VEGF-ablation therapy reduces drug delivery and therapeutic response in ECM-dense tumors

F Röhrig1,2,3,9, S Vorlová2,9, H Hoffmann1,2,3,9, M Wartenberg4, FE Escorcia5, S Keller1, M Tenspolde1, I Weigand1, S Gätzner2, K Manova6, O Penack1, DA Scheinberg5, A Rosenwald4, S Ergün1, Z Granot8 and E Henke1,2,3

The inadequate transport of drugs into the tumor tissue caused by its abnormal vasculature is a major obstacle to the treatment of cancer. Anti-vascular endothelial growth factor (anti-VEGF) drugs can cause phenotypic alteration and maturation of the tumor's vasculature. However, whether this consistently improves delivery and subsequent response to therapy is still controversial. Clinical results indicate that not all patients benefit from antiangiogenic treatment, necessitating the development of criteria to predict the effect of these agents in individual tumors. We demonstrate that, in anti-VEGF-refractory murine tumors, vascular changes after VEGF ablation result in reduced delivery leading to therapeutic failure. In these tumors, the impaired response after anti-VEGF treatment is directly linked to strong deposition of fibrous extracellular matrix (ECM) components and high expression of lysyl oxidases. The resulting condensed, highly crosslinked ECM impeded drug permeation, protecting tumor cells from exposure to small-molecule drugs. The reduced vascular density after anti-VEGF treatment further decreased delivery in these tumors, an effect not compensated by the improved vessel quality. Pharmacological inhibition of lysyl oxidases improved drug delivery in various tumor models and reversed the negative effect of VEGF ablation on drug delivery and therapeutic response in anti-VEGF-resistant tumors. In conclusion, the vascular changes after anti-VEGF therapy can have a context-dependent negative impact on overall therapeutic efficacy. A determining factor is the tumor ECM, which strongly influences the effect of anti-VEGF therapy. Our results reveal the prospect to revert a possible negative effect and to potentiate responsiveness to antiangiogenic therapy by concomitantly targeting ECM-modifying enzymes.

Oncogene (2017) 36, 1–12; doi:10.1038/onc.2016.182; published online 6 June 2016

INTRODUCTION

Treating solid tumors is complicated by the inadequate transport of systemically administered drugs into the tumor tissue.1 A wide range of antitumor agents, both cytotoxic drugs and newer targeted therapies, has been shown to accumulate at much higher concentrations in non-target organs than in the tumor.2–6 Furthermore, the distribution of drugs within the tumor is heterogeneous, leaving many tumor cells protected from therapeutically effective drug concentrations.7–11

The abnormal vasculature observed in many tumors has been discussed as a major factor in reducing tumor drug delivery.12,13 Tumor blood vessels are often tortuous, leaky, strongly ramified, poorly supported by pericytes and lack a hierarchical structure, which causes an often insufficient and uneven blood flow. This aberrant vessel phenotype results from an imbalanced, strongly pro-angiogenic microenvironment. Agents that dampen the angiogenic signaling, such as inhibitors of the vascular endothelial growth factor (VEGF) pathway, improve blood vessel functionality. In preclinical models, increased pericyte coverage, reduced interstitial pressure and diminished vessel leakiness are observed after antiangiogenic therapy.14–16 The improved vessel quality should indeed increase drug delivery into the tumor, and this ‘vessel normalization’ hypothesis may explain why antiangiogenic drugs that show little efficacy as monotherapy in patients improve response to concomitant chemotherapies.17 However, antiangiogenic drugs were developed to inhibit the formation of the tumor vasculature, and reduction in vessel density is also a well-documented result of antiangiogenic treatment.16,18,19 Reduced vessel density decreases transport into the tumor and increased tumor hypoxia is commonly reported after VEGF ablation.19 Few studies have carefully examined the consequences of antiangiogenic therapy not on surrogate markers but on tumor drug delivery, with the contrarian results that the effects could be either positive or negative.20–25 It can be assumed that whether the ‘improved vessel quality’ or the ‘reduced vessel density’ effect predominates after antiangiogenic therapy may be affected by many factors. Among those are: the dosage of the antiangiogenic drug, the duration of the treatment, and characteristics of the particular tumor. However, to date, we still lack stratifying criteria to predict how antiangiogenic therapy affects drug delivery in individual tumors.

To be effective, an antineoplastic drug must not only be transported into the tumor but it also has to reach the individual tumor cell at a critical concentration. Although the effectiveness of drug transport into the tissue is determined by vascular characteristics, transport within the tissue occurs by diffusion.

