Landscape analysis of adjacent gene rearrangements reveals BCL2L14–ETV6 gene fusions in more aggressive triple-negative breast cancer

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Triple-negative breast cancer (TNBC) accounts for 10 to 20% of breast cancer, with chemotherapy as its mainstay of treatment due to lack of well-defined targets, and recent genomic sequencing studies have revealed a paucity of TNBC-specific mutations. Recurrent gene fusions comprise a class of viable genetic targets in solid tumors; however, their role in breast cancer remains underappreciated due to the complexity of genomic rearrangements in this cancer. Our interrogation of the whole-genome sequencing data for 215 breast tumors catalogued 99 recurrent gene fusions, 57% of which are cryptic adjacent gene rearrangements (AGRs). The most frequent AGRs, BCL2L14–ETV6, TTC6–MIPOL1, ESR1–CCDC170, and AKAP8–BRD4, were preferentially found in the more aggressive forms of breast cancers that lack well-defined genetic targets. Among these, BCL2L14–ETV6 was exclusively detected in TNBC, and interrogation of four independent patient cohorts detected BCL2L14–ETV6 in 4.4 to 12.2% of TNBC tumors. Interestingly, these fusion-positive tumors exhibit more aggressive histopathological features, such as gross necrosis and high tumor grade. Amid TNBC subtypes, BCL2L14–ETV6 is most frequently detected in the mesenchymal entity, accounting for ~19% of these tumors. Ectopic expression of BCL2L14–ETV6 fusions induce distinct expression changes from wild-type ETV6 and enhance cell motility and invasiveness of TNBC and benign breast epithelial cells. Furthermore, BCL2L14–ETV6 fusions prime partial epithelial–mesenchymal transition and endow resistance to paclitaxel treatment. Together, these data reveal AGRs as a class of underexplored genetic aberrations that could be pathologic in breast cancer, and identify BCL2L14–ETV6 as a recurrent gene fusion in more aggressive form of TNBC tumors.

BCL2L14–ETV6  |  gene fusion  |  triple-negative breast cancer  |  recurrent adjacent gene rearrangements  |  chemoresistance

Recurrent gene fusions that result from chromosome translocations comprise a critical class of genetic cancer-causing aberrations, which have fueled modern cancer therapeutics. In the past decade, the discovery of novel gene fusions in epithelial tumors have generated great therapeutic impact in recent years. This is represented by the discovery of an EML4–ALK fusion in ~4% of lung cancer and the FGFR–TACC fusion in ~3% of glioblastomas that have culminated in effective targeted therapies in these tumors (1, 2). Most recently, larotrectinib targeting the NTRK gene fusions accounting for up to ~1% of solid tumors have received Food and Drug Administration approval for pan-cancer use, which is considered as the first targeted therapy with tissue-agnostic indication (3). Although low in percentages, these neoplastic gene fusions will likely move toward genetic subtyping of solid tumors that may be possibly curable by fusion-targeted therapies.

In our previous study, analysis of a Cancer Genome Atlas (TCGA) RNA-sequencing (RNA-seq) dataset identified a recurrent gene fusion between the 5′ region of ESR1 and the coding region of the adjacent CCDC170 gene, which was subsequently verified by several other studies (4–8). This fusion represents a cryptic class of genomic rearrangements between adjacent genes (genes within 500-kb distance), which we termed as adjacent gene rearrangements (AGRs). ESR1–CCDC170 is detected in 6 to 8% of luminal B breast tumors and promotes increased aggressiveness (9), which suggests that AGRs can meaningfully contribute to breast cancer development, pathogenesis, and resistance to cancer therapies. Nonetheless, AGRs have been frequently overlooked by fusion detection tools based on RNA-seq data due to the overwhelming number of adjacent chimeras resulting from intergenic splicing events. In addition, such cryptic genomic changes cannot be detected by conventional cytogenetic assays, such as spectral karyotyping or fluorescence in situ hybridization due to the proximity of the rearranged DNAs.

Significance

Identification of triple-negative breast cancer (TNBC)-specific genetic events that could guide the treatment decisions represents an unmet clinical need, whereas recent genomic sequencing studies have revealed a paucity of TNBC-specific mutations. In this study, analysis of whole-genome sequencing data for 215 breast tumors catalogued 99 recurrent gene fusions. Among different types of rearrangements, we found that a special class of cryptic adjacent gene rearrangements (AGRs) may occur more frequently than realized in breast cancer. The most frequent AGRs are preferentially found in more aggressive forms of breast cancers, among which BCL2L14–ETV6 is exclusively detected in TNBC. BCL2L14–ETV6 is enriched in high-grade, necrotic, mesenchymal TNBC tumors, and endows increased invasiveness and paclitaxel resistance via inducing partial epithelial mesenchymal transition.

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Competing interest statement: M.T.L. is a founder and limited partner in StemMed Ltd., and a manager in StemMed Holdings, its general partner. He is a founder and equity stake holder in Tvardi Therapeutics Inc.

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Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession nos. GSE120919 for the RNA-seq data on BT20 models and GSE123917 for combined BT20 expression profile data).

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and the limited resolutions of these assays. For these reasons, AGRs remain an underexplored area of breast cancer genetics.

Here, we performed a landscape study of adjacent gene rearrangements in breast cancer catalogued by whole-genome sequencing (WGS) data, and identified a recurrent fusion, BCL2L14-ETV6, that is preferentially present in triple-negative breast cancer (TNBC). The fusion partners, an ETS family transcription factor gene (ETV6) and an apoptosis facilitator Bcl-2-like protein 14 gene (BCL2L14) are neighboring genes of HER2, and PI3KCA mutations in TCGA 92 breast tumors. The AGRs detected in at least two TCGA tumors and using Fisher's exact test (two-tailed).

