron Microscopy Sciences), and permeabilized for 15 min at room temperature with 1X PBST. Cells were blocked for 30 min at room temperature using SuperBlock® (PBS) Blocking Buffer (Thermo Fisher) and incubated with primary antibody overnight at 4°C. Slides were washed with 1X PBST, separated from the chambers, and incubated with fluorescent-labeled secondary antibody and Hoechst 33258 (Thermo Fisher) for 30 min at room temperature in the dark. Cells were washed with PBST and rinsed with PBS prior to mounting with coverslips using ProLong™ Gold Antifade Mountant (Thermo Fisher). Slides were imaged using an LSM 780 inverted confocal microscope (Zeiss, Oberkochen, Germany) and ZEN Black software (Zeiss). Antibody information can be found in Supplemental Table 5.

Cell fractionation and co-immunoprecipitation

For cell fractionation, cells were plated in replicates of four in 15 cm dishes and allowed to adhere for 24 h. Cells were treated as indicated for 2 h, medium was removed, cells were washed with ice cold PBS, and pellets were collected in 1X PBS for subsequent nuclear, cytoplasmic, or whole cell lysate preparation. Whole cell lysates were prepared in the same manner as for western blotting. Cytoplasmic protein was extracted by incubating cells with cytoplasmic lysis buffer (10 mM Tris-HCl pH 8.0, 10 mM KCl, 0.1 mM EGTA, 0.1 mM EDTA) containing PI for 15 min on ice. Ten percent NP-40 was added, followed by vortexing and centrifugation at 14,000 rpm for 15 min at 4°C. Supernatants were saved as cytoplasmic extracts. Remaining pellets were washed twice with cytoplasmic lysis buffer, then resuspended in nuclear lysis buffer (20 mM Tris-HCl pH 8.0, 0.4 M NaCl, 0.1 mM EGTA, 0.1 mM EDTA) containing PI for 30 min on ice. Following incubation, lysates were centrifuged at 14,000 rpm for 15 min at 4°C and the supernatant was saved as nuclear extract.

Protein concentrations were determined using the Pierce™ BCA Protein Assay Kit. Five hundred micrograms of protein were used for overnight immunoprecipitation at 4°C. Following immunoprecipitation, protein complexes were captured using 40 µl Protein G Dynabeads™ (Thermo Fisher) for 2 h at 4°C with rotation. Beads were washed three times with NETN buffer and protein was eluted via boiling with 2X Laemmli Sample Buffer (Bio-Rad) containing β-mercaptoethanol for 5 min. Immunoprecipitated samples were subjected to western blotting, along with 40 µg of nuclear extract that was not subjected to immunoprecipitation as an input loading control. Antibody information can be found in Supplemental Table 5.

Chromatin immunoprecipitation followed by PCR and sequencing

Cells were plated in 10 cm dishes and treated in triplicate as indicated above for 5 days, followed by fixation for 10 min at room temperature with 1% parafomaldehyde and quenched with 125 mM Glycine (Sigma-Aldrich) for 5 min at room temperature. Nuclear extracts were prepared and immunoprecipitation with specific antibodies was carried out as previously described. ChIP-PCR was performed using the same conditions described for RT-qPCR using the specified ChIP-specific primers (Supplemental Tables 6) and 2µl chromatin solution (diluted 1:400).
For H3K27me3 and H3K27ac ChIP-seq, samples were prepared in an identical fashion. For p65 and RNA Pol II phospho-Ser2 ChIP-seq, fixed cell pellets were processed as previously described\(^{40}\) by the Mayo Clinic Epigenomics Development Laboratory (Rochester, MN, USA). ChIP-seq libraries were prepared from immunoprecipitated chromatin solutions and input DNA using the TruPLEX® DNA-seq Kit V2 (Rubicon Genomics, Ann Arbor, MI, USA). Libraries were sequenced using 50 base pair paired-end sequencing on an Illumina HiSeq 4000. Raw sequencing reads were analyzed using the HiChip pipeline\(^{41}\) to generate library-size normalized signal tracks for visualization and a list of peaks. Briefly, paired-end reads were mapped to the human reference genome (hg38) by Burrows-Wheeler Alignment (BWA)\(^{42}\) with default settings, and only pairs with at least one of the ends being uniquely mapped were retained for further analysis. Alignments were position sorted and duplicates were removed using the Picard tools (https://broadinstitute.github.io/picard/). Peaks were called using the MACS2 algorithm at FDR $\leq 1\%$. Visualization tracks and heat maps were generated by deepTools 2.0.

**Statistical analyses**

A Fisher’s exact test was used to assess differences between those with ERβ−–TNBC and those with ERβ+–TNBC with respect to patient and disease characteristics at primary diagnosis (Table 1). $P$ values $\leq 0.05$ were considered to be statistically significant. Cox modeling was performed to assess whether overall survival differed with respect to ERβ expression after adjusting for known prognostic factors and administration of adjuvant chemotherapy. Analyses were carried out using SAS 9.3. All in vitro experiments were conducted in biological replicates of at least three and with 3–6 technical replicates per assay. Representative data sets are shown. Student’s t-tests, one-way ANOVAs, and Wilcoxon Rank tests were used to determine statistically significant differences between treatments as indicated. $P$ values $\leq 0.05$ were considered statistically significant. Graphs and analyses were generated using GraphPad Prism 8 (GraphPad Software, San Diego, CA, USA).

**Reporting summary**

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**DATA AVAILABILITY**

All RNA and ChIP sequencing data that support the findings of this study have been deposited in the National Center for Biotechnology Information Gene Expression Omnibus (GEO) and are accessible through the GEO Series accession numbers GSE108981, GSE155684 and GSE155685. All other relevant data are included in the manuscript or available from the corresponding author upon request.

**CODE AVAILABILITY**

No custom codes were used to analyze the data presented in this manuscript and all codes are publicly available.

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ADDITIONAL INFORMATION

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