Hexokinase 2-driven glycolysis in pericytes activates their contractility leading to tumor blood vessel abnormalities

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Defective pericyte-endothelial cell interaction in tumors leads to a chaotic, poorly organized and dysfunctional vasculature. However, the underlying mechanism behind this is poorly studied. Herein, we develop a method that combines magnetic beads and flow cytometry cell sorting to isolate pericytes from tumors and normal adjacent tissues from patients with non-small cell lung cancer (NSCLC) and hepatocellular carcinoma (HCC). Pericytes from tumors show defective blood vessel supporting functions when comparing to those obtained from normal tissues. Mechanistically, combined proteomics and metabolic flux analysis reveals elevated hexokinase 2 (HK2)-driven glycolysis in tumor pericytes, which up-regulates their ROCK2-MLC2 mediated contractility leading to impaired blood vessel supporting function. Clinically, high percentage of HK2 positive pericytes in blood vessels correlates with poor patient overall survival in NSCLC and HCC. Administration of a HK2 inhibitor induces pericyte-MLC2 driven tumor vasculature remodeling leading to enhanced drug delivery and efficacy against tumor growth. Overall, these data suggest that glycolysis in tumor pericytes regulates their blood vessel supporting role.
Pericytes have an important regulatory role in stabilizing blood vessel wall, controlling vessel dilation, and regulating vascular perfusion\(^1\)-\(^3\). Unlike other host cells, pericyte expresses specific surface markers/receptors such as platelet derived growth factor receptor \(\beta\) (PDGFR\(\beta\)) and CD146 (also known as the melanoma cell adhesion molecule (MCAM)), and respond to specific growth factors including PDGF, transforming growth factors and angiopoietins\(^2\),\(^4\). During angiogenesis, endothelial cells secrete PDGF to recruit pericytes to newly formed vessels via activation of the PDGFR\(\beta\) signaling\(^1\),\(^12\). Unlike PDGFR\(\beta\), CD146 was originally identified as an endothelial cell marker during angiogenesis\(^6\), which has been recently shown to highly express in pericytes too, and important for regulating pericyte-endothelial cell interaction during blood–brain barrier development\(^8\). Therefore, these two surface proteins are routinely used as positive markers for pericyte identification.

Just as tumor endothelial cells differ from quiescent endothelial cells in normal tissues, tumor-derived pericytes appear to be abnormal or dysfunctional when compared with pericytes in normal tissues\(^7\),\(^8\). It is unclear whether cancer cells instruct the pericyte fate changes to facilitate tumor growth. Indeed, defective pericyte-endothelial cell interaction is observed in many cancers, which is characterized by compromised blood vessel integrity\(^9\),\(^10\). For instance, the loss or detachment of pericytes from tumor blood vessels is frequently observed in some cancers and is believed to promote cancer metastasis\(^11\),\(^12\). On the other hand, pericytes have been shown to protect endothelial cells from anti-angiogenic drug (i.e. bevacizumab), and therefore combination treatment with PDGFR\(\beta\) tyrosine kinase inhibitor (i.e. eliminating PDGFR\(\beta\) positive pericytes) and VEGF inhibition more efficiently block tumor angiogenesis than anti-VEGF therapy alone in several animal models\(^13\),\(^14\). However, the combination therapy has failed to provide any therapeutic benefit in the clinical trial of renal carcinoma patients and even shows additive toxicity in the patients\(^14\),\(^15\). Therefore, it is a clinically unmet need to explore the molecular mechanism of pericyte-endothelial cell interaction in preclinical setting. Due to the lack of a cellular model for studying human pericyte biology, the exact cause of abnormal pericyte behavior in tumors is unknown. Previous studies show that the regulation of metabolic pathways determines the cell state and fate decisions in endothelial cells\(^16\). For instance, Inhibition of the glycolytic activator PFKFB3 in endothelial cells can improve pericyte coverage and blood vessel function, which therefore enhances the delivery and efficacy of chemotherapeutics drugs in preclinical animal models\(^17\). However, the role of metabolic rewiring in human pericyte biology still remains largely unknown.

In this study, we establish the role of pericyte-hexokinase 2 (HK2) driven glycolysis in remodeling tumor vasculature. By developing a method for isolation and culture of pericytes from normal adjacent tissues (NPC) and tumors (TPC) derived from NSCLC (non-small-cell lung cancer) and HCC (hepatocellular carcinoma) patients respectively, we showed that TPC had abnormal blood vessel supporting functions as compared with NPC in two different cancers. Combined proteomics and metabolic flux analysis revealed an increased HK2-driven glycolysis in TPC as compared with NPC, which prohibited its blood vessel supporting role via activation of the ROCK2-MLC2 mediated contractility. Importantly, depletion of HK2/ROCK2 restored the blood vessel supporting function of NPC, while treatment of HK2 inhibitor induces MLC2-driven tumor vasculature remodeling to enhance chemotherapy delivery and efficacy against tumor growth. The role of pericyte-HK2 expression in this process was further illustrated by the fact that administration of chemotherapy reduced tumor growth in mice that were co-injected with tumor cells and HK2-depleted TPC as compared with mice that were co-injected with tumor cells and scramble transfected TPC.

Overall, our work develops a cellular model for studying pericyte biology and uncovers that pericyte-HK2-driven glycolysis induces tumor blood vessel abnormality by activating ROCK2-MLC2 mediated contractility.

**Result**

**Development of the microbead activated vascular cell sorting enrichment method coupled with FACS.** Although previous studies have managed to isolate pericytes from mouse brain tissues\(^18\),\(^19\), there is no effective protocol for isolation and culture of human pericytes from paired normal adjacent tissues and tumors. Due to the fact that both lung and liver tissues contain a rich population of vascular cells, they were therefore chosen as a source of human pericyte isolation in this study (Supplementary fig. 1). Since anti-human CD146 antibody conjugated with FITC and anti-FITC magnetic microbeads are commercially available, we proposed to employ a CD146\(^+\)–FITC-microbead activated cell sorting (MACS) method to enrich the vascular cell population from tissue samples first, and then used positive and negative selection markers to isolate pericyte population via fluorescence activated cell sorting (FACS) (Supplementary fig. 1). In order to carry out our pericyte purification method, we first examined whether CD146 was expressed on vascular endothelial cells and pericytes in normal adjacent tissue and tumors derived from NSCLC and HCC patients. Based on the fact that CD34 and \(\alpha\)-SMA/PDGFR\(\beta\) is known as a vascular endothelial cell and pericyte marker respectively\(^7\),\(^20\), they were used for defining the CD146 staining in this study. We then performed triple immunofluorescent staining of these markers with paired normal adjacent tissues and tumors derived from NSCLC and HCC patients respectively, showing that CD146 was expressed in both CD34 positive (+ve) vascular endothelial cells and \(\alpha\)/SMA/PDGFR\(\beta\)+ve pericytes within normal adjacent tissues and tumors derived from NSCLC and HCC patients, while ~65–72% percentage of blood vessels were CD146 and \(\alpha\)-SMA double positive in both normal adjacent tissues and tumors derived from NSCLC and HCC patients respectively (Fig. 1a–f and Supplementary fig. 2a, b). No significant difference in the percentage of CD146 and \(\alpha\)-SMA double positive blood vessels was observed between normal adjacent tissues and tumors (Fig. 1c, f). In addition, we showed that \(\alpha\)-SMA was also expressed in PDGFR\(\beta\)+ve pericytes around CD34 positive blood vessels (Supplementary fig. 2c, d), which was consistent with the previous finding\(^8\). Therefore, our results suggested that CD146–FITC-microbead activated cell sorting method could be used to enrich the vascular cell population from tissue samples derived from NSCLC and HCC patients.

To isolate pericytes from normal adjacent tissues and tumors, the dead cells and debris were first removed from mechanically minced and enzymatically digested tissues. The single cell suspension was then labeled with FITC-conjugated CD146 antibody and then enriched by using anti-FITC magnetic microbeads. The enriched vascular cell population was subsequently labeled with antibodies specific for endothelial cells (CD31), immune cells (CD45), and pericyte markers (PDGFR\(\beta\)). The labeled cells were then collected by using FACs (Supplementary fig. 1). FACs plots showed the gating strategy for the separation of endothelial cells (EC) and pericytes (PC) from normal adjacent tissues (NEC and NPC) and tumor tissues (TEC and TPC) derived from NSCLC and HCC (Fig. 1g–j and Supplementary fig. 2e–h). Importantly, our MACS enrichment method successfully improved the efficacy of isolating human pericytes from paired normal adjacent

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\(\beta\) - PDGFR\(\beta\)

\(\alpha\) - \(\alpha\)-SMA

\(+\) - positive

\(-\) - negative

\(\beta\) - PDGFR\(\beta\)
tissues and tumors derived from NSCLC and HCC patients almost by 7–13 folds as compared to FACS method alone (Fig. 1k, l). Compared with FACS method alone, our protocol enhanced the successful chance of isolating enough pericytes mainly due to MACS enrichment method and temperature-controlled incubation with optimized amount of collagenase and DNase I.

