KMT2C mediates the estrogen dependence of breast cancer through regulation of ERα enhancer function

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Received: 26 March 2017 / Revised: 30 January 2018 / Accepted: 28 March 2018 / Published online: 14 May 2018
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Abstract
Estrogen receptor alpha (ERα) is a ligand-activated nuclear receptor that directs proliferation and differentiation in selected cancer cell types including mammary-derived carcinomas. These master-regulatory functions of ERα require trans-acting elements such as the pioneer factor FOXA1 to establish a genomic landscape conducive to ERα control. Here, we identify the H3K4 methyltransferase KMT2C as necessary for hormone-driven ERα activity and breast cancer proliferation. KMT2C knockdown suppresses estrogen-dependent gene expression and causes H3K4me1 and H3K27ac loss selectively at ERα enhancers. Correspondingly, KMT2C loss impairs estrogen-driven breast cancer proliferation but has no effect on ER- breast cells. Whereas KMT2C loss disrupts estrogen-driven proliferation, it conversely promotes tumor outgrowth under hormone-depleted conditions. In accordance, KMT2C is one of the most frequently mutated genes in ER-positive breast cancer with KMT2C deletion correlating with significantly shorter progression-free survival on anti-estrogen therapy. From a therapeutic standpoint, KMT2C-depleted cells that develop hormone-independence retain their dependence on ERα, displaying ongoing sensitivity to ERα antagonists. We conclude that KMT2C is a key regulator of ERα activity whose loss uncouples breast cancer proliferation from hormone abundance.

Introduction
Cancer-specific transcriptional programs are foundational to the development of oncogenic phenotypes. For example, oncogenes such as MYC and BRAF drive unique gene expression signatures that have been shown to be essential for transformation and cancer maintenance [1–5]. From a therapeutic vantage, reversal of these transcriptional programs is critical to the efficacy of most forms of targeted therapy.

Recent large-scale genomic analyses have identified key chromatin modifications permissive of such tissue-specific and cancer-specific transcriptional programs [6, 7]. Among the most significant of these chromatin modifications is methylation at histone H3 lysine 4 (H3K4me), generally marking regions of active and poised transcription. H3K4 histone methyltransferases mono-methylate H3K4, di-methylate H3K4, or tri-methylate H3K4 via their enzymatically active SET domain. Trimethylation (H3K4me3) of these residues is observed to be more abundant at promoter regions while monomethylation (H3K4me1) is more abundant at enhancer regions [8].
Interestingly, data from several large-scale cancer sequencing studies have identified \textit{KMT2C} (also referred to as \textit{MLL3}), which encodes an H3K4 histone methyltransferase, as one of the most commonly mutated genes in breast cancer with a mutation frequency of approximately 8\% \cite{9–11}. KMT2C, which is partially redundant with its most closely related homolog, KMT2D, is thought to be an H3K4me1/2 methyltransferase active predominantly at enhancer regions of the genome \cite{12, 13}. The high prevalence of \textit{KMT2C} mutation suggests that KMT2C may have important functions in breast cancer, which is so often characterized by its dependence upon the transcription factor, ER\textalpha. Here, we investigated the role of KMT2C in breast cancer pathogenesis and found it to be an essential ER\textalpha coactivator.

**Results**

**KMT2C is one of the most frequently mutated genes in breast cancer**

Within the H3K4 methyltransferase family, \textit{KMT2C} is by far the most commonly mutated member with a frequency of approximately 8\% in TCGA breast cancer samples \cite{10} (Fig. 1a). To assess the presence and persistence of \textit{KMT2C} mutation in breast cancer pathogenesis, we examined \textit{KMT2C} mutation frequency in metastatic tumors. \textit{KMT2C} mutations were found in 9.8\% of over 700 metastatic breast tumors analyzed, placing it among the most commonly mutated genes in breast cancer (Fig. 1b). The majority of \textit{KMT2C} mutations are frameshift, truncation or missense mutations, with a substantial proportion predicted to interfere with expression of the carboxy-terminal SET domain (Fig. 1c). The mutations appear to occur in all breast cancer subtypes with a relatively even distribution (Fig S1). Together this data suggests that KMT2C may act as a breast cancer tumor suppressor and might be a candidate regulator of H3K4me in these tumors.

**KMTC2 knockdown reduces cell proliferation of ER\textsuperscript{+} breast cancer cells**

In order to test the function of KMT2C in breast cancer, we cloned two short hairpins against \textit{KMT2C} \cite{14} (shKMT2C). We found that both short hairpins, shKMT2C\#1 and shKMT2C\#2, reduce expression of KMT2C by 60–70\% without affecting expression of its most closely related homologs, \textit{KMT2A, B,} or \textit{D} (Fig. S2A). We stably expressed shKMT2C\#1 and shKMT2C\#2 in a panel of cell lines representative of the clinical subtypes of breast cancer and found similar degrees of knockdown across the panel (Fig.
S2B). All cell lines used underwent next-generation sequencing and showed no clear deleterious mutation in KMT2C (Supplementary Table 1). To confirm sufficient knockdown of KMT2C protein levels, we used MCF7 cells that have been engineered to express HA at the C-terminal end of an endogenous KMT2C allele (KMT2C-HA cells). Expression of shKMT2C #1 and shKMT2C #2 in these cells resulted in knockdown of KMT2C-HA by immunoblotting (Fig. S2C), while not affecting protein levels of KMT2A, B or D (Fig. S2D).

We subsequently used the shKMT2C-expressing breast cancer models to assay the effects of KMT2C loss on cell

Fig. 2 KMT2C loss inhibits proliferation of ER+ cells. a–d Breast cancer cells stably expressing either shRenilla or shKMT2C were assayed for proliferation using the alamarBlue cell viability assay. Values correspond to the mean of six experimental replicates ± s.e.m. All cells are cultured in full serum containing media e. Parental and MCF7 KMT2C CRISPR cells were assayed for proliferation using the alamarBlue cell viability assay. Values correspond to the mean of six experimental replicates ± s.e.m. Data correspond to one representative assay from a total of two or three independent assays. f–i Indicated breast cancer cells stably expressing either shRenilla, shKMT2C or shKMT2D were assayed for proliferation using the alamarBlue cell viability assay. Values correspond to the mean of six experimental replicates ± s.e.m.
proliferation. KMT2C knockdown resulted in a 40–70% reduction in proliferation selectively in the three ER+ HER2− cell lines examined, MCF7, T47D and Cama-1 (Fig. 2a). In contrast to the effects seen in the ER+ HER2− cells lines, KMT2C knockdown had no effect on the proliferation of the ER+ HER2+ cell line BT474, the ER-HER2+ cell lines, SKBR3 and HCC1954 and the triple negative cell lines, MDA-MB-231, MDA-MB-468, MCF10A and...
KMT2C might be regulating signaling by ERα given the effects of KMT2C loss on estrogen-driven breast cancer models, we next examined whether and how KMT2C loss suppresses ERα activity by placing these marks at ERα target genes in the Kmt2c knockout mice to levels that were approximately 40–90% below levels seen in control mice (Fig. 3h). In addition, we assayed for nuclear expression of the PR, a known ERα-target gene, by IHC and found significant reductions (Fig. 3i). These data confirm that KMT2C regulates key aspects of the estrogen response program while not globally controlling ERα expression or activity.

