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EPHB2 carried on small extracellular vesicles induces tumor angiogenesis via activation of ephrin reverse signaling

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Abstract

Angiogenesis is a key process that allows nutrient uptake and cellular trafficking and is co-opted in cancer to enable tumor growth and metastasis. Recently, extracellular vesicles (EVs) have been shown to promote angiogenesis; however, it is unclear what unique features EVs contribute to the process. Here, we studied the role of EVs derived from head and neck squamous cell carcinoma (HNSCC) in driving tumor angiogenesis. Small EVs (SEVs), in the size range of exosomes (50-150 nm), induced angiogenesis both \textit{in vitro} and \textit{in vivo}. Proteomic analysis of HNSCC SEVs revealed the cell-cell signaling receptor EPHB2 as a promising candidate cargo to promote angiogenesis. Analysis of patient data further identified EPHB2 overexpression in HNSCC tumors to be associated with poor patient prognosis and tumor angiogenesis, especially in the context of overexpression of the exosome secretion regulator cortactin. Functional experiments revealed that EPHB2 expression in SEVs regulates angiogenesis both \textit{in vitro} and \textit{in vivo} and that EPHB2 carried by SEVs stimulates ephrin-B reverse signaling, inducing STAT3 phosphorylation. A STAT3 inhibitor greatly reduced SEV-induced angiogenesis. These data suggest a novel model in which EVs uniquely promote angiogenesis by transporting Eph transmembrane receptors to non-adjacent endothelial cells to induce ephrin reverse signaling.
Introduction

Head and neck squamous cell carcinoma (HNSCC) is an invasive cancer that originates in the oral cavity, larynx, and pharynx, in association with alcohol and tobacco use or exposure to human papillomavirus (1-3). While much of the morbidity of HNSCC is caused by local invasion into nerve and vascular tissues of the neck, mortality is driven primarily by metastasis to regional lymph nodes and distant organs (4-7). A key event that facilitates this metastasis is tumor-associated lymph and blood vessel formation (Fig. 1A).

Extracellular vesicles (EVs), including exosomes and other small EVs (SEVs) and larger EVs (LEVs) such as microvesicles, are secreted from cells and mediate cell-cell communication via protein, lipid and nucleic acid cargoes (8). EVs are key mediators of cellular functions such as survival, proliferation, motility, and apoptosis. Recently, many reports have shown that tumor-derived EVs play a large role in tumor progression (9). Many of these functions are due to paracrine and distant signaling to non-cancer cells, including induction of cancer-associated fibroblasts, regulation of tumor immunity, and premetastatic niche formation.

Among the paracrine activities, a key reported function of tumor EVs is angiogenesis (10-14). Tumor-derived EVs may also promote lymphangiogenesis (15, 16). Despite the number of studies, implicating both RNA (11, 17, 18) and protein (10, 19, 20) cargoes, a clear and universal mechanism has not emerged for the apparently critical role of EVs in angiogenesis. It is also not clear whether the same mechanisms will be used for different types of blood vessels or by different tumor types.

Current angiogenesis therapy focuses on soluble secreted molecules, especially vascular endothelial growth factor (VEGF). However, despite the use of anti-VEGF therapy in some cancers as well as in “wet” age-related macular degeneration (21-25), its utility has been more limited than was originally anticipated (26-28). Therefore, identifying unique mechanisms of angiogenesis is of interest both biologically and therapeutically. Since EVs constitute a fundamentally different form of carrier, transporting either internal cytoplasmic cargoes or transmembrane or lipid-linked surface molecules, EV-induced angiogenesis is likely to represent a distinct mode of action from VEGF and other soluble pro-angiogenesis mediators.

In this study, we investigated the role of EVs released from HNSCC cells on angiogenesis and lymphangiogenesis (Fig. 1A). In vivo tumor-associated angiogenesis correlated with the in vitro SEV production rate of several HNSCC cell lines. Furthermore, SEVs purified from HNSCC cells induced angiogenesis, both in vitro and in vivo. Proteomic analysis of SEVs purified from a panel of HNSCC cell lines revealed Ephrin-type receptors as candidate angiogenic protein cargoes. Blocking and genetic inhibition experiments validated Ephrin type B receptor 2 (EPHB2) as a key SEV cargo that promotes HNSCC-mediated
angiogenesis both \textit{in vitro} and \textit{in vivo}. Mechanistic experiments indicate that SEV-induced ephrin-B reverse signaling through STAT3 is critical for EV-induced HNSCC angiogenesis.
Results

Cellular SEV production correlates with HNSCC tumor-induced angiogenesis.

To investigate the relationship between EVs and tumor angiogenesis, we correlated the rate of in vitro SEV release by 5 HNSCC cell lines with the capacity of those same cell lines to promote in vivo angiogenesis (Fig. 1). SEVs were purified by a cushion density gradient method (29) in order to minimize EV aggregation and enhance EV separation from protein aggregates. EV number, size, marker status and morphology were characterized by nanoparticle tracking, Western blot, and transmission electron microscopy analyses in accordance with current guidelines (30) (Suppl. Fig. 1A-C). For tumor induction, human (SCC61, OSC19, Detroit 562) and mouse (MOC1, MOC2) HNSCC cell lines were injected orthotopically into the tongues of nude or syngeneic mice. The tumors were harvested, fixed, and immunostained for the endothelial cell marker CD31. Analysis of CD31 area/tumor area revealed that Detroit 562 and MOC2 tumors had significantly higher blood vessel density than tumors derived from the other cell lines (Fig. 1B, 1C). Detroit 562 and MOC2 cells also exhibited higher rates of in vitro SEV secretion (Fig. 1D). Comparison of the association between tumor blood vessel density and SEV secretion rate across all the cell lines by linear regression revealed a strong correlation (Fig. 1E, R squared value = 0.78).

