## Article

# **Cell Reports**

## A P53-Independent DNA Damage Response Suppresses Oncogenic Proliferation and Genome Instability

## **Graphical Abstract**



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## In Brief

The origins of genome instability in cancer remain poorly understood. Fagan-Solis et al. reveal a p53-independent genome integrity checkpoint pathway mediated by Mre11 that protects against genome instability in breast cancer. Mre11 dysfunction in breast cancer models induces a genomic loss signature and vulnerability to PARP and ATR inhibitors.

## **Highlights**

- Mre11-mediated tumor suppression is p53 independent
- Mre11 suppresses DNA damage at oncogene-induced R loops
- Breast cancers with Mre11 dysfunction exhibit a genomic loss signature
- Mre11 dysfunction induces sensitivity to PARP and ATR inhibitors



## A P53-Independent DNA Damage Response Suppresses Oncogenic Proliferation and Genome Instability

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## **SUMMARY**

The Mre11-Rad50-Nbs1 complex is a DNA doublestrand break sensor that mediates a tumor-suppressive DNA damage response (DDR) in cells undergoing oncogenic stress, yet the mechanisms underlying this effect are poorly understood. Using a genetically inducible primary mammary epithelial cell model, we demonstrate that Mre11 suppresses proliferation and DNA damage induced by diverse oncogenic drivers through a p53-independent mechanism. Breast tumorigenesis models engineered to express a hypomorphic Mre11 allele exhibit increased levels of oncogene-induced DNA damage, R-loop accumulation, and chromosomal instability with a characteristic copy number loss phenotype. Mre11 complex dysfunction is identified in a subset of human triplenegative breast cancers and is associated with increased sensitivity to DNA-damaging therapy and inhibitors of ataxia telangiectasia and Rad3 related (ATR) and poly (ADP-ribose) polymerase (PARP). Thus, deficiencies in the Mre11-dependent DDR drive proliferation and genome instability patterns in p53-deficient breast cancers and represent an opportunity for therapeutic exploitation.

## INTRODUCTION

Structural chromosomal instability (CIN) is a frequent hallmark of clinically aggressive cancers, such as triple-negative (estrogen receptor, progesterone receptor, and HER2-negative) breast cancer (TNBC) and high-grade serous ovarian cancer (HGSOC),

yet its etiology remains poorly understood (Cancer Genome Atlas, N. and Cancer Genome Atlas Network, 2012; Jiang et al., 2010; Jonkers et al., 2001). Cancers with high levels of structural CIN are characterized by nearly universal disruption of the p53 pathway and frequent genetic aberrations that drive a hyper-proliferation phenotype (e.g., amplification of *c-Myc* or *Cyclin E* and/or *Rb1* deletion). Cancers with high levels of CIN also have frequent perturbation of DNA damage response (DDR) pathway genes (Knijnenburg et al., 2018), although their relevance to genome instability and therapeutic sensitivity in p53-deficient cancers remains unclear.

Oncogene-induced hyper-proliferation stimulates DNA replication stress (Bartkova et al., 2005, 2006; Di Micco et al., 2006; Halazonetis et al., 2008), resulting in accumulation of single- and double-strand breaks (DSBs) during S phase (Gaillard et al., 2015; Hills and Diffley, 2014; Macheret and Halazonetis, 2015) and activation of the DDR. The etiology of oncogene-induced replication stress has been extensively studied. Proposed mechanisms include nucleotide depletion, oxidative stress, misregulated replication origin firing, re-replication, perturbed replication fork kinetics, and under-replicated genomic DNA (Gaillard et al., 2015; Hills and Diffley, 2014; Kotsantis et al., 2018). Recent evidence suggests that oncogene expression stimulates genome-wide activation of ectopic intragenic origins, which results in replication stress due to a higher rate of transcription replication conflicts (TRCs) (Macheret and Halazonetis, 2018). Furthermore, depletion of RNA-DNA hybrids (i.e., R-loops) by RNase H overexpression reduces the burden of oncogene-induced DSBs (Kotsantis et al., 2016). Despite these advances, the relevance of pathways that regulate R-loop-mediated genome instability in tumorigenesis models remains poorly understood.

The Mre11-Rad50-Nbs1 complex is a DSB sensor that lies at the nexus between DNA repair and DDRs. The Mre11 complex is critical for ataxia telangiectasia mutated (ATM) activation at

DSBs and downstream activation of G2/M and p53-dependent G1/S cell cycle checkpoints (Oh and Symington, 2018; Stracker and Petrini, 2011; Syed and Tainer, 2018). The nuclease and structural functions of the Mre11 complex promote the resection of DSBs to generate 3' overhangs, which are a prerequisite for homologous recombination (HR)-mediated repair and replication fork stability (Hashimoto et al., 2011; Lemaçon et al., 2017; Trenz et al., 2006). Although complete deficiency is lethal, hypomorphic alleles of Mre11 complex genes are causative for ataxia-telangiectasia-like disorder and Nijmegan breakage syndrome and have helped to establish critical roles for this pathway in suppression of replication-associated DSBs and response to exogenous clastogens. Recent findings also implicate the Mre11 complex and its closely associated nuclease, Sae2/ CtIP, in the resolution of R-loops in mammals (Chang et al., 2018; Makharashvili et al., 2018).

Accumulating evidence supports a tumor-suppressive function for the Mre11 complex. Individuals with rare germline variants in Mre11 complex genes are at elevated risk of developing breast cancer (Damiola et al., 2014; Heikkinen et al., 2006). Although somatic mutations in Mre11 complex genes are infrequent in cancer (~3%; Zehir et al., 2017), aberrantly reduced protein expression has been identified in subsets of bladder, colorectal, breast, and ovarian cancers (Bartkova et al., 2008; Brandt et al., 2017; Choudhury et al., 2010; Situ et al., 2019; Zaki et al., 2014). Mice expressing the hypomorphic Mre11<sup>ATLD1</sup> allele also exhibit accelerated tumorigenesis driven by p53 mutation (Theunissen et al., 2003), Chek2 deficiency (Stracker et al., 2008), and mammary-specific Her2 expression (Gupta et al., 2013). Her2-driven breast cancers engineered to express a hypomorphic Mre11 complex exhibit increased genome instability and greater metastatic potential (Gupta et al., 2013). The mechanisms for these diverse tumor-suppressive effects have not been elucidated and may entail ATM- and p53-independent pathways. Indeed, ATM-independent tumor suppression was recently demonstrated using mice with knockin of a hypomorphic Nbs1 allele (Nbs1<sup>4B</sup>) (Balestrini et al., 2016). Here, Mre11 complex dysfunction was associated with common fragile site instability and dramatically increased genomic instability in the induced lymphomas. Collectively, these observations highlight an important role for the Mre11 complex in tumor suppression and protection against genome instability. A greater understanding of these Mre11-dependent effects may have important implications for classification of human tumors and identification of potential therapeutic vulnerabilities.

The goal of this study was to characterize the effects of Mre11 complex dysfunction on oncogenic proliferation, DNA damage, and genome instability in p53-proficient and p53-deficient inducible breast cancer models. We use single-cell, whole-genome sequencing to characterize the effect of Mre11 dysfunction on the landscape of Her2-induced genomic instability in mammary epithelial hyperplasia. We further establish p53-independent effects of Mre11 in suppressing oncogene-induced cell proliferation, DNA damage, and R-loops induced by diverse oncogenic drivers. We characterize a murine model of  $Rb1^{-/-}Trp53^{-/-}$  breast cancers with Mre11 hypomorphism that reveals abundant expression of R-loops and a marked increase in genomic deletions relative to other types of chromo-

somal rearrangements. Finally, drug sensitivity analyses of engineered p53-deficient breast tumors with or without Mre11 dysfunction identify therapeutic vulnerabilities that may be clinically exploitable.

## RESULTS

### Early Induction of Chromosomal Instability by Oncogenic Stress in Primary Mammary Epithelial Cells

We previously demonstrated that breast cancers initiated by Her2/Neu expression (abbreviated here as "Her2") develop with shorter latency and increased levels of CIN in mice expressing a hypomorphic allele of Mre11 (Mre11<sup>ATLD1/ATLD1</sup>; abbreviated Mre11<sup>ATLD</sup>; Gupta et al., 2013). To investigate whether Mre11 hypomorphism alters CIN patterns during early stages of oncogenic hyperplasia, we established a system for propagating primary murine mammary epithelial cells (pMMECs) on lethally irradiated LA7 mammary stem cell feeder cells (Ehmann et al., 1984; Jechlinger et al., 2009). Consistent with our previously reported in vivo findings, we observed significantly greater Her2-induced proliferation in Mre11ATLD pMMECs relative to wild-type (WT) pMMECs (Figure 1A). Because oncogeneinduced CIN during preneoplasia is stochastic and precedes clonal expansion, analysis of CIN patterns requires a single cell-based approach. Accordingly, we performed low-depth (~0.2×), single-cell, whole-genome sequencing (scWGS) in WT and Mre11<sup>ATLD</sup> pMMECs 14 days after lentiviral transduction with EGFP alone or EGFP+Her2 (Figure 1A). Copy number alterations (CNAs) were detected at single-cell resolution by pooling mapped reads into genomic bins with average size of 25 kb, using a previously described analysis pipeline (Garvin et al., 2015; Martelotto et al., 2017). The CNA profiles of 24 single cells from each genotype (WT+EGFP, WT+Her2, Mre11ATLD+EGFP, and Mre11<sup>ATLD</sup>+Her2) are depicted as a clustered heatmap, with the total fraction of altered genome for each cell shown on the right as a bar graph (Figure 1B). Significantly, WT pMMECs expressing EGFP had very few CNAs, corroborating the low level of background signal associated with the scWGS methodology and analysis pipeline employed in this study. In contrast, there is a substantial increase in the fraction of genome altered per cell after both Her2 expression and Mre11 perturbation (Figure 1B, bar graph on right). Unsupervised clustering was also performed but did not reveal any clonally related individual cells (Figure 1B), which is consistent with the relatively short time period after transduction when the cells were analyzed. Although the majority of the observed CNAs are non-recurrent and thus are indicative of a sporadic etiology, there are some focal CNAs that were recurrently seen in multiple cells within the same genotype. An interesting example is a focal region in chromosome 1 (boxed region in Figure S1) that has increased ploidy in 4/24 WT+Her2 pMMECs and 5/24 Mre11<sup>ATLD</sup>+Her2 pMMECs, but not in the other genotypes analyzed. This minimal region of CNA overlap contains Parp1, which is instrumental for singlestrand break repair (Ray Chaudhuri and Nussenzweig, 2017) and is frequently overexpressed in human HER2+ breast cancers (Stanley et al., 2015). Furthermore, a recent study suggests that Parp1 activity reduces the rate of replication fork progression (Maya-Mendoza et al., 2018). Consistent with these prior