Results

AGRs Comprise the Most Frequent Form of Intergenic Rearrangements in Breast Cancer. To provide a systematic picture of AGR events in breast cancer, we first analyzed the full spectrum of experimentally confirmed somatic translocations in 9 breast cancer cell lines and 15 breast tumors catalogued from WGS data in a previous study.

Fig. 1. Landscape of recurrent adjacent gene rearrangements in breast cancer revealed by WGS data. (A) Frequency chart of experimentally validated interchromosome, intrachromosome distant, and intrachromosome adjacent translocations in nine breast cancer cell lines and 15 breast tumors revealed by WGS data (13). Among 9,408 confirmed somatic translocations, about half are intrachromosomal translocations within 500-kb distance. (B) CIRCOS plot showing the landscape of 99 recurrent gene rearrangements detected in 215 breast tumors based on WGS data from the ICGC. The histogram inside the CIRCOS plot represents the recurrence of the gene rearrangements in the chromosome position, indicating the number of patients that harbor the gene fusion. (C) Genomic hotspots of colinear and noncolinear AGR or intrachromosomal gene rearrangements. Adjacent and intrachromosomal rearrangements in the genomes are displayed in rainfall plot, with each dot representing a respective positive sample. The x axis shows 24 chromosomes in the human genome, and y axis shows the distance between the rearrangement points (base pairs at log2 scale). The horizontal line indicates the cutoff for adjacent gene rearrangements (500 kb in distance). (D) Scatter plot showing the incidences of 99 recurrent gene rearrangements and their concept signature scores, which were detected from 215 ICGC breast tumors profiled by WGS. The x axis indicates the incidence of gene rearrangements in the cohort. The y axis indicates the maximum ConSig scores of 5’ or 3’ partner genes. (E) Tile plot showing the top recurrent AGRs and the known breast cancer oncogenes, including ER, PR, HER2, and PIK3CA mutations in TCGA 92 breast tumors. The AGRs detected in at least two TCGA tumors and >1% of all ICGC tumors are shown in the figure. Group-wise mutual exclusivity test using discrete independence statistics called “Discover” that take into account the distribution of all somatic gene rearrangements, suggests that there are significant number of tumors that harbor only one of these AGRs (11) [i.e., ETV6–NTRK3 fusion in secretory breast carcinoma (12)]. In this study, we further investigated the pathological role of BCL2L14–ETV6 in TNBC.
Our analysis include the development of fusion genes underlying cancer based on their association analysis of these lead recurrent AGRs revealed their preferential presence in the more aggressive forms of breast cancers, including basal-like and luminal B cancers (Fig. 1). As shown in Fig. 1A, a majority of the intrachromosomal translocations are within 500 kb apart, with a median distance of ~100 kb. This suggests that AGRs may be a more frequent genetic event than realized. Although most of these rearrangements are likely the consequence of genomic instability, it is plausible that a subset of them could be recurrent genetic events that are pathological in breast cancer.

To discover AGRs in breast cancer systematically, we further analyzed the somatic structural mutations catalogued by the International Cancer Genome Consortium (ICGC) based on WGS data for 215 breast tumors. The somatic structural mutations were first mapped with the human exome to reveal genes and exons affected by the rearrangements. The fusion partners were determined based on the strands and genomic regions retained in the rearrangements. To explore if the intergenic rearrangements are enriched in specific breast cancer subtypes, we isolated the 92 ICGC breast tumors contributed by TCGA that have detailed histopathological data from a recent report (14) (SI Appendix, Fig. S1). Overall Her2 and Basal subtypes show significantly higher total number of rearrangements compared to luminal A tumors. Luminal B tumors also exhibit a trend of increased total rearrangements than luminal A tumors. In addition, the breast tumors with high nuclear pleomorphism show significantly higher number of rearrangements. Next, we identified the recurrent gene fusions and classified them into AGRs (local rearrangements involving genes less than 500 kb apart), distant intrachromosomal rearrangements (involving partner genes more than 500 kb apart), or interchromosomal rearrangements (Fig. 1B). In total, we identified 99 recurrent gene fusions that occur in at least two breast tumors, including 57 adjacent gene fusions (57.6%), 35 intrachromosome fusions (35.4%), and 7 interchromosome fusions (7.1%) (Dataset S1). The AGR events spread throughout the genome, with some genomic regions harboring higher incidence of recurrent gene rearrangements (Fig. 1C). Among the 57 recurrent AGRs, 20 are between colinear genes with 5′ located upstream of the 3′ partner (35.1%) and 35 are between noncolinear genes with the 5′ partner located downstream of the 3′ partner (61.4%), which are likely the results of intergenic deletions or tandem duplications, respectively.

