

Tumor distribution and efficacy of antiangiogenic receptor tyrosine kinase inhibitors

Ph.D. Dissertation

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2. ABBREVIATIONS

AC: alternate current

ACN: acetonitrile

ADME: adsorption, distribution, metabolism and elimination

AGC: automatic gain control

AI: angiogenesis inhibitor

AKT: protein kinase with transforming capabilities developed in the Ak strain of mice

ALK: activin receptor-like kinase

Ang: angiopoietin

ANOVA: analysis of variance

ARNT: aryl hydrocarbon receptor nuclear translocator

ARRIVE: animal research: reporting of in vivo experiments

BAMBI: BMP and activin membrane-bound inhibitor homologue

BCR-ABL: break point cluster- abelson

BM: basement membrane

BMP: bone morphogenic proteins

CA4P: combretastatin A-4 phosphate

CSC: cancer stem cells

CBP: CREBB binding protein

c-FMS: colony-stimulating factor-1 receptor

CHCA: α -cyano-4-hydroxycinnamic acid

CID: collision-induced dissociation

c-KIT: cellular homolog of the feline sarcoma viral oncogene v-kit

c-MYC: cellular myelocytomatosis transcription factor

COX: cyclooxygenase

CRC: colorectal cancer

CSF-1R: colony stimulating factor 1 receptor

CSL: CBF1/Su(H)/Lag-1) transcription factor

C-Trap: curved linear trap

CXCR4: SDF receptor

DC: direct current

DFG motif: Asp-Phe-Gly motif
APE motif: Ala-Pro-Glu motif
DHB: 2,5-dihydroxybenzoic acid
DLL: delta-like ligand
DTC: differentiated thyroid carcinoma
EC: endothelial cell
ECM: extracellular matrix
EGF/EGFR: epidermal growth factor/ epidermal growth factor receptor
EMA: European Medicines Agency
EPAS: endothelial PAS domain-containing protein
EPC: endothelial progenitor cell
ErbB: erythroblastic leukemia viral oncogene homologue
ERK: extracellular signal-regulated kinase
FAK: focal adhesion kinase
FDA: Food and Drug Administration
FGF/FGFR: fibroblast growth factor/ fibroblast growth factor receptor
FLK: fetal liver kinase
FLT: fms-like tyrosine kinase
FT-MS: Fourier transform-mass spectrometry
GDF: growth and differentiation factor
GIST: gastrointestinal stromal tumor
HCC: hepatocellular carcinoma
HCD: high energy collisional dissociation
HE: haematoxylin&eosin
HER: human epidermal growth factor receptor
HGF: hepatocyte growth factor
HIF: hypoxia-inducible factor
HPLC: high-performance liquid chromatography
HRE: hypoxia-response element
HSP27: heat shock protein 27
IC50: the half maximal inhibitory concentration
IGF/IGFR: insulin-like growth factor/ insulin-like growth factor receptor

IL: interleukin
ITK: interleukin-2-inducible T-cell kinase
JNK: c-Jun N-terminal kinase
KDR: kinase domain-containing receptor
LCK: lymphocyte-specific protein tyrosine kinase
LC-MS: liquid chromatography-mass spectrometry
LYN: Lck/Yes novel tyrosine kinase
LTQ: linear trap quadrupole
mAB: monoclonal antibody
MALDI: matrix-assisted laser desorption ionization
MAPK: mitogen-activated protein kinase
MEK: MAPK/ERK Kinase
MIS: mullerian inhibitory substance
MMP: matrix metalloproteinases
MS: mass spectrometry
MSI: mass spectrometry
mTOR: mammalian target of rapamycin
MVD: microvessel density
m/z: mass to charge ratio
NCE: normalized collision energy
NF- κ B: nuclear factor kappa-light-chain-enhancer of activated B cells
NICD: Notch intracellular domain
NRP: neuropilin
NSCLC: non-small cell lung cancer
PA/PAI: plasminogen activator /plasminogen activator inhibitor
PDGF/PDGFR: platelet-derived growth factor/ platelet-derived growth factor receptor
PHD: prolyl-hydroxylase domain-containing protein
PI3K: phosphatidylinositol 3-kinase
PIP2: phosphatidylinositol 4,5-bisphosphate
PKB: protein kinase B
PKC: protein kinase C
PLC γ : phospholipase C gamma