Purified HNSCC SEVs promote endothelial tube formation

To directly test the angiogenesis-promoting effects of HNSCC-derived SEVs, we performed an in vitro tube formation assay. Human umbilical vein endothelial cells (HUVECs) were treated for 12 hours with conditioned medium or EVs purified from 2 different human HNSCC cell lines, OSC19 and Detroit 562, before obtaining phase contrast images for analysis of total tube length and number of junctions. Compared to control media, conditioned medium from HNSCC cells significantly increased endothelial tube network formation. By contrast, conditioned medium that was depleted of EVs by ultracentrifugation for 2 hours at 100,000 x g did not increase tube formation over that of control media (Fig. 1F). Finally, the addition of purified HNSCC SEVs to the control medium induced formation of a significantly more extensive tube network (Fig. 1F). We tested two concentrations of SEVs, 5 x 10^7 - which we estimate to be approximately the number of SEVs contained in the 400 µl of OSC19 conditioned medium used in the assay - and 1x10^8, which had slightly increased efficacy for Detroit 562 EVs but not for OSC19 EVs. In contrast, LEVs at the same concentrations did not affect tube formation (Suppl. Fig. 2A). Purified SEVs from 2 other HNSCC cell lines also induced HUVEC tube formation, suggesting that this is a common property of HNSCC EVs (Suppl. Fig. 2B). To examine the effects of HNSCC-derived EVs on lymphangiogenesis, we performed the same assay using lymphatic endothelial cells (HMVEC-
Proteomics and transcriptomics analysis identifies EPHB2 as a candidate angiogenic EV cargo

To determine whether HNSCC SEVs carry angiogenic proteins, we performed proteomic analysis of SEVs purified from 3 different human HNSCC cell lines. One of the datasets was published previously for SEVs from SCC61 cells (31). In addition, we performed proteomics analyses on SEVs purified by cushion density gradient ultracentrifugation from OSC19 and Detroit 562 cells, identifying respectively 1016 and 1237 unique proteins (Suppl. Table 1, 2). From these three datasets, we identified a number of proteins that have been shown to regulate angiogenesis and/or lymphangiogenesis (Table 1)(32-52). Notably, multiple ephrin and ephrin receptor (Eph) family proteins were detected, suggesting that this class of proteins could drive angiogenesis in HNSCC tumors. Furthermore, EPHB2 and EPHA2 were detected in all three HNSCC SEV samples (Table 1).

To prioritize our results, we analyzed publicly available RNA-Seq data for HNSCC patient tumors in the TCGA database to determine whether RNA expression levels of EPHB2 or EPHA2 correlate with patient outcome. This analysis revealed that overexpression of EPHB2 - but not EPHA2 - significantly correlates with decreased survival of HNSCC patients (Fig. 2A). EPHB2 is also an interesting cargo, since it can promote ephrin-B reverse signaling (53, 54) and studies in genetically engineered mouse models have shown that ephrin-B reverse signaling plays a critical role in both developmental and tumor angiogenesis (32, 55, 56). In addition, further analysis of the TCGA RNA-Seq data using gene set enrichment analysis revealed that the Hallmark Angiogenesis gene set is closely correlated with EPHB2 mRNA expression in HNSCC patient tumors (Fig. 2B).

We previously showed that the HNSCC-overexpressed protein cortactin promotes exosome secretion (31). Cortactin overexpression had been previously found in small datasets to correlate with poor patient prognosis (57-59). Analysis using the TCGA RNA-Seq dataset confirmed that cortactin overexpression significantly correlates with decreased survival of HNSCC patients (Fig. 2C). Based on these data, we hypothesized that key exosome cargoes might synergize with cortactin overexpression to drive poor patient prognosis. Indeed, analysis of TCGA RNA-Seq data revealed that co-overexpression of cortactin with EPHB2 further stratified patients into a subset with greatly decreased survival, compared with patient tumors with overexpression of only cortactin, only EPHB2, or no overexpression of either gene (Fig. 2D and
Suppl. Table 3). Conversely, analysis of the relationship between cortactin and EPHB2 expression with the Hallmark Angiogenesis gene set revealed that only EPHB2 expression correlated with expression of genes in that gene set (Fig. 2D). These data are consistent with cortactin as a general regulator of exosome secretion and EPHB2 as a specific angiogenic cargo. Immunofluorescent analysis of a tissue microarray with HNSCC samples from 103 patients further revealed that EPHB2 and cortactin co-overexpression is closely related to regional lymph node and distant organ metastasis, especially for undifferentiated tumors (Fig. 2E and Suppl. Fig. 3). Based on these data, we focused on EPHB2 as a promising candidate to test for its role in SEV-mediated angiogenesis.

**EPHB2 in SEVs promotes endothelial tube formation**

In Western blot analyses, EPHB2 protein was expressed in SEVs from multiple human HNSCC cell lines (Fig. 3A). As it was expressed at the highest levels in OSC19 cells, we focused on this cell line for most further mechanistic studies. Consistent with the ability of OSC19 SEVs but not LEVs to induce tube formation, we detected very little EPHB2 expression in LEVs in comparison to SEVs (Fig. 3A). EPHB2 can induce angiogenesis by binding to B-type ephrins on endothelial cells and inducing ephrin reverse signaling, a process that is thought to require cell-cell contacts (32, 55). We hypothesized that SEVs could substitute for neighboring cells and induce ephrin reverse signaling at a distance. To test this possibility, we examined phosphorylation of ephrin-B in HUVECs after incubation with OSC19 SEVs. As a positive control, we added recombinant EPHB2 (Fc-EPHB2). As shown in Fig. 3B, both Fc-EPHB2 and purified OSC19 SEVs induced ephrin-B phosphorylation in HUVECs, compared to no treatment control. To test the role of SEV-associated EPHB2 in angiogenesis, we blocked the ability of EPHB2 to bind to ephrin-B ligands on cells by preincubating the OSC19 SEVs with recombinant ephrin-B2 protein (Fc-ephrin-B2) for 10 minutes before adding the mixture to endothelial cells (see schema in Fig. 3C). Compared to control SEVs, uptake of SEVs preincubated with Fc-ephrin-B2 by HUVEC cells was decreased (Suppl. Fig. 4), suggesting that EPHB2-ephrinB interactions mediate binding of a subset of SEVs with cells. Likewise, preincubation of Fc-ephrin-B2 with SEVs reduced their ability to induce HUVEC tube formation in a dose-dependent manner (Fig. 3D). A similar result was observed using SEVs purified from Detroit 562 cells (Fig. 3D). Recombinant Fc-ephrin-B2 protein alone had only a small effect in the opposite direction at the highest concentration.

We also tested the effect of SEVs purified from EPHB2-knockdown (KD) or EPHB2-overexpressed (OE) OSC19 cells on angiogenesis. Similar to the results with Fc-ephrin-B2 blocking of EPHB2 on SEVs, tube formation of HUVECs treated with EPHB2-KD SEVs was significantly reduced compared to those treated with control SEVs (Fig. 4A, B). Conversely,
EPHB2-OE SEVs increased HUVEC tube formation (Fig. 4C, D). The enhanced tube formation by EPHB2-OE SEVs was statistically significant compared to control SEVs when a lower number of SEVs (2.5x10^7) was used as opposed to the higher number (5x10^7) used in the other tube formation assays. These data indicate that both the number and EPHB2 cargo content of SEVs impact endothelial tube formation.