Systematic Discovery of Recurrent AGRs in Breast Cancer. We ranked the recurrent gene rearrangements based on their incidence in the ICGC breast tumor patient cohort, and their concept signature (ConSig) scores (Fig. 1D and Dataset S1). The ConSig scores were developed in our previous study to compute the functional relevance of fusion genes underlying cancer based on their associations with molecular concepts associated with known cancer causal genes (15). The top four most frequent gene fusions identified by our analysis include BCL2L14–ETV6, TTC6–MIPOL1, ESRI–CCDC170, and AKAP8–BRD4, all of which are AGRs (Fig. 1D). To test if the top recurrent AGRs could be a function of genomic instability, we isolated the 92 TCGA breast tumors that have available DNA damage repair (DDR) deficiency scores (16), and sorted these tumors by their genomic instability index (GII) (SI Appendix, Fig. S2). Overall these fusions showed modest enrichment in the tumors with high GII scores, suggesting that DDR deficiency may facilitate formation of a subset of the rearrangements generating these fusions. Further clinicopathological association analysis of these lead recurrent AGRs revealed their preferential presence in the more aggressive forms of breast cancers, including basal-like and luminal B breast cancers (Fig. 1E and Dataset S2). Among the top AGRs, BCL2L14–ETV6 and AKAP8–BRD4 are exclusively found in basal-like breast cancers, while TTC6–MIPOL1 and ESRI–CCDC170 (9) are preferentially present in luminal B tumors. While basal-like and luminal B tumors tend to have a higher number of rearrangements, the specific enrichment of these fusions in either of these subtypes but not in all genomically unstable entities implies their potential function in these tumors. To test if the lead recurrent AGRs display alteration patterns in which most tumors only have one of these fusions, we performed mutual exclusivity tests using discrete independence statistics, called “Discover,” that account for the heterogeneous rearrangement rates across tumors (17). A group-wise mutual exclusivity test for the top recurrent AGRs suggests that there are significant number of tumors that harbor only one of these rearrangements ($P < 0.001$) (Fig. 1E). This suggests that these recurrent AGRs tend not to cooccur in the same tumor, as opposed to typical DDR-driven rearrangements coexisting in DDR-deficient tumors. Next, we surveyed the incidences of rearrangements based on fusion partner genes and stratified these incidences based on TCGA clinicopathological features (SI Appendix, Fig. S3). The result revealed that most of the lead fusion genes are preferentially present in high-grade tumors, except for TENM4, SHANK2, and TPM3P9. Among these lead fusion genes, we detected several kinase fusion genes, such as DLG2, BRD4, and TNIK. Taken together, the preferential presence of these recurrent AGRs in specific aggressive forms of breast tumors and their tendency not to coexist in genomically unstable tumors imply their potential pathological roles in breast cancer.

Characterization of the Lead Recurrent AGRs in Breast Cancer Samples. To explore if the most frequent gene rearrangements are significantly associated with specific histopathological features, we analyzed the detailed histopathological data of TCGA breast tumors available from a recent report (14). Interestingly, our analysis revealed that BCL2L14–ETV6 and AKAP8–BRD4 tend to occur in breast tumors with gross necrosis (particularly, extensive necrosis), higher tubule formation score, and higher nuclear pleomorphism (SI Appendix, Fig. S4 and Dataset S3). Tumor necrosis is defined as the morphological changes following cell death (18). The presence of necrosis in breast cancer indicates more aggressive tumors that are associated with early recurrence, poor prognosis (19), and ~35% of TNBC tumors present necrosis features (20). To further verify the above histopathological associations in a larger cohort of TNBC tumors, we analyzed the somatic rearrangements detected by WGS data in 516 breast tumors, which are provided by the Catalogue of Somatic Mutations in Cancer (COSMIC) (21, 22). From a total of 162 TNBCs in this cohort, we detected 10 BCL2L14–ETV6+ cases, but there is no AKAP8–BRD4 case (Table 1 and Dataset S4). In both TCGA and COSMIC cohorts of TNBC tumors, the BCL2L14–ETV6+ tumors tend to have a higher level of ETV6 expression than fusion-negative cases, but not all ETV6-overexpressing tumors harbor the BCL2L14–ETV6 fusion (SI Appendix, Fig. S5). The BCL2L14–ETV6 fusions are exclusively detected in TNBC, and correlate with more aggressive features, including presence of necrosis, high mitotic and nuclear pleomorphism scores, advanced tumor stage, and high pathology grade, consistent with the above findings (Fig. 1F). In addition, among TNBC subtypes, BCL2L14–ETV6 fusions most frequently present in the mesenchymal subtype characterized by enriched cell motility and epithelial-to-mesenchymal transition (EMT) pathways, accounting for ~19.2% of these tumors in the TCGA+COSMIC cohort (Fig. 1F). In addition, BCL2L14–ETV6 is also detected in 11.6% of the basal-like 1 (BL1) tumors characterized by enriched cell cycle and cell-division pathways (23).

We then proceed to validate the lead recurrent AGR fusions, including BCL2L14–ETV6, TTC6–MIPOL1, and AKAP8–BRD4, in a panel of breast cancer cell lines and human breast cancer tissues by RT-PCR. The validation of the most frequent AGR, BCL2L14–ETV6, will be detailed below. Since TTC6–MIPOL1 is preferentially expressed in luminal breast tumors, we first screened...
this fusion in 141 estrogen receptor (ER)\textsuperscript{+} breast tumors from the University of Pittsburgh (Pitt) cohort using primers located on the first exon of \textit{TTCT} and the last exon of \textit{MIPOL1}, which identified one positive case in this cohort (\textit{SI Appendix, Fig. S6}4). In addition, we also detected this fusion in one luminal cell line (MDA-MB-361) (\textit{SI Appendix, Fig. S6B} and verified the presence of the \textit{AKAP8–BRD1} fusion in one patient-derived xenograft (PDX) tumor through screening a panel of 34 TNBC PDX tumors (24, 25) (\textit{SI Appendix, Fig. S7A}).

