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Anti-EGFR VHH-armed death receptor ligand–engineered allogeneic stem cells have therapeutic efficacy in diverse brain metastatic breast cancers

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Basal-like breast cancer (BLBC) shows brain metastatic (BM) capability and overexpresses EGFR and death-receptors 4/5 (DR4/5); however, the anatomical location of BM prohibits efficient drug-delivery to these targetable markers. In this study, we developed BLBC-BM mouse models featuring different patterns of BMs and explored the versatility of stem cell (SC)–mediated bi-functional EGFR and DR4/5-targeted treatment in these models. Most BLBC lines demonstrated a high sensitivity to EGFR and DR4/5 bi-targeting therapeutic protein, E_vDR_L [anti-EGFR VHH (E_v) fused to DR ligand (DR_L)]. Functional analyses using inhibitors and CRISPR-Cas9 knockouts revealed that the E_v domain facilitated in augmenting DR4/5-DR_L binding and enhancing DR_L-induced apoptosis. E_vDR_L secreting stem cells alleviated tumor-burden and significantly increased survival in mouse models of residual-tumor after macrometastasis resection, perivascular niche micrometastasis, and leptomeningeal metastasis. This study reports mechanism based simultaneous targeting of EGFR and DR4/5 in BLBC and defines a new treatment paradigm for treatment of BM.

INTRODUCTION

Breast cancer (BC) is the second most common cancer that can metastasize to the brain and, in fact, brain metastasis (BM) is a major cause of cancer-related deaths in patients with BC. Approximately 15 to 30% of patients with metastatic BC develop BM (1, 2). Along with the increase in the incidence of BC (3), the occurrence of BC-BM has also increased in recent years owing to improved extra-cranial disease control and poor central nervous system (CNS) penetration of drugs (4). Among the four main intrinsic subtypes of BC, basal-like breast cancer (BLBC) has a 70 to 80% overlap with triple-negative breast cancer (TNBC) and constitutes 12 to 15% of BC (5). BLBC has the poorest prognosis and the shortest survival among the BC subtypes (6), owing to the unavailability of specific therapeutic options including hormonal or molecular-targeted therapy. BLBC metastasizes to the brain more frequently than the other subtypes (7, 8), shortening patient survival (9).

BM continues to represent a formidable challenge in the clinical management of patients with cancer. Currently, stereotactic radiosurgery (SRS), surgical resection, and whole-brain irradiation are the most common treatment options for BM; however, these tumors are generally resistant to systemic chemotherapy because of the blood-brain barrier (BBB). For detectable tumors up to 3 cm, there is a favorable indication for SRS (10). However, residual invasive cancer

cells following surgical resection of large tumors that were not eligible for SRS, undetectable dormant tumor cells in the perivascular niche (PVN), and leptomeningeal metastasis, also known as meningeal carcinomatosis, are challenging conditions and lack effective treatment options. Tumors in the eloquent areas of the brain are unresectable, and residual tumor cells after resection eventually cause recurrence. Because, compared to other cancers, BC is known to feature later recurrences (11), patients who have undergone treatment remain at a persistent risk even for decades (11, 12). Repeated chemotherapy is often unable to kill the PVN-dwelling cancer cells as they are known to establish a strict localization outside the vasculature (13, 14). Leptomeningeal metastasis is another devastating condition observed in 11 to 20% of patients with CNS metastasis of BC (15, 16). The lack of effective treatments leads to extremely short survival (median survival: 4 to 6 weeks) (17). Although intrathecal (IT) administration of anticancer agents is often attempted, there is no evidence for improvement in survival (18). Given the multistep and complex biological nature of BM, tumor models that recapitulate metastatic brain tumor features are limited. In this study, we first developed imageable mouse models for clinically challenging BLBC-BM conditions, including single intracranial metastasis resection, PVN micrometastasis, and leptomeningeal metastases, and used these models to investigate in detail the efficacy and mechanism of stem cell–based bifunctional BM treatments targeting over-expressed epidermal growth factor receptor (EGFR) and death receptors 4/5 (DR4/5).

EGFR is up-regulated in BLBC (19), and anti-EGFR therapies have been adopted (20). Given that EGFR is one of the most important mediators of BM in BC (21), and EGFR positive tumors are more frequent in BC-BM than primary tumors (22), targeting EGFR has the potential to be beneficial in BC-BM. However, in contrast to other EGFR–up-regulated cancers, EGFR-targeting therapies have not been successful in BLBC (20). On the other hand,

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BLBC cell lines are sensitive to DR4/5-induced death signaling triggered by tumor necrosis factor-related apoptosis-inducing ligand (TRAIL, herein as DR_L) binding, which induces apoptosis selectively in cancer cells (23). However, DR_L alone is not sufficient to treat BLBC (24), and even enhancing modifications of DR_L have not translated into remarkable treatment benefits in patients (25). In a preliminary screen of BLBC-BM patient samples, EGFR and DR4/5 displayed a concurrent up-regulation in the tumor area. Therefore, it is of great interest to explore the strategy to simultaneously evaluate targeting EGFR and DR4/5 in BLBC-BM. A variable domain of camelid heavy-chain-only antibody (VHH), also known as nanobody is a small molecule, consisting solely of the antigen binding domain (26). We have previously engineered bivalent anti-EGFR VHH (E_V) and shown their efficacy in mouse tumor models (27). We have recently developed a bi-functional molecule E_V fused to DR_L (E_VDR_L) that simultaneously targets EGFR and DR4/5, but its mechanism-based efficacy has not been fully understood.

Given the challenges related to systemic delivery of a majority of therapeutic agents across the BBB and short half-life and high hepatotoxicity of DR_L (28), engineered stem cells offer an excellent platform to target CNS tumors. We and others have previously established use of neural stem cells (NSCs) and mesenchymal stem cells (MSCs) engineered to express tumor-specific biomolecules for treating primary brain neoplasms. Especially for tumors in the CNS, stem cell administration in the resection cavity has been shown to improve drug delivery (29). For micrometastasis at PVN, arterial delivery of therapeutic stem cells offers an advantage owing to the BBB penetration capability of stem cells (30–32). For leptomeningeal metastasis, the primary reason for the failure of IT drug administration is the difficulty of infusing drugs continuously and the incessant turnover of cerebrospinal fluid (CSF) that clears the drug (17). IT delivery of therapeutic stem cells offers an advantage as stem cells can survive in the CSF space and continuously secrete therapeutic molecules; however, there are no reports on stem cell therapy for leptomeningeal metastasis as a secondary CNS tumor. In this study, we characterized in detail the anti-BLBC efficacy of E_VDR_L and assessed the therapeutic efficacy of stem cell-delivered E_VDR_L in different mouse tumor models of breast to BM.

RESULTS

EGFR and DR4/5 are up-regulated in BLBC-BM

We analyzed the dataset from The Cancer Genome Atlas (TCGA) (33) and showed that BLBC [typically triple negative for estrogen receptor, progesterone receptor, and HER2 (TNBC); fig. S1A] has significantly higher expression of *EGFR* and *DR4/5* mRNA compared to the other BC subtypes (Fig. 1A). In addition, a cohort of cell lines from TCGA (34) provided further evidence that BLBC cell lines have significantly higher expression of *EGFR* and *DR5* mRNA compared to the non-BLBC subtypes (Fig. 1B). To confirm these results, we tested cellular and cell surface expression of EGFR and DR4/5 by Western blot and flow cytometry, respectively, in 15 human BC cell lines (HER2-enriched: SKBR3 and MDA-MB-453; luminal A: MCF7, HCC1500, ZR75-1, and HCC1428; luminal B: BT474, T47D, and MDA-MB-175VII; basal-like: BT549, Hs578T, SUM159, MDA-MB-231, MDA-MB-436, and MDA-MB-468) and three patient-derived BLBC-BM cell lines (BMET02, BMET05, and BMET15), which were confirmed by mRNA microarray (fig. S1B) (35). BLBC

cell lines showed significantly higher expression of EGFR compared to the other BC subtypes (Fig. 1, C and D). Although higher expression of DR5 was observed in BLBC as compared to non-BLBC, the expression levels were not significant, most likely due to the insufficient number of cell lines tested (Fig. 1, C and D). In addition, immunohistochemistry of TNBC patient samples showed that BM tissue displayed a significantly higher expression of EGFR compared to the primary tumor (Fig. 1E and fig. S1C). Together, these data revealed that EGFR and DR4/5 are promising targets in BLBC-BM.

Development and characterization of mouse tumor models of BLBC-BM

We developed mouse models representing three major clinically relevant forms of BM: macrometastasis, micrometastasis, and leptomeningeal metastasis (Fig. 2A). First, we generated a patient-derived, BMET02 line expressing a bimodal firefly luciferase (Fluc)-mCherry (FmC) fusion protein (fig. S2A). We confirmed a direct correlation between Fluc signals and implanted BMET02-FmC cell numbers in vivo and show that BMET02-FmC has similar growth rate as parental BMET02 (fig. S2, B and C). Intracardiac injection, the most common method for development of experimental BM models, can lead to widespread tumor formation (36). Even “standard” intracarotid arterial (ICA) injection may reduce the rate of aberrant (nonbrain) metastasis, this route routinely requires the ligation of the external carotid artery, thereby prohibiting repeated injections. Moreover, when using the standard ICA injection technique with ligation of the external carotid artery, we still observed extracranial metastasis, likely as a result of tumor cell distribution into small arterial feeder branches, such as the occipital artery (OA), pterygopalatine artery (PPA), and superior thyroid artery (STA) (fig. S3, A to C) (37–39). Therefore, we established a modified ICA injection technique, in which we ligated these feeder arteries, thereby greatly reducing the formation of extracranial tumors (fig. S3A and movie S1). In addition, partial preservation of the external carotid artery enabled multiple ICA injections from the same side; this allows us to efficiently test cell-based therapies in this model (movie S1). We injected BMET02-FmC into mice using this modified ICA injection technique and monitored tumor development by bioluminescence imaging (BLI). BLI signals of day 0 demonstrated the successful ICA injection of viable BMET02-FmC cells in mouse; however, the BLI signals then quickly dropped to undetectable levels, most likely due to only a very small portion of tumor cells completing extravasation and surviving in the brain (Fig. 2B). Chronological brain samples from ICA-injected BMET02-bearing mice showed multiple tumors in the brain parenchyma (Fig. 2C). Immunohistochemistry of brain blood vessels revealed that BC cells extravasated in the early phase (day 7), stayed alongside the blood vessel for a while (day 13), and then started growing along the vessels (day 20) (Fig. 2D). Immunohistochemistry showed late phase BLBC-BM tumors were highly proliferative and hypervascularized and surrounded by astrocytes (fig. S3D).