PIGF: placental growth factor
pNET: pancreatic neuroendocrine tumors
PTEN: phosphatase and tensin homolog
RAS: rat sarcoma viral oncogene homolog gene
RAF: rapidly accelerated fibrosarcoma
RCC: renal cell cancer
RET: rearranged during transfection
RTK/RTKI: receptor tyrosine kinase/ receptor tyrosine kinase inhibitors
SAPK: stress-activated kinase
s.c.: subcutan
SDF: stromal derived factor
SH2-domain: Src homology 2 domain
SEM: secondary electron multiplier
SEM: standard error of the mean
SMA: smooth muscle actin
SNP: single nucleotide polymorphism
Src: sarcoma
STAT: signal transducer and activator of transcription
TFA: trifluoroacetic acid
TGF: transforming growth factor
TIC: total ion current
TIE: tunica intima endothelial receptor
TNF: tumor necrosis factor
TOF: time of flight
uPA/uPAR: urokinase plasminogen activator/ urokinase plasminogen activator receptor
VDA: vascular disrupting agents
VEGF/VEGFR: vascular endothelial growth factor/ vascular endothelial growth factor
receptor
VHL: von Hippel-Lindau
VSMC: vascular smooth muscle cell

3. INTRODUCTION

3.1 Angiogenesis

The word “angiogenesis” was first mentioned in 1787 in the work of John Hunter, an English surgeon who studied the process in the growing antlers of deers (1). In contrast to vasculogenesis, which means the development of the vascular system during embryogenesis, angiogenesis is the process when new blood vessels are formed from preexisting ones.

Under physiological conditions, angiogenesis is activated in response to low oxygen level. The process is regulated by the HIF (hypoxia-inducible factor) complex. HIFs are basic helix-loop-helix DNA binding transcription factor proteins of the Per-Arnt-Sim family, and function as heterodimers consisting of an oxygen-regulated alpha subunit and a stably expressed beta subunit (2). The alpha subunits are encoded by three genes in mammals: HIF1 α , HIF2 α (EPAS1: Endothelial PAS domain-containing protein 1) and HIF3 α (2-4). Also three forms of the beta subunit have been identified until now, the HIF1 β (aryl hydrocarbon receptor nuclear translocator: ARNT), ARNT2 and ARNT3 (5,6). The structure, regulation and function of all HIFs seem to be similar, but compared to their homologs (HIF1 α and ARNT) the members of the HIF2 and -3 family have been reported to have a more restricted pattern of expression and thus may play more specialized roles in oxygen delivery than the HIF1 subunits (6,7).

Under normal oxygen tension, the alpha subunits are hydroxylated at the prolyl residues, a process that is catalyzed by the prolyl-hydroxylase domain-containing protein (PHD). This hydroxylation promotes interaction with the von Hippel-Lindau (VHL) ubiquitin ligase and, consequently, they undergo proteasomal degradation (8,9). Thus the half-life of alpha subunits is measurable in minutes and the protein is hardly detectable at all (10). The degradation process is regulated not only by the VHL protein, but p53 as well (11).

In case of hypoxia, the HIF alpha and beta subunits form a complex, recruits its co-activator, p300/CBP (CREBB Binding Protein) and binds to the Hypoxia Responsive Element (HRE) of the target genes, thus modifying their transcription (12). Several of these target genes are responsible for inducing angiogenesis in order to increase the oxygen delivery of the tissue, such as vascular endothelial growth factor (VEGF),

VEGF receptor 1, tunica intima endothelial receptor 2 (TIE2), angiopoietin 2 (Ang2), erythropoietin, Insulin-like growth factor 2 (IGF2), Transforming Growth Factor β (TGF β 3), c-MET, adrenomedullin, NO Synthase 2, plasminogen activator inhibitor-1 (PAI1) (3,13-21). Regulation of oxygenation via the HIF molecule is depicted in **Figure 1**.