\textit{BCL2L14–ETV6} Is Exclusively Detected in \textit{TNBC}. Next, we focused our study on the \textit{BCL2L14–ETV6} rearrangements, which were identified in 12.2% and 6.2% of TNBC cases in the TCGA and COSMIC cohorts, respectively (Fig. 1F and Table 1). We first detected \textit{BCL2L14–ETV6} fusion transcripts by RT-PCR in 134 TNBC tumors from two available patient cohorts. To detect most fusion variations, we designed a pair of primers located on exon 2 of \textit{BCL2L14} and the last exon of \textit{ETV6}, respectively. This primer set detected \textit{BCL2L14–ETV6} fusion in 4 of the 89 TNBC tumors from the Pitt cohort (Fig. 2A), and 2 of the 45 TNBC tumors from the Baylor College of Medicine (BCM) cohort (Fig. 2B). The clinopathological features for all of the 134 TNBC patients from Pitt and BCM cohorts are provided in Dataset S5. The fusion-positive cases were subsequently verified by capillary sequencing. Next, we tested the expression of \textit{BCL2L14–ETV6} in a panel of 44 breast cancer cell lines and 34 TNBC PDX tumors. We detected one PDX tumor that expresses \textit{BCL2L14–ETV6} but not in the cell lines tested (\textit{SI Appendix, Fig. S8}). The most common variant detected is the fusion between exon 4 of \textit{BCL2L14} and exon 2 of \textit{ETV6} (referred to as E4E2) that present in two patient cases and one PDX tumor. We also tested \textit{BCL2L14–ETV6} by RT-PCR in 200 ER\textsuperscript{+} breast tumor tissues from the BCM cohort but did not detect any fusion-positive ER\textsuperscript{+} tumors, which supports its TNBC specificity (\textit{SI Appendix, Fig. S9}).

To assess if the \textit{BCL2L14–ETV6}\textsuperscript{+} tumors present the histopathological features discussed above, we performed histopathological evaluations for the four index tumors from the Pitt cohort for which the tissue sections are available. All four tumors are reported as grade 3 tumors with a high nuclear pleomorphism score and high mitotic count score (Dataset S6). In addition, two of four fusion-positive tumors present extensive necrosis and the remaining two fusion-positive tumors present focal necrosis (\textit{SI Appendix, Fig. S10}), consistent with the above findings.

Characterization of \textit{BCL2L14–ETV6} Genomic Rearrangements and Protein Products. To verify the genomic origin of \textit{BCL2L14–ETV6} in the positive cases, we performed genomic PCR using tiling primers designed specifically for \textit{BCL2L14} or \textit{ETV6} intron regions predicted to harbor the rearrangement based on the fusion variants detected in the index cases from the BCM cohort. This assay successfully amplified the genomic fusion points in both of the \textit{BCL2L14–ETV6}\textsuperscript{+} tumors in the BCM cohort (Fig. 2C). The breakpoint junctions in the genomic DNA were further verified by capillary sequencing. Next, we explored the association of \textit{BCL2L14–ETV6} with copy number aberrations in the TCGA cohort. Copy number data revealed frequent somatic tandem duplications in the \textit{ETV6}/\textit{BCL2L14} loci, which are present in four of the five positive TCGA tumors detected by WGS data (\textit{SI Appendix, Fig. S11A}). In addition, copy number data also revealed tandem duplications delineating the \textit{ETV6}/\textit{BCL2L14} loci in the TCGA tumors that were not profiled by WGS, suggesting these as potential positive cases. These data suggest that \textit{BCL2L14–ETV6} fusions may be the result of either tandem duplications or reciprocal rearrangements that generate both \textit{BCL2L14–ETV6} and \textit{ETV6–BCL2L14} fusions (Fig. 1E), as with the \textit{ESR1–CCDC17O} fusion we identified previously (9).

Next, we investigated the structure of \textit{BCL2L14–ETV6} proteins. Among five variants detected, three variants (E2E3, E2E6, and E4E2) encode chimeric proteins containing the amino terminus (N terminus) of \textit{BCL2L14} and the carboxyl-terminus (C terminus) of \textit{ETV6} (Fig. 3A). The \textit{ETV6} protein contains an N-terminal pointed (PNT) domain responsible for protein partner binding, and a C-terminal DNA-binding (ETS) domain critical for DNA binding-dependent transcriptional repressor function. Both the most common variant, E4E2, and the E2E3 variants retain the PNT domain and ETS domain, whereas the E2E6 protein only retains the ETS domain. E4E3 and E5E5, on the other hand, do not translate the protein sequence of \textit{ETV6} due to a frameshift after the fusion junction, resulting in expression of C-terminus–truncated \textit{BCL2L14} proteins.

Next, we ectopically expressed the open-reading frames (ORFs) of the fusion variants E2E3, E4E3, and E4E2 in the fusion-negative MCF10A breast epithelial cell line and the BT20 basal-like breast cancer cell line, both of which are triple-negative in (ER, progesterone receptor [PR], and HER2) receptor expression (26). Cells transduced with the vector containing the \textit{lacZ} vector or the vector containing the wild-type (wt) \textit{ETV6} ORF were used as controls. Western blot using polyclonal antibodies against the C terminus of \textit{ETV6} or the N terminus of \textit{BCL2L14} detected strong expression of the E2E3 (62 kDa) and E4E2 (74 kDa) proteins in the transduced BT20 and MCF10A cells (Fig. 3B and C). The 27-kDa E4E3 fusion protein was detected by the \textit{BCL2L14} antibody, but not by the \textit{ETV6} antibody, suggesting that this variant encodes a truncated \textit{ETV6} protein, which does not contain the \textit{ETV6} protein sequence. Next, we sought to detect the endogenous \textit{BCL2L14–ETV6} fusion protein in the PDX tumor expressing the E4E2 variant (BCM-2147). Western blot using the \textit{ETV6} antibody detected the same-sized band of E4E2 protein as in the engineered BT20 cells (Fig. 3D).