To develop a leptomeningeal metastasis model, we IT injected BMET02-FmC into the cisterna magna (fig. S3E). BLI showed tumor growth around the CNS (Fig. 2E). Brain and spine samples showed widely disseminated tumors in various areas of CSF space across the CNS (Fig. 2F). The tumor resection model was developed as we previously reported (Fig. 2G) (40). In short, following establishment of a cranial window, tumor cells were directly injected into

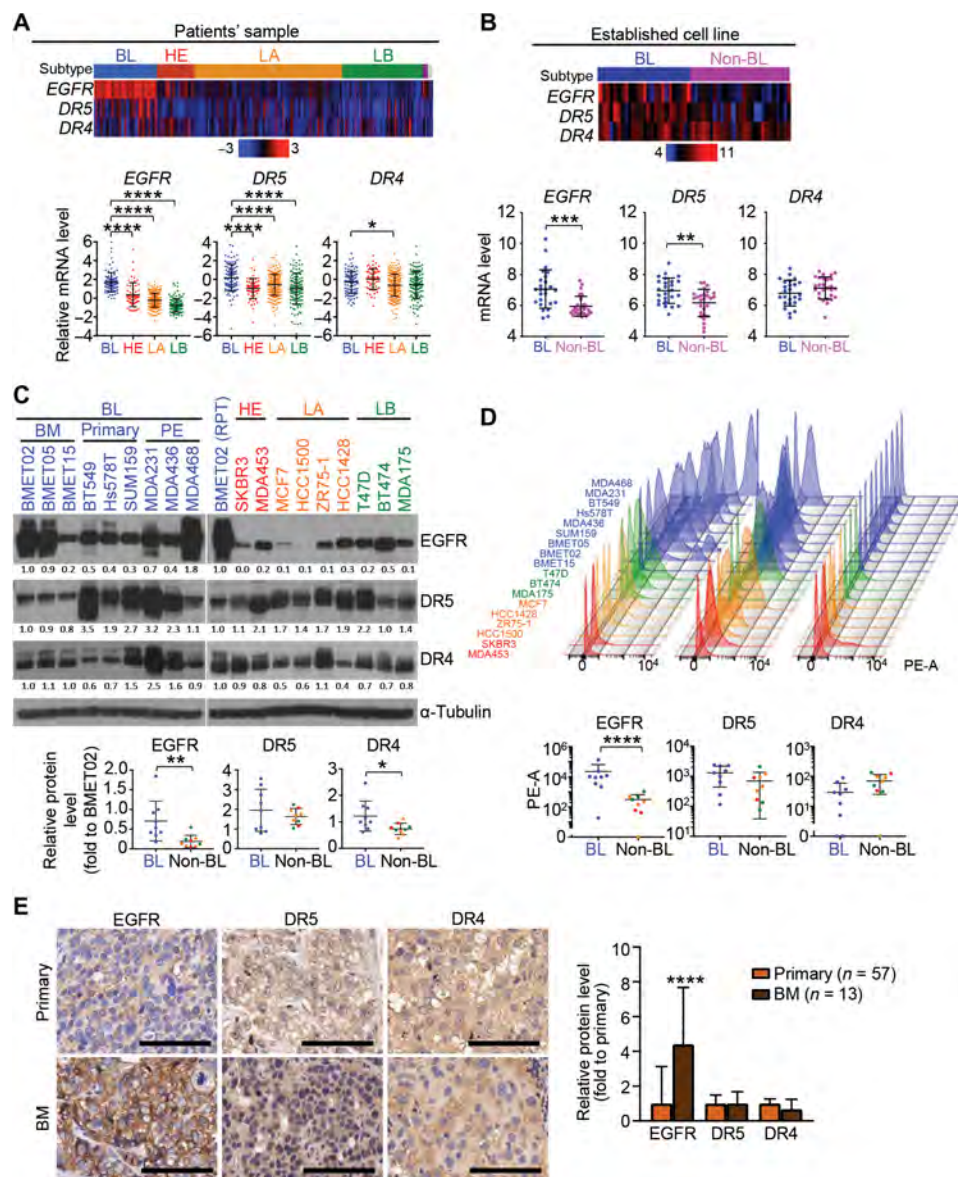


Fig. 1. EGFR and DR4/5 are up-regulated in BLBC-BM. (A) Top: Heatmap of mRNA levels of *EGFR*, *DR5*, and *DR4* in patient samples of four subtypes (BL, basal-like; HE, HER2-enriched; LA, luminal A; LB, luminal B) of BC from TCGA database ($n = 526$). Bottom: Comparison of *EGFR*, *DR5*, and *DR4* mRNA levels between subtypes. (B) Top: Heatmap of mRNA levels of *EGFR*, *DR5*, and *DR4* in cell lines of BLBC or non-BLBC from TCGA database ($n = 52$). Bottom: Comparison of *EGFR*, *DR5*, and *DR4* mRNA levels between subtypes. (C) Top: Western blot (WB) of *EGFR*, *DR5*, and *DR4* in 18 BC cell lines (PE, pleural effusion; RPT, repeat; loading control-adjusted ratios are provided under blots). Bottom: Relative expression of *EGFR*, *DR5*, and *DR4* in BLBC and non-BLBC. (D) Top: Cell surface protein levels of *EGFR*, *DR5*, and *DR4* analyzed by flow cytometry in 18 BC cell lines. Bottom: Comparison of cell surface expression of *EGFR*, *DR5*, and *DR4* in BLBC and non-BLBC. PE-A, phycoerythrin-area. (E) Left: Representative micrograph of immunohistochemistry of *EGFR*, *DR5*, and *DR4* in primary tumors and BM of TNBC. Scale bars, 100 μ m. Right: Quantifications of immunohistochemical staining densities by ImageJ (primary, $n = 57$; BM, $n = 13$).

the superficial brain parenchyma followed by bioluminescent imaging of tumor growth and microscopically assisted tumor resection. To confirm that orthotopically injected BMET02-FmC leads to a similar pattern of metastatic tumor formation as we had previously observed with ICA-injected tumor cells, we performed ex vivo brain sectioning followed by immunohistochemical staining for vascularization and surrounding astrocytes (fig. S3F). Fluorescence images and BLI showed that resection substantially reduced tumor size (Fig. 2, H and I). Together, these data demonstrate the establishment of three clinically relevant mouse models of BLBC-BM,

which have the potential to facilitate development and preclinical testing of the next generation of BM therapies.

EGFR induces apoptosis in BLBC cells

Clinical tissue obtained from patients with TNBC-BM and tumor tissues from ICA-injected patient-derived BMET02-FmC-bearing mice expressed both of *EGFR* and *DR4/5* as revealed by immunofluorescence (Fig. 3A and fig. S4A). In addition, our flow cytometry analysis showed that each single tumor cell has both of *EGFR* and *DR5* using a BLBC cell line (fig. S4B). To simultaneously target cell

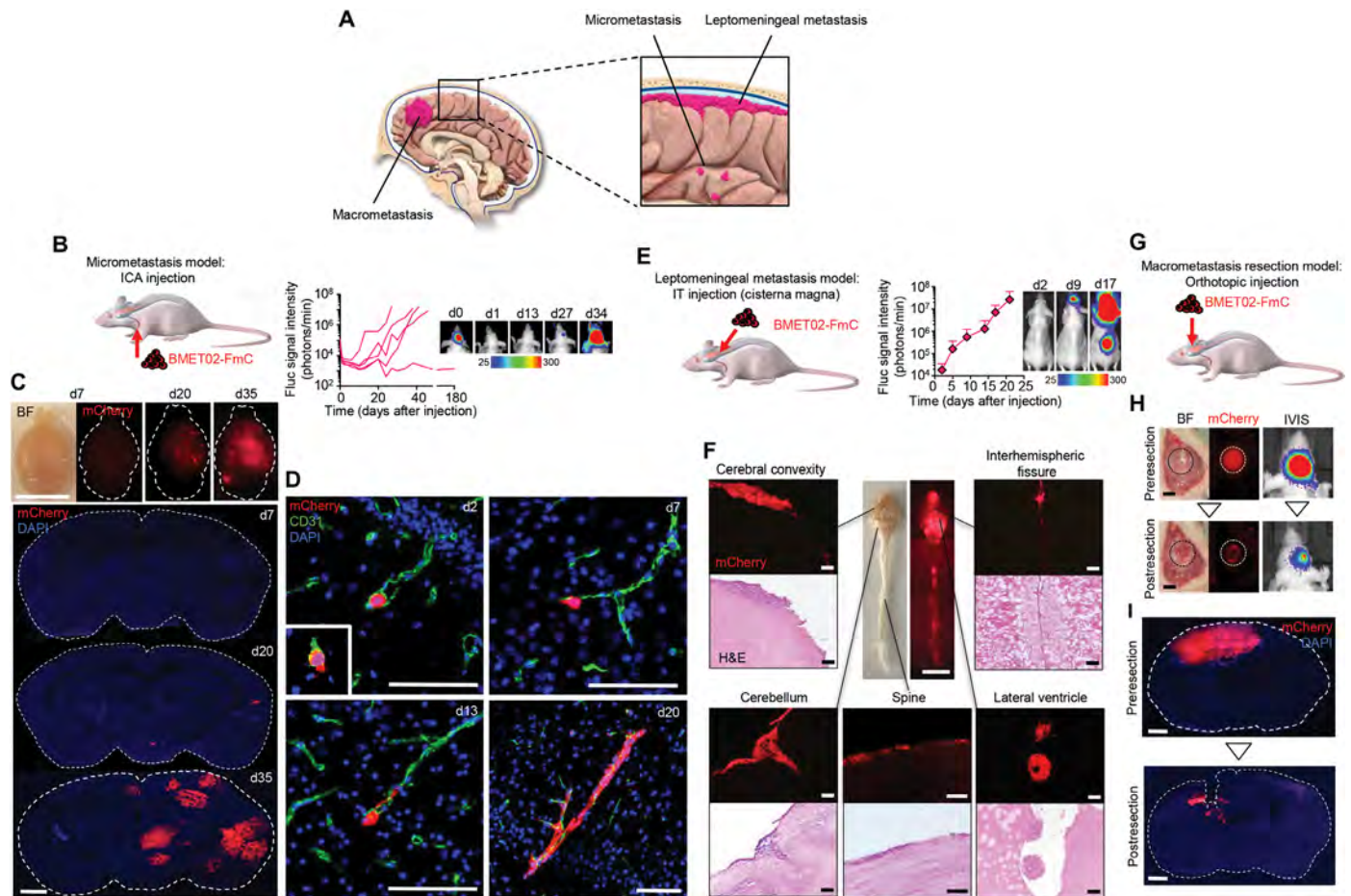


Fig. 2. Developing and characterizing clinically relevant mouse models for BLBC-BM. (A) Schematic representation of three clinical scenarios of BM. (B) Left: Schematic of micrometastasis model. Right: BLI signal curves of each mouse after ICA injection of BMET02-FmC and representative pictures. (C) Top: Chronological representative bright-field (BF) and fluorescence photographs of brain samples from ICA-injected BMET02-FmC-bearing mice. Scale bar, 10 mm. Bottom: Photomicrograph of coronal sections of the samples. Scale bar, 1 mm. (D) Chronological photomicrographs of immunohistochemistry of CD31 in brain sections that had ICA injection of BMET02-FmC. Scale bars, 100 μ m. Critical moment of extravasation of cancer cells was observed on day 2 (inset of d2). (E) Left: Schematic of leptomeningeal metastasis model. Right: BLI signal curve of IT-injected BMET02-FmC-bearing mice ($n = 2$) and representative photographs. (F) Center: Representative photograph of brain and spine samples from mice 23 days after IT injection of BMET02-FmC. Scale bar, 10 mm. Surrounding: Representative microphotograph of fluorescence and hematoxylin and eosin (H&E) staining of the brain and spine samples. Scale bars, 100 μ m. (G) Schematic of macrometastasis resection model. (H) Left: Representative intraoperative BF and fluorescence photographs of brain of pre- and postresection of BMET02-FmC tumor. Scale bars, 1 mm. Right: Representative pictures of BLI. (I) Representative photomicrograph of brain sections of pre- and postresection of tumor. Scale bars, 1 mm. Photo credit: Yohei Kitamura, Brigham and Women's Hospital. DAPI, 4',6-diamidino-2-phenylindole.

surface EGFR and DR4/5, we used two bi-functional proteins encoded by cDNAs for anti-EGFR VHH (E_V) or anti-EGFR ScFv (E_S) fused to DR_L (E_V DR_L and E_S DR_L, respectively) (Fig. 3B and fig. S4C) and tested their efficacy in patient-derived BLBC-BM lines. Cell viability and protein assays showed that E_V DR_L induced significant cytotoxicity and caspase-mediated apoptosis compared to E_S DR_L (Fig. 3, C and D). We further assessed the antitumor effects of E_V DR_L in 18 BC cell lines. Most BLBC cell lines responded to DR_L and E_V DR_L but not to E_V . E_V DR_L consistently mediated more potent antitumor effects than DR_L (Fig. 3E and fig. S4D). A correlation between the expression levels of DR5 (see Fig. 1D) and the efficacy of DR_L was observed (Fig. 3F), and therapeutic effects of E_V DR_L relative to those of DR_L were correlated with EGFR expression levels in BLBC cells (Figs. 1, C and D, and 3G and fig. S4E). These results suggest that the therapeutic sensitivity to E_V DR_L of BC cell is mainly determined by the expression levels of DR5 and EGFR (Fig. 3H).