Characterization of the isolated pericytes from normal adjacent tissues and tumors derived from NSCLC and HCC patients. To confirm the purity of our isolated pericytes derived from normal adjacent tissues and tumors, we performed flow
both NSCLC/HCC-derived NPC and TPC strongly expressed fi
addition, no signifi
can be observed in tissue section, which possessed a cell body with a prominent nucleus and a small
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migration ability. Phase contrast microscope analysis revealed that NPC and TPC had the morphological characters of
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similar doubling times and could be cultured up to 12 passages (Fig. 3a, b). Further RT-PCR ana-
lysis confirmed that these pericytes either expressed relatively low or undetectable levels of endothelial cell markers CD31 and CD34, immune cell marker CD45, or fibroblast marker FAP (fibroblast associated protein) (Fig. 2a, b and Supplementary fig. 3a, b). Further RT-PCR ana-
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Next, we examined their cell morphology, proliferation, doubling time, passage number, senescence, viability, and migration ability. Phase contrast microscopy analysis revealed that both NPC and TPC had the morphological characters of pericytes as observed previously in tissue section, which possessed a cell body with a prominent nucleus and a small content of cytoplasm with several long processes (Fig. 2b and Supplementary fig. 3b). Functional studies revealed that the NPC and TPC derived from NSCLC or HCC patients expressed PDGFRα and CD146 as well as newly identified pericyte markers desmin and CD13, but did not express endothelial cell marker CD31 and CD34, immune cell marker CD45, or fibroblast marker FAP (fibroblast associated protein) as compared with human umbilical vein endothelial cells (HUVEC), peripheral blood mononuclear cell (PBMC), and human fibro-
blasts respectively (Fig. 2c and Supplementary fig. 3c). In contrast, both NSCLC/HCC-derived NPC and TPC strongly expressed pericyte markers PDGFRB and CD146 as compared to fibroblasts and HUVEC respectively (Fig. 2c and Supplementary fig. 3c). In addition, no significant difference in the expression level of pericyte markers between either NPC and TPC, or at passage 3 and 12 was observed (Fig. 2a–c and Supplementary fig. 3a–e), suggesting that our cultivating method could maintain the pericyte identity of the isolated cells over passages.

To evaluate the blood vessel supporting role of NPC and TPC in vivo, we performed in vivo matrix plug assays by embedding NPC/TPC/fibroblasts/SMC in matrigel-fibrin matrix with HUVEC, and then implanted them into nude mice subcuta-
newously. Matrix gel plug assays indicated that co-transplantation of HUVEC and NSCLC/HCC-derived-NPC and-TPC signifi
cantly increased the number of blood vessels as compared with HUVEC transplant alone (Fig. 3g, h). Further histological examination showed that co-transplantation of HUVEC with NPC, but not TPC, supported the formation of functional blood vessels (containing red blood cells) with strong basement membrane marker collagen IV expression (Fig. 3g–j), while smaller and irregular shaped blood vessels, reduced collagen IV staining were observed in HUVEC co-transplanted with TPC (Fig. 3g–j). Previous studies have shown that the expression of basement membrane marker collagen IV has been implicated in blood vessel integrity and perfusion.17 As expected, both smooth muscle cells and fibroblasts had a very limited effect on tumor blood vessel formation and collagen IV expression (Fig. 3g, h, j), while SMC reduced the lumen diameter of blood vessels most significantly as compared with NPC (Fig. 3g, i). Finally, neither NPC, TPC, fibroblast, or SMC showed difference in tube coverage (Fig. 3k).

Since pericytes have been implicated in regulating capillary dilation and contraction,1,14 we therefore assessed the contractile properties of NPC and TPC by performing time-lapse microscopy studies with them after treated with carbachol. After the treatment, NSCLC/HCC-derived TPC strongly contracted and

endothelial cell proliferation and migration, we exposed HUVEC cells to conditioned medium from NPC/TPC derived from NSCLC or HCC patients, indicating that the conditioned media from either NPC or TPC increased the proliferation of HUVEC (Fig. 3a), while neither of them showed any effect on HUVEC migration in a wound healing assay (Fig. 3b). To examine the functional role of NPC and TPC in blood vessel formation, we fluorescently labeled NPC and TPC as well as fibroblasts and smooth muscle cells (SMC), and then co-cultured HUVEC with either of them on matrigel. After co-culturing, we evaluated their effect on total tube length, branch points, and numbers (Fig. 3c–f and Supplementary fig. 4a–d). Although NPC, TPC, fibroblasts, and smooth muscle cells aligned along tubes when added to HUVECs, they had different effects on tube lengths and branching. Strikingly, co-culturing HUVEC with TPC reduced total tube length and branch points and number as compared to HUVEC co-cultured with either NPC, fibroblasts, or HUVEC alone, while normal fibroblasts had no effect on all of these (Fig. 3c–f and Supplementary fig. 4a–d). Unlike NPC, TPC or normal fibroblast, SMCs pulled tubes apart following alignment and caused a rapid regression of the tubular network within 24 h (Fig. 3c–f and Supplementary fig. 4a–d).

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showed up to a 30% change in surface area, while NSCLC/HCC-derived NPC displayed a modest change in their surface area (Supplementary fig. 4e, f). In contrast, no significant change in the surface area of HUVEC/fibroblast was observed post-treatment (Supplementary fig. 4e, f). Together, these results showed the functional difference between NPC and TPC in vitro and in vivo, indicating that TPC had poor BV supporting functions compared with NPC, which may be linked to its increased contractility phenotype.

**Combined proteomics and metabolic flux analysis reveals an elevated HK2-driven glycolysis in TPC.** Metabolic reprogramming determines cell proliferation, migration, and contractility\(^{21}\). Indeed, endothelial cells undergo PFKFB3 dependent glycolytic
switch to match their high energy demand for migration and proliferation during angiogenesis. However, the role of metabolic state in determining pericyte’s BV supporting function remains unknown. In addition, it is unclear whether NPC and TPC express different glycolytic gene signature. We therefore performed RT-PCR analysis of the NPC and TPC isolated from NSCLC or HCC patients to compare their glycolytic gene pathway expression, indicating that NPC and TPC had distinct difference in the expression of genes associated with glycolysis (Fig. 4a, b). In both NSCLC and HCC, the expression of glycolytic gene, hexokinase 2 (HK2) was significantly up-regulated in TPC as compared to NPC (Fig. 4a, b). Although previous studies indicated that inhibition of PFKFB3 in human placenta pericytes prohibited their cell migration and proliferation abilities, we did not observe any difference in the PFKFB3 expression between NPC and TPC (Fig. 4a, b). In addition, proteomics analysis indicated that the expression of HK2, but not PFKFB3 as well as other glycolytic enzymes and glucose transporters, was significantly increased in TPC as compared to NPC (Fig. 4c). Western blot analysis confirmed the up-regulation of HK2 protein expression in NSCLC/HCC-derived TPC as compared with NPC (Fig. 4d). Based on these findings, we next examined the glycolytic capacity of NPC and TPC by performing glucose uptake assay, lactate assay, Seahorse analysis as well as metabolic flux analysis. As expected, NPC and TPC showed no difference in the glucose uptake, while the lactate production was significantly increased in TPC as compared with NPC (Fig. 4e, f). Further Seahorse assay indicated an increase in extracellular acidification rate (ECAR) but decreased oxygen consumption rate (OCR) in TPC as compared that to NPC (Fig. 4g, h and Supplementary fig. 5a, b), implying that TPC relied on glycolysis for their energy production. To obtain the mechanistic underpinnings of the deregulated TPC metabolism, we carried out 13C6-glucose labeled metabolic flux analysis using U-13C6-glucose to estimate metabolic fluxes in TPC and NPC derived from NSCLC or HCC patients. Metabolic tracking revealed the relative abundance of glycogenolysis metabolite M3 lactate was increased in TPC as compared to NPC, while no significant difference in the relative abundance of TCA (tricarboxylic acid) cycle metabolites directly derived from U-13C6-glucose, such as M2 citrate, M2 ketoglutarate, M2 succinate, M2 fumarate, and M2 malate, was observed between NPC and TPC (Fig. 4i and Supplementary fig. 5c).

