EphrinB2 drives perivascular invasion and proliferation of glioblastoma stem-like cells

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Abstract Glioblastomas (GBM) are aggressive and therapy-resistant brain tumours, which contain a subpopulation of tumour-propagating glioblastoma stem-like cells (GSC) thought to drive progression and recurrence. Diffuse invasion of the brain parenchyma, including along preexisting blood vessels, is a leading cause of therapeutic resistance, but the mechanisms remain unclear. Here, we show that ephrin-B2 mediates GSC perivascular invasion. Intravital imaging, coupled with mechanistic studies in murine GBM models and patient-derived GSC, revealed that endothelial ephrin-B2 compartmentalises non-tumourigenic cells. In contrast, upregulation of the same ephrin-B2 ligand in GSC enabled perivascular migration through homotypic forward signalling. Surprisingly, ephrin-B2 reverse signalling also promoted tumourigenesis cell-autonomously, by mediating anchorage-independent cytokinesis via RhoA. In human GSC-derived orthotopic xenografts, EFNB2 knock-down blocked tumour initiation and treatment of established tumours with ephrin-B2-blocking antibodies suppressed progression. Thus, our results indicate that targeting ephrin-B2 may be an effective strategy for the simultaneous inhibition of invasion and proliferation in GBM.

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Introduction Glioblastoma (GBM) is the most common and aggressive type of primary brain tumour and one of the most lethal types of cancer (Chen et al., 2012b). Current therapies consist of maximally safe surgical resection followed by radio and chemotherapy. However, these are largely ineffective and
GBM invariably recur following treatment, resulting in a median survival of <15 months (Weathers and Gilbert, 2014).

A leading cause of GBM recurrence is the diffuse infiltration of the tumour cells into the surrounding brain parenchyma, which limits efficacy of surgery and radiotherapy (Cuddapah et al., 2014; Lamszus et al., 2003). In addition, GBM contain subpopulations of cells with stem cell properties termed glioblastoma stem-like cells (GSC), which are able to self-renew, differentiate into tumour-bulk cells and reconstitute a phenocopy of the original lesion upon transplantation (Venere et al., 2011). It is therefore increasingly recognized that GSC are critical players in tumour initiation and progression. Furthermore, GSC were shown to be intrinsically resistant to chemo- and radiotherapy and more invasive than non-stem tumour cells (Chen et al., 2012a; Cheng et al., 2011; Bao et al., 2006; Venere et al., 2011; Sadahiro et al., 2014). This suggests that GSC might be the primary cells within GBM responsible for infiltration and tumour recurrence following therapy and that GSC-targeting therapies should be beneficial for GBM treatment.

GBM invasion occurs along three main routes: the white matter tracts, the interstitial space of the brain and the perivascular space surrounding blood vessels (Scherer, 1938; Cuddapah et al., 2014). Invasion along the perivascular space is a favourable migration route because endothelial cells secrete chemoattractants, which actively recruit tumour cells to the vasculature (Montana and Sontheimer, 2011; Cuddapah et al., 2014). In addition, the perivascular space is enriched in migration-promoting ECM components and is fluid-filled, thereby opposing little physical resistance to invading tumour cells (Gritsenko et al., 2012; Cuddapah et al., 2014). Importantly, within GBM, GSC are particularly prone to perivascular invasion, likely due to their similarities with normal neural progenitors, which preferentially migrate along blood vessels during development and after injury in the adult (Cuddapah et al., 2014; Watkins et al., 2014). In agreement with this, GSC reside in perivascular niches and the majority of invading cells migrate along the host vasculature in xenograft GSC models of both mouse and human origin (Farin et al., 2006; Zagzag et al., 2008; Baker et al., 2014; Calabrese et al., 2007; Winkler et al., 2009; Watkins et al., 2014).
Besides infiltration, GBM/vascular interactions underlie two additional key tumourigenic mechanisms. First, tumour cell migration along pre-existing normal blood vessels is an important tumour vascularisation mechanism, known as vascular co-option, by which tumours gain access to oxygen and nutrients independent of angiogenesis (Donnem et al., 2013). Vascular co-option plays crucial roles during initial tumour growth, seeding of satellite lesions and tumour recurrence following therapy. Second, association with the perivascular niche provides GSC with important self-renewal and survival signals, which support GSC tumour-propagating abilities and therapeutic resistance (Charles and Holland, 2010). Therefore, interactions with the vasculature, particularly within the GSC compartment, are critical throughout gliomagenesis and were indeed proposed as promising therapeutic targets for GBM treatment, but the mechanisms remain poorly defined (Cuddapah et al., 2014; Vehlow and Cordes, 2013).

GSC share many properties with normal neural stem cells (NSC), such as stem cell markers expression (Sox2, Nestin, CD133, ALDH1, etc.) and multilineage differentiation and, importantly, extensive evidence indicates that NSC themselves can be cells of origin in GBM (Chen et al., 2012b; Venere et al., 2011; Chen et al., 2012a). However, GSC also significantly differ from their normal counterparts, in that they harbour transforming mutations that drive their tumourigenic properties, including deregulated proliferation and increased invasiveness (Engstrom et al., 2012). Therefore, the comparison of GSC carrying known mutations to otherwise genetically matched normal NSC, should inform disease mechanisms, enable genotype-phenotype correlation and may identify new therapeutic targets for GBM.

In this study, we generated a murine GBM model by sequentially introducing oncogenic lesions relevant to the human disease into normal NSC and compared vascular interactions of the resultant transformed GSC-like cells and immortalised parental cells to interrogate mechanisms of perivascular invasion. Using this system, we identified ephrin-B2 as a critical driver of perivascular invasion. ephrin-B2 is a member of the Eph/ephrin family of receptor tyrosine kinases and their membrane-bound ligands, a fundamental cell communication system with widespread roles in tissue development, maintenance and disease (Pasquale, 2010). Activation of Eph receptors by ephrin ligands on adjacent cells modulates cell behaviour, including migration, proliferation and stemness, by eliciting forward signalling downstream of Ephs and reverse signalling downstream of ephrins (Kullander and Klein, 2002). Deregulation of the Eph/ephrin system contributes to the pathogenesis of many types of cancer, including GBM (Pasquale, 2010). Indeed, EphA2 and EphA3 were shown to drive GSC self-renewal, whereas EphA2, EphA4, EphB2, ephrin-B2 and ephrin-B3 have all been linked to GBM invasion, through incompletely understood mechanisms (Day et al., 2014; Tu et al., 2012; Binda et al., 2012; Day et al., 2013; Nakada et al., 2010; 2011).

We found that ephrin-B2 expressed on vascular endothelial cells inhibits the migration of non-tumourigenic cells, resulting in cell confinement. In contrast, transformation to GSC-like cells overrides this tumour-suppressive mechanism to drive perivascular invasion. We show that this is caused by upregulation in GSC-like cells of the same ephrin-B2 ligand, which desensitizes the cells to vascular confinement by constitutively activating Eph forward signalling non-cell-autonomously. Furthermore, we discovered that ephrin-B2 reverse signaling also elicits tumour cell proliferation in the absence of normal anchorage signals by driving Rho-A-dependent cytokinesis in a cell-autonomous manner. Consistent with these important roles, ephrin-B2 overexpression was sufficient to fully transform immortalised NSC to the same extent as oncogenic Ras. In human GBM specimens, high Ephrin-B2 levels were detected in perivascular tumour cells with GSC features at the infiltrative tumour margin, indicative of a role in the GSC compartment in primary tumours. Remarkably, EFNB2 knockdown in primary human GSC isolated from patient material or treatment of established tumours derived from these GSC with anti-ephrin-B2 single chain blocking antibodies strongly suppressed tumourigenesis, by concomitantly inhibiting vascular association and proliferation. Thus, ephrin-B2 may be an attractive therapeutic target for the treatment of GBM.
Results

Endothelial ephrin-B2 compartmentalises immortalized, but not transformed, neural stem cells

To investigate mechanisms of GSC/vascular interactions in the context of syngeneic, immuno-competent brains, we sequentially introduced mutations commonly found in human GBM (RTK activation, p53 and RB inactivation) in primary murine SVZ NSC to generate fully transformed, GSC-like cells and genetically-matched immortalised NSC (Network, 2008). We used two complementary strategies for this. First, we used a ‘classical’ transformation paradigm previously shown to drive gliomagenesis in vivo, whereby NSC were immortalised with SV40 large-T antigen (imNSC1) and transformed with RasV12 (herein referred to as GSC1) to inactivate Trp53 and Rb, and mimic the increased Ras signalling that results from Nf1 loss, respectively (Blouw et al., 2003; Hahn et al., 1999; Sonoda et al., 2001; Huszthy et al., 2012). This approach allowed us to readily test candidate effectors by transforming NSCs isolated from mice carrying the specific mutation, as previously reported (Blouw et al., 2003). In the second approach, we induced transformation by defined genetic changes in the same pathways to rule out artifacts of oncogene overexpression. Nf1fl/fl NSCs were immortalised with p53 shRNAs and ectopic CDK4 to inactivate p53 and the p16/RB axis, respectively (imNSC2), and transformed by Cre-mediated Nf1 deletion (herein referred to as GSC2).

Unlike previously reported for SVZ NSC in vitro (Wang et al., 2012), increased Ras signalling did not cause premature glial differentiation of the NSC in vitro in either model (Figure 1—figure supplement 1a and Tables 1 and 2 and Supplementary files 1 and 2). In contrast, GSC1 and GSC2 retained stem cell properties in vitro as judged by high clonal efficiency in neurosphere culture and differentiation into glial and neuronal lineages upon mitogen withdrawal (Figure 1—figure supplement 1a–c). Furthermore, both cell types (but not their immortalised controls) formed colonies in soft agar (Figure 4c,d) and gave rise to highly aggressive tumours upon intracranial transplantation in immunocompromised mice (5/5 animals, 100% penetrance, median survival 24d for GSC1 and 38.5 for GSC2), whereas GSC1 also did so in syngeneic animals (5/9 animals, 56% penetrance, median survival 73d), indicative of a more aggressive phenotype. Consistent with their stem-like properties, clonal dilution experiments revealed that as little as 100 cells of either line was sufficient to generate aggressive tumours (Figure 1—figure supplement 1d). Importantly, the tumours recapitulated the histopathology and gene expression signatures of human GBM, including presence of necrosis, neovascularisation, nestin and Sox2 expression and a strong enrichment in the Verhaak mesenchymal subtype gene signature (Figure 1a–d and Figure 1—figure supplement 1e) (Klei et al., 2010; Verhaak et al., 2010). The tumours presented diffuse borders with the majority of invading cells migrating along blood vessels and displacing astrocyte endfeet and pericytes to come in direct contact with endothelial cells, as previously reported for both murine and human GSC (Figure 1b and Figure 1—figure supplement 1f) (Zagzag et al., 2008; Farin et al., 2006; Nagano et al., 1993; Watkins et al., 2014). Thus, GSC1 and 2 resemble mesenchymal glioblastoma stem-like cell lines and are highly similar, interchangeable model systems.