To explore candidate factors besides EGFR and DRs that might influence BLBC's sensitivity to E_V DR_L, we first analyzed the difference in apoptosis-related genes among the subtypes based on the data from TCGA (fig. S5A). Among them, down-regulation of *BCL2* and *Bcl-xL* (anti-apoptotic Bcl genes) and up-regulation of *BID* and *BAX* (pro-apoptotic Bcl gene) were found in BLBC (fig. S5A). Western blot analysis on different BLBC cell lines revealed differential expression of *BCL2*, *Bcl-xL*, and *BID*. However, no significant correlation between the protein expression of *BCL-2*, *Bcl-xL*, *BID*, and efficacy of DR_L was observed (fig. S5B). Decoy death receptors, DcR1 and DcR2, are known to influence the sensitivity of cells to DR_L (41). TCGA revealed down-regulation of *DcR1* and up-regulation of *DcR2* in BLBC (fig. S5A), and Western blot analysis of BLBC lines showed low expression of DcR1 and varying expression levels of DcR2 (fig. S5C). Furthermore, *MYC*, which is known to be related to both of apoptosis pathway and EGFR signaling pathway (42, 43),

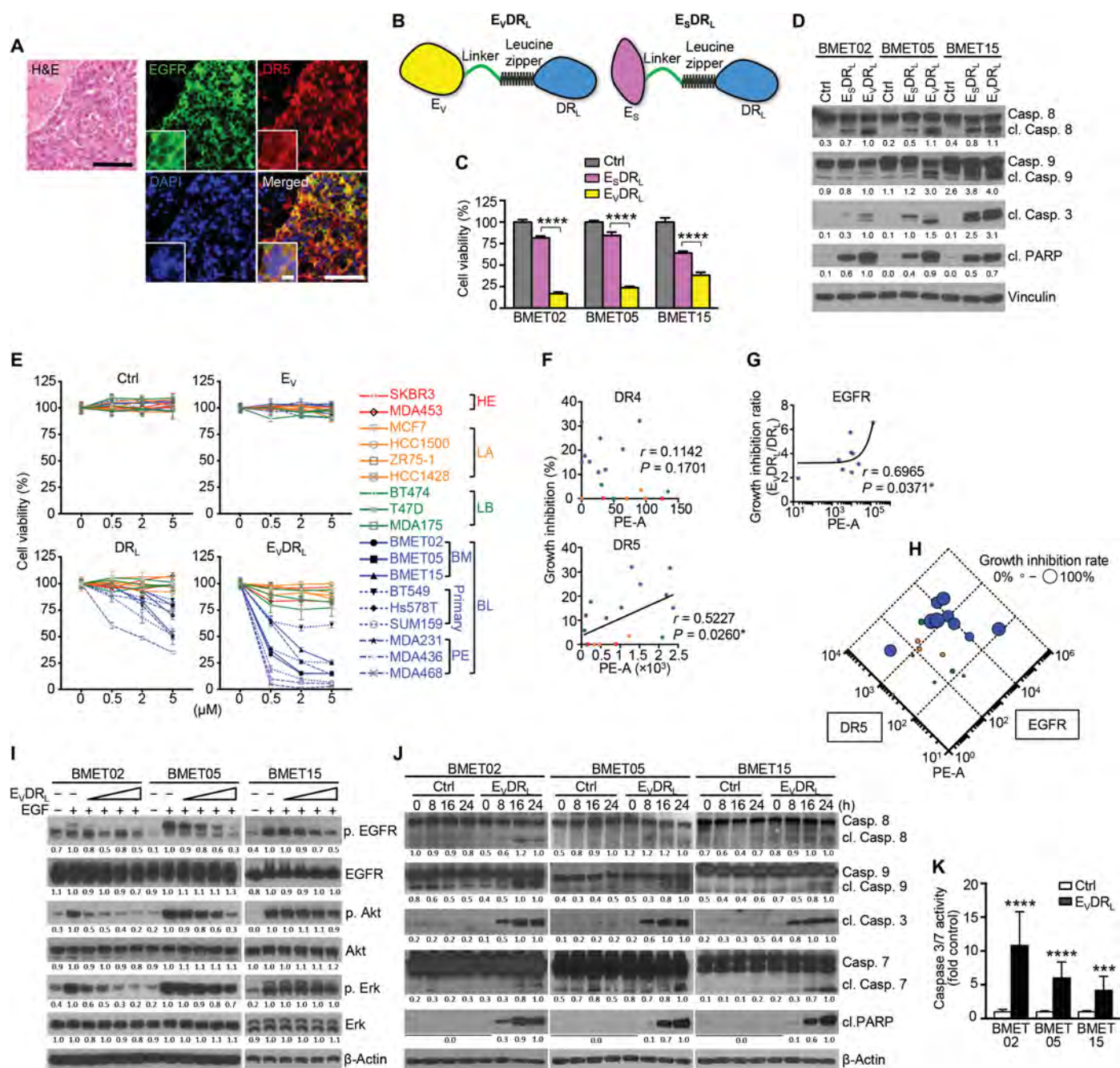


Fig. 3. BLBC is sensitive to E_vDR_L, and E_v domain of E_vDR_L enhances apoptosis-inducing effect of DR_L depending on tumor cell surface EGFR expression. (A) Representative microphotograph of low and high (the insets) magnifications of H&E and immunohistochemistry of EGFR and DR4/5 in patient samples of TNBC-BM. Scale bars, 100 μ m (main images) and 10 μ m (insets). (B) Schematic showing the construction of anti-EGFR VHH-DR_L (E_vDR_L) and anti-EGFR scFv-DR_L (E_sDR_L) proteins. (C) Cell viability of BLBC-BM lines after 72-hour treatment with control media (Ctrl), E_sDR_L, or E_vDR_L ($n=3$, technical replicates). (D) WB showing cleavage of caspases and poly(ADP-ribose) polymerase (PARP) in BLBC-BM lines after 8-hour treatment with Ctrl, E_sDR_L, or E_vDR_L ($n=3$, technical replicates). (Loading control-adjusted ratios are provided under blots; only cleaved part was quantified). (E) Cell viability of 18 BC cell lines after 72-hour treatment with different concentrations of Ctrl, E_v, DR_L, or E_vDR_L ($n=3$, technical replicates). (F) Correlation between cell surface DR4/5 expression and growth inhibition effect of DR_L at the time point of 24 hours. (G) Correlation between cell surface EGFR expression and growth inhibition ratio between DR_L and E_vDR_L at the time point of 24 hours. (H) Correlation between cell surface DR5 and EGFR expression and the growth inhibition efficacy of E_vDR_L. (I) WB showing phosphorylation of EGFR and its downstream elements in BLBC-BM lines with EGF treatment after pretreatment with various concentrations of E_vDR_L ($n=3$, technical replicates). (J) WB showing cleavage of caspases and PARP in BLBC-BM lines after 8-, 16-, and 24-hour treatment with Ctrl or E_vDR_L ($n=3$, technical replicates). (K) Caspase-Glo 3/7 assay of BLBC-BM lines after 8-hour treatment with Ctrl or E_vDR_L ($n=3$, technical replicates).

was down-regulated in BLBC cell lines (fig. S5A). Western blot analysis showed a negative correlation between protein expression of Myc and efficacy of E_VDR_L in the cell lines tested (fig. S5D). These results suggest that sensitivity of BLBC cells to E_VDR_L is mainly determined by EGFR and DR expression; however, other factors, such as apoptosis-related factors, might also influence their sensitivity to E_VDR_L.

As expected, we observed that E_VDR_L inhibited EGFR signaling (Fig. 3I) and induced caspase-mediated apoptosis in BLBC-BM tumor cells (Fig. 3, J and K). We also confirmed that E_V alone and E_SDR_L also inhibited EGFR signaling (fig. S5, E and F). These data showed that E_VDR_L simultaneously targets EGFR and DR4/5 and consistently induces apoptosis in a cohort of BLBC lines.

E_V domain of E_VDR_L enhances DR_L-induced apoptosis

We initially hypothesized that the main function of E_V domain of E_VDR_L would be to block EGFR signaling. However, E_V alone showed marginal effects on the cell viability of BLBC tumor cells expressing high levels of EGFR (Fig. 3E), and the differences in efficacy between DR_L and E_VDR_L were apparent very early ~24 hours after treatment (fig. S4D). This did not support our initial hypothesis and suggested that the E_V domain has another mechanism beyond blocking EGFR signaling to enhance therapeutic efficacy of E_VDR_L. To identify the mechanism, we tested whether E_V could sensitize tumor cells to DR_L by modulating interactions between apoptosis pathways and EGFR downstream elements. However, we did not observe any changes when the cells were treated with E_V (fig. S6A). Next, we combined E_V and DR_L to test whether this recapitulated the effects of E_VDR_L in BLBC-BM lines. Treatment with E_V + DR_L had a lower efficacy compared to E_VDR_L (Fig. 4, A and B), suggesting that fusing E_V with DR_L is necessary to enhance efficacy. Next, we tested the efficacy of E_VDR_L on another BLBC cell line, MDA-MB231-FmC, which has relatively low EGFR and high DR4/5. We observed similar findings in this line as well (fig. S6, B and C). Next, we assessed the proximity between EGFR and DR5 before and after treatment with E_VDR_L using real-time Förster resonance energy transfer (FRET) imaging on BMET02 and NIH-3T3 cells expressing recombinant EGFR–yellow fluorescent protein (YFP) and DR5–cyan fluorescent protein (CFP) (Fig. 4C). Both receptors were localized to the cell surface, and typical trace showed that treatment with E_VDR_L correlated with an increase in detectable FRET in the cells (Fig. 4D). These data suggest that EGFR and DR5 are in close proximity on the cell surface, and E_VDR_L binding further increases their association.