To prove the clinical significance of our observation, we performed an immunofluorescence study with normal adjacent tissues and tumors derived from NSCLC and HCC patients, indicating that the percentage of pericyte-HK2 positive blood vessels was significantly up-regulated in tumors as compared to normal adjacent tissues (Fig. 4j and Supplementary fig. 5d). We next performed triple immunostaining of PDGFRα, CD34, and α-SMA with our own NSCLC and HCC patient cohorts, indicating that high percentage of pericyte-HK2 positive blood vessels (i.e. when more than the mean number of tumor blood vessels are pericyte-HK2 positive blood vessels across individual patient samples in our NSCLC and HCC cohorts respectively) was associated with poor overall survival in NSCLC and HCC patients respectively (Fig. 4k and Supplementary fig. 5e). Collectively, our findings establish a major metabolic reprogramming of TPC, evidently through up-regulated HK2-driven glycolysis in TPC as compared to NPC.

**Pericyte-HK2-driven glycolysis inhibits the blood vessel supporting function of pericyte via ROCK2-MLC2 mediated contractility.** Next, the function relevance of glycolytic switch in TPC was examined. We pre-treated TPC with placebo or HK2 inhibitor for 15 min and then co-cultured them with HUVEC cells in tube formation assays. In both NSCLC and HCC, we showed that pre-treatment with HK2 inhibitor enhanced the total tube length, branch points, and number in HUVEC co-cultured with TPC, when compared with HUVEC co-cultured with placebo pre-treated TPC (Fig. 5a-d). To test its effect in vivo, we embedded HUVEC with TPC pre-treated with placebo or HK2 inhibitor 3-bromopyruvate (3-BP) in matrigel fibrin matrix, which were then implanted into nude mice and given either placebo or HK2 inhibitor treatment before harvesting. Strikingly, treatment with HK2 inhibitor did not affect the capacity of TPC for promoting blood vessel formation, while it enhanced blood vessel diameter and collagen IV expression in HUVEC co-transplanted with TPC as compared to HUVEC co-transplanted with placebo pre-treated TPC (Fig. 5e-h). No significant difference in percentage of coverage between these two groups (Fig. 5i). Importantly, treatment with HK2 inhibitor did not affect blood vessel density, blood vessel diameter, and collagen IV expression in HUVEC transplant alone as compared with placebo treated group (Fig. 5e-h). Further metabolic study showed that pre-treatment with HK2 inhibitor had no effect on glucose uptake in TPC as compared with placebo treated TPC (Supplementary fig. 6a), while it significantly inhibited lactate production in TPC (Supplementary fig. 6b). In addition, pre-treatment with HK2 inhibitor significantly reduced carbachol-induced cell contraction in TPC as compared to placebo pre-treated TPC (Supplementary fig. 6c). These results demonstrated that the elevated HK2 expression in TPC contributed to its defect in cell contractility and blood vessel supporting function, and that was determined by glycolysis.

Consistent with these findings, Rho-associated coiled-coil containing protein kinase (ROCK) signaling pathway has been implicated in regulating pericyte shape and contractility, while these processes require energy supply mainly from glycolysis. Non-muscle myosin II is a hexameric actin-binding protein that is formed by two heavy chains and two regulatory light chains, which has an important role in cell contractility. Interestingly, ROCK can directly phosphorylate MLC2 to increase contraction.
Fig. 3 Tumor-derived pericytes display defective blood vessel supporting functions in vitro and in vivo. a CCK8 proliferation assay showed that exposure with conditioned medium from NSCLC/HCC-derived NPC or TPC significantly enhanced the proliferation of HUVEC as compared to untreated HUVEC (n = 5 experimental repeats). b Wound healing assay of the HUVEC after exposed with condition medium from NSCLC/HCC-derived NPC or TPC. Representative pictures of wound healing assay in each group are given (n = 3 independent experiments). c HUVEC were either cultured alone or co-cultured with CFSE (green fluorescent dye) labeled NSCLC-derived-NPC/-TPC, fibroblast or smooth muscle cells (SMC) in a tube formation assay. d–f Bar charts show the relative total tube length, branch points and number in each group (n = 3 independent experiments). g Representative images of the serial sections of matrix plugs after immunostained with CD34, PDGFRβ, or collagen IV antibody in each group are given. h–k Violin plots show the quantification of the total number of blood vessels, lumen diameter, collagen IV level and percentage of coverage in each group (n = 3 mice per group). Results are given as means ± SEM. a Two-way ANOVA, b, d–f, h–k One-way ANOVA. Scale bars in b represents 100 μm, c 200 μm, g 50 μm.
Importantly, administration of HK2 inhibitor significantly enhanced total tube length, branch point, and number (Fig. 5l). Furthermore, pre-treatment with ROCK inhibitor also prohibited carbachol-induced cell contraction in NSCLC/HCC-derived TPC pre-treated with either placebo or ROCK inhibitor GSK429286A in matrigel, indicating that pre-treatment with ROCK inhibitor significantly down-regulated the expression of ROCK2 and p-MLC2 in TPC (Fig. 5k). To determine the role of ROCK2 in regulating the activity of myosin II during cell contraction\textsuperscript{26}. As expected, western blot analysis indicated that the expression of ROCK2 downstream effector MLC2 (Myosin-light-chain 2) was increased in TPC as compared to that in NPC (Fig. 5j).
TPC (Supplementary fig. 6c). Western blot analysis indicated that treatment with ROCK inhibitor down-regulated the phosphorylation of MLC2 in TPC as compared with placebo-treated TPC (Supplementary fig. 6d). Clinically, the pericryte expression of ROCK2 was up-regulated in NSCLC/HCC-derived tumors as compared with normal adjacent tissues (Supplementary fig. 6e), while high ratio of ROCK2 to pericyte marker expression correlated with poor survival in NSCLC and HCC patients respectively (Supplementary fig. 6f, g).

Since human ROCKs consist of two isoforms, ROCK1 and ROCK2, we further examine whether HK2 regulates TPC’s blood vessel supporting role via ROCK1 or ROCK2. We first stably knock-downed HK2, ROCK1, or ROCK2 expression in NSCLC/HCC-derived TPC, and also overexpressed ROCK2 in HK2-depleted TPC, which was confirmed by the RT-PCR and western blot analysis (Supplementary fig. 6h–k). Interestingly, depletion of HK2 down-regulated ROCK2, but not ROCK1 expression, in NSCLC/HCC-derived TPC (Supplementary fig. 6h). Further tube formation assays showed that stable knock-down of pericycle-HK2 expression or depleting pericycle-ROCK2 by shRNA/ROCK inhibitor HA-1077 (i.e. a more potent ROCK2 inhibitor) increased the total tube length, branch points, and number in HUVEC co-cultured with TPC as compared with HUVEC co-cultured with scrambled transfectant TPC, while silencing ROCK1 had no significant effect on tube formation (Supplementary fig. 6l–s). Furthermore, overexpression of ROCK2 in HK2-depleted TPC reduced the enhanced total tube length, branch points, and number observed in HUVEC co-cultured with HK2-depleted TPC (Supplementary fig. 6l–s). The above results showed that ROCK2, but not ROCK1, was the downstream effector of HK2 mediated pericycle’s blood vessel supporting function.

Accumulating evidence suggest that tumor cells can secrete paracrine factors to affect their surrounding stromal cell metabolism. To determine whether the difference in metabolic states between NPC and TPC was driven by tumor cells, we exposed NPC to conditioned medium from either NSCLC cell line A549 or HCC cancer cell line HepG2 and then used them for western blotting, glucose uptake, lactate concentration, and tube formation assays. Strikingly, exposure with the conditioned medium (CM) from cancer cells did not affect the glucose uptake in NSCLC/HCC-derived NPC (Supplementary fig. 6t), while it increased the expression of HK2, ROCK2, and p-MLC2 as well as lactate production in NPC as compared with untreated NPC (Fig. 5p and Supplementary fig. 6u). When these CM treated cells co-cultured with HUVEC in a tube formation assay, it significantly decreased total tube length, branch points, and number as compared to HUVEC co-cultured with untreated NPC (Fig. 5q–t).

Recent studies suggested that tumor-derived hyaluronan (HA) fragments up-regulated glycolytic gene expression (including HK2) to modulate the pro-tumor function of monocytes, which could be rescued by using a HA antagonist (Pep-1). Interestingly, our flow cytometry analysis indicated that NSCLC/HCC-derived NPC expressed the HA receptor CD44 on their cell surface (Supplementary fig. 6v). We next examine the role of tumor cell-derived HA fragments in regulating pericyte metabolism by exposing NSCLC/HCC-derived NPC with conditioned medium from A549/HepG2 cells in the presence of HA antagonist (Pep-1) or control peptide (Cpep). Strikingly, our results showed that blockade of HA fragments by a HA antagonist (Pep-1) inhibited the up-regulation of HK2 and ROCK2 in NPC after treated with conditioned medium from A549/HepG2 cells as compared with the group treated with control peptides, while it had no effect on ROCK1 expression (Supplementary fig. 6w). To confirm ROCK2 as the downstream target of HK2 mediated glycolysis in TPC, we therefore pre-treated NSCLC/HCC-derived TPC with ROCK inhibitor, which were then used for lactate concentration measurement and seahorse assays. As expected, inhibition of ROCK did not affect the lactate concentration, ECAR and OCR in TPC as compared to placebo treated group (Supplementary fig. 7a–e), while treatment of ROCK inhibitor had no effect on HK2 expression in TPC (Supplementary fig. 7f, g). Overall, these data suggested that tumor-derived paracrine signal induced pericycle-HK2-driven glycolysis to dysregulate pericycle’s blood vessel supporting role by up-regulating ROCK2-MLC2 mediated contractility.

