Increased Vascular Permeability in the Bone Marrow Microenvironment Contributes to Disease Progression and Drug Response in Acute Myeloid Leukemia

Graphical Abstract

Highlights

- Functional imaging of the BM vascular niche using intravital 2P microscopy
- Increased vascular permeability is caused by robust molecular alterations in ECs
- Persistent high NO after chemotherapy is associated with poor prognosis in patients
- Inhibition of EC-derived NO normalizes vasculature and improve response to Ara.C

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In Brief

Passaro et al. show that acute myeloid leukemia, mainly via nitric oxide (NO), causes bone marrow vascular abnormalities and that failure to restore normal vasculature after induction chemotherapy is associated with a poor prognosis. Importantly, inhibition of NO production improves treatment response in vivo.

Data Resources

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Increased Vascular Permeability in the Bone Marrow Microenvironment Contributes to Disease Progression and Drug Response in Acute Myeloid Leukemia

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SUMMARY

The biological and clinical behaviors of hematological malignancies can be influenced by the active crosstalk with an altered bone marrow (BM) microenvironment. In the present study, we provide a detailed picture of the BM vasculature in acute myeloid leukemia using intravitral two-photon microscopy. We found several abnormalities in the vascular architecture and function in patient-derived xenografts (PDX), such as vascular leakiness and increased hypoxia. Transcriptomic analysis in endothelial cells identified nitric oxide (NO) as major mediator of this phenotype in PDX and in patient-derived biopsies. Moreover, induction chemotherapy failing to restore normal vasculature was associated with a poor prognosis. Inhibition of NO production reduced vascular permeability, preserved normal hematopoietic stem cell function, and improved treatment response in PDX.

INTRODUCTION

Acute myeloid leukemia (AML) is the most common acute leukemia in adults (Sant et al., 2010). AML is a hematological malignancy arising from the occurrence of genetic mutations in hematopoietic progenitors, which cause a blockage in the maturation and an uncontrolled growth of leukemic blasts in the bone marrow (BM). While the clinical presentation of AML is quite uniform, it is a highly heterogeneous disease at the molecular level. There is a consistent effort aimed to characterize each molecular subgroup and design the best therapeutic strategy (Grimwade et al., 2010). Besides a few cases (Wang and Chen, 2008), the common clinical practice remains induction therapy with cytarabine (AraC) (Buchner et al., 2012; Mayer et al., 1994). However, therapy resistance and relapse remain a major clinical challenge.

Another approach aims to identify and target common features within this complex disease. One possibility could be the BM microenvironment, which is the site where leukemic cells arise, expand, and eventually develop resistance to therapy. The BM microenvironment is constituted by different types of cells, with a predominant vascular component responsible for nutrient and metabolite turnover, the ingress and egress of different cells, and the regulation of normal hematopoietic...
Table 1. Patient-Derived Samples Used for Xenograft Studies

| AML ID | Cytogenetics | Cyto Risk | NPM | FLT3 |
|--------|--------------|-----------|-----|------|
| AML1   | ND\textsuperscript{a} | ND        | mutant | ND   |
| AML2   | t(9,6) \textsuperscript{b} | intermediate | ND | ND   |
| AML3   | normal | intermediate | WT | ITD\textsuperscript{c} |
| AML4   | 11q23/MLL | ND | ND | ND   |
| AML5   | complex | poor | WT | ND   |
| AML6   | normal | intermediate | mutant | ITD |
| AML7   | normal | intermediate | mutant | ITD |
| AML8   | Del 7q+8 + 1 | poor | WT | WT   |
| AML9   | normal | intermediate | mutant | ITD |
| AML10  | complex | poor | ND | ND   |

\textsuperscript{a}Not determined. \textsuperscript{b}Internal tandem duplication.

stem cell (HSC) function (Boulais and Frenette, 2015; Mendez-Ferrer et al., 2015). The introduction and development of intravital two-photon (2P) microscopy has allowed direct, high-resolution, dynamic imaging of the calvarium BM (Colmone et al., 2008; Foster et al., 2015; Lassailly et al., 2013; Lo Celso et al., 2009; Sipkins et al., 2005), providing a powerful tool for the functional characterization of the BM microenvironment.

Recent findings indicate that myeloid malignancies also affect the function of the BM niche, pointing to the existence of an active crosstalk between leukemic cells and the microenvironment (Dong et al., 2016; Frisch et al., 2012; Hanoun et al., 2014; Kim et al., 2008; Kode et al., 2013; Raaijmakers et al., 2010; Schepers et al., 2013; Schmidt et al., 2011; Zhang et al., 2012). Similar to what has been observed in solid cancers, AML has been associated with an increase in microvascular density (MVD) and production of pro-angiogenic factors, notably vascular endothelial growth factor (VEGF) (Chand et al., 2016; Hussong et al., 2000; Kampen et al., 2013). However, whether increased MVD and VEGF constituted a prognostic factor for AML treatment response remains unclear (Aref et al., 2005; Pedro et al., 2000; Reddy and Moreb, 2000).

Moreover, despite promising preclinical studies (Zhu et al., 2003), clinical trials incorporating anti-VEGF inhibitors have not produced encouraging results (Fiedler et al., 2003, 2005; Giles et al., 2006; Mattison et al., 2015; Roboz et al., 2006; Zahiragic et al., 2007), suggesting that targeting pro-angiogenic cytokines may not be the best strategy for disrupting the crosstalk between AML and the vascular niche. A thorough analysis of the status of the vasculature in the BM is required to dissect the complexity of the vascular phenotype associated with AML. Thus, we aim to study the in vivo picture of the abnormalities associated with the BM vasculature induced by AML engraftment and unravel common pathologic processes, which could represent potential targets in AML.

RESULTS

AML Engraftment Alters Vascular Architecture and Function

To provide a detailed picture of the BM vasculature in AML, we studied the status of the vascular niche in human AML patient-derived xenografts (PDX). Recipient mice were left unconditioned, given the toxic effect derived from the irradiation or myelosuppressive treatment on the vasculature (Hooper et al., 2009; Kopp et al., 2005; Shirio and Tavassoli, 1991; and data not shown). We observed an expansion of the endothelial compartment among the non-hematopoietic stroma upon human AML engraftment (Figure 1A and Table 1). Importantly, this effect was specific to AML, as no such expansion was observed in mice engrafted with normal human hematopoietic stem/progenitor cells (HSPCs) derived from umbilical cord blood (CB) (Figure 1A). Moreover, the percentage of endothelial cells (ECs) was positively correlated to the leukemic engraftment of human AML cell lines and patient-derived samples (Figure S1A), suggesting a gradual pathologic process. Not only did the percentage of ECs increase, but there was also a real expansion of the endothelial compartment in terms of absolute number, specifically upon human AML engraftment (Figure 1B). We also observed an increased MVD, as shown by the higher number of vessel sprouts quantified by immunofluorescence (Figures S1B and S1C), similar to what is observed in patient-derived tumors (Chand et al., 2016; Pedro et al., 2000). The existence of...

Figure 1. AML-Derived Alteration of the BM Vascular Architecture and Function

(A) Quantification of CD31\textsuperscript{+} ECs in the BM (shown as percentage of CD45\textsuperscript{+} BM cells) of non-transplanted mice (ctrl) and mice transplanted with HSPCs (CB) or AML patient-derived samples, as depicted. Each dot represents an individual mouse. Ctrl, n = 22; CB, n = 29; AML patients (AML1, 2, 3, 5, 6, 7, 8, 9), n = 30. Data are shown as mean ± SEM.

(B) Absolute number of CD31\textsuperscript{+} ECs in the BM (2 femurs, 2 tibias, and 2 iliac crests) of non-transplanted mice (ctrl) and mice transplanted with CB-derived HSPCs (CB) or AML patient-derived samples, as depicted. Each dot represents an individual mouse. Ctrl, n = 17; CB, n = 26; AML patients (AML1, 2, 6, 7, 8, 9), n = 23. Data are shown as mean ± SEM.

(C) Frequency of Sca-1\textsuperscript{high} and Sca-1\textsuperscript{low} CD31\textsuperscript{+} cells in the BM of non-transplanted mice (ctrl) and mice transplanted with AML patient-derived samples. Ctrl, n = 19; AML patients (AML2, 3, 6, 8, 9), n = 16. Data are shown as mean ± SEM.

(D) Representative 3D reconstruction of BM vasculature of the calvarium BM of non-transplanted mice (ctrl) and mice transplanted with CB-derived HSPCs (CB) or AML patient-derived samples, as depicted, imaged via 2P microscopy 1 min after injection of 655-conjugated NT-Qtracker as vessel-pooling agent. White arrows pointing at sinusoids. Data are representative of triplicates in 4 independent experiments. Scale bars represent 70 μm.

(E) Quantification of vascular mean diameter in the calvarium BM of non-transplanted mice (ctrl) and mice transplanted with CB-derived HSPCs (CB) or human AML-derived samples, as depicted, using IMARIS filament tool. Dots represent the diameter of vascular fragments in the z stack of the calvarium of at least 3 mice per group. Ctrl, n = 235; CB, n = 232; HL60, n = 91; ML1, n = 148; U937, n = 85; AML6, n = 130; AML8, n = 94. Red lines represent the mean ± SEM.

(F) Representative 3D reconstruction of BM hypoxia imaged via intravital microscopy using the HypoXis probe together with vasculature (dextran) and Nestin\textsuperscript{+} cells in non-transplanted mice or mice transplanted with AML6 patient-derived cells, as depicted. Scale bars represent 50 μm.

(G) Distribution and relative frequency of vessel distances to hypoxic areas in the BM of non-transplanted mice (ctrl) or mice transplanted with AML6 patient-derived cells, as depicted.
specific endothelial cell markers defining distinct BM vascular niches has recently been highlighted (Itkin et al., 2016). We thus analyzed the expression of these markers in the context of AML disease in PDX. We observed a significant loss of ECs associated with sinusoids (CD31+ Sca1low) as well as an increased number of ECs associated with arterioles (CD31+ Sca1high) (Figures 1C and S1D). We next analyzed the architecture of the BM vasculature by 2P microscopy using a vessel-pooling agent to visualize the vascular tree in the calvarium BM. Although vascular architecture appeared highly heterogeneous among different PDX (Figures 1D and S1E), we noticed some common abnormalities. First, the regularity of sinusoidal structures, which are preserved with normal human engraftment, was lost in human AML xenografts (Figure 1D, white arrows pointing at sinusoids). Second, the mean vascular diameter of vessels was reduced (Figure 1E), a pathologic phenotype previously reported in tumor angiogenesis as a result of solid stress applied to vessels by overgrowing tumor cells (Padera et al., 2004; Stylianopoulos and Jain, 2013). Vessel compression was also highlighted by H&E staining in long bones (Figure S1F, dashed circles indicating vessel lumen). To study BM perfusion, we injected isolectin B4 (IB4), a pan-endothelial marker (Lassally et al., 2013), and analyzed its distribution on the BM vasculature by 2P microscopy. In control mice, we observed a homogeneous IB4 perfusion rate, allowing the visualization of ECs surrounding the arteriolar and sinusoidal vasculature (Figure S1G, ctrl). In contrast, we observed the presence of many poorly perfused areas in the BM of AML xenografts (Figure S1G). We next tested whether AML engraftment also affected BM oxygenation, by measuring the BM hypoxia. While in non-transplanted mice we observed a heterogeneous staining with Hypoxyprobe, indicating a physiological spread distribution of hypoxic areas, human AML engraftment increased the hypoxia homogeneously throughout the bones (Figure S1H). Quantification of Hypoxyprobe staining in BM cells by flow cytometry confirmed the significant increase of BM hypoxia upon human AML engraftment compared with normal human engraftment (Figure S1I). Of note, at early stage of engraftment hypoxia was localized in close proximity to AML cells (Figures S1J–S1M), whereas at high engraftment the BM was overall hypoxic (Figures S1N–S1P). We next visualized the hypoxic state of the BM via intravital microscopy using HypoxiSense probe (Bao et al., 2012). Similar to what was observed with Hypoxyprobe, we observed increased hypoxia in the BM upon AML engraftment with this alternative method (Figures 1F, S1Q, and S1R). To evaluate the hypoxic state of the vasculature, we measured the distance of each vessel to hypoxic areas. Whereas in control BM this distance was widely distributed, in the presence of AML most of the vessels were close to hypoxic areas (Figures 1G and S1S). Together these results showed that AML is associated with a structurally and functionally abnormal vasculature in the BM.