**HNSCC-derived EVs induce in vivo angiogenesis**

To test the role of EV-delivered EPHB2 in in vivo angiogenesis, we performed a Matrigel plug assay (60-62) in which SEVs purified from OSC19 cells were mixed with growth factor reduced Matrigel and implanted subcutaneously in nude mice. After 21 days, the plugs were harvested and analyzed for blood vessel content (Fig 4E). Quantitation of CD31-positive vessel area/image area in the plugs showed that control EVs induced blood vessel formation. By contrast, in the absence of EVs or in the presence of EPHB2-KD or Fc-ephrin-B2-pretreated EVs, very few blood vessels were formed (Fig. 4E). We validated that the Matrigel plugs indeed contained blood vessels by Hematoxylin and Eosin staining as well as co-staining of some samples for both CD31 and hemoglobin (Suppl. Fig 5A).

To determine whether EPHB2 expression in HNSCC tumors regulates angiogenesis, we injected OSC19 cells into the tongues of nude mice. While there was no effect of EPHB2 expression on tumor size, the vessel density in EPHB2-OE OSC19 tumors was significantly increased compared with control and EPHB2-KD tumors (Fig. 4F). EPHB2-KD did not lead to reduced angiogenesis in OSC19 tumors, possibly due to the relatively low basal level of angiogenesis in this cell line (Fig 1E). Indeed, when we assessed angiogenesis in Detroit 562 tumors that have higher baseline levels of angiogenesis and higher SEV secretion rates (Fig 1E), EPHB2-KD reduced angiogenesis (Suppl. Fig. 5B). In addition, EPHB2-OE in Detroit 562 cells led to enhanced tumor blood vessel density (Suppl. Fig. 5B). Although OSC19 is not a metastatic cell line in this model, we analyzed lymph nodes from these mice. We observed one mouse with lymph node metastasis in the EPHB2-OE tumor group, but no metastasis from the other groups (Fig. 4G). For the Detroit 562 cells, KD or OE of EPHB2 respectively decreased and increased the percent of mice with lymph node metastases; however the differences were not statistically significant (Suppl. Fig. 5C).

**miRNAs are not responsible for HNSCC SEV-mediated angiogenesis**

A number of reports have implicated miRNAs as EV cargoes regulating angiogenesis (17, 18, 63). In addition to inducing ephrin reverse signaling, EPHB2 on SEVs could potentially drive angiogenesis by enhancing binding and uptake of SEVs by target cells, to deliver miRNAs. To
determine whether miRNAs are part of the EV-driven angiogenesis we observed, we inhibited miRNA maturation in OSC19 cells by depletion of DICER with shRNA (Suppl. Fig. 6A) and purified SEVs from those cells. To confirm the effect of DICER-depletion on the levels of mature miRNAs in SEVs, we quantitated total RNA as well as candidate mature and pre-miRNA levels in SEVs purified from control and DICER-KD cells. For candidate miRNAs, we chose miR-155 and miR-21, based on their known expression in HNSCC (64, 65) and their previous association with angiogenesis (66, 67). Consistent with the >90 % KD of DICER in cells (Suppl. Fig. 6A), mature miRNA was decreased and pre-miRNA was increased in EVs purified from DICER-KD cells (Suppl. Fig. 6B). Analysis of the total tube length and numbers of junctions of HUVECs treated with EVs from DICER-KD cells did not show significant changes compared to that of HUVECs treated with control EVs (Suppl. Fig. 6C). These data suggest that delivery of miRNA is unlikely to be the mechanism by which EPHB2 on HNSCC SEVs promotes angiogenesis.

**EV-carried EPHB2 promotes angiogenesis by inducing ephrin reverse signaling.**
Our initial characterization of OSC19 SEVs showed that they can induce ephrin-B phosphorylation (Fig. 3B), an early step in ephrin reverse signaling. To further determine whether EPHB2 carried on SEVs induces ephrin reverse signaling, HUVECs were incubated with scrambled control, EPHB2-KD or EPHB2-OE SEVs purified from OSC19 cells for 30 min followed by Western blot analysis for downstream signaling events. Compared to PBS, control SEVs stimulated ephrin-B phosphorylation (Fig. 5A). Consistent with an important role of EPHB2, incubation of HUVECs with SEVs purified from EPHB2-OE and -KD cells respectively increased and decreased phospho-ephrin-B levels in HUVECs. Phosphorylation of STAT3, which is known to be stimulated by ephrin-B reverse signaling (68) was also increased by SEVs in an EPHB2-dependent manner (Fig. 5A). Interestingly, phosphorylation of VEGFR2 was not increased by SEV treatment (Fig. 5A). Since STAT3 phosphorylation was strongly induced by SEVs, dependent on their EPHB2 content, we tested whether inhibition of STAT3 affects SEV-induced tube formation. Indeed, inhibition with the small molecule inhibitor Stattic (69) greatly reduced both tube length and junction number (Fig. 5B). These data indicate that EPHB2 carried on SEVs induces angiogenesis by activation of ephrin reverse signaling and downstream STAT3 activation.

**Discussion**
In this study, we found that a key Eph receptor is exported in HNSCC SEVs and drives tumor-associated angiogenesis. Thus, blocking EphB-ephrin-B interactions by preincubation of SEVs with Fc-ephrin-B2 or by KD of EPHB2 inhibited SEV-induced angiogenesis, both in vitro and in
vivo. Furthermore, EPHB2 expression in HNSCC cells promotes tumor angiogenesis in an orthotopic model. Mechanistically, EPHB2-positive SEVs induce ephrin reverse signaling, as assessed by phosphorylation of ephrin-B and STAT3. Finally, inhibition of STAT3 activity greatly reduced SEV-induced angiogenesis. These data suggest a model in which EV-carried EPHB2 can induce signaling at a distance to recruit blood vessels into tumors, where they facilitate tumor survival and metastasis (Fig. 5C).

Our model, in which key transmembrane or membrane-linked molecules are transported by EVs, may help to explain the unique role that EVs appear to play in driving angiogenesis. Thus, EV-transported membrane-linked molecules could synergize with soluble, diffusible secreted factors. For example, ephrin-B reverse signaling is known to synergize with VEGF signaling, but traditional models would limit Eph-ephrin signaling to cell-cell contacts. By contrast, EVs secreted from the center of a hypoxic tumor could reach the tumor edge and beyond to stimulate angiogenic sprouting towards the source of secreted factors. Gradients of EVs could provide stable cues that synergize with soluble angiogenic factors such as VEGFs or angiopoietins.