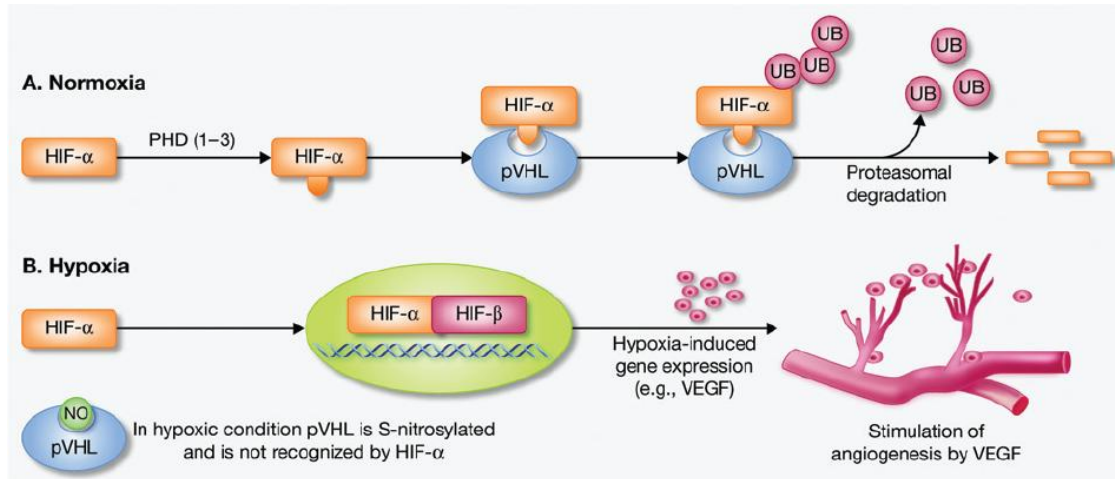


Figure 1. Regulation of oxygenation via the HIF molecule (22). Degradation of HIF α via ubiquitination in normoxia (A.) and angiogenesis stimulation in hypoxia (B.).

VEGF is considered to be the key hypoxia dependent cytokine for endothelial sprouting (23). Besides, many other factors are also able to positively influence the process of angiogenesis such as platelet-derived growth factors (PDGFs) (24-26), fibroblast growth factors (FGFs) (27,28), placental growth factor (PIGF) (29), angiopoietins (30), Jagged (31), epidermal growth factor (EGF) (32), hepatocyte growth factor (HGF) (33) and interleukin-8 (IL-8) (34). These ligands bind to their receptors on the surface of endothelial cells and act in an autocrin or paracrin manner. Binding the growth factor leads to the activation of signaling cascades, influencing the survival, proliferation and migration of endothelial cells and thus the maintenance of existing- and development of new vessels. Moreover, these pathways regulate further processes involved in angiogenesis, such as secretion of additional growth factors (35), upregulation of angiogenic receptors (36), alterations in cell-cell and cell-matrix interactions by matrix metalloproteinases (MMPs) (37), and activation of the members of endothelial cell (EC) adhesion molecule family (38). After the new capillary is formed by ECs, PDGF-BB and bFGF secreted by the endothelium recruits PDGFR β or FGFR expressing pericytes and vascular smooth muscle cells (VSMCs) (36,39). These supporting cells provide

structural stability for the new blood vessel, promote EC survival and regulate blood flow via influencing vasoconstriction and -dilatation. Moreover, via the secretion of VEGF pericytes also promote EC sprouting and survival (40). By binding to their receptor, TGF β 1 (41), Ang1 (42) and sphingosine-1-phosphate (43) stabilize the interaction between mural cells and ECs.

The healthy body controls angiogenesis by balancing a series of angiogenesis stimulating and inhibiting factors. Thus, in a healthy adult ECs have long half-life, and angiogenesis is only activated under special conditions, such as the reproducible processes of women or wound healing.