© 2017 Macmillan Publishers Limited, part of Springer Nature. All rights reserved 0950-9232/17
www.nature.com/onc
The rate of diffusion within the tissue depends on the characteristics of the extracellular matrix (ECM). Tumors are often characterized by increased deposition of ECM proteins. Intriguingly, the ECM can compromise up to 60% of the tumor volume. Furthermore, subsequent enzymatic modification can turn the tumor ECM more rigid. The negative impact of the dense and rigid tumor ECM on the diffusion of macromolecular agents has been established.26,27 Less is known on its effect on small-molecule drugs. Although composition, quantity and modification of the ECM differs widely between different tumors, experimental data on how this influences drug efficacy and outcome of therapy are limited. Moreover, the likely interconnection between vascular parameters, which can be affected by antiangiogenic treatment, and diffusion controlling ECM parameters has not been studied in detail.

Here we describe a model of syngeneic, implantable fibrosarcoma that is resistant to monoclonal antibody mG6-31-driven VEGF-ablation therapy.28 Blockade of VEGF in this model reduces microvessel density, increases tumor hypoxia and impairs drug delivery leading to reduced responsiveness to the chemotherapeutic drugs. We demonstrate that these resistant tumors express high levels of lysyl oxidases, resulting in a high degree of ECM crosslinking and impeded drug permeation by increasing the ECMs’ barrier function. These results highlight a strong interconnection between vascular and ECM parameters in their influence of drug transport in tumors. We show for the first time that inhibition of lysyl oxidases substantially improves drug delivery and potentiates the therapeutic response to drugs used to treat malignant diseases.

Furthermore, alteration of the ECM via lysyl oxidase inhibition reverts the potential negative effect of anti-VEGF therapy on drug delivery, and combination of lysyl oxidase inhibition with anti-VEGF therapy synergistically improves response to cytotoxic treatment.

RESULTS
MT6 sarcomas are resistant to VEGF ablation
In contrast to the situation in patients, most murine tumors are responsive to VEGF ablation and react with significant growth reduction. Only few examples for murine tumors that react poorly to anti-VEGF therapy are known.29,30 Interpretation of results from combination treatments in responsive models is difficult as both anti-VEGF and the chemotherapeutic are individually effective in reducing tumor growth. Thus an increased effectiveness by combining both is impossible to pinpoint to a specific mechanism. Consequently, the effects of anti-VEGF therapy on drug delivery and efficacy have to be studied in models that are unresponsive to VEGF-ablation monotherapy.

Murine fibrosarcomas established by implanting MT6 cells into C57Bl/6 J mice31 failed to respond to anti-VEGF treatment with mG6-31—an antibody that binds both the human and the murine form of VEGF-A.28 When given at 5 mg/kg body weight (BW) every 6 days, mG6-31 did not significantly reduce tumor growth (Figures 1a and b). An effective doubling of the dosage (7.5 mg/kg BW, q4d) did not improve the outcome (Supplementary Figure S1). Although tumor growth was not significantly affected, mG6-31 treatment...

Figure 1. MT6 sarcomas are resistant to VEGF ablation but react with vascular changes. (a, b) Treatment of established MT6 tumors with mG6-31 (5 mg/kg BW i.p., on days 12 and 18 indicated by black arrowheads) did not affect tumor growth, measured using a caliper or by weighting excised tumors. (c) Sections of tumors were stained for pan endothelial cell antigen (PanEC). Treated tumors display drastically reduced vessel density. Amount of PanEC-positive vessels was counted in whole tumor sections (n = 4). Scale bar: 1.00 mm. (d) After mG6-31 treatment, remaining vessels showed increased coverage with NG2-positive pericytes, which was again quantified after imaging whole double immunofluorescent (PanEC and NG2) stained tumor sections (n = 5). (e) Treated tumors showed stronger staining for CA IX, indicating increased hypoxia (scale bar: 50 μm, n = 5).
strongly reduced microvessel density (Figure 1c), improved vessel maturation (indicated by increased pericyte coverage, Figure 1d) and increased staining for the hypoxia marker carbonic anhydride IX (CA IX; Figure 1e). Taken together, these results demonstrate that the mG6-31 antibody is effective in the MT6 tumors at causing the expected vascular changes, though the tumors fail to react with a reduction in growth. This mimics the situation observed in patients and makes the MT6 tumors one a rare good model to follow the effect of anti-VEGF treatment on drug delivery.