KMT2C loss suppresses ERα function at enhancer sites

It has been speculated that H3K4me5 may help demarcate regions for ERα binding and activity [15]. Given the known enzymatic activity of KMT2C in generating H3K4me5 in other cell types [16], we hypothesized that KMT2C might regulate ERα activity by placing these marks at ERα binding sites. We first examined total H3K4me1 and H3K4me3 levels by immunoblotting and found no change.
following KMT2C loss, implying that KMT2C is not the sole methyltransferase responsible for these marks in ER+ cells (Fig. 4a). We next examined the pattern of genome-wide H3K4me1 and H3K4me3 following KMT2C loss using ChIP-sequencing. Whereas KMT2C loss had no meaningful effect upon H3K4me3 across the genome (data...
not shown), it appeared to selectively alter H3K4me1 at 869 affected loci, or 1% of the total 84,484 sites of H3K4me1 occupancy in control cells (Fig. 4b, Fig. S9).

To analyze the functional significance of these sites of H3K4me1 loss, we integrated these data with the RNA sequencing data (Fig. 3a) and found a strong enrichment for sites with H3K4me1 loss with sites whose nearby genes showed diminished expression with KMT2C knockdown (Fig. 4c). To be sure, we also included assessment of decreased H3K4me1 at promoter regions and found that it also correlated with decreased gene expression (Fig. S10).

However, the 73 promoter sites that correlated with reduced gene expression accounted for a small portion of overall KMT2C-mediated H3K4me1 loss. This is again consistent with previous studies indicating that H3K4me1 demarcates activity at enhancers as opposed to promoters. To confirm these effects upon ERα enhancers, we examined H3K4me1 at known ERα enhancer sites and found decreases in H3K4me1 from 15% at the PGR enhancer to 80% at the CA2 enhancer (Fig. 4d), given the prior data that ERα regulates gene expression through its actions at both promoter and enhancer sites, our integrated H3K4 methylation and RNA sequencing data points to KMT2C predominantly altering ERα activity via its effects upon ERα enhancer sites.

As the H3K4me1 mark is known to demarcate both regions of active transcription and regions “poised” for transcription, histone H3 K27 acetylation (H3K27ac) can be used to more strictly delineate those sites that are actively engaged in transcription [17]. To determine if KMT2C was regulating active enhancers, we assayed for H3K27ac at the AGR3, CA2, and PGR enhancer sites, where we saw loss of H3K4me1. We found significant reduction of H3K27ac at these sites (Fig. 4e). We went on to perform H3K27ac ChIP-sequencing on KMT2C knockdown MCF7 cells and found that H3K27ac was coordinately reduced at sites of H3K4me1 loss (Fig. 4f, Fig. S11). These data further establish a model in which KMT2C is responsible for depositing H3K4me1 selectively at active ERα enhancers thereby facilitating a key aspect of the ERα regulon.

H3K4me1 may facilitate enhancer function by promoting binding of any of a number of components necessary for site-specific transcription. It has been speculated that DNA-binding of ERα itself may require H3K4 methylation and/or pioneer factor binding [15, 18, 19]. To test this hypothesis, we examined ERα ChIP-sequencing in KMT2C knockdown cells. Unlike the case with H3K4me1/H3K27ac, we did not observe major loss of ERα binding at sites of changes in gene expression (Fig. S12A). Not surprisingly, we did find a handful of sites with decreased ERα binding and concomitant decreases in proximal gene expression (Fig. S12B). However, the 12 genes accounted for by loss of ERα binding were only 0.3% of the total gene expression changes observed. Moreover, we observed no change in levels of ERα at >75% of ERα binding sites proximal to sites we found to have KMT2C dependent H3K4me1 (Fig. S12A). As such, the data points to KMT2C regulating ERα enhancer function perhaps via its facilitation of essential ERα coactivators, but not via recruitment of ERα itself.

In order to nominate potential coactivators involved in mediating the effects of KMT2C upon ERα, we performed de novo motif analysis upon a 1kB radius around sites where H3K4me1 had decreased in shKMT2C cells. We found that these regions were significantly enriched over randomly selected background sequences for the well-known ERα pioneer factor, FOXA1 [15] as well as the ERα coactivator, AP-1 [20] (Fig. S13). While a number of motifs were considered to be significant in this analysis, it is notable that these two are both known factors necessary for ERα action. To further screen for potential intermediaries between KMT2C and ERα activity, we looked for changes in the ERα interactome. Given the large number of potential interactions, we sought to restrict this search to interactions occurring on DNA. To this end, we employed rapid immunoprecipitation mass spectrometry of endogenous proteins (RIME) [21, 22] with shKMT2C#2 MCF7 cells cultured in heavy SILAC media and shRenilla MCF7 cells cultured in light SILAC media. We compared our list of peptides differentially bound to ERα with previously published ERα-binding partners [23]. These data revealed only modest changes with most interactions comparable between cells (Fig. 4h). However, they did reveal reduced binding of ERα to MCM4, FOXA1, and TRPS1 by over 1/3 in shKMT2C cells and no proteins showing enhanced binding (Fig. 4h, full list Supplemental Table 2). Taken together, the findings point to a possible role for KMT2C in regulation of
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Fig. 5 KMT2C loss promotes hormone independent outgrowth. a Cells were assayed for proliferation in CSS media using the alamarBlue viability assay. Values correspond to the mean of six experimental replicates ± s.e.m. Data correspond to one representative assay from a total of two independent assays. b Supervised analysis of the 8622 differentially expressed genes between shKMT2C#2 cells in CSS media for 5 days and shKMT2C-R. CSS, charcoal stripped media. c Supervised analysis of the 9398 differentially expressed genes between shRenilla cells in CSS media for 5 days and shKMT2C-R. d GSEA of 4347 genes significantly downregulated in b as compared to the Hallmark Estrogen Response Early Geneset. e GSEA of 4559 genes significantly downregulated in c as compared to the Hallmark Estrogen Response Early Geneset. f MCF7 shKMT2C-R cells were treated with DMSO, estradiol (E2), fulvestrant (Fulv.), AZD9496 (9496), ARN1917 (1917), GDC927 (927), or RU58668 (58668) at indicated doses. Proliferation assayed using the alamarBlue viability assay. Values correspond to mean of 6 experimental replicates ± s.e.m. g Percent of cases with mutations detected in patient samples (MSKCC [50]) compared to those in TCGA Luminal A/B [10]. h Progression free survival for ER+ breast cancer patients on single agent aromatase inhibitor (AI)
the interaction between ERα and FOXA1, a well-established pioneer factor essential for ERα activity.