#### Figure 1. Oncogene Expression Rapidly Induces Copy Number Aberrations in Individual pMMECs

(A) Schematic representation of the single-cell, whole-genome sequencing process. WT or  $Mre11^{ATLD}$  pMMECs were transduced with EGFP or Her2-EGFP. Growth curves on an irradiated feeder layer are shown. The mean of three biological replicates is shown with error bars depicting the standard error of the mean (SEM). \*\*\*p < 0.001, calculated using a two-tailed t test on log transformed day 14 data. On day 14, fluorescence-activated cell sorting (FACS) was performed to sort single EGFP+ cells into a 96-well microtiter plate. These cells were then processed for sequencing as described in the methods.

(B) Heatmap showing copy number of individual segments for each cell. The individual cells are clustered by geometric distance, which does not reveal any clonal relationships. The histograms at the right show the total fraction of aberrant genome per cell. p values were calculated by two-tailed Mann-Whitney test. \*\*\*\*p < 0.0001; \*p < 0.05.

See also Figure S1.

studies, our findings are suggestive that *Parp1* copy number gains may confer a growth advantage in the setting of Her2-induced mammary preneoplasia.

## Mre11 Dysfunction Results in a Copy Number Loss Phenotype

We observed a significant increase in CNAs per cell after Her2 expression in both *WT* and *Mre11*<sup>*ATLD*</sup> pMMECs (Figure 2A), indicating that oncogenic stress stimulates CIN early in neoplasia and does not require DDR deficiency. In fact, we did not observe an increase in CNAs per cell in *Mre11*<sup>*ATLD*</sup> pMMECs expressing Her2, suggesting that oncogene expression is the major driver of CNA development in this model. Consistent

with a role for Mre11 in suppressing spontaneous genomic instability,  $Mre11^{ATLD}$  pMMECs expressing EGFP also had a modest increase in CNAs compared to the respective WT control (Figure 2A). We next evaluated the size distribution of observed CNAs in the different genotypes. Notably, the CNA size distributions in both EGFP and Her2-expressing  $Mre11^{ATLD}$  pMMECs were significantly larger than the CNA size distribution observed in WT+Her2 pMMECs (Figure 2B). This difference was even more striking when evaluating CNAs that give rise to genomic gains versus losses. Although the size distribution of CNA gains and losses ( $\leq$  5,000 kb) was equivalent in the WT pMMEC genotypes, we observed a highly significant (p < 0.0001) enrichment for larger size genomic loss CNAs



#### Figure 2. Mre11 Dysfunction Alters the Profile of Copy Number Aberrations to Favor Genomic Loss

(A) Scatterplot showing total aberrations per cell in the different genotypes of pMMECs (mean  $\pm$  SEM). \*\*p < 0.01; \*\*\*\*p < 0.001 by two-tailed Mann-Whitney test. (B) Violin plots depicting the size distribution of aberrant copy number regions in each cell type. The line represents the median value. \*\*\*p < 0.001; \*\*\*\*p < 0.0001 by two-tailed Mann-Whitney test.

(C) Scatterplot CNA segment size for deletions and insertions, demonstrating a bias toward genomic deletions in *Mre11<sup>ATLD</sup>* genotypes. Error bars are median ± 95% confidence level. \*\*\*\*p < 0.0001 by two-tailed Mann-Whitney test.

(D) Cumulative size distribution CNA deletions and insertions for WT+EGFP (green), WT+Her2 (purple), Mre11<sup>ATLD</sup>+EGFP (blue), and Mre11<sup>ATLD</sup>+Her2 (red) pMMECs.

(E) Breakpoint regions in  $Mre11^{ATLD}$ +Her2 cells are associated with the largest genes (>300 kb). Violin plots show an empirical null distribution of overlaps generated by shuffle permutation of the observed CNA data, as described in the methods. Black line in violin plots is the median value, and the black dot is the observed value. p values are approximated from the empirical null distribution. \*p = 0.03–0.05; \*\*p = 0.01–0.03; \*\*\*p < 0.01.

(F) Expression of Her2 increases the number of R-loop DNA-RNA hybrids, detected by S9.6 immunofluorescence staining of WT pMMECs transduced with EGFP control or Her2. Scatterplot of the number of S9.6 foci per nucleus in control and Her2-expressing cells is shown. On the right are representative images of S9.6 nuclear foci used to generate the scatterplot. \*\*\*\*p < 0.0001 using a two-tailed t test. Scale bar represents 5 μm. See also Figure S2.

in both EGFP and Her2-expressing Mre11<sup>ATLD</sup> pMMECs (Figure 2C). The enrichment for genomic loss CNAs in Mre11ATLD pMMECs is also visually evident in the scWGS heatmaps shown in Figure 1B. To better characterize the relationship between CNA size and the observed enrichment for genomic losses, we plotted the cumulative frequency distribution of CNA gains and losses according to individual aberration size (Figure 2D). This analysis demonstrated that the enrichment for genomic loss was evident for CNAs larger than 1,000 kb in Mre11<sup>ATLD</sup>+EGFP cells and for CNAs greater than 500 kb in Mre11<sup>ATLD</sup>+Her2 cells. In contrast, no enrichment for CNA losses was observed in WT+Her2 pMMECs. The enrichment for genomic losses in the setting of Mre11 hypomorphism is reminiscent of copy number losses that result from accumulation of under-replicated DNA (UR-DNA) in settings of elevated replication fork stress in other model organisms (Salim et al., 2017; Yarosh and Spradling, 2014). Thus, our findings are consistent with replication fork instability caused by Mre11 dysfunction in

the setting of oncogenic stress, resulting in an accumulation of UR-DNA. Furthermore, because copy number losses are one mechanism by which loss of heterozygosity (LOH) in cancer may arise, an enrichment for larger sized deletions in  $Mre11^{ATLD}$  pMMECs may be consistent with the high levels of LOH that are observed in human breast and ovarian cancers with homologous recombination deficiency (Wang et al., 2012).

## Chromosomal Aberrations Are Associated with Large Genes

Next, we investigated whether there were any distinguishing features of the genomic regions where the oncogene-induced CNAs were observed. The sparseness of our scWGS data precluded single-base-pair resolution of CNA breakpoints. In contrast, we approximated the genomic region of the chromosomal aberration breakpoint as contained within the two genomic bins that span the transition in copy number (Figure S2A; STAR Methods). We next evaluated whether these

breakpoint-containing regions were enriched in previously annotated genomic features and used a shuffle permutation of the observed CNA profiles for each of the genotypes as a means of establishing statistical significance of the enrichment (Supplemental Methods). We did not observe any significant enrichment or depletion of satellite, simple repeat, long terminal repeats (LTRs), long interspersed nuclear elements (LINEs), short interspersed nuclear elements (SINEs), or CpG island sequences among the breakpoint regions for any of the genotypes (Figure S2B). In contrast, there was significant enrichment of breakpoint-containing regions with genes. Genes were further subdivided according to size into four bins of <50 kb, 50-150 kb, 150-300 kb, and >300 kb. There was a significant enrichment of chromosomal breakpoint regions within large genes (>300 kb) relative to random chance for Her2-expressing Mre11<sup>ATLD</sup> pMMECs (Figure 2E). In contrast, chromosomal breakpoint regions in WT+Her2 pMMECs were relatively enriched in moderately sized genes between 50 kb and 300 kb. Large genes are known to take longer than one cell cycle to transcribe and have been associated with R-loops, chromosomal fragile sites, and recurrent deletions in many cancers (Glover et al., 2017; Helmrich et al., 2006, 2011; Le Tallec et al., 2013; Wilson et al., 2015). Recent work has also demonstrated an increase in transcription-replication conflicts (TRCs) upon oncogene expression (Macheret and Halazonetis, 2018), which may be due to greater transcriptional activity and/or persistence of R-loops (Kotsantis et al., 2016). We thereby assessed global R-loop levels in our pMMEC model using immunofluorescence with the monoclonal S9.6 antibody and observed a significant increase in R-loops 3 days after transduction with lentivirus expressing Her2-EGFP in pMMECs. relative to control pMMECs transduced with EGFP alone (Figure 2F). Collectively, these findings indicate that oncogenic stress in early mammary neoplasia stimulates R-loop expression and CNAs enriched in large genes, which in the setting of Mre11 dysfunction becomes strongly enriched in a copy number loss phenotype.