Previous studies have also shown that EVs derived from both cancer and normal cells (including mesenchymal stem cells, fibroblasts, microglia, and endothelial progenitor cells) play a key role in angiogenesis (70-73). Some previously reported angiogenic protein EV cargoes include IL-8, IGFBP1, IGFBP3, carbonic anhydrase 9, annexin II, myoferlin, WNT4, VEGFA and VEGFD (13, 19, 20, 74-76). From our proteomics analysis of HNSCC EVs, IL-8, IGFBP1, carbonic anhydrase 9, WNT4, and VEGFR were not observed (Table 1). VEGFA, VEGFC, IGFBP3, Annexin II and myoferlin were observed in at least one HNSCC EV sample, and VEGFA and IGFBP3 have been previously reported to promote angiogenesis in HNSCC (77, 78). In this manuscript, we focused on receptor-ligand transmembrane molecules that could not be transported in any other way than via EVs and that could specifically target the EVs to endothelial cells. Furthermore, in separate antibody array experiments, we did not observe significant amounts of VEGFs or IGFBP3 associated with HNSCC SEVs (not shown), suggesting that only minimal amounts associate with SEVs or that they are lost during our purification process. However, it is certainly possible that additional EV cargoes may promote angiogenesis in HNSCC tumors and some cytoplasmic cargoes could even be carried in the same vesicles as Ephrin receptors and ligands.

In addition to protein cargoes, a number of EV-derived miRNAs have been shown to promote angiogenesis (9, 79). Using DICER-KD to prevent maturation of pre-miRNAs in HNSCC cells, we detected no effect of KD EVs on endothelial tube formation. While our result does not rule out EV-delivered miRNAs being important angiogenic regulators in other systems, it does
suggest that it is not a major driver in our system.

Eph receptors are a subfamily of receptor protein-tyrosine kinases (80) and were first identified as regulators of cell–cell interactions in the nervous system (81, 82), including axon guidance and branching, and development of neuronal growth cones. More recently, knockout mouse studies identified EphB-ephrin-B signaling as a key regulatory control for development of blood and lymph vessels, due to interactions between EPHB4 and ephrin-B2 on venous and arterial endothelial cells, respectively (32, 55). In one of those studies, ephrin-B reverse signaling was also shown to be important for tumor angiogenesis (55). Additional EphB receptors, including EPHB2, may act in a similar manner in remodeling the embryonic vascular system (34).

Ephrins are also involved in cell migration, cell fate, and immunity (83, 84).

Our study is the first to our knowledge to identify a role for EphB-ephrin-B interactions in HNSCC-induced tumor angiogenesis. Furthermore, our study is the first to identify Ephrin family molecules on EVs as key angiogenesis regulators. Consistent with our finding that SEVs can induce ephrin reverse signaling in endothelial cells, a recent study showed that exosomal EPHB2-ephrin-B1 reverse signaling can induce neuronal repulsion (85). These results suggest that EV-based ephrin signaling may be a general paradigm that could mediate ephrin signaling in a variety of contexts, including axon guidance, angiogenesis, and potentially immune cell interactions (86-89).

Consistent with our TCGA analysis (Fig. 2A), EPHB2 overexpression in several other cancer types has been reported to correlate with decreased patient survival (90, 91). We further found that the correlation of EPHB2 overexpression in HNSCC tumors with poor prognosis was even higher if the exosome secretion regulator cortactin (31) was also overexpressed, suggesting that exosomal EPHB2 may drive tumor aggressiveness. Our findings suggest that induction of tumor angiogenesis could be an important component of that process. Future studies should further elucidate the role of EV-carried EPHB2 and other EV-associated Ephrin family molecules in driving near and long distance signaling in cancer.

Anti-angiogenesis therapy has thus far been employed in a subset of cancers and largely consists of anti-VEGF antibody or kinase inhibitor therapy against VEGFR signaling (22, 92-98). In HNSCC, a number of trials testing anti-angiogenesis therapies are either completed or ongoing; however, the effects have been limited (99-104). In the most recent phase 3 trial, conventional therapy with the monoclonal anti-VEGF antibody bevacizumab did not show significant improvement in overall or disease-free survival compared to conventional therapy alone (105). In fact, in the latest national comprehensive cancer network guidelines for HNSCC therapy, no anti-VEGF reagents were recommended (106), despite strong association of numerous angiogenesis regulators with poor prognosis in HNSCC (107, 108). These data suggest that new targets are
needed. One possibility is to test whether EV-based targets such as EPHB receptors or ligands that should synergize with VEGF would lead to greater efficacy.

In summary, we find that EPHB2 carried on HNSCC-secreted EVs can induce ephrin-B reverse signaling and tumor angiogenesis. This mechanism may be important for understanding of the HNSCC microenvironment and may also be a potential target for future cancer therapy.

Methods
Antibodies
phosphorylated ephrin-B (#3481, rabbit polyclonal), EPHB2 (D2X2I, #83029, rabbit monoclonal), phosphorylated STAT3 (Tyr705, D3A7, #9145, rabbit monoclonal), STAT3 (124H6, #9139, mouse monoclonal), phosphorylated VEGFR2 (Ty1175, 19A10, #2478, rabbit monoclonal), and VEGFR2 (55B11, #2479, rabbit monoclonal) were purchased from Cell Signaling Technology (Danvers, MA). ephrin-B2 (EPR10072(B), ab150411, mouse monoclonal), CD31 (EPR3094, ab76533, rabbit monoclonal), Cortactin (4F11, ab33333, mouse monoclonal) were purchased from Abcam (Cambridge, MA). EPHB2 (AF467, goat polyclonal) and goat IgG HRP conjugated antibody (HAF017, rabbit polyclonal) were purchased from R & D systems (Minneapolis, MN). EPHB4 (3D7G8, #37-1800, mouse monoclonal), Alexa Fluor 750 goat anti-rabbit secondary antibody (A21039), Alexa Fluor 647 goat anti-mouse secondary antibody (A32728), Alexa Fluor 488 goat anti-rabbit secondary antibody (A27034) were purchased from Thermo Fisher Scientific (Waltham, MA). and β-actin (AC-74, A5316, mouse monoclonal) was purchased from Sigma-Aldrich (St Louis, MO). Anti-rabbit IgG (H+L) HRP Conjugate (W4011) and anti-mouse IgG (H+L) HRP Conjugate (W4021) were purchased from Promega (Madison, WI). Hemoglobin β/γ/δ/ε Antibody (A-8) Alexa Fluor (A-8, sc-390668, mouse monoclonal) was purchased from Santa Cruz Biotechnology (Dallas, TX). For western blot, antibodies were diluted 1:2000-1:10,000. For immunohistochemistry or immunofluorescent staining, antibodies were diluted 1:200-1:1000.