To confirm that E_VDR_L binding to EGFR and DR4/5 at the cell surface results in internalization of these receptors, we assessed the colocalization of EGFR with early, Rab5, and late, Rab7, regulators of endocytosis, posttreatment of cells with E_VDR_L. EGFR colocalized with early endosomal protein, Rab5, within 5 min and with late endosomal protein, Rab7, at 15-min time after E_VDR_L treatment as compared to control treated or in EGFR-negative cells, implying that EGFR was endocytosed into the cell cytoplasm (fig. S6D). Because this phenomenon did not take place when cells were treated with control media or when the treated cells had very low levels of EGFR, these findings imply that EGFR-bound E_VDR_L was endocytosed into the cell.

Next, we used cetuximab and erlotinib to block EGFR. Cetuximab, an anti-EGFR monoclonal antibody, blocks E_VDR_L-EGFR binding, extracellularly; and erlotinib, a receptor tyrosine kinase

inhibitor, inhibits phosphorylation of EGFR, intracellularly (Fig. 4E). Consistent with previous reports (44), cetuximab and erlotinib alone had no effects on BLBC-BM cell proliferation (fig. S6E). As TNBC cell resistance to EGFR inhibitors could simply arise from relatively lower EGFR expression in TNBC than the other EGFR inhibitor-sensitive cancers, we compared EGFR expression of patients' TNBC-BM samples with BM of non-small cell lung carcinoma (NSCLC), which is an EGFR inhibitor-sensitive cancer type. Similar levels of EGFR expression were seen in both TNBC-BM and NSCLC-BM (fig. S6F). In addition, a cell line cohort from TGCA database (34) showed that TNBC cell lines have the same or even higher *EGFR* mRNA level compared to NSCLC, pancreatic cancer, and colorectal cancer, which are all considered EGFR inhibitor-sensitive cancers (fig. S6G). Cetuximab significantly blocked E_VDR_L-mediated reduction of cell viability and induction of apoptosis in BLBC-BM lines but did not affect the effect of DR_L (Fig. 4, F and G, and fig. S6H). Coimmunoprecipitation (Co-IP) studies showed that cetuximab interfered with formation of an EGFR-E_VDR_L-DR4/5 complex (Fig. 4H). After confirming that erlotinib sufficiently inhibits EGFR phosphorylation (fig. S6I), we treated erlotinib-pretreated BLBC-BM cells with E_VDR_L. Erlotinib pretreatment did not affect the efficacy of E_VDR_L (Fig. 4I). These results suggest that EGFR binding is critical for E_VDR_L therapeutic effects, but its efficacy is not mainly via blocking EGFR signaling.

To validate DR4/5 as the targets of E_VDR_L treatment, we generated DR4, DR5, and DR4/5 knockout (KO) BMET02 lines using CRISPR-Cas9 gene editing and treated them with DR_L or E_VDR_L (Fig. 4J). We confirmed that the growth rate of BMET02-DR4/5^{KO} is similar to its parental BMET02 cell line (fig. S2B). Both BMET02-DR4^{KO} and BMET02-DR5^{KO} lines were significantly less sensitive to DR_L and E_VDR_L than the control line. KO of both DR4 and DR5 induced almost complete resistance to DR_L and E_VDR_L (Fig. 4K and fig. S6J). These results suggested that DR4/5 expression is essential for E_VDR_L activity, which is enhanced by the interaction between E_V domain and EGFR.

We tested whether DR_L binding to DR4/5 differed when cells were exposed to E_VDR_L and E_V + DR_L. Co-IP assays showed 1.3- to 1.5- and 1.7- to 2.0-fold higher binding of E_VDR_L to DR4 and DR5, respectively, compared to DR_L (Fig. 4L). These results indicated that the E_V domain of E_VDR_L enhances the apoptosis-inducing function by increasing its binding to DR4/5 (Fig. 4M). E_SDR_L has lesser binding to DR5 compared to E_VDR_L resulting in its reduced apoptosis-inducing effect (fig. S6K).

E_VDR_L-secreting stem cells kill BLBC cells in vitro and in vivo

We generated human MSC (hMSC) expressing E_VDR_L and confirmed continuous secretion of E_VDR_L for 120 hours (Fig. 5A). hMSC expressed substantial levels of DR4/5 and high levels of DcR2 expression (fig. S5C). Coculture with BMET02-FmC showed that hMSC-E_VDR_L had more potent tumor-killing ability than hMSC-DR_L (Fig. 5B and movie S2). In parallel, we also generated DR_L- and E_VDR_L-secreting mouse NSC (mNSC), mouse MSC (mMSC), and induced pluripotent stem cell (iPSC)-derived NSCs (iPSC-NSC). All the E_VDR_L-secreting stem cells tested had significantly higher tumor cell killing activity compared with those secreting only DR_L (Fig. 5C and fig. S7). Next, we encapsulated E_VDR_L-secreting stem cells in synthetic extracellular matrix (sECM) that enabled us to prevent “wash out” of therapeutic stem cells and retain them within the tumor resection cavity based on our previous work with



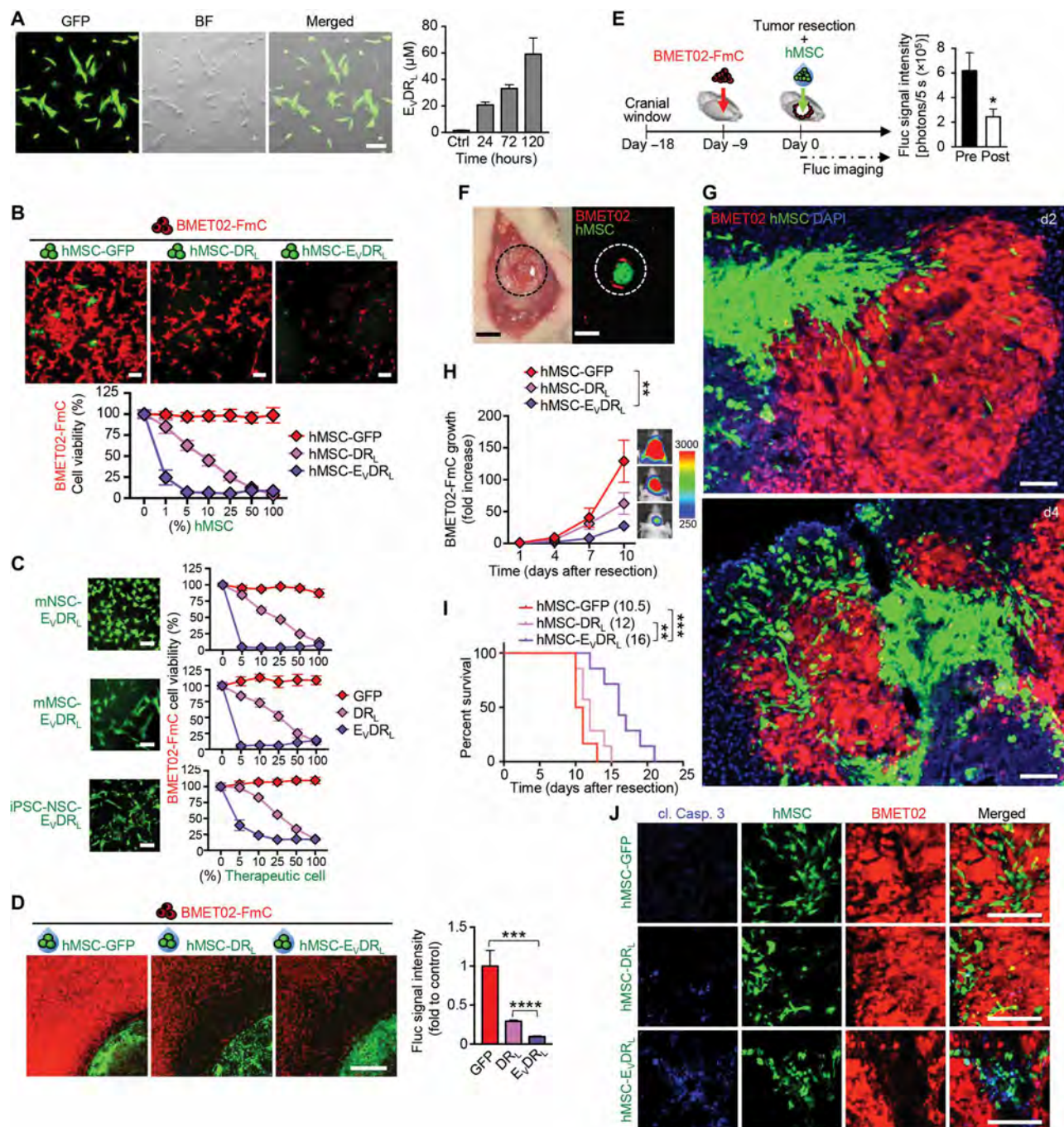


Fig. 5. E_vDR_L -secreting stem cells have antitumor effects against BLBC in vitro and in vivo. (A) Left: Photomicrograph of E_vDR_L -secreting hMSC. Scale bar, 100 μ m. Right: Concentration of E_vDR_L in culture media of hMSC- E_vDR_L quantified by enzyme-linked immunosorbent assay (ELISA) ($n=2$, technical replicates). (B) Top: Photomicrographs of BMET02-FmC cocultured with hMSC-GFP/ DR_L / E_vDR_L for 72 hours. Scale bars, 100 μ m. Bottom: Cell viability of BMET02-FmC after 72-hour coculture with increasing percentages of hMSC-GFP, hMSC- DR_L , or hMSC- E_vDR_L ($n=3$, technical replicates). (C) Photomicrographs of different engineered stem cells (left) (scale bars, 100 μ m) and cell viability of BMET02-FmC cocultured with increasing percentages (0 to 100) of the stem cells (right) ($n=3$, technical replicates). (D) Left: Photomicrograph of BMET02-FmC cocultured with sECM-encapsulated hMSC-GFP, hMSC- DR_L , or hMSC- E_vDR_L . Scale bar, 1 mm. Right: Relative number of BMET02-FmC cells 72-hour following coculture with sECM-encapsulated hMSC-GFP, hMSC- DR_L , or hMSC- E_vDR_L ($n=3$, technical replicates). (E) Left: Experimental outline for testing efficacy of sECM-encapsulated hMSC- E_vDR_L in BMET02-FmC-bearing mice. Right: BLI signals before and after resection ($n=20$). (F) Intraoperative photographs of light and fluorescence of mice implanted sECM-hMSC into the resection cavity of BMET02-FmC tumor. Scale bars, 1 mm. (G) Representative photomicrographs of brain section from mice 2 and 4 days after resection of BMET02-FmC tumor and implantation of sECM-hMSC. Scale bars, 100 μ m. (H) Estimate of relative tumor volume after resection in treatment groups based on Fluc signal intensity of BMET02-FmC (hMSC-GFP, $n=6$; hMSC- DR_L , $n=7$; hMSC- E_vDR_L , $n=7$). (I) Kaplan-Meier survival curves of the mice with median survival (days) indicated in the legend. (J) Immunohistochemistry of cleaved caspase-3 of brain sections from treated and control mice. Scale bars, 100 μ m. Photo credit: Yohei Kitamura, Brigham and Women's Hospital

glioblastoma resection model (40) and showed that they have efficient tumor killing abilities of BMET02-FmC cells in vitro (Fig. 5D and fig. S8A). We also confirmed that hMSC are viable in the mouse brain for at least 2 weeks after implantation (fig. S8B). In addition, we confirmed that hMSC-E_VDR_L treatment did not induce any significant toxicity, as indicated by stable body weight of treated mice as well as unremarkable histology of CNS and major organs (fig. S8, C to E).