VEGF ablation impedes efficacy of chemotherapy in MT6 tumors. Because growth of the MT6 sarcomas is not affected by mG6-31 treatment, it is a suitable model to study the effect of anti-VEGF treatment on drug delivery and efficacy. The different cancers of mesenchymal origin, typically grouped together as sarcomas, respond poorly to systemic treatment with chemotherapies (reviewed in Constantinou et al.32). Addition of VEGF ablation to the chemotherapeutic regimen could potentially improve the situation of sarcoma patients. We devised a combination treatment of well-established MT6 tumors with mG6-31 and doxorubicin, the most commonly used therapeutic in the management of sarcomas.33 In contrast to the expectations, combination of mG6-31 (5 mg/kg BW, q6d) with doxorubicin (5 mg/kg BW, on days 3 and 5 of a 6-day cycle) was less effective in reducing tumor growth than chemotherapy alone (Figures 3a and b). We tested the vascular changes caused by VEGF ablation reduces the dye dispersal in tumors after mG6-31 treatment (Figures 2g and h). H33342 distribution analysis revealed reduced accumulation in treated tumors (Figure 2f). When the experiment was repeated on day 22 postimplantation used as free base or as liposomal formulation (Figures 2d and e).

To further elucidate the reasons for the differential effect of mG6-31 pretreatment in the two tumor models, we first verified that there were no intrinsic differences in sensitivity to the chemotherapeutic between 4T1 and MT6 tumor cells (Supplementary Figure S3). The observed differential effect on drug delivery can also be caused by differences in vascular density or vessel permeability in the tumor models. Thus mG6-31 treatment of 4T1 and MT6 tumors was repeated in parallel, and the tumors were excised 4 days after a single dose of mG6-31. Staining for the vessel marker CD31 revealed more positive staining in MT6 tumors (Figures 3f and g). However, microvessel density was not different (Figure 3h) as vessels were larger in the MT6 tumors than in 4T1 tumors. A detailed 3D analysis of perfused blood vessels stained with isoclectin GS-IB4 showed that blood vessels in MT6 tumors are larger, more tortuous and more ramified (Figures 3j and k). Treatment with mG6-31 led in both models to a pruned down, less chaotic vasculature. Per fused vessel density was similarly decreased in both tumor models after VEGF ablation (Supplementary Figure S4). Thus neither the vascular parameters before treatment nor the changes affected on these parameters by VEGF ablation indicated a cause for the differential effect on drug delivery in the two tumor models. However, a striking difference between 4T1 and MT6 tumors was the extent of H33342 penetration into the tumor parenchyma (Figures 3i and m). In untreated 4T1 tumors, the dye penetrated much deeper from the vessels into the surrounding tissue (21.30 ± 2.005 μm) than in MT6 tumors (10.99 ± 0.7584 μm). Consequently, the tumor volume that was supplied with H33342 in relationship to the vessel surface area was much higher in 4T1 tumors (Figure 3n). mG6-31 treatment improved this ratio in 4T1 tumors even further, while the changes in the MT6 tumors were insignificant. In summary, mG6-31 strongly reduces the perfused vessel density in MT6 sarcomas without significantly improving drug penetration into the tumor parenchyma. This results in less tumor cells being reached and affected by the delivered cytotoxic drug, reducing treatment efficacy. In the 4T1 model, better tissue penetration is further enhanced after mG6-31 treatment, explaining why after mG6-31 treatment a larger proportion of the tumor is supplied with drugs despite reduced vessel density. These results indicate that tissue permeability is one of the factors that determine whether VEGF ablation improves delivery and response to concomitantly applied drugs.

The ECM of non-responsive MT6 tumors is less permeable for small-molecule drugs. Possible causes for the different tissue permeability are differences in the ECM amount or its composition between MT6 and 4T1 tumors. Trichrome and picrosirius red staining of MT6 tumor sections showed homogeneous deposition of collagenous matrix forming a fine interconnected network surrounding individual cells (Figure 4a). Conversely, in 4T1 tumors collagen is often associated with blood vessels, the network forming large irregular clusters of cells. We next isolated ECM from both tumors. Cellular components were removed by high salt extraction (Figure 4b) and the remaining mainly insoluble material was re-suspended in buffer, yielding a suspension of the complete, native ECM components (cECM).
Figure 2. VEGF ablation leads to therapeutic failure in MT6 fibrosarcoma. (a–c) Established MT6 sarcomas were treated with mG6-31 (5 mg/kg BW, i.p.) or a control antibody (IgG) and consecutively with two rounds of either doxorubicin or doxil (note: data from a six-armed study was divided into two graphs for clarity). Tumors did not react significantly to mG6-31 treatment alone, and animals pretreated with mG6-31 did respond less to cytotoxic treatment. Treatment days are indicated with black (mG6-31) and red (doxorubicin/doxil) arrowheads. (d–f) MT6 tumors treated with mG6-31 and injected with doxorubicin, doxil or 3H-paclitaxel. mG6-31 pretreatment reduced accumulation of all three drugs in the tumor. (g) 3D angiography using Alexa647-labeled Isolectin (IL-GS-IB4) shows reduction in perfused vessel density in mG6-31-treated tumors. (h) Distribution of H33342 in whole tumor sections was also reduced after mG6-31 treatment (n = 4). * indicates statistical significance of doxorubicin/doxil treatment groups versus double treatment groups, *P < 0.05, **P < 0.01. All error bars: ± s.e.m.