## Mre11 Suppresses Oncogenic Proliferation Independently of p53 and ATM

We next investigated the effect of Mre11 hypomorphism in transgenic mammary hyperplasia models induced by *c-Myc* (Myc) overexpression, Rb1 deletion, and/or Trp53 deficiency-genetic aberrations that are prevalent in human TNBCs and basal-like breast cancer (Cancer Genome Atlas, N. and Cancer Genome Atlas Network, 2012). Because the Mre11ATLD allele is a premature stop codon in the C-terminal region of Mre11, we hypothesized that a comparable hypomorphic allele could be generated using CRISPR/Cas9 gene targeting in pMMECs, as has previously been demonstrated in murine embryonic fibroblast cell lines (Wyatt et al., 2016). pMMECs isolated from Rosa26<sup>LSL-Cas9-EGFP</sup> (abbreviated R26<sup>Cas9</sup>) mice (Platt et al., 2014) were transduced with lentivirus expressing Cre recombinase and single guide RNA (sgRNA) targeting the C-terminal region of Mre11 (Cre-sgMre11) or a control non-coding region on chromosome 2 (Cre-sgControl; Figures 3A and 3B). Effective target site mutagenesis was confirmed by Sanger sequencing of 10 cloned amplicons each from R26<sup>Cas9</sup> pMMECs transduced with Cre-sgControl or Cre-sgMre11 (Figure S3A). CRISPRdirected mutations corresponding to the expressed sgRNA were identified, which resulted in an Mre11 frameshift mutation in all 10 clones analyzed from sgMre11-transduced pMMECs (Figure S3A). Furthermore, pMMECs transduced with CresgMre11 had reduced expression of Mre11 (Figure S3B), phenocopying the destabilizing effect of the *Mre11*<sup>ATLD</sup> allele on Mre11 complex protein stability (Theunissen et al., 2003) These findings validated the pMMEC platform to evaluate phenotypes associated with oncogene expression and/or Mre11 hypomorphism.

We next interbred the  $R26^{Cas9}$  mice with transgenic mice containing a Cre-inducible Myc overexpression cassette also at the Rosa26 locus ( $Rosa26^{LSL-MycOE-hCD2}$  or  $R26^{Myc}$ ) to generate combination transgenic  $R26^{Myc/Cas9}$  mice (Figures 3A and 3B). pMMECs derived from these mice were transduced with lentivirus expressing Cre recombinase and either sgControl or sgMre11, revealing greater Myc-induced proliferation in cells with a hypomorphic Mre11 complex (Figure 3C). Notably, expression of Cre-sgMre11 in  $R26^{Cas9/Cas9}$  pMMECs did not result in a substantial increase in proliferation, indicating that this anti-proliferative effect of Mre11 is oncogene specific.

To investigate whether suppression of oncogene-induced proliferation by the Mre11 complex is p53 dependent, we generated R26<sup>Myc/Cas9</sup>; Trp53<sup>fl/fl</sup> mice. Cre recombinase expression in pMMECs isolated from these mice induces deletion of Trp53 exon 2 in conjunction with activated expression of Cas9 and Myc (Figures 3B and S3). Mre11 complex hypomorphism induced by expressing Cre recombinase and sgMre11 resulted in significantly increased proliferation relative to the R26<sup>Myc/Cas9</sup>; Trp53<sup>fi/fl</sup> pMMECs transduced with Cre and sgControl (Figure 3D). We also generated and analyzed R26<sup>Cas9</sup>Trp53<sup>fl/fl</sup>Rb1<sup>fl/fl</sup> mice, representing another model of human TNBC (Figure 3B; Cancer Genome Atlas, N. and Cancer Genome Atlas Network, 2012; Jiang et al., 2010; Jonkers et al., 2001). Once again. R26<sup>Cas9</sup>Trp53<sup>fl/fl</sup>Rb1<sup>fl/fl</sup> pMMECs transduced with Cre recombinase and sgMre11 demonstrated significantly increased proliferation relative to cells expressing sgControl (Figure 3D). These findings strongly argue that Mre11-mediated suppression of oncogenic proliferation is p53 independent. Given that the Mre11 complex has both ATM-dependent and independent effects in the response to DNA damage and replication stress (Rein and Stracker, 2014), we sought to determine the ATM dependency of Mre11-mediated suppression of oncogenic proliferation. R26<sup>Cas9</sup>Trp53<sup>fl/fl</sup>Rb1<sup>fl/fl</sup> pMMECs were transduced with either sgControl or sgMre11 and then treated with ATM inhibitor Ku55933 or vehicle control. ATM inhibition (Figure S3D) did not increase proliferation of Rb1-/-Trp53-/pMMECs (Figure 3E). A possible explanation for these observations is that growth suppression by ATM is p53 dependent, consistent with emerging clinical evidence that ATM and TP53 may be epistatic in breast cancer (Weigelt et al., 2018). Conversely, Mre11 hypomorphism stimulated oncogenic proliferation irrespective of ATM inhibition (Figure 3E). These observations collectively indicate that Mre11-mediated suppression of oncogenic proliferation is both p53 and ATM independent.



## Figure 3. Mre11 Suppresses Oncogenic Proliferation in pMMECs Independently of *Trp53* and ATM

(A) Graphic representing how pMMECs are harvested from the mice and manipulated *in vitro* to assess growth rates.

(B) Description of compound transgenic mice used for pMMEC experiments and resulting genotypes after introduction of Cre recombinase.

(C) pMMEC growth curves examining the effect of Myc overexpression and/or *Mre11* hypomorphic mutation. Cell counts are normalized to their respective day 0 counts.

(D) Significant effect of *Mre11* mutation on oncogenic growth induced by Myc overexpression or *Rb1* deletion in p53-deficient pMMECs. Cell counts are normalized to their respective day 0 counts.

(E) ATM inhibitor Ku55933 does not phenocopy the growth-stimulating effects of *Mre11* mutation in  $Rb1^{-/-}Trp53^{-/-}$  pMMECs. Statistical significance in (C)–(E) was determined by two-tailed t test on log-transformed data comparing day 12. The p values were adjusted for multiple comparisons by the method of Holm-Sidak. Data are represented as mean  $\pm$  SEM.

See also Figure S3.

## Mre11 Suppresses Oncogene-Induced R-Loops and DNA Damage

The Mre11 complex suppresses the accumulation of spontaneous, replication-associated DSBs (Oh and Symington, 2018). In contrast, some studies have demonstrated that Mre11 nuclease activity can also promote destabilization of stalled replication forks (Malacaria et al., 2019; Schlacher et al., 2011). To determine whether Mre11 promotes or suppresses oncogene-induced DSBs in pMMECs, we measured yH2AX foci, 53BP1 foci, and neutral COMET tails (Figures 4A, 4B, and S4A). Mre11 hypomorphism induced by CRISPR/Cas9-mediated targeting of Mre11 resulted in significantly higher levels of nuclear DSBs in both p53-proficient and p53-deficient pMMEC models. Similarly, Mre11 hypomorphism increased the levels of single-stranded DNA damage, as measured by pRPA2 immunofluorescence and alkaline COMET assay (Figures 4B and S4B). The increase in pRPA2 foci was most significant in cells that were no longer in S phase (Figure S4C), indicative of a deficiency in resolution/repair of replication stress in Mre11 mutant pMMECs. These findings argue that Mre11 suppresses the accumulation of oncogenic DNA damage (single-stranded breaks and DSBs) in both p53-proficient and p53-deficient models of mammary preneoplasia.

Due to our prior finding that oncogenic stress in the setting of Mre11 hypomorphism induces copy number aberrations in genomic regions known to have high levels of R-loops (see Fig-

orescence. As previously demonstrated with Her2 expression (see Figure 2F), we again found that oncogenic stress stimulates R-loop foci in both R26<sup>Myc</sup> and R26<sup>Cas9</sup>Rb1<sup>-/-</sup>Trp53<sup>-/-</sup> pMMECs (Figure 4C). Significantly, we also found R-loop levels increased substantially further upon Mre11 dysfunction, in both p53-proficient and p53-deficient models (Figure 4C). These observations are consistent with a recent study implicating the Mre11 complex as a critical mediator of transcription-replication conflicts in yeast and mammals (Chang et al., 2019). To establish whether the increase in R-loops mediates the increased levels of oncogenic DNA damage in pMMECs with hypomorphic Mre11, we modified our Cre-sgControl/sgMre11 lentiviral construct to also co-overexpress RNase H1 (Figure S4D). As expected, co-overexpression of RNase H1 abolished the increase in R-loops observed in the setting of hypomorphic Mre11 (Figure 4D). RNase H1 expression also rescued the increase in oncogenic DSBs and single-stranded DNA breaks induced by Mre11 dysfunction (Figures 4E and 4F). These findings were also confirmed by neutral and alkaline COMET assays (Figures S4E and S4F). Thus, an aberrant accumulation of R-loops in pMMECs with Mre11 dysfunction is necessary for the observed increase in oncogenic DNA damage. Collectively, these findings support a model wherein Mre11 mitigates DNA damage incurred at sites of oncogeneinduced transcription-replication conflicts (TRCs) (see Figure 7).

ure 2E), we assessed global R-loop levels using S9.6 immunoflu-



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## Mre11-Deficient Mammary Tumors Have Increased R-Loops and a Genomic Loss Phenotype

To extend and validate these results *in vivo*,  $R26^{Cas9}Rb1^{fl/}$  $f^{f}Trp53^{fl/fl}$  and  $R26^{Myc/Cas9}$  female mice were administered intraductal injections of lentivirus expressing Cre-sgControl or Cre-sgMre11 into mammary gland 4. Two weeks after injection, the mice were euthanized and gland 4 (injected) and gland 5 (uninjected control) were processed for histopathology. There was a qualitative (Figure 5A) and quantitative (Figure 5B) increase in mammary epithelial hyperplasia after intraductal injection with Cre-sgMre11, relative to glands injected with Cre-sgControl. Thus, Mre11 suppresses oncogenic hyperplasia *in vivo* in response to diverse oncogenic drivers and in the setting of induced p53 deficiency.