**AML Engraftment Increases Vascular Permeability in the BM**

To study the vascular barrier function in the BM of human AML xenografts, we used intravital 2P microscopy to image the calvarium of unconditioned NSG mice engrafted with different human AML cell lines or patient-derived samples. Human AML cell lines were engineered to express GFP and luciferase to facilitate the monitoring of BM engraftment via bioluminescence (Figure S2A) and BM aspirate (Figure S2B). Prior to imaging, tetramethylrhodamine isothiocyanate (TRITC)-dextran was injected intravenously to label the vasculature (Egawa et al., 2013; Fukumura and Jain, 2008) (Figure 2A). Strikingly, we observed an important leakage of the dextran outside the vasculature in the BM of mice engrafted with human AML cell lines compared with control non-transplanted mice (Figures 2B and S2C). This observation was corroborated by H&E staining of long bones showing the loss of vascular barriers and release of erythrocytes in the BM parenchyma (Figure S2D, red arrows pointing at erythrocytes). To quantify the vascular leakiness, we used a strategy allowing reliable measurement independent of the fluorescence intensity of the probe (detailed in Figure S2E and STAR Methods). The intravital observation of the calvarium BM vasculature at different time points after injection of the vessel-pooling agents showed an increased vascular leakiness starting between 6 and 9 min post injection (Figure 2C). We quantified vascular permeability in xenografts of different human AML samples 10 min after injection of the vessel-pooling agents (Figures 2D and 2E). All the samples used in our study, which represented different molecular subtypes of human AML, induced increased vascular leakiness in the BM. A different method using in vivo Hoechst permeability in BM cells (Wong et al., 2015) was used to confirm the AML-associated vascular phenotype in the BM (Figures S2F and S2G). Of note, AML-engrafted areas appeared to be much more leaky than...
Figure 3. Persistence of Increased Vascular Permeability after Chemotherapy

(A) Schematic of the experiment. Patient-derived AML cells were injected intravenously into NSG mice and engraftment was monitored via BM aspirate. Once BM engraftment (>40%) assessed, 10 mg/kg/day of AraC or solvent was administrated subcutaneously for one or two alternate weeks, depending on the patient sample. Seven days after the last administration of AraC, vessel-pooling agents were administrated intravenously. Mice were euthanized 10 min later and calvaria imaged via 2P microscopy.

(B) Representative 3D reconstruction of the BM vasculature in the calvarium of mice engrafted with different patient-derived samples treated with solvent (top) or AraC (bottom). Scale bars represent 70 μm.

(C) Vascular leakiness in the BM of mice transplanted with different human AML patient-derived samples, treated with solvent or AraC, as depicted. Three or more replicates were used for each condition. Data are shown as whiskers minimum-to-maximum plots, the line inside the box representing the mean, the top and the bottom lines representing the 75% and 25% percentiles, respectively, and the lines above and below the box representing the SD.

(D) Absolute number of CD31+ ECs in the BM (n = 2 femurs) of mice transplanted with AML6 (top) or AML9 (bottom) patient-derived cells, treated with solvent or AraC, as depicted; n = 3 in each condition. Data are shown as mean ± SEM.

(E) Hypoxia represented as HypoxiSense signal intensity/voxel in the calvaria BM of non-transplanted mice (ctrl) and mice transplanted with CB-derived HSPCs (CB) or AML patient-derived samples, treated or not with AraC, as depicted. Ctrl, n = 9; CB UT, n = 7; CB AraC, n = 8; AML patients (AML6 and 9) UT, n = 16; AraC, **p < 0.01.

(legend continued on next page)
non-engrafted areas (Figures S2H and S2I). In normal BM, Nestin-associated vessels, characterized by higher expression of Sca-1 (Itkin et al., 2016), show less permeability than Nestin-<sup>-</sup> vessels (Figure S2J). On the contrary, in the presence of leukemic engraftment, both Nestin-<sup>+</sup> and Nestin-<sup>-</sup> vessels were associated with increased leakiness (Figures S2J and S2K), highlighting a substantial functional difference between normal and leukemic Nestin-associated Sca-1<sup>-</sup> vessels. Of note, reduction of normal periocyte coverage was also observed in the BM of AML PDX (Figure S2L). These results show an altered vascular permeability in the BM associated with AML engraftment, observed commonly throughout different AML subtypes.

Chemotherapy Does Not Rescue Normal Vascular Permeability in the BM

It is still debated whether AML-derived angiogenesis, in terms of MVD in the BM and VEGF quantification, is reduced in patients upon chemotherapy (Aref et al., 2005; Chand et al., 2016; Pedro et al., 2000; Reddy and Moreb, 2000). Thus, we analyzed the level of VEGF in BM biopsies obtained at diagnosis and post treatment from a cohort of AML patients (Tables S1 and S2). We found an increase in VEGF in the majority of patients’ post-treatment biopsies (Figures S3A and S3B). This suggests a persistent pro-angiogenic environment after chemotherapy. We thus studied the vascular niche state in AML PDX after chemotherapy. We set up a preclinical treatment protocol, which closely followed the clinical practice, by treating AML xenotransplanted mice daily with 10 mg/kg of AraC during one or two alternate weeks, depending on xenografts’ response (Figure 3A). This treatment significantly reduced the leukemic burden in the BM (Figure S3C) and induced a “remission-like” phase, during which the remaining leukemic cells persist and give rise to relapse after few weeks (not shown). Notably, residual AML cells after AraC treatment localized in close proximity to the vasculature (Figure S3D). We observed the BM vasculature with 2P microscopy during the “remission-like” phase in different cohorts of AML PDX treated with this protocol. Although the leukemic engraftment was significantly reduced, the vascular permeability was still very high, similar to mice treated with the control solvent, which were highly engrafted with leukemic cells (Figures 3B and 3C). Moreover, the number of ECs was not reduced in AML PDX after AraC treatment (Figure 3D), suggesting that the abnormal vascular phenotype was maintained. This effect was not due to an unspecific toxicity of the chemotherapy on the vascular niche, as non-transplanted mice or mice engrafted with normal HSPCs and subjected to the same treatment did not show abnormal vascular permeability (Figures S3E and S3F), frequency, or number of ECs in the BM (Figures S3G and S3H). These results show that in AML disease, the vascular barrier functionality is still compromised after chemotherapeutic treatment. Furthermore, we observed that BM hypoxia was not normalized upon removal of leukemic cells (Figures 3E and 3F).

Notably, similarly to what was observed in presence of high leukemic infiltration, most of the vessels were still close to hypoxic areas upon AraC treatment (Figure 3G). These data highlight a persistent poorly functional vasculature in the BM of AML PDX after chemotherapy.

AML-Induced Molecular Signature in Vascular ECs

AML cells have been extensively studied at the transcriptional level, with many reports assessing the production of angiogenic factors (Aref et al., 2002; Hatfield et al., 2009; Ribatti et al., 2007). To identify common downstream effectors of this angiogenic stimulation in the vascular niche, we studied the transcriptome of BM-derived ECs retrieved from mice engrafted with different AML patient-derived cells via RNA sequencing (RNA-seq), and compared it with ECs retrieved from non-engrafted mice or mice engrafted with normal HSPCs. As shown by the hierarchical clustering, despite the heterogeneity found among the groups, we identified a common deregulated signature associated with human AML engraftment (Figure 4A). Gene set enrichment analysis (GSEA) highlighted several altered processes directly associated with the abnormal vascular phenotype observed above, including vasculature development, angiogenesis, and response to hypoxia (Figure 4B). Several deregulated pathways associated with angiogenic and hypoxic stimulation were identified (Figures 4B and S4A). Interestingly, the expression of adhesion molecules changed dramatically upon AML engraftment, with upregulation of integrins associated with Fak pathway activation in ECs (Figure S4B) inducing migration, growth, and survival of newly forming vessels during angiogenesis (Muether et al., 2007; Stenzel et al., 2011; Weis and Cheresh, 2011), as well as initiation of VEGF-mediated vascular permeability (Chen et al., 2012; Elcei et al., 2002; Rodrigues and Granger, 2015). We noticed also reduced expression of tight junction components, e.g., Cldn1 responsible for maintaining endothelial layer integrity (Rodrigues and Granger, 2015) or Vcam1 required for HSC homing and retention to the niche (Lewandowski et al., 2010; Ulyanova et al., 2007). Interestingly, we found among the common highest up-regulated genes in ECs upon AML engraftment Nos4, a NAPDH oxidase expressed in the vasculature and involved in the response to hypoxia via production of reactive oxygen species (ROS), activation of nitric oxide synthase 3 (NOS3), and release of nitric oxide (NO) (Craigie et al., 2011) (Figure 4C). Of note, in response to AML engraftment, Nos4 expression in ECs (Figure 4D) and ROS production in the BM (Figure 4E) were both increased. Moreover, the upregulated expression of Nos4 in ECs and the high production of ROS in the BM were maintained after AraC treatment (Figures 4F and 4G). These data show that the AML-induced hypoxic environment alters the molecular signature of vascular ECs, activating several pro-angiogenic genes.
Figure 4. AML-Induced Transcriptional Signature in BM ECs

(A) Heatmap from hierarchical clustering of GSEA top 100 ranked genes between untransplanted mice (ctrl group), mice engrafted with human CB-derived HSPCs (CB group), and mice engrafted with different patient-derived samples (AML group) derived BM ECs, using RNA-seq data. Red indicates higher expression values and blue indicates lower expression values (log2 FPKM [fragments per kilobase per million mapped] scale).