Figure 3. Effect of VEGF ablation on drug distribution and drug tissue penetration is context dependent. (a) 4T1 tumors were treated with mG6-31 or a control IgG and consecutively treated with two rounds of doxorubicin. Treatment days are indicated with black (mG6-31) and red (doxorubicin/doxil) arrowheads. (b) Weight of 4T1 tumors excised 26 days after implantation and 14 days of treatment. (c, d) Pretreatment with mG6-31 did not change amounts of doxorubicin or 3H-paclitaxel delivered to 4T1 tumors and normal organs. Drug accumulation assays were performed on day 14 after tumor implantation, 2 days after the initial mG6-31 treatment. (e) Distribution of H33342 in whole tumor sections was improved after mG6-31 treatment. (f) Immunostaining for CD31 in 4T1 and MT6 tumors treated with mG6-31 or an IgG control antibody (scale bar: 100 μm). (g, h) Quantification of relative area stained positive for CD31 and quantification of microvessel density in MT6 and 4T1 tumors treated with mG6-31 or control IgG. Isotopic GS-48 staining of perfused vessels (2-projections: 45 slides each with 0.98-μm spacing, scale bar: 100 μm) showed a more tortuous and convoluted vessel network in the MT6 tumors, mG6-31 reduced the degree of ramification in MT6 tumors. (k–m) Quantification of H33342 penetration depth from vessel surface and tissue volume supplied with detectable amounts of H33342 in correlation to vessel surface area in MT6 and 4T1 tumors treated with mG6-31 or control IgG. * indicates statistical significance versus control, # indicates statistical significance versus single treatment groups, *P < 0.05; **P < 0.01; ***P < 0.001. All error bars: ± s.e.m., n = 4 if not otherwise indicated. NS, not significant.
Alternatively, we extracted the ECM precipitate with urea buffer, to obtain the urea soluble fraction of the tumors’ ECM. MT6 tumors yielded significantly higher amounts of cECM proteins (Figure 4c); however, the extracts from 4T1 tumors proved to be more soluble in the urea buffer. We coated membranes of transwell inserts with these extracts and tested the permeability for doxorubicin.

**Figure 4:**
- **a:** Time course of MT6 tumor growth. Doxorubicin was given as indicated. Error bars indicate standard errors. 
- **b:** Tumor weight of antibody-treated and control tumors. Error bars indicate standard errors. 
- **c:** Comparison of tumor weight gain of antibody-treated and control tumors. Error bars indicate standard errors. 
- **d:** Percentage of CD31 positive area. Error bars indicate standard errors. 
- **e:** Percentage of CD31 positive area. Error bars indicate standard errors. 
- **f:** Immunohistochemistry for CD31 in MT6 and 4T1 tumors. 
- **g:** Immunohistochemistry for IL-GS-IB4 in MT6 and 4T1 tumors. 
- **h:** Percentage of CD31 positive area. Error bars indicate standard errors. 
- **i:** Immunohistochemistry for IL-GS-IB4 in MT6 and 4T1 tumors. 
- **j:** Percentage of CD31 positive area. Error bars indicate standard errors. 

© 2017 Macmillan Publishers Limited, part of Springer Nature. Oncogene (2017) 1 – 12
Lysyl oxidase inhibition improves drug delivery into tumors

The reported redundancy of the lysyl oxidase family members in their biological function and substrate spectra, as well as our results that expression of individual family members are highly diversely expressed in tumors, suggested that a successful treatment strategy by reducing ECM crosslinking should be based on an inhibitor that is effective against all five lysyl oxidases. Thus, to test whether inhibition of lysyl oxidase activity could improve drug delivery in vivo, we treated established tumors with the pan-lysyl oxidase inhibitor \( \beta \text{APN} \). The treatment was performed by intraperitoneal (i.p.) injections with 100 mg/kg BW \( \beta \text{APN} \) in saline\(^{38} \) and did not affect the growth of MT6 tumors, whereas the growth of 4T1 tumors was reduced (Figure 5a).