If tissues cannot produce adequate amounts of angiogenic growth factors, blood vessel growth becomes inadequate leading to improper circulation and eventually to necrosis. Insufficient angiogenesis occurs in diseases such as coronary artery disease, stroke and chronic wounds. In these pathological features therapeutic angiogenesis is aimed to stimulate the sprouting of new vessels with growth factors being developed to treat these conditions.

On the other hand, by overexpressing the angiogenesis stimulating factors, blood vessel growth becomes highly intensive in some pathological conditions, such as endometriosis, atherosclerosis, psoriasis, rheumatoid arthritis, inflammation, ischaemia, ocular neovascularization and cancer (44).

3.2 Tumor-induced angiogenesis

After the experiments of Hunter, more than a century had passed until an interest in the angiogenesis of tumors was piqued. Until the 1930-40s these investigations were restricted mostly to the observation of the morphology of tumor blood vessels (45). Later the process of neovascularization was also observed, but it had not become the focus of research until 1971, when Judah Folkman postulated that above a certain size (about 1-2 mm diameter) intratumoral diffusion is no longer sufficient and tumors are incapable of growing and metastasizing, unless they develop their own blood supply to ensure the necessary oxygen and nutrient level of the cells. He also assumed that tumor cells secrete growth factors to facilitate the process of angiogenesis, ie. capillary growth is induced by the communication between tumor blood vessels and the tumor tissue (46).

This communication is mediated by the HIF α proteins, which are stabilized in hypoxic tumor cells, and thus facilitate the expression of angiogenic proteins. Moreover, the expression level of HIF1 α is controlled not only by oxygen tension, but also by reactive oxygen species, that are secreted in response to carcinogens (47,48).

Moreover, oncogenes, such as activated EGFR (49), ErbB-2/Her2 (50) mutant RASv12 (51), mTOR (52) and Src (53) also augments HIF mediated gene expression. On the other hand, some tumor suppressor genes also play an important role in the regulation of HIF1 α , therefore loss of function of these gene products may influence angiogenesis in tumor tissues. The product of the tumor suppressor gene PTEN attenuates hypoxia mediated HIF1- α stabilization through the inhibition of AKT (54), while pVHL (9) and p53 (11) regulate the ubiquitination and degradation of the HIF complex, thus mediating the expression of angiogenic molecules.

Furthermore, solid tumors and their microenvironment secrete and often overexpress a range of growth factors, cytokines and hormones that coordinate the complex series of events of new capillary growth. Among them ie. VEGFA (55), IL1 β , insuline (56), heregulin (50), EGF (57), IGF1,-2 (13) TGF β (58), Tissue Necrosis Factor α (TNF α) (59) enhance HIF expression, activating a vicious circle in the angiogenic process.

HIF independent regulation of the VEGF pathway also occurs in tumors, as some oncogenes and tumor suppressor genes (cSrc (60), BCR-ABL (61), RAS (62), p53 (63)); cellular receptors (activated EGFR (64), IGF-1R (65) and overexpressed HER2 (66)); and cytokines (COX-2 (67), PDGF-AA (68)) can also influence VEGF production. These changes in the HIF-VEGF protein system drive the activity of angiogenesis in tumors, which can be further modulated by the expression of other angiogenic growth factors.

The schematic process of tumor-induced angiogenesis is depicted in **Figure 2**.