Since gene fusions tend to translocate to abnormal cellular compartments (9), we investigated the cellular localization of the fusion proteins compared to wt\textit{ETV6} protein in the transduced \textit{BCL2L14} in the engineered BT20 and MCF10A cells. Due to the lack of specific antibody

### Table 1. Incidence of \textit{BCL2L14–ETV6} gene fusion detected in four different patient cohorts of 942 breast tumors

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Method</th>
<th>Total</th>
<th>Non-TNBC</th>
<th>TNBC</th>
<th>Necrotic TNBC</th>
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<tr>
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<td>WGS</td>
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<td>0/48 (0)</td>
<td>5/41 (12.2)</td>
<td>3/23 (13.0)</td>
</tr>
<tr>
<td>COSMIC</td>
<td>WGS</td>
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<td>0/345 (0)</td>
<td>10/162 (6.2)</td>
<td>4/48 (8.3)</td>
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<tr>
<td>Pitt</td>
<td>RT-PCR</td>
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<td>4/89 (4.5)</td>
<td>4/47 (10.1)</td>
</tr>
<tr>
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<td>RT-PCR</td>
<td>245</td>
<td>2/45 (4.4)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td>942</td>
<td>21/337 (6.2)</td>
<td>7/71 (9.9)</td>
<td>5/43 (11.6)</td>
</tr>
</tbody>
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*BL1 and BL2, basal-like 1 and 2; M, mesenchymal; LAR, luminal androgen receptor.

†Only four \textit{BCL2L14–ETV6} cases from the Pitt cohort are analyzed for pathological features, which are not counted in the overall frequencies in necrotic TNBC.

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Fig. 2. Characterization of the BCL2L14-ETV6 fusions in 134 triple-negative breast tumors from two different patient cohorts. (A) RT-PCR analyses of BCL2L14-ETV6 fusion and wtETV6 in triple-negative tumors from the Pitt cohort (n = 89). A five-donor normal breast pool (NB) was used as a negative control. Representative gel images are shown. Fusion-positive cases are marked with red asterisks. The chromatograms (Lower) show the junction sequences of BCL2L14-ETV6 fusion variants detected in Pitt-TN49, Pitt-TN134, Pitt-TN138, and Pitt-TN144 tumors. (B) RT-PCR analyses of BCL2L14-ETV6 fusion, wild-type BCL2L14, wild-type ETV6, and GAPDH in 45 triple-negative breast tumors from the BCM cohort. A five-donor normal breast pool (NB) was used as a negative control. Fusion-positive cases are labeled with red asterisks. Chromatograms (Lower) show the junction sequences of BCL2L14-ETV6 fusion variants detected in BCM-TN13 and BCM-TN35 tumors. (C) Genomic PCR analysis of the BCL2L14-ETV6 TNBC tumor samples from the BCM cohort (BCM-TN13 and BCM-TN35) identified the precise genomic fusion points. (Left) Schematic of the genomic breakpoints identified in BCM-TN13 and BCM-TN35 tumors. (Right) Gel images and chromatograms of BCL2L14-ETV6 genomic PCR products. Genomic DNA from MCF10A cells was used as a negative control.

against BCL2L14-ETV6 that can be used for immunofluorescence, we performed fractionation of the fusion overexpressing cells and detected the fusion protein localizations by Western blots. Interestingly, the E2E3 and E4E2 fusion proteins tend to be enriched in the cytoplasmic fraction, while wtETV6 mainly presents in the nucleus, in line with its role as a transcription factor. The E4E3 fusion that expresses the truncated BCL2L14 protein was found to be enriched in the cytoplasm as well (Fig. 3E). Differential localization of the fusion proteins from wtETV6 suggests that BCL2L14-ETV6 fusion proteins may
function in a distinct cellular mechanism compared to wtETV6. It is possible that the BCL2L14 portion of the fusion variants may promote cytoplasm localization of the fusion proteins.

**BCL2L14–ETV6 Endows Enhanced Invasiveness and Paclitaxel Resistance.** We sought to examine the function of the BCL2L14–ETV6 fusion in the engineered BT20 and MCF10A cell lines. Among TNBC cell lines, BT20 is a nonmetastatic, chemo-sensitive line (27, 28) overexpressing E-cadherin (29). We thus selected this line for studying the more aggressive and chemo-resistant phenotypes driven by this fusion. MCF10A is an immortal but untransformed human mammary epithelial cell (HMEC) line. Both MCF10A and BT20 cell lines express endogenous ETV6 and BCL2L14 proteins (Fig. 3 B and C).

Transwell migration and invasion assays revealed that ectopic expression of the E2E3, E4E3, or E4E2 fusion variants but not wtETV6 significantly enhanced cell motility and invasion in BT20 cells, when compared to vector control (Fig. 4A). Similarly, enhanced cell motility and invasion (Fig. 4B) were also observed in the engineered MCF10A cells expressing these fusion variants.