To explore the effect of E_VDR_L-secreting hMSC in vivo, we first used a macrometastasis mouse model. Nine days after stereotactic implantation of BMET02-FmC in the brain, the tumor was resected partially, and hMSCs expressing green fluorescent protein (GFP), DR_L, or E_VDR_L were encapsulated in sECM and implanted in the resection cavity (Fig. 5, E and F). Brain histology showed that implanted sECM-encapsulated hMSCs migrated toward the tumor (Fig. 5G). BLI and histology of tissues revealed a significant decrease in tumor volumes in hMSC-E_VDR_L-treated mice as compared to the control group (Fig. 5H and fig. S8F). Also, the hMSC-E_VDR_L-treated group showed significantly longer survival time than hMSC-GFP- and hMSC-DR_L-treated groups (Fig. 5I). Immunofluorescence showed that hMSC-E_VDR_L induced increased caspase-3 cleavage in tumor cells compared to hMSC-GFP or hMSC-DR_L (Fig. 5J). These data reveal that stem cells engineered to simultaneously target EGFR and DR4/5 have potent efficacy in mouse tumor models of BM resection.

ICA-injected E_VDR_L-secreting stem cells effectively control PVN micrometastasis

To explore the therapeutic potential of stem cell delivery of E_VDR_L across BBB to micrometastatic cells within PVN, we used ICA injection of E_VDR_L-secreting stem cells. Since there is limited literature on the fate of ICA-injected stem cells, we first investigated the fate of five different types of stem cells (hMSC, hNSC, mNSC, mMSC, and iPSC-NSC) after ICA injection either in the presence or absence of tumors. Among the different stem cells tested, mNSCs survived in the brain for the longest period (fig. S9A). Since an extended survival of stem cells may increase their potential of tumor formation, we engineered mNSCs to coexpress E_VDR_L and a kill switch, prodrug-converting enzyme HSV-TK (TK) and confirmed that mNSC-TK and mNSC-E_VDR_L-TK cells were eradicated by ganciclovir (GCV) treatment (fig. S9B). Coculture with BMET02-FmC did not show additional therapeutic benefit of TK in mNSC-E_VDR_L-TK presumably because the highly efficient tumor killing by E_VDR_L masked other effects, but when cocultured with BT549-FmC, a relatively less sensitive BLBC line, an enhanced tumor killing effect was observed with mNSC-E_VDR_L-TK (fig. S9C). Next, we used BMET02-FmC micrometastatic models and ICA-injected mNSC-GFP or mNSC-E_VDR_L-TK and the BLI signal and monitored animal survival (Fig. 6A). Histology showed the presence of widely distributed stem cells along the brain vasculature that colocalized with BMET-02-FmC tumor cells (Fig. 6B). mNSC-E_VDR_L-TK treatment significantly extended both macrometastasis-free survival (Fig. 6, C and D) and overall survival (Fig. 6E) compared to mNSC-GFP. Further, GCV administration in mNSC-E_VDR_L-TK-treated mice resulted in similar therapeutic benefit compared with mNSC-E_VDR_L-TK alone, indicating enough safety of such therapeutic stem cells in a preclinical BM model, as well as little benefit of TK to mNSC-E_VDR_L in this cohort. Together, these studies reveal that ICA injection of E_VDR_L-secreting stem cells has therapeutic efficacy in mouse models of BC brain micrometastases.

IT-injected E_VDR_L-secreting stem cells effectively treat leptomeningeal metastasis

To explore the efficacy of stem cell delivery of E_VDR_L for leptomeningeal metastasis, we tested IT injection of hMSC secreting E_VDR_L. First, we confirmed that hMSCs survived in the CSF space for at least 2 weeks after IT injection in naïve mice and were gradually cleared out (Fig. 7A). In addition, we confirmed that IT injection of stem cells did not affect mice body weight (fig. S10A). Next, we showed that IT-injected hMSC homed to the tumors of IT-injected BMET02-FmC that were growing in the CSF space of different CNS regions (Fig. 7B). hMSC-E_VDR_L secreted a substantial amount of E_VDR_L into the CSF (Fig. 7C). hMSCs engineered to express GFP and renilla luciferase (Rluc) (hMSC-GRI) or hMSC-E_VDR_L were IT-injected twice to BMET02-FmC-bearing mice and evaluated by BLI and survival monitoring (Fig. 7D). Rluc imaging detected accumulation of hMSC-GRI after two implantations, and Fluc imaging of tumor cells and harvested brain and spinal cord samples showed that hMSC-E_VDR_L significantly suppressed tumor growth (Fig. 7, E to G, and fig. S10B). Compared to the control, treatment with hMSC-E_VDR_L resulted in significant improvement in survival of tumor-bearing mice (Fig. 7H). We also tested the efficacy of hMSC-E_VDR_L on the tumors generated from another BLBC cell line, MDA231-BrM2-FmC (fig. S10C) (21). Mice bearing MDA231-BrM2-FmC tumors treated with IT-injected hMSC-E_VDR_L demonstrated a significant survival benefit (Fig. 7, I and J). Immunofluorescence of cleaved caspase-3 showed apoptosis induction in hMSC-E_VDR_L-treated tumors (Fig. 7K). These results clearly demonstrated the therapeutic efficacy of IT-injected hMSC-E_VDR_L in mouse models of BC leptomeningeal metastases.

DISCUSSION

In this study, we developed different imageable mouse models of BLBC-BM and explored the versatility of stem cell-mediated bi-functional EGFR and DR4/5 therapeutics in these models. Our results show that the E_V domain of E_VDR_L enhances DR_L-induced apoptosis in a broad spectrum BLBC lines, and the stem cell-secreted E_VDR_L targets PVN micrometastasis and leptomeningeal metastasis, thus offering a promising therapeutic strategy for BLBC-BM.

Clinically relevant metastatic BM models are usually developed by intracardiac administration of cancer cells; however, this leads to widespread extracranial tumor formation (36, 45, 46). Standard ICA injections also cause extracranial tumors in the face (37–39). When using sagittal imaging as we show in fig. S3 (A to C), we observed extracranial tumor formation in about 80% of cases following standard ICA injections. Nevertheless, a number of previous studies have disregarded these findings. In these models, signal emitted from extracranial (often facial) tumors substantially confounds BLI evaluation of brain tumor progression and treatment benefits. In this study, we refined the technique of ICA injection, and our modified approach reduced the rate of extracranial tumor formation to less than 20% of cases, enabling accurate BLI monitoring of intracranial tumors and evaluation of treatment efficacy. Although creating our model requires advanced technical skills and is time consuming, we believe that creating and optimizing mouse tumor models of metastasis confer multiple advantages in advancing BM preclinical research.

Although promising, there may be some potential limitations in the models we developed and used in this study. Since BM is

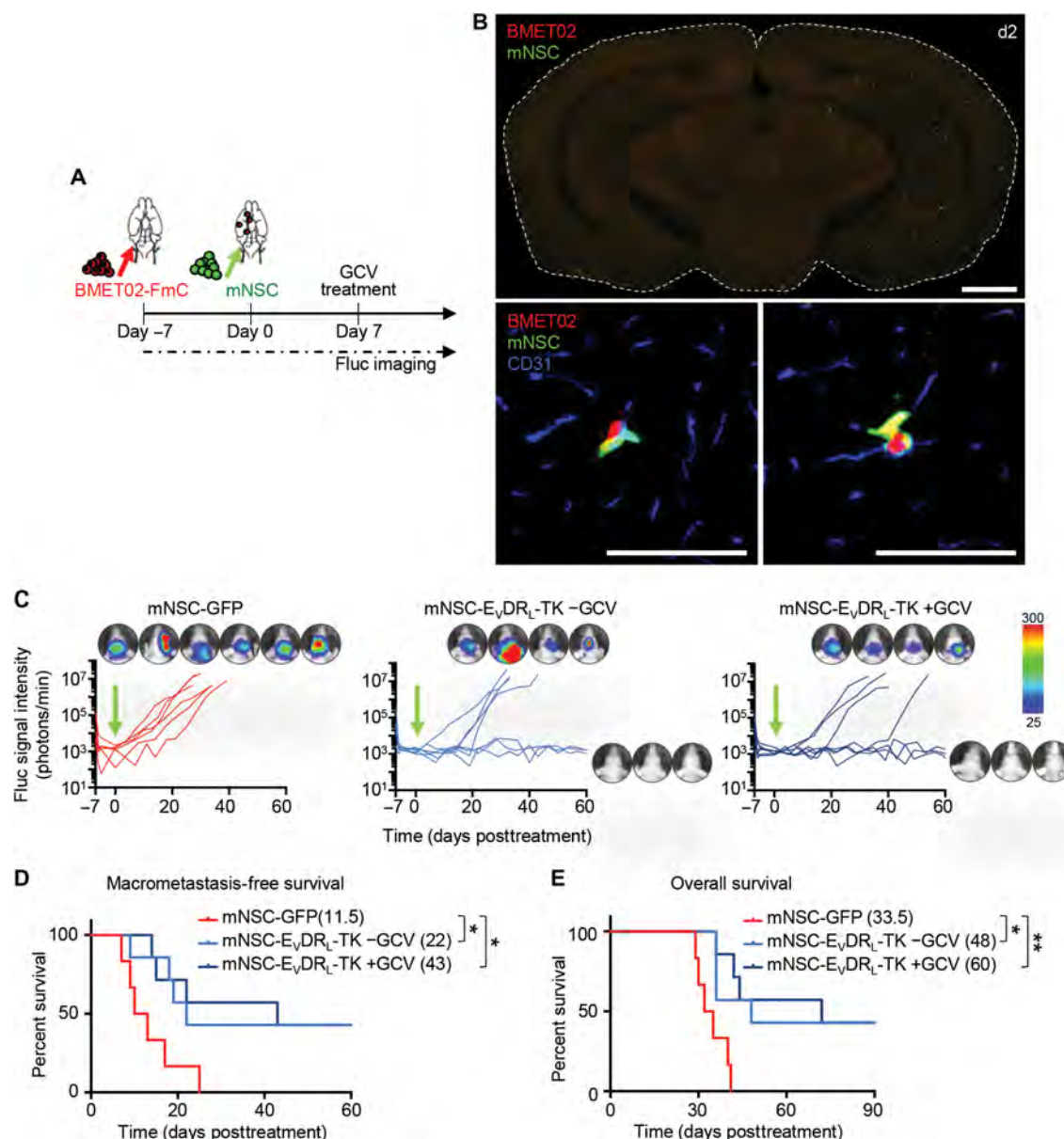


Fig. 6. E_vDR_L-secreting stem cells show antitumor efficacy for micrometastasis of BLBC. (A) Experimental outline for testing efficacy of ICA injection of mNSC-E_vDR_L-TK in mice that had ICA injection of BMET02-FmC 7 days before. (B) Top: Representative photomicrograph of whole brain section from ICA-injected BMET02-FmC-bearing mice 2 days after ICA injection of mNSC. Scale bar, 1 mm. Bottom: Representative photomicrograph of immunohistochemistry of CD31 of the brain section. Scale bars, 100 μ m. (C) BLI signal curves and photographs of mice treated with mNSC-GFP/E_vDR_L-TK +/- GCV (mNSC-GFP, $n = 6$; mNSC-E_vDR_L-TK -GCV, $n = 7$; mNSC-E_vDR_L-TK +GCV, $n = 7$). (D) Kaplan-Meier curves of macrometastasis-free survival. The presence of macrometastasis was judged from the substantial BLI signal around 1×10^4 photons/min. Median macrometastasis-free survivals (days) are indicated in the legend. (E) Kaplan-Meier curves of overall survival of mice with median overall survival (days) indicated in the legend.

clinically developed by the cells via arterial extravasation, arterial injection is regarded as an optimal way to develop BM models (47). As ICA injection does not reproducibly induce resectable intracranial tumor formation (single large tumor at surgically accessible location), we had to use orthotopic BM cell line injections for developing BM resection model. Although not optimal, this model has some advantages, e.g., easier to develop models uniform in size and timing, especially for testing some therapeutic efficacy in limited living materials. Although we observed similar microenvironmental findings to arterially injected models, there is no doubt that

arterial injection is the preferred way to create BM models for study when tumor resection is not considered.