Administration of HK2 inhibitor enhances blood vessel function, doxorubicin delivery, and efficacy against tumor growth, while reducing hypoxia without affecting blood vessel number and coverage. Previous studies from other laboratories and our
enhance chemotherapy delivery and efficacy against tumor growth and progression. However, these works mainly focused on targeting endothelial cells, but not on pericytes. We next tested whether combination treatment with HK2 inhibitor and chemotherapeutic agent doxorubicin enhanced its efficacy against lung and liver tumor growth. Strikingly, our results indicated that co-treatment with HK2 inhibitor and doxorubicin significantly inhibited the growth of subcutaneous lung (i.e. LLC, A549) or liver (i.e. mouse HCC cell line Hepa 1-6) tumors as compared to the group treated with either placebo, doxorubicin, or HK2 inhibitor alone (Fig. 6a–d and Supplementary fig. 8a–h). Notably, treatment with HK2 inhibitor or doxorubicin alone showed limited/modest effect on tumor growth as compared to
placebo treated group (Fig. 6a–d and Supplementary fig. 8a–h). Analysis of lung tumor vasculature showed that any treatment including 3-BP had no significant effect on total blood vessel number and pericyte coverage as compared with other treatment groups (Fig. 6c–g and Supplementary fig. 8i–k), while it significantly enhanced tumor blood vessel diameter, basement membrane collagen IV staining, and blood vessel perfusion (Fig. 6h–j and Supplementary fig. 8l, m). Moreover, mice injected intravenously with Hoechst dye showed a significant increase in intratumoral Hoechst dye levels, relative to blood vessel number, after treatment with HK2 inhibitor alone, and with a combination of doxorubicin when compared with placebo or single doxorubicin treated mice (Fig. 6k). No significant difference in the weight of LLC/A549 tumor-bearing mice during treatment was observed (Supplementary fig. 8n, o). Importantly, co-immunostaining analysis confirmed that LLC tumor pericytes expressed HK2 (Supplementary fig. 8p), while the expression of pericyte-p-MLC2 was reduced in LLC/A549 tumors after treated with HK2 inhibitor as compared with placebo (Supplementary fig. 8q). These results demonstrate that the pro-vascular functional effect of HK2 inhibitor was apparent in vivo, and corresponded with improved doxorubicin efficacy.

Since previous studies have shown a strong correlation between blood vessel perfusion/dilation and chemotherapeutic drug delivery30,31, we next sought to examine whether the increase in blood vessel perfusion and dilation was sufficient to improve doxorubicin delivery in our animal models. Ex vivo two photon imaging showed that LLC tumor blood vessel perfusion, vascular diameter, and doxorubicin delivery were all increased in mice treated combination group as compared with doxorubicin alone treated groups (Fig. 6l–n and Supplementary movie 1, 2), implying that HK2 inhibitor treatment induced MLC2-driven tumor vasculature remodeling. Further confocal microscopy analysis confirmed that the intratumor doxorubicin level was increased in mice treated with HK2 inhibitor and doxorubicin when compared to mice treated with doxorubicin alone (Supplementary fig. 8r). Importantly, Doppler ultrasound analysis of microbubble perfusion revealed that treatment with HK2 inhibitor enhanced LLC tumor blood flow and perfusion in mice as compared with the groups treated with either placebo or doxorubicin alone (Fig. 6o, p). Notably, the increase in these dynamic flow and perfusion readouts was observed across the whole tumor, including the tumor core (Fig. 6o, p). Since tumor hypoxia has been linked with poor prognosis in cancer patients30, we next examined the tumor expression of hypoxia indicator glucose transporter (Glut1) in all treatment groups, indicating that the HK2 inhibitor or HK2 inhibitor and doxorubicin combination treatment significantly reduced tumor hypoxia (Supplementary fig. 8s), suggesting that the improved blood vessel function reduced tumor hypoxia in mice treated with HK2 inhibitor. To further examine whether the observed synergistic inhibitory effect of HK2 inhibitor and doxorubicin on tumor growth in vivo was due to direct inhibition on tumor cells or through modulating TPC function, we treated LLC cells with either placebo, 3-BP, doxorubicin or 3-BP, and doxorubicin combination in vitro, indicating that co-administration of 3-BP and doxorubicin had no synergistic inhibitory effect on LLC tumor cell growth in vitro (Supplementary fig. 8t). To determine whether 3-BP or/and doxorubicin could affect tumor cells mediated pericyte’s glycolytic change, we examined the effect of conditioned medium harvested from cancer cells pre-treated with 3-BP or/and doxorubicin on pericyte-HK2 and ROCK2 expression. Both RT-PCR and western blot analysis indicated that the conditioned medium of 3-BP or/and DOX pre-treated tumor cells could still up-regulate HK2 and ROCK2 expression in NSCLC/HCC-derived NPC as compared to untreated NPC group (Supplementary fig. 9a–e). Altogether, these data suggested that the synergistic inhibitory effect of 3-BP and doxorubicin on tumor growth in vivo was through modulating TPC’s blood vessel supporting function.

To confirm the role of pericyte-HK2 in modulating tumor vasculature and subsequent drug delivery, we subcutaneously co-injected A549/MHCC-LM9 (i.e. human HCC cell line) cells with either stable HK2-depleted TPC, stable ROCK2 overexpressing HK2-depleted TPC or scramble transfected TPC at 10:1 ratio into nude mice and then treated the tumor-bearing mice with placebo or doxorubicin. Strikingly, administration of doxorubicin reduced tumor growth in the mice that were co-injected with A549/MHCC-LM9 cells and HK2-depleted cells as compared to the mice that were co-injected with A549/MHCC-LM9 cells and scramble transfected TPC (Supplementary fig. 10a–h), while co-injection of A549/MHCC-LM9 cells with ROCK2 overexpressing HK2-depleted pericytes reduced the enhanced doxorubicin efficacy against tumor growth observed in mice co-injected with A549/MHCC-LM9 cells and HK2-depleted cells (Supplementary fig. 10a–h). No significance difference in total blood vessel number and pericyte coverage was observed between groups (Supplementary fig. 10i–k). Interestingly, the tumor blood vessel diameter, basement membrane collagen IV staining, blood vessel perfusion, and intratumoral Hoechst were all increased in mice that were co-injected with A549 cells and HK2-depleted TPC as compared to mice co-injected with A549 cells and scramble transfected TPC, while the increased blood vessel functions observed were reduced in mice that were co-injected with A549
cells and ROCK2 overexpressing HK2-depleted TPC (Supplementary fig. 10l–o). Furthermore, the expression of pericyte-p-MLC2 was reduced in tumors arising from the mice that were co-injected with A549 cells and HK2-depleted TPC as compared to A549 cells co-injected with scramble transfected TPC, whilst its expression was increased in the tumors derived from co-injection of A549 cells and ROCK2 overexpressing HK2-depleted TPC (Supplementary fig. 10p). Importantly, ex vivo two photon imaging showed that the tumor blood vessel perfusion, vascular diameter, and doxorubicin delivery were all increased in the mice that were co-injected with A549 cells and HK2-depleted TPC as compared with the mice that were co-injected with A549 cells and scramble transfected TPC, while the increased blood vessel perfusion, vascular diameter, and doxorubicin delivery observed
Fig. 6 Administration of HK2 inhibitor remodels tumor vasculature to enhance blood flow, tumor perfusion, doxorubicin delivery, and efficacy against tumor growth. a C57/BL6 mice were subcutaneously injected with 1 × 10⁶ LLC cells and treated with placebo, 3-bromopyruvate (3-BP), doxorubicin (DOX), 3-BP, and DOX combination (n = 5 mice per group). Representative gross image of tumors from each treatment group is given. b Tumor growth curves showed that co-treatment of 3-BP and doxorubicin inhibited the growth of LLC tumors as compared with placebo, doxorubicin or 3-BP treated group. c, d Bar charts indicate the tumor weight and volume in each group. e–i Midline sections of LLC subcutaneous tumors were double immunostained for CD34 (green) and α-SMA/collagen IV (red) for blood vessel quantification, percentage of pericyte coverage, blood vessel diameter, and collagen IV intensity analysis. j LLC tumor bearing mice were intravenously injected with a FITC-PECAM antibody (green) to detect perfused blood vessels. k Bar charts show the relative area of Hoechst dye uptake into perivascular tumor cells to area of FITC-PECAM in each treatment group. l–n Ex vivo two photon microscopic imaging of FITC-PECAM antibody perfused LLC tumors after treated with DOX alone or 3-BP and DOX combination. Bar charts show the quantification of vessel diameter and doxorubicin in each group. Statistical tests were two-sided. a–p Doppler ultrasonography in live mice provides relative blood flow (peak enhancement, PE) and tumor perfusion (Wash-in perfusion index, WiPI) in subcutaneous LLC tumors. Representative ultrasound images. Bar charts, quantitation across the whole tumors, including tumor cores. Results are given as means ± SEM. b Two-way ANOVA. c, d, f–k, p One-way ANOVA. m, n Student’s t test. Scale bars in a, o represents 1 cm, e, j, k, l 50 μm, i, f 20 μm.