Cohorts of Cre-sgControl and Cre-sgMre11-injected R26<sup>Cas9</sup>Rb1<sup>fl/fl</sup>Trp53<sup>fl/fl</sup> mice were monitored for mammary tumor development. Tumor initiation frequency was higher in glands injected with Cre-sgMre11 (~70%) versus Cre-sgControl  $(\sim 50\%)$ , and there was a trend toward a shorter latency period (Figure 5C). Tumor-free survival rates, however, did not achieve statistical significance (p = 0.052), suggesting that tumor latency in the R26<sup>Cas9</sup>Rb1<sup>fl/fl</sup>Trp53<sup>fl/fl</sup> model may also be dependent on the stochastic acquisition of secondary genetic events. Mammary tumor lines were established from the induced tumors, and as expected, expression of the entire Mre11 complex (Mre11, Rad50, and Nbs1) was substantially reduced in tumors derived from mice injected with Cre-sgMre11 (Figure 5D). Consistent with our observations with pMMECs, sgMre11 mammary tumor lines had a higher proliferation rate (Figure 5E) and significantly higher levels of R-loops (Figure 5F) relative to sgControl mammary tumor lines.

We also performed matched tumor-normal whole-genome sequencing (~25× mean depth of coverage) of three Cre-sgControl and three Cre-sgMre11-induced mammary tumors in *R26<sup>Cas9</sup>Rb1<sup>fl/fl</sup>Trp53<sup>fl/fl</sup>* mice. Site-specific mutagenesis of the Mre11 locus was confirmed in all of the Cre-sgMre11 mammary tumors (data not shown). Single-nucleotide variant mutation signatures were not statistically different between the Cre-sgControl and Cre-sgMre11 tumors (Figure S5). Notably, there was no increase in COSMIC signature 3, which has been associated with homologous-recombination-deficient cancers (Alexandrov et al., 2013; Riaz et al., 2017; Rosenthal et al., 2016). A similar lack of COSMIC signature 3 enrichment has been

observed in human breast cancers with ATM deficiency (Weigelt et al., 2018). Structural variants (SVs) were observed in both sgControl and Cre-sgMre11  $Rb1^{-/-}Trp53^{-/-}$  mammary tumors (Figure 5G). Significantly, the observed SVs in Mre11 hypomorphic tumors were highly enriched for deletions relative to control tumors (Figures 5G and 5H). Thus, the genomic loss phenotype observed in early oncogenic hyperplasia induced in the setting of Mre11 hypomorphism (see Figures 2C and 2D) is also preserved upon tumorigenesis in a distinct p53-deficient breast cancer model. These observations indicate that Mre11 complex dysfunction is causative for a genomic scar phenotype consisting of an enrichment for copy number losses/genomic deletions.

## Breast Cancers with Mre11 Dysfunction Are Hypersensitive to DNA-Damaging Therapy and Inhibitors of ATR and PARP

Our finding that the Mre11 complex has p53-independent, tumor-suppressive functions leads to a hypothesis that Mre11 complex dysfunction may be selected for in p53-deficient breast cancers. Indeed, a prior study has demonstrated downregulation of Mre11 complex proteins in TNBC (Bartkova et al., 2008), 90% of which harbor p53 pathway deficiency (Cancer Genome Atlas, N. and Cancer Genome Atlas Network, 2012). To validate these findings, we performed immunohistochemistry for Mre11 and Nbs1 on a tissue microarray of 254 early-stage TNBCs that were treated at Memorial Sloan Kettering Cancer Center. Ninety-three percent of the patients received adjuvant chemotherapy. We identified 23 samples (9%) with less than 10% of tumor nuclei staining for Mre11 or Nbs1 protein, while infiltrating stromal cells retained normal expression (Figure 6A). Notably, we found that all tumors with low Mre11 expression also had low Nbs1 expression, consistent with underexpression or destabilization of the Mre11 complex. The lack of immunoreactivity for Mre11 and Nbs1 was also confirmed on whole tissue sections from the original tumor blocks. These "Mre11 complex low" cancers presented with similar T- and N- stage at presentation relative to cancers with normal Mre11 complex expression (Figure S6). Notably, patients with Mre11 complex low TNBC had a significantly better breast-cancer-specific survival compared to the remainder of TNBCs with normal Mre11 complex expression (Figure 6B).

Because disease-free survival of patients with TNBC is highly associated with improved chemotherapeutic responses

### Figure 4. Mre11 Suppresses Oncogene-Induced DNA Damage and R-Loops in p53-Proficient and p53-Deficient Models

(A) Mre11 suppresses oncogene-induced  $\gamma$ H2AX foci formation in both p53-proficient and p53-deficient pMMECs. Bar graphs show quantification of the percent of nuclei containing  $\geq$ 5  $\gamma$ H2AX foci in the different genotypes. Representative images (right) of the nuclei containing  $\gamma$ H2AX foci are shown. White bar indicates 5  $\mu$ m.

(D) S9.6 (R-loop) foci after RNase H1 overexpression in R26<sup>Cas9</sup>Rb1<sup>n///</sup>Trp53<sup>n///</sup> pMMECs transduced with Cre-sgControl or Cre-sgMre11.

(E and F) Additionally, RNase H1 overexpression counteracts the increase in (E)  $\gamma$ H2AX and (F) p-RPA2 foci seen in Mre11 hypomorphic R26<sup>Cas9</sup>Rb1<sup>n/n</sup>Trp53<sup>n/n</sup> pMMECs.

\*\*\*p < 0.001; \*\*\*\*p < 0.0001. p values are calculated using a two-tailed Mann-Whitney test. See also Figure S4.

<sup>(</sup>B) Bar graphs depicting the fold change in tail DNA percent for both alkaline (left) and neutral (right) COMET assays in pMMECs with the genotypes shown post-Cre-sgRNA transduction. Representative images of alkaline and neutral COMETs in *R26<sup>Cas9</sup>*+sgControl, *R26<sup>Cas9/Myc</sup>*+sgControl, and *R26<sup>Cas9/Myc</sup>*+sgMre11 pMMECs are shown. Data are represented as mean ± SEM.

<sup>(</sup>C) Mre11 suppresses oncogene-induced R-loop formation independently of *Trp53*. Scatterplot shows a quantification of the nuclear S9.6 foci in pMMECs from the genetic backgrounds shown after transduction with Cre-sgControl versus Cre-sgMre11. Representative images (right) of the nuclei containing S9.6 foci are shown. White bar indicates 5 µm.



## Figure 5. Cell Proliferation, R-Loops, and Genomic Loss Signature Are Elevated in *Rb1<sup>-/-</sup>Trp53<sup>-/-</sup>* Mammary Tumors Engineered with Mre11 Hypomorphism

(A) Representative images of mammary hyperplasia 2 weeks after intraductal injection of R26<sup>Cas9</sup>Rb1<sup>#/#</sup>Trp53<sup>#/#</sup> or R26<sup>Cas9/Myc</sup> mice with Cre-sgControl or Cre-sgMre11-expressing lentivirus.

(B) Quantification of mammary hyperplasia in *R26<sup>Cas9</sup>Rb1<sup>fl/fl</sup>Trp53<sup>fl/fl</sup>* mice as shown in (A). The fifth, non-injected, mammary gland serves as a negative control. Data are represented as mean ± SEM. Significance was determined using two-tailed Mann-Whitney test.

(C) Kaplan-Meier tumor-free survival plot of  $R26^{Cas9}Rb1^{n/n}Trp53^{n/n}$  mice after mammary intraductal injection of either Cre-sgControl or Cre-sgMre11 lentivirus. p value was calculated using a two-tailed Gehan-Breslow-Wilcoxon test.

(D) Mre11, Rad50, and Nbs1 protein levels in tumors from mice in (C), validating protein destabilization induced by CRISPR/Cas9-mediated Mre11 mutagenesis. (legend continued on next page)

(Cortazar et al., 2014), we hypothesized that Mre11 hypomorphism may promote hypersensitivity to DNA-damaging therapeutics in breast cancer. To directly address this possibility, we used breast tumor lines derived from R26<sup>Cas9</sup>Trp53<sup>fl/fl</sup>Rb1<sup>fl/fl</sup> mice injected with Cre-sgControl or Cre-sgMre11 to evaluate the effect of Mre11 hypomorphism on sensitivity to DNA-damaging therapeutics. We found that breast cancer cells expressing hypomorphic Mre11 were hypersensitive to several DNA-damaging therapeutics commonly used to treat breast cancer, but not to the anti-microtubule agent Taxol (Figure 6C). We also evaluated the sensitivity of these mammary tumor lines to inhibitors of ATR (VE-821) and PARP (BMN-673), particularly because both of these pathways are components of the replication stress response and the ATR pathway has been implicated in resolution of R-loops (Cristini et al., 2018; Hodroj et al., 2017; Nguyen et al., 2018). We observed significantly greater hypersensitivity of Mre11 hypomorphic Rb1-/-Trp53-/- breast tumor lines to both VE-821 and BMN-673, relative to control Rb1<sup>-/-</sup>Trp53<sup>-/-</sup> breast tumor lines (Figure 6D). Collectively, these findings indicate that Mre11 complex dysfunction is evident in a subset of TNBC patients and associated with improved clinical outcomes after DNA-directed cancer therapy.

## DISCUSSION

Seminal studies (Bartkova et al., 2005, 2006; Di Micco et al., 2006; Gupta et al., 2013; Halazonetis et al., 2008) have described an Mre11-dependent DDR activated by oncogeneinduced replication stress in preneoplasia that mediates physiologically significant tumor suppression. Some of these studies have claimed that the oncogene-induced DDR exerts these effects primarily through p53 activation. To our knowledge, ours is the first study to directly assess the effects of DDR perturbation in p53-deficient models of oncogenic preneoplasia. We demonstrate that Mre11-mediated suppression of oncogenic proliferation, DNA damage, and genome instability are p53 independent. Thus, our findings challenge current models of the oncogene-induced DDR and its relation to p53: rather than being epistatic to p53, the Mre11-dependent DDR mediates p53independent effects that regulate oncogenic phenotypes, including proliferation, genome instability, and therapeutic sensitivity.