(B) Enrichment plots for Processes (top) and Pathways (bottom) from GSEA between CB and AML groups using RNA-seq gene expression data indicate enrichment of these pathways in AML group. Normalized enrichment score (NES) and nominal p value are shown.

(C) Normalized expression values of Nox4 in CB versus AML groups analyzed via RNA-seq. Data are shown as whiskers minimum-to-maximum plots, the line inside the box representing the mean, the top and the bottom lines representing the 75% and 25% percentiles, respectively, and the lines above and below the box representing the SD.

(D) Nox4 expression analyzed by qRT-PCR in CD31 BM cells retrieved from the depicted groups of mice. CB, n = 5; AML cell lines, n = 8; AML patients (AML1, 2, 9), n = 13. Data are shown as mean ± SEM.

(E) Cellular ROS levels in BMC retrieved from mice of depicted groups; n = 3 per group. Data are shown as mean ± SEM.

(F) Nox4 expression analyzed by qRT-PCR in CD31 BM cells retrieved from mice engrafted with AML6 and treated or not with AraC. n = 3 per group. Data are shown as mean ± SEM.

(G) Cellular ROS levels in BMC retrieved from mice engrafted with AML6 and treated or not with AraC; n = 3 per group. Data are shown as mean ± SEM. ns, not significant; *p < 0.05, **p < 0.01. See also Figure S4.
pathways and positively regulating Nox4 expression and the response to hypoxia.

**AML Engraftment Is Associated with Increased Nitric Oxide Levels in the BM**

As NO is the major mediator of vascular permeability in both physiological and several pathological conditions (Aramoto et al., 2004; Fukumura et al., 2001, 2006; Mayhan, 1999), we investigated whether the AML-derived upregulation of Nox4 in ECs influenced NO production in the BM vascular niche. We found increased levels of NO in the BM of AML xenografts compared with non-transplanted mice or mice engrafted with normal HSPCs (Figure 5A). A slight but significant increase of NO levels was also observed in BM of PDX with low engraftment (not shown). This effect seemed to be specific to BM (Figure S5A). Of note, as already observed for vascular permeability and Nox4 expression in ECs, chemotherapy failed to normalize the NO levels in the BM of AML PDX (Figure 5B).

To translate our observations to the human disease, we analyzed the NO levels in our cohort of AML patient-derived BM biopsies and compared them with those of healthy donors (Tables S1 and S2). We observed a significant increase in the NO level in BM aspirates of patients at diagnosis, which were similar to those observed in xenografts, compared with healthy donors (Figure 5C). NO levels were higher in the BM compared with the blood of the same patient at diagnosis (Figure S5B), similar to what was observed in mice with AML xenografts. Importantly, also similar to what was observed in AML xenografts, the NO levels remained elevated in the majority of post-treatment BM samples analyzed (Figures 5D and 5E). Moreover, failure in reducing NO levels appeared to be associated with a higher incidence of treatment failure (Figure S5C), with a probability of 73% in the cohort of patients studied herein (Figure S5D). This probability increases to 100% if we restrict the analysis to the intermediate-risk group samples, characterized by a more controversial prognostic assessment (Figure S5E). Our data suggest that persistence of high NO levels in the BM of patients after therapy could represent a biomarker for poor prognosis, particularly in patients with intermediate-risk AML. The upregulation of Nox4 in ECs, indicating an overactivation of NOS3 in this compartment, likely is responsible for the increased NO levels in the BM. We therefore analyzed the expression of NOS3 in the two major components of the vascular niche, endothelial (CD31\(^{+}\)) and mesenchymal perivascular (Nestin\(^{+}\) cells (Figure S5F). We observed that ECs represented the major compartment expressing NOS3 both at mRNA (Figure S5G) and protein (Figure 5F) levels. Moreover, we observed that NOS3 protein expression was significantly increased in ECs retrieved from mice engrafted with human AML patient-derived samples (Figure 5G). Of note, not only the total NOS3 protein level but also its activation was increased, as shown by increased phosphorylation of Ser1177 (Figure 5H (Kolluru et al., 2010)). In contrast, NOS3 was not increased in the mesenchymal compartment upon AML engraftment (Figure S5H). NOS2 protein expression is not changed in either the endothelial or the mesenchymal compartment upon AML engraftment (Figure S5I). Analysis of different endothelial subpopulations revealed that in normal BM, CD31\(^{+}\)Sca-1\(^{\text{high}}\) ECs exhibit lower ROS and NO levels than CD31\(^{+}\)Sca-1\(^{\text{low}}\) ECs (Figure 5I), and AML engraftment caused an increase in both ROS and NO levels in the CD31\(^{+}\)Sca-1\(^{\text{high}}\) population (Figure 5J). We investigated the consequences of long-term higher production of NO and ROS in ECs, and observed that in heavily engrafted AML xenografts ECs displayed increased apoptosis in both Sca-1\(^{\text{high}}\) and Sca-1\(^{\text{low}}\) ECs (Figure S5J), whereas cell-cycle arrest was restricted to the Sca-1\(^{\text{high}}\) EC fraction (Figure S5K). We next evaluated whether the leukemic compartment might also contribute to the NO overproduction. Human AML cells do not express higher mRNA levels of NOS2 or NOS3 compared with normal hematopoietic cells residing in the BM, as reported in publicly available datasets (HemaExplorer/BloodSpot, not shown). Moreover, the mRNA levels of these enzymes in AML patient-derived samples were similar to those from healthy donors (Figures S5L and S5M). There was no significant upregulation of these enzymes after chemotherapy (Figures S5N and S5O), or in AML xenografts (P1) compared with their primary counterpart (P0) (Figure S5P). The NOS3 protein level in AML was much lower than that in the endothelial compartment (Figure S5Q). Our results show an overproduction of NO in the BM in the presence of AML at diagnosis as well as after therapy, produced by an abnormally activated NOS3 in BM ECs.

**Targeting Vascular Permeability Cooperates with Chemotherapy to Improve AML Treatment Response**

To test whether the exogenous production of NO has a role in leukemia transplantation and progression, we transplanted leukemic cells from two models of murine AML (MLL-AF9 and MLL-ENL) in Nos3-knockout (Nos3-KO) recipient mice and studied the status of the vasculature and disease progression. Interestingly, MLL-AF9 leukemia occurrence was significantly delayed in Nos3-KO mice (Figure S6A, top), Nos3-KO mice were active and healthy at the time wild-type (WT) mice showed clear signs of disease and were euthanized. Similar results were obtained using two distinct clones of MLL-ENL AML transplanted in WT or Nos3-KO recipient mice, showing significantly reduced disease penetrance (Figure S6A, bottom), reduced leukemic engraftment in the BM and spleen (Figures S6B and S6C), and amelioration of the anemic phenotype caused by AML (Figure S6D). Moreover, Nos3 inactivation cooperated with AraC treatment to reduce leukemic engraftment in both MLL-AF9 and MLL-ENL models (Figure 6A) and reduced the vascular leakiness in the BM of AML-engrafted mice (Figure 6B). Together these data suggest that exogenous NO production by NOS3 has a role in AML engraftment. We next investigated whether the observed difference was likely to be attributed to a delay in engraftment or also to a reduction in disease progression. For that we used the PDX model and allowed AML cells to engraft the animals before inhibiting NO production with NOS inhibitors. We set up a similar treatment strategy to the one used before, including NOS inhibitors during and after the chemotherapy, to normalize the vascular permeability (Figure 6C). Treatment of mice xenografted with human ML1 cell line with NOS inhibitors in combination with chemotherapy significantly inhibited leukemic progression, as shown by reduced number of leukemic cells in the BM and the spleen (Figures S6E and S6F). The sole administration of NO inhibitors in heavily engrafted leukemic mice did not produce any significant decrease of leukemic load in the BM (Figure S6E), likely due to
Figure 5. AML Engraftment Is Associated with Increased Nitric Oxide Levels in the BM

(A) Nitric oxide levels shown as DAF fluorescence intensity (normalized mean fluorescence intensity [MFI]) in BM cells of non-transplanted mice (ctrl) and mice transplanted with CB-derived HSPCs (CB) or AML patient-derived samples, as depicted. Ctrl, n = 13; CB, n = 24; AML patients (AML1, 2, 6, 8, 9), n = 35. Data are shown as minimum-to-maximum box plots, the line inside the box representing the mean.

(B) Nitric oxide levels (normalized MFI) in BM cells of mice transplanted with AML patient-derived cells, treated with solvent or AraC, as depicted. AML patients: AML6 and 9. UT, n = 7; AraC, n = 7. Data are shown as minimum-to-maximum box plots, the line inside the box representing the mean.

(C) Nitric oxide levels in BM cells of healthy donors or AML patients at diagnosis. Healthy ctrl, n = 8; AML patients, n = 19. Error bars represent mean ± SEM.

(D) Nitric oxide levels in BM cells of AML patients at diagnosis (D) and post treatment (PT); n = 18. Error bars represent mean.

(E) Pie chart showing the percentage of patients with decreased or stable/increased levels of nitric oxide after treatment.

(F) NOS3 expression measured via flow cytometry in Nestin+ and CD31+ cells, as depicted. Data are representative of triplicates.

(G) Percentage of NOS3+ ECs measured by flow cytometry in the BM of non-transplanted mice and mice engrafted with CB-derived HSPCs (CB) or AML patient-derived samples, as depicted. Ctrl, n = 8; CB, n = 12; AML patients (AML2, 6, 8, 9), n = 24. Data are shown as mean ± SEM.

(H) Activated NOS3 (measured by flow cytometry with an antiphosphoSer1177 antibody) in CD31+ ECs of the BM of mice engrafted with CB-derived CD34+ cells (CB) or AML patient-derived samples, as depicted. CB, n = 4; AML patients (AML2, 6, 9), n = 9. Data are shown as mean ± SEM.

(I) Cellular ROS (left) and nitric oxide (right) levels in different populations of BM-derived ECs, as depicted, in non-transplanted mice (ctrl). Data are representative of triplicates.

(J) Cellular ROS (left) and nitric oxide (right) levels in different populations of BM-derived ECs, as depicted, in mice engrafted with AML6 patient-derived cells. Data are representative of triplicates.

ns, not significant; *p < 0.05, **p < 0.01, ****p < 0.0001. See also Figure S5.
the absence of a specific effect of NO on leukemic cells. Indeed, treatment of leukemic cells (ML1 or primary patient samples) in vitro with NO inhibitor or NO donor did not affect their survival or proliferation ability (Figures S6G and S6H). NO inhibition combined with AraC treatment reduced NOS3 activation (Figure 6D), vascular leakiness (Figure S6I), and the BM hypoxia (Figure 6E).

Importantly, the combined treatment was more powerful than the chemotherapy alone in reducing leukemic progression in the BM (Figures 6F and S6J) and the spleen (Figure S6K) of mice engrafted with two different patient-derived samples (Figure S6L). The combined treatment also significantly extended the “remission-like” phase of the disease compared with chemotherapy alone (Figure 6G). These data show that increased NO levels contribute to the vascular leakiness in AML-engrafted mice, and restoring normal vascular function in the BM by NOS inhibition improves the therapeutic response compared with standard chemotherapy.