Fig. 7 Schematic diagram represents the role of pericyte-HK2-driven glycolysis in regulating tumor blood vessel abnormality via ROCK2-MLC2 mediated contractility. TPC up-regulates HK2-driven glycolysis to induce ROCK2-MLC2 mediated contractility, which in turn negatively regulates its blood vessel supporting role. Administration of HK2 inhibitor 3-BP induces MLC2-driven tumor vasculature remodeling to increase blood flow, tumor perfusion, blood vessel diameter as well as chemotherapeutic drug delivery and efficacy against tumor growth, while reducing tumor hypoxia without affecting blood vessel density and pericycle coverage.

was decreased in mice that were co-injected with A549 cells and ROCK2 overexpressing HK2-depleted TPC (Supplementary fig. 10q and Supplementary movie 3–5). Doppler ultrasound analysis of microbubble perfusion revealed that depletion of HK2 in TPC enhanced A549 tumor blood flow and perfusion in mice as compared with scramble control group (Supplementary fig. 10r), while overexpression of ROCK2 in HK2-depleted TPC reduced the enhanced tumor blood flow and perfusion observed (Supplementary fig. 10r). These results demonstrated the important role of pericyte-HK2-driven ROCK2-MLC2 mediated contractility in tumor vasculature remodeling and drug delivery.

Overall, our findings show that treatment with HK2 inhibitor or depletion of pericyte-HK2 by shRNA can enhance blood flow and perfusion leading to increased intratumoral drug delivery and efficacy against tumor growth, whilst decreasing tumor hypoxia without affecting blood vessel density and pericycle coverage (Fig. 7). These results indicate that targeting pericyte-HK2 expression/activity could be an effective treatment strategy for cancer patients.

Discussion

Our method successfully purified a population of pericytes from normal adjacent tissues and tumors derived from NSCLC or HCC patients. We showed an elevated HK2-driven glycolysis in TPC as compared with NPC, which contributed to its abnormal blood vessel supporting function by activating ROCK2-MLC2 mediated contractility. Depletion of HK2/ROCK2 by shRNA or inhibitors reduces TPC contractility to remodel vasculature, while combined treatment of HK2 inhibitor and doxorubicin enhanced the efficacy of chemotherapeutic drug against tumor growth in both lung and liver cancer animal models. This is an unexpected result and call for a reconsideration of our current strategy targeting pericytes for cancer treatment.

Pericytes, also known as contractile cells, play an important role to modulate angiogenesis, blood vessel diameter, blood flow, vascular perfusion, and integrity mainly via their contractile properties1,7,10,32. Interaction between pericyte-endothelial cell in the blood vessel wall is a fundamental process in regulating vascularization, vasoconstriction, vasodilation, and stabilization9,14, while defective interaction between two cell types causes human pathological conditions, including dementia syndrome and tumor angiogenesis1,12. Indeed, genetic knock-out of PDGFB or PDGFRβ both causes vascular dysfunction and perinatal death5. Due to the lack of an effective human pericyte isolation protocol, most of the in vitro pericyte studies were performed by using either bovine/mouse retinal pericytes or mouse brain pericytes1,18,33. Although pericytes can potentially differentiate into other cell types34, the retinal pericytes were neither isolated by FACS, nor fully characterized by pericyte marker expression or cultured in medium supplemented with pericyte growth factors in previous studies32,35. Furthermore, it remains debating about the mesenchymal stem cell property of pericytes in vivo36. Contrary to these studies, we developed a MACS method to enrich vascular cell population from the dissociated tissues and then subjected the cell population for subsequent FACS to purify pericytes, which were then cultured in commercially available medium dedicated for human pericytes. By
performing a series of characterization experiments, we confirmed the pericyte identity of our isolated cells derived from NSCLC and HCC, while there was no significant difference in basic function between NPC and TPC. In addition, our culturing method can maintain the pericyte identity of the isolated cells over passages.

We next sought to explore their roles in regulating vasculature in vitro and in vivo. Previously, McLroy et al.37, showed that exposure of immortalized HUVEC with bovine retina pericyte conditioned medium can inhibit its migration as compared with HUVEC exposed with serum-free medium or endothelial cell conditioned medium, while it also repressed the tube formation ability of HUVEC. On the other hand, Kumar et al.38 found that mesenchymoangioblast-derived pericytes promoted blood vessel formation and stability in vitro and in vivo. The difference amongst these studies may be due to different sources/species of pericytes being used in the tube formation assays. Nevertheless, our data indicated that TPC exerted a stronger repressive effect on the total tube length, branch points, and number as compared with NPC, while in vivo co-transplantation of HUVEC with TPC formed smaller and irregular shaped blood vessels with poor basement membrane structure as compared with HUVEC co-transplanted with NPC. In addition, our cell contraction assays showed that NSCLC/HCC-derived TPC displayed a stronger carbachol-induced cell contraction as compared with NPC, fibroblast, or HUVEC. Overall, our study showed that TPC had defective blood vessel supporting functions as compared with NPC.

Although the contribution of specific metabolic programs to the regulation of cell state/functions have been extensively studied in endothelial cells, the role of metabolic state in regulating pericyte function remains unclear. Transcriptional profiling study showed a consistent increase in glycolytic gene hexokinase 2 expression in NSCLC/HCC-derived TPC as compared with NPC. Since HK2 plays a key role in catalyzing the first step of the glycolytic pathway where glucose is phosphorylated to glucose-6-phosphate39, we demonstrated that the lactate production was increased in TPC as compared to NPC. Importantly, metabolic flux analysis indicated the relative abundance of lactate, but not the TCA cycle metabolites, was increased in TPC as compared to NPC. These findings were supported by the clinical observation that the pericyte expression of HK2 was up-regulated in tumors as compared with normal adjacent tissues, while high percentage of pericyte-HK2 positive blood vessels correlated with poor overall survival in NSCLC and HCC patients respectively. We next explored the role of pericyte-HK2 in regulating its blood vessel supporting function. Our results showed that pre-treatment with HK2 inhibitor enhanced the total tube length, branch points, and number in HUVEC co-cultured with TPC in a tube formation assay as compared with HUVEC co-cultured with placebo pre-treated TPC. In addition, HK2 inhibitor pre-treated NPC formed smaller and irregular shaped blood vessels with poor basement membrane structure as compared with NPC co-transplanted with NPC. While administration of HK2 inhibitor reduced ROCK2 and p-MLC2 expression in TPC as compared with placebo pre-treated TPC. Importantly, depletion of HK2 or ROCK2, but not ROCK1, enhanced the total tube length, branch point and numbers in HUVEC co-cultured with TPC in tube formation assays as compared with HUVEC co-cultured with scramble transfected TPC, while overexpression of ROCK2 in HK2-depleted TPC reduced the enhanced effect observed. These results suggest that pericyte-HK2-driven glycolysis contributes to vascular dysfunction via activation of the ROCK2–MLC2 mediated contractility.

We next sought to explore the molecular control of HK2 expression in TPC. Since previous work showed that cancer cells can affect surrounding stromal cell metabolism via paracrine signaling28, we therefore exposed NPC with conditioned medium from cancer cells, indicating that exposure of NPC with conditioned medium from cancer cells up-regulated HK2 and ROCK2 expression as compared to untreated NPC. Indeed, recent studies showed that tumor-derived paracrine signal up-regulates pericyte-HK2-driven glycolysis to dysregulate its blood vessel supporting role by up-regulating ROCK2 mediated pathway.