Next, as only GSC1 formed tumours in syngeneic mice, we used this model to assess interactions with the normal brain vasculature in real time in vivo using 2-photon microscopy. GFP-labeled GSC1 or imNSC1 were injected at a mimimum depth of 150–200 μm into the cortex of syngeneic recipients under a chronic cranial window and 5–7 days later their migration in relation to Dextran-TxRed-labeled blood vessels was imaged over a period of 6 hr (Figure 1e,f and Figure 1—figure supplement 1g) (Holtmaat et al., 2009). Although imNSC do not form tumors long-term, the cells were still present, viable and migratory at this time-point, as indicated by morphology and motility outside of the perivascular space. Consistent with previous reports, both cell types readily associated to the vasculature (Kokovay et al., 2010; Baker et al., 2014; Farin et al., 2006; Nagano et al., 1993; Watkins et al., 2014; Winkler et al., 2009). However, while individual GSC1 migrated out of the tumour mass along blood vessels, resulting in cell scattering, imNSC remained in stationary groups (Figure 1e,f). GSC1 perivascular migration was independent of their position relative to the tumour bulk or cell density, with single cells migrating from the main tumour as efficiently and at similar speed as GSC1 migrating at a distance from the tumour margin (Figure 1e and Figure 1—figure supplement 1h). Similarly, rare sparse imNSC further away from the tumour margin failed to migrate along blood vessels (Figure 1—figure supplement 1h). Importantly, this effect was not due to a general impairment in imNSC migration, as these cells migrated to a similar extent as GSC in vitro.
Figure 1. The vasculature compartmentalises immortalised neural but not glioma stem cells through endothelial ephrin-B2. (a) Representative H&E staining of GSC1 tumours 60 days after intracranial injection into nude mice. Neovascularisation (i), focal necrosis (ii) and increased cellular density (iii) Figure 1 continued on next page
can be observed. (b) Representative fluorescent images of GFP-labeled GSC1 tumours stained for the vascular marker CD31 (red) and GFP to identify tumour cells (green, left), the stem cell markers nestin (green, middle) and Sox2 (red, right), as indicated. The arrows indicate examples of GFP-labeled GSC1 that have migrated away from the tumour mass and are invading along the vasculature as single cells or in small groups. Scale bars = 50 μm. (c) Heatmap showing Verhaak subtype classification of 4 GSC1 and 3 GSC2-derived tumours in vivo. Colour scale with corresponding normalised enrichment scores is shown on the right. All tumours classified as mesenchymal with a nes >2.6. (d) Mean GSEA enrichment plot for the Verhaak mesenchymal gene signature in the GSC lines. Both nom p-val and FDR q-val = 0. (e) Intravital 2-photon micrographs of GFP-labeled imNSC1 and GSC1 cells injected into the cortex of wildtype (Efnb2 WT EC) and endothelial specific Efnb2 knockout mice (Efnb2 -/- EC) and imaged 7 days later over 6 hr through an intracranial window at a depth of 200 μm. The vasculature was labeled by tail-vein injection of Tx-red conjugated Dextran (3000 MW). Arrowheads indicate representative perivascular migration patterns for each genotype. Scale bar = 50 μm. (f) Quantification of the migrated distance of the cells depicted in (c). Each dot represents one cell. n indicates number of animals imaged. One way ANOVA with Tukey post hoc test. (g) Left: schematic representation of the experimental set up for in vitro migration assays with endothelial cells. Middle: merged fluorescent and phase contrast still images taken from time-lapse microscopy experiments of GFP-labeled imNSC1 and GSC1 (green) migrating towards brain microvascular endothelial cells (bmvEC, unlabeled) at the indicated time points. Right: quantification of boundary length at 60 hr. Students t-test. Scale bar = 500 μm. (h) Still fluorescence and phase contrast merged images of GFP-labeled imNSC2 and GSC2 migrating towards bmvEC (unlabeled, left) for 60 hr and quantification (right) as in (e). Error bars denote s.e.m., Students t-test. Scale bar = 500 μm. (i) Schematic representation of the experimental set up for in vitro migration assays toward recombinant ephrin-B2-Fc (left), phase contrast images (middle) and quantification (right) of imNSC1 and GSC1 migration against coated ephrinB2-Fc pre-clustered with fluorescently-labelled anti-Fc antibodies at 60 hr. Error bars denote s.e.m., Students t-test. Scale bar = 500 μm. Green dots denote boundary of ephrin-B2 coating identified by fluorescence. (j) Still images (left) and quantification (right) of GFP-labeled imNSC (GFP) migrating towards bmvEC (unlabeled) treated with control siRNA (Scr) or siRNA against Efnb2 (siEfnb2) for 60 hr. Scale bar = 500 μm. Error bars denote s.e.m., Students t-test. For this and later figures dots indicate individual data points and ***p<0.001; **p<0.01 and *<0.05. See also Figure 1—figure supplement 1 and Figure 1—source data 1.

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The following source data and figure supplements are available for figure 1:

Source data 1. Raw data for all quantification of NSC/GSC migrated distance and boundary length shown in Figure 1.

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Figure supplement 1. GSC1/2 resemble glioma stem-like cells.

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Figure supplement 1—source data 1. Raw data for all quantitative analyses shown in Figure 1—figure supplement 1.

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This suggests that signals from the vasculature compartmentalises non-tumourigenic cells to restrict their migration, whereas transformation overrides these signals to enable perivascular spread.

To test this more directly and identify potential effectors, we developed an in vitro cell migration assay that mimics the response of infiltrating GBM cells to initial contact with endothelial cells. We seeded both imNSC/GSC models and primary brain microvascular endothelial cells (bmvEC) in separate wells of culture inserts and assessed migration of the two cell types towards each other following insert removal by time-lapse microscopy. Remarkably, this assay closely recapitulated in vivo migratory patterns, in that endothelial cells strongly repelled and compartmentalised imNSC forming a sharp boundary (Figure 1 and Video 1). Similar effects were observed with normal NSC, indicating that compartmentalisation is not caused by immortalisation, but rather reflects the response of normal cells to the vasculature (not shown). In contrast, GSC1 were refractory to compartmentalisation and migrated over the endothelial monolayer, giving rise to an uneven and longer boundary (Figure 1g and Video 2). The behavior of GSC2 was identical to GSC1, confirming that it is a

Table 1. Gene set enrichment analysis FDR q-values for astrocyte, oligodendrocyte, OPC and neuron signatures in GSC lines vs NSC.

| Gene set enrichment analysis | FDR q-value (FDR) | GSC vs NS | GSC1 vs NS | GSC2 vs NS | Cell type term gene number |
|-----------------------------|------------------|-----------|------------|------------|---------------------------|
| GAGE analysis               | oligo            | 1         | 1          | 1          | 497                       |
|                             | neuron           | 1         | 1          | 1          | 497                       |
|                             | astro            | 1         | 1          | 1          | 486                       |
|                             | opc              | 1         | 1          | 1          | 490                       |

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general property of transformed cells (Figure 1h). Eph/ephrin signalling is one of most important mediators of cell-cell contact-dependent boundary formation and we reported that endothelial ephrin-B2, the most abundant ephrin-B2 in the endothelial cells in vivo and in vitro (Figure 1—figure supplement 1j), induces cell sorting of normal NSC (Cayuso et al., 2014; Ottone et al., 2014; Gale et al., 2001). We therefore tested the role of ephrin-B2 in endothelial-induced compartmentalisation by assessing imNSC1 migration towards recombinant ephrin-B2-Fc. As shown in Figure 1i and Videos 3 and 4, ephrin-B2-Fc compartmentalised imNSCs, but not GSC, to the same extent as endothelial cells. This effect was highly specific and was not due to general differences in migration between imNSC and GSC cells because imNSC migrated onto Fc peptides, ephrin-A1 or ephrin-A5-Fc ligands to a similar extent as GSC cells (Figure 1—figure supplement 1i). Conversely, Efnb2 knock-down in endothelial cells disrupted boundary formation against imNSC, indicating that endothelial ephrin-B2 is both necessary and sufficient for compartmentalisation (Figure 1j and Figure 1—figure supplement 1j).

To assess the role of vascular ephrin-B2 in vivo, we performed intravital imaging of imNSC implanted in inducible endothelial-specific conditional Efnb2 knock-out mice (Efnb2ΔEC), following postnatal tamoxifen-mediated recombination (Ottone et al., 2014). Strikingly, selective deletion of ephrin-B2 in the endothelium elicited robust perivascular migration of imNSC1, confirming that vascular ephrin-B2 compartmentalises non-tumourigenic cells in vivo (Figure 1e,f).

**Upregulation of ephrinB2 drives perivascular invasion of GSC**

As we reasoned that changes in Eph/ephrin levels might underlie the ability of GSC to escape ephrin-B2-mediated vascular compartmentalisation, we interrogated expression levels of all Ephs and ephrins in imNSC/1 and GSC/1/2 by qRT-PCR (Figure 2a and Figure 2—figure supplement 1a). Normal NSC controls were included to rule out p53 and Rb-dependent changes unrelated to perivascular migration. We found that p53 and Rb inactivation did not change Eph/ephrin expression

| GAGE analysis q-value (FDR) | NSvsGSC | NSvsGSC1 | NSvsGSC2 | Cell type term gene number |
|-----------------------------|---------|----------|----------|---------------------------|
| astro                       | 4.06848E-22 | 2.84603E-23 | 2.04565E-19 | 490 |
| opc                         | 1.57235E-16 | 7.01306E-17 | 3.24522E-14 | 486 |
| neuron                      | 1.51727E-06 | 4.55938E-07 | 4.58202E-05 | 497 |
| oligo                       | 0.064920265 | 0.06708047 | 0.222744258 | 497 |

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**Table 2.** Gene set enrichment analysis FDR q-values for astrocyte, oligodendrocyte, OPC and neuron signatures in NSC vs GSC lines.

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**Video 1.** Endothelial cells compartmentalise imNSC. Merged fluorescent and phase contrast movie from time-lapse microscopy experiments of cell-tracker labelled imNSC1 (green) migrating towards brain microvascular endothelial cells (bmvEC, unlabelled cells). Images were taken every 10 min for 60 hr.

**Video 2.** GSC escape endothelial compartmentalisation. Merged fluorescent and phase contrast movie from time-lapse microscopy experiments of cell-tracker labelled GSC1 (green) migrating towards brain microvascular endothelial cells (bmvEC, unlabelled cells). A slight decrease in fluorescence was observed due to higher proliferation rate of the cells. Images were taken every 10 min for 60 hr.
substantially. In contrast, elevated Ras signalling, triggered by either RasV12 expression or Nf1 loss, strongly downregulated three Ephs (Epha4, Ephb1 and Ephb2) and upregulated two ephrins (Efna5 and Efnb2) in both models. However, western analysis indicated that of these genes, only Ephb1, Ephb2 and Efnb2 changed significantly at the protein level (Figure 2b) and were thus further assessed in migration assays against ephrin-B2-Fc. Surprisingly, we found that re-introduction of Ephb1 or Ephb2 by overexpression in GSC1 (Figure 2—figure supplement 1b) did not affect migration towards recombinant ephrin-B2-Fc, indicating that changes in the complement of Eph receptors do not underlie unimpeded migration, as in other systems (Figure 2c) (Astin et al., 2010). Instead, genetic deletion of Efnb2 in GSC1 (Figure 2—figure supplement 1c) fully rescued boundary formation in response to ephrin-B2-Fc and endothelial cells in vitro and blocked perivascular migration in vivo (Figure 2c–e). Conversely, Efnb2 overexpression (Figure 2—figure supplement 1d) conferred imNSC1 with the ability to migrate over recombinant ephrin-B2 and a cultured endothelium and to escape vascular compartmentalisation in vivo (Figure 2c–e). Together, these results show that ephrin-B2 upregulation downstream of Ras underlies GSC evasion of ephrin-B2-mediated endothelial repulsion and invasion along the vasculature.

**Elevated ephrin-B2 desensitizes GSC to vascular repulsion by constitutively activating homotypic forward signalling**

To understand the mechanisms involved in ephrin-B2-driven perivascular invasion, we characterised the early events of imNSC1 and GSC1 migration towards ephrin-B2-Fc in greater detail. We noticed that, although all GSC cells eventually migrated over ephrin-B2-Fc, a proportion of the cells (~20%) are repelled on first contact (Figure 3a). Repelled cells invariably migrated towards ephrin-B2-Fc either as single cells or in small groups (<2 contacts), whereas non-repelled cells migrated in larger groups (>3 contacts) (Figure 3b). This was not the case in imNSC cultures, where all cells were equally repelled, regardless of the number of homotypic cell-cell interactions at initial contact. This suggested that homotypic cell-cell interactions among GSC, mediated by intrinsic ephrin-B2, might underpin evasion of extrinsic ephrin-B2 repulsion. To test this, we repeated the migration assay under conditions that disrupt GSC cell-cell contacts by inhibiting cadherin-based junctions (Ottone et al., 2014). Both culture in low Ca²⁺ media and dominant-negative N-cadherin overexpression completely disrupted cell-cell junctions without affecting cell motility (Figure 3—figure supplement 1a,b) and blocked GSC migration over ephrin-B2 ligands, indicating that homotypic cell-cell interactions are specifically required (Figure 3c).
Figure 2. Upregulation of ephrin-B2 in GSCs enables perivascular invasion (a) Quantitative RT-PCR analysis of indicated Eph receptors and ephrin ligands. Error bars denote s.e.m., p values of differences in expression between imNSC1 and GSC1 are shown. Multiple t-test analysis. (b) Western analysis of levels of the indicated proteins in normal neural stem cells (NSC), imNSC1 and GSC1. n = 3. (c) Kymographs from time-lapse experiments of the indicated cell types migrating against coated ephrinB2-Fc over 60 hr. Quantification of the kymographs is shown on the right. Error bars denote s.e. Figure 2 continued on next page.
We next asked whether homotypic ephrin-B2 signalling drives migration through activation of forward or reverse signalling within the GSC population. Ectopic expression of a C-terminal truncated form of ephrin-B2 (ΔCEfnb2), which lacks reverse signalling (Pfaff et al., 2008), promoted imNSC migration over ephrin-B2-Fc to the same extent as full-length Efnb2, demonstrating that forward signalling between neighboring cells is responsible (Figure 3d and Figure 2—figure supplement 1d). Indeed, western analysis (Figure 3e) indicated that while imNSC controls displayed low basal levels of activated Eph receptors (p-Eph), GSC exhibited constitutively high levels of p-Eph, despite their reduced levels of EphB1 and EphB2 (Figure 2b) and no increase in other ephrin-B2 receptors in these cells (Figure 2a and not shown). High basal p-Eph could be mimicked by ephrin-B2 overexpression in imNSC and reduced to basal control levels by culture in low Ca\(^{2+}\) (Figure 3e). In addition, the high phosphorylation levels of Ephs in GSC appeared to be saturating, as, in contrast to imNSC, co-culture with endothelial cells or treatment with recombinant ephrin-B2 failed to further activate Eph receptors in these cells (Figure 2a and not shown). High basal p-Eph could be mimicked by ephrin-B2 overexpression in imNSC and reduced to basal control levels by culture in low Ca\(^{2+}\) (Figure 3e). In addition, the high phosphorylation levels of Ephs in GSC appeared to be saturating, as, in contrast to imNSC, co-culture with endothelial cells or treatment with recombinant ephrin-B2 failed to further activate Eph receptors in these cells (Figure 3f). Importantly, Eph desensitization was again entirely dependent on endogenous ephrin-B2 because genetic deletion of Efnb2 in GSC fully restored Eph stimulation by exogenous ephrin-B2. Consistent with the known ability of Ephs to trigger repulsion (Astin et al., 2010; Pasquale, 2010), such high basal levels of activated Eph resulted in GSC undergoing much greater scattering and faster migration than imNSC following homotypic collisions in sparse monocultures, in an ephrin-B2-dependent manner (Figure 3—figure supplement 1c,d). Thus, high ephrin-B2 levels in GSC result in constitutive activation of Eph forward signalling and desensitization to exogenous stimulation.