However, doxorubicin concentrations after a single bolus injection were found to be significantly improved in both tumor models after treatment, having nearly doubled in the treated MT6 tumors (1.596 ± 0.3361 versus 0.8257 ± 0.1391 \( \mu \text{g} / \text{g} \) in control animals, Figure 5b). In other organs, doxorubicin accumulation was not significantly changed after \( \beta \text{APN} \) treatment. Injected Hoechst 33342 was more evenly distributed throughout the tumor after \( \beta \text{APN} \) treatment (Figures 5c and d). Improved oxygenation resulted in reduced expression of the hypoxia-inducible factor 1a-regulated genes CA IX and VEGF-A (Figure 5e). Recently, it was reported that LOX positively regulates VEGF-A expression and thereby stimulates angiogenesis.\(^{39} \) However, in contrast to this reported direct effect of LOX activity on VEGFA expression, in vitro neither 4T1 nor MT6 cells reacted to \( \beta \text{APN} \) with changed VEGF-A mRNA expression levels (Supplementary Figure S8). These results demonstrated that the reduced levels VEGF-A in the treated tumors are indeed an effect of improved oxygenation. Finally, improved supply of the tumors was indicated by a markedly reduction in central necrosis after \( \beta \text{APN} \) treatment (Figures 5f and g). To examine whether the improvement in drug and oxygen supply to the tumors was indeed caused by a better permeability of the ECM after LOX(L) inhibition, we isolated cECM from both tumors after treatment. Total interference reflection microscopy\(^{40} \) demonstrated the expected strong reduction of crosslinked collagen fibrils in the cECM of treated tumors (Supplementary Figure S7). In transwell-coating experiments, cECM preparations showed significantly better permeability for doxorubicin after \( \beta \text{APN} \) treatment, indicating that lysyl oxidases have a critical role in forming a chemo-protective barrier (Figure 5h).

Lysyl oxidase inhibition reverses the negative impact of VEGF ablation on drug delivery in MT6 sarcomas

After having established that inhibition of lysyl oxidases improved drug delivery and distribution, we tested whether this affects therapeutic success in a combination therapy setup and might even reverse the negative effect of VEGF ablation on drug delivery in MT6 fibrosarcomas. Animals with established MT6 tumors were pretreated with \( \beta \text{APN} \) before starting doxorubicin-based therapy (Figure 6a). Treatment with \( \beta \text{APN} \) significantly improved tumor response toward chemotherapy (average tumor size 270.9 ± 57.8 versus 518.6 ± 75.1 \( \text{mm}^3 \)). Addition of mG6-31 to the \( \beta \text{APN} / \text{doxorubicin} \) regimen further improved
therapeutic efficacy (92.8 ± 37.2 mm³). In an additional experiment, we verified that combination of mG6-31 and βAPN did not affect tumor growth (Supplementary Figure S7). In addition to excluding a simple additive effect in the triple treatment group, the results of the study also demonstrated that the refractoriness of the MT6 tumors towards VEGF ablation is not caused by their dense, highly crosslinked ECM.

To test whether the improved response to chemotherapy after combining lysyl oxidase inhibition with antiangiogenic therapy is caused by improved drug delivery, we treated established MT6 sarcomas for the short period of 4 days with βAPN in combination with a single dose of mG6-31 (Figure 6b) and examined distribution of Hoechst 33342 within the tumor. In tumors treated with the combination of βAPN
and mG6-31, the Hoechst dye indeed permeated significantly deeper into the tumor tissue than in control tumors and in tumors from mice that received βAPN alone (Figure 6c).

Doxorubicin given as a bolus dose also accumulated significantly stronger in the tumors treated with the βAPN/mG6-31 combination (Figure 6d).