Single-cell, whole-genome sequencing of premalignant mammary epithelial cells revealed important clues into the critical functions of the Mre11-dependent DDR in response to oncogenic stress. We found that oncogene expression was sufficient to rapidly induce widespread CNAs in otherwise completely normal primary mammary epithelial cells, independently of Mre11 status. Recent genomic analyses of human breast preneoplasia have also suggested an early origin of chromosomal instability, which does not seem to require DDR deficiency (Martelotto et al., 2017; Rane et al., 2015). In the setting of Mre11 dysfunction, we observed a striking enrichment in genomic losses rather than gains, which was further exacerbated by oncogene expression. A similar enrichment in genomic deletions was seen in Rb1-/-Trp53-/- mammary tumors engineered to express hypomorphic Mre11 alleles, indicating that this genomic scar signature of Mre11 dysfunction persists during tumorigenesis. A possible explanation for the observed enrichment in genomic losses may be the accumulation of UR-DNA in oncogene-expressing cells with Mre11 dysfunction. Coupled with the observation that Mre11 dysfunction results in higher levels of unrepaired oncogene-induced DSBs, the increased levels of UR-DNA in this setting may be due to a higher rate of replication fork collapse (Figure 7).

The location of oncogene-induced chromosomal aberrations in the setting of Mre11 dysfunction were non-randomly distributed and significantly enriched in large genes (>300 kb). These genomic regions are known to be occupied by R-loops, depleted in replication origins, and associated with chromosomal fragility (Helmrich et al., 2011). Thus, our findings reinforce prior reports of fragile site instability induced by oncogenic stress (Miron et al., 2015) and complement the recent demonstration of oncogeneinduced TRCs that stimulate intragenic dormant origin firing (Macheret and Halazonetis, 2018). Our results suggest that Mre11 promotes replication fork stability at oncogene-induced TRCs (Figure 7). R-loops can either be persistent structures that are stabilized by DNA secondary structure or transient intermediates of gene transcription. Further work to determine whether one of these types of R-loops is a greater contributor to oncogeneinduced CIN is warranted. We observed a striking increase in R-loop expression in mammary hyperplasias and tumors with Mre11 dysfunction. Mitigation of R-loops by RNase H1 expression in Mre11 hypomorphic cells diminished oncogenic DNA damage. A causative role for R-loops in the etiology of oncogenic DNA damage can be explained in the setting of replication fork collisions, which can give rise to fork uncoupling and/or collapse. Thus, our findings support a model wherein the Mre11 complex has critical functions in mitigating DNA damage at TRCs in oncogene-expressing cells (Figure 7). Similar conclusions were drawn in a recent analysis of yeast and human

<sup>(</sup>H) Analysis of structural aberrations identified from tumor-normal WGS from three *WT* and three *Mre11* hypomorphic mammary tumors from the  $R26^{Cas9}Rb1^{fl/t}Trp53^{fl/t}$  model. Deletions (>50 bp) are highly over-represented in Mre11 hypomorphic  $Rb1^{-/-}Trp53^{-/-}$  mammary tumors. Data are represented as mean  $\pm$  SEM. Q value was estimated using a two-tailed t test with false discovery rate correction using Graphpad Prism v8. All other comparisons were not statistically significant (i.e., Q < 0.05).



<sup>(</sup>E) Cells from the sgMre11 tumors grow faster *in vitro* than cells from the sgControl tumors. Three independent tumor lines are shown for each genotype. Growth curves were acquired using the Incucyte Live Cell Analysis system. Data are represented as mean ± SEM.

<sup>(</sup>F) Cells from sgMre11-derived tumors have more R-loops than cells from sgControl tumors. Three independent tumor lines of each genotype were evaluated by immunofluorescence using the S9.6 antibody. Data are represented as mean ± standard deviation. Significance was determined using a two-tailed Mann-Whitney test.

<sup>(</sup>G) Representative circos plot of structural variations identified by whole-genome sequencing of WT or Mre11 hypomorphic mammary tumors derived from the R26<sup>Cas9</sup>Rb1<sup>fl/fl</sup>Trp53<sup>fl/fl</sup> model. Purple lines indicate translocations between different chromosomes. Blue lines indicate intra-chromosomal rearrangements.



Figure 6. Favorable Clinical Outcomes and Therapeutic Vulnerabilities of Mre11 Hypomorphic p53-Deficient Breast Cancers

(A) Immunohistochemistry for (1) Mre11 and (2) Nbs1 performed on tissue microarrays of primary triple-negative breast cancers (n = 254). In both cases (1 and 2), the left panel represents an example of a "normal" expressor and the right case represents an example of a "low" expressing tumor.

(B) Kaplan-Meier breast-cancer-specific survival of patients with TNBC that had normal expression of Mre11 and Nbs1 or patients with TNBC that had low expression of both Mre11 and Nbs1 (i.e., Mre11 complex low). p value was calculated by a two-tailed log rank test.

(C) *In vitro* sensitivity of sgControl and sgMre11 *Rb1<sup>-/-</sup>Trp53<sup>-/-</sup>* mammary tumor lines treated with the indicated doses of cisplatin, doxorubicin, camptothecin, and Taxol. Percent confluence was measured using an Incucyte Live-Cell Analysis System 7 days after drug exposure.

(D) Similar drug sensitivity assays using the ATR inhibitor VE-821 and PARP inhibitor BMN-673. The data shown represent the average of at least two independent cell lines for each genotype with three replicates each. Significance was determined using a two-tailed t test at the highest drug dosage. Data are represented as mean  $\pm$  SEM.

See also Figure S6.

cell line models (Chang et al., 2019). A more detailed examination of how the Mre11 complex promotes genome integrity at TRCs—particularly in consideration of head-on versus codirectional collisions (Hamperl et al., 2017)—may provide additional clues regarding the etiology of genome instability during tumorigenesis.

The observation that Mre11 mutant cells have increased proliferation despite elevated levels of unrepaired DNA damage points to an oncogene-induced DNA damage checkpoint that is mediated by Mre11. The nature of the Mre11-mediated checkpoint in response to oncogenic stress remains to be further elucidated. Our finding that this checkpoint remains operative in p53-deficient cells suggests a possible role for an Mre11mediated G2/M checkpoint in response to oncogenic stress. We found that ATM inhibition, however, did not phenocopy the effect of Mre11 hypomorphism in promoting oncogenic proliferation, suggesting that non-canonical mechanisms may also be involved.

Reduced expression of Mre11 complex proteins has been identified in a variety of human malignancies (Bartkova et al., 2008; Brandt et al., 2017). Our data provide direct evidence supporting a role for Mre11 in mediating resistance to DNA-damaging therapy as well as PARP and ATR inhibitors in TNBCs. Indeed, low expression of Mre11 and Nbs1 was



Figure 7. Model Depicting Genome Stabilizing Functions of Mre11 at Oncogene-Induced Transcription-Replication Conflicts Mre11 dysfunction during oncogenic breast neoplasia results in accumulation of R-loops, replication-associated DSBs, under-replicated DNA, an enrichment of genomic deletions, and uncontrolled proliferation. See the Discussion for further details.

associated with improved clinical outcomes in a cohort of TNBC patients who were treated with adjuvant chemotherapy. The finding that these cancers may also be hypersensitive to targeted DDR pathway inhibitors represents opportunities for targeted therapy that may reduce the need for highly toxic combination chemotherapeutic regimens in a subset of TNBC patients.

The Mre11-dependent DNA damage response is evolutionarily more primitive than p53 and functions as a mechanism to preserve genome integrity in response to endogenous and exogenous genotoxic stresses. The observation that DDR gene perturbations are prevalent across many human cancer types (Knijnenburg et al., 2018) suggests that aspects of the DDR are also tumor suppressive. Our findings reveal a p53-independent checkpoint function of the Mre11 complex that suppresses genome instability and uncontrolled proliferation induced by oncogenic mutations. Disruption of this genome integrity checkpoint may drive the catastrophic loss of chromosomal stability that is observed in a variety of clinically aggressive cancers. Elucidating additional mediators of this tumor-suppressive pathway may reveal new opportunities for personalized therapy of human cancers, including those with p53 deficiency.

## **STAR**\***METHODS**

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## SUPPLEMENTAL INFORMATION

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### **AUTHOR CONTRIBUTIONS**

Conceptualization, G.P.G. and J.H.J.P.; Methodology, G.P.G.; Software, D.A.S. and L.E.M.; Formal Analysis, D.A.S. and N.U.R.; Investigation, K.D.F.-S., D.A.S., R.J.K., L.G.M., and Y.H.W.; Resources, G.P.G., A.Y.H., S.N.P., J.S.P., J.S.R.-F., and J.H.J.P.; Writing – Original Draft, G.P.G., K.D.F.-S. and D.A.S.; Writing – Review & Editing, all authors; Funding Acquisition, G.P.G., K.D.F.-S., and J.H.J.P.; Supervision, G.P.G.

#### **DECLARATION OF INTERESTS**

G.P.G. has ownership interest (including patents) in and is a consultant/advisory board member for Naveris, Inc., outside the scope of the present study. J.S.R.-F. reports personal/consultancy fees from VolitionRx, Page.AI, Goldman Sachs, Grail, Ventana Medical Systems, Invicro, Roche Diagnostics, and Genentech, outside the scope of the present study. J.H.J.P. is a consultant for Ideaya Biosciences, Novus Biologicals, and Atropos Therapeutics, outside the scope of the present study.