To further test whether this desensitization mechanism underlies the ability of GSC to override ephrin-B2-mediated repulsion, we pre-stimulated imNSC with recombinant ephrin-B2-Fc to strongly activate Eph forward signalling prior to their migration out of the insert (Figure 3f) and assessed migration towards coated ephrin-B2 in boundary assays. Parallel Fc-treated cultures served as controls. Remarkably, we found that ephrin-B2-Fc (but not Fc) pretreatment enabled migration over the ephrin-B2 boundary, confirming that activation of Eph forward signaling on first contact with exogenous ephrin-B2 is the mechanism involved. In agreement with these findings, immunofluorescence analysis of GSC1-derived orthotopic tumours revealed high pEph levels in cells migrating along blood vessels at the tumour edge, indicating that constitutive forward signaling also contributes to perivascular invasion in vivo (Figure 3—figure supplement 1e).

We conclude that GSC escape from endothelial compartmentalisation depends on continuous activation of Eph forward signalling elicited by elevated ephrin-B2 through homotypic cell-cell interactions within the tumour cell population. This, in turn, desensitises the receptors to further activation by extrinsic ephrins, thereby overriding the repulsion by endothelial ephrin-B2 and enabling unimpeded perivascular migration.
Figure 3. ephrin-B2 upregulation drives perivascular migration by saturating Eph forward signalling through homotypic cell-cell interactions. (a) Quantification of cell behavior upon initial contact with coated ephrin-B2. n = 3, error bars denote s.e.m. (b) Left: representative images taken from Kruschke et al. eLife 2016;5:e14845. DOI: 10.7554/eLife.14845.
Figure 3 continued

videos of GSC1 cells as they first come in contact with coated ephrin-B2 (green). Right: quantification of the number of homotypic GSC1 cell–cell contacts at time of initial interaction with ephrin-B2. A minimum of 50 cells were counted in each experiment. Error bar denotes St.D, Student t-test. (c) Representative kymographs (left) and quantifications (right) of GSC1 migrating towards ephrin-B2 that were either cultured in normal media (ctrl) or low Ca\[^{2+}\] conditions (low Ca\[^{2+}\]), or transduced with GFP control (GFP) or dominant negative N-Cadherin (DN-Ncad) adenoviral constructs. Dotted lines demarcate coated ephrin-B2 boundary. n = 3, error bars denote s.e.m., Student t-test. (d) Kymographs and quantifications of the responses of imNSC1 and imNSC1-\(\Delta\text{Efnb2}\) to coated ephrin-B2 ligands. n = 3, error bars denote s.e.m., Student t-test. (e) Western analysis of the levels of activated Eph receptors (p-Eph) in the indicated cell types cultured in either normal growth media (ctrl) or calcium depleted (low Ca\[^{2+}\]) conditions. β-tubulin served as loading control. n = 3 (f) Western blots of p-Eph levels in indicated cells cultured either with control proteins (Fc), endothelial cells (Endo) or ephrin-B2-Fc (ephrin-B2-Fc) for 18 hr. β-tubulin is used as loading control. n = 3 (g) Phase contrast images (left) and quantification (right) of imNSC1 migration towards coated ephrin-B2-Fc following pre-treatment with either control (Fc) or clustered ephrin-B2-Fc to activate Eph forward signalling. Experiments were stopped at 48 hr to assure maximal stimulation of the cells throughout the assay. Green dots denote boundary of ephrin-B2 coating identified by fluorescence. n = 3, error bars denote s.e.m., Students t-test. Scale bar = 250 μm. See also Figure 3—figure supplement 1 and and Figure 3—source data 1.

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The following source data and figure supplements are available for figure 3:

Source data 1. Raw data for all quantifications of NSC/GSC migration assays shown in Figure 3.

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Figure supplement 1. Increased repulsion between ephrin-B2 expressing cells leads to greater migration velocity and distance.

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Figure supplement 1—source data 1. Raw data for all quantitative analyses shown in Figure 3—figure supplement 1.

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ephrin-B2 transforms imNSC

Given these important roles in invasion, we asked whether ephrin-B2 might also affect tumorigenesis by performing tumourigenicity studies of luciferase-tagged imNSC1, imNSC1-\(\text{Efnb2}\), GSC1 and GSC1-\(\text{Efnb2}\)\(^{-/-}\) implanted orthotopically in nude mice. We used immunocompromised mice for these experiments because the incomplete tumour penetrance of GSC in syngeneic animals precludes rigorous assessment of their tumourigenicity in the syngeneic model. Strikingly, quantitative imaging and survival analysis revealed that, while imNSC1 did not form tumours and GSC1 formed tumours at full penetrance, \(\text{Efnb2}\) deletion strongly suppressed GSC1 tumour growth and, conversely, \(\text{Efnb2}\) overexpression was sufficient to fully transform imNSC (Figure 4a,b). Specifically, imNSC1-\(\text{Efnb2}\) tumours resulted in a similar median survival as GSC1 (28d). In contrast, all imNSC animals survived tumour-free beyond 200 days and 7 out of 9 animals of the GSC1-\(\text{Efnb2}\)\(^{-/-}\) cohort developed lesions with much slower kinetics, while the remaining 2 mice remained tumour-free, as confirmed by post-mortem examination. These effects were not due to a general loss of stem-like properties due to ephrin-B2 deletion, as clonal efficiency remained unaffected in GSC1-\(\text{Efnb2}\)\(^{-/-}\) (Figure 4c). To rule out the possibility that the transforming ability of ephrin-B2 might be a peculiarity of the IgT-Ras model, we repeated the soft agar assays with imNSC2, imNSC2-\(\text{Efnb2}\), imNSC2-\(\Delta\text{Efnb2}\), GSC2 and GSC2 knock-down for \(\text{Efnb2}\) and obtained identical results, confirming the generality of these findings and the functional equivalence of the two GSC lines (Figure 4d and Figure 4—figure supplement 1b).

We next asked how ephrin-B2 overrides anchorage checkpoints by comparing cell cycle progression of immortalised and transformed cells in suspension. We seeded imNSC1, imNSC1-\(\text{Efnb2}\),
**Figure 4.** Ephrin-B2 is a glioblastoma oncogene controlling anchorage independent proliferation. (a) Representative bioluminescent images of nude mice 20 days after intracranial injection of imNSC1, imNSC1-Efnb2, GSC1, GSC1-Efnb2−/−. (b) Kaplan-Meier survival plots of the mice depicted in (a). 

Figure 4 continued on next page.
Significance is given relative to imNSC1 for imNSC1-Efnb2 and to GSC1 for GSC1-Efnb2−/−. Log Rank Mantel Cox. (c, d) Left: Representative micrographs of GSC1 (q) and GSC2 (d) cells of indicated genotype grown in soft agar for 10d. Right: quantification of number of colonies formed in soft agar in all cultures, expressed as percentage over total number of seeded cells. n = 3, error bars depict s.e.m. One way ANOVA with Tukey correction. (e) Left: representative FACS profiles of cells grown in attachment or methylcellulose for 72 hr, showing DNA content by propidium iodide (PI) staining. Right: quantification of cell cycle phases from the FACS profiles. n = 3–5 as indicated by the dots. Error bars depict s.e.m., p values indicate significance of changes in G2/M phase. (f) Representative PI FACS profiles and quantifications of imNSC and GSC1 isolated from brain tissue 7 days after intracranial injection. n = 3. Error bars depict s.e.m. One way ANOVA with Tukey post hoc test shown for G2/M phase. See also Figure 4—figure supplement 1 and Figure 4—source data 1.

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The following source data and figure supplements are available for figure 4:

Source data 1. Raw data for Kaplan Meier analysis, number of colonies formed in soft agar and cell-cycle analysis presented in Figure 4
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Figure supplement 1. Anchorage-independent proliferation is independent of homotypic cell-cell contacts.
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Figure supplement 1—source data 1. Raw data for all quantitative analyses shown in Figure 4—figure supplement 1
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GSC1 and GSC1 Efnb2−/− in adhesion or methylcellulose culture for 72 hr and measured their PI-FACS profiles (Cremona and Lloyd, 2009). As anticipated, all cells proliferated efficiently in attachment (Figure 4e). By contrast, their behavior in suspension was very distinct. While GSC1 continued to proliferate in methylcellulose with similar kinetics, suspended imNSC did not. This was not due to cell death as judged by activated caspase3+ staining (Figure 4—figure supplement 1c). Instead, imNSC arrested with a 4n DNA content. This indicated that while imNSC progress normally through the G1 and S phases of the cell-cycle, their progression through G2/M is blocked by an anchorage checkpoint, which is overridden by activated Ras.

Importantly, Ras-mediated progression through G2/M was again dependent on ephrin-B2 reverse signalling, because Efnb2 deletion in GSC re-instated the G2/M arrest (without additional apoptosis, Figure 4—figure supplement 1c), and the cell-cycle arrest of imNSC could be rescued by overexpression of full-length, but not ΔC, Efnb2 (Figure 4e). To assess the relevance of these findings to in vivo tumourigenesis, 7d after intracranial implantation imNSC1, imNSC1-Efnb2, GSC1 and GSC1Efnb2−/−, were recovered from the injected brains and their PI profiles analysed (Figure 4f). Strikingly, the cell-cycle profiles of all cells were indistinguishable from the corresponding methylcellulose cultures, confirming that ephrin-B2 drives gliomagenesis in vivo by promoting proliferation in the absence of normal anchorage signals.

**ephrin-B2 drives anchorage-independent cytokinesis**

A previous study reported that human fibroblasts cultured in suspension arrest at cytokinesis and that oncogenic Ras can bypass this arrest (Thullberg et al., 2007). We therefore explored the role of cytokinesis in our system by pulsing all attached and suspended cultures with EdU to distinguish G2/M arrested from cycling cells completing mitosis and staining for phalloidin to detect cortical actin. As shown in Figure 5a and b, we found that all methylcellulose cultures devoid of ephrin-B2 reverse signalling (imNSC1, imNSC1-ΔCEfnb2 and GSC1 Efnb2−/−) contained a much larger proportion of EdU+ binucleated cells with decondensed chromatin compared to adherent conditions, indicative of a cytokinesis block (Thullberg et al., 2007). In contrast, cells with intact ephrin-B2 reverse signalling (imNSC1-Efnb2 and GSC1) had similar low percentages of EdU+ binucleated cells in both suspension and attachment culture. Thus, ephrin-B2 reverse signalling drives anchorage-independent cytokinesis.

Of the known effectors of ephrin-B2, Src and Rhoa have been previously linked to anchorage independent proliferation, with Rhoa also regulating constriction of the contractile ring during cytokinesis (Daar, 2012; Desgroisselier et al., 2009; Jordan and Canman, 2012). We therefore assessed Src and Rhoa in ephrin-B2-mediated anchorage-independent cytokinesis by measuring levels of their activated forms (phosphorylated Src, p-Src, and GTP-bound Rhoa) in suspension by western blot and pull-down assays, respectively. p-Src was undetectable in suspended imNSC1, imNSC-Efnb2 and GSC1 and, similarly, levels of activated FAK, a critical Src substrate, were also reduced (Figure 5—figure supplement 1a), indicating that Src does not play a role in our system. In stark
Figure 5. ephrin-B2 mediates cytokinesis in the absence of anchorage through RhoA. (a) Representative fluorescent images of binucleated cells grown in suspension for 24 hr and stained with phalloidin (green) to label cortical actin and EdU (red) to distinguish cycling from arrested cells. Scale bar = 20 \mu m. (b) Quantification of binucleated cells in indicated attached and suspended cultures. Values represent percentages over total number of cells. n = 4, error bars denote s.e.m. Two way ANOVA with Tukey post hoc test. (c) Western analysis of RhoA pulldown assays showing levels of activated RhoA (RhoA-GTP) and total RhoA levels in indicated cells grown in suspension for 24 hr. Bottom graph shows quantifications of activated RhoA levels from the western blots. n = 3. Error bars indicate s.e.m. (d) Quantification of the PI cell cycle profile as measured by FACS. Cells transfected with either constitutively active RhoA (V14) or dominant negative RhoA (N19) were grown in suspension for 24 hr. n = 3, error bars depict s.e.m. One way ANOVA with Tukey post hoc test was used to calculate p values for differences in G2/M phase of each suspended culture relative to corresponding attached cultures. See also Figure 5—figure supplement 1 and Figure 5—source data 1.