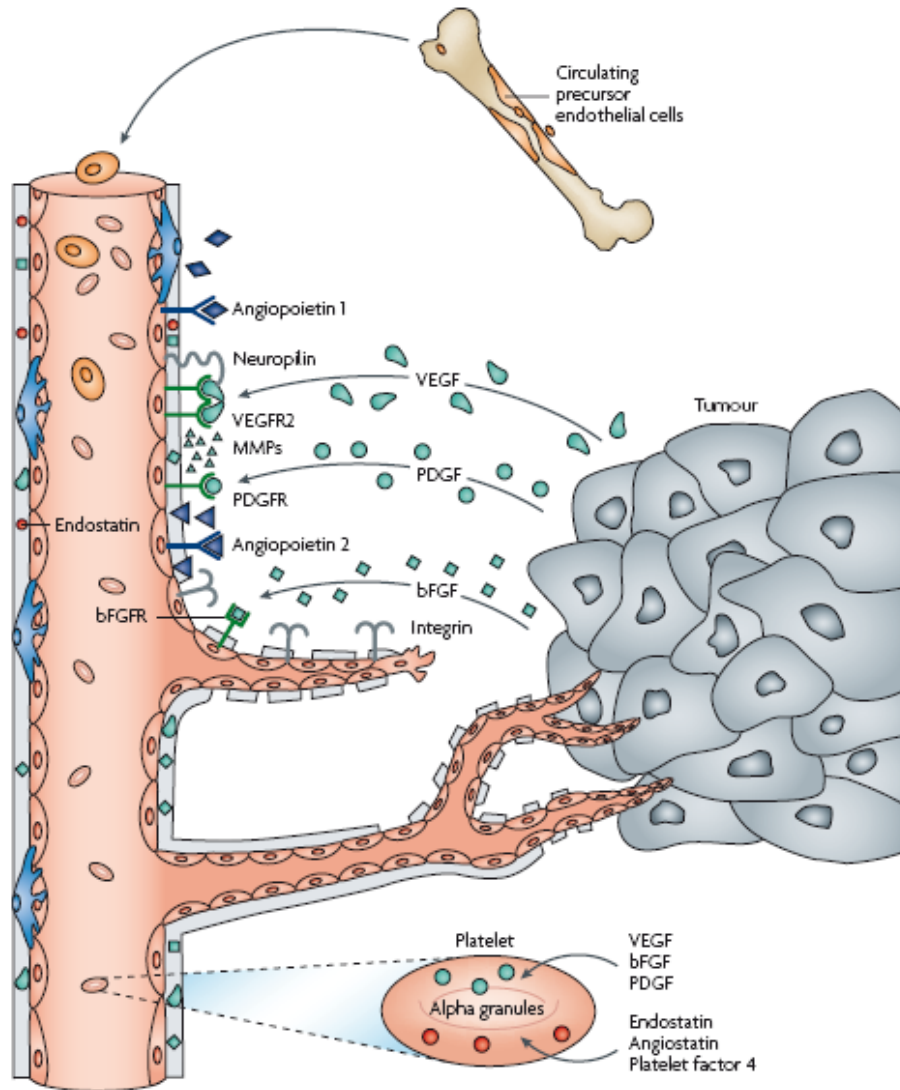


Figure 2. The process of tumor-induced angiogenesis (69). Main angiogenic growth factors (VEGF, PDGF, bFGF) are secreted by tumor cells or platelets and bound to their transmembrane receptors on ECs. The process is further regulated by integrins, MMPs and intrinsic angiogenesis inhibitors, such as endostatin, angiostatin or platelet factor-4.

3.2.1 Main receptor families and signaling pathways in tumor-induced angiogenesis

Kinases are phosphotransferase enzymes that transmit the terminal phosphate group of a high-energy donor molecule, such as adenosine triphosphate (ATP), to their substrate. Protein kinases can be divided into a larger group of serin/threonine kinases and a smaller tyrosine kinase group, based on the amino acid they target. Besides, atypical kinases also exist.

Another classification is based on the localization of the kinase. Receptor kinases are membrane bound and essential for the transduction of extracellular signals into the cell. Non-receptor kinases are responsible for intracellular communication. Angiogenesis is mainly regulated by receptor tyrosine kinases (RTKs).

RTKs have a variable extracellular part for ligand binding, a transmembrane region to anchor the molecule to the cell membrane, and they all share a conserved intracellular secondary structure. This consists of (i) a bi-lobed catalytic core, responsible for binding ATP in a deep cleft, located between the lobes (hinge region); (ii) a substrate binding site, where interacting proteins can bind and (iii) an activation loop, containing Asp-Phe-Gly (DFG) and Ala-Pro-Glu (APE) motifs at the N- and C-terminal part of the loop, respectively.