Figure 5. Lysyl oxidase inhibition improves drug accumulation and distribution within tumors. (a) Treatment of established MT6 and 4T1 tumors with βAPN (100 mg/kg BW i.p., treatment days are indicated by arrowheads). (b) Doxorubicin accumulation in βAPN-treated MT6 and 4T1 tumors versus control tumors and in normal organs of βAPN-treated MT6-bearing C57Bl/6 J mice versus organs from control animals. (c) 3D confocal micrographs of MT6 and 4T1 tumors injected with H33342 and IL-IB4-A647 after βAPN treatment (Z-projections: 30 slides each with 0.90-μm spacing, SB: 100 μm). (d) Quantification of the H33342-positive area in MT6 tumors after βAPN treatment (n = 4). (e) mRNA levels for the hypoxia marker CA IX and for VEGF-A in MT6 and 4T1 tumors after βAPN-treatment (n = 5). (f) Whole mount sections of MT6 and 4T1 tumors display a strong reduction of central necrosis after APN treatment (NA = necrotic area, SB = 1.00 mm). (g) Quantification of necrotic areas in MT6 and 4T1 tumor sections (n = 8). (h) Relative permeability of usECM extracts obtained from control and APN treated MT6 and 4T1 tumors. Transwell inserts were coated with 3 μg/mm² of respective usECM extracts and tested for doxorubicin permeability (n = 3). * indicates statistical significance versus control. All error bars: ± s.e.m.
DISCUSSION

Combining drugs that inhibit the VEGF pathway, such as Avastin, with chemotherapeutic treatment regimens is meanwhile an integral part in the management of various malignant diseases. Nevertheless, the therapeutic benefit of adding anti-VEGF agents observed in large patient cohorts is only moderate, indicating that not all patients benefit equally from these agents. We need a better understanding of how these agents influence transport and...
efficacy of cytotoxic drugs and in which context they are most effective. This will enable us to predict which patients will benefit from adding vascular-targeted therapies to their treatment and in which patients antiangiogenic therapy might be detrimental. The refractoriness of MT6 tumors to anti-VEGF therapy does not only reflect the situation in patients but also permits unperturbed observation of secondary effects, such as drug transport and effectiveness. Recently, Van der Veldt et al. have demonstrated that bevacizumab (Avastin) leads to decreased perfusion and delivery of docetaxel in non-small-cell lung cancer patients. However, the experimental setup in this study did not allow for assessment of whether therapeutic efficacy was also reduced or whether non-small-cell lung cancer patients would still benefit from the combination by a possible additive effect. The MT6 model displays the same response to anti-VEGF that is observed in these patients, and our results indicate for the first time that such a reduction in drug delivery after anti-VEGF therapy might indeed compromise therapeutic activity. If VEGF ablation leads to impaired delivery and distribution of cytotoxic drugs but has no measurable antitumor effect by itself, combination of both might indeed be less effective than the cytotoxic drug alone. However, if the antiangiogenic drug itself inhibits tumor growth, a combination with a cytotoxic drug may still prove beneficial, as Cesca et al. reported for vandetanib that also reduced paclitaxel transport into xenografts, but a combination of both was still better than each therapeutic alone.

In our experiments, the vascular changes after VEGF ablation improved response to chemotherapy in 4T1 breast carcinomas. The varying results using anti-VEGF/cytotoxic combination therapy indicate that the effect of antiangiogenic therapy is indeed strongly context dependent.

Our results experimentally underline the inter-relationship between vessel characteristics that control tumor perfusion and matrix properties that regulate drug diffusion within the tissue. In the MT6 model, the key factor leading to reduced accumulation and effectiveness of cytotoxic drugs after mG6-31 treatment was clearly the dense, highly crosslinked ECM. The MT6 matrix was altered nearly impermeable to small-molecule drugs by high lysyl oxidase levels, an effect further corroborated in several in vitro experiments. Consequently, only in combination with lysyl oxidases inhibition, VEGF ablation was able to improve drug delivery into MT6 tumors. In addition to lysyl oxidases, it is likely that also other ECM-modifying enzymes and the composition and quantity of ECM components themselves have a strong impact on drug permeability and thereby on the effect of antiangiogenic therapeutics. Besides the drastically different lysyl oxidase levels, the differences in ECM content and composition probably contributed to the differential drug permeability observed in 4T1 and MT6 tumors. Hyaluronic acid has been shown to impede drug diffusion by a very different mechanism: as it binds large amounts of water, the freshly formed ECM, it will alter its characteristics, including its remodeled. When lysyl oxidases are inhibited and cannot crosslink the ECM’s quality only after more chronic inhibition. It is known that lysyl oxidase activity is necessary to maintain functional and mature connective tissue. Its inhibition by substances such as βAPN can cause diseases, such as lathyrism. Targeting tumor-specific upstream regulators of lysyl oxidase expression might be an alternative way to improve ECM characteristics. Recently, Liu et al. reported that inhibition of transforming growth factor-β in experimental mammary carcinomas improved delivery of doxorubicin and doxil. Transforming growth factor-β is a known positive regulator of all lysyl oxidase family members.