Cell lines and constructs.
HUVECs (Lonza, Walkersville, MD) were maintained in EBM-2 complete endothelial growth medium (Cambrex Corporation) containing 2% fetal bovine serum (FBS), human recombinant VEGF (rhVEGF), bFGF, human epidermal growth factor (hEGF), insulin growth factor (IGF-I), hydrocortisone, ascorbic acid, heparin, and GA-1000 (gentamicin and amphotericin B, 1 μg/mL) according to supplier instructions. HMVECs-dLyAd (Lonza) were maintained in EBM-2 Basal Medium (Lonza), 5% FBS, hydrocortisone, hFGF, VEGF, IGF-1, ascorbic Acid, hEGF, and GA-1000 according to supplier instructions. HUVECs and HMVECs-dLyAd were used at passage 6.
or less. OSC19, Detroit 562, and SCC61 cells were a gift from Dr. Wendell Yarbrough and were maintained in DMEM with 20% FBS with 0.4 μg/ml hydrocortisone. MOC1 and MOC2 cells were obtained from Dr. Young Kim and were cultured in Iscove’s modified Dulbecco’s medium / F12 at a 2:1 mixture with 10% FBS, 5 ng/mL epidermal growth factor, 400 ng/mL hydrocortisone, and 5 mg/mL insulin (Sigma-Aldrich). Cells were cultured on 100 mm tissue culture plates and grown under 5% CO2 at 37°C. Medium was renewed every 3 days, and cells were split every 4 days. Stable knockdown or stable expression of genes was achieved using the ViraPower Lentiviral expression system (Thermo Fisher Scientific) according to the manufacturer’s protocol. Gene manipulation from polyclonal stable cell lines was checked by Western blot, and cells were passaged for no longer than a month before discarding and thawing early-passage cultures. Cells were periodically checked for mycoplasma contamination. Lentiviral shRNA constructs for DICER (KD1: TRCN0000051260, 5’-CCGGCCACACATCTTCAAGACTTAAGTGTTTTTTG-3’, and KD2: TRCN0000051262, 5’-CCGGGCTGGCTGTAAAGTACGACTACTCGAGTAGTCGTACTTTACAGCCAGCTTTTG-3’), and EPHB2 (KD1: TRCN0000006423, 5’-CCGGGCTAGACAAGATGATCCGCAACTCGAGTGCGGATCATCTTGTCTAGCTTTTT-3’, and KD2: TRCN0000006426, 5’-CCGGCGGGAGTTTGCCAAGAAATTCTCGAGAATTTCCTTGGCAAACTCCCGTTTTT-3’) were from Millipore-Sigma. The stable EPHB2 overexpression construct was made by cloning the open reading frame of EPHB2 downstream of the CMV promoter into pLOC lentiviral vector (Dharmacon).

**Isolation of EVs from conditioned medium and EV characterization**

Cells were cultured to 80% confluence, and washed with PBS before culturing for 48 h in OptiMEM, which is a serum-free but growth factor-containing medium. Conditioned medium was subjected to serial centrifugation to respectively sediment live cells (300 x g for 5 min), dead cells (2,500 x g for 25 min), and LEVs (10,000 x g for 30 min in Ti 45 rotor). To obtain SEVs, we used a cushion-density gradient protocol (29). Briefly, 30 mL of the supernatant from the 10,000xg centrifugation step was overlaid onto 2 ml of OptiPrep (60% wt/vol aqueous iodixanol; Axis-Shield PoC, Oslo, Norway) and centrifuged at 100,000 x g for 4 h in an SW32 rotor. After the spin, 1 ml medium from the bottom of the tube was mixed with 2 ml OptiPrep to make a SEVs-Optiprep mixture with a final 40% wt/vol concentration of iodixanol. This mixture was then subjected to density gradient ultracentrifugation by adding it into the bottom of a 14 × 89-mm tube and overlaying OptiPrep solutions (20% wt/vol, 10% wt/vol, and 5% wt/vol) diluted with 0.25 M sucrose/10 mM Tris, pH 7.5. A continuous gradient from 40%-5% was made through
centrifugation at 100,000 x g (24,000 rpm in SW40 rotor) for 18 h. 12 density gradient fractions were collected and diluted in PBS before pelleting through another round of centrifugation at 100,000 x g for 3 h, followed by washing and resuspension in PBS.

Analysis of size and protein concentration of EVs
All EV preparations were analyzed for size and concentration by nanoparticle tracking with a Zetaview analyzer (Particle Metrix GmbH, Meerbusch, Germany) as recommended by the company. EV protein levels were determined using the microBCA Protein Assay Reagent kit (Pierce) according to the manufacturer’s instructions.

Transmission electron microscopy analysis
Transmission electron microscopy (FEI/Philips CM200 FEG, Amsterdam, Netherlands) was performed on purified EVs. For negative staining of purified EVs, Formvar carbon film–coated grids (FCF-200-Cu; Electron Microscopy Sciences) were washed in Milli-Q water (MilliporeSigma), followed by 100% ethanol. For each step, excess liquid was removed by wicking with filter paper. 10-μl samples were added to grids for 30 min. Grids were immediately stained with 2% phosphotungstic acid, pH 6.1, for 20s and allowed to dry overnight. Grids were imaged using a FEI Tecnai T12 transmission electron microscopy (120 kV LaB6 source), Gatan cryotransfer stage, and AMT XR41-S side-mounted 2K × 2K CCD camera, 2102 SC. All SEV and LEV images were collected at 110,000 x with the CCD camera.