On the other hand, ectopic expression of the fusion variants in BT20 cells did not result in significant changes in cell viability or cell cycle progression, whereas the wtETV6-expressing BT20 cells showed decreased viability and increased G0/G1 phase (SI Appendix, Fig. S12).

Taxane-based chemotherapy remains the cornerstone for the treatment of TNBC patients; however, the effectiveness is severely limited by intrinsic and acquired resistance. Since BCL2L14–ETV6 most frequently present in the mesenchymal TNBC tumors that are relatively resistant to chemotheraphy (30), we explored the role of BCL2L14–ETV6 in chemoresistance. We first treated the engineered BT20 cells with various doses of paclitaxel, a widely used taxane drug for TNBC patients. BCL2L14–ETV6 fusion-expressing BT20 cells displayed modest reduced sensitivity to paclitaxel following short-term (72 h) treatment, compared to the vector- or wtETV6-expressing cells (SI Appendix, Fig. S13A). We then tested the effect of low-dose prolonged paclitaxel treatment to observe acquired resistance. Following paclitaxel treatment for 1 mo, BT20 cells expressing wtETV6 or vector control were almost eradicated, whereas all
Thus examined the changes in apoptosis biomarkers following paclitaxel treatment. The BT20 cells overexpressing BCL2L14–ETV6 fusions did not show evident reduced apoptosis compared to wtETV6-expressing cells (SI Appendix, Fig. S13B). This suggests that the paclitaxel resistance driven by this fusion may not be attributed to the apoptotic pathway.

**BCL2L14–ETV6 Fusions Induce Distinctive Expression Changes from wtETV6.** To systematically profile the expression changes induced by BCL2L14–ETV6, we performed transcriptome sequencing of BT20 cells stably expressing the vector, wtETV6, or BCL2L14–ETV6 variants. Principal component analysis (PCA) revealed that the vector- and wtETV6-expressing cells form distinctive and independent clusters, whereas the BT20 cells expressing the different fusion variants are clustered together far from both the vector- and wtETV6-expressing cells (Fig. 5A). Furthermore, hierarchical clustering analysis revealed that the engineered BT20 cells were clustered into two main clusters, with the vector control or wtETV6-expressing cells as one major cluster and fusion-expressing cells as the other major cluster (Fig. 5B). These data suggest that BCL2L14–ETV6 fusions induced distinct gene-expression changes from wtETV6 and vector control in BT20 cells. It is interesting to note that while the E4E3 fusion variant encodes the C-terminus–truncated BCL2L14 protein, this variant induced a similar pattern of expression changes as the E2E3 and E4E2 variants that encode chimeric BCL2L14–ETV6 protein, suggesting that these distinct fusion variants may play a coherent functional role. To identify the pathways characteristic of BCL2L14–ETV6-expressing BT20 cells, we performed gene set enrichment analysis (GSEA) comparing the three fusion variants with the vector control in pairwise. Interestingly, the EMT pathway, known to promote paclitaxel resistance and invasiveness, is among the top up-regulated pathways in BT20 cells expressing BCL2L14–ETV6 (SI Appendix, Fig. S14 and Datasets S7 and S8). Among the core enrichment genes, 73 EMT pathway genes were up-regulated in the fusion-expressing BT20 cells (Fig. 5C and Dataset S9). These results indicate that BCL2L14–ETV6 fusions may induce up-regulation of EMT gene signature. To investigate the transcriptional regulatory mechanisms that regulate the EMT gene signature driven by BCL2L14–ETV6, we constructed breast cancer cell line BT20-specific transcriptional regulatory network using the ARACNe algorithm (31), and performed master regulator analysis (MRA). Among the 13 predicted master regulator candidates, SNAI2 is an established EMT-inducing transcription factor (SI Appendix, Fig. S15). The snail family genes SNAI1 (also denoted as SNAIL) and SNAI2 (also denoted as SLUG) are known to activate EMT and repress epithelial genes in tumors, including in breast cancer (29, 32).

**BCL2L14–ETV6 Fusions Prime EMT.** Next, we explored the expression of EMT biomarkers in the engineered MCF10A cells and BT20 cells by Western blots, including E-cadherin, N-cadherin, and vimentin. Loss of E-cadherin represents the first step of EMT transition (33). Both MCF10A cells and BT20 expressing vector control strongly express E-cadherin, suggesting their epithelial states (Fig. 5D–F). In fusion-expressing MCF10A cells, the expression level of E-cadherin is repressed, whereas the expression level of vimentin, an end-stage marker in EMT (34), and SNAI2 and its family member SNAI1 in fusion-expressing MCF10A cells. In the engineered BT20 cells, E-cadherin is repressed in all fusion-expressing models, but not in the wtETV6 model. Up-regulation of N-cadherin and SNAI1/ SNAI2 were also observed in fusion-expressing BT20 cells; however, there is no induction of vimentin following fusion overexpression (Fig. 5E). The fusion-specific induction of SNAI1/2 transcriptional factors and EMT markers became