DR_L is a well-investigated molecule, which induces cancer-specific apoptosis. It is also known that BLBC is sensitive to DR_L (23); however, the underlying mechanism has been unclear. We showed that the increased DR_L sensitivity of BLBC is closely associated with up-regulated DR4/5 expression. However, DR-targeted therapies have shown poor efficacy (24, 25). Enhancement of the efficacy of DR_L has been attempted by many researchers via approaches such as molecular modifications (to stabilize it or prolong its longevity), combination with other molecules, and

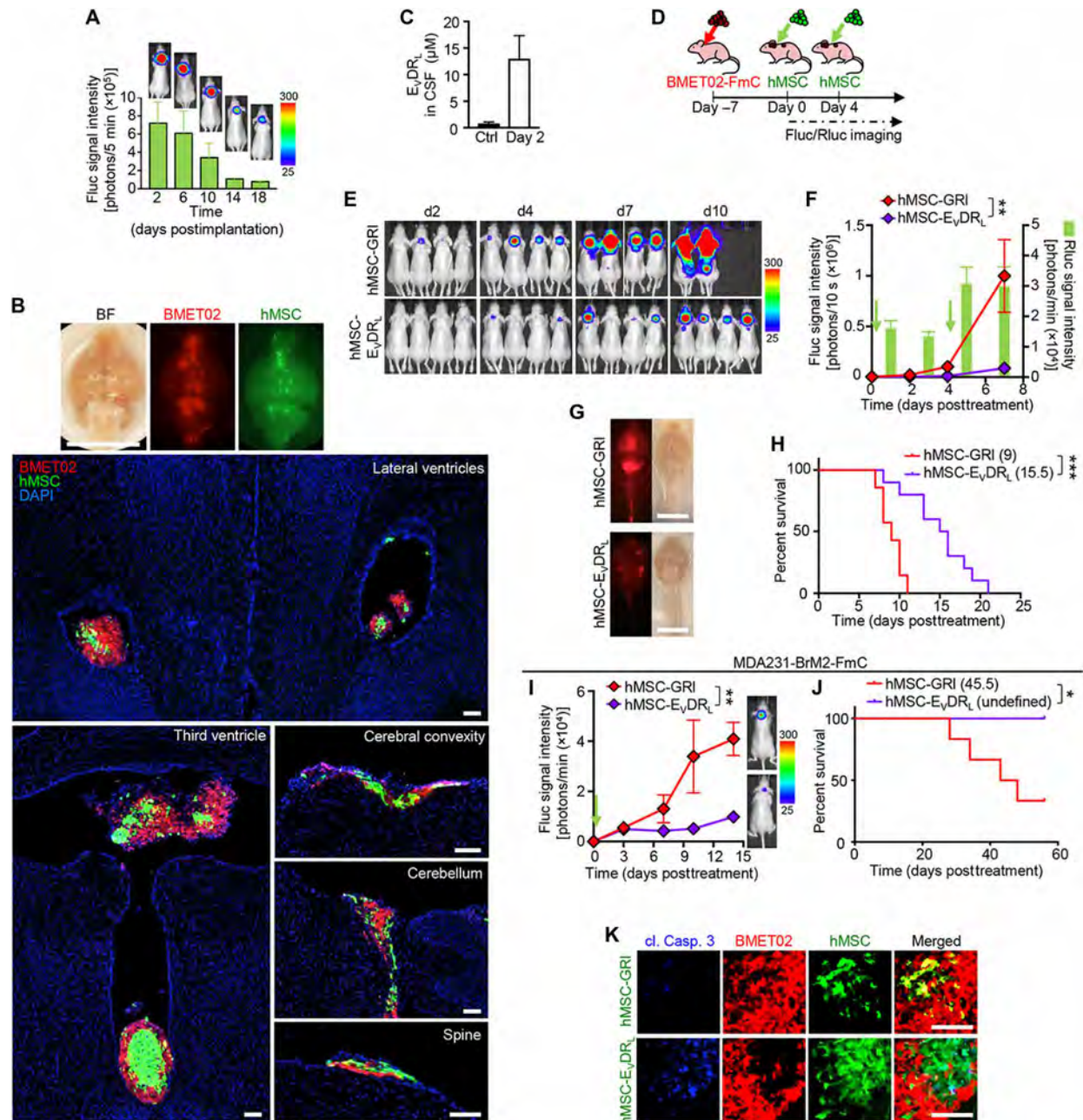


Fig. 7. E_VDR_L -secreting stem cells have antitumor effects in leptomeningeal metastasis of BLBC. (A) BLI signal and photographs of IT-injected hMSC-GFP-Fluc (GFI)-bearing mice ($n=3$). (B) Top: Representative photograph of whole-brain sample of an IT-established, BMET02-FmC leptomeningeal metastasis-bearing mice, 2 days after IT injection of hMSC. Scale bar, 10 mm. Bottom: Representative photomicrographs of brain and spine sections from the mice. Scale bars, 100 μ m. (C) Concentration of E_VDR_L in CSF from mice before and 2 days after IT injection of hMSC- E_VDR_L quantified by ELISA ($n=3$). (D) Experimental outline for testing efficacy of IT injections of hMSC- E_VDR_L in mice that had IT injection of BMET02-FmC 7 days before. (E) Representative BLI pictures of IT-injected BMET02-FmC-bearing mice treated with hMSC-GRI or hMSC- E_VDR_L . (F) Fluc signal curves of BMET02-FmC treated with hMSC-GRI or hMSC- E_VDR_L and Rluc signals of injected hMSC-GRI (hMSC-GRI, $n=7$; hMSC- E_VDR_L , $n=10$). (G) Representative photographs of whole-brain sample of IT-injected BMET02-FmC-bearing mice treated with hMSC-GRI or hMSC- E_VDR_L for 7 days. Scale bars, 10 mm. (H) Kaplan-Meier curves of overall survival of mice. Median survivals (days) are indicated in the legend. (I) Fluc signal curves and representative BLI images of mice bearing MDA231-BrM2-FmC tumors treated with hMSC-GRI or hMSC- E_VDR_L (hMSC-GRI, $n=6$; hMSC- E_VDR_L , $n=5$). (J) Kaplan-Meier curves of overall survival of mice with median survival (days) indicated in the legend. (K) Immunohistochemistry of cleaved caspase-3 in brain sections from IT-injected BMET02-FmC-bearing mice treated with hMSC-GRI or hMSC- E_VDR_L . Scale bars, 100 μ m.

sensitizing the target site to DR_L (25, 48). Here, since our initial aim of fusing the E_V domain to DR_L was to block EGFR signaling, it was an unexpected discovery that the E_V domain augmented DR_L - $DR4/5$ binding and thereby enhanced the efficacy of E_VDR_L .

Because EGFR is up-regulated in BLBC (19) and is a marker of poor prognosis (49), it is a promising target for BLBC. However, EGFR-targeting therapy has failed to show any survival benefits (20). Consistently, BLBC cell lines were resistant to two EGFR inhibitors

herein. However, we showed that the E_V domain of E_VDR_L significantly improved the efficacy of DR_L, suggesting that EGFR can be used as an anchor to increase therapeutic targeting of other cell surface molecules in BLBC and other EGFR-expressing cancers. We also showed that EGFR and DR5 expression levels are significantly related to the efficacy of E_VDR_L. These results imply that analysis of resected tumor tissue from patients could allow a prediction of efficacy of this treatment and thus offer personalized treatments. We show that VHH of anti-EGFR was more effective than scFv when fused to the N terminus of DR_L. VHH composes of a single domain and is more stable and robust than scFv, which has a linker connecting two domains requiring a supramolecular assembly. Also, VHH has a unique epitope that is longer and contains a more flexible complementary determining region, which increases the affinity to corresponding receptors (50). These differences between VHH and scFv might contribute to the higher efficacy of E_VDR_L.

Our findings indicate that BLBC's sensitivity to E_VDR_L is mainly determined by the expressions of EGFR and DRs. Although there are likely other apoptosis-related factors involved in BLBC's sensitivity to E_VDR_L, e.g., Myc, our studies did not demonstrate how these factors could influence the sensitivity to E_VDR_L. Further investigations are needed to completely understand the influence of these factors on BLBC's sensitivity to E_VDR_L. In this study, we have not compared the difference in efficacy between stem cell-delivered E_VDR_L therapy and systemic venous injection of E_VDR_L; however, given the short half-life of DR_L and the inability of a majority of drugs to cross BBB, we anticipate that stem cell-delivered E_VDR_L will be a more effective treatment paradigm for BM.

BC often presents metastasis years or even decades after treatment and apparent good disease control (11, 12). This suggests that disseminated tumor cells of BC stays dormant within the PVN in the distant organs for a long period; these cancer cells outside the vasculature are difficult to target with systemically administered drugs (13, 14). We successfully developed cancer models with BC cells residing in the brain PVN and used stem cells to deliver therapeutic molecules. However, the treatment for dormant cancer cells in PVN is difficult due to the inability to detect such scattered cells in patients and drug delivery beyond BBB. Given the ability of engineered stem cells to migrate to cancer cells and penetrate BBB (30–32), our stem cell-based treatment has the potential to overcome that. Our results indicate that clinical trials of this therapy using selective arterial administration by neuroendovascular devices for patients with BC who had initial radical treatment are a feasible option to erase dormant cancer cells before growth, prevent future macrometastasis in the brain, and should be considered. Recent studies reported the existence of micrometastatic cancer cells in other organs, such as the lung, bone, and liver, and recurrence in these organs long periods after initial therapy is a considerable challenge (12). Our approach of stem cell delivery of potent therapeutics offers an immense potential of killing cancer cells dormant at the PVN in these organs.

We also showed that stem cell delivery is a promising approach for treating leptomeningeal metastasis, which is considered a terminal condition without any effective therapeutic strategies. We showed that IT-injected stem cells could stay alive in the niche for weeks and secrete therapeutic molecules into CSF without affecting the general health of the mouse. Clinical trials of this therapy should be considered for patients with BLBC leptomeningeal metastasis in the future. IT stem cell therapy has already been established as a safe

treatment and tested in patients of trauma (51), stroke (52), epilepsy (53), and neurodegenerative diseases (54, 55). Two animal studies have reported IT stem cell therapies for disseminated primary brain tumor—glioma and medulloblastoma (56, 57). However, to our knowledge, there is no previous report showing efficacy of stem cell therapy for leptomeningeal metastasis, a secondary CNS tumor. The same strategy might be effective for leptomeningeal disease originating from other types of primary and metastatic cancers.