Western blot analysis.
Cells were lysed in 50 mmol/L Tris-HCl (pH 7.4), 1% NP40, 0.25% sodium deoxycholate, 150 mmol/L NaCl, 1 mmol/L EDTA, 1 mmol/L phenylmethylsulfonyl fluoride (PMSF), 1 μg/mL pepstatin, 1 mmol/L sodium orthovanadate. EVs were lysed in 1% SDS. For Western blot analysis, 40 μg (for cell lines) or 5 μg (for EVs) protein was resolved on 6 or 10% SDS-polyacrylamide gels and transferred to nitrocellulose membranes. Blots were detected with Pierce Western Blot Signal Enhancer (Thermo Fisher Scientific) and imaged on a chemiluminescence imager (Amersham imager 600, GE healthcare Life Sciences, Pittsburgh, PA)

Endothelial tube formation assay
HUVECs or HMVECs-dLyAd (5 × 10^4 per well) were suspended in 400 μL of Opti-MEM medium containing 10% EBM-2 medium without serum, and gently added to Matrigel-coated wells in a 24-well plate. Cells were treated with 5 × 10^7 per well of tumor-derived SEVs or LEVs in 5 μl of PBS or PBS alone. For the experiments using Fc-ephrin-B2, HUVECs were cocultured with 5 × 10^7
purified HNSCC EVs with or without pre-conjugation (for 10 minutes) of various concentrations of recombinant human ephrin-B2-Fc chimera protein (R & D). As a positive control for ephrin-B phosphorylation, HUVECs were cocultured with 15 μg recombinant human EPHB2-Fc chimera protein (R & D). For inhibition of STAT3 activity, cells were treated with 200 nM Statistic (#2798, Tocris, Minneapolis, MN). After 12 hours, images of cells were captured using phase contrast microscopy (x 10/0.25-N.A. LPlan PH2 objective lens, EVOS XL Core Imaging System, Thermo Fisher Scientific, Waltham, MA). Total tube length and total junction number were calculated automatically using the angiogenesis analyzer plugin in ImageJ (http://image.bio.methods.free.fr/).

Animals
7-week-old female athymic nude mice and C57Bl/6 mice were purchased from Charles River Laboratory (Boston, MA) and kept in a pathogen-free facility approved by the American Association for the Accreditation of Laboratory Animal Care that met all current regulations and standards of the U.S. Department of Agriculture, U.S. Department of Health and Human Services, and the National Institutes of Health. Mice were fed irradiated standard mouse chow and autoclaved, reverse osmosis treated water. Animal procedures were carried out according to a protocol approved by The Vanderbilt University’s Institutional Animal Care and Use Committee.

Orthotopic nude mouse model of HNSCC
We used an orthotopic nude mouse model of HNSCC (109, 110). Cells were harvested from subconfluent cultures by trypsinization and washed with PBS. Oral tongue tumors were established by injecting cells (1 × 10^5 for OSC19, Detroit 562, SCC61 and MOC1; 5 × 10^4 for MOC2) suspended in 30 μl of serum-free DMEM into the tongues of nude (OSC19, Detroit 562, SCC61) or C57BL/6 (MOC1, MOC2) mice as described previously (109). Mice were examined twice a week for tumor size and weight loss. Tumor volume was calculated as \((A)(B^2)\pi/6\), where A is the longest dimension of the tumor and B is the dimension of the tumor perpendicular to A. Mice were sacrificed after 4 weeks (SCC61, OSC19 and Detroit 562) or 2 weeks (MOC1 and MOC2), and tongues were fixed, sectioned and stained. Imaging of fluorescent CD31 staining, Hemoglobin staining, and Hematoxylin-Eosin staining in tumors was performed using an Aperio Versa 200 scanner (Leica) in the Vanderbilt Digital Histology Shared Resource. Lymph node metastasis in Hematoxylin-Eosin stained cervical lymph nodes tissue was assessed using light microscopy.

Assessment of blood vessel density from tumor tissue
After scanning CD31 stained tumor tissue specimens, whole tumor area and CD31 stained area in the tumors was calculated using ImageJ. CD31 stained area / whole tumor area x 100 was assessed as blood vessel density.

**Matrigel plug assay**
The Matrigel plug assay was performed essentially as previously described (61). Briefly, growth factor-reduced Matrigel (350 μl) was mixed in liquid form at 4°C with 50 μl of EVs (30 μg). The Matrigel-SEV mixture was injected subcutaneously into the flanks of 6 to 8-week-old athymic nude mice. After 21 days, mice were sacrificed and the Matrigel plugs were excised and fixed in 4% formaldehyde in phosphate buffer. Plugs were embedded in paraffin, sectioned, and immunostained for CD31. Images were acquired with an A1R Confocal laser Microscope system (Nikon) equipped with a × 20/1.40 NA Plan Apo objective lens (Nikon), and analyzed using ImageJ.

**Tumor tissue microarray**
After appropriate institutional review board approval for retrospective study of patient specimens and records, a tissue microarray (TMA) of 120 primary oral cavity SCCs was created from previously untreated patients with no history of prior cancer. After creation of the array, patient identifiers were purged so the specimens were deidentified. The array was assembled retrospectively from 1997 to 2011 from patients from databases from the Departments of Radiation Oncology, Otolaryngology, and Pathology at Washington University. For each patient, 2 (or 4) two millimeter punches of representative blocks from their primary tumors were taken and placed on the array depending on the amount of primary tumor tissue available. Diagnoses were confirmed by one reviewing pathologist (JSL). Pathology data was garnered from review of the pathology database original reports and clinical follow up from departmental databases and review of the electronic medical records, including date of death where applicable. Multiplex immunofluorescent staining of a 6 μm section of the tissue microarray was performed with antibodies for cortactin and EPHB2 in conjunction with collagen (CNA35) (111) and a nuclear dye (DAPI). Images were acquired by whole slide scanning. Histological staining in differentiated and undifferentiated regions of the tumor tissue was scored for the % of cells positive (0-3) and the intensity of fluorescent staining (1-4) with the final score calculated as a multiplier of these two scores (0-12). Quantitation for each protein was determined for differentiated and undifferentiated portions of the tissue from each patient. Tumor tissue staining scores were subsequently subjected to unsupervised clustering of staining for EPHB2 and cortactin was performed in R to identify possible groups of patients with distinct staining patterns that might have corresponding changes in tumor dissemination.
Mass spectrometry and liquid chromatography MS/MS analysis

Shotgun proteomic analysis of purified EVs was performed by first partially resolving the proteins by about 1.5 cm in a 10% Novex precast gel, excising the protein region, and then performing in-gel tryptic digestion to recover peptides. The peptides were analyzed via Multidimensional Protein Identification Technology essentially as described earlier (112, 113). Briefly, digested peptides were loaded onto a biphase pre-column consisting of 4 cm of reversed phase material followed by 4 cm of strong cation exchange reversed phase material. Once loaded, this column was placed in line with a 20 cm reversed phase analytical column packed into a nanospray emitter tip directly coupled to a linear ion trap mass spectrometer. A subset of peptides was eluted from the SCX material onto the reversed phase analytical via a pulse of volatile salt, those peptides were separated by an reversed phase gradient, and then ionized directly into the mass spectrometry. This proceeded for a total of 11 salt elution steps over the course of approximately 22 hours of data acquisitions. Both the intact masses (mass spectrometry [MS]) and fragmentation patterns (MS/MS) of the peptides were collected in a data-dependent manner utilizing dynamic exclusion to maximize depth of proteome coverage. Resulting peptide MS/MS spectral data were searched against the human protein database to which common contaminants had been appended using Sequest (114) and identifications were collated and filtered using Scaffold (www.proteomesoftware.com).