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**Fig. 4.** Ectopic expression of BCL2L14–ETV6 endows increased cell migration, invasion, and paclitaxel resistance. (A and B) Ectopic expression of BCL2L14–ETV6 fusion variants in BT20 TNBC cells (A) and MCF10A benign mammary epithelial cells (B) significantly enhanced cell migration as revealed by Boyden chamber assay (Left), and increased cell invasion as revealed by transwell Matrigel assay (Right), relative to the vector control. Results were summarized from experimental triplicates. (C) BCL2L14–ETV6 fusions endows clonal resistance in BT20 cells following prolonged paclitaxel treatment for one month as shown by clonogenic assay. Here a low dosage of 5 nM paclitaxel is used for treatment to observe long-term treatment effect. (D) BCL2L14–ETV6 fusions endows clonal resistance in MCF10A cells following prolonged paclitaxel treatment for 1 mo as shown by clonogenic assay. Here 15 nM paclitaxel is used for treatment since MCF10A is less sensitive to paclitaxel. The quantitative results in the Upper panels of C and D are based on two replicates of each condition. The vehicle-treated cells were harvested in 14 d for the BT20 model, and 7 d for the MCF10A model, while the PTX-treated cells were harvested in 1 mo due to their different growth rates. The comparing cell models (i.e., vector, wtETV6, fusion variants) were harvested at the same time point. PTX: paclitaxel; Vehicle: 0.1% dimethyl sulfoxide. *P < 0.05; **P < 0.01; ***P < 0.001, significance was determined using Student's t test (two-tailed) and error bars reflect mean ± SD.
more obvious when the BT20 cells were treated with TGFβ-1 and EGF, known to induce EMT (35) (Fig. 5F). Loss of the epithelial marker E-cadherin and gain of one of the mesenchymal markers, N-cadherin or vimentin, in MCF10A or BT20 cells suggest that the cells are likely having partial instead of full activation of EMT.

Since EMT is often associated with stemness properties (36) known to promote clonal chemoresistance (37), we examined the expression of the known stemness biomarkers for breast cancer, CD44 and ALDH1A3 (38), in the BT20 models. Our RNA-seq data revealed increased expression of CD44 and ALDH1A3 in fusion-expressing BT20 cells compared to vector or wtETV6-expressing BT20 cells (SI Appendix, Fig. S16A). Consistently, flowcytometry analysis revealed higher number of CD44+/ALDH1A3+ cell populations in fusion-expressing BT20 cells, compared to vector or wtETV6 controls (SI Appendix, Fig. S16 B and C). Together, these results support the role of BCL2L14–ETV6 in inducing partial EMT in TNBC cells.
Discussion

TNBC comprises 10 to 20% of all breast cancers. Due to lack of well-defined molecular targets, treatment of TNBC tumors relies on taxane and platinum-based chemotherapies. Despite the distinctive receptor status, recent genomic sequencing studies have revealed a paucity of TNBC-specific mutations, apart from a distinctive mutational enrichment pattern from other breast cancers, such as more frequent TP53 mutations and less frequent PIK3CA mutations (39). While recent transcriptomic and genomic sequencing studies have revealed oncogenic gene fusions in TNBC patients, some of these may be non-recurrent and might be considered individual fusions, such as MAGI3–AKT3 and FGFR3–TACC3 (40–42), whereas others tend to fuse with promiscuous partners, such as Notch and MAST fusions, which may be considered as gene family fusions (43). To date, canonical gene fusions of the same fusion partners that recur in a significant subset of TNBC patients have not been reported. Identification of TNBC-specific genetic events that could guide the treatment decisions in this aggressive subtype of breast cancer represents an unmet clinical need.

Despite the complexity and heterogeneity of structural rearrangements in breast cancer (7, 13), our systematic analyses of somatic structural rearrangements based on WGS data cataloged 99 recurrent gene fusions in breast cancer. Among the different types of rearrangements, we found that AGR represents a special type of cryptic rearrangement that may occur more frequently than realized in breast cancer. Such cryptic genomic changes are hardly detectable by conventional cytogenetic assays or by transcriptome sequencing. For these reasons, AGRs can only be confidently detected from WGS datasets. Further studies revealed that the top recurrent AGRs are more frequently enriched in specific more aggressive forms of breast cancer that lack well-defined drivers, such as basal or luminal B breast cancer. These AGRs tend not to aggregate in the genomically unstable tumors, suggesting that they are potential pathological events instead of merely the consequence of genomic instability. Among the top four confirmed recurrent gene rearrangements (BCL2L14–ETV6, AKA8P–BRD4, TTC6–MIPOL1, and ESR1–CCDC170) BCL2L14–ETV6 is frequently and specifically detected in TNBC, and with which we chose to perform further functional studies. For the TTC6–MIPOL1 rearrangement, while the tandem duplication delineating this fusion encompasses the immediately proximal FOXA1 gene, it is unlikely that one copy number gain can significantly enhance FOXA1 expression. In addition, two of four TTC6–MIPOL1+ TCGA tumors do not exhibit copy number changes in the FOXA1 locus (SI Appendix, Fig. S1B). Future studies will be required to further evaluate the function of this fusion in luminal breast cancer.

Next, we performed in-depth functional studies on the BCL2L14–ETV6 fusion. We first experimentally validated this fusion in two independent TNBC patient cohorts, which identified 6 BCL2L14–ETV6+ cases of a total of 134 TNBC cases. Taking together WGS data and RT-PCR validation results, this fusion was detected in 4.4 to 12.2% of TNBC tumors (with an average of 6.2%) from four independent patient cohorts (Table 1). Further investigation of histopathological associations in the TCGA and COSMIC cohorts revealed that this fusion is preferentially present in the TNBC tumors with gross necrosis and more aggressive histopathological features, such as marked nuclear pleomorphism, numerous mitoses, and high tumor grade (Fig. 1F). Such association is further verified by evaluating pathological slides for the fusion-positive cases from the Pitt cohort. All of these cases are grade III TNBCs with extensive or focal necrosis. It is interesting to note that RT-PCR of wtETV6 also revealed ETV6 exon duplications in TNBC cell lines or PDX tumors. These include exon 2 duplication of ETV6 in two PDX tumors and in HCC1187, and exon 4 duplication of ETV6 in one PDX tumor (SI Appendix, Fig. S8). This suggests that ETV6 genetic aberrations could involve both intergenic and intragenic rearrangements. Future studies will be required to explore the function of ETV6 exon duplications in TNBC.