The ATP-binding site is divided into the following subregions (**Figure 3.**): adenine region, sugar region and phosphate binding region, based on the part of the ATP it binds. The sugar- and phosphate binding regions form a hydrophilic channel. Besides, a hydrophobic pocket (selectivity pocket) is also formed near the ATP binding region by the activation loop in the inactive conformation. This is not used by ATP, but ensures selectivity of receptor tyrosine kinase inhibitors. Moreover, ATP does not use the hydrophobic channel either, which provides a slot to open for solvents and it can be exploited to gain binding affinity.

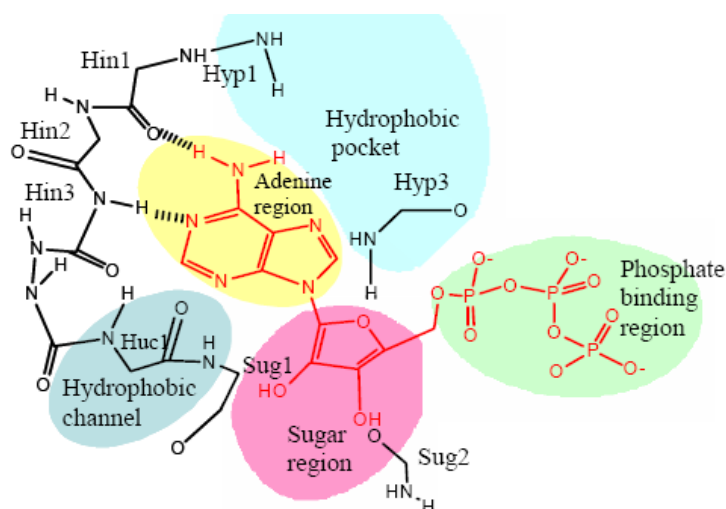


Figure 3. Binding of ATP to receptor tyrosine kinases (70). The ATP-binding site is divided into the following subregions: adenine region, sugar region and phosphate binding region, based on the part of the ATP it binds. The sugar- and phosphate binding regions form a hydrophilic channel. Note that the hydrophobic pocket and the hydrophobic channel are not used by ATP, but ensures binding affinity and selectivity of RTKIs.

The activation loop can form a number of conformations. In the "out" conformation it creates a hydrophobic pocket near the ATP-binding cleft, thus blocking the accessibility of the receptor for ATP. Ligand binding of the extracellular domain triggers receptor dimerization, resulting in transphosphorylation of specific tyrosine residues in the activation loop, juxtamembrane- and C terminal regions. As a result, the receptor turns to the "in" conformation, which opens the hinge region for ATP. ATP binds in the cleft via the adenine ring, which forms 2 hydrogen bonds with the kinase 'hinge'. The ribose and triphosphate group of ATP bind in a hydrophilic channel that extends to the substrate binding site, which is essential to the catalysis of autophosphorylation. After autophosphorylation, the receptor recruits interacting proteins that bind to certain phosphorylation sites, which subsequently phosphorylate other proteins. These activation signaling pathways eventually lead to biological responses, such as cell activation, proliferation, differentiation, migration, survival and vascular permeability, (70,71). Activation of RTKs is depicted in **Figure 4**.

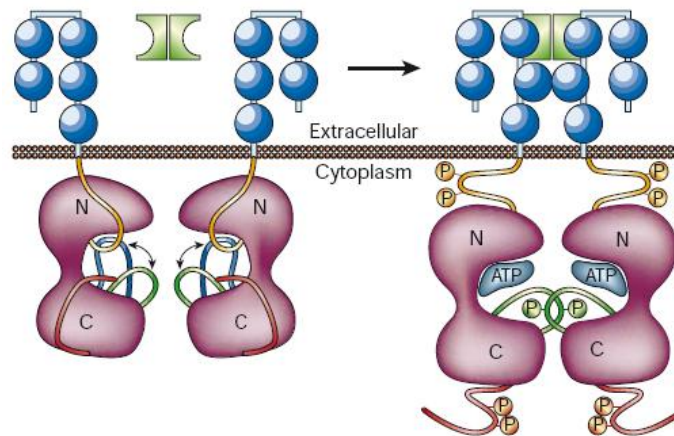


Figure 4. Activation of RTKs (71). Ligand binding of the extracellular domain triggers receptor dimerization, resulting in transphosphorylation of specific tyrosine residues in the activation loop, juxtamembrane- and C terminal regions. As a result, the receptor turns to the "in" conformation, which opens the hinge region for ATP.