Bioinformatic and Statistical analyses

Statistical analysis of overall survival and gene expression was assessed for the Cancer Genome Atlas (TCGA) patient data using Linkedomics (115) (http://linkedomics.zhang-lab.org/admin.php). Single-sample Gene set enrichment analysis (ssGSEA) was performed on RNA-seq data of HNSCC patients from TCGA database using from Msigdb hallmark angiogenesis gene set (116). List of the gene set was shown in Supplementary Table 3. We calculated normalized angiogenesis enrichment score across the patients and compared with patients categorized into CTTN -high and -low (greater than average CTTN mRNA expression, CTTN-H; lower than average mRNA expression, CTTN-L; respectively) and EPHB2 -high and -low (greater than average EPHB2 mRNA expression, EPHB2-H; lower than average mRNA expression, EPHB2-L; respectively) concurrently. Kaplan–Meier survival curves were evaluated using log-rank tests using gene expression dichotomized at the median. The association of clinical parameters with survival was computed by univariate Cox regression. The synergistic effect of two genes were analyzed by comparing median expression based dichotomized patient groups with each other. Hazard ratios were assessed by Cox regression method in R (https://www.R-project.org) using package survival
For experimental data analysis, GraphPad Prism 8 software (GraphPad Software, San Diego, CA) was used. For all tube formation assays, the statistical analysis was based on the biological replicate average values, not on the technical replicate values. Normality assumption was tested using a one-way ANOVA for all tube formation assays, analysis of CD31 staining area, EV secretion rates, miR-qPCR results, tumor volumes, and Western blot results. Tukey-Kramer method was used for multiple comparison in the analysis of tube formation assays, CD31 staining area, EV secretion rates, and tube formation assays (Figures 1C, 1D, 4B, and 4D). Dunnet's test was used to compare each of a number of treatments with a single control in the analysis of tube formation assays, CD31 staining areas, and Western blot results (Figures 1F, 3D, 4E, 4F, and 5A). P value was evaluated by t-test in Figures 2A, 2C and 2D. P values <0.05 were considered significant for all statistical analysis.

Study Approval
Animal procedures were carried out according to a protocol approved by The Vanderbilt University’s Institutional Animal Care and Use Committee (approved protocol number: M1800027-00). For the TMA array analysis, The Washington University’s institutional review board approved for retrospective study of patient specimens and records (approved IRB number: 201109193).

Author Contributions
S.S. and A.W. designed the experiments. S.S. conducted the majority of experiments. A.E. and A.Z. stained and analyzed the TMA sections. S.V. and B.Z. carried out bioinformatic and statistical analysis. Y.K. and J.L. supplied clinical materials and information. S.S. and A.W. wrote the manuscript with input and editing from all coauthors.

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Competing interests
The authors declare no competing financial interests.