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The following source data and figure supplement are available for figure 5:

Source data 1. Raw data for quantifications of binucleated cells and cell cycle analysis presented in Figure 5.
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Figure supplement 1. Src and FAK do not mediate anchorage-independent proliferation downstream of ephrin-B2.
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contrast, levels of active RhoA were low in suspended cells that lacked ephrin-B2 reverse signalling (i.e. imNSC, imNSC1-ΔEfnb2 and GSC1Efnb/−) but greatly increased in cells that express high levels of full-length ephrin-B2 (i.e. imNSC1-Efnb2 and GSC1), suggesting the involvement of RhoA (Figure 5c). To further test this, we overexpressed constitutively active (RhoA-V14, which increases levels of active GTP-bound RhoA, Figure 5—figure supplement 1b) and dominant negative (RhoAN19, which inhibits activation of endogenous RhoA, Figure 5—figure supplement 1b) (Qi et al., 1995) forms of RhoA in imNSC1/GSC1Efnb2/− and imNSC1-Efnb2/GSC1, respectively, and assessed the ability of the cells to proliferate anchorage-independently by FACS. Remarkably, RhoA-V14 could rescue the cell-cycle arrest of imNSC1 and GSC1Efnb2/−, whereas RhoA-N19 arrested imNSC1-Efnb2 and GSC1 in G2/M (Figure 5d and Figure 5—figure supplement 1c). Together, these results demonstrate that ephrin-B2 drives anchorage-independent cytokinesis of GSCs through RhoA-mediated reverse signalling.

**Ephrin-B2 drives invasion and anchorage-independent proliferation of human mesenchymal GSC**

We have shown that ephrin-B2 drives two key aspects of tumour formation in our murine models: GSC perivascular invasion and proliferation. We thus sought to determine the relevance of these findings to human GBM (hGBM). To this end, we first examined Ephrin-B2 levels at the infiltrative margin of 10 GBM patient specimens and correlated them with GSC marker expression (Figure 6a, Figure 6—figure supplement 1a and Table 3). Two of these tumours were the original lesions from which G19 and G26 human GSC lines used below have been isolated. Serial sections were stained with H&E and with antibodies against Ephrin-B2 and ALDH1, a stem cell marker which labels GSC in perivascular and hypoxic niches (Rasper et al., 2010). All tumours presented cytoplasmic and membranous Ephrin-B2 expression in 25 to 90% of the tumour cells, as well as neurons, inflammatory cells and vascular endothelial cells, as reported (Gale et al., 2001; Sawamiphak et al., 2010; Ottone et al., 2014). In contrast, ALDH1 expression was less abundant, with expression detected predominantly in neoplastic cells surrounding blood vessels, in perinecrotic regions and at the infiltrative tumour margin, as expected from a stem cell marker (Rasper et al., 2010). Weaker ALDH1 expression was also detected in reactive astrocytes and endothelial cells. Importantly, many perivascular ALDH1+ neoplastic cells at the infiltrative tumour margin co-expressed ephrin-B2. Thus, ephrin-B2 is expressed in stem-like cells invading along blood vessels in primary human tumours.

We next assessed EFNB2 expression in a panel of 8 well-characterised primary human GSC lines isolated from independent patient tumours, including two of the tumours analysed by IHC above (Caren et al., 2015). These lines recapitulate the transcriptional subtypes of primary GBM and predominantly cluster into proneural and mesenchymal subtypes, as previously reported for GSC (Figure 6 b and Figure 6—figure supplement 1b) (Bhat et al., 2013; Mao et al., 2013). Consistent with our mouse models, RNA-sequencing analysis revealed frequent upregulation of EFNB2 in GSCs and a significant correlation between EFNB2 and mesenchymal gene expression levels (Figure 6b). In addition, analysis of 402 GBM from the TGCA dataset classified according to Verhaak et al. and corrected for IDH1 status, indicated that EFNB2 expression levels are highest in mesenchymal and classical GBM subtypes (Figure 6—figure supplement 1c) (Verhaak et al., 2010). When tumours were divided into two groups on the basis of EFNB2 levels relative to median expression within subtypes, EFNB2 correlated with decreased survival in mesenchymal GBM (Figure 6—figure supplement 1d). Together, these results are indicative of a functional role for ephrin-B2 in GSC tumorigenesis within human GBM, specifically of mesenchymal subtype.

To test this more directly, we introduced control scrambled shRNAs or shRNAs to EFNB2 in 3 of the mesenchymal GSC lines described above (Figure 6—figure supplement 1e) and assessed effects of Ephrin-B2 depletion on invasion and proliferation in vitro. As shown in Figure 6c–h, we found that, similar to the murine GSC models, in the absence of ephrin-B2, GSC (but not SCR shRNA-transduced controls) lost the ability to migrate over coated ephrin-B2 and failed to proliferate anchorage-independently, resulting in a G2/M cell-cycle arrest. These effects were not caused by an impairment of GSC self-renewal following ephrin-B2 downregulation and were highly specific, as overexpression of an shRNA-resistant EFNB2 construct in G26 shEFNB2 cells fully rescued their anchorage-independent proliferation (Figure 6—figure supplement 1f–h).
Figure 6. Ephrin-B2 is expressed in human GSC and drives their invasion and proliferation in vitro. (a) Haematoxylin-eosin (H&E), ALDH1 and Ephrin-B2 HRP staining of infiltrative margins of patient tumours. Three different glioblastomas of 10 analysed are shown. Atypical tumour cells expressing both

*Figure 6 continued on next page*
EFNB2 silencing or anti-ephrinB2 scFv Ab treatment suppress human GSC tumourigenesis

To test effects of ephrin-B2 on hGSC tumourigenicity in vivo, we used two complementary strategies. First, we xenotransplanted luciferase-tagged SCR or shEFNB2 G26 cells intracranially in nude mice and monitored tumour growth and disease-free survival by bioluminescence imaging and survival analysis (Figure 7a–c). We found a dramatic impairment of tumour growth in the EFNB2 knock-down group relative to control, in that vector-transduced G26 formed GBM in all animals (as previously reported (Stricker et al., 2012) whilst EFNB2 knock-down cells gave rise to slower growing tumours in only 2 out of 10 animals, with the other 8 mice remaining tumour-free for >1 year. To investigate the mechanisms responsible and confirm the generality of these findings, we repeated these experiments using both G26 and G19 lines and examined vascular association and proliferation of the GSC by immunofluorescence and FACS analysis at 10 days following implantation, a time point at which knock-down cells could still be detected by bioluminescence (Figure 7b). SCR GSC associated with the vasculature at this early time point (and at later stages of tumour growth), resulting in vascular co-option, as reported (Watkins et al., 2014). In contrast, vascular contact was severely compromised in knock-down cells.

Table 3. Patient information for tumours 1–8 used for Ephrin-B2 IHC

| name  | gender | age  | site              | IDH1/2 | MGMT | ATRX | Grade |
|-------|--------|------|-------------------|--------|------|------|-------|
| case 1| M      | 69   | left temporal     | wt     | 0    | retained | IV    |
| case 2| M      | 45   | left temporal     | wt     | 0    | retained | IV    |
| case 3| M      | 54   | right temporo-parietal | wt     | <10% | retained | IV    |
| case 4| M      | 55   | right temporal    | wt     | 0    | retained | IV    |
| case 5| F      | 61   | right parietal    | wt     | 25%  | retained | IV    |
| case 6| M      | 57   | right frontal     | G395A  | 0    | lost   | IV (secondary) |
| case 7| F      | 73   | left frontal / crossed midline | wt | 0    | retained | IV    |
| case 8| F      | 39   | left temporo-parietal | wt     | 0    | retained | IV    |

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The following source data and figure supplements are available for figure 6:

Source data 1. Raw data for quantifications of kymographs, number of colonies formed in soft agar and cell-cycle analysis of human GSC presented in Figure 6.

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Figure supplement 1. EphrinB2 expression is increased and correlates inversely with patient survival in mesenchymal human GBM.

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Figure supplement 1—source data 1. Raw data for all quantitative analyses shown in Figure 6—figure supplement 1.

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Figure 7. EFNB2 silencing abolishes hGSC tumorigenicity. (a) Representative bioluminescent images of nude mice injected at d0 with $10^5$ luciferase-labeled G26 cells, stably expressing scrambled shRNA (SCR, top) or shRNA directed against EFNB2 (shEFNB2, bottom). (b) Quantification of luciferase bioluminescence measured at the indicated time points in both groups. n = 9 for ctrl and 10 for shEFNB2, error bars denote s.e.m. Two-way ANOVA with Tukey correction. (c) Kaplan-Meier survival plots of the mice depicted in (a). n = 5 for SCR, 10 for shEFNB2. Log Rank Mantel Cox test. (d) Representative fluorescence images (left) and quantification (right) of GSC/vascular interactions in tumors derived from G19 SCR and G19 shEFNB2 cells at 10 days post-implantation. Sections were stained for GFP to identify tumor cells and CD31 (red) to label pre-existing blood vessels. Arrows indicate vascular association and co-option in SCR tumours, which is reduced in shEFNB2 tumours. (e) Quantification of percentages of Ki67+ cells over Figure 7 continued on next page.
Furthermore, similar to GSC1Efnb2−/− murine cells, EFNB2 downregulation resulted in a marked decrease in proliferation, accompanied by an arrest in the G2/M phase of the cell-cycle, indicative of a cytokinesis block (Figure 7e,f). Thus, Ephrin-B2 drives tumour initiation by mediating vascular association and proliferation of human GSC.

Second, we asked whether Ephrin-B2 inhibition could suppress tumourigenesis of pre-established tumours. For this, we took advantage of an Ephrin-B2 blocking scFv antibody fragment (B11) we previously developed (Abengozar et al., 2012). Migration and methylcellulose assays in vitro confirmed that B11 effectively inhibits Ephrin-B2 signalling in GSC1 (Figure 8—figure supplement 1a,b). Luciferase-tagged G26 cells were implanted intracranially in immunocompromised mice and 13d later, once sizeable, exponentially growing tumours had formed, but prior to the onset of neoangiogenesis, B11 was administered intravenously for a total of 9 consecutive days, as reported (Abengozar et al., 2012; Binda et al., 2012). This treatment regimen enabled us to assess direct effects of B11 on the tumour cells in the absence of confounding anti-angiogenic effects known to result from ephrin-B2 inhibition (Sawamiphak et al., 2010; Abengozar et al., 2012). Efficient delivery of the scFv across the blood/tumour barrier to the tumour cells was confirmed in parallel animals using Alexa-680 labelled B11 (Figure 8—figure supplement 1c). Remarkably, B11 strongly suppressed tumour growth in all animals without any evidence of toxicity, with three of six animals showing complete regression, as judged by quantitative bioluminescence imaging, survival analysis and post-mortem examination of the injected brains (Figure 8a–c). Analysis and quantification of the tumours immediately following treatment also revealed that, in contrast to vehicle-treated control tumours, invading tumour cells failed to associate with and coopt the vasculature (Figure 8d). Furthermore, proliferation was reduced in B11 tumours relative to controls, with B11 samples containing many Ki67+ multinucleated cells, indicative of a cytokinesis defect (Figure 8e). Therefore, B11 suppresses G26 tumourigenicity through inhibition of ephrin-B2-dependent perivascular invasion and anchorage-independent proliferation, independent of angiogenesis. We conclude that Ephrin-B2 plays an important role in the pathogenesis of human GBM and its inhibition might be an effective strategy for curtailing GBM progression and recurrence.

Discussion

Perivascular invasion is a critical mechanism of GBM growth and infiltration, which greatly contributes to the marked therapeutic resistance of these tumours (Cuddapah et al., 2014; Scherer, 1938). Here, using a novel mouse ‘progression series’ that mimics transformation of normal NSC to mesenchymal GSC, we identified ephrin-B2 as a critical driver of GSC perivascular invasion. Interestingly, a role for ephrin-B2 and its phosphorylation in the tumour cells has been previously linked to GBM invasiveness, albeit not in the context of GSC. This suggests that ephrin-B2 might promote invasion both by generally enhancing cell-intrinsic invasive potential through reverse signalling (Nakada et al., 2010; Tu et al., 2012) and, also, by specifically enabling perivascular spread through forward signalling as we have shown here.