KMT2C loss promotes hormone-independent ER+ breast cancer cell proliferation

Given the necessity for KMT2C in estrogen-driven growth, we hypothesized that loss of KMT2C might facilitate a hormone-independent growth state as might occur in the context of hormone deprivation therapy. To assess the role of KMT2C in this context, we took advantage of the ability of MCF7 cells to grow in hormone-deprived conditions after 3–4 months [24, 25]. Using similar conditions, we observed that KMT2C-deficient MCF7 cells displayed a much more rapid time to growth with cells emerging in as little as 2 weeks (Fig. 5a). Consistent with these differences, control shRenilla cells continued to be growth arrested for 10–12 weeks beyond the time KMT2C knockdown cells grew. By contrast, we found that this rapid outgrowth was not observed in KMT2D knockdown cells (Fig. S14).

To investigate the basis of hormone-independent growth in this context, we established a hormone-independent MCF7 shKMT2C#2 cell line referred to as shKMT2C-R. Next-generation DNA sequencing (IMPACT) of the shKMT2C-R cells revealed no obvious acquired mutations compared to parental cells that could explain the rapid outgrowth in hormone-deprived media (Supplemental Table 3).

We assayed for KMT2C mRNA levels in these resistant cells and found that KMT2C levels remained suppressed (Fig. S15A, B). We also performed H3K4me1 ChIP-sequencing in these cells and found that the original 869 sites with reduced H3K4me1 continued to have reduced levels of H3K4me1 (Fig. S15C, D). These data suggested that the development of hormone-independent growth was not merely through replacement of KMT2C activity upon ERα enhancers by upregulating a different H3K4 methyltransferase (or suppressing the demethylase).

To interrogate whether there might be any unique dependencies of these cells, we performed RNA sequencing on the shKMT2C-R cells and compared their gene expression levels to that of parental shKMT2C#2 or shRenilla MCF7 cells maintained in hormone-deprived media for 5 days (Fig. 5b, c). We found that genes downregulated in shKMT2C-R cells were enriched for ERα target genes (Fig. 5d, e) and genes implicated in responses to growth factor stimulation (Fig S16). We also examined genes upregulated in the shKMT2C-R cells and found enrichment for genes involved in metabolic pathways including fatty acid, inositol phosphate, glucose and amino acid metabolic pathways. However, no individual pathway appeared significant when we corrected for multiple hypothesis testing (data not shown).

As the gene expression data suggested an ongoing suppression of ERα target genes, we examined whether these cells had become insensitive to the action of estrogen. We tested the response of shKMT2C-R to estradiol (E2) and found that the shKMT2C-R cells continued to respond to E2 by increasing their proliferative rate (Fig. 5f). These data suggest an ongoing responsiveness to the estrogen stimulated growth program, but a capability to proliferate in the absence of ERα ligands. To determine if unliganded ERα was still promoting growth in these cells, we examined their response to ERα antagonism. Similar to E2 response, shKMT2C-R cells were still dependent on ERα as they were sensitive to inhibition by multiple selective ERα degraders including fulvestrant, AZD9496, ARN1917, SRN927, and RU58668 (Fig. 5f). These data together suggested that the shKMT2C-R cells continue to be responsive to an E2-driven growth program while unliganded ERα maintains essential roles under hormone-deprived conditions.

Given the dual effects of KMT2C in promoting hormone-dependent tumor growth while suppressing hormone-independent tumor growth, we examined the clinical outcomes of patients with KMT2C loss. First, we applied the gene expression signatures derived from our shKMT2C cells (Fig. 3a) and our shKMT2C-R cells (Fig. 5c, d) and compared them to gene expression data from 1209 Luminal A and B breast cancer patients in the METABRIC data set [26]. We were able to separate the patients into three groups (high, mid, low) based on the similarity of their gene expression data to the derived gene expression signatures (Fig. S17). Those with a high score had a gene expression profile most similar to that of KMT2C knockdown. We found that patients with high scores were more likely to have poorer overall survival and be Luminal B in subtype than Luminal A (Fig. S17).

Together these data suggest that whereas KMT2C loss suppresses estrogen-driven tumor growth, it can also facilitate emergence of growth in hormone-depleted conditions. In keeping, we noted that KMT2C mutation prevalence was increased to over 30% among patients with metastatic, hormone-refractory breast cancer compared to the ~9% mutation frequency observed in primary, untreated Luminal A/B TCGA breast cancer (Fig. 5g, S18). In support, we went on to look at patients with metastatic breast cancer who received single agent aromatase inhibitor and compared patients wildtype or mutant for KMT2C (truncation mutations and deletions). We found that these patients with unambiguously pathogenic KMT2C mutations had significantly shorter progression-free survival (PFS) on hormone deprivation therapy than patients with wildtype KMT2C, further supporting a role for KMT2C in hormone resistance (Fig. 5h).
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**Figure:**

- **Panel a:** Illustration of ERα binding sites with shKMT2C-R and shRenilla.
- **Panel b:** Motif Name and their corresponding p-values for different elements.
- **Panel c:** Heatmap showing ERα binding at 10Kb downstream of MUC1 TSS.
- **Panel d:** Bar chart showing H3K4me1 at 10Kb downstream of MUC1 TSS.
- **Panel e:** Heatmap showing ERα binding at 20Kb downstream of IRX4 TSS.
- **Panel f:** Bar chart showing H3K4me1 at 20Kb downstream of IRX4 TSS.
- **Panel g:** Heatmap showing ERα binding at 3.5Kb upstream of CMYC TSS.
- **Panel h:** Bar chart showing H3K4me1 at 3.5Kb upstream of CMYC TSS.
ERα is reprogrammed in the shKMT2C-R cells