**NOS Inhibition Potentiates Normal HSPC Function**

Increased levels of NO have been associated with increased HSPC motility and egress from the BM, leading to a lower repopulation capacity of the BM stem cell pool after injuries (Aicher et al., 2003; Gur-Cohen et al., 2015). Moreover, recent findings indicate that loss of vascular integrity promotes HSPC mobilization and loss of the HSC pool in the BM (Itkin et al., 2016). We thus speculated that increased NO and vascular leakiness in AML could affect normal HSC. In the BM of AML xenografts, the residual murine HSC displayed higher levels of ROS (Figure 7A) and were mobilized to the periphery, as shown by increased SLAM* cells in the blood of mice with AML xenografts compared with those without, as well as a reduced proportion of HSC in the BM (Figures 7B and 7C). The absolute number of SLAM* cells in the BM was also decreased in mice with AML xenografts and in mice transplanted with murine MLL-ENL leukemic cells (Figures S7A and S7B). Therefore, we tested whether normalization of the vascular niche with NOS inhibitors would be beneficial for normal HSC function. We observed that NOS inhibitors in combination with AraC treatment were more efficient than chemotherapy alone in re-establishing the normal number of SLAM* cells in the BM of PDX (Figure 7D). Not only the number but also the stem cell function of these SLAM* cells was increased, as shown by their repopulation ability in secondary recipients (Figures S7C and 7E–7G). These data suggest that overproduction of NO by altered vascular niche affects normal residual HSC in the BM and that NOS inhibition may improve AML therapy by restoring normal stem cell function. We next tested the effect of NOS inhibitors on HSPCs. Hence, we treated mice engrafted with normal HSPCs with NOS inhibitors and analyzed the effect on the stem cell pool (Figure 7H). Interestingly, we observed reduced HSPC egress to the blood (Figure 7I), an effect specific for this compartment as no significant difference was observed on the egress of total hCD45* cells into the blood (Figure S7D). This was accompanied by a specific increase of HSPCs in the BM (Figures 7J and S7E). This effect was not due to an increase in proliferation of the stem pool, as shown by cell-cycle analysis (Figure S7F), thus excluding a possible exhaustion of the compartment. Indeed, NO inhibition not only increased the number of HSPCs in the BM but also their stem cell function, as shown by their increased repopulation ability in secondary transplantation assays (Figure 7K). We next tested whether NO inhibition would be beneficial for residual normal HSC to outcompete AML cells during the relapse process. To achieve this, and mimic the competition between AML cells and normal HSCs into the BM niche, we injected simultaneously normal human CB-derived HSPCs and AML patient-derived cells intravenously into non-irradiated NBSGW mice (McIntosh et al., 2015) and monitored the normal versus malignant engraftment in the presence or absence of long-term NOS inhibitors (Figure 7L). Analysis of the BM engraftment at the end of the treatment showed that NOS inhibition favored the normal over leukemic engraftment in the BM (Figures 7M and 7N). Together, these results suggest that the use of NO inhibitors will not only reverse the vascular permeability observed during AML progression and thus improve chemotherapy efficiency, but will also restore a normal stem cell vascular niche, likely protecting normal residual HSCs and allowing them to outcompete the leukemic cells.

Altogether, our results provide strong evidence for the importance of the alteration of the vascular niche in AML progression and relapse, via increased NO production by ECs, and call for clinical trials incorporating NOS inhibitors to target the abnormal vascular niche and improve the treatment response.
The image contains various graphs and data representations from a scientific document. The text includes references to HSPC (hematopoietic stem/progenitor cells) and AML (acute myeloid leukemia), with data on engraftment, transplantation analysis, and various biological parameters such as the percentage of donor CD45 cells in the blood and absolute numbers of cells in BM (bone marrow) and spleen. The graphs show results from different treatments and comparisons, including NOS inhibitors and CB P1. The data is presented with statistical significance markers (*, **, ***). The figures are labeled A through N, with each graph depicting different aspects of the study's findings. The legend on the next page is indicated to provide additional context for the graph symbols and data.
Tumor angiogenesis is a pathological complication of several types of solid cancers, which has been extensively studied and characterized. The angiogenic process is mostly initiated by pro-angiogenic factors derived from tumor cells, which in turn activates a pro-inflammatory response in the microenvironment, associated with the initiation of irregular and uncontrolled vascular growth. Newly formed tumor vessels appear abnormal and leaky in most of the solid cancers. This structurally aberrant network affects the local blood flow, metabolite exchange, and oxygenation, leading to poor drug delivery, increased tumor growth, and metastatic potential (Carmeliet and Jain, 2011a, 2011b; Jain, 2005).

In the AML context, leukemic cells have been shown to modify ECs in vivo via several pathways, mainly mediated by pro-angiogenic factors such as VEGF family members (Kampen et al., 2013; Poulos et al., 2014). Moreover, publicly available datasets as well as several studies reported that VEGF is not the only pro-angiogenic factor produced by AML cells, which may be a major reason why clinical trials incorporating VEGF inhibitors have not been very successful. AML cells also express high levels of hepatocyte growth factor (Aref et al., 2002), fibroblast growth factor (Ribatti et al., 2007), and other factors (Chand et al., 2016; Hatfield et al., 2009; Lee and Kim, 2014). However, no extensive analysis of the AML vascular niche in vivo has been reported.

In the present study, we provide a detailed picture of the vasculature in AML. We not only confirm that AML engraftment induces hypoxia and angiogenesis in the BM but also show that it alters the normal vascular architecture and function. ECs involved in the angiogenic process are highly metabolically active and consume a high level of oxygen (Potente and Carmeliet, 2017). As a result, these newly formed abnormal vessels are incapable of supplying nutrients and oxygen (as well as drugs) in a homogeneous way in the BM tissue, likely contributing to maintain the overall hypoxia. At a late stage of disease, after facing an exponential and uncontrolled growth and being exposed to high levels of NO/ROS production, ECs display an increased apoptotic phenotype, likely associated with a toxic process of vascular regression (Lobov et al., 2011).

The crosstalk between leukemic cells and ECs in the BM has not been extensively studied at the molecular level. The available data in the literature rely on the in vitro co-culture system, which does not faithfully reproduce the complexity of the in vivo environment. Our analysis of the transcriptome of vascular ECs upon human AML engraftment provides important insights into the field. First, it confirms at the molecular level the observed toxic phenotypes. Second, it highlights commonly deregulated genes and pathways, which can represent potential targets for the disruption of the AML-microenvironment crosstalk. Interestingly, the NOX4-NOS3 axis, already reported previously to act in response to stress induced by hypoxic conditions (Craigie et al., 2011; Drummond et al., 2000; Mittal et al., 2007; Thomas et al., 2002; Zhang et al., 2008), has a major role in orchestrating the AML-induced vascular leakiness.

Persistent vascular leakiness and increased NO levels in AML xenografts after induction therapy suggest that the abnormal and poorly functional vessels participate to maintain a hypoxic vascular environment during the remission phase, likely contributing to treatment failure. Our hypothesis is strengthened by the analysis of NO in an independent cohort of patients which, although relatively small, faithfully represents the high diversity of the AML genetic landscape. Our analysis argues for NO as a potential prognostic BM marker, as stable/increased levels of NO post-induction therapy seems to be associated with a poor clinical outcome.

Increased vascular leakiness can lead to poor drug delivery and to a microenvironment favoring resistance to therapy and...
relapse. The participation of vascular leakiness to the pathophysiology of AML can be exerted at different levels. First, increased vascular permeability deregulates the balance of blood supply to the BM and, consequently, drug delivery. This leads to the formation of areas with low perfusion rate, where leukemic cells can hide and escape therapy. Restoring a normal vascular function would allow more homogeneous chemotherapeutic delivery. Second, high vascular permeability (Itkin et al., 2016) and high NO levels (Gur-Cohen et al., 2015) have recently been associated with increased HSC activation and egress from the BM in mice, with a consequent reduction of their repopulating activity. Our data showing increased ROS levels and higher egress of BM-derived HSCs are in line with these findings, further extending them to the human HSPC compartment. Indeed, we show that inhibition of NO results in an enriched number and activity of BM HSPCs. Conversely, the presence of AML engraftment induces the generation of an exogenous overproduction of NO in the BM, which likely affects the HSC motility and activity, bringing a rapid exhaustion of their functionality. In this scenario, inhibition of NO would produce multiple benefits to AML patients. NOS inhibitors are already in clinical trials (Granados-Principal et al., 2015; Ozkor et al., 2015) and specific inhibitors of NOS3 have been developed (Bucci et al., 2000; Gratton et al., 2003). Thus, our preclinical data using both genetic and pharmacologic approaches to inhibit NO in combination with AraC treatment show that targeting vascular permeability would be a successful strategy to normalize the BM vascular functionality, improve therapeutic responses, and preserve normal HSC function.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- **KEY RESOURCES TABLE**
- **CONTACT FOR REAGENT AND RESOURCE SHARING**
  - Human Primary Samples
  - Mouse Models
  - Drug Treatments In Vivo
  - Intravitral Two-Photon Microscopy
  - Images Processing
  - Bioluminescence Imaging
  - Cell Lines
  - Transduction and Culture of Human Primary Samples
  - Transduction and Culture of Murine Primary Samples
  - AML and HSPCs Transplantation Assay In Vivo
  - Bone Marrow Cell Isolation
  - HO Diffusion
  - Hypoxiprobe Assessment
  - Flow Cytometry Analysis and Cell Sorting
  - Drug Treatments In Vitro
  - Reverse Transcriptase and Real Time Quantitative PCR (RT-qPCR)
  - Histological Processing and Immunostaining
  - Sample Preparation for RNA-Sequencing
  - RNA-Sequencing Analysis Methods
  - Statistical Analysis
- **DATA AND SOFTWARE AVAILABILITY**
  - Data Resources

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures and two tables and can be found with this article online at [http://dx.doi.org/10.1016/j.celrep.2017.08.001](http://dx.doi.org/10.1016/j.celrep.2017.08.001).

AUTHOR CONTRIBUTIONS

D.P. designed, conducted, and performed the experiments and wrote the manuscript. A.d.T., A.A., K.R.-P., K.F., F.L., L.A.M., and B.M. performed some experiments. L.B. prepared the libraries and P.C. performed the statistical analysis of RNA-seq data. G.D. performed statistical analysis on patients’ cohorts. J.G. provided patient samples and clinical information. D.B. designed experiments and wrote the manuscript.

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REFERENCES

Aicher, A., Heeschen, C., Mildner-Rihm, C., Urbich, C., Ihling, C., Technau-Ihling, K., Zeiher, A.M., and Dimmel, S. (2003). Essential role of endothelial nitric oxide synthase for mobilization of stem and progenitor cells. Nat. Med. 9, 1370–1376.

Aramoto, H., Breslin, J.W., Pappas, P.J., Hobson, R.W., 2nd, and Duran, W.N. (2004). Vascular endothelial growth factor stimulates differential signaling pathways in in vivo microcirculation. Am. J. Physiol. Heart Circ. Physiol. 287, H1590–H1598.