We found that ephrin-B2 expressed on vascular endothelial cells compartmentalises non-transformed cells. A similar role for Eph/ephrin signalling in constraining migration of premalignant cells was reported in colorectal cancer, where ephrin-B ligands in the surrounding normal tissue inhibit invasion of incipient lesions through activation of EphB receptors in the tumour cells (Cortina et al.,...
Figure 8. Treatment with anti-EphrinB2 ScFv blocking antibodies suppresses progression of established GBMs. (a,b) Representative images (a) and quantification (b) of bioluminescence radiance of PBS or anti-Ephrin-B2 scFv antibody (B11) injected mice. n = 4 for PBS ctrl and 6 for B11 treatment groups. (c) Kaplan-Meier survival plots of the mice depicted in a and b. n = 4 for PBS-treated, 6 for B11-treated G26 tumours. Log Rank Mantel Cox test. (d) Representative immunofluorescence images (left) of PBS and B11-treated G26 tumours 5 days after the first B11 injection, stained for GFP.
(green) to identify tumour cells and CD31 (red) to label the endogenous vasculature. Quantification of vascular association is shown on the right. Arrows indicate co-opted blood vessels in PBS-treated tumours. Scale bar 50 μm. Error bars denote s.e.m. Student t-test. (e) Representative immunofluorescence images (left) PBS and B11-treated G26 tumours stained for GFP and the proliferation marker Ki67. Quantification of the percentage of Ki67+ cells over total number of GFP+ tumour cells is shown on the right. Note the presence of multinucleated cells (arrows and inset) in B11-treated tumours. Scale bar~25 μm, n = 3, error bars denote s.e.m. Students t-test. See also Figure 8—figure supplement 1 and Figure 8—source data 1.

**Source data 1.** Raw data for quantifications of tumour growth by bioluminescence analysis, survival by Kaplan-Meier analysis, tumour cell interactions with the vasculature and Ki67 labelling of human GSC-derived tumours presented in Figure 8.

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The following source data and figure supplements are available for figure 8:

**Figure supplement 1.** B11 crosses the blood/tumour barrier and inhibits Ephrin-B2 on tumour cells.

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**Figure supplement 1—source data 1.** Raw data for all quantitative analyses shown in Figure 8—figure supplement 1.

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Our study is also consistent with work by Astin et al., which reported that ephrin ligands on stromal cells repel normal and non-metastatic prostate cancer cells (Astin et al., 2010). Thus, the present work further strengthens the notion that ephrins expressed in normal tissues act as tumour-suppressors during early tumourigenesis by promoting tumour confinement and identifies the vasculature as a critical mediator of these effects. It is tempting to speculate that ephrin-B2-mediated vascular compartmentalisation might be a more general tumour-suppressive mechanism, limiting the spread of premalignant lesions across many cancer types.

In contrast, malignant transformation overrides vascular compartmentalisation to enable stereotypical GBM invasion along the perivascular space (Baker et al., 2014; Farin et al., 2006; Watkins et al., 2014; Winkler et al., 2009). Mechanistically, we found this to be dependent on the upregulation of ephrin-B2 in the GSC themselves, which elicits constitutive activation of Eph forward signalling in the tumour through homotypic cell-cell interactions. Consequently, this promotes perivascular spread in two ways. First, constitutive Eph activation increases repulsion between the tumour cells within the tumour bulk, thereby promoting GSC motility and scattering of single cells away from the tumour, a hallmark of GBM migration (Vehlow and Cordes, 2013). Second, it desensitizes the cells to extrinsic ephrin-B2 repulsion, thereby circumventing vascular confinement and permitting cells that have detached from the tumour mass to continue their migration along the vasculature (Figure 8—figure supplement 1d). Thus, invasive GSC augment their tumourigenic potential by ‘hijacking’ the same signalling pathway through which the vasculature suppresses tumourigenesis in normal tissues. Intriguingly, the mechanism of invasion we identified here, differs from what has been reported for colorectal and prostate cancer in the studies mentioned above (Astin et al., 2010; Cortina et al., 2007). Indeed, while in those systems tumour spread depended on the downregulation of Eph receptors in a cell-autonomous manner, in our GBM model non-cell-autonomous constitutive activation of Eph forward signalling is responsible, suggesting that the mechanisms that evade ephrin repulsion during tumour progression might be critically dependent on cancer type.

We further report that ephrin-B2 is also a critical effector of the ‘Ras-transformed phenotype’, capable of both transforming immortalised cells and mediating, at least in part, Ras-dependent transformation. Indeed, IgT-immortalised stem cells overexpressing Efnb2 formed aggressive tumours with nearly identical kinetics to Ras-transformed cells and genetic deletion of Efnb2 in GSCs significantly delayed tumourigenesis. We demonstrate that these effects depend on the surprising and previously unknown ability of ephrin-B2 to drive anchorage-independent proliferation through the activation of cytokinesis in a RhoA-dependent manner. It is of note that, in contrast to perivascular invasion, these ephrin-B2 effects are mediated by reverse signalling and independent of cell-cell contact, indicating that ephrin-B2 drives two key aspects of GBM tumourigenesis through a complex interplay of cell-autonomous and non-cell-autonomous mechanisms (Figure 8—figure supplement 1d). Previous studies in endothelial and smooth muscle cells have reported a similar cell-cell contact- and Eph-independent role for ephrin-B2 reverse signalling in driving acto-myosin contractility and cell spread, respectively (Bochenek et al., 2010; Foo et al., 2006). We currently do not know how
RhoA becomes activated downstream of ephrin-B2. A potential candidate is Dishevelled, which mediates activation of Rho/ROCK by ephrin-B1 (Tanaka et al., 2003). It would be interesting to explore the role of Dishevelled in our system in the future.

Our results in GBM specimens and primary human GSC lines strongly support a critical role for ephrin-B2 in the human disease through the same mechanisms that we identified in the murine models. Indeed, EFNB2 knock-down prior to implantation abrogated tumour initiation and treatment of pre-existing intracranial tumours with Ephrin-B2 blocking antibodies, under conditions that mimic human therapeutic paradigms, strongly reduced the growth and expansion of pre-formed tumours. Significantly, both approaches curtailed tumour growth by impairing both vascular association and cytokinesis. In patient samples stratified by GBM subtype, EFNB2 levels were highest in mesenchymal and classical tumours and correlated, inversely, with survival specifically in mesenchymal GBM. This is in agreement with our mesenchymal mouse models and analysis of human GSC lines, which revealed a robust correlation between EFNB2 and mesenchymal gene expression levels. This recurrent correlation is particularly noteworthy, as EFNB2 was previously identified as a component of the core mesenchymal gene network (Carro et al., 2010). Furthermore, increasing evidence suggests that conversion to the mesenchymal signature is associated to progression, radioresistance and recurrence and is thus a fundamental driver of GBM malignancy (Ozawa et al., 2014; Bhat et al., 2013).

GSCs are thought to be largely responsible for tumour infiltration and propagation (Chen et al., 2012a; Venere et al., 2011). As our work is based on murine and human GSC models of GBM, and Ephrin-B2 levels are high in the GSC compartment in primary tumours (Figure 6a), our findings indicate that anti-ephrin-B2-based therapies would target the most critical subset of cells within these lesions. As such, by harnessing the complexity of the Eph/ephrin system, inhibition of ephrin-B2 would in itself be an effective ‘combinatorial therapy’, capable of suppressing two critical GSC-intrinsic properties, namely perivascular invasion and proliferation, and might thus represent an attractive strategy for blocking GBM progression and recurrence. In addition, as ephrin-B2 levels are robustly increased in GSC compared to normal neural stem cells and other tissues, such therapies should be relatively tumour-specific and non-toxic.

Materials and methods

Animals

All animal work was carried out in accordance with the regulations of the Home Office and the ARRIVE guidelines. Efnb2ΔEC mice and recombination protocols were described previously (Wang et al., 2010; Ozawa and James, 2010; Ottone et al., 2014). Tamoxifen-injected Efnb2fl/fl littermates were used as controls. C57Bl6 and CD-1 nude mice for tumourigenicity analysis were obtained from Charles River. For GSC1 tumorigenic studies and B11 treatments of G26 cells, sample size was calculated by power analysis using PASS software based on pilot studies assessing the tumorigenicity of the parental line. For experiments with shEFNB2 human G26 cells sample size was set based on previous studies (Pollard et al., 2009).

Craniotomies and intravital imaging

Craniotomies were performed as previously described (Holtmaat et al., 2009). Briefly 6–8 week old Efnb2 ΔEC or Efnb2ΔEC mice were anaesthetized with ketamine-xylazine intraperitoneal injection (0.083 mg/g ketamine, 0.0078 mg/g xylazine). The animals were then injected with 0.02 ml 4 mg/ml intramuscular dexamethasone to limit an inflammatory response and subcutaneous bupivacaine (1 mg/kg) a local anaesthetic. Once the skull was exposed a few drops of lidocaine (1% solution) were applied on its surface. Before covering the burr hole with a glass coverslip, 2x10^4 cells were injected into the cortex at a depth of 150–300 μm using a picospritzer. Mice were left to recover for 7 days and then imaged using a purpose built microscope equipped with a tunable Coherent Ti:Sapphire laser and PrairieView acquisition software. Mice were anaesthetized with isoflurane and secured to a fixed support under the microscope. The eyes were coated with Lacri-lube (Allergan) to prevent dehydration. Depth of anaesthesia was closely monitored. To visualize blood vessels a 3000MW dextran-Texas Red.
conjugate was injected intravenously prior to imaging. A pulsed laser beam with a wavelength of 910 nm was used to ensure that both GFP-tumour cells and Texas Red showed sufficient signal intensity. Each imaging session lasted for no longer than 60 min and mice were imaged up to four times daily with cells imaged every 20 min. After image acquisition individual frames were aligned and the displacement of single cells measured using ImageJ software.

Orthotopic xenografts and in vivo imaging
5x10⁴ luciferase expressing cells were injected into 6–8 week old CD-1 nude mice into the right putamen (1 mm rostral to bregma, 2 mm lateral and 2.5 mm depth) as previously described (Ozawa and James, 2010). Tumour cells were loaded into the syringe just prior to injection and the needle kept in place for a further 5 min to ensure minimal reflux of the material along the needle tract. Tumour formation, growth and volume were indirectly calculated by sequential images taken with an IVIS Spectrum in vivo imaging system (Perkin Elmers). Following administration of 120 mg/kg D-luciferin (Intrace medical) by intraperitoneal injection, mice were anaesthetized (3% isoflurane) and imaged under continuous exposure to 2% isoflurane. Luminescent measures were performed once a week starting 5 days after cell implantation until day 40. Bioluminescence was detected by the IVIS camera system, integrated, digitised, and displayed. Pseudocolor scale bars were consistent for all images of dorsal views in order to show relative changes at tumour site over time. Tumours were quantified by calculating total flux (photons/s/cm²) using Living Image software (Xenogen, Caliper Life Sciences).

Animals were sacrificed when they showed signs of distress or weight loss. For treatment experiments, PBS or B11 administration was started once the tumours reached a minimum signal intensity of 1x10⁶ photons/s/cm². Mice were then randomised into two groups prior to intravenous injection of 5 doses of either anti ephrinB2-scFv B11 (total dose 20 mg/kg) or PBS control over a period of 9 days. Survival curves were estimated using the Kaplan-Meier method. Significance was calculated using the log-rank Mantel-Cox test. To determine efficiency of delivery, B11 was labeled with Alexa Fluor 680 using the SAIVI rapid antibody kit (Invitrogen) according to the manufacturer’s instructions.

Cell culture, gene delivery and constructs
Primary mouse microvascular endothelial cells were obtained from Caltag Medsystems and subcultured according to the suppliers recommendations (cells were primary cells isolated directly from normal mouse brain, no further authentication performed by the authors, mycoplasma negative as tested by Mycoalert kit, Lonza).

Human GSC lines were described previously (Pollard et al., 2009; Caren et al., 2015). Cells were originally isolated from patient tumours and are maintained in serum free cultures on laminin (Pollard et al., 2009, no further authentication performed by the authors, mycoplasma negative as tested by Mycoalert kit, Lonza). Co-culture experiments with endothelial cells for assessment of p-Eph levels were performed as previously described (Ottone et al., 2014). Briefly, endothelial cells were seeded at confluence on PLL-coated dishes and imNSC or GSC cells seeded on top the following day. The cells were separated by differential trypsinisation. For analysis of clonal efficiency, single cells were sorted into individual wells of a low attachment 96 well plate using a FACS Aria III cell sorter and cultured in neural stem cell media for 7d after which percentage of neurosphere formed was calculated. Transient transfections and viral transductions were performed as previously reported (Ottone et al., 2014). The following plasmids were used: FUGW was used to label all imNSC and GSC cells (Lois et al., 2002). Constitutively active RhoA-V14 and dominant negative RhoA-N19 constructs were a kind gift of Anne Ridley (Ridley and Hall, 1992). Plasmid encoding full-length cDNAs of human EphB1 was purchased from OriGene Technologies, Inc (RC214301) and mouse EphB2 was a kind gift of E. Batlle.