While it remains to be addressed whether DNA repair deficiency may promote the formation of this fusion, our biological studies suggest that BCL2L14–ETV6 fusions appear to enhance cell mobility and invasiveness, and promote paclitaxel resistance when ectopically expressed in basal-like HMEC cell line and nonmetastatic chemo-sensitive TNBC cell line models. In addition, transcriptome sequencing revealed that despite encoding distinct protein products, the three fusion variants induced a coherent transcriptional program that is distinctive from wtETV6. Of note, while TCGA copy number data suggest genomic amplifications of the ETV6 genomic loci in a subset of breast tumors harboring BCL2L14–ETV6 tandem duplications (SI Appendix, Fig. S1A), ectopic overexpression of wtETV6 did not elicit increased cell migration, invasion, or paclitaxel resistance in TNBC cells (Fig. 4). It is possible that the observed genomic amplifications could be secondary events following formation of this fusion to enhance its function. In our previous study we leveraged the secondary amplifications of oncogenic fusions to identify pathological recurrent fusions (44).

Furthermore, our data suggest that the breast cancer cells overexpressing BCL2L14–ETV6 show a characteristic enrichment of EMT signature. EMT is known to confer stemness features and thus induce invasiveness and chemoresistance in TNBC (32, 45). Interestingly, our data suggest that BCL2L14–ETV6 fusion proteins may prime for partial EMT instead of full activation of EMT. Tumor cells in partial EMT state are in a state of plasticity that favors metastasis and chemoresistance (46), and are frequently observed in TNBC (47). Consistently, BCL2L14–ETV6 fusions are most frequently detected in the mesenchymal subtype of TNBC tumors that is closely associated with EMT (23, 30). In this study, we compared the function of BCL2L14–ETV6 with wtETV6 as the major fusion variant E4–E2 and E2–E3 retain most of the ETV6 domains, whereas the C-terminal–truncated BCL2L14 portion lacks intact BCL2-like domain. Furthermore, the paclitaxel resistance driven by this fusion does not seem to be attributable to the changes in apoptosis signaling (SI Appendix, Fig. S1B). Nonetheless, future studies will be required to elucidate the function of the BCL2L14 portion in the fusion.

While it will be interesting to study the endogenously expressed fusion protein in the BCM-2147 PDX model, technical difficulties exist for genetic inhibition studies in many PDX tumors, including BCM-2147. First, the knockdown studies will require rescue experiments to verify the specificity of the siRNAs, which need to be performed on stable cell lines. There are no fewer than six laboratories attempting to generate cell lines from our BCM PDX models, including laboratories that have generated stable cell lines from primary tissue previously. Thus far, it has not been possible to generate cell lines from any PDX model tested. Although we have established methods for lentiviral transduction for shRNA-mediated knockdown in PDX, the transduction rate is about 30 to 50%, unlike established cell lines where the infection rate typically exceeds 95%. Given this low transduction rate, shRNA-mediated knockdown and genome editing with CRISPR is very inefficient. Furthermore, whereas a majority PDX models will retransplant under all of the dissociation conditions tested, future studies will be required to overcome these technical challenges and elucidate the function of the endogenous BCL2L14–ETV6 in vivo.

In summary, this study revealed adjacent gene rearrangements as a class of cryptic genetic events that is more frequent than realized in breast cancer. Future investigation of molecular mechanisms of the pathological AGRs, such as BCL2L14–ETV6, may pave the way to new precision medicine against these genetic targets and improve the clinical outcome. Furthermore, immune checkpoint inhibition was most recently found to

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significantly prolong the survival of metastatic TNBC patients (48). The potential benefit of immunotherapy in the BCL2L14-ETV6\(^{\ddagger}\) patient cohort merits future investigations.

### Materials and Methods

To systematically characterize recurrent AGRs in breast cancer, we analyzed the somatic structural mutation data catalogued by the ICGC based on WGS data for 215 breast tumors. To detect BCL2L14-ETV6 fusion transcripts, we designed a pair of primers located on exon 2 of BCL2L14 and the last exon of ETV6, respectively, and performed RT-PCR on 134 triple-negative breast tumors, including 45 tumors procured from the Tumor Bank at Baylor College of Medicine and 89 tumors procured from the Health Sciences Tissue Bank of University of Pittsburgh. The primer sequences and PCR conditions are provided in Dataset S10. The full-length cDNAs of BCL2L14-ETV6 fusion variants (E233, E433, and E462) were amplified from fusion-positive tumors, and engineered into a lentiviral pLent7.3 vector (Invitrogen). BCL2L14-ETV6 protein products were detected by Western blots and the antibodies are provided in Dataset S11. Transwell migration and Matrigel invasion assays were performed to assess cell invasiveness, and clonogenic assays were performed to assess cell viability following paclitaxel treatment. Transcriptome sequencing of the engineered BT20 cells was performed on the NovaSeq 6000 system. The RNA-seq data are made available through Gene Expression Omnibus (GSE120919). The protocols, codes, and materials used in this study are available upon request to the corresponding author.

More detailed methods are provided in SI Appendix, Supplementary Materials and Methods.

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