Our findings for shKMT2C-R cells demonstrated that they no longer required estrogen to support growth but maintained a dependence of ERα expression and reactivated ERα gene expression programs. To examine the site of action of ERα in this context, we performed ERα ChIP-sequencing of these cells. Whereas we continued to identify many sites seen in the shRenilla cells (n = 6598), we also identified over 10,000 unique ERα binding sites (Fig. 6a). We performed motif analysis on these distinct sites and found motifs of FOXA1 and AP-1 to be enriched (Fig. 6b, S19). AP-1, in particular, has been previously shown to play a major role in the hormone-independent outgrowth of ER+ breast cancer [27–33]. To test the relevance of AP-1 in shKMT2C-R cells, we looked for ERα binding nearby to target genes, IRX4 and MUC1, whose expression is known to be regulated by AP-1 dependent ERα activity [20]. We also assayed ERα binding at a known AP-1 dependent ERα binding site upstream of c-MYC, a key driver of hormone-independent growth [34]. We found significant increases in ERα binding downstream of IRX4 and MUC1 and at the known AP-1-dependent ERα binding site in shKMT2C-R cells as compared to control cells (Fig. 6c–h). These data raise the possibility that AP-1 may, in part, be responsible for reprogramming ERα under conditions of KMT2C loss.

Discussion

In this report, we utilize tumor genome sequencing analyses to help uncover the H3K4 methyltransferase KMT2C as a critical regulator of hormone-dependent ERα activity. Although it is unlikely that KMT2C will prove to be the only histone modifier that regulates ERα function, the findings in this report suggest it to be a key regulator of the estrogen responsive phenotype with major implications for hormone therapy of breast cancer.

The initial suggestion for a role for KMT2C in estrogen dependence comes from its frequent mutation in primary breast cancer. As most mutations appear to be deleterious, we modeled KMT2C loss in breast cancer cell lines. We were surprised to find that KMT2C loss led to a selective proliferative defect in ER+ driven breast cancers. These data were remarkable as no breast cancer model we tested showed an obvious growth advantage from loss of KMT2C. Given previous observations about the interaction between H3K4me and nuclear receptor activities [16, 35–45], we speculated that KMT2C might be an essential regulator of ERα.

To examine the mechanisms of KMT2C-mediated regulation of ER+ breast cancer proliferation, we analyzed the effect of its loss upon established gene expression signatures of ligand-activated ERα. KMT2C loss down-regulated estrogen responsive gene expression. These effects of KMT2C loss are notable in light of the previously reported effects of KMT2C loss in other contexts, including hematopoietic cells and stem cells, where its loss is predominantly associated with an increase in stem-like properties and chemotherapy resistance [20, 46, 47]. Specifically, previous work studying KMT2C in mouse mammary stem cells suggested it to be tumor-promoting [47], while our own work on human ER+ breast cancer cells suggest KMT2C to be either tumor-suppressive or tumor-promoting, depending on hormone availability. In support of KMT2C acting as a tumor suppressor, KMT2C loss has been shown to contribute to urothelial carcinomas by forming a tumor suppressive complex with p53 [45]. This disparity in KMT2C function across multiple tissue types points to the importance of cellular context in understanding the functions of KMT2C. Whereas this gene is among the most frequently mutated genes in all of cancer [5], the dominant effects of its loss appear to be cell type specific. Whether there are key shared functions of KMT2C across different tissue types or malignancies remains to be seen.

Early analyses of KMT2C biochemical activities and KMT2C knockout cells pointed to an essential role for this protein in generating the H3K4me1 mark associated more frequently with enhancer sites than with promoters [16]. Analysis of H3K4me1 in cancer cells suggested that KMT2C might be an essential regulator of ERα binding to enhancer sites. However, more recent analyses of KMT2C function in mouse models of cancer demonstrated that KMT2C might influence ERα binding at a subset of enhancer sites, leading to increased ERα activity in this context [16].

To understand the basis for how H3K4me1 might influence ERα activity, we conducted ERα ChIP-sequencing and found no significant loss of ERα binding despite the loss of expression of proximal ERα target genes. How then does H3K4me1 influence ERα activity at these enhancer sites? We note that in examining the region surrounding these bindings sites, it appears that sites predicted...
for ERα binding partners, such as FOXA1 and AP-1, are prevalent. In addition, we observed that KMT2C loss led to decreases in ERα interactions with multiple known binding factors including MCM4, FOXA1 and TRPS1. Finally, as independent proof of potential interactions of KMT2C and FOXA1, Jozwik et al. recently uncovered a direct interaction between KMT2C and FOXA1. How FOXA1, AP-1, and other specific factors ultimately coordinate KMT2C-dependent ERα activity will be an area for further investigation.

Finally, we sought to understand the basis for tumors selecting for loss of KMT2C. We note that not all KMT2C mutations seen in breast cancer will recapitulate loss of function but given that over 55% of KMT2C mutations are truncation mutations and given the lack of a specific hotspot mutation, our KMT2C knockdown cells provided a key model to study the functional role of KMT2C in breast cancer. We queried KMT2C mutation frequency at different stages of breast cancer progression and observed a ~7% mutation rate in primary breast cancer, a similar rate (~10%) in metastatic breast cancer and a higher rate in hormone-independent breast cancer (~30%). These data were in some contrast to the proliferative defects we observed in ER+ cell lines growing in hormone-containing media. We investigated whether there might be a selective advantage for its loss in estrogen-depleted conditions. Here, we indeed observed that KMT2C loss led to markedly accelerated outgrowth of cells. These data were further corroborated by examining how loss of KMT2C impacted patient survival. Among these patients, KMT2C loss was correlated with poorer survival outcomes and shortened PFS on aromatase inhibitor therapy. In accordance, it was recently published that low KMT2C expression was significantly associated with poor outcome [48]. Together, these data suggest that KMT2C loss is likely to have complex effects on the clinical behaviors of breast cancers. We anticipate that under some circumstances the loss of KMT2C will slow estrogen-driven tumor growth and potentially be associated with favorable outcomes. In other contexts, loss of KMT2C may be permissive for tumors developing an estrogen-independent growth program, perhaps only in selected genotypes where another growth program is sufficiently active. Moreover, even in this context, our data would suggest ERα may still be targetable as our shKMT2C-R cells were still sensitive to SERD-mediated ERα antagonism. Indeed, our analyses of these shKMT2C-R cells suggest that KMT2C loss was permissive for large-scale reprogramming of ERα such that it acquired thousands of new binding loci. The function of these new binding sites is not yet known, but the sensitivity of the cells to ERα inhibitors suggest there may be key oncogenic functions gained. It will thus be imperative to further understand the growth requirements for different KMT2C-mutated tumors to better utilize this biomarker in selecting the types of endocrine therapy most appropriate for this common clinical entity.