The most important angiogenic RTKs are VEGFRs, PDGFRs, FGFRs, and the TIE receptors. Beside tyrosine kinases, serine-threonine kinases also regulate the process of neovascularization. These act similar to RTKs. Main examples are the members of TGF receptor family. Other receptors exert their effect not by phosphorylating partner molecules, but by regulating potential transcriptional activators. Of these, the Notch receptor family is the predominant member of angiogenic signalling.

3.2.1.1 VEGFR family

The VEGF family consists of the following growth factors in mammals: VEGFA (VEGF), VEGFB, VEGFC, VEGFD and PlGF. They are homodimeric polypeptides, although naturally occurring heterodimers of VEGFA and PlGF have also been described (72,73). Splicing and processing regulates the ability of the growth factors to bind to the appropriate receptors or co-receptors. Ligands can bind to three different receptors, VEGFR1-3 (FLT1: fms-like tyrosyl kinase-1, FLK1/KDR: Fetal liver kinase-1/Kinase Domain-containing Receptor and FLT4 respectively) (74). The receptors have a ligand-binding extracellular domain consisting of seven immunoglobulin-like loops, a transmembrane domain, a juxtamembrane domain, and a split tyrosine kinase domain followed by a C-terminal tail (75). As co-receptors neuropilin 1-2 (NRP1, -2) and heparan sulphate play a role in the regulation of angiogenesis by increasing the binding affinity of specific ligands to the receptors (76-78).

VEGFA is the ligand of both VEGFR1 and VEGFR2, but while VEGFR1 has much higher affinity for binding the growth factor, than VEGFR2, its kinase activity is weakly induced (79). Moreover, beside being highly expressed on immune cells, hematopoietic cells, vascular endothelial and -smooth muscle cells and several tumor cells (79-84), this receptor exists partly in a soluble form, thus it rather acts as a „trap receptor” for the angiogenic growth factors (85). As VEGFB and PlGF bind just to VEGFR1, their proangiogenic role is limited (86).

VEGFR2 binds VEGFA and proteolytically processed VEGFC, -D (87-89). VEGFR2 is highly expressed not only on lymphatic and vascular endothelial cells but on hematopoietic and tumor cells as well (90-94). Soluble antiangiogenic form of the receptor also exists, but as its binding affinity is much smaller than that of VEGFR1, its trap function is also less important (95).

VEGFC binds with a higher affinity to VEGFR3 than to VEGFR2, while VEGFD has similar affinity for both receptors in human, but does not bind to VEGFR2 in mouse (96,97). VEGFR3 is expressed in the venous endothelial cells in the cardinal vein during the later stages of embryogenesis, which gives rise to lymphatics, thus in adults VEGFR3 can be found mainly in lymphatic endothelium and to a lesser extent in vascular endothelial cells (98). It is disputed, whether it can be found on tumor cells (99). Soluble VEGFR3 suppresses lymphangiogenesis and lymphatic metastasis in

different cancer types (100,101). Consequently, the VEGFA-VEGFR2 axis is the predominant mediator of tumor-induced angiogenesis.

Alternative mRNA splicing of VEGFA gives rise to several distinct isoforms. The splice variants are noted as VEGF_{xxx} (xxx means the number of amino acids present in the proteins without the signal peptide) (102). These isoforms differ not only in their expression pattern but their biochemical and thus biological properties as well. Also exon 8b containing less angiogenic or antiangiogenic variants of VEGF_{xxx} are documented (103), and seem to be able to inhibit neovascularization. Moreover, they might be responsible for the failure of antiangiogenic therapies targeting VEGFs (104).

Figure 5. shows the main ligands and receptors of the VEGF superfamily.