Generation of imNSC and GSC lines
Primary NSCs were isolated directly from the SVZ of postnatal or adult Cdh5(Pac)-CreERT2 or NF1fl/fl transgenic animals and cultured as described previously (Wang et al., 2010; Zhu et al., 2001; Ottone et al., 2014). To generate imNSC and GSC lines the following plasmids were used: pBabe-largeT + pLXSN-hRasV12 (Cremona and Lloyd, 2009), shp53 pLKO.1 (Godar et al., 2008), pCMV-
Cdk4 (van den Heuvel and Harlow, 1993) (addgene 1874). Recombination of the \textit{NF1}^{fl/fl} allele was achieved using Adeno-Cre viruses at an MOI of 100, as reported (Ottone et al., 2014). All experiments were performed on a minimum of two independent batches of GSC1 and 2 generated as described above from independent primary preparations (no further authentication performed by the authors, mycoplasma negative as tested by Mycoalert kit, Lonza). Cells were used within the first 10–15 passages from infection.

**Immunoblotting, immunocytochemistry and immunohistochemistry**

Western blots and immunocytochemistry were performed as reported previously (Ottone et al., 2014). The RhoA activation assay kit (Millipore) was used according to manufacturers instructions. For immunohistochemical analysis of tumours, mice were perfused with 4% PFA and the brain post-fixed in 4% paraformaldehyde for 2 hr, placed in a 30% sucrose solution over night, embedded in OCT and snap frozen. Immunohistochemistry was performed on 30 µm cryostat sections. Tissue sections were stained overnight at 4°C with antibodies diluted in 10% goat serum. Hematoxylin and Eosin (H&E) staining was performed on 3 µm paraffin embedded sections. The following commercial primary antibodies were used: β-Tubulin (Sigma T8328 1:5000), BrdU (Roche 11170376001 1:400), cleaved caspase-3 (Cell Signaling #9664 1:500), EphA4 (abcam ab641 1:1000), EphB1 (abcam ab129103 1:1000), EphB2 (abcam ab76885 1:1000), EphB3 (abcam ab133742), EphB4 (R&D AF446), ephrinA5 (R&D AF3743 1:500), ephrinB2 (R&D AF496 1:250), CD31 (BD 550274 1:400), GAPDH (abcam ab9483 1:1000), GFAP (abcam 54554 1:400), GFP (Invitrogen A-21311 1:400), Nestin (Millipore MAB353 1:400), pan-pEph (abcam ab61791 1:500), pFAK (Cell signalling #3283 1:500), p-Src (Cell Signaling 2101s 1:1000), O4 (R&D MAB1326 1:400), Sox2 (Cell signalling 3728s 1:250), SSEA1 (BD Pharmingen 560079 1:400), Tuj1 (Covance MMS-435P 1:500) and NG2 (Chemicon, 1:250).

**Migration assays and quantifications**

For migration assays, endothelial cells and imNSC/GSCs were plated at a density of 2x10⁴ cells into adjacent compartments of cell-culture silicon inserts (Ibidi) separated by a 500 µm gap. Alternatively one well of the insert was coated over night with 4 µg/ml of recombinant ephrin-B2-Fc fusion proteins or Fc controls (R&D) clustered at a molar ratio of 1:2 with fluorescently labelled anti-Fc antibody for 1.5 hr at room temperature. After removal of the insert the cells were cultured in medium supplemented with 1 % Matrigel (Invitrogen) and live cell imaging was performed in a heated and CO₂ controlled chamber for 60 hr. For stimulation of imNSCs, recombinant ephrinB2-Fc ligands were preclustered with anti-human Fc antibodies as above and added to imNSCs for 24 hr before removal of the insert at a final concentration of 10 µg/ml. Analysis of ephrinb2-stimulated imNSC migration was terminated at 48 hr to achieve maximal stimulation of forward signaling throughout the experiments. Migration was quantified by tracing the boundary between GFP positive NSCs and live cell imaging was performed in a heated and CO₂ controlled chamber for 60 hr. For analysis of anchorage independent proliferation, cells were seeded for up to 72 hr in SVZ culture medium containing 1.8% dissolved Methylcellulose (Sigma) to form a semisolid hydrogel as previously described. To retrieve cells, the suspension was diluted fivefold with DMEM and centrifuged at 500 g for 10 min. The pellet was then washed twice with ice-cold PBS. For analysis of binucleated cells, all cells were incubated with EdU (Life technology) for 4 hr and placed on PLL-coated cover-slips 15 min before fixation and staining. As the metaphase in mammalian cells typically lasts less than an hour this experimental set up enabled us to distinguish between cycling (EdU positive) and
cytokinesis arrested (EdU negative) cells. To assess colony formation in soft agar 5x10^3 cells/6 well were seeded into 0.5% agar on top of a bottom layer of 1% agar (Sigma). Total number of colonies formed was counted after 10 days.

**Flow cytometry**

Cell pellets retrieved from methylcellulose were resuspended, fixed in 70% EtOH for 4 hr and stained with propidium iodide (Sigma). A minimum of 10,000 cells were counted for each condition using a BD LSR II. The PI profile was then analysed using the cell-cycle module of FlowJo X. Cells from tumours were isolated as follows: The GFP positive tumour tissue was dissected using a fluorescent dissection microscope and digested using papain according to the manufactures recommendation (Worthington Biochemical Corporation). Cells were stained with PI as above prior to FACS analysis. All FACS stainings were performed on at least 3 independent cultures/tumours.

**RNA sequencing and Bioinformatic analysis**

GFP-labeled GSC1 and GSC2 tumours were dissected and digested as above. RNA was isolated by using RNAeasy Plus Mini Kit (Qiagen) and RNA sequencing libraries were constructed using the NEBNext Ultra RNA Library Prep Kit for Illumina (NEB). RNA-seq data from adult brain of normal mice was obtained from GEO [1] (accession numbers GSM1055111, GSM1055112 and GSM1055113). Sequencing reads from tumour samples were aligned with the Tophat splice junction mapper (Kim et al., 2013), version 2.0.11 against GRCm37/ mm9 and transcript annotations. All parameters were set to default except inner distance between mate pairs (r = 100) and library type (fr-firststrand). The normal brain data was also aligned using Tophat with default parameter values except for distance between mature pairs (r = 200). The DESeq2 Bioconductor package (version 1.4.5) (Love et al., 2014) was used to perform differential gene expression analysis on read counts obtained with HTSeq-count (version 0.5.3p9) (Anders et al., 2015), and p-values were adjusted for multiple testing with the Benjamini-Hochberg procedure (Love et al., 2014). Genes with adjusted p-value <= 0.05 were deemed to be differentially expressed.

Sequencing reads for NS and GSC cell lines were aligned to mouse genome build GRCm38/mm10 with STAR 2.5.2a (Dobin et al., 2013) using the two-pass method for novel splice detection (Engstrom et al., 2012). Read alignment was guided by gene annotation from Ensembl release 84 (Cunningham et al., 2015) with optimal splice junction donor/acceptor overlap settings. Transcripts were quantified with HTSeq-count (Anders et al., 2015) based on feature coordinates from Ensembl 84. Gene set enrichment analysis was carried out with the GAGE Bioconductor package (version 2.18.0) (Luo et al., 2009). Gene sets used corresponded to either biological process (BP) terms from the Gene Ontology, or derived from CNS cell type RNA-seq data described by the Barres group (http://web.stanford.edu/group/barres_lab/brainseq2/brainseq2.html) using mean log2 FPKM values for astrocyte, neuron, OPC and oligodendrocyte classes. The top 500 genes for each class were identified by the ratio of individual class average expression to the maximum non-class average expression values. Log2 fold-change values computed by DESeq2 (version 1.8.2) were used as the input for GSA analysis. Data visualised in Figure 1—figure supplement 1a is derived from log2 FPKM data for the left panel and variance stabilised read counts using the rlog function in DESeq2.

**GBM subtype analysis**

Subtype analysis of human GSC lines was completed using methods and microarray data described by Caren et al. (Caren et al., 2015). Sample log2-transformed expression values for the signature centroid genes were produced by taking the mean expression across sample replicates. Centroid genes that could not be assigned to annotated genes were also omitted from further analysis. To ensure the accuracy of subtype expression estimates only subtype genes with high variance across the GNS dataset were carried forward. Subtype scores per sample were computed from mean Z-score transformed levels of overexpressed centroid genes for each subtype. Samples were then classified as belonging to the subtype associated with the highest mean Z-score or mixed subtype when presenting a similarly high expression of another subtype’s mean Z-score.

Gene set enrichment analysis (GSEA) (Subramanian et al., 2005) was applied to mouse tumor samples to test for Verhaak et al. subtype enrichment using the pre-ranked GSEA tool. DESeq2 computed T statistics from each tumor to control mouse sample differential expression results were used.
to rank the gene lists. Pre-defined genesets pertaining to Verhaak et al. (Verhaak et al., 2010) were obtained from the Molecular Signatures Database. TCGA data analysis was performed as described previously (Binda et al., 2012). Survival and expression analysis was performed on the publicly available TCGA data and relative mRNA expression obtained from the TCGA data portal (http://cancer-genome.nih.gov/dataportal/data/about). The analysis was performed using samr R package. A two class unpaired model was used with the following parameters: delta 0.01, number of permutation 100.

Quantitative RT-PCR and siRNA/shRNA knockdown
Quantitative RT-PCR analysis and siRNA transfections were performed as previously described (Ottone et al., 2014). See Supplementary file 3 for primer details. For knockdown of EFNB2 in human G26 cells, lentiviral shRNA constructs were used (Invitrogen). Control cells were infected with control vectors. Both cell types were selected with puromycin and knock-down efficiency validated by qPCR and immunoblotting prior to tumourigenic studies.

Neuropathological assessment of Ephrin-B2 expression in human GBMs
EphrinB2 expression pattern was investigated in brain tissue samples of eight patients operated of supratentorial hemispheric glioblastoma at Imperial College Healthcare Trust between February and June 2015. None of patients had radiotherapy or chemotherapy prior to surgery or any relevant comorbidities. Patients consent was given for all samples. The project received ethical approval from the Imperial College Healthcare Tissue Bank committee on behalf of MREC (ICHBT HTA licence: 12275, REC Wales approval: 12/WA/0196). All 8 tumours were wild type for IDH1 and 2 mutations and ATRX status with the exception of case 6, which was a secondary glioblastoma bearing the common G395A amino acid substitution in IDH1 and showing loss of ATRX expression.

Two additional GBM samples were the original lesions from which G19 and G26 cells have been isolated (Pollard et al., 2009). Preoperative structural MRI was available in all patients; the T2-weighted FLAIR sequences were reviewed to assess the extent of invasion into normal tissue. All patients underwent maximally safe surgical debulking and the isolated tissues were extensively sampled in order to represent the bulk of the tumour along with its infiltrative component in the surrounding grey matter. Specifically, areas representative of the invasive front of the tumour were chosen from each case. The tissues were fixed in 4% buffered formalin for 24 hr and then processed using standard method to produce paraffin sections. Sequential three-micron sections cut from each selected block and were used for immunoperoxidase immunohistochemistry. Immunostainings were performed following a standard protocol. Briefly, the sections were dewaxed in xylene and rehydrated in decreasing alcohols to distilled water. Antigen retrieval was performed incubating the sections for 20 min in steam-heated sodium citrate buffer (10 mM Sodium Citrate, 0.05% Tween 20, pH 6) at 90°C or 1 mM EDTA at pH 8. Following antigen unmasking, endogenous peroxidase was quenched with 1% hydrogen peroxide in PBS at -20°C for 15 min. After rinsing in PBS, the sections were incubated overnight at room temperature with the following primary antibodies: anti-EphrinB2 (clone EFR-163M hybridoma supernatant 1:4, CNIO) (Abengozar et al., 2012), anti-ALDH1 (monoclonal, BD Bioscience, dilution 1:500), anti-ALDH1 (mouse, clone 44, BD biosciences). The ephrin-B2 antibody gave robust membrane and cytoplasmic staining and labelled the endothelium, confirming specificity. Nuclear labelling was however detected in some tumour cases, a feature not present in control samples. Only membranous and cytoplasmic staining was scored as positive in the analysis of the tumours. The SuperSensitive IHC detection system from BioGenex (Fremon, CA, USA) was used to visualise antibody binding following the manufacturer’s instructions and counterstained with Mayer’s Haemalum.

Statistics
Statistical analysis was performed using GraphPad Prism statistical analysis software. The precise tests are stated in the figure legends. Shapiro-Wilk test was used to confirm normal distribution of the data. All experiments for which quantifications were performed were carried out a minimum of three times as indicated in the figure legends.
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Ethics
Animal experimentation: All animal work was carried out in accordance with the regulations of the Home Office and the ARRIVE guidelines and performed under the authority of Project License 70-7292. Patients consent was given for all human GBM samples. The project received ethical approval from the Imperial College Healthcare Tissue Bank committee on behalf of MREC (ICHTB HTA licence: 12275, REC Wales approval: 12/WA/0196).