Similar to the defective vasculature, the dense tumor ECM impairs transport and distribution. Both characteristics thereby simultaneously protect the tumor from treatment and contribute to a more malignant phenotype. Auxiliary drugs that change these characteristics could improve efficacy of the whole array of antitumor drugs currently at our disposal. Moreover, a better understanding of the intricate mechanisms that protect the tumor from the negative impact of our treatment efforts could lead to a strategic approach toward cancer therapy that aims at defusing those protective traits and making tumors more vulnerable.

**MATERIAL AND METHODS**

If not otherwise indicated, chemicals were purchased from Sigma Aldrich (Munich, Germany) or Carl Roth (Karlsruhe, Germany). βAPN was purchased from Sigma Aldrich. Protein concentrations were determined with the Pierce BCA Kit (Thermo Fisher, Rockford, IL, USA) using a 30-min incubation time at 60 °C. The anti-VEGF antibody mG6-31 and a control IgG (anti-Ragweed) were provided by Genentech (San Francisco, CA, USA).

**Cell culture**

MT6 (CRL-2805), 4T1 (CRL-2539), MDA-MB-231 (HTB-26), MDA-MB-435S (HTB-129), MCF7 (HTB-22), ZR-75-1 (CRL-1500) and LLC (CRL-1642) cells were obtained from ATCC (Manassas, VA, USA). AC755, EMT6 and 38290 TTT cells have been purchased from NCI Tumor Repository (http://ncifrederick.cancer.gov/Services/NcifRepositories.aspx). E0771 cells have been purchased from Tebu-Bio (Offenbach, Germany). All tumor cells were maintained in Dulbecco’s modified Eagle’s medium (Gibco/Thermo Fisher Scientific, Waltham, MA, USA) with 10% fetal bovine serum and Penicillin/Streptomycin at 37 °C, 5% CO2. Cell lines were tested for mycoplasma contamination.

**Tumor models and treatment**

All experiments involving animals were reviewed and approved by the institutional animal care and use committee at MSKCC or by the Regional Administration of Unterfranken, Würzburg, Germany. All animal experiments were performed according to ethical standards and guidelines.

**Tumor engraftment.** MT6 (106 cells in phosphate-buffered saline) fibrosarcomas were generated by subcutaneous injection in the dorsal region of female C57Bl/6 J mice. 4T1 (106 cells in phosphate-buffered saline) breast adenocarcinomas were generated by injection of cells into the inguinal mammary fat pad of female Balb/c mice. Balb/c mice were purchased from Charles River (Sulzfeld, Germany) and C57Bl/6J from Jackson Laboratories (Bar Harbor, ME, USA). All animals in the individual experiments were of the same age and sex. For each experiment, tumor-bearing mice were randomly assigned to the different treatment groups just prior to the start of treatment. In treatment
studies where tumor growth was a critical outcome, assessment of tumor size was performed blinded by a second researcher.

Exclusion of data. Animals that never developed tumors owing to take rate <100% were excluded from the studies. All data from animals that died or had to be killed prior to the scheduled termination of the experiment were excluded.

Tumor treatment. Treatment of fully established tumors started on day 12 after implantation. mG6-31 or an anti-Ragweed control antibody were given at 5 mg/kg every 6 days on days 12, 18 and 24 by i.p. injection. Doxorubicin/doxil (both at 5 mg/kg BW free doxorubicin) was administered by i.p. injection on days 14, 16, 20, 22, 26, 28, 32 and 34. Control substance for doxorubicin was 0.9% NaCl. 3-APN was administered at 100 mg/kg or 30 mg/kg BW in 0.9% NaCl by daily i.p. injection. Control substance was 0.9% NaCl.

Tumor growth was followed by measuring perpendicular diameters of the tumors with a vernier caliper. Tumor volume was calculated using the equation \( V = \pi/6 \times l \times w^2 \). In addition, tumors were excised postmortem and weighted. Only tumors that could be excised completely without additional invaded tissue were used for weight measurements.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

We thank Robert Benezra (Cancer Biology and Genetics Program, Memorial Sloan-Kettering Cancer Center) for helpful discussion; Thomas Jarchau (Institute for clinical Biochemistry and Pathobiochemistry, Univeritätsklinikum Würzburg) for a thorough review of the project; Dorothea Schwering, Katharina Schneider, Sandra Theisen, and Christiane Bruns for help with experiments. Funding was provided by the Deutsche Forschungsgemeinschaft (DFG Grant Nos. HE3655/1–1, HE3655/2–1 and HE3655/3–1 to EH).