**Methods**

**Plasmids**

SGEP (pRRL) was a kind gift from C. Sawyers. psPAX2 and pVSVG vectors were kind gifts from P. Chi.

**Cell Lines**

All cell lines were maintained at 37 °C and 5% CO₂ in a humidified atmosphere. SKBR3, BT-474, Cama-1, T47D, MCF10A HCC1954 and MDA-MB-231 MDA-MB-468 and HCC1806 cell lines were obtained from the American Type Culture Collection, and MCF7 Tet-On cells were obtained from Clontech. HEK293CT cells were a kind gift from P. Chi. SK-BR-3, Cama-1, HCC1954 and MDA-MB-468 and MCF7 cells were grown in DMEM-F12, T47D and HCC1806 and MDA-MB-231 cells were grown in RPMI, and HEK293CT cells were grown in DMEM, supplemented with 10% heat-inactivated FBS, 100 µg/ml penicillin, 100 mg/ml streptomycin and 4 mM glutamine. MCF10A cells were grown in DMEM-F12 supplemented with 5% heat-inactivated horse serum, 100 µg/ml penicillin, 100 mg/ml streptomycin, 4 mM glutamine, 20ng/mL EGF, 500 µg/ml Hydrocortisone, 100ng/mL Cholera Toxin, and 10 µg/ml Insulin. All cell lines tested negative for mycoplasma.

No cell line used is in the database of commonly misidentified cell lines. MCF-7, SKBR3 and MDA-MB-231 were authenticated by STR profiling. Cama-1, and T47D and MCF10A, MDA-MB-468, HCC-1806, HCC-1954 were authenticated by next-generation (IMPACT) sequencing.

**Clinical samples (MSKCC set)**

We previously described use of a 410-gene targeted capture-based sequencing platform (MSK-IMPACT [49]) to analyze 39 tumors with either recurrence of disease after receiving adjuvant therapy or World Health Organization-defined progression of metastatic disease on therapy for genomic alterations [50]. Informed consent was received from all patients.

**Sequencing breast cancer cell lines for the presence of KMT2C alterations**

MCF7, T47D, Cama-1, SKBR3, BT-474, MCF10A, MDA-MB-468, HCC-1954, HCC-1806 and MDA-MB-231 cells
were sequenced using the next-generation sequencing (IMPACT) assay as previously described [49].

Generation of MCF7 KMT2C-HA by CRISPR/Cas9

The following guide RNA sequence targeting KMT2C exon 60 was selected using the Optimized CRISPR Design tool (http://crispr.mit.edu [51]): AGCCGCCCAGTACGACGGTACGCTAGCA.

DNA oligonucleotides were purchased from IDT and cloned into px458-GFP vector [52]. For homologous recombination, we purchased a custom IDT Ultramer 200 bp repair template with the HA sequence (TACCGATGTCCAGATTACGCT) directly upstream the KMT2C stop codon. This 200 bp fragment contained two silent mutations, one to remove the PAM site (TCC > TTC) and another to introduce a HincII restriction enzyme (GTGAAC > GTTAAC) site upstream of the HA tag. The 200 bp fragment also contained 87 bps of homology to KMT2C exon 60 upstream of the HA tag and 83 bp of homology to the KMT2C 3′UTR downstream of the stop codon. 12µg of the targeting construct and 240 nM repair template were nucleofected into MCF7 cells using the Lonza Nucleofector V kit and Program P020 on the Nucleofector device. Nucleofected cells were single cell sorted based on GFP positivity 48 h following nucleofection. Clones were screened for the presence of successful homologous recombination, we purchased a custom IDT Ultramer 200 bp repair template with the HA sequence (TACCGATGTCCAGATTACGCT) directly upstream the KMT2C stop codon. This 200 bp fragment contained two silent mutations, one to remove the PAM site (TCC > TTC) and another to introduce a HincII restriction enzyme (GTGAAC > GTTAAC) site upstream of the HA tag. The 200 bp fragment also contained 87 bps of homology to KMT2C exon 60 upstream of the HA tag and 83 bp of homology to the KMT2C 3′UTR downstream of the stop codon. 12µg of the targeting construct and 240 nM repair template were nucleofected into MCF7 cells using the Lonza Nucleofector V kit and Program P020 on the Nucleofector device. Nucleofected cells were single cell sorted based on GFP positivity 48 h following nucleofection. Clones were screened for the presence of successful HA insertion by KMT2C exon 60 PCR and subsequent restriction enzyme digest with HincII. Positive clones show two digested products while negative clones only had a single undigested band. A single positive clone, called KMT2C-HA, containing the HA coding sequencing downstream KMT2C was selected to carry out further studies. As a control, we used one of the negative clones from the screening process, called KMT2C crispr control.

Lentiviral Infections

We plated HEK293CT packaging cells at 2.5 × 10⁷ cells/tissue culture dish (10 cm in diameter) and transfected them with 2.25µg of the SGEV lentiviral vector (encoding shRenilla, shKMT2C #1 or shKMT2C #2), 2.25 µg of psPAX2 and 0.5 µg of pVSVG with X-tremeGENE HP (Roche) according to the manufacturer’s protocol. Conditioned medium containing recombinant lentivirus was collected and filtered through non-pyrogenic filters with a pore size of 0.45 µm (Millipore). Samples of these supernatants were applied immediately to target cells, which had been plated 18 h before infection at a density of 3 × 10⁶ cells/tissue culture dish (10 cm in diameter). Polybrene (Sigma) was added to a final concentration of 8 µg/ml, and supernatants were incubated with cells for 12 h. After infection, cells were placed in fresh growth medium and cultured as usual. Selection with 5–7 µg/ml puromycin was initiated 48 h after infection.

Proliferation assays

About 500–1000 cells in 200 µL of media were seeded per well of a 96-well plate with six replicates per sample. If necessary, cells were treated the following day, (Day 0). On days when the plates were measured, 25 µL of Resazurin (R&D Systems AR002) was added per well followed by 4 h incubation at 37 °C. Plates were then read using a SpectraMax M5 (Molecular Devices) and results were analyzed with Softmax Pro 6.2.2 software with an Endpoint Readtype (Excitation: 560 nM, Emission: 590 nM). Results were normalized to blank media with no cells.

For long-term estrogen deprived proliferation assay (Fig. 2f), 5000 cells were seeded in 200 µL of charcoal stripped media. Media was replenished twice a week during the course of the proliferation assay.