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## **STAR\*METHODS**

## **KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Chicken anti-Mouse IgG (H+L) Cross- Adsorbed Secondary Antibody, Alexa Fluor 488, (1:10,000 for IF)	Thermo Fisher Scientific	Cat# A-21200, RRID:AB_2535786
Chicken anti-Rabbit IgG (H+L) Cross- Adsorbed Secondary Antibody, Alexa Fluor 488, (1:10,000 for IF)	Thermo Fisher Scientific	Cat# A-21441, RRID:AB_2535859
Mouse Anti-beta-Actin Monoclonal Antibody, Unconjugated, Clone AC-15 (1:10,000 for WB)	Sigma-Aldrich	Cat# A1978, RRID:AB_476692
Rabbit Anti-Phospho-p53 (Ser15) Antibody (1:1000 for WB)	Cell Signaling Technology	Cat# 9284, RRID:AB_331464
Mouse Anti-DNA-RNA Hybrid [S9.6] Antibody (1:500 for IF)	Kerafast	Cat# ENH001, RRID:AB_2687463
Mouse Anti-Human CD2 Monoclonal Antibody, Clone LT2 (1:10 for Flow)	Miltenyi Biotec	Cat# 130-091-115, RRID:AB_244321
Rabbit Anti-RPA2, (phospho Ser4, Ser8) Polyclonal, Unconjugated antibody (1:500 for IF)	Novus	Cat# NBP1-23017, RRID:AB_1726226
Rabbit Anti-53BP1 Polyclonal Antibody (1:500 for IF)	Bethyl	Cat# A300-272A, RRID:AB_185520
Goat anti-Hamster IgG Secondary Antibody, HRP (1:10,000 for WB)	Thermo Fisher Scientific	Cat# PA1-29626, RRID:AB_10985385
Goat Anti-Rabbit IgG, IRDye® 680LT Conjugated antibody (1:10,000 for WB)	LI-COR Biosciences	Cat# 926-68021, RRID:AB_10706309
Goat Anti-Mouse IgG, IRDye® 800CW Conjugated antibody (1:10,000 for WB)	LI-COR Biosciences	Cat# 926-32210, RRID:AB_621842
Mouse Anti-p53 (1C12) mAb Antibody (1:1000 for WB)	Cell Signaling Technology	Cat# 2524, RRID:AB_331743
Rabbit Anti-phosphorylated Histone H2AX (γ-H2AX) Polyclonal Antibody (1:500 for IF)	Trevigen	Cat# 4418-APC-100
Rabbit Anti-KAP1 Polyclonal Antibody, Unconjugated (1:500 for WB)	Abcam	Cat# ab10484, RRID:AB_297223
Rabbit Anti-KAP1 (phospho S824) antibody, (1:500 for WB)	Abcam	Cat# ab70369, RRID:AB_1209417
Mouse Anti-NBS1 Monoclonal Antibody, Unconjugated (1:10,000 for WB)	Novus	Cat# NB100-221, RRID:AB_10001212
Rabbit Anti-RAD50 Polyclonal Antibody, Unconjugated (1:10,000 for WB)	Novus	Cat# NBP2-20054
Armenian Hamster Anti-Mre11 Monoclonal Antibody, Unconjugated (1:500 for WB)	Novus	Cat# NBP2-59677
Mouse Anti-RNaseH1 (H-4) Monoclonal Antibody, Unconjugated (1:1000 for WB)	Santa Cruz Biotechnology	Cat# sc-376326, RRID:AB_10987730
Bacterial and Virus Strains		
Endura DUOs Electrocompetent Cells	Lucigen	60242-2
Chemicals, Peptides, and Recombinant Proteins		
Liberase Blendzyme 2	Roche	11988425001
Cultrex 3D-Culture Matrix	Irevigen	3447-020-01
EpiCult-B Mouse Medium Kit	Stem Cell Technologies	05610

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
HuMEC Ready Medium (1X)	GIBCO	12752010
Dulbecco's Modified Eagle Medium:	GIBCO	11320082
	GIBCO	15630080
		10000000 407700 50MC
Denicillin Streptomycin (10,000 L1/mL)		407709-5000G
	Marthinsten	15140122
	Chara Call Taskaslasias	07010
		07913
Privetkalarining Linear (MM/ 85 000)	GIBCO	25200-056
Polyethylenimine, Linear (MW 25,000)	Polysciences	
	Fisher Scientific	BP9706-160
	Fisher Scientific	BP26481
Ampicillin	Fisher Scientific	B1760-25
Talazoparib (BMN-673)	Selleck Chemicals	S7048
KU-55933 (ATM Kinase Inhibitor)	Selleck Chemicals	S1092
VE-821 (ATR Kinase Inhibitor)	Sigma-Aldrich	SML1415
Mechlorethamine hydrochloride (HN2)	Sigma-Aldrich	122564
Paclitaxel	Sigma-Aldrich	T7402
cis-Diammineplatinum(II) dichloride	Sigma-Aldrich	P4394
Doxorubicin hydrochloride	Sigma-Aldrich	D1515
(S)-(+)-Camptothecin	Sigma-Aldrich	C9911
Hexadimethrine bromide (Polybrene)	Sigma-Aldrich	107689
Corning® Cell-Tak and Tissue Adhesive	Corning	354240
Critical Commercial Assays	, ,	
PlasmoTest	Invivogen	REP-PT1
RNAeasy Plus Mini Kit	QIAGEN	74136
Comet Assay Kit	Trevigen	4250-050-K
Q5® Hot Start High-Fidelity 2X Master Mix	New England Biolabs	M0494S
NEBuilder® HiFi DNA Assembly Master Mix	New England Biolabs	E2621L
EdU-Click 594	Baseclick	BCK-Edu594
TOPO® TA Cloning® Kit for Sequencing	Invitrogen	450030
T4 DNA Ligase	New England Biolabs	M0202S
Experimental Models: Cell Lines		
HEK293T/17	ATCC	CRL-11268
LA-7	ATCC	CRL-2283
Experimental Models: Organisms/Strains		
FVB;129-Rb1 <sup>tm2Brn</sup> /Nci Referred to in this	Frederick National Laboratory for	01XC1
manuscript as "Rb1 <sup>FL</sup> "	Cancer Research	
FVB.129P2- <i>Trp53<sup>tm1Brn</sup>/</i> Nci Referred to in this manuscript as <i>"Trp53<sup>FL</sup>"</i>	Frederick National Laboratory for Cancer Research	01XC2
C57BL/6N-Gt(ROSA) 26Sor <sup>tm13(CAG-MYC,-CD2')Rsky</sup> /J Referred to in this manuscript as "R26 <sup>MycOE</sup> "	The Jackson Laboratory	020458
B6;129- <i>Gt(ROSA)</i> <i>26Sor<sup>tm1(CAG-cas9*,-EGFP)Fezh/</sup></i> J Referred to in this manuscript as " <i>R26<sup>Cas9</sup>"</i>	The Jackson Laboratory	024857
Oligonucleotides		
sgControl: CTGATTTGAATAATGATGCC	Generated in this study	N/A
sgMre11: TGGAGATCACTACTCGAGGC	Generated in this study	N/A
	· · · · · · · · · · · · · · · · · · ·	

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
Recombinant DNA		
pLV-Cre_LKO1	Addgene	12106
LentiCRISPR V2	Addgene	52961
psPAX2	Addgene	12260
pMD2.G	Addgene	12259
pEGFP-RNASEH1	Addgene	108699
LentiCRISPR-Cre-V2-sgControl-LumiFluor	Generated in this study	N/A
LentiCRISPR-Cre-V2-sgMre11-LumiFluor	Generated in this study	N/A
Lentiviral_pRRL-EF1a-GpNLuc	Gift from Antonio Amelio, Ph.D.	N/A
Lentiviral_pRRL-EF1a-NeuT-LumiFluor	Generated in this study	N/A
LentiCRISPR-Cre-V2-sgControl-RNaseH1	Generated in this study	N/A
LentiCRISPR-Cre-V2-sgMre11-RNaseH1	Generated in this study	N/A
Software and Algorithms		
BowTie2 v2.3.4.1	Langmead and Salzberg, 2012	https://sourceforge.net/projects/ bowtie-bio/files/bowtie2/2.3.4.1
Samtools v1.6.0	Li et al., 2009	http://www.htslib.org/download/
BedTools v2.26.0	Quinlan and Hall, 2010	https://github.com/arq5x/bedtools2
Python $\geq$ v3.5	https://www.python.org/	N/A
Ginkgo	Garvin et al., 2015	http://qb.cshl.edu/ginkgo/?q=
Völur algorithm	Generated in this study	https://github.com/pkMyt1/Volur
GNU Gzip v1.5	N.A.	https://www.gnu.org/software/gzip/
Graphpad Prism v8	N.A.	https://www.graphpad.com/
Python-Levenshtein Library v0.12.0	N.A.	https://github.com/ztane/ python-Levenshtein
Fiji	Schindelin et al., 2012	https://imagej.net/Fiji#Downloads
SnapGene software v4.3.4	GSL Biotech	https://www.snapgene.com
Open Comet v1.3.1	Gyori et al., 2014	http://www.cometbio.org
Other		
Mouse Reference Sequence GRCm38	http://www.Ensembl.org//useast.ensembl. org/?redirectsrc=//www.ensembl.org%2F	http://www.ensembl.org//useast. ensembl.org/Mus_musculus/Info/ Index?redirectsrc=//www.ensembl. org%2FMus_musculus%2FInfo% 2FIndex
Microsatellites, CpG Islands, Simple Repeats, SINE Elements, LINE Elements, LTR	UCSC Genome Browser	https://genome.ucsc.edu/cgi-bin/ hgTables
Genome annotation tables	http://www.Ensembl.org//useast.ensembl. org/?redirectsrc=//www.ensembl.org%2F	Ensembl v91

## LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Gaorav Gupta (gaorav\_gupta@med.unc.edu). All unique/stable reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

## EXPERIMENTAL MODEL AND SUBJECT DETAILS

## **Cell Lines**

HEK293T/17 cells were obtained from American Type Culture Collection (ATCC, CRL-11268) and were cultured according to manufactures' specifications. LA-7 cells obtained from American Type Culture Collection (ATCC, CRL-2283). LA-7 cells were maintained in DMEM/F12 medium supplemented with 10% fetal bovine serum, 20 mM HEPES, and 10 ug/ml Insulin ("LA-7 medium"). Prior to use as feeder cells, LA-7's were lethally irradiated with 70 Gy ionizing radiation using a Rad Source RS2000 irradiator.