In conclusion, we demonstrate the efficacy of a stem cell-delivered therapeutics against EGFR and DR4/5 in mouse models representing three clinically challenging BM conditions. Our findings provide a scientific rationale that supports clinical trials of this strategy in patients with BLBC-BM.

MATERIALS AND METHODS

Antibodies and reagents

The following antibodies and reagents were used in this study. Antibodies against β-actin (#4970), phospho-AKT (Ser⁴⁷³, #4060), AKT (#9272), caspase-7 (#9492), caspase-8 (#9746), caspase-9 (#9508), cleaved caspase-3 (#9661), EGFR (#2646 and #4267), phospho-EGFR (Tyr¹⁰⁶⁸, #3777), cleaved poly(ADP-ribose) polymerase (PARP; #9541), phospho-p44/42 mitogen-activated protein kinase (MAPK) (ERK1/2) (Thr²⁰²/Tyr²⁰⁴, #9101), p-44/42 MAPK (ERK1/2) (#9102), Fas-associated death domain protein (#2782), Bcl-2 (#2872), Bcl-xL (#2764), XIAP (#2042), cIAP2 (#3130), phospho-signal transducers and activators of transcription 3 (STAT3) (Tyr⁷⁰⁵, #9145), STAT3 (#4904), HER2 (#2242), horseradish peroxidase (HRP) anti-rabbit (#7074), Rab5 (#46449), Rab7 (#95746) (Cell Signaling Technology), anti-nuclear factor κB (#ab16502), anti-TRAIL (#ab9959), anti-CD31 (#ab28364), HRP anti-mouse (#ab205719) (Abcam), anti-α-tubulin (#T5168), anti-Vinculin (#V4505), NeuN (#MAB377), glial fibrillary acidic protein (GFAP) (#MAB3402) (Sigma-Aldrich), Alexa Fluor 488 anti-EGFR antibody (#352908), anti-DR4 (#1139), anti-DR5 (#2019) (ProSci), anti-DR4 (#sc-32255), anti-DR5 (#sc-166624), anti-cIAP1 (#sc-271419), normal mouse IgG (#sc-2025) (Santa Cruz), anti-Ki-67 (#180191Z), anti-GFAP (#180063), Alexa Fluor anti-rabbit 405 (#A-31556), Alexa Fluor anti-rabbit 488 (#A-11008), Alexa Fluor anti-mouse 555 (#A-21422), Alexa Fluor anti-rabbit 647 (#A-21244), Phycoerythrin (PE) anti-DR4 (#12-6644-42), PE anti-DR5 (#12-9908-42), PE mouse IgG isotype (#12-4714-42) (Invitrogen), Cetuximab (ImClone Systems), Erlotinib (#SYN-1039, Selleck Chemicals), human recombinant EGF (R&D Systems), PE anti-EGFR (#352903, BioLegend), and IBA1 (#019-19741, FUJIFILM).

Immunohistochemistry for patient tissue samples

TNBC patient tissue samples were obtained from Massachusetts General Hospital as approved by institutional review board (IRB) at Harvard Medical School, Keio University Hospital as approved by IRB of Keio University School of Medicine, and US Biomax Inc. (#BR1901). Immunohistochemical analyses of patient samples were performed by Servicebio Inc. and iHisto Inc., using antibodies for EGFR (#GB13804), estrogen receptor (#GB11205), progesterone receptor (#GB11262) from Servicebio Inc. and the antibodies listed above.

Cell lines

Patient-derived BLBC-BM cell lines (BMET02, BMET05, BMET15) were obtained by dissociation of brain tumor samples from patients

with metastatic breast carcinoma diagnosis and cultivated as described below. Brain tumor samples were obtained as approved by IRB at Harvard Medical School. Established BC cell lines MDA-MB-453, MCF7, HCC1500, HCC1428, ZR75-1, BT474, T47D, MDA-MB-175VII, and SUM159 were provided by A. Tilston-Lunel, Bob Varelas laboratory, Boston University. SKBR3 was provided by N. Wang, Massachusetts General Hospital. MDA-MB-231, MDA-MB-231-BrM2, MDA-MB-436, and MDA-MB-468 were provided by J. Massagué, Memorial Sloan Kettering Cancer Center. BT549, Hs578T, and NIH-3T3 were purchased from American Type Culture Collection. The immortalized hMSC line, hASC-TS, was a gift from L. Balducci (58). Immortalized mNSC line, C17.2, was provided by E. Y. Snyder (59). Bone marrow-derived mMSC line was obtained from D. Prockop, University of Texas. Immortalized human fetal NSC (hNSC) line, hNSC100, was provided by A. Martínez-Serrano, Autonomous University of Madrid. Mouse iPSC-NSC was generated from mouse embryonic fibroblasts as previously described (60).

BMET02, BMET05, BMET15, MDA-MB-453, MCF7, MDA-MB-175VII, MDA-MB-231, MDA-MB-231-BrM2, MDA-MB-436, and MDA-MB-468 were grown in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% (vol/vol) fetal bovine serum (FBS) and 1% (vol/vol) penicillin/streptomycin. Hs578T was grown in DMEM supplemented with 10% (vol/vol) FBS, insulin (0.01 mg/ml), and 1% (vol/vol) penicillin/streptomycin. SKBR3 was grown in McCoy's 5a medium with 10% (vol/vol) FBS and 1% (vol/vol) penicillin/streptomycin. HCC1500, HCC1428, ZR75-1, BT474, and BT549 were grown in RPMI 1640 with 10% (vol/vol) FBS and 1% (vol/vol) penicillin/streptomycin. T47D was grown in RPMI 1640 with 10% (vol/vol) FBS, 1% (vol/vol) penicillin/streptomycin, and insulin (0.2 U/ml). SUM159 was grown in Ham's F-12 with 5% (vol/vol) FBS, insulin (0.005 mg/ml), hydrocortisone (1 µg/ml), and 1% (vol/vol) penicillin/streptomycin. NIH-3T3 were cultured in DMEM supplemented with 10% NCS, penicillin (100 U/ml), and streptomycin (100 µg/ml). hMSC was grown in DMEM/F-12 supplemented with 10% (vol/vol) FBS, 1% (vol/vol) L-glutamine, 1% (vol/vol) penicillin/streptomycin, and recombinant human fibroblast growth factor (FGF) (40 ng/ml; R&D Systems, Minneapolis, MN). mNSC was grown in DMEM supplemented with 10% (vol/vol) FBS, 1% (vol/vol) L-glutamine, and 1% (vol/vol) penicillin/streptomycin. mMSC was grown in low-glucose DMEM supplemented with 15% (vol/vol) FBS, 1% (vol/vol) L-glutamine, 1% (vol/vol) nonessential amino acid solution, and 1% (vol/vol) penicillin/streptomycin. hNSC was cultured in 4:1 culturing medium [DMEM/F-12 (Invitrogen), 0.6% D-glucose (Sigma-Aldrich), 0.5% albumax (Invitrogen), 0.5% glutamine (Invitrogen), recombinant human FGF (40 ng/ml; R&D Systems), recombinant human EGF (40 ng/ml; R&D Systems), N2 supplements (Invitrogen), and 1% nonessential amino acid solution (Cellgro; Mediatech)] and growth medium [DMEM with 5% FBS, 1 mM sodium pyruvate (Cellgro; Mediatech), and 26 mM sodium bicarbonate]. iPSC-NSC was grown in NeuroCult basal medium (Stem Cell Technologies) supplemented with EGF (20 ng/ml; R&D Systems), FGF2 (20 ng/ml; R&D Systems), N2 supplement, Heparin, and 1% (vol/vol) penicillin/streptomycin on Geltrex (Fisher Scientific)-precoated flask.

Cell viability and caspase assays

Tumor cells were plated in 96-well plates and treated with different doses of anti-EGFR VHH (E_V), DR ligand (DR_L), E_VDR_L , or anti-EGFR

scFv-TRAIL (E_SDR_L) and control media for 24, 48, and 72 hours. To obtain conditioned media containing these proteins, lentiviral plasmid vectors coding for E_V , DR_L , E_VDR_L , and E_SDR_L were transfected into 293T cells. Medium was changed the next day, collected 40 hours after transfection, concentrated using centrifugal filter (#UFC901024, MilliporeSigma), and stored at -80°C until future use. Their concentrations were quantified by enzyme-linked immunosorbent assay (ELISA). Control media were made from GFP control-transduced cells transduced in parallel with E_V , DR_L , E_VDR_L , and E_SDR_L . Cell viability was measured using an adenosine triphosphate-dependent luminescent reagent (CellTiter-Glo, #G755A, Promega; Glomax, Promega) according to the manufacturer's instructions for non-Fluc-expressing cells or with D-luciferin (#122799, PerkinElmer) and coelenterazine h (#760506, PerkinElmer) for Fluc- and Rluc-expressing cells, respectively. Caspase-3/7 activity was determined using a DEVD-aminoluciferin assay (Caspase-Glo 3/7, #G8091, Promega) according to the manufacturer's instructions. All experiments were performed in triplicates.

Western blot analysis

After treatment, cells were washed with cold phosphate-buffered saline (PBS) and then lysed with cold NP-40 lysis buffer (#BP-119, Boston BioProducts) supplemented with protease inhibitor (#A32965, Thermo Fisher Scientific) and phosphatase inhibitors (#P5726 and #P0044, Phosphatase Inhibitor Cocktail 2,3 from Sigma-Aldrich). Cells were scraped into tubes and centrifuged at 4°C at 13,000g for 10 min. Supernatant protein concentrations were determined using the Bio-Rad DC Protein Assay Kit (#500-0113, #500-0114, and #500-0115). The 6× SDS sample buffer (#BP-111R, Boston BioProducts) was added to the samples, which were then boiled for 5 min. Ten to forty micrograms of protein was loaded on SDS-polyacrylamide gel electrophoresis gel (#456-1086 and #456-1093, Bio-Rad), transferred to polyvinylidene difluoride membrane (#IPVH00010, Merck Millipore), and probed with primary antibodies overnight. After wash, the membrane was probed with secondary antibodies and developed with enhanced chemiluminescence (#1863096, #1863097, and #34095, Thermo Fisher Scientific).

Flow cytometry analysis of cell surface receptors

Cells were trypsinized, washed, and resuspended in stain buffer (#554657, BD Biosciences). Cells were stained with PE-conjugated anti-human EGFR, DR4, or DR5 antibodies in solution at 4°C for 30 min. For the double staining, PE-conjugated anti-DR5 antibody and Alexa Fluor 488-conjugated anti-EGFR antibody were used. Rinses were performed with stain buffer at 4°C . PE-conjugated isotype-specific IgG was used as a control. Flow cytometry was performed using FACSARIA II (BD) cell sorter, and data were analyzed using FlowJo (BD).

Coimmunoprecipitation

After treatment, cells were washed with PBS twice and then lysed with radioimmunoprecipitation assay (RIPA) buffer (#BP-115, Boston BioProducts) supplemented with protease inhibitor (#A32965, Thermo Fisher Scientific). Cells were scraped into tubes and centrifuged at 4°C at 13,000g for 20 min. Supernatant protein concentrations were determined using the Bio-Rad DC Protein Assay Kit. A mixture of 1 mg of protein, 20 µl of anti-human DR4/5 antibodies, and 30 µl of Protein A/G agarose (#sc-2003, Santa Cruz) was incubated overnight at 4°C . After washing with RIPA buffer, samples were boiled

with 6× SDS sample buffer (#BP-111R, Boston BioProducts) for 8 min. Samples were then centrifuged at 4°C at 13,000g for 2 min, and the supernatant was used for Western blot.