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Figure 1. Small extracellular vesicles (SEVs) derived from HNSCC cells drive tumor angiogenesis. (A) Hypothesis for study. EVs mediate angiogenesis and lymphangiogenesis to respectively drive distant and locoregional metastasis. (B) Representative images of tongue tumors derived from SCC61, OSC19, Detroit 562, MOC1 and MOC2 cells stained for CD31 (black, identifies endothelial cells). SCC61: n=4, OSC19, Detroit 562, MOC1 and MOC2: n=5. 10 images for each tumor. Scale bar = 100 µm. (C) Plot of CD31-positive vessel area per total tumor area in tongue tumors. SCC61: n=4, OSC19, Detroit 562, MOC1 and MOC2: n=5. Total tumor area and CD31-stained area were calculated using ImageJ. (D) SEV secretion rate of cell lines, calculated from nanoparticle tracking analysis of purified vesicles obtained from a known final number of cells over 48 h. SCC61: n=4, OSC19: n=7, Detroit 562: n=5, MOC1: n=11 and MOC2: n=8. (E) Linear regression models were performed to analyze relationship between SEV secretion rates and blood vessel density in tumors for various cell lines. Adjusted R^2 of the linear regression model = 0.7822. (F) HUVECs were incubated for 12 h in serum-free media + PBS (control), conditioned media of cancer cells (CM), CM ultracentrifuged to remove SEVs (CM-SEV), or serum free media + purified cancer cell SEVs (5 X 10^7 and 1 X 10^8). (Left) Representative images of HUVEC cells cultured with PBS (Control) or OSC19/Detroit 562 SEVs (5 X 10^7). ImageJ outlining of vessels for automated analysis is shown. Scale bar = 500 µm. (Right) Quantification of relative total tube length and junction numbers for the indicated groups. Plots show the average of ≥3 technical replicates per condition for each n value from ≥3 independent experiments. For (C), (D) and (F), box and whisker plots show median and 25th-75th %ile. Tukey-Kramer method was used in (C) and (D), and Dunnet method was used in (F) for statistical analysis. *, P < 0.05; **, P < 0.01; ***, P < 0.001.
Figure 2. EPHB2 overexpression in HNSCC tumors correlates with patient survival and tumor angiogenesis. (A)-(D) Analysis of TCGA RNA-Seq expression data, provisional HNSCC tumor dataset. (A) Kaplan-Meier analysis of relationship between EPHB2 or EPHA2 expression in HNSCC tumors to overall survival. The association of clinical parameters with survival was computed by univariate Cox regression. n=528. (B) Positive correlation between the Hallmark Angiogenesis gene set and EPHB2 mRNA expression in tumors. (Pearson correlation r = 0.402, P = 1.227x10^{-21}). n=528. (C) Relationship between cortactin (CTTN) RNA expression in HNSCC tumors and overall survival. The association of clinical parameters with survival was computed by univariate Cox regression. n=528. (D) (Left) Synergistic effect of EPHB2 and CTTN expression in HNSCC tumors on overall patient survival. The association of clinical parameters with survival was computed by univariate Cox regression. n=528. (Right) Correlation of EPHB2 and CTTN mRNA expression with the Hallmark Angiogenesis gene set in HNSCC tumors. Y-axis is the normalized enrichment score (NES) of hallmark angiogenesis gene set in respective groups. Single-sample Gene set enrichment analysis (ssGSEA) was performed for statistical analysis. n=528. (E) Multiplex staining for EPHB2 (green) and CTTN (red) in a TMA generated from surgical HNSCC specimens (103 patients). Representative staining for patient samples in clusters I and II is shown for differentiated (left panel) and undifferentiated (right panel) HNSCC areas, see also Suppl. Fig. 3 for unmerged representative images. Scale = 100 µm. Cumulative data of intensity scoring (see methods) from the TMA is displayed as a heatmap with unsupervised clustering revealing three distinct clusters. The ratios in cluster I to the other clusters were compared separately using Fisher's test. The two-sided P values are Regional I vs II: 0.07, I vs III: 0.001; II vs III: 0.20; Distant I vs II: 0.03, I vs III: 0.0001, II vs III: 0.08. *, P < 0.05; ***, P < 0.001 shown in the table is in comparison to cluster I.
Figure 3. SEVs carry EPHB2 and induce ephrin reverse signaling. (A) (Left) Representative Western blots of EPHB2 in whole cell lysates (WCL) and SEVs of OSC19, Detroit 562, and SCC61. (Right top) Western blot comparing EPHB2, CD63 and TSG101 in OSC19 whole cell lysates and SEVs (loading 5 μg in each lane). (Right, bottom) Western blot comparing EPHB2, HSP70 and TSG101 in OSC19 SEVs and LEVs (loading 5 μg in each lane). (B) (Top) Representative Western blots of phospho-ephrin-B (p-ephrin-B), ephrin-B2, and β-actin in cell lysates from HUVECs treated with Fc-EPHB2, PBS (-) or 5 X 10^7 OSC19 SEVs (+), as indicated. (Bottom) p-ephrin-B/ephrin-B2 ratio from 3 independent experiments (median and 25th-75th %ile). Dunnet method was used for statistical analysis. (C) Schematic of blocking experiments in which preincubation of Fc-ephrin-B2 with SEVs binds EPHB2 and blocks its binding to ephrin-B ligands on HUVECs. (D) Tube formation assay. HUVECs were cultured with PBS (control), OSC19 SEVs with or without preincubation with recombinant ephrin-B2 (Fc-ephrin-B2), or with Fc-ephrin-B2 alone. (Top) Representative images. Scale bar = 500 μm. (Bottom) Quantification of relative total tube length and junction number. Dot plots represent median and 25th-75th %ile. Tukey-Kramer method was used for statistical analysis using the average of ≥3 technical replicates per condition for each n value from ≥3 independent experiments. *, P < 0.05; ***, P < 0.001 when compared to no added SEVs condition; #, P <0.05; ### P< 0.001 when compared to +SEVs condition.
Figure 4. EPHB2 on SEVs drives in vitro tube formation and in vivo angiogenesis. (A) Representative Western blot of EPHB2 in parental (control), scrambled control (SCR) and EPHB2-KD OSC19 whole cell lysates and SEVs. β-actin and TSG101 serve as loading controls. (B) Tube Formation Assay (Top) Representative images. Scale bar = 500 µm. (Bottom) Quantification of relative total tube length and junction number. Plots show the average of ≥3 technical replicates per condition for each n value from ≥3 independent experiments. ***, P < 0.001 comparing PBS control to other groups. ###, P < 0.001 comparing +Scr SEVs to +KD SEVs groups. (C) Western blot analysis of EPHB2, β-actin and TSG101 in parental, empty vector control (Cont) and EPHB2-overexpression (EPHB2OE) OSC19 whole cell lysates and SEVs. (D) Analysis of tube formation assay from HUVECs treated with the indicated numbers and types of SEVs. Plots show the average of ≥3 technical replicates per condition for each n value from ≥3 independent experiments. (E) Matrigel plug assay in which SEVs or PBS were mixed with Matrigel and implanted subcutaneously in nude mice before harvesting and staining for blood vessels (CD31, green) or nuclei (Hoechst 33342, red). (Top) Representative images of stained Matrigel plug tissue. (Bottom) Quantification of CD31-positive vessel area per total tumor area in Matrigel plugs. Dot plot shows mean ± SE. Dunnet method was used for statistical analysis. n≥15 images per condition in 5 samples from 2 independent experiments. ***, P < 0.001 comparing PBS control to other groups. (F)&(G) Control (Scr), EPHB2-KD and -OE HNSCC cells were implanted in the tongues of nude mice. (F) Top: Representative images and quantitation of CD31-staining (black) in tongue tumors. Scale bar = 100 µm. (Bottom) Quantitative analysis of tumor volume and blood vessel density in each group. n = 5 images (1 image each from 5 tumors, whole tumor sections scanned for image quantitation). (G) Frequency of lymph node metastasis. For (B), (D), (E) and (F), box and whisker plots show median and 25th-75th %ile. Dunnet method was used in (B) and (E), and Tukey-Kramer method was used in (D) and (F) for statistical analysis. *, P<0.05 ***, P < 0.001. n.s. not significant.
Figure 5. EPHB2 on SEVs drives an ephrin-B-STAT3 angiogenic signaling cascade. (A) Representative western blots of phosphorylated (p-ephrin-B, p-STAT3, p-VEGFR2) and total (ephrin-B2, STAT3, VEGFR2, β-actin) proteins in cell lysates from HUVECs treated with PBS or SEVs, as indicated. Graphs show normalized quantitative analyses of the ratio of phosphorylated to total proteins from n=3 Western blots. Box and whisker plots show median and 25th-75th %ile. Dunnet method was used for statistical analysis. * p<0.05, ***p<0.001. (B) Tube formation assay of HUVECs treated with SEVs in the presence or absence of the Stat3 inhibitor Stat3. Representative images and analysis. Scale bar = 500 μm. Box and whisker plots show median and 25th-75th %ile. Tukey-Kramer method was used for statistical analysis using the average of ≥3 technical replicate per condition for each n value from ≥3 independent experiments. *** P < 0.001. (C) Model of SEV-induced angiogenesis pathway. EPHB2 on cancer cell-derived SEVs binds to ephrin-B2 on endothelial cells and activates ephrin-B2 reverse signaling with downstream phosphorylation and activation of STAT3. While phosphorylation of VEGFR2 downstream of ephrin-B2 reverse signaling has been reported (55), we did not detect changes due to SEV treatment, suggesting selective activation of ephrin reverse signaling pathways by SEVs.
Table 1. Angiogenesis-related proteins in human HNSCC derived SEVs

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