## Primary Murine Mammary Epithelial Cells (pMMECs)

pMMECs were derived by harvesting the 4<sup>th</sup> and 5<sup>th</sup> mammary glands from 6-12-week-old female transgenic mice with the desired genotype. Glands were incubated in Liberase digestion medium (EpiCult-B Mouse Medium Kit (Stem Cell Technologies, 285 Units Collagenase Type 3 (Worthington), 20mM HEPES (GIBCO), 20 ug/mL Liberase Blendzyme 2 (Roche) and shaken (vertically) at 37°C overnight. The resulting digestion was spun down and resuspended in 3 mls trypsin with EDTA and 1000U DNase and incubated at 37°C for 5 min. LA-7 medium (DMEM-F12 media, 10% FBS, 20 mM HEPES, 10 µg/mL Insulin, 1X L-glutamine, 1X Penicillin-Streptomycin) was added to neutralize the trypsin. Cells were spun down and resuspended in 10U Dispase (Stem Cell Technologies) and 1000U DNase I (Worthington Biochemical) and incubated at 37°C for 5 min. Cells were washed twice with LA-7 medium and the resulting cells were resuspended in EpiCult-B Mouse Medium Kit (Stem Cell Technologies) and seeded onto Cultrex 3D-Culture Matrix (Trevigen) coated 6 well plates. For longer term cell growth experiments, pMMECs were seeded on lethally irradiated LA-7 cells and cultured in LA-7 medium. All cells were cultured to 80% confluence then passaged by trypsinization. Cells were tested monthly for mycoplasma using PlasmoTest Kit.

## **Transgenic Mouse Models**

Mice used in this study were housed in the Division of Comparative Medicine at the University of North Carolina at Chapel Hill, a facility accredited by the Association for the Assessment and Accreditation of Laboratory Animal Care International (AAALAC). R26<sup>LSL-Cas9</sup> (JAX#024857) and R26<sup>LSL-MycOE</sup> (JAX#020458) transgenic mouse strains were obtained from the Jackson Laboratory. Rb1<sup>fl/fl</sup> and Trp53<sup>fl/fl</sup> mouse strains were generously provided by the Perou laboratory, and originally obtained from the Frederick National Laboratory for Cancer Research (Strains #01XC1 and #01XC2). A subset of interbred mouse strains used in this study were analyzed by the Mouse Universal Genotyping Array (MUGA) from Neogen Genomics, and determined to be > 90% FVB with a minor contribution from C57BL/6J. For mammary tumor induction studies, six to twelve-week-old female R26<sup>Cas9/Cas9</sup>; Rb1<sup>fl/fl</sup>;  $Trp53^{fl/fl}$  mice received bilateral intraductal injections, into the fourth mammary gland, containing 5 × 10<sup>5</sup> transduction units (TU) of either LentiCRISPR-Cre-V2-sgControl (Cre-sgControl) or LentiCRISPR-Cre-V2-sgControl (Cre-sgMre11) lentivirus. Mouse cohorts were palpated for the development of mammary tumors twice weekly, and three times weekly after mammary tumors had formed. Mice were euthanized using humane experimental endpoints in accordance with UNC Institutional Animal Care and Use Committee (IACUC) guidelines. At necropsy, mammary tumors were harvested and sectioned into four pieces. Two pieces were immediately flash frozen for RNA and DNA extraction. One piece with any remaining glands (4th and 5th) were fixed in 4% paraformaldehyde and processed for paraffin embedding and H&E staining (Histoserv Inc.). One piece was taken for creation of tumor lines. Briefly, tumor pieces were incubated in digestion medium (DMEM, 10% FBS, 1mg/ml Collagenase Type 3, 1mg/ml Hyaluronidase) and shaken (horizontally) at 37°C for four hours. The resulting digestion was spun down and resuspended in trypsin with DNase and incubated at 37°C for 5min. LA-7 medium was added to neutralize the trypsin. Cells were spun down and resuspended in Dispase and deoxyribonuclease and incubated at 37°C for 5min. Cells were washed twice with LA-7 medium and passed through a 70 um filter. The resulting cells were resuspended in LA-7 media and seeded into co-culture with irradiated LA-7 feeder cells. The cells were cultured over 10 passages and analyzed via flow for GFP expression to monitor tumor cell outgrowth. Tumor lines were then utilized for downstream assays when GFP expression reached < 90%.

## **METHOD DETAILS**

## Cloning

### LentiCRISPR-Cre-V2-sgRNA LumiFluor plasmid

This plasmid was created by using restriction enzymes (Xbal and BgIII) to cut the Cre sequence from the pLV-Cre\_LKO1 plasmid and swapping it for the Cas9 sequence in lentiCRISPR V2 using restriction digest and T4 ligation. In order to get rid of the Bsmbl site within Cre, Gibson cloning was used (HiFi DNA Assembly Master Mix; NEB) to change the sequence of a Valine residue from GTC to GTA, thus removing the site while preserving the protein sequence. Using the remaining BsmB1 sites, the sequences for sgControl (53bp1 intron sequence) and sgMre11 were inserted into the sgRNA scaffolding region.

### LentiCRISPR-Cre-V2-sgRNA-RNasH1

This plasmid was created via Gibson cloning of the RNaseH1 gene from the pEGFP-RNASEH1 plasmid (Addgene #108699) to replace the LumiFluor gene in the LentiCRISPR-Cre-V2-sgControl LumiFluor and LentiCRISPR-Cre-V2-sgMre11 LumiFluor plasmids.

## Lentiviral\_pRRL-EF1a-NeuT-LumiFluor

This plasmid was created via Gibson cloning of the Lentiviral\_pRRL-EF1a-GpNLuc plasmid to insert the *NeuT* gene (constitutively active truncation mutant of *Neu*, the rodent ortholog of *Her2*) and an autocleavage P2A sequence upstream of the luciferase gene. *NeuT*-P2A was synthesized as a gene block (IDT), using pSV2-NeuT (Addgene #10919) as the reference sequence. Because *NeuT* is commonly used to recapitulate Her2-initiated tumorigenesis, we refer to this construct as "Her2" in the text and figures of the manuscript.

All plasmids created were confirmed by Sanger sequencing (Eton Bioscience Inc.). LentiCRISPR-Cre-V2-sgControl-Lumifluor, LentiCRISPR-Cre-V2-sgMre11-Lumifluor, LentiCRISPR-Cre-V2-sgControl-RNaseH1, LentiCRISPR-Cre-V2-sgMre11-RNaseH1, LentiViral\_pRRL-EF1a-GpNLuc, and LentiViral\_pRRL-EF1a-NeuT-LumiFluor will be made available by the Lead Contact.

## **Topo cloning**

pMMECs were infected twice with either Cre-sgControl or Cre-sgMre11. Upon confirmation of at least 80% viral efficiency, Topo Cloning was performed per manufactures protocol. Sequencing was performed by Eton Biosciences and analysis performed using Bioedit software.

## Immunofluorescence

pMMECs were infected twice with either Cre-sgControl, Cre-sgMre11, GFP, or NeuT virus. Upon confirmation of at least 80% viral efficiency, cells were seeded onto 3D-matrix coated coverslips, were treated with EdU for 10 min, and subsequently fixed by cold Methanol:Acetone (1:1) incubation at -20C for 10 mins. Cells then underwent EdU detection using the EdU detection kit (Baseclick) in accordance with kit instructions. Cells were then blocked in PBS + 5% FBS for 1 hour, followed by incubation in the appropriate primary antibody for 1 hour (p- $\gamma$ H2AX; 53bp1; p-RPA2; or S9.6), secondary antibody for 30 min, then DAPI for 1 min. Coverslips were then mounted onto slides with Prolong Gold mounting medium, cured for 2 hours and stored at 4C in the dark until imaging. Coverslips were examined on an Olympus BX61 upright wide field microscope. Resulting foci were analyzed using Fiji software (Schindelin et al., 2012).

## **Comet assay**

pMMECs were infected twice with either Cre-sgControl, Cre-sgMre11, GFP, or NeuT virus and confirmation of at least 80% viral efficiency was determined by Flow Cytometry (Attune NxT) for GFP expression. The presence of SSBs and DSBs were analyzed via Alkaline (SSBs/DSBs) and Neutral (DSBs) comet assay using the Trevigen comet assay kit according to manufacturer's protocol. Comet images were captured by fluorescence microscopy using Olympus BX61 upright wide field microscope. The tail DNA percent was quantified using the ImageJ software with OpenComet plug-in (Gyori et al., 2014).

## **Growth assays**

pMMECs were infected twice with either Cre-sgControl or Cre-sgMre11 virus then seeded into 12 well plates at a density of 3 X 10<sup>4</sup> cells/well onto LA-7 feeder cells. Duplicate samples were harvested every 2-3 days for 15 days. Total cells/well were counted, cells were fixed in 3% PFA and subjected to flow analysis (Attune NxT) for the presence of GFP. Prior to ATMi growth assays, the dose of ATMi that resulted in 50% and 100% inhibition of the ATM pathway was determined. Wild-type mouse embryonic fibroblasts (MEFs) were treated with 0, 1, 3, 5, 7, 10, 15, or 20 uM of the ATM inhibitor Ku55933 for 30 min. They were then subjected to 10 Gy, incubated under normal culture conditions and then cell lysate was collected for western blotting analysis of the phosphorylated form of the ATM downstream protein, Kap1. For ATMi growth assays pMMECs were treated as above but with the additional presence of either 5 or 10 uM of ATMi.