Lentiviral transductions and engineering of stable cell lines

Lentiviral vector of E_VDR_L was constructed by inserting cDNA encoding extracellular domain of DR_L into LV-anti EGFR VHH (E_V). Lentiviral vector of E_SDR_L was constructed by replacing E_V domain of E_VDR_L with cDNA encoding anti EGFR scFv (E_S). We used previously described lentiviral vectors of GFP, E_V, and DR_L (27, 61). Lentiviral packaging was performed by transfection of 293T cells as previously described (62), and cells were transduced with lentiviral vectors at multiplicity of infection of 2 in medium containing protamine sulfate (2 µg/ml). For BLI, cells were transduced with LV-Pico2-Fluc-mCherry, LV-Pico2-Rluc-mCherry, LV-Pico2-Fluc-GFP, or LV-Pico2-Rluc-GFP. They are selected by fluorescence-activated cell sorting (FACS) using a BD FACSAria Fusion cell sorter or by puromycin selection (1 µg/ml) in culture. GFP or mCherry expression was visualized by fluorescence microscopy.

Coculture and GCV treatment experiments

BMET02-FmC cells (2 × 10³ cells per well) were cocultured with different numbers of therapeutic stem cells in 96-well plates. After 72 hours, the relative number of BMET02-FmC cells was determined by Fluc luminescence (Glomax, Promega). For coculture experiments with encapsulated stem cells, 5 × 10⁴ of hMSC-GFP/DR_L/E_VDR_L cells were encapsulated with 10-µl sECM (HyStem-C Hydrogels, #GS313, BioTime) and added at the center of the well of a 24-well plate. After 30 min, BMET02-FmC was seeded around the gel. After 72 hours, the relative number of BMET02-FmC cells was counted by Fluc luminescence (IVIS Lumina, PerkinElmer). For in vitro GCV treatment, cells were treated with GCV (5 µg/ml) for 96 hours, and the relative cell number of them was quantified by CellTiter-Glo (Promega).

CRISPR KO of DR4 and DR5

CRISPR KO of DR4 and DR5 was conducted as previously described (61, 63). To establish KO lines, cells were transduced with lentiviral Cas9 expression vectors coding for either tetracycline-inducible or constitutively expressed Cas9 protein as previously described (64, 65). Confirmed Cas9 lines were engineered with lentiviral single guide RNA (sgRNA) expression vector pLKO.DEST.hygro containing the sgRNA target sequences described above for DR4 or DR5, followed by selection with hygromycin (200 to 500 µg/ml). For generating the double KO lines, confirmed Cas9 lines were co-engineered with pLKO.DEST.hygro and pLKO.DEST.egfp lentiviral expression vectors to express both DR4 and DR5 targeting sgRNAs followed by selection with hygromycin and FACS sorting for GFP.

ELISA of DR_L

Concentrations of DR_L released from therapeutic stem cell lines in culture medium or in CSF of mice that had IT injection of stem cells were quantified using a human-specific TRAIL antigen capture ELISA kit (#ab46074, Abcam).

Subtyping of cell lines

Genomic RNA from cell lines was extracted using the RNeasy Mini Kit (#74104, Qiagen). mRNA array was performed using a PAM50 plate (NanoString), and the results were analyzed by the company

(35). Subtypes were decided on the basis of the algorithm from the company.

Real-time FRET imaging

Confocal images were acquired with a Leica TCS SP8 Falcon system equipped with a 440-nm pulsed SMD diode laser and a tunable white light laser (WLL). YFP constructs were imaged with 512-nm WLL excitation, and emissions were collected over the range of 533 to 565 nm. CFP constructs were imaged with a 440-nm excitation laser and an emission of 465 to 495. For sensitized emission FRET experiments, BMET02 or NIH-3T3 cells were plated on 22 × 50 mm glass coverslips and cotransfected with LV-EGFR-YFP and LV-DR5-CFP using Lipofectamine LTX transfection reagent. Coverslips were mounted onto a JG-23W HP flow-through perfusion chamber (Warner Instruments), and cells were maintained in Ringer's solution containing 125 mM NaCl, 25 mM Hepes, 10 mM glucose, 5 mM K₂HPO₄, 1 mM MgSO₄, and 1 mM CaCl₂ (pH 7.40). Real-time FRET imaging experiments were performed using fluorescence ratio imaging systems built around a Nikon Eclipse TE2000-U inverted epifluorescence microscope equipped with an Andor Ultra 888 EM-CCD camera and a 60× Plan Apo TIRF (total internal reflection fluorescence) (numerical aperture, 1.45) oil immersion objective. Filter wheels (Sutter Instruments) were placed in the excitation and emission path, and image acquisition parameters were controlled by Metafluor software (Molecular Devices). FRET emission ratios (480 nm/535 nm; 440-nm excitation) were acquired every 10 s. After establishing baseline FRET levels for 3 to 10 min, E_VDR_L (2 µM) in Hepes-buffered Ringer's solution was added manually to the chamber with a pipette, and FRET changes were followed for 30 to 40 min.

Immunofluorescence analyses of endocytosis proteins

BMET02 cells were treated with control media or E_VDR_L (2 µM) after 4-hour starvation and fixed on the plates 5 and 15 min after treatment. Cells were stained with primary antibodies (EGFR and Rab5/Rab7) and counter-stained with secondary antibodies (Alexa Fluor anti-rabbit 488 and Alexa Fluor anti-mouse 555, respectively).

Animal models

All in vivo procedures were approved by the Subcommittee on Research Animal Care at Brigham and Women's Hospital. Mice that died or were euthanized for ethical reasons before defined experimental end points were excluded. Animals were randomly allocated to cages and experimental groups.

Macrometastasis resection model

Female nude mice (6 to 8 weeks of age, 20 to 25 g, Envigo) were immobilized on a stereotactic frame 9 days before tumor implantation. Using a stereomicroscope (SZX10, Olympus), a small circular portion of the skull covering the right cerebral hemisphere (3 mm by 5 mm) was removed to create a cranial window for subsequent tumor cell implantation and tumor debulking. Nine days later, the mice were again immobilized on a stereotactic frame, the previously established cranial window was exposed, and BMET02-FmC (5 × 10⁴ cells per mouse) in 4 µl of PBS was superficially implanted into the right frontal cerebral cortex (2-mm lateral from bregma, 0.5-mm deep) using a microsyringe (Hamilton). Nine days after the implantation of tumor cells, the mice underwent fluorescence-guided tumor resection followed by implantation of hMSC into the resection cavity as previously described. hMSC (5 × 10⁵ cells per

mouse) was encapsulated in 10 μ l of HyStem-C Hydrogels (#GS313, BioTime) 20 min before implantation to allow gel formation. Mice were then followed up for survival and sacrificed when neurological symptoms became apparent. Mice whose BLI signal disappeared completely after resection were excluded from this study.

Micrometastasis model

The detailed technique of ICA injection of tumor/stem cells is demonstrated in movie S1. Female nude mice (6 to 8 weeks of age) were anesthetized with ketamine-xylazine and fixed on the stage of a stereomicroscope (SZX10, Olympus). Midline skin incision was made to expose the right carotid arteries. Using 8-0 sutures, right OA, PPA, STA, and external carotid artery were ligated to prevent cells from going to extracranial parts. Internal and common carotid arteries were then ligated, and a catheter (#18000-10, Fine Science Tools) connected to a 1-ml syringe (Henke-Sass Wolf) was inserted into the external carotid artery. After releasing blood flow of common and internal carotid arteries, BMET02-FmC (5×10^4 cells per mouse) suspended in 100 μ l of PBS was slowly injected through the catheter. After injection, external carotid artery was permanently ligated. Mice with apparent extracranial tumors that could disturb the evaluation were excluded from the study. Seven days after tumor injection, stem cells were injected into the same artery using the same technique as above without ligations of OA, PPA, and STA. mNSC (4×10^5 cells per mouse) in 100 μ l of PBS was slowly injected.

Leptomeningeal metastasis model

IT injection of tumor was performed on the basis of previous literature (66) with slight modifications. Female nude mice (6 to 8 weeks of age) were immobilized on a surgical platform after anesthesia with ketamine-xylazine. Midline skin incision was made behind the neck, and occipital muscles were dissected. The dura mater between skull and atlas vertebra was exposed. Under observation of cerebellum and brainstem through the dura mater, a catheter connected to microsyringe (Hamilton) was inserted into cisterna magna. BMET02-FmC (2.5×10^4 cells per mouse) or MDA231-BrM2-FmC (5×10^4 cells per mouse) in 4 μ l was injected slowly through the catheter. The hole in the dura mater was closed with a small muscle piece immediately after removing catheter. hMSC (5×10^5 cells per mouse) was injected in a similar manner via the same hole from the previous injection.

Orthotopic injection

For testing the fate of stem cells, and for creating tumor-bearing mice for screening of ICA-injected stem cells, orthotopic injection into brain was performed. Mice were immobilized on a stereotactic frame, BMET02-RmC cells (1×10^5 cells per mouse) or hMSC (5×10^5 cells per mouse) in 4 μ l of PBS were implanted into the right frontal cerebrum (2-mm lateral and 1-mm anterior from bregma, 2.5-mm deep) using stereotactic frame.

In vivo imaging

BLI was used to follow in vivo growth of Fluc- or Rluc-engineered implanted tumor cells over time using a PerkinElmer IVIS Lumina system. For Fluc imaging, mice were imaged 7 min after intraperitoneal injection of D-luciferin (#122799, PerkinElmer). For Rluc imaging, mice were imaged 1 min after intravenous injection of coelenterazine h (#760506, PerkinElmer).

In vivo GCV treatment

For in vivo experiments involving mNSC elimination via the inducible suicide system HSV-TK, mice were treated daily with intraperitoneal injection of GCV (10 mg/kg) for 2 weeks starting 7 days after mNSC injection.

Tissue processing and immunohistochemistry of mouse samples

Tumor-bearing mice were perfused with PBS and subsequently with 4% paraformaldehyde. Brains and spines were harvested, followed by sectioning for histological analyses. Brain and spine sections on slides were washed in PBS and mounted with aqueous mounting medium (#H1000 and #H1200, Vector Laboratories) to be visualized with confocal microscopy (Axio Observer.Z1, Zeiss). For fluorescence immunohistochemistry, sections were incubated with primary antibodies overnight at 4°C. After wash, secondary antibodies were probed and detected by confocal microscope. For hematoxylin and eosin (H&E) staining, sections were incubated with H&E Y dye (1% alcohol), dehydrated with 70, 95, and 100% ethanol, and mounted in xylene-based mounting medium.

Statistical analyses

Data were analyzed by Student's *t* test for comparison between two groups and by Pearson's test for correlation. Data were plotted as means with SD for all in vitro data except Fig. 3E and fig. S4D, and with SEM for all in vivo data, Fig. 3E and fig. S4D. Survival curves were compared using the log-rank test. Analyses were done using Prism 7.0a and 8.3.1 (GraphPad). **P* $\leq$ 0.05, ***P* $\leq$ 0.01, ****P* $\leq$ 0.001, *****P* $\leq$ 0.0001.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <http://advances.sciencemag.org/cgi/content/full/7/10/eabe8671/DC1>

[View/request a protocol for this paper from Bio-protocol.](#)

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