Aref, S., Mabed, M., Sakrana, M., Goda, T., and El-Sherbiny, M. (2002). Soluble hepatocyte growth factor (sHGF) and vascular endothelial growth factor (sVEGF) in adult acute myeloid leukemia: relationship to disease characteristics. Hematology 7, 273–279.

Aref, S., El Sherbiny, M., Goda, T., Fouda, M., Al Askalany, H., and Abdalla, D. (2005). Soluble VEGF/sFlt1 ratio is an independent predictor of AML patient outcome. Hematology 10, 131–134.

Bao, B., Groves, K., Zhang, J., Handy, E., Kennedy, P., Cuneo, G., Supuran, C.T., Yared, W., Rajopadye, M., and Peterson, J.D. (2012). In vivo imaging and quantification of carbonic anhydrase IX expression as an endogenous biomarker of tumor hypoxia. PLoS One 7, e50860.

Boulais, P.E., and Frenette, P.S. (2015). Making sense of hematopoietic stem cell niches. Blood 125, 2621–2629.

Bucci, M., Gratton, J.P., Rudic, R.D., Acevedo, L., Roviezzo, F., Cirino, G., and Sessa, W.C. (2000). In vivo delivery of the caveolin-1 scaffolding domain inhibits nitric oxide synthesis and reduces inflammation. Nat. Med. 6, 1362–1367.
Buchner, T., Schienk, R.F., Schaich, M., Dohner, K., Krahl, R., Krauter, J., Heil, G., Krug, U., Sauerland, M.C., Heinecke, A., et al. (2012). Acute myeloid leukaemia (AML): different treatment strategies versus a common standard arm—combined prospective analysis by the German AML Intergroup. J. Clin. Oncol. 30, 3604–3610.

Carmeliet, P., and Jain, R.K. (2011a). Molecular mechanisms and clinical applications of angiogenesis. Nature 473, 298–307.

Carmeliet, P., and Jain, R.K. (2011b). Principles and mechanisms of vessel normalization for cancer and other angiogenic diseases. Nat. Rev. Drug Discov. 10, 417–427.

Chand, R., Chandra, H., Chandra, S., and Verma, S.K. (2016). Role of microvessel density and vascular endothelial growth factor in angiogenesis of hematological malignancies. Bone Marrow Res. 2016, 5043483.

Chen, X.L., Nam, J.O., Jean, C., Lawson, C., Walsh, C.T., Goka, E., Lim, S.T., Zhou, M., Broxmeyer, H.E., Scadden, D.T., and Qu, C.K. (2016). Transcriptional and posttranscriptional regulation of endothelial nitric oxid synthase activation. Circulation 124, 731–740.

Dobin, A., Davis, C.A., Schlesinger, F., Drenkow, J., Zaleski, C., Jha, S., Batut, P., Chaisson, M., and Gingeras, T.R. (2013). STAR: ultrafast universal RNA-seq aligner. Bioinformatics 29, 15–21.

Dong, L., Yu, W.M., Zheng, H., Loh, M.L., Bunting, S.T., Pauly, M., Huang, G., Zhou, M., Broxmeyer, H.E., Scadden, D.T., and Qu, C.K. (2016). Leukaemogenic effects of Ptpn11 activating mutations in the stem cell microenvironment. Nature 539, 304–308.

Drummond, G.R., Cai, H., Davis, M.E., Ramosamy, S., and Harrison, D.G. (2000). Transcriptional and posttranscriptional regulation of endothelial nitric oxide synthase expression by hydrogen peroxide. Circ. Res. 86, 347–354.

Egawa, G., Nakamizo, S., Natsuaki, Y., Doi, H., Miyachi, Y., and Kabashima, K. (2013). Intravital analysis of vascular permeability in mice using two-photon microscopy. Sci. Rep. 3, 1932.

Eliceiri, B.P., Puente, X.S., Hood, J.D., Stupack, D.G., Schlaepfer, D.D., Huang, X.Z., Sheppard, D., and Cheresh, D.A. (2002). Src-mediated coupling of focal adhesion kinase to integrin alpha(v)beta5 in vascular endothelial growth factor signaling. J. Cell Biol. 157, 149–160.

Fiedler, W., Mesters, R., Tinnefeld, H., Loges, S., Staub, P., Duhrensen, U., Flasshove, M., Ottmann, O.G., Jung, W., Cavalli, F., et al. (2003). A phase 2 clinical study of SU5416 in patients with refractory acute myeloid leukemia. Blood 102, 2763–2767.

Fiedler, W., Serve, H., Dohner, H., Schwittay, M., Ottmann, O.G., O’Farrell, A.M., Bello, C.L., Alfred, R., Manning, W.C., Cherrington, J.M., et al. (2005). A phase 1 study of SU11248 in the treatment of patients with refractory or resistant acute myeloid leukemia (AML) or not amenable to conventional therapy for the disease. Blood 105, 986–993.

Foster, K., Lassailly, F., Anjos-Afonso, F., Currie, E., Rouault-Pierre, K., and Bonnet, D. (2015). Different motile behaviors of human hematopoietic stem versus progenitor cells at the osteoblastic niche. Stem Cell Rep. 5, 690–701.

Frisch, B.J., Ashton, J.M., Xing, L., Becker, M.W., Jordan, C.T., and Calvi, L.M. (2012). Functional inhibition of osteoblastic cells in an in vivo mouse model of myeloid leukemia. Blood 119, 540–550.

Fukumura, D., and Jain, R.K. (2008). Imaging angiogenesis and the microenvironment. APIMS 116, 695–715.

Fukumura, D., Gohongi, T., Kadambi, A., Izumi, Y., Ang, J., Yun, C.O., Buerk, D.G., Huang, P.L., and Jain, R.K. (2001). Predominant role of endothelial nitric oxide synthase in vascular endothelial growth factor-induced angiogenesis and vascular permeability. Proc. Natl. Acad. Sci. USA 98, 2604–2609.

Fukumura, D., Kashigawagi, S., and Jain, R.K. (2006). The role of nitric oxide in tumour progression. Nat. Rev. Cancer 6, 521–534.

Giles, F.J., Bellamy, W.T., Estrov, Z., O’Brien, S.M., Verstovsek, S., Ravandi, F., Beran, M., Bycott, P., Pitavalla, Y., Steinfeldt, H., et al. (2006). The angiogenesis agent, AG-013736, has minimal activity in elderly patients with poor prognosis acute myeloid leukemia (AML) or myelodysplastic syndrome (MDS). Leuk. Res. 30, 801–811.

Granados-Principal, S., Liu, Y., Guevara, M.L., Blanco, E., Choi, D.S., Qian, W., Patel, T., Rodriguez, A.A., Cusimano, J., Weiss, H.L., et al. (2015). Inhibition of iNOS as a novel effective targeted therapy against triple-negative breast cancer. Breast Cancer Res. 17, 25.

Gratton, J.P., Lin, M.I., Yu, J., Weiss, E.D., Jiang, Z.L., Fairchild, T.A., Iwakiri, Y., Groszmann, R., Claffey, K.P., Cheng, Y.C., and Sessa, W.C. (2003). Selective inhibition of tumor microvascular permeability by cavantrin blocks tumor progression in mice. Cancer Cell 4, 31–39.

Grimwade, D., Hills, R.K., Moorman, A.V., Walker, H., Chatters, S., Goldstone, A.H., Wheatley, K., Harrison, C.J., and Burnett, A.K.; National Cancer Research Institute Adult Leukaemia Working Group (2010). Refinement of cytogenetic classification in acute myeloid leukaemia: determination of prognostic significance of rare recurring chromosomal abnormalities among 5876 younger adult patients treated in the United Kingdom Medical Research Council trials. Blood 116, 354–365.

Gur-Cohen, S., Itkin, T., Chakraborty, S., Graf, C., Kollet, O., Ludin, A., Golan, K., Kalinkovich, A., Ledergor, G., Wong, E., et al. (2015). PAR1 signaling regulates the retention and recruitment of EPCR-expressing bone marrow hematopoietic stem cells. Nat. Med. 21, 1307–1317.

Hanou, M., Zhang, D., Mizoguchi, T., Pinho, S., Pierce, H., Kunisaki, Y., Lacombe, J., Armstrong, S.A., Duhrensen, U., and Frenette, P.S. (2014). Acute myelogenous leukaemia-induced sympathetic neuropathy promotes malignancy in an altered hematopoietic stem cell niche. Cell Stem Cell 15, 365–375.

Hattfield, K., Oyan, A.M., Ersvaer, E., Kalland, K.H., Lassailly, F., Gjertsen, B.T., and Bruder, O. (2009). Primary human acute myeloid leukaemia cells increase the proliferation of microvascular endothelial cells through the release of soluble mediators. Br. J. Haematol. 144, 53–68.

Hooper, A.T., Butler, J.M., Nolan, D.J., Kranz, A., lida, K., Kobayashi, M., Kopp, H.G., Shido, K., Petit, I., Yang, K., et al. (2009). Engraftment and reconstitution of hematopoiesis is dependent on VEGFR2-mediated regeneration of sinusoidal endothelial cells. Cell Stem Cell 4, 263–274.

Horton, S.J., Grier, D.G., McGonigle, J.G., Thompson, A., Morey, M., De Silva, I., Moulding, D.A., Kioussis, D., Lappin, T.R., Brady, H.J., and Williams, O. (2005). Continuous DLL-ENL expression is necessary to establish a “Hox Code” and maintain immortalization of hematopoietic progenitor cells. Cancer Res. 65, 9245–9252.

Horton, S.J., Walf-Vorderwulbecke, V., Chatters, S.J., Sebire, N.J., de Boer, J., and Williams, O. (2009). Acute myeloid leukaemia induced by DLL-ENL is cured by oncogene ablation despite acquisition of complex genetic abnormalities. Blood 113, 4922–4929.

Huysung, J.W., Rodgers, G.M., and Shami, P.J. (2000). Evidence of increased angiogenesis in patients with acute myeloid leukaemia. Blood 95, 309–313.

Itkin, T., Gur-Cohen, S., Spencer, J.A., Schajnowitz, A., Ramosamy, S.K., Kusumbe, A.P., Ledergor, G., Jung, Y., Milo, I., Poulos, M.G., et al. (2016). Distinct bone marrow blood vessels differentially regulate haematopoiesis. Nature 532, 323–328.

Jain, R.K. (2005). Normalization of tumor vasculature: an emerging concept in antiangiogenic therapy. Science 307, 58–62.

Kampen, K.R., Ter Elst, A., and de Bont, E.S. (2013). Vascular endothelial growth factor signaling in acute myeloid leukaemia. Cell Mol. Life Sci. 70, 1307–1317.

Karolisich, D., Hinrichs, A.S., Furey, T.S., Roskin, K.M., Sugnet, C.W., Haussler, D., and Kent, W.J. (2004). The UCSC Table Browser data retrieval tool. Nucleic Acids Res. 32, D493–D496.