Additional files

Supplementary files
- Supplementary file 1. List of significantly enriched GO terms in GSC vs NSC listing FDR q-values

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• Supplementary file 2. List of significantly enriched GO terms in NSC vs GSC listing FDR q-values
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• Supplementary file 3. Mouse Primers used for qRT-PCR
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Major datasets
The following previously published datasets were used:

| Author(s)            | Year | Dataset title                                                                 | Dataset URL                                                                 | Database, license, and accessibility information |
|----------------------|------|-------------------------------------------------------------------------------|------------------------------------------------------------------------------|--------------------------------------------------|
| Paul Bertone         | 2015 | Expression profiling of normal and glioblastoma-derived neural stem cells     | http://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-3864/                   | Publicly available at EBI ArrayExpress            |
| Alexey Fushan        | 2015 | Gene Expression Defines Natural Changes in Mammalian Lifespan                  | http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1055111                 | Publicly available at NCBI Gene Expression Omnibus (accession no: GSM1055111) |
| Alexey Fushan        | 2015 | Gene Expression Defines Natural Changes in Mammalian Lifespan                  | http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSM1055112                 | Publicly available at NCBI Gene Expression Omnibus (accession no: GSM1055112) |
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References
Abéngozar MA, de Frutos S, Ferreiro S, Soriano J, Perez-Martinez M, Olmeda D, Marenchino M, Cañamero M, Ortega S, Megias D, Rodríguez A, Martínez-Torre cuadra J. 2012. Blocking ephrinB2 with highly specific antibodies inhibits angiogenesis, lymphangiogenesis, and tumor growth. *Blood* **119**:4565–4576. doi: 10.1182/blood-2011-09-380006

Anders S, Pyl PT, Huber W. 2015. HTSeq—a Python framework to work with high-throughput sequencing data. *Bioinformatics* **31**:166–169. doi: 10.1093/bioinformatics/btu638

Astin JW, Batson J, Kadir S, Charlet J, Persad RA, Gillatt D, Oxley JD, Nobes CD. 2010. Competition amongst Eph receptors regulates contact inhibition of locomotion and invasiveness in prostate cancer cells. *Nature Cell Biology* **12**:1194–1204. doi: 10.1038/ncc2122

Baker GJ, Yadav VN, Motsch S, Koschmann C, Calinescu AA, Mineharu Y, Camel o-Piragua SI, Orringer D, Bannykh S, Nichols WS, deCarvalho AC, Mik kelsen T, Castro MG, Lowenstein PR. 2014. Mechanisms of glioma formation: iterative perivascular glioma growth and invasion leads to tumor progression, VEGF-independent vascularization, and resistance to antiangiogenic therapy. *Neoplasia* **16**:543–561. doi: 10.1016/j.neo.2014.06.003

Bao S, Wu Q, McLendon RE, Hao Y, Shi Q, Hjelmeland AB, Dewhirst MW, Bigner DD, Rich JN. 2006. Glioma stem cells promote radioreistance by preferential activation of the DNA damage response. *Nature* **444**:756–760. doi: 10.1038/nature05236

Bhat KP, Balasubramaniyan V, Vaillant B, Ezhilarasan R, Hummelink K, Hollingsworth F, Wani K, Heathcock L, James JD, Goodman LD, Conroy S, Long L, Lelic N, Wang S, Gumim J, Raj D, Kodama Y, Raghunathan A, Olar A, Joshi K, et al. 2013. Mesenchymal differentiation mediated by NF-kB promotes radiation resistance in glioblastoma. *Cancer Cell* **24**:331–346. doi: 10.1016/j.ccr.2013.08.001

Binda E, Visioli A, Giani F, Lamorte G, Copetti M, Pitter KL, Huse JT, Caja llo L, Zanetti N, DiMeco F, De Filippis L, Mangiola A, Maira G, Anile C, De Bonis P, Reynolds BA, Pasquale EB, Vescovi AL. 2012. The EphA2 receptor drives self-renewal and tumorigenicity in stem-like-tumor-propagating cells from human glioblastomas. *Cancer Cell* **22**:765–780. doi: 10.1016/j.ccr.2012.11.005

Blouw B, Song H, Tihan T, Bosje T, Ferrara N, Gerber HP, Johnson RS, Bergers G. 2003. The hypoxic response of tumors is dependent on their microenvironment. *Cancer Cell* **4**:133–146. doi: 10.1016/S1535-6108(03)00194-6

Bochenek ML, Dickinson S, Astin JW, Adams RH, Nobes CD. 2010. Ephrin-B2 regulates endothelial cell morphology and motility independently of Eph-receptor binding. *Journal of Cell Science* **123**:1235–1246. doi: 10.1242/jcs.061903

Calabrese C, Poppleton H, Kocak M, Hogg TL, Fuller C, Hamner B, Oh EY, Gaber MW, Finklestein D, Allen M, Frank A, Bayazitov IT, Zakharenko SS, Gajjar A, Davidoff A, Gilbertson RJ. 2007. A perivascular niche for brain tumor stem cells. *Cancer Cell* **11**:89–92. doi: 10.1016/j.ccr.2006.11.020
Carcen H, Stricker SH, Bulstrode H, Gagrca S, Johnstone E, Bartlett TE, Feber A, Wilson G, Teschendorff AE, Bertone P, Beck S, Pollard SM. 2015. Glioblastoma Stem Cells Respond to Differentiation Cues but Fail to Undergo Commitment and Terminal Cell-Cycle Arrest. Stem Cell Reports 5. doi: 10.1016/j.stemcr.2015.09.014

Carro MS, Lim WK, Alvarez MJ, Boilo RJ, Zhao X, Snyder EY, Sulman EP, Anne SL, Doetsch F, Colman H, Lasorella A, Aldape K, Califano A, Iavarone A. 2010. The transcriptional network for mesenchymal transformation of brain tumours. Nature 463:318–325. doi: 10.1038/nature08712

Cayuso J, Xu Q, Wilkinson DG. 2015. Mechanisms of boundary formation by Eph receptor and ephrin signaling. Developmental Biology 401. doi: 10.1016/j.ydbio.2014.11.013

Charles N, Holland EC. 2010. The perivascular niche microenvironment in brain tumour progression. Cell Cycle 9:3012–3021. doi: 10.14616/j.cc.15.12335

Chen J, Li Y, Yu TS, McKay RM, Burns DK, Kernie SG, Parada LF. 2012. A restricted cell population propagates glioblastoma growth after chemotherapy. Nature 488:522–526. doi: 10.1038/nature11287

Chen J, McKay RM, Parada LF. 2012. Malignant glioma: lessons from genomics, mouse models, and stem cells. Cell 149:36–47. doi: 10.1016/j.cell.2012.03.009

Cheng I, Wu Q, Guruyanova OA, Huang Z, Huang Q, Rich JN, Bao S. 2011. Elevated invasive potential of glioblastoma stem cells. Biochemical and Biophysical Research Communications 406:646–648. doi: 10.1016/j.bbrc.2011.02.123

Cortina N, Palomo-Ponce S, Iglesias M, Fernandez-Masip JL, Vivancos A, Whissell G, Humã M, Peiró N, Gallego L, Jonkheer S, Davy A, Lloreta J, Sancho E, Battie E. 2007. EphB-ephrin-B interactions suppress colorectal cancer progression by compartmentalizing tumor cells. Nature Genetics 39:1376–1383. doi: 10.1038/ng.2007.11

Cremona CA, Lloyd AC. 2009. Loss of anchorage in checkpoint-deficient cells increases genomic instability and promotes oncogenic transformation. Journal of Cell Science 122:3272–3281. doi: 10.1242/jcs.047126

Cuddapah VA, Robel S, Watkins S, Sontheimer H. 2014. A neurocentric perspective on glioma invasion. Nature Reviews. Neuroscience 15:455–465. doi: 10.1038/nn3765

Cunningham F, Amode MR, Barrett D, Beal K, Bills K, Brent S, Carvalho-Silva D, Clapham P, Coates G, Fitzgerald S, Gil L, GIRON CG, Gordon L, Hourlier T, Hunt SE, Janacek SH, Johnson N, Juettemann T, KAHARI AK, Keenan S, et al. 2015. Ensembl 2015. Nature 43:662–669. doi: 10.1039/nar/2015

Daar IO. 2012. Non-SH2/PDZ reverse signaling by ephrins. Seminars in Cell & Developmental Biology 23:65–74. doi: 10.1016/j.semcdb.2011.10.012

Day BW, Stringer BW, Al-Ejeh F, Ting MJ, Wilson J, Ensbey KS, Jamieson PR, Bruce ZC, Lim YC, Offenhauser C, Charmsaz S, Cooper LT, Ellacott JK, Harding A, Leveque L, Inglis P, Allan S, Walker DG, Lackmann M, Osborne et al. 2013. EphA3 maintains tumorigenicity and is a therapeutic target in glioblastoma multiforme. Cancer Cell 23:238–248. doi: 10.1016/j.ccr.2013.01.007

Day BW, Stringer BW, Boyd AW. 2014. Eph receptors as therapeutic targets in glioblastoma. British Journal of Cancer 111:1265–1261. doi: 10.1038/bjc.2014.73

Desgrosellier JS, Barnes LA, Shields DJ, Huang M, Lau SK, Prévost N, Tarin D, Shattil SJ, Cheresh DA. 2009. An integrin alpha(3)(beta)(3)-c-Src oncogenic unit promotes anchorage-independence and tumor progression. Nature Medicine 15:1163–1169. doi: 10.1038/nm.2009

Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M, Gingeras TR. 2013. STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29:15–21. doi: 10.1093/bioinformatics/bts635

Donnem T, Shin SI. 1974. Cellular tumorigenicity in nude mice: correlation with cell growth in semi-solid medium. Cell 3:355–359. doi: 10.1002/667-4940050-6

Farin A, Suzuki SO, Weiker M, Goldman JE, Bruce JN, Canoll P. 2006. Transplanted glioma cells migrate and proliferate on host brain vasculature: a dynamic analysis. Glia 53:799–808. doi: 10.1002/glia.20334

Fog SS, Turner CJ, Adams S, Compagni A, Aubyn D, Kogata N, Lindblom P, Shani M, Zicha D, Adams RH. 2006. Ephrin-B2 controls cell motility and adhesion during blood-vessel-wall assembly. Cell 124:161–173. doi: 10.1016/j.cell.2005.10.034

Freedman VH, Shin SI. 1974. Cellular tumorigenicity in nude mice: correlation with cell growth in semi-solid medium. Cell 3:355–359. doi: 10.1002/667-4940050-6

Gale NW, Baluk P, Pan L, Kwan M, Holash J, DeChiara TM, McDonald DM, Yancopoulos GD. 2001. Ephrin-B2 selectively marks arterial vessels and neovascularization sites in the adult, with expression in both endothelial and smooth-muscle cells. Developmental Biology 230:151–160. doi: 10.1006/dbio.2000.0112

Godar S, Ince TA, Bell GW, Feldser D, Donaher JL, Bergh J, Liu A, Miu K, Watnick RS, Reinhardt F, McAllister SS, Barsky T, Weinberg RA. 2008. Growth-inhibitory and tumor- suppressive functions of p53 depend on its repression of CD44 expression. Cell 134:62–73. doi: 10.1016/j.cell.2008.06.006

Gritsenko PM, Ilina O, Friedl P. 2012. Intestinal guidance of cancer invasion. The Journal of Pathology 226:185–199. doi: 10.1002/path.3031

Gürsel DB, Shin BJ, Burkhardt JK, Kesavabhotla K, Schlaff CD, Boockvar JA. 2011. Glioblastoma stem-like cells—biology and therapeutic implications. Cancers 3:2655–2666. doi: 10.3390/cancers3022655

Hahn WC, Counter CM, Lundberg AS, Beijersbergen RL, Brooks MW, Weinberg RA. 1999. Creation of human tumour cells with defined genetic elements. Nature 400:464–468. doi: 10.1038/22780
van den Heuvel, S. & Harlow, E. 1993. Distinct roles for cyclin-dependent kinases in cell cycle control. Science 262: 2050–2054. doi: 10.1126/science.8266103

Holtmaat, A., Bonhoeffer, T., Chow, D.K., Chuckowree, J., De Paola, V., Hofer, S.B., Hübener, M., Keck, T., Knott, G., Lee, W.C., Mostany, R., Mrsic-Flogel, T.D., Nedivi, E., Portera-Cailliau, C., Svoboda, K., Trachtenberg, J.T., Wilbrecht, L. 2009. Long-term, high-resolution imaging in the mouse neocortex through a chronic cranial window. Nature Protocols 4:1128–1144. doi: 10.1038/nprot.2009.89

Huszthy, P.C., Daphu, I., Niclou, S.P., Stieber, D., Nirgo, J.M., Sakariassen, P.D., Miletic, H., Thorsen, F., Bjerkkvig, R. 2012. In vivo models of primary brain tumors: pitfalls and perspectives. Neuro-Oncology 14: 979–993. doi: 10.1093/neuonc/noq135

Jordan, S.N., Cannan, J.C. 2012. Rho GTPases in animal cell cytokinesis: an occupation by the one percent. Cytoskeleton 69: 919–930. doi: 10.1002/cm.21071

Kim, D., Perteau, G., Trapnell, C., Pimentel, H., Kelley, R., Salzberg, S.L. 2013. TopHat2: accurate alignment of transcriptomes in the presence of insertions, deletions and gene fusions. Genome Biology 14: R36. doi: 10.1186/gb-2013-14-4-r36

Kleiheues, P., Cavenee, W.K. 2000. Pathology and Genetics of Tumours of the Nervous System. Lyon, France: IARC Press.