Immunoblotting

Cells were washed once with cold PBS and scraped off the plate with a rubber policeman. The cell suspension was briefly centrifuged to pellet cells, PBS was removed, and the cell pellet was stored at −80 °C until lysis. For cell lysis, pellets were resuspended in a non-denaturing lysis buffer (Cell Signaling Technology) supplemented with protease and phosphatase inhibitors (Pierce) and were sonicated briefly for 30 s. Lysates were cleared by centrifugation at 14,000g for 10 min, and protein concentration was determined using the BCA kit (Pierce), which measures the reduction of Cu2+ to Cu1+ by protein in an alkaline medium. For each sample, 25 µg of protein lysate was loaded onto 4–12% SDS-PAGE minigels (Invitrogen) for electrophoresis and immunoblotting against H3K4me1 (Cell Signaling 5326), H3K4m3 (Cell Signaling 5587), pERα (Cell Signaling 8757), β-actin (Cell Signaling 4970).

Quantitative RT-PCR

RNA was extracted from cells 48 h after transfection or from the mammary glands of 12-week old virgin females using the RNeasy Mini kit (Qiagen) according to the manufacturer’s protocol. cDNA was synthesized with 1 µg of RNA from each sample using the qScript cDNA SuperMix (Quanta Biosciences) according to the manufacturer’s protocol. Synthesized cDNA was diluted with one volume of DEPC-treated water, and 2 µl of the mixture was added to TaqMan PCR Master Mix (Applied Biosystems) along with primers. Relative quantification for each mRNA
was performed using the comparative CT method with a ViiA 7 Real-Time PCR system (Applied Biosystems). Samples were run in triplicate, and mRNA levels were normalized to those of RPLP0 or Gapdh for each reaction. TaqMan primers were all purchased from Applied Biosystems. Catalog numbers are as follows: AGR3 (Hs00411286_m1), BANK1 (Hs01099378_m1), CA2 (Hs01070108_m1), CACNA1G (Hs00367969_m1), CAV1 (Hs00971716_m1), CD44 (Hs01075861_m1), CXCL12 (Hs03676656_mh), ESRI (Hs00174860_m1), GDNF (Hs01931883_s1), IGFBP4 (Hs01075900_m1), KMT2A (Hs00610538_m1), KMT2B (Hs00207065_m1), KMT2C (Hs01005521_m1), KMT2D (Hs00231606_m1), KRT13 (Hs02558881_s1), PEG10 (Hs00248288_s1), PGR (Hs01556707_m1), RPLP0 (Hs00999902_m1), SERPINA1 (Hs00165475_m1), SOX5 (Hs00753050_s1), TFF3 (Hs00902278_m1), TMPRSS3 (Hs00917537_m1), Kmnc2 (Mm01156942_m1), Pgr (Mm00435628_m1), Gapdh (Hs99999902_m1), IGFBP3 (Hs00902278_m1), KMT2C, SERPINA1, and ESRI were all purchased from Applied Biosystems. Chromatin Immunoprecipitation

Complexes were purified using H3K4me1 (abcam 8895), H3K27ac (abcam 4729), and ERα (sc-453) -specific antibodies and DNA was purified using the PCR Purification Kit (Qiagen). ChIP primers are as follows CA2 enhancer forward: 5′-GCAATCCTGAGAAAATCTGAA-3′, reverse: 5′-GCTCTGGCTCAAACTATC-3′, PGR enhancer forward: 5′-GAGGGCTGTGAAACAAACA-3′, reverse: 5′-TGCTGAGATCACACCTAGACAA-3′, AGR2 enhancer forward: 5′-ACGAAGCCTGCTTCTGAAC-3′, reverse: 5′-GTTTACAAGACATCAACACATGA-3′, IRX4 enhancer forward: 5′-AGGGATCACTCCAGACTCCT-3′, reverse: 5′-CTGAGTGGCAAGTGGCTTC-3′, MUC1 enhancer forward: 5′-TCCTCGAGAGAGCAACTCC-3′, reverse: 5′-CCGAGATCCGGTGGTATG-3′, CMYC enhancer forward: 5′-AAGACTGCTCCTCGTCTGT-3′, reverse: 5′-TGCTGCTGCTGCTGTAGAA-3′.

Immunohistochemistry

Tissue were fixed in 10% neutral buffered formalin for 48 h, were processed in xylene and ethanol, embedded in paraffin, sectioned at 4 µm thickness, and stained with hematoxylin and eosin or with an antibody against the PR (Santa Cruz sc-538) and then detected with SignalStain Boost IHC Detection Reagent (Cell Signaling, 8114). Slides were examined by a board-certified pathologist (S. Monette, MSKCC).

Rapid immunoprecipitation mass spectrometry of endogenous proteins (RIME)

RIME was performed as previously described [21, 22]. Briefly, MCF7 cells were grown in SILAC Protein Quantification Kit DMEM:F12 (Thermo Fischer Scientific 88439) supplemented with dialyzed FBS (Life Technologies 88440) with either l-Lysine-2HCl (Thermo Scientific 88429) and l-Arginine-HCl (Thermo Scientific 88427) for light-labeled media or l-Arginine-HCl 13C6 (Cambridge Isotope Laboratories 13E-430) and l-Lysine-2HCl 13C6, 15N2 (Cambridge Isotope Laboratories 14F-832) for heavy-labeled media. A total of 60 million cells total with 30 million light labeled shRenilla cells and 30 million heavy labeled shKMT2C#2 cells was used for each replicate. Ten micrograms of antibody against ERα (Santa Cruz sc-543) were used for each replicate. There were three replicates done in total.