Cancer cells rely on glycolysis to meet their energy demand, which is known as the Warburg effect. Since HK2 is a key mediator of glycolysis, its inhibitor, 3-bromopyruvate, has been extensively tested as a single anticancer agent for treating pre-clinical tumor models41,42. Unfortunately, it has so far provided modest treatment outcomes in different types of cancers43. Indeed, we showed that treatment with HK2 inhibitor 3-bromopyruvate alone showed modest effect on subcutaneous lung or liver tumor growth, while co-administration of HK2 inhibitor and doxorubicin significantly inhibited the growth of the tumors by inducing MLC2-driven vasculature remodeling, while reducing tumor hypoxia without affecting tumor blood vessel density and pericyte coverage. By performing ex vivo two photon microscopy and Doppler ultrasound imaging, we proved that administration of 3-BP induced tumor vasculature remodeling to enhance blood flow, perfusion, and chemotherapy delivery. Interestingly, treatment with 3-BP and doxorubicin showed no synergistic inhibitory effect on LLC cell growth in vitro, while exposure of NPC with conditioned medium from 3-BP or and doxorubicin pre-treated tumor cells could still up-regulate HK2 and ROCK2 expression as compared with untreated NPC. Together with the fact that the efficacy of chemotherapy delivery has been shown to be linked with tumor blood vessel perfusion and diameter31, these findings suggest that the synergistic repressive effect of 3-BP inhibitor and doxorubicin on tumor growth in vivo relies on modulating pericyte's BV supporting role, but not through tumor cell inhibition. Additionally, the role of pericyte-HK2 in regulating MLC2-driven tumor vasculature remodeling and drug delivery was further illustrated.
Methods
demonstrate, using strategy distinct from anti-angiogenesis or blood vessel function and drug delivery and efficacy against lung and liver cancer growth.

Overall, our work develops a cellular model for studying human pericyte biology and shows that pericyte-HK2-driven glycogenesis induces tumor blood vessel abnormality by activating ROCK2-MLC2 mediated contractility. Importantly, our data demonstrate, using strategy distinct from anti-angiogenesis or targeting pericyte coverage, that a radical approach of modulating the metabolic status of pericytes to remodel tumor vasculature for enhancing chemotherapy efficacy in cancer treatment.

Methods

Clinical specimens. Human non-small-cell lung cancer and hepatocellular carcinoma paraffin-embedded samples and fresh tissue samples were obtained from Sun Yat-sen Memorial hospital with complete clinical data. The NSCLC patients and HCC patients who underwent curative freshly resected tissue enrolled for pericyte isolation were based on the following main criteria: (1) patients were absence of anticoagulation therapies prior to the operation. (2) No concurrent autoimmune disease, HIV, or syphilis. All human non-small lung cancer and hepatocellular carcinoma paraffin-embedded samples and fresh tissue samples were anonymously coded in accordance with local ethical guidelines (as requested by the Declaration of Helsinki) with written informed consent and a protocol approved by the review board of Sun Yat-sen Memorial hospital (Guangzhou, China).

Correlation study between gene expression and patient survival using the Kaplan–Meier plotter database. Both the RNA-seq data and clinical data including outcome and clinicopathological information of human NSCLC and HCC patients were obtained from the Kaplan–Meier plotter database (KM plotter). All the survival studies were done according to the web tool developer’s instruction (http://kmplot.com). The expression of genes of interest were selected from the KM plotter database, which were then used to plot Kaplan–Meier survival curves in order to determine the correlation between their expression and patient survival in NSCLC and HCC, using auto best-selected cutoff or median setting. The relationship between the ratio of ROCK2 to PDGFRβ/Cd44/ACTA2 expression and patient survival data was analyzed using the KM plotter database.

Cell culture. Human umbilical vein endothelial cells (HUVEC) were isolated and cultured in endothelial cell medium (ScienCell, 1001) according to the manufacturer’s instruction. Human primary dermal fibroblasts (HFF-1, SCSP-109) were purchased from National Infrastructure of Cell Line Resource and cultured in Dulbecco’s Modified Eagle Medium (DMEM) (GIBCO, C11995500BT), supplemented with 10% fetal bovine serum (FBS) (Biological Industries, 0-400-1A). Human normal adjacent tissue derived pericytes and tumor-derived pericytes were isolated as follow and cultured in pericyte medium (ScienCell, 1201). Peri-vascular mononuclear cells (PMBC) were isolated from buoyous coats derived from the blood of healthy donors by Ficoll density gradient. The human primary pulmonary artery smooth muscle cells (SMC, PCS-100-023) were purchased from ATCC and cultured in vascular cell basal medium (PCS-100-030, ATCC) supplemented with vascular smooth muscle growth kit components (PCS-100-042, ATCC). For HUVEC and pericytes culture, 0.1% gelatin (Sigma, 9000-70-8) containing 0.7% type I collagen (Sigma, C8676-1) was coating on plate or flask prior to seeing cells. Lewis lung carcinomas (LLC), A549, Hepa-1 and HepG2 cells (all from ATCC) as well as MEL-LC-LM9 cells (a gift from Professor Shimei Zhuang) were cultured in RPMI1640/DMEM medium supplemented with 10% fetal bovine serum and 1% penicillin and streptomycin.

Immunohistochemistry and immunofluorescent staining. Formalin-fixed, paraffin-embedded tissues were cut into 4-μm sections and subjected to Immunohistochemistry (IHC) and Immunofluorescence (IF) staining. Sections were stained in the IHC and IF staining included: rabbit mAb against human CD34 (ZSGB-BIO, ZM-564907). To apply the antibodies which were originated from the same host, Alexa Fluor® 488 Tyramide Reagent (Invitrogen™, A21082, 1:500) were used to detect human antigens, and the sections were counterstained with 4′-6′-diamidino-2-phenylindole (DAPI, BD PharmingenTM, 49407). To apply the antibodies which were originated from the same host, Alexa Fluor® 488 Tyramide Reagent (Invitrogen™, B40953) was used to remove the antibody but preserving the fluorescent signal.

Coverslip cell staining. Isolated pericytes (NPC, TPC), normal fibroblasts, HUVEC, and smooth muscle cells were first solidified onto coverslips in 12-well plates overnight, which were then fixed with 4% formaldehyde in PBS for 10 min at room temperature. Cells were blocked, permeabilized, and incubated with rabbit mAb against human FAP (E1V9V) (CST, 61012, 1:100), mouse mAb against human desmin (Merck Millipore, MAB3430, 1:100), or indirect mAbs. Alexa Fluor–conjugated secondary antibodies and DAPI were used as above.

RNA extraction and real-time PCR. Total RNA was extracted from the indicative cells using Trizol reagent (Invitrogen, AM9738). Aliquots (2 μg) of total RNA were reverse transcribed into cDNA using 5x All-In-One RT MasterMix (Abm, 4949) followed by quantitative real-time-PCR (qPCR) using HieffTM qPCR SYBR®
Green Master Mix (No RoX) (YEASEN, N12015E08). All reactions were run in triplicate and performed on a Roche Light Cycler 480 Instrument II (Roche). The cycle threshold (Ct) values did not differ by >0.2 among the triplicate and performed on a Roche Light Cycler 480 Instrument II (Roche) article.

**Cell proliferation assay, doubling time measurement, and drug testing.** CCK-8 proliferation assay was done according to the Jiangsu KeyGENBioTECH Corp., Ltd. Briefly, 1500 HUVECs per well were seeded onto a 96-well plate and the cell proliferation was detected at indicated time points. For the conditioned medium experiment, 2000 HUVECs per well were seeded onto a 96-well plate. After 24 h, the cells were incubated with fresh endothelial cell medium containing conditioned medium from NPC/TPC (7 to 3 ratio). CCK-8 reagent was added to the culturing medium 4 h prior to each time point. Doubling time for pericytes at each passage was calculated by the formula: \( T (\text{hours}) = \ln (2) \times \text{duration of culture (hours)}/\ln (\text{output cell number/input cell number}) \). For in vitro drug testing, 1500 LLC cells per well were first plated on a 96-well plate overnight and treated with either 200 μM 3-BP, 2.5 μM doxorubicin or 3-BP, and doxorubicin combination for 48 h.

**Wound healing assay.** Pericytes or HUVECs were seeded on 24-well plates at a density that reached 90% confluence after 24 h. A 200 μl pipette tip was used to scratch a straight line across the wells. Afterwards, the HUVECs were incubated with fresh EC medium containing conditioned medium derived from NPC/TPC (7 to 3 ratio). Images were taken under bright field using a phase contrast microscope (Nikon, Ni-U) at indicated time points. The cell migration was quantified by calculating the migrated area in each group.

**Trans-well migration assay.** An endothelial cell dependent recruitment model was developed by using a trans-well system. Briefly, 4×10^4 HUVECs were seeded onto a 24-well plate overnight, while 5×10^5 NPC or TPC were seeded on the upper chamber of a trans-well insert in a 24-well plate overnight. Afterwards, the insert was then transferred to the 24-well plate containing HUVECs for 6 h. The migrated cells on the bottom part of the insert were then stained with crystal violet and counted by using a phase contrast microscope. 