Kim, Y.W., Koo, B.K., Jeong, H.W., Yoon, M.J., Song, R., Shin, J., Jeong, D.C., Kim, S.H., and Kong, Y.Y. (2008). Defective Notch activation in microenvironment leads to myeloproliferative disease. Blood 112, 4628–4638.
Kode, A., Manavalan, J.S., Mosialou, I., Bhagat, G., Rathinam, C.V., Luo, N., Khiabanian, H., Lee, A., Murty, V.V., Friedman, R., et al. (2014). Leukaemogenesis induced by an activating beta-catenin mutation in osteoblasts. Nature 506, 240–244.

Kolluru, G.K., Siamwala, J.H., and Chatterjee, S. (2010). eNOS phosphorylation in health and disease. Biochimie 92, 1186–1198.

Kopp, H.G., Avecilla, S.T., Hooper, A.T., Shmelkov, S.V., Ramos, C.A., Zhang, F., and Rafii, S. (2005). Tie2 activation contributes to hemangiogenic regener-1513–1517.

Krause, D.S., Fulzele, K., Catic, A., Sun, C.C., Dombkowski, D., Hurley, M.P., Lezeau, S., Attar, E., Wu, J.Y., Lin, H.Y., et al. (2013). Differential regulation of myocardial leukemoidiasis by the bone marrow microenvironment. Nat. Med. 19, 1513–1517.

Lassailly, F., Foster, K., Lopez-Onieva, L., Currie, E., and Bonnet, D. (2013). Multimodal imaging reveals structural and functional heterogeneity in different bone marrow compartments: functional implications on hematopoietic stem cells. Blood 122, 1730–1740.

Lee, J.Y., and Kim, H.J. (2014). (Lymph)angiogenic influences on hematopoietic cells in acute myeloid leukemia. Exp. Mol. Med. 46, e122.

Lewandowski, D., Barroca, V., Duonge, F., Bayer, J., Van Nhye, J.T., Pestourie, C., Fouchet, P., Tavitian, B., and Romeo, P.H. (2010). In vivo cellular imaging pinpoints the role of reactive oxygen species in the early steps of adult hematopoietic reconstitution. Blood 115, 443–452.

Li, B., and Dewey, C.N. (2011). RSEM: accurate transcript quantification from RNA-seq data with or without a reference genome. BMC Bioinformatics 12, 323.

Lo Celso, C., Fleming, H.E., Wu, J.W., Zhao, C.X., Miake-Lye, S., Fujisaki, J., Lezeau, S., Attar, E., Wu, J.Y., Lin, H.Y., et al. (2013). Differential regulation of hematopoietic cells in acute myeloid leukemia. Exp. Hematol. 41, 976–986.e3.

R Development Core Team. (2015). A Language and Environment for Statistical Computing (R Foundation for Statistical Computing), 327–334.

Roboz, G.J., Giles, F.J., List, A.F., Cortes, J.E., Carlin, R., Kowalski, M., Bilić, S., Masson, E., Rosamilia, M., Schuster, M.W., et al. (2006). Phase 1 study of PTK787/ZK 222584, a small molecule tyrosine kinase receptor inhibitor, for the treatment of acute myeloid leukemia and myelodysplastic syndrome. Leukemia 20, 952–957.

Reddy, V., and Moreb, J. (2009). Prognostic correlation of increased angiogenesis in acute myeloid leukemia with cytogenetics. Blood 113, 1617–1618.

Ribatti, D., Vaccara, A., Rusnati, M., and Presta, M. (2007). The discovery of basic fibroblast growth factor/fibroblast growth factor-2 and its role in haematological malignancies. Cytokine Growth Factor. Rev. 18, 327–334.

Sipkins, D.A., Wei, X., Wu, J.W., Runnels, J.M., Cote, D., Means, T.K., Luster, A.D., Scadden, D.T., and Lin, C.P. (2005). In vivo imaging of specialized bone marrow endothelial microdomains for tumour engraftment. Nature 435, 969–973.

Stenzel, D., Lundkvist, A., Sauvaget, D., Busse, M., Graupera, M., van der Flier, A., Wijelath, E.S., Murray, J., Sobel, M., Costell, M., et al. (2011). Loss or inhibition of stromal-derived PGF prolongs survival of mice with imatinib-resistant Bcr-Abl(+) leukemia. Cancer Cell 19, 740–753.

Subramanian, A., Tamayo, P., Mootha, V.K., Mukherjee, S., Ebert, B.L., Gillette, M.A., Paulovich, A., Pomeroy, S.L., Golub, T.R., Lander, E.S., and Mesirov, J.P. (2005). Gene set enrichment analysis: a knowledge-based
approach for interpreting genome-wide expression profiles. Proc. Natl. Acad. Sci. USA 702, 15545–15550.

Thomas, S.R., Chen, K., and Keaney, J.F., Jr. (2002). Hydrogen peroxide activates endothelial nitric-oxide synthase through coordinated phosphorylation and dephosphorylation via a phosphoinositide 3-kinase-dependent signaling pathway. J. Biol. Chem. 277, 6017–6024.

Ulyanova, T., Priestley, G.V., Nakamoto, B., Jiang, Y., and Papayannopoulou, T. (2007). VCAM-1 ablation in nonhematopoietic cells in MxCre+ VCAM-1f/f mice is variable and dictates their phenotype. Exp. Hematol. 35, 565–571.

Wang, Z.Y., and Chen, Z. (2008). Acute promyelocytic leukemia: from highly fatal to highly curable. Blood 111, 2505–2515.

Weis, S.M., and Cheresh, D.A. (2011). alphaV integrins in angiogenesis and cancer. Cold Spring Harbor Perspect. Med. 1, a006478.

Wong, P.P., Demircioglu, F., Ghazaly, E., Alrawashdeh, W., Stratford, M.R., Scudamore, C.L., Cereser, B., Cmogorac-Jurcevic, T., McDonald, S., Elia, G., et al. (2015). Dual-action combination therapy enhances angiogenesis while reducing tumor growth and spread. Cancer Cell 27, 123–137.

Zahiragic, L., Schliemann, C., Bieker, R., Thoennissen, N.H., Burow, K., Kramer, C., Zuhladorf, M., Berdel, W.E., and Mesters, R.M. (2007). Bevacizumab reduces VEGF expression in patients with relapsed and refractory acute myeloid leukemia without clinical antileukemic activity. Leukemia 21, 1310–1312.

Zhang, Q., Malik, P., Pandey, D., Gupta, S., Jagnandar, D., Belin de Chantemele, E., Banfi, B., Marrero, M.B., Rudic, R.D., Stepp, D.W., and Fulton, D.J. (2008). Paradoxical activation of endothelial nitric oxide synthase by NADPH oxidase. Arterioscler. Thromb. Vasc. Biol. 28, 1627–1633.

Zhang, B., Ho, Y.W., Huang, Q., Maeda, T., Lin, A., Lee, S.U., Hair, A., Holyoake, T.L., Huetther, C., and Bhatia, R. (2012). Altered microenvironmental regulation of leukemic and normal stem cells in chronic myelogenous leukemia. Cancer Cell 21, 577–592.

Zhu, Z., Hattori, K., Zhang, H., Jimenez, X., Ludwig, D.L., Dias, S., Kussie, P., Koo, H., Kim, H.J., Lu, D., et al. (2003). Inhibition of human leukemia in an animal model with human antibodies directed against vascular endothelial growth factor receptor 2. Correlation between antibody affinity and biological activity. Leukemia 17, 604–611.
# STAR METHODS

## Key Resources Table

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| CD45 30-F11 1 in 400 Mouse | eBioscience | Cat: 45-0451-82; RRID: AB_1107002 |
| TER119 Ter-119 1 in 400 Mouse | eBioscience | Cat: 12-5921-82; RRID: AB_466042 |
| CD31 390 1 in 400 Mouse | eBioscience | Cat: 25-0311-82; RRID: AB_469616 |
| Endomucin V.7C7 1 in 100 Mouse | Santa Cruz Biotechnology | Cat: sc-65495; RRID: AB_2100037 |
| Sca-1 D7 1 in 400 Mouse | eBioscience | Cat: 17-5981-83; RRID: AB_496488 |
| Lin - 1 in 400 Mouse | Biologend | Cat: 133311; RRID: AB_11203535 |
| Kit 2B8 1 in 400 Mouse | BD Bioscience | Cat: 553354; RRID: AB_394805 |
| CD48 HM48-1 1 in 400 Mouse | Biologend | Cat: 103432; RRID: AB_256163 |
| CD150 TC15-12F12.2 1 in 400 Mouse | Biologend | Cat: 115914; RRID: AB_439797 |
| IB4 - 100ug/mouse Mouse | Life Technology | Cat: i21411; RRID: AB_2314662 |
| Sca-1/Ly6A/E EPR3355 1 in 100 Mouse | Abcam | Cat: ab109211; RRID: AB_10862573 |
| Ki67 SolA15 2 in 100 Mouse | eBioscience | Cat: 12-5698-82; RRID: AB_11150954 |
| Ki67 20Raj1 2 in 100 Human | eBioscience | Cat: 50-5699-82; RRID: AB_2574237 |
| rAnnexin V - 2 in 100 Mouse/Human | eBioscience | Cat: BMS306; RRID: AB_2687874 |
| CD45 PD7/26 + 2B11 1 in 100 Human | Dako | Cat: M0701; RRID: AB_2314143 |
| CD34 581 1 in 25 Human | BD Bioscience | Cat: 555821; RRID: AB_396150 |
| CD45 201 1 in 25 Human | BD Bioscience | Cat: 47-0459-42; RRID: AB_1944368 |
| CD38 HIT2 1 in 25 Human | BD Bioscience | Cat: 555462; RRID: AB_1994368 |
| Ki67 20Raj1 1 in 25 Human | eBioscience | Cat: 50-5699; RRID: AB_2574237 |
| CD45 HI30 20ug/mouse Human | Biologend | Cat: 304017; RRID: AB_389314 |
| NOS3-pSER1177 C9C3 1 in 100 Mouse/Human | Cell Signaling | Cat: 9570S; RRID: AB_823493 |
| NOS3-eNOS 1 in 20 Mouse/Human | BD Bioscience | Cat: 560103; RRID: AB_1645506 |
| NOS2 CXNFT 1 in 100 Mouse/Human | eBioscience | Cat: 17-5920-80; RRID: AB_2573243 |
| **Bacterial and Virus Strains** | | |
| pHAGE2-EF1aFull-MCS-ires-Tomato | Kind gift of G. Mostoslavsky; modified from J Immunol Methods. 2008;329:31-44 | N/A |
| pTWG-CMV-LUC-eGFP | Modified from pTWG-eGFP (Addgene) and N/A pGL4-50 luciferase reporter (Promega) | N/A |
| **Biological Samples** | | |
| See Tables 1, S1, and S2 | N/A | N/A |