Immunoprecipitated samples were washed as previously described and digested overnight with trypsin, peptides desalted using C18 zip tips, and then dried by vacuum centrifugation. Each sample was reconstituted in 10 µL 0.1% (vol/vol) formic acid and 4 µL, analyzed by microcapillary liquid chromatography with tandem mass spectrometry using the NanoAcuity (Waters) with a 100-µm-inner-diameter × 10-cm-length C18 column (1.7 µm BEH130, Waters) configured with a 180-µm × 2-cm trap column coupled to a Q-Exactive Plus mass spectrometer (Thermo Fisher Scientific). Peptides were eluted with a linear gradient of 0–35% acetonitrile (0.1% formic acid) in water (0.1% formic acid) over 150 mins with a flow rate of 300 nL/min. The QE Plus was operated in automatic, data-dependent MS/MS acquisition mode with one MS full scan (380–1800 m/z) at 70,000 mass resolution and up to ten concurrent MS/MS scans for the ten most intense peaks selected from each survey scan. Survey scans were acquired in profile mode and MS/MS scans were acquired in centroid mode at 17,500 resolution and isolation window of 1.5 amu and normalized collision energy of 27. AGC was set to 1 × 10^6 [6] for MS1 and 5 × 10^4 [4] and 100 ms IT for MS2. Charge exclusion of unassigned and greater than 6 enabled with dynamic exclusion of 15 s. All MS/MS samples were analyzed using MaxQuant (Max Planck Institute of Biochemistry, Martinsried, Germany; version 1.5.3.3) at default settings with a few modifications. The default was used for first search tolerance and main search tolerance: 20 and 4.5 ppm, respectively. Labels were set to Arg6 and Lys8. MaxQuant was set up to search the reference human proteome database downloaded from Uniprot on September 6, 2016. Maxquant performed the search assuming trypsin digestion with up to two missed cleavages. Peptide, Site, and Protein FDR were all set to 1%. One unique peptide was required for high-confidence protein identifications and a minimum ratio count of two peptides (one unique and one razor) were required for SILAC ratio determination. The following modifications were used as variable modifications for identifications and included for protein quantification: oxidation of methionine, acetylation of the protein N-
A total of $2 \times 10^6$ MCF7 cells stably expressing either shRenilla, shKMT2C #1 and shKMT2C #2 using the RNeasy Mini kit (Qiagen) according to the manufacturer’s protocol. RNA size, concentration and integrity were analyzed using Agilent 2100 Bioanalyzer. Libraries were generated using Illumina’s TruSeq RNA sample Prep Kit v2, following the manufacturer’s protocol. Sequencing was done on the HiSeq2500 sequencer as 50-bp paired-read with 30–40 million reads per sample by the NYU Genome Technology Center.

The raw data were analyzed using Basepair software (http://www.basepairtech.com/) with pipelines including the following steps: raw reads were aligned to the transcriptome derived from UCSC genome assembly hg19 using STAR [53] with default parameters. Read counts for each transcript were measured using featureCounts [54]. Differentially expressed genes were determined using DESeq2 [55] and a cut-off of 0.05 on adjusted p-value (corrected for multiple hypotheses testing) was used for creating lists and heatmaps, unless otherwise stated.

**Chromatin Immunoprecipitation followed by sequencing**

A total of $2 \times 10^6$ MCF7 cells stably expressing either shRenilla, shKMT2C #1 or shKMT2C 2 were fixed with 1% formaldehyde at room temperature, lysed and sonicated leading to a DNA average size of 200 bp. 1ug of H3K4me1 (abcam 8895), H3K27ac (abcam 4729) and ERα (Santa Cruz HC-20)-specific antibodies was added to the samples and incubated overnight at 4 °C. The complexes were purified using ChIP grade protein-A/G magnetic beads (Thermo Scientific QB210263) followed by elution from the beads and reverse cross-linking. DNA was purified using PCR purification columns (QIAGEN). H3K4me1, H3K4me3 and ERα ChIP-seq libraries were prepared using 5ng of DNA and KAPA Hyper Prep Kit for Illumina, according to the manufacturer’s protocol. Libraries were validated using the Agilent 2200 Tapestation and KAPA library quantification qPCR assay and sequenced on a HiSeq2500 sequencer as 50-bp single-end reads by the NYU Genome Technology Center.

The raw data were analyzed using Basepair software (http://www.basepairtech.com/) with pipelines including the following steps: The raw fastq data were trimmed using trim_galore to remove low-quality ends from reads (quality <15) and adapter sequences. The trimmed data was aligned using Bowtie2 [56] to UCSC genome assembly hg19. Duplicate reads were removed using Picard and bigwig files were created for visualization. Peaks were identified with Macs1.4 [57] and transcription factor binding motifs were detected with Homer. Motifs were identified as enriched over 48,246 randomly selected background sequences with matched %GC content. Peaks overlapping with Satellite repeat regions were discarded and remaining filtered peaks were annotated using custom scripts based on UCSC Flat data [58], where peaks between $-2500$ and $2500$ bp of a transcription start site were marked as Promoter, the overlapping gene body were marked as Genebody and the rest were marked as Intergenic. For intergenic peaks, a gene was considered a target if it was within 1 Mb of the peak.

**Generation of conditional KMT2C knockout mice**

Mice with the Kmt2c floxed (F) allele were generated by Biocytogen (Worcester, MA) in a C57BL/6 background. ES cells were verified using Southern blotting. F/F mice were bred to transgenic mice expressing the Cre enzyme in the mammary tissue under the control of the MMTV long-terminal repeat (mice were a kind gift from S. Lowe). Cre-mediated excision of the Kmt2c gene was verified by genotyping PCR of DNA extracted from toe clips or mammary fat pads. Genotyping for the MMTV-cre transgenes were done using generic Cre primers, forward (oIMR1084): 5’-GGGTCTGGCAGTAAAAACTATC-3’, reverse (oIMR1085): 5’-GTGAAACAGCATTGCTCATT-3’, as well as internal positive controls, forward (oIMR7338): 5’-CTAGGCCACAGAATTTGAAATCT-3’, reverse (oIMR7339): 5’-GTAAGTGGAAAATCTTAGCAT-3’. KMT2C floxed allele was genotyped using A1 primers, forward: 5’-GCAAGATCAAGGAGCTGTCTGTTCAAGTATT-3’, reverse: 5’-TGCTCTTAGGTCACCGTAGCTACT-3’. Cre mediated excision of KMT2C exon 3 was verified by the presence of a 3’LoxP band at 372 bp using the following primers, forward: 5’-GGAGGCTGCTCGTCCAAGATT-3’, reverse: 5’-GTGGCTAGGTATCCACAAG-3’. For the in vivo experiments, 12-week were used. All mice were housed in the MSKCC animal facility.

**METABRIC [26] patients and tumor samples**

A total of 1209 Luminal A and B breast cancer patients with Illumina HumanHT-12 V3 expression beadchip array and complete clinical annotation from the METABRIC data set were included in the analysis.

**Scoring for shKMT2C signatures with ssGSEA in primary METABRIC patients**

ssGSEA [2] was used to compute a set of different scores based on in vitro-derived shKMT2C expression signatures.
For each sample independently, an ssGSEA score was calculated after gene expression levels were rank-normalized and ordered, using the empirical cumulative distribution functions of genes in the signature and the remaining genes as previously described [1, 2, 7] using the bioconductor package GSVA [6]. Patients were stratified into high, low and intermediate scorers using population quartiles for the numerical ssGSEA score.