**Senescence detection.** NPC and TPC at passage 3 and 12 were measured by using senescent cells histochemical staining kit (Sigma, CS0030). The pictures were captured using a fluorescence microscope (Nikon, Ni-U).

**Cell viability/apoptosis assay.** Primary pericytes were harvested at 90% confluence density. Then the percentage of viable/apoptotic cells were evaluated by using FITC Annexin V apoptosis detection kit with PI (BioLegend, 640991). 

**Cell contraction assay.** Cell contraction assay was done by adding 1 mM Car-bachol (Sigma, PRH1511) into 80% confluent monolayer of NPC, TPC, HUVEC, or fibroblast for 30 min. For the inhibition assay, TPC were pre-treated with 200 μM 3-BP or 0.4 μM GSK429286A inhibitors at room temperature for 15 min. The inhibitors were then removed by centrifuging. For the conditioned medium experiment, NPC were first pre-treated with conditioned medium from A549 or HepG2 cells for 6 h. Then these cells were then co-cultured with HUVEC as described above and were taken by using a fluorescence microscope 24 h after co-culturing (Nikon, Ni-U). The total tube length, branch points, and number were measured in each group.

**Matrix plug assay.** Matrix plug assay was done by mixing HUVEC with either NPC, untreated TPC, TPC pre-treated with placebo/HK2 inhibitor 3-bromopyruvate (3-BP), fibroblast, smooth muscle cells (2:1), which were then cultured with conditioned medium derived from A549/TPC for 12 h. Then the plugs were performed as above. The cellular glucose level was detected by using a glucose assay kit (Cytyc, 200-00) in conditioned medium derived from tumor cells (7 to 3 ratio) for 12 h. Then these cells were performed as above. The cellular glucose level was detected by using a glucose assay cell-based assay kit (Cytox, 600470) according to the manufacturer’s instruction. 

**Glucose uptake assay.** In total, 3×10^4 NPC or TPC/well in 100 μl culture medium were seeded on a 96-well plate and grown overnight. The next day, the cells were treated with or without the indicative inhibitors for 15 min. Then cells were washed and cultured with glucose-free medium for 4 h. In all, 20 min before flow cytometric assay, 100 μg/ml 2-NBDG was added into the culturing medium. In some experiments, NPC were incubated with fresh PC medium containing conditioned medium derived from tumor cells (7 to 3 ratio) for 12 h. Then these cells were performed as above. The cellular glucose level was detected by using Glucose Uptake Cell-Based Assay Kit (Cayman, 600470) according to the manufacturer’s instruction.

**l-lactate detection assay.** In total, 3000 NPC or TPC were plated into a 96-well plate and incubated with 120 μl of percyte medium overnight. The next day, the cells were treated with or without the indicative inhibitors for 15 min at room temperature. The cells were then washed and cultured with serum free medium for 12 h. In some experiments, NPC were incubated with fresh PC medium containing conditioned medium derived from tumor cells (7 to 3 ratio) for 12 h. In these cells were performed as above. The l-lactate concentration of culturing medium was detected by using a glycolysis cell-based assay Kit (Cayman, 600450) according to the manufacturer’s instruction.

**Proteomics analysis.** For the sample preparation, NPC and TPC were cultured in a 100-mm culture plate at 37 °C in a 5% CO2 waterlogged atmosphere using percyte medium. Confluent cells were harvested by scraping with 800 μl of RIPA buffer. The pellets were collected and resuspended in 200 μl of lysis buffer. A 20 μl sample was loaded onto a 20% PAGE, and the gel was stained with Coomassie Blue R-250. After destaining, the gel was scanned using a GS-700 Imaging Densitometer (Bio-Rad). The 2-D DIGE analysis was performed using the same sample set. The images were analyzed using the ImageMaster 2D Platinum software. The average of 3 replicates was considered for the final results. The proteins were excised from the gels and subjected to in-gel digestion with trypsin. The resulting peptides were extracted and submitted for nanoLC-MS/MS analysis. The resulting MS/MS spectra were searched against the SwissProt database using MS-Frag software. The protein identification was considered significant when the peptide and protein scores were above the threshold of 0.05.
digested in-solution with trypsin. Briefly, the samples were initially precipitated with acetone, then resuspended with 50 Mm urea buffer, reduced with dithiothreitol (1.5 h, 2 ° C), alkylated with iodoacetamide (1 h, 25 ° C for 40 min), and then were diluted to 60 mM urea for overnight digestion with trypsin (Sequencing Grade Modified Trypsin, V5111, Promega) at 37 °C. Samples were then diluted 2-fold again and digested overnight with trypsin at 37 °C. Tryptic peptides were desalted using a HLB column (Oasis HLB, 160000383, Waters), evaporated to dryness. Then, 300 μg of dried pericyte protein digest were dissolved with 10 μl of 0.1% formic acid with H2O47.

The mass spectrometer was operated in positive ionization mode with an easy nano flex (Nanospray ionization, NSI) with spray voltage set at 2.3 kV and source temperature at 100 °C. The instrument was operated in data-dependent acquisition mode, with full MS scans over a mass range of m/z 300–2000. In each cycle of data-dependent acquisition analysis, following each scan survey, the most intense ions above a threshold ion count of 2.2e4 were selected for fragmentation at normalized collision energy of 30% (HCD). The number of selected precursor ions for fragmentation was determined by the “Top Speed” acquisition algorithm and a dynamic exclusion of 60 s. Fragment ion spectra were acquired in the linear ion trap (IT) or the Orbitrap (OT, 70 K resolution) depending on the method, with an AGC of 4e5 and a maximum injection time of 50 ms for ion trap MS2 detection, and an AGC of 5e5 and a maximum injection time of 50 ms for Orbitrap MS2 detection. All data were searched with Xcalibur software v4.0. Peptide ions were injected into a nano-UPLC system (EASY-nLC 1200 liquid chromatograph) equipped with a 50-cm C18 column (EASY-Spray; 75-75-μm (PepMap RSLC C18, 3μm coating) and connected to a Thermo ISQ MS. GC/MS was performed using a Thermo 1300 equipped with a DB-35MS capillary column (30 m, 0.25 mm i.d., 0.25 μm coating) and hyphenated to an Thermo Scientific Orbitrap Fusion Tribrid Mass Spectrometer. Chromatographic gradient lengths 180 min were tested for peptide separation. Mobile phase: A, H2O: HCOOH (100: 0.01); B, ACN: H2O: HCOOH (100:0.01:0.01). Gradient elution: 0–2 min, 5% B; 2–5 min, 10% B; 5–152 min, 10%; 152–172 min, 28–38%B; 172–175 min, 38–100%B, followed by equilibration of the column with mobile phase A as described previously48. Proteome Discoverer software suite (v2.3, Thermo Fisher Scientific) was used for peptide identification and quantification. The data was searched against the Swiss-Prot human database. At the MS1 level, a precursor ion mass tolerance of 2 ppm was used, and up to three missed cleavages were allowed. The fragment ion mass tolerance was set to 20 ppm for the Orbitrap MS2 detection methods and 0.5 Da for the linear ion trap MS2 detection methods. Oxidation of methionine, Cys(terminal) and Cys(N-term) oxidation, and oxidation at N-term of peptide sequence were allowed. Oxidation of methionine, carbamidomethylation of Cys(terminal) and Cys(N-term) of peptide sequence were allowed. For peptide identiﬁcation, Xcalibur software v4.046 was used. For the metabolic study, A549/HepG2 cells were ﬁrst pre-treated with either placebo, 200 μM 3BP, 2.5 μM doxorubicin or 3BP and doxorubicin combination for 8 h and then washed off and incubated with fresh culturing medium for 16 h. The conditioned medium was then harvested, which were then mixed with fresh pericyte medium (7 to 3 ratio) and subsequently incubated with NSCLC/HCC-derived NPC for 24 h before harvesting for RT-PCR and western blotting analysis.