## **Viral production and infection**

HEK293T/17 cells were transfected, using Polyethylenimine (PEI), with viral packaging plasmids, psPax2 and pMD2.G, and either LentiCRISPR-Cre-V2-sgControl-Lumifluor, LentiCRISPR-Cre-V2-sgMre11-Lumifluor, Lentiviral\_pRRL-EF1a-GpNLuc, or Lentiviral\_pRRL-EF1a-NeuT-LumiFluor, LentiCRISPR-Cre-V2-sgControl-RNaseH1, or LentiCRISPR-Cre-V2-sgMre11-RNaseH1 plasmids. Twenty-four hours post transfection, cells were washed and refed with fresh medium. Viral containing media was collected for 3 days. Collected media was filtered (0.45 um) then spun down for 2 hours at 16C at 21,000 rpm. Virus containing pellet was resuspended in PBS and incubated at  $4^{\circ}$ C for 24 hours then aliquoted and stored at -80C. For lentiviral infections, cells were transduced with the appropriate virus combined with  $4\mu$ g/ml Polybrene overnight. Cells were refed with fresh viral containing medium and incubated overnight two to three times (2-3 back to back infections). Following the last infections cells were washed three times with PBS and cultured with MEGM. For testing viral efficacy, a small sample of cells were fixed with 3% Paraformaldehyde (PFA) and were assessed via flow cytometry (Attune NxT) for the presence of GFP indicating Cas9 expression or were stained with anti-CD2-PE indicating Myc expression.

## Chemotherapeutic and inhibitor sensitivity assays

Tumor cells were seeded at a density of 1500 cells per well of a 96 well dish and allowed to attach overnight. After the cells were attached, various drugs or DMSO control were added to the media to the final concentrations shown in the figures. Each treatment was done in triplicate. Cells underwent IncuCyte live cell imaging (S'artorius), scanned every two hours to determine cell confluence and growth rates over seven days.

## Automated analysis of mouse mammary tissues

The process of quantitative image analysis begins with the acquisition of high-resolution digital slides. FFPE sections of mouse mammary tissue stained with hematoxylin and eosin were scanned on an Aperio ScanScope XT (Leica Biosystems). Images were then uploaded to eSlide Manager and visualized with ImageScope 12.3 (Leica Biosystems). Separate tissue sections on each image were annotated by KFS. Mouse mammary glands four and five were differentiated based on the location of the lymph node which separates the glands. The annotated images were then imported to Definiens Architect XD 2.7.0 Build 60765 x64 for analysis with Tissue Studio version 4.4.2. Using the Tissue Studio portal, the annotated images were preselected for region-of-interest (ROI) detection. The Definiens Composer algorithm was used to segment the tissue into different ROIs: Brown Adipose, Epithelium, RBCs, Stroma, Glass, and White Adipose. This algorithm was trained on representative regions to classify all the tissue within the ROIs in the final analysis. The program then calculated the total tissue area and the area percentages for each of the ROIs. In addition, the program was configured to detect and score nuclei within each of the ROIs, based on a hematoxylin stain threshold and average size set by the analyst. All nuclei were counted and classified as small (<  $20 \,\mu$ m<sup>2</sup>), medium ( $20 - 40 \,\mu$ m<sup>2</sup>), or large (>  $40 \,\mu$ m<sup>2</sup>). These values were used to calculate a histological score equal to (1 x % nuclei small) + (2 x % nuclei medium) + (3 x % nuclei large). The analysis output included all quantitative results as well as screen captures of the ROI detection plus overlays and the cellular analysis (nuclei). Slide scanning and tissue quantification was performed by the Bentley R. Midkiff at the Translational Pathology Lab at UNC.

## Human breast cancer tissue microarrays

Tissue microarrays were constructed from 271 patients with non-metastatic TNBC (ER/PR < 1%; HER2 0/1+, or HER2 2+/FISH not amplified) who underwent surgical resection of their primary tumor at our institution between 2002 and 2007. Exclusion criteria were < 1 cm primary tumor size, prior breast radiation, inflammatory breast cancer, and neoadjuvant chemotherapy. Immunohisto-chemistry (IHC) was performed using rabbit polyclonal anti-sera generated against human Mre11 (1:3000) and human Nbs1 (1:3000). Tumors were classified as low-expressors of the Mre11 complex if < 10% of cancer cells had detectable nuclear protein expression of Mre11 and/or Nbs1 relative to background staining levels, and determined by two independent reviewers. 254 out of 271 cases were deemed evaluable. Lack of immunoreactivity was confirmed by repeating the IHC using whole tissue sections in a subset of Mre11 complex low-expressors.

## **QUANTIFICATION AND STATISTICAL ANALYSIS**

### Single cell whole genome sequencing

Mammary epithelial cells for single cell sequencing were derived from littermate *WT* and *Mre11*<sup>ATLD1</sup>/ATLD1</sup> female mice. Lentivirus expressing EGFP or EGFP+HER2 was added to the cells for 24 hours. Following this infection, the cells were transferred to 10 cm tissue culture dishes containing irradiated LA7 cell feeder layers. The cells were maintained in culture for 2 weeks at which time they were dissociated into a single cell suspension and sorted for EGFP positive cells into a 96 well plate. These were used for the whole genome single cell sequencing described below.

Single cell sequencing libraries were prepared using the WGA4 kit (Sigma-Aldrich) as previously described (Garvin et al., 2015; Martelotto et al., 2017). Half of the individual cell libraries (48/96) were pooled into each of two library pools. Each pool was run on a single lane for 100-cycle single end sequencing (HiSeq2500, Illumina). The resulting FASTQ files and a bed file defining the indices for the pool, were used as input into Völur (https://github.com/pkMyt1/Volur). The pipeline trimmed 27 nucleotides from the end of each read, aligned the reads to the GRCm38 reference with BowTie2 (Langmead and Salzberg, 2012), converted the SAM files to BAM with SamTools (Li et al., 2009), and output gzipped, demultiplexed BED files suitable for use with Ginkgo (Garvin et al., 2015). Ginkgo was run using a variable 25 Kb segment size, 76 base pair bowtie simulated reads, and global (sample with lowest LOD) segmentation. All other parameters were left as the default. Once Ginkgo completed the segment copy data file was downloaded. This file was modified by removing the chromosome-Y elements, masking alignment errors by setting the copy number in those regions to 2, and normalizing any cell called triploid to diploid. Chromosome Y was removed because these were female mice. These mice are not the same genetic background as the GRCm38 reference. This resulted in regions of copy gain or loss common to all samples.

## **Quantification and statistical analysis**

Since we did not provide a known diploid sample for Ginkgo to use as a reference, any cell that was scored as being triploid by Ginkgo was adjusted by subtracting 1 from the copy value of each segment. Segments observed in the *Mre11* <sup>ATLD1</sup> cells that were likely mapping errors because of the mouse strain used were masked by setting those regions to a copy number of 2 creating a no triploid, *Mre11*<sup>ATLD1</sup> masked segment copy bed file. Ginkgo defines the copy number based on how many reads mapped to each predefined segment. The mapped reads for this determination may cross the segment boundaries. Because of this the copy number transition cannot be assigned to the segment boundaries. To account for this uncertainty, we use a breakpoint for our analysis. We define a breakpoint region as the start coordinate of the preceding segment through the stop coordinate of the segment containing the copy number change. Using the masked segment copy file, the number and location of each breakpoint was derived for each cell type. The unique breakpoints were then intersected with bed files of interest using BedTools (Quinlan and Hall, 2010) via the Py-thon pybedtools API within Völur.

## Statistical significance of breakpoint region intersects

The determination that an observed set of breakpoint regions is enriched for overlap with annotated genome regions (e.g., genes, obtained from http://www.Ensembl.org//useast.ensembl.org/?redirectsrc=//www.ensembl.org%2F) was based on generation of an empirical null distribution dataset by randomly shuffling the genomic locations of the individual breakpoint regions for each cell and then determining the frequency of intersects with the genome feature of interest. This process was repeated 10,000 times for each intersect set. The observed frequency of intersection value was considered significantly different than the randomly shuffled dataset if it was < 5% or > 95% of this empirical null distribution. Specific association with a target set was defined as being those target sets that were not enriched for the control cells but were enriched for the oncogene expressing or  $Mre11^{ATLD1}$  cells.

## **Tumor whole genome sequencing**

Whole genome sequencing services were provided by Novogene. Reads were trimmed using SeqPurge (Sturm et al., 2016), aligned to the mm10 mouse reference using bwa-mem (Li, 2013) and subsequently realigned with ABRA2 (Mose et al., 2019). The resultant BAM files were sorted and duplicate marked using biobambam2 (Tischler and Leonard, 2014). Small variants were called using Strelka2, Mutect2 and Cadabra (Cibulskis et al., 2013; Kim et al., 2018; Mose et al., 2019). Mutect2 calls were filtered using the GATK's best practices whereas the Strelka2 and Cadabra default filters were applied. Quality thresholds were used to filter calls using values of QSS\_NT > = 70 and QSI\_NT > = 40 for Strelka2, SNV TLOD > = 9 and Indel TLOD > = 10 for Mutect2, and QUAL > = 15 for Cadabra. Variants were additionally filtered using a panel of normals constructed using the Mutect2-GATK best practices. Structural variants were called using Manta (Chen et al., 2016) and copy number variants were called with CNVKit (Talevich et al., 2016). Mutation signature analysis was performed using deconstructSigs (Rosenthal et al., 2016). Circos plots were created with Circa (http://omgenomics.com/circa).

## **Statistical Analyses**

Two-tailed statistical tests for experimental data were conducted using Graphpad Prism version 8. The specific test used in each analysis is indicated in the corresponding figure legend.

## DATA AND CODE AVAILABILITY

The custom algorithms developed for this study are available at https://github.com/pkMyt1/Volur. Whole genome sequencing reads for murine breast cancers with matched normal liver have been uploaded to NCBI Sequence Read Archive (SRA) with accession number PRJNA595908. Single cell whole genome sequencing BAM files and copy number segment files have not been uploaded to SRA due to data format incompatibility, and can be requested from the Lead Contact.