**Chemicals, Peptides, and Recombinant Proteins**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Hypoxyprobe Plus Kit | Natural Pharmacia International | HP2-1000Kit |
| HypoxiSense 680 Fluorescent Imaging Agent | PerkinElmer | NEV11070 |
| DAF-FM DA solution | Sigma-Aldrich | D2321 |
| CellROX Deep Red Reagent | Thermo Fisher Scientific | C10422 |
| N^2^-Methyl-L-arginine acetate salt (NMMA) | Sigma-Aldrich | M7033 |
| Caveolin-1 scaffolding domain peptide (Cavtratin) | EnzolifeSciences | ALX-153-064-M005 |
| S-Nitroso-N-acetyl-DL-penicillamine (SNAP) | Sigma-Aldrich | N3398 |

(Continued on next page)
Further information and requests for resources and reagents should be directed to the Lead Contact Dominique Bonnet at dominique.bonnet@crick.ac.uk

Human Primary Samples
Umbilical Cord Blood (CB) samples were obtained from normal full-term deliveries after signed informed consent. AML samples used for xenografts were obtained after informed consent at St Bartholomew’s Hospital (London, UK). Details are listed in Table 1. BM specimens from 19 adult patients with newly diagnosed, untreated AML were also obtained after informed consent at St Bartholomew’s Hospital (London, UK). Patient characteristics are depicted in Table S1. Additional BM biopsies were available from these same 19 patients following chemotherapy (details in Table S2). To establish controls, BM biopsies were obtained either from healthy donors for bone marrow transplantation or from patients with diffuse large cell lymphoma without BM infiltration (Table S2).
Cells were enriched for CD34+, using an EasySep Human CD34 Positive Selection kit (StemCell Technologies, Vancouver, Canada) according to the manufacturer’s instructions, with a purity of 85 to 99%. T cells were depleted from all AML samples using the anti-CD3 mAb OKT-3 (West Lebanon, USA).

Mouse Models

All animal experiments were performed under the project license (PPL 70/8904) approved by the Home Office of UK and in accordance to The Francis Crick institute animal ethics committee guidelines. NESTIN-GFP mice were a kind gift from Dr G. Enikolopov. NOD-SCID IL2Rγnull (NSG), C57BL/6, NG2-DsRed, Nos3-KO, Rag2-KO/GammaC-KO and NOD.B6-SCID c-Kit+/−/Wt (NBSGW) strains were obtained from Jackson Laboratory, Bar Harbor, Maine, USA and were bred at The Francis Crick Institute Biological Resources Facility in individually vented cages (IVCs) under Specific Pathogen Free (SPF) conditions. NSG-NESTIN-GFP and NSG-NG2-DsRed mice were obtained by back-crossing the original lines into the NSG background (generation 8 or more).

Drug Treatments In Vivo

AraC was administrated subcutaneously at the indicated doses (10 mg/kg/day for NSG mice and 60 mg/kg/day for C57BL6 mice) at specific time points, for 7 days, as depicted. To inhibit NOS activity, mice received intraperitoneal injection of NMMA (20 mg/kg/day; Sigma-Aldrich, Gillingham, Dorset, UK) or Cavitratin (2.5 mg/kg/day, Enzo LifeScience), or vehicle, for 2 weeks, 5 days/week. All animal work was in accordance to the Home Office and The Francis Crick guidelines.

Intravital Two-Photon Microscopy

For intravital imaging in time-lapse, mice were anesthetized with 2.5% Isoflurane, the head shaved and held in a stereotaxic skull holder. A skin incision revealed the calvaria and methylcellulose (4%, Sigma-Aldrich, Gillingham, Dorset, UK) was applied to prevent the tissue drying. For imaging at a single time point, mice were euthanized by cervical dislocation, the skin was removed, and the intact skull was immerged in PBS for direct examination of the calvarium under the microscope. Images were obtained on a Zeiss 710 NLO laser scanning multiphoton microscope with a 20x 1.0 NA water immersion lens. The microscope was equipped with a MaiTai “High Performance” fully automated 1-box 517 mode-locked Ti:Sapphire laser with DeepSee dispersion compensation (Spectra-Physics), tuned to 890 nm excitation wavelength. To assess the hypoxia with hypoxisense probe, mice received an i.v. injection of 0.4 mg of human normal immunoglobulin twice (48 and 24 hours prior to imaging) (Gammalplex, Bio Products Laboratory, Hertfordshire, UK). The HypoxySense 680 probe (ref: NEV11070, PerkinElmer, Massachusetts, US) was injected 24 hours prior imaging (according to the manufacturer instructions), and on the day of imaging each mouse was intravenously injected with 100 μl of human CD45 AF488 antibody (Biolegend, Cat:304017) 30 minutes prior sacrifice, or with 150 kDa TRITC-dextran in case of NSG-Nestin-GFP mice. Mice were imaged as described above. Bone signal (Second Harmonic Generation, SHG) was collected at 380-485nm, hCD45 or Nestin-GFP at 500-550nm, TRITC-dextran at 555-625nm and Hypoxisense at 640-690nm with a 750 nm Infra red filter by not descanned detectors. To assess vascular perfusion, 100 μg of IB4-AF488 (Life Technology, Cat: I21411) were i.v. injected into mice 30 min prior imaging. 15 μl of Qtracker® 655 Vascular Labels (Thermo Fisher Scientific, Cat: Q21021MP) were injected i.v. 1 min prior imaging. SHG was collected at 380-485nm, IB4-AF488 at 500-550nm, and Qtracker at 640-690nm by not descanned detectors. To assess the hypoxic state of the vasculature, NSG-Nestin-GFP mice were used, and TRITC-dextran (150 kDa) was intravenously injected 1 min before imaging to label the vascular tree. SHG was collected at 380-485nm, Nestin-GFP at 500-550nm, TRITC dextran at 555-625nm and Hypoxisense at 640-690nm with a 750 nm Infra red filter by not descanned detectors. To image the vascular leakiness, we used the strategy detailed in Figure S2E. Briefly, two vessel-pooling agents of different molecular weights were intravenously injected simultaneously: Qtracker® 655 Vascular Labels (Thermo Fisher Scientific, Cat: Q21021MP), high molecular weight and 65-85 kDa TRITC dextran, medium molecular weight (Sigma-Aldrich, Gillingham, Dorset, UK). Mice were imaged at different time points (Figure 2C) after probes injection, or 10 min after probes administration in all other cases. Calvarium was imaged. SHG was collected at 380-485nm, GFP, 65-85 kDa TRITC dextran at 555-625nm, and 655- Qtracker at 640-690nm by not descanned detectors. Each z stack of images (100-150 μM) was rendered in 3D using Imaris software (Bitplane). At least 9 images representing different areas of bone were taken per group.

Images Processing

To measure vessel mean diameter, the “filament” algorithm was applied using the signal intensity derived from the NT-Qdot in the BM vasculature (Imaris, Bitplane). To measure the distance to hypoxic areas of BM vessels, a volumetric iso-surface was generated using the signal intensity derived from the Hypoxisense. A map of distances from the Hypoxisense iso-surface was calculated with “Distance Transformation” tool using Imaris XT extension. A volumetric iso-surface to define the vessels was then generated using the intensity derived from the dextran. The distance between each vessel and the closest hypoxprobe pixel was measured (Bitplane).

To assess the vascular leakiness, the signal intensity derived from the Qtracker was used to generate a volumetric iso-surface of the BM vasculature, using the standard surface algorithm of IMARIS. “Background Subtraction” (5-10% based on local contrast) was performed with manual threshold. The iso-surface was then used to segment the signal intensity (SUM) derived from the dextran in “IN” and “OUT” the vasculature, using the “Mask Settings” function. The ratio between these two intensities (OUT/IN) represents...
the leakiness of the vasculature at a given time point (Imaris, Bitplane). To measure the distance of AML cells to vessels, a volumetric iso-surface was generated using the signal intensity derived from the dextran. A map of distances from the Hypoxisense iso-surface was calculated with “Distance Transformation” tool using Imaris XT extension. The spot function was used to identify hCD45+ cells. The distance between each spot and the closest vascular pixel was measured (Bitplane). To quantify the hypoxia, in each z-stack the hypoxisense signal intensity and the total volume of the z-stack (voxel) were calculated. The hypoxia was expressed as signal intensity/voxel (Imaris, Bitplane).

**Bioluminescence Imaging**

Before imaging, each mouse was given an intraperitoneal injection of 125 mg/kg luciferin (Caliper Life Science). General anesthesia was induced with 2% isoflurane and the mouse was placed in a light-tight heated chamber. After acquiring bright-field images of each mouse, anterior and posterior luminescent images were then successively acquired with 1- to 5-min exposure time. Optical images were displayed and analyzed with the IVIS Living Image software (Caliper Life Science). Regions of interest were manually drawn around the bodies of the mice to assess the intensity of the bioluminescence signal for each mouse. Optical signal was expressed as photon count.

**Cell Lines**

293 HEK were grown in DMEM; HL60, ML1 and U937 were grown in RPMI1640, MS5 in IMDM. All cell lines were tested for mycoplasma prior to commencing experiments. All cell lines came originally ATCC (distributor LGC standards, UK) and were grown by our cell service at the Institute. Before use these lines, they were authenticated using the Short Tandem Repeat (SRF) profiling. All media were supplemented with 10% FBS and 1x Penicillin-Streptomycin and all reagents were from Gibco®-Life Technologies (Paisley, UK).

**Transduction and Culture of Human Primary Samples**

T-cell depleted hAML MNCs were pre-stimulated in StemSpam SFEM (StemCell Technologies, Vancouver, Canada) supplemented with 25 ng/ml G-CSF or 100ng/ml SCF, 100ng/ml Flt3-L, 60ng/ml IL-3 and 10ng/ml TPO (PrepoTech, London, UK), respectively for 4 to 6 hours. CD34-enriched HSPCs were pre-stimulated in StemSpam SFEM (StemCell Technologies, Vancouver, Canada) supplemented with 25 ng/ml G-CSF, 150ng/ml SCF, 150ng/ml Flt3-L, 10ng/ml IL-6 and 20ng/ml TPO (PrepoTech, London, UK) for 4 to 6 hours. Lentiviral supernatant (GFP-Luciferase vector for human AML MNCs and tomato vector (Pan et al., 2008) for HSPCs) was added at a multiplicity of infection (MOI) of 50 in the presence of 5μg/mL polybrene for 12 hours. The cells were then washed and expanded in culture. The efficiency of transduction was analyzed after 4 days by GFP expression. Primary hAML cells were co-cultured on a layer of primary human MSC for one week. Transduced AML cells were co-cultured on MS-5 cells for 4 days before sorting on the human CD45 and GFP fraction and injected into mice. Cells were grown in Myelocult H5100 medium (StemCell Technologies, Vancouver) supplemented with 20ng/ml IL-3, 20ng/ml G-CSF and 20ng/ml TPO (PeproTech, London, UK). Transduced HSPC cells were grown in StemSpam SFEM supplementer with 150 ng/ml SCF, 150 ng/ml Flt-3 and 20 ng/ml TPO for 4 days before sorting on the human CD45 and tomato fraction and injected into mice.