**Metabolite extraction and derivatization.** Tracer chemically deﬁned medium was ﬁrst prepared for metabolic ﬂux analysis, which consisted of glucose-free DMEM medium (Gibco, 11960025) supplemented with [13 C] glucose (Cambridge Isotope Laboratories, CLM-1396-1). Pericytes were seeded onto six-well plate and incubated with the tracer chemically deﬁned medium at 37 °C for 4 h. Afterwards, the metabolite extraction was done as described previously50. Polar metabolites (including organic acids and sugars) were extracted using methanol/water/chloroform. Briefly, spent medium was removed, and cells were rinsed with 0.9% (w/v) saline, and 250 μl of ~80 °C methanol was added to quench metabolism. In total,100 μl of ice-cold water containing 1 mg norvaline internal standard was added to each well. Both solution and cells were collected via scraping. Cell lysates were then freeze-dried, and 200 μl of 30 ° C chloroform sample tubes, vortexing and centrifugation, the top aqueous layer (polar metabolites) and bottom organic layer were removed and dried under air. Derivatization of polar metabolites was performed using the gerstel multi-purpose sampler (MPS 2XL). Dried polar metabolites were dissolved in 20 μl of 2% (w/v) methoxamine hydrochloride (MP Biomedicals) in pyridine and held at 37 °C for 60 min. Subsequent conversion to their tert-butylidimethylsilyl (tBDMS) derivatives for organic acids was accomplished by adding 30 μl of N-methyl-N-(tert-butyldimethylsilyl) triﬁluoroacetamide supplemented with 1% tert-butylidimethylchlorosilane (Regis Technologies) and incubating at 37 °C for 30 min.

**Gas chromatography/mass spectrometry analysis.** GC/MS analysis was performed using a Thermo 1300 equipped with a DB-35MS capillary column (30 m, 0.25 mm i.d., 0.25 μm coating) and connected to a Thermo IQ MS. GC/MS was operated in positive ionization mode. Sample injection was performed in splitless mode at 270 °C, using helium as the carrier gas at a flow rate of 1.2 mL/min. For measurement of organic and amino acids, the GC oven temperature was held at 80 °C for 2 min and increased to 255 °C at 3.5 °C/min, and then ramped to 330 °C at 15 °C/min for a total run time of ~60 mins with 5 mins of delay time. The MS source was held at 300 °C, and the detector was run in scanning mode, using ion abundance in the range of 100–400 m/z.

**Metabolite quantification and isotopomer spectral analysis.** All results were based on chromatographic peak areas that received from targeted data evaluation in the Thermo Scientific (TraceFinder) and analyzed as described previously50. Slight smoothing as implemented in peak detection by Tracefinder (5 points) was applied to improve automated peak integration. Mole percent enrichment (MPE) of isotopes was calculated as the percent of all atoms within the metabolite pool that were labeled.

**Measurement of ECAR and OCR by Seahorse analysis.** Seahorse assays were performed according to the manufacturer’s instructions. Briefly, 2,4 × 10^5 cells/mL. NPC or TPC were seeded onto a 96-well Agilent Seahorse XF culture microplates (Agilent) and incubated with human pericyte medium. For the ROCK inhibitor experiments, NSCLC/HCC-derived TPC was first pre-treated with 0.4 μM ROCK inhibitor GSK429286A for 1 h and then washed off before carrying on the ECAR and OCR measurement. A standard manufacturer-recommended two-step seeding procedure was utilized. After achieving cell adherence, the microplates were placed overnight in a 37 °C, 5% CO2 incubator. Afterwards, the incubation medium for each well was aspirated, while the adherent cells were washed and then incubated with the Seahorse XF Base Medium (w/o phenol red) supplemented with 1 M glutamine (final concentration of 1 mM), 1 M glucose (final concentration of 10 mM) of OCR, 1 M glutamine (final concentration of 1 mM) of ECAR (adjusted the pH to 7.4 with 1.0 N NaOH) at a 37 °C, non-CO2 incubator for 1 h before being transferred onto the microplate stage of a Agilent Seahorse XF96 ﬂux analyzer (Agilent). ECAR and OCR readings were taken using a 2-mins mix, 1-min wait, and 2-mins read cycling protocol. Various compounds were injected to alter the metabolic environment or to help determine the effects of an altered metabolic environment. These compounds included oligomycin, carbonyl cyanide-3-(trifluoromethoxy)phenylhydrazone (FCCP) and rotenone/antimycin of OCR, and glucose, 2-deoxyglucose (2-DG), and oligomycin of ECAR. OCR-ECAR data are presented as absolute rates or as percent changes from a stable baseline. For comparisons of data points between cells maintained under different conditions, group means were compared by Student’s t-tests, with P-values <0.05 considered significant. All wells were normalized with protein concentration by using BCA assay.

**Mouse tumor models.** In all, 4–6-weeks-old C57/BLK6 mice or nude mice were subcutaneously injected with either 1 × 10^5 LLC cells, 3 × 10^5 A549 cells, or Hepa 1–6 cells into the flank for subcutaneous tumor growth. For the co-injection experiments, the nude mice were co-injected with 3 × 10^5 A549/HMHC-LM9 cells and either 3 × 10^5 HK2-depleted TPC, scrambled transfect TPC or ROCK2 overexpressing HK2-depleted TPC subcutaneously (101:1 ratio). Tumor growth was measured by using callipers every 3 days. Tumor volume (V) was calculated: V = (length × width^2) / 2. Once the tumor size reached 100 mm^3, the tumor-bearing mice were treated with either placebo, HK2 inhibitor 3-BP (4 mg/kg, intraperitoneal Injection (IP)), and HK2 inhibitor 3-BP (4 mg/kg, intravenous injection (IV)), and HK2 inhibitor (3-BP) and doxorubicin combination via IP/IV injection every 3 days for up to 15–18 days. When the treatment finished, animals were culled, tumor excised and either fixed in 4% formaldehyde in PBS overnight, or snap frozen in liquid nitrogen for subsequent immunostaining analysis. All animal procedures were approved by the Institutional Animal Care and Use Committee (IACUC) of Sun Yat-sen University.

**Blood vessel density and pericyte coverage.** The number of CD34-positive blood vessels per field in tumor sections from each treatment group was counted. For pericyte coverage, the percentage of a-SMA and CD34-double positive blood vessels over the total number of CD34-positive blood vessels was counted.
**Doxorubicin quantification by confocal microscopy.** For doxorubicin quantification, tumor-bearing mice were injected with 4 mg/kg doxorubicin 1 h prior to culling. The tumors were harvested and snap-frozen and the tumor sections were fixed and imaged by using confocal microscopy. The fluorescence signal of DOX from five different fields was analyzed by ImageJ software.

**Ex vivo two photon microscopic imaging.** For the quantification of doxorubicin and vascular perfusion, LLC tumor-bearing mice were injected with 4 mg/kg doxorubicin 1 h and 20 μg of rat monoclonal anti-FITC-PECAM antibody 10 min prior to culling, respectively. Tumor were harvested and then imaged by using the 25 x water immersion objective (Olympus, NA 1.05) of a confocal microscope (Olympus) equipped with a tunable pulsed chameleon infrared multiphoton laser (MaiTai eHODS and INSIGHT X3) and PMT (GaAsp). Z-stack images were taken up to 100 μm with 191 slices.

**Ultrasound analysis of blood flow and tumor perfusion.** High-resolution ultrasound imaging of LLC tumors using the Vevo 2100 system with a MS250 transducer was performed according to the manufacturer’s instruction (Visualsonics). For blood flow and tumor perfusion measurement, contrast-enhanced ultrasound imaging was carried out according to manufacturer’s instructions (Vevo 2100 system). Briefly, LLC tumor bearing mice were administered with 100 μL of bolus Vevo Micromarker suspension (VisualSonics) via tail vein using a tail vein Micromarker™ catheter. The ultrasound images were processed and analyzed by using VevoQTM contrast quantification software according to the manufacturer’s instruction.

**Quantification and statistical analysis.** Student’s t test, Log-rank (Mantel-Cox) test and one/two-way ANOVA were performed for the statistical analysis and the details can be found in the figure legends: test used, exact value of n and dispersion measures. Statistical analysis was performed using GraphPad Prism.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

The mass spectrometry proteomics data generated in this study has been deposited to the Proteome X consortium via the PRIDE partner repository with the dataset identifier PXD020603. The source data underlying Figs. 1c, f, k, l, 2c – 18 and Proteome X consortium via the PRIDE partner repository with the dataset identifier PXD020603. The mass spectrometry proteomics data generated in this study has been deposited to the PRIDE partner repository with the dataset identifier PXD020603. The source data underlying Figs. 1c, f, k, l, 2c – 18 and Proteome X consortium via the PRIDE partner repository with the dataset identifier PXD020603. The mass spectrometry proteomics data generated in this study has been deposited to the PRIDE partner repository with the dataset identifier PXD020603. The source data underlying Figs. 1c, f, k, l, 2c – 18 and Proteome X consortium via the PRIDE partner repository with the dataset identifier PXD020603.

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Author contributions
Y-M.M., X.J. and X.Z. carried out majority of the experiments and contributed equally to the paper; Q.M., S.W., Y.C., X.K., X.Q., L.S. and C.H. assisted with the experiments; M.W. and C.L. provided clinical samples; P.-P.W conceived the study, supervised the research, and wrote the manuscript.

Competing interests
The authors declare no competing interests.

Additional information

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