Kokovay, E., Goderie, S., Wang, Y., Lotz, S., Lin, G., Sun, Y., Roysam, B., Shen, Q., Temple, S. 2010. Adult SVZ lineage cells home to and leave the vascular niche via differential responses to SDF1/CXCR4 signaling. Cell Stem Cell 7: 163–173. doi: 10.1016/j.stem.2010.05.019

Kullander, K., Klein, R. 2002. Mechanisms and functions of Eph and ephrin signalling. Nature Reviews. Molecular Cell Biology 3: 475–486. doi: 10.1038/nrm856

Lamszus, K., Kunkel, P., Westphal, M. 2003. Invasion as limitation to anti-angiogenic glioma therapy. Acta Neurochirurgica. Supplement 88: 169–177. doi: 10.1007/978-3-7091-6090-9_23

Lois, C., Hong, E.J., Pease, S., Brown, E.J., Baltimore, D. 2002. Germline transmission and tissue-specific expression of transgenes delivered by lentiviral vectors. Science 295: 868–872. doi: 10.1126/science.1067081

Love, M.I., Huber, W., Anders, S. 2014. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology 15. doi: 10.1186/s13059-014-0550-8

Luo, W., Friedman, M.S., Shedden, K., Hankenson, K.D., Woolf, P.J. 2009. GAGE: generally applicable gene set enrichment for pathway analysis. BMC Bioinformatics 10: 161. doi: 10.1186/1471-2105-10-161

Mao, P., Joshi, K., Li, J., Kim, S., Li, P., Santana-Santos, L., Luthra, S., Chandran, U., Benos, P.V., Smith, L., Wang, M., Hu, B., Cheng, S.Y., Sobol, R.W., Nakano, I. 2013. Mesenchymal glioma stem cells are maintained by activated glycolytic metabolism involving aldehyde dehydrogenase 1A3. Proceedings of the National Academy of Sciences of the United States of America 110: 8644–8649. doi: 10.1073/pnas.1221478110

Nagano, N., Sasaki, H., Aoyagi, M., Hirakawa, K. 1993. Invasion of experimental rat brain tumor: early morphological changes following microinjection of C6 glioma cells. Acta Neuropathologica 86: 117–125. doi: 10.1007/BF00334878

Nakada, M., Anderson, E.M., Demuth, T., Nakada, S., Reavie, L.B., Drake, K.L., Hoelzinger, D.B., Berens, M.E. 2010. The phosphorylation of ephrin-B2 ligand promotes glioma cell migration and invasion. International Journal of Cancer 126: 1155–1165. doi: 10.1002/ijc.24849

Nakada, M., Hayashi, Y., Hamada, J. 2011. Role of Eph/ephrin tyrosine kinase in malignant glioma. Neuro-Oncology 13: 1163–1170. doi: 10.1093/neuonc/nor102

Cancer Genome Atlas Research Network. 2008. Comprehensive genomic characterization defines human glioblastoma genes and core pathways. Nature 455: 1061–1068. doi: 10.1038/nature07385

Ottone, C., Krusche, B., Whitby, A., Clements, M., Quadrato, G., Pitulescu, M.E., Adams, R.H., Parrinello, S. 2014. Direct cell-cell contact with the vascular niche maintains quiescent neural stem cells. Nature Cell Biology 16: 1045–1056. doi: 10.1038/ncb3045

Ozawa, T., James CD. 2010. Establishing Intracranial Brain Tumor Xenografts with Subsequent Analysis of Tumor Growth and Response to Therapy Using Bioluminescence Imaging. Journal of visualized experiments: JoVe . doi: 10.3791/1986

Ozawa, T., Riester, M., Cheng, Y.K., Huse, J.T., Squatrito, M., Helmy, K., Charles, N., Michor, F., Holland, E.C. 2014. Most human non-GCIMP glioblastoma subtypes evolve from a common proneural-like precursor glioma. Acta Neuropathologica 128: 1128–1144. doi: 10.1007/s00401-014-1511-7

Pasquale, E.B. 2010. Eph receptors and ephrins in cancer: bidirectional signalling and beyond. Nature Reviews. Cancer 10: 165–180. doi: 10.1038/nrc2806

Pfaff, D., Heroult, M., Riedel, M., Reiss, Y., Kirmse, R., Ludwig, T., Korf, T., Hecker, M., Augustin, H.G. 2008. Involvement of endothelial ephrin-B2 in adhesion and transmigration of EphB-receptor-expressing monocytes. Journal of Cell Science 121: 3842–3850. doi: 10.1242/jcs.030627

Pollard, S.M., Yoshikawa, K., Clarke, I.D., Danovi, D., Stricker, S., Russell, R., Bayani, J., Head, R., Lee, M., Bernstein, M., Squire, J.A., Smith, A., Dirks, P.B. 2009. Glioma stem cell lines expanded in adherent culture have tumor-specific phenotypes and are suitable for chemical and genetic screens. Cell Stem Cell 4: 568–580. doi: 10.1016/j.stem.2009.03.014

Qiu, R.G., Chen, J., McCormick, F., Symons, M. 1995. A role for Rho in Ras transformation. Proceedings of the National Academy of Sciences of the United States of America 92: 11781–11785. doi: 10.1073/pnas.92.25.11781

Rasper, M., Schäfer, A., Piontek, G., Teufel, J., Brockhoff, G., Ringel, F., Heindl, S., Zimmer, C., Schlegel, J. 2010. Aldehyde dehydrogenase 1 positive glioblastoma cells show brain tumor stem cell capacity. Neuro-Oncology 12: 1024–1033. doi: 10.1093/neuonc/noq070
Ridley AJ, Hall A. 1992. The small GTP-binding protein rho regulates the assembly of focal adhesions and actin stress fibers in response to growth factors. *Cell* **70**:389–399. doi: 10.1016/0092-8674(92)90163-7

Sadahiro H, Yoshikawa K, Ideguchi M, Kajiwara K, Ishii A, Ikeda E, Owada Y, Yasumoto Y, Suzuki M. 2014. Pathological features of highly invasive glioma stem cells in a mouse xenograft model. *Brain Tumor Pathology* **31**:77–84. doi: 10.1007/s10143-013-0149-x

Sawamiphak S, Seidel S, Essmann CL, Wilkinson GA, Pitulescu ME, Acker T, Acker-Palmer A. 2010. Ephrin-B2 regulates VEGFR2 function in developmental and tumour angiogenesis. *Nature* **465**:487–491. doi: 10.1038/nature08995

Scherer HJ. 1938. Structural Development in gliomas. *American Journal of Cancer* **34**:333–351.

Sonoda Y, Ozawa T, Hirose Y, Aldape KD, McMahon M, Berger MS, Pieper RO. 2001. Formation of intracranial tumors by genetically modified human astrocytes defines four pathways critical in the development of human anaplastic astrocytoma. *Cancer Research* **61**:4956–4960.

Montana V, Sontheimer H. 2011. Bradykinin promotes the chemotactic invasion of primary brain tumors. *The Journal of Neuroscience* **31**:4858–4867. doi: 10.1523/JNEUROSCI.3825-10.2011

Subramanian A, Tamayo P, Mootha VK, Mukherjee S, Ebert BL, Gillette MA, Paulovich A, Pomeroy SL, Golub TR, Lander ES, Mesirov JP. 2005. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. *Proceedings of the National Academy of Sciences of the United States of America* **102**:15545–15550. doi: 10.1073/pnas.0506580102

Tanaka M, Kamo T, Ota S, Sugimura H. 2003. Association of Dishevelled with Eph tyrosine kinase receptor and ephrin mediates cell repulsion. *The EMBO Journal* **22**:847–858. doi: 10.1093/emboj/cdg088

Thulberg M, Gad A, Le Guyader S, Strömblad S. 2007. Oncogenic H-Ras V12 promotes anchorage-independent cytokinesis in human fibroblasts. *Proceedings of the National Academy of Sciences of the United States of America* **104**:20338–20343. doi: 10.1073/pnas.0706609105

Tu Y, He S, Fu J, Li G, Xu R, Lu H, Deng J. 2012. Expression of EphrinB2 and EphB4 in glioma tissues correlated to the progression of glioma and the prognosis of glioblastoma patients. *Clinical & Translational Oncology* **14**:214–220. doi: 10.1007/s12094-012-0786-2

Vehlow A, Cordes N. 2013. Invasion as target for therapy of glioblastoma multiforme. *Biochimica Et Biophysica Acta* **1836**:236–244. doi: 10.1016/j.bbacli.2013.07.001

Venere M, Fine HA, Dirks PB, Rich JN. 2011. Cancer stem cells in gliomas: identifying and understanding the apex cell in cancer’s hierarchy. *Glia* **59**:1148–1154. doi: 10.1002/glia.21185

Verhaak RG, Hoadley KA, Purdom E, Wang V, Qi Y, Wilkerson MD, Miller CR, Ding L, Golub T, Mesirov JP, Alexe G, Lawrence M, O’Kelly M, Tamayo P, Weir BA, Gabriel S, Winckler W, Gupta S, Jakkula L, Feiler HS, et al. 2010. Integrated genomic analysis identifies clinically relevant subtypes of glioblastoma characterized by abnormalities in PDGFRA, IDH1, EGFR, and NF1. *Cancer Cell* **17**:98–110. doi: 10.1016/j.ccr.2009.12.020

Wang Y, Nakayama M, Pitulescu ME, Schmidt TS, Bochenek ML, Sakakibara A, Adams S, Davy A, Deutsch U, Luthi U, Barberis A, Benjamin LE, Mäkinen T, Nobes CD, Adams RH. 2010. Ephrin-B2 controls VEGF-induced angiogenesis and lymphangiogenesis. *Nature* **465**:483–486. doi: 10.1038/nature09002

Wang Y, Kim E, Wang X, Novitch BG, Yoshikawa K, Chang LS, Zhu Y. 2012. ERK inhibition rescues defects in fate specification of NF1-deficient neural progenitors and brain abnormalities. *Cell* **150**:816–830. doi: 10.1016/j.cell.2012.06.034

Watkins S, Robel S, Kimbrough IF, Robert SM, Ellis-Davies G, Sontheimer H. 2014. Disruption of astrocyte-vascular coupling and the blood-brain barrier by invading glioma cells. *Nature Communications* **5**: doi: 10.1038/ncomms5196

Weathers SP, Gilbert MR. 2014. Advances in treating glioblastoma. *F1000prime Reports* **6**: doi: 10.12703/P6-46

Winkler F, Kienast Y, Fuhrmann M, Von Baumgarten L, Burgold S, Mittregerger G, Kretzschmar H, Herrns J. 2009. Imaging glioma cell invasion in vivo reveals mechanisms of dissemination and peritumoral angiogenesis. *Glia* **57**:1306–1315. doi: 10.1002/glia.20850

Zagzag D, Esencay M, Mendez O, Yee H, Smirnova I, Huang Y, Chiriboga L, Lukyanov E, Liu M, Newcomb EW. 2008. Hypoxia- and vascular endothelial growth factor-induced stromal cell-derived factor-1alpha/CXCR4 expression in glioblastomas: one plausible explanation of Scherer’s structures. *The American Journal of Pathology* **173**:545–560. doi: 10.2353/ajpath.2008.071197

Zhang Y, Chen K, Sloan SA, Bennett ML, Scholze AR, O’Keeffe S, Phatnani HP, Guarnieri P, Caneda C, Ruderisch N, Deng S, Liddelow SA, Zhang C, Daneman R, Maniatis T, Barres BA, Wu JQ. 2014. An RNA-sequencing transcriptome and splicing database of glia, neurons, and vascular cells of the cerebral cortex. *The Journal of Neuroscience* **34**:11929–11947. doi: 10.1523/JNEUROSCI.1860-14.2014

Zhu Y, Romero MI, Ghosh P, Ye Z, Charnay P, Rushing EJ, Marth JD, Parada LF. 2001. Ablation of NF1 function in neurons induces abnormal development of cerebral cortex and reactive gliosis in the brain. *Genes & Development* **15**:859–876. doi: 10.1101/gad.862101