**Study Population, clinical annotation and prospective sequencing for patients with aromatase inhibition treatment**

A total of 671 breast tumor specimens from 466 patients with ER+/HER2− metastatic breast cancer who received aromatase inhibitors in metastatic setting underwent prospective clinical genomic profiling between April 2014 and March 2017. For each treatment line, the time of biopsy collection for MSK-IMPACT testing was compared with the treatment start and stop dates. We excluded samples that were collected after the start of CDK4/6 inhibitor therapy (n = 263 samples) and only included samples that were collected prior to therapy initiation. For the patients with multiple sequenced specimens we only included one pre-treatment sample and selected the sample that was collected within a shorter time interval prior to initiation of aromatase inhibitor therapy. The final cohort included 347 pre-treatment samples from 347 patients.

For all 347 patients, tumor and patient-matched normal DNA samples were extracted from either representative formalin-fixed paraffin embedded tumor biopsy samples or mononuclear cells from peripheral blood. All specimens underwent massively parallel next-generation sequencing in a CLIA-certified FDA-approved laboratory using MSK-IMPACT, a hybridization capture-based next-generation sequencing assay, which analyzes all protein-coding exons and selected intronic and regulatory regions of 341 to 468 cancer-associated genes, all as previously described [4, 21]. Somatic mutations, DNA copy number alterations, and structural rearrangements were identified as previously described [21] and all mutations were manually reviewed. The oncogenic KMT2C alterations were assessed using the latest versions of the OncoKB [5] [www.oncokb.org] and cancer hotspots [6] [www.cancerhotspots.org] knowledge bases and only alterations that are considered “oncogenic” or “likely oncogenic” were included in the analysis.

**Generation of MCF7 KMT2C CRISPR by CRISPR/Cas9nickase**

The following guide RNA sequences targeting KMT2C exon 6 were selected using the Optimized CRISPR Design tool (http://crispr.mit.edu [51]): gRNA1 CTAGTGACCACCTCCA CACAACGG, gRNA2 AGAACCATTGTTAGTGACCTGG.

DNA oligonucleotides were purchased from IDT and cloned into pX335-GFP vector [52] to generate targeting constructs that were subsequently co-transfected in an equimolar ratio into MCF7 cells using Lipofectamine HP (Thermo Fischer Scientific). Seventy-two hours after transfection, cells were sorted using a MoFlo cell sorter (Beckman Coulter) for cells expressing Cas9 nickase (GFP-positive cells) and left to recover for 1 week before sorting for single cells and allowing colonies to form. Indels within the KMT2C exon 6 were detected using the Surveyor Mutation Detection kit (IDT) and confirmed by subcloning and sequencing of KMT2C exon 6. One clone showing knockout of all 2/3 KMT2C alleles was used for subsequent analysis.

**Statistics**

Patient data was collected retrospectively; therefore, we did not predetermine a sample size. Sample size estimates for animal studies were determined based on previous reports using similar sample sizes [59]. Samples were placed into their experimental groups according to their type, and so randomization was not used. Investigators were not blinded to the experimental groups. Only female mice were considered as predetermined by the fact that we were interested in phenotypic effects to the mammary gland. qPCR data and qChIP were obtained from independent technical replicates (n values indicated in the corresponding figure legends). Normal distribution and equal variance was seen in the majority of data and, so, we assumed normality and equal variance for all samples. On this basis, we used the Student’s t-test (two-tailed, unpaired) to estimate statistical significance. Analysis was conducted using Prism v7.0c (GraphPad Software).

Survival for the METABRIC patient data was reported using Kaplan–Meier curves, and significance was estimated with the log-rank test. Statistical analysis for survival curves of patients with aromatase inhibitor treatment used univariate Cox proportional hazard models to determine the association between oncogenic KMT2C alterations and PFS with disease progression on aromatase inhibitors or patient death. For patients with multiple lines of therapy, only the first treatment line from that class that was started after the MSK-IMPACT biopsy was included in the analysis. We tested the proportionality assumption of the Cox regression model through time-dependency analysis of selected genetic alterations (cox.zph function of the R package survival). We rejected the null hypotheses with a two-sided α = 0.05. Statistical significance was determined by log-rank test stratified by the treatment.
Study approval

All mice were treated with procedures approved by the Institutional Animal Care and Use Committee. Access to the METABRIC [26] cohort was obtained with consent from the MSKCC institutional review board for anonymized evaluation of genomic associations. Study involving patients treated with aromatase inhibitors was approved by the Memorial Sloan Kettering Cancer Center Institutional Review Board and all patients provided written informed consent for tumor sequencing and review of patient medical records for detailed demographic, pathologic, and treatment information (NCT01775072). Detailed treatment history data were obtained for each patient and included all lines of systemic therapy from time of diagnosis of invasive carcinoma to the study data lock in May 2017. The exact regimen as well as the dates of start and stop of therapy were recorded for each treatment line.

Data deposition

Our ChIP sequencing and RNA sequencing data has been deposited in NCBI GEO

Accession number: GSE100328.

Acknowledgements We would like to thank P. Chi for plasmids and cells, C. Sawyers for plasmids, M. Jasins and S. Lowe for mice and S. Monette for assistance with mouse pathology. We would like to thank J. Reis-Filho, W. Toy, Z. Li, A. Smith, R. Levine, N. Rosen and P. Chi for their helpful discussions. This work was funded by generous support of the Anna Fuller Foundation, the Cancer Couch Foundation, and the NCI Cancer Center Support Grant (CCSG, P30 CA08748). SC is funded by the Susan G. Komen Foundation. KG is funded by the NCI Cancer Center Support Grant (CCSG, P30 CA08748). AS is supported of the Anna Fuller Foundation, the Cancer Couch Foundation, for their helpful discussions. This work was funded by generous support of the Anna Fuller Foundation, the Cancer Couch Foundation, and the NCI Cancer Center Support Grant (CCSG, P30 CA08748). SC is funded by the Susan G. Komen Foundation. KG is funded by the Geoffrey Beene Cancer Research Center and a Grayer Fellowship. AP is a Mildred–Scheel Postdoctoral Research Fellow of the Deutsche Krebshilfe e.V.

Author contributions SC, KG, AS, PR and JH conceived and designed the experiments. KG, AS, QL, PR, FS, AP, YC and SC conducted the experiments. KG, AS, FS, AP, NS, MB, RH, and SC analyzed the data. KG, AS, FS, AP, OAW, RH, and SC wrote the manuscript.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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