AUTHOR CONTRIBUTIONS

FR, HH, MW, SV, FEE, SK, IW, MT, SG, ZG and EH conducted the experiments. OP, KM, DAS, AR, ZG and EH planned the experiments.

REFERENCES

1 Kratz F. Drug delivery in oncology – challenges and perspectives. In: Kratz F, Steinhagen H, Senter P (eds). Drug Delivery in Oncology – Challenges and Perspectives in Drug Delivery in Oncology – from Research Concepts to Cancer Therapy, vol. 1: VCM; Weinheim, Germany; 2011, pp LIX-LXXV.
2 Gangloff A, Hueser WA, Kesner AL, Kiesewetter DD, Pio BS, Pegram MD et al. Estimation of paclitaxel biodistribution and uptake in human-derived xenografts in vivo with 18F-fluoroplaclitaxel. J Nucl Med 2005; 46: 1866–1871.
3 Kesner AL, Hueser WA, Hett NL, Pio BS, Czernin J, Pegram MD et al. Biodistribution and predictive value of 18F-fluorocyclophosphamide in mice bearing human breast cancer xenografts. J Nucl Med 2007; 48: 2021–2027.
4 Staffhorst RW, van der Born K, Erkelens CA, Hamelers IH, Peters GJ, Boven E et al. Vascular changes after anti-VEGF therapy. Proc Natl Acad Sci USA 2010; 107: 2909–2914.
5 Chung AS, Wu X, Zhuang G, Ng H, Kasman I, Zhang J et al. An interleukin-17-mediated paracrine network promotes tumor resistance to anti-angiogenic therapy. Nat Med 2013; 19: 1114–1123.
6 Phillips PG, Yalcin M, Cui H, Abdel-Nabi H, Sajjad M, Bernacki R et al. TGF-beta blockade improves the distribution and efficacy of therapeutics in breast carcinoma by normalizing the tumor stroma. Proc Natl Acad Sci USA 2012; 109: 16618–16623.
7 Minko T, Kopeckova P, Pozharov V, Jensen KD, Kopecek J. The influence of cytotoxicity of macromolecules and of VEGF gene modulated vascular permeability on the enhanced permeability and retention effect in resistant solid tumors. Pharm Res 2000; 17: 505–514.
8 Liu J, Liao S, Diop-Frimpong B, Chen W, Goel S, Naxerova K et al. TGF-beta blockade improves the distribution and efficacy of therapeutics in breast carcinoma by normalizing the tumor stroma. Proc Natl Acad Sci USA 2012; 109: 16618–16623.
Doxorubicin induces apoptosis and CD95 gene expression in human primary endothelial cells through a p53-dependent mechanism. J Biol Chem 2002; 277: 10883–10892.

Modulation of lysyl oxidase-like 2 enzymatic activity by an allosteric antibody inhibitor. J Biol Chem 2010; 285: 20964–20974.

Variations in LOXL1 associated with exfoliation glaucoma do not affect amine oxidase activity. Mol Vis 2012; 18: 265–270.

Cellular fibronectin binds to lysyl oxidase with high affinity and is critical for its proteolytic activation. J Biol Chem 2005; 280: 24690–24697.

Doxorubicin induces apoptosis and CD95 gene expression in human primary endothelial cells through a p53-dependent mechanism. J Biol Chem 2002; 277: 10883–10892.

Hyaluronan cation of collagen networks promotes sarcoma angiogenesis and invasion. Int Orthop 2012; 36: 627–638.

Enzymatic targeting of the stroma ablates physical barriers to treatment of pancreatic ductal adenocarcinoma. Cancer Cell 2012; 21: 418–429.

Antiangiogenesis strategies revisited: from starving tumors to alleviating hypoxia. Cancer Cell 2014; 26: 605–622.

Combining two strategies to improve perfusion and drug delivery in solid tumors. Proc Natl Acad Sci USA 2013; 110: 18632–18637.

Allosteric inhibition of lysyl oxidase-like-2 impedes the development of a pathologic microenvironment. Nat Med 2010; 16: 1009–1017.

Vascular changes after anti-VEGF therapy
F Röhrig et al

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/

Supplementary Information accompanies this paper on the Oncogene website (http://www.nature.com/onc)