**Transduction and Culture of Murine Primary Samples**

Primary murine AML carrying the MLL-AF9 or MLL-ENL translocations were generated as reported (Horton et al., 2009). Leukemic cells retrieved from BM or spleen of primary leukemic mice (MLL-AF9 #1, MLL-ENL #1 and #2) were cultured in RPMI supplemented with 10% FCS, 10 ng/ml IL3, 10 ng/ml IL6, 100 ng/ml SCF (Sigma-Aldrich, Gillingham, Dorset, UK). Cells were infected by spinoculation with a lentivirus expressing GFP-Luciferase (MLL-AF9 #1 and MLL-ENL #1) or tomato (MLL-ENL #2) (centrifugation at 700g, 25°C, 45 minutes) in the presence of 5μg/mL polybrene (Sigma-Aldrich, Gillingham, Dorset, UK), as previously described (Horton et al., 2005).

**AML and HSPCs Transplantation Assay In Vivo**

**Murine AML**

Murine AML cells from culture were transplanted into sub-lethally irradiated C57BL6 mice. Secondary AML cells retrieved from these animals were transplanted into unconditioned WT or Nos3-KO mice at the indicated concentrations.

**Murine Normal HSC**

For HSC in vivo functionality assays, murine CD45.1 cells retrieved from NSG mice were isolated using an EasySep murine CD45 Positive Selection kit (StemCell Technologies, Vancouver, Canada) and 250000 cells were transplanted i.v. into sub-lethally irradiated Rag2-KO/GammaC-KO secondary recipients.

**Xenografts**

For xenografts assays, human HSC lines, AML patient-derived samples and HSPC were transplanted into 8- to 12-week old unconditioned NSG or NSG-NESTIN-GFP and NSG-NG2-DSRED mice by i.v. injection, as indicated. For competitive transplantation experiments, 5,000 tomato+ HSPCs were injected together with 100000 untransduced AML6 cells i.v. into unconditioned NBSGW mice (Jackson Laboratory, BarHarbor, Maine, USA). BM engraftment was assessed by BM aspirate or bioluminescence, when indicated.
Bone Marrow Cell Isolation

At the end of each experiment, animals were euthanized by cervical dislocation and the 6 rear bones collected in cold PBS. To retrieve BM endothelial and mesenchymal cells, bones were dissected in small pieces of 2-3 mm diameter and incubated 30 min at 37 degrees in a HBSS digestion solution containing 2% FBS, DNase I (10 µg/ml), Collagenase (1 mg/ml), Dispase II (6 mg/ml) and Heparin (20 U/ml), all from Sigma-Aldrich, Gillingham, Dorset, UK. Bone pieces were next crushed with mortar and pestle in the same digestion solution and incubated at 37 degrees for 30 min. Cell suspension was then homogenized with a micropipette and filtered with a 100 micron cell strainer.

HO Diffusion

Mice were administrated with two injections of 0.4 mg HO (Sigma-Aldrich, Gillingham, Dorset, UK) at 5 minutes interval, and euthanized 5 minutes later. After cervical dislocation, bones were quickly dissected and immediately immersed in ice-cold PBS containing inhibitors of the APC-efflux pump (Reserpine, Verapamil; Sigma-Aldrich, Gillingham, Dorset, UK) to avoid HO extrusion. Cells were recovered and processed as described above and stained with specific surface antibodies. HO uptake was calculated by dividing the geometric mean fluorescence intensity of the HO blue channel by the same analysis carried out on the same population of cells from the same organ but from an animal not injected with HO (negative control). This corresponds to the specific fluorescence intensity and reflects the actual uptake of HO by the population of interest.

Hypoxiprobe Assessment

Pimonidazole (PIM; Hypoxyprobe) was purchased from Natural Pharmacia International, Inc. Mice were i.v. injected with 125 mg/kg PIM or saline solution and euthanized after 2 hours. For flow cytometry analysis, bones were rapidly harvested and kept in PBS at 4 °C. Cells were recovered and processed as described above, and stained with specific surface antibodies. Next, they were permeabilized (BD Cytofix/Cytoperm kit) and stained with specific FITC-MAb1, following manufacturer instructions. Cells from PIM non-injected animals but stained with the same concentration of relevant antibody after permeabilization were used as a negative control to calculate the specific geometric mean fluorescence intensity as previously reported (Ito et al., 2013). For IHC analysis, samples were processed as described below, and stained with primary FITC-MAb1 and secondary anti-FITC-HRP following manufacturer instructions.

Flow Cytometry Analysis and Cell Sorting

Flow cytometry analysis was performed using a Fortessa flow cytometer (BD Biosciences, Oxford, UK). AML transduced cells were identified based on their GFP expression. Dead cells and debris were excluded from the analysis using 4,6-diamidino-2-phenylindole (DAPI) staining. Murine HSC (SLAM) were defined by the expression of these surface markers: CD45+LIN-Kit+CD48-CD150+, with the identification based on their GFP expression. Flow cytometry analysis was performed using a Fortessa flow cytometer (BD Biosciences, Oxford, UK). AML transduced cells were injected animals but stained with the same concentration of relevant antibody after permeabilization were used as a negative control to calculate the specific geometric mean fluorescence intensity.

Drug Treatments In Vitro

Human ML1 cells were seeded alone or on a confluent layer of MS-5 cells for 24 hours before starting the treatments. NMMA was used at a concentration of 5 mM for 48 hours before assessing apoptosis. Primary AML-derived samples were seeded on a confluent layer of MS-5 cells for 4 days before starting the treatment. Drugs were added to the culture (NMMA at 5 mM and SNAP at 10 µm; Sigma-Aldrich, Gillingham, Dorset, UK) for 5 days before assessing apoptosis and cell cycle.

Reverse Transcriptase and Real Time Quantitative PCR (RT-qPCR)

Total RNA from sorted cell samples was isolated using the RNeasy Mini Kit (Qiagen, Crawley, UK) according to the manufacturer’s instructions. mRNA was reverse transcribed by Superscript III Reverse Transcriptase (Invitrogen) with an oligoDT primer (Sigma-Aldrich, Gillingham, Dorset, UK). RT-qPCR was performed with an ABI 7500 FAST Thermal Cycler (Applied Biosystems) using SYBR Green dye (Applied Biosystems). RNA was quantified with the Comparative CT Method and GAPDH was used as a housekeeping gene. The CT values used are the result of triplicates. The primers used are described in the Key Resources Table.

Histological Processing and Immunostaining

Murine samples. Harvested humerus samples were fixed o/n in 10% neutral buffered formalin and then decalcified with 17% EDTA (Osteosoft, Millipore) during 7 days. Samples were processed, paraffin embedded and sectioned (5 µm) for histological studies. Hematoxylin/eosin was performed first to assess quality of the sections of BM samples and visualize vessel integrity. For
immunofluorescence (IF) studies heat antigen retrieval was performed in all cases. Primary antibodies are listed in the key resources table. Goat anti-mouse/rabbit/rat secondaries antibodies coupled to AF-488, AF555 and AF647 were used (all from Invitrogen). DAPI was included in the mounting media to label the nuclei. Images were obtained with a Zeiss LSM710 upright confocal microscope. At least 3 images were taken per sample group. To quantify micro-vascular density (MVD), the number of vessel sprouts/mm² was counted (Bitplane), and data are show as fold over ctrl.

Sample Preparation for RNA-Sequencing
The quality and concentration of total RNA were determined on Agilent 2100 Bioanalyzer using the Eukaryote Total RNA Pico Assay. Most of the total RNA has average RIN number 5-8, with a concentration at least 4 pg/µl. Some of the samples were concentrated on a speedvac without heat to obtain a final volume of 5 µl. 3.5 ng of Total RNA in 5 µl volume was used to generate cDNA synthesis with Nugen Ovation® RNA-Seq System V2 kit (part No. 7102). The resulting SPIA-cDNA were normalized to 100 ng in 15 µl based on Qubit DNA HS assay. Fragmentation was done using 8 microTUBE-15 AFA Beads Strip V2 (PN 520159) on Covaris E-series at setting 20% DF, 18W, 200 burst, 20C temp, 120s treatment time and no Intensifier. Only 10-30 ng in 10 µl of fragmented cDNA was used starting from the End Repair step of the Ovation® Ultralow Library Systems V2 (part No. 0344NB) protocol, with 10 cycles PCR. The RNAseq libraries were quality checked on Qubit DNA HS assay, Bioanalyser and Illumina Ecoreal QPCR followed by Illumina PE51 sequencing on HiSeq 2500 V3 reagents.

RNA-Sequencing Analysis Methods
FastQ files were generated using CASAVA BCL to FastQ (version 2.16). Sequencing yield was typically ~25 million strand-specific paired-end reads. The RSEM package (version 1.2.29) (Li and Dewey, 2011) in conjunction with the STAR alignment algorithm (version 2.5.1b) (Dobin et al., 2013) was used for the mapping and subsequent gene-level counting of the sequenced reads with respect to mm10 Ensembl genes downloaded from the UCSC Table Browser 15 on 14th April 2016 (Karolchik et al., 2004). The “--forward-prob” parameter was set to “0” and all other parameters were kept as default. Differential expression analysis was performed with the DESeq2 package (version 1.10.1) (Love et al., 2014) within the R programming environment (version 3.2.3) (R Development Core Team, 2015). Genes with FDR <0.05 were judged to be differentially expressed. Genes from each given pairwise comparison were ranked using the Wald statistic. GSEA pre-ranked analysis was performed with respect to MSigDB (version 5.1) (Subramanian et al., 2005) gene sets C2 canonical pathways and C5 GO biological process. All parameters were kept as default except for enrichment statistic (weighted), min size (5) and max size (50000).

Statistical Analysis
Statistics were performed with Prism (GraphPad). Unpaired Student’s t-test was used to compare two groups, except in Figure 5E and S3B and S5B, where paired Student’s t-test was used. Log-rank test was used for survival curves. Details of each analysis are in figure legends. For probabilities distribution in our cohort of patients, we have considered the joint probability distribution of two Boolean variables: NO normalization and response to therapy. We applied Bayesian inference to estimate the probabilities of each of the four possible combinations of these variables from the cohort of patients. These probabilities are related to the occurrence of each combination of NO normalization and response to therapy, according to a multinomial distribution parameterized by the four probabilities. We estimated these parameters from their posterior distributions obtained by updating a uniform prior using the data available. The posterior distributions for the four probabilities have been obtained by generating a sample of the parameters of the multinomial (N=1.10⁶), weighted according to importance sampling. The generated sample was then used to calculate relevant conditional probabilities and their corresponding posterior distributions, which allow the calculation of maximum a posteriori (MAP) estimates and 95% credible intervals (CIt95). The analysis was performed using custom-made software written in R (R Development Core Team, 2015). A similar analysis was performed including only intermediate risk group patients.

DATA AND SOFTWARE AVAILABILITY
Data Resources
The accession number for the transcriptome sequencing data generated in this study are deposited on Geo bank (reference number: GSE88905).