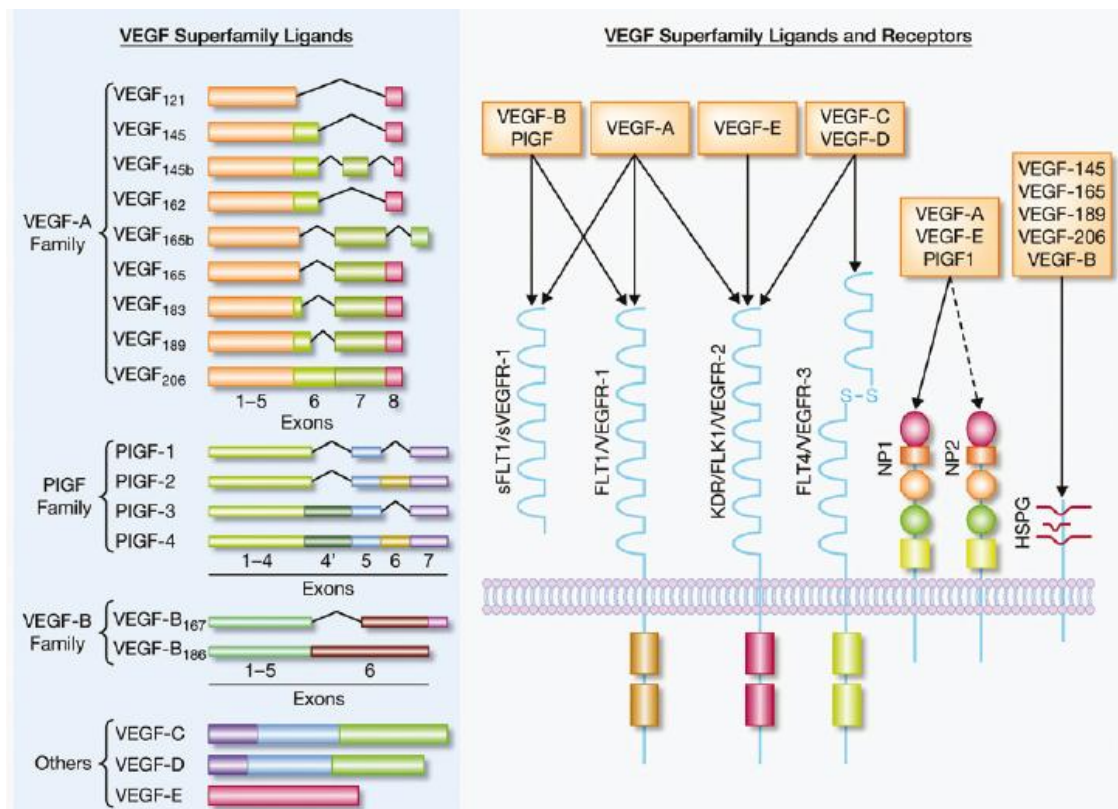


Figure 5. VEGF Superfamily Ligands and receptors (22).

VEGFs can either bind to the receptors freely or be presented by co-receptors. Ligand binding results in receptor homo-, or heterodimerization, followed by changes in the intracellular domain conformation. This leads to the liberalization of the ATP binding site of the receptor and binding of ATP results in the auto- or transphosphorilation of tyrosine residues on the receptor and downstream signal transducers (105).

The VEGF ligands are produced by most parenchymal cells, and paracrine VEGF signaling is essential for the angiogenic cascade, proliferation, survival, permeability responses and endothelial differentiation. Moreover, autocrine VEGF signaling also exists, but only conveys survival signals (106).

After ligand binding the receptor dimerizes and activates the PLC γ /Ras/Raf/MEK/ERK pathway, leading to the regulation of gene expression and cell proliferation (89). With PI3K activation they modulate AKT signaling and thus the survival of receptor expressing cells (107). Via the elevation of intracellular Ca²⁺ or NO level, induced by PI3K or PLC they can also enhance vascular permeability (108). Cell migration is regulated by the PKC/p38/MAPK2-3/HSP27 pathway and the FAK or the STAT cascade (109). Signaling pathways activated by VEGFR2 are shown in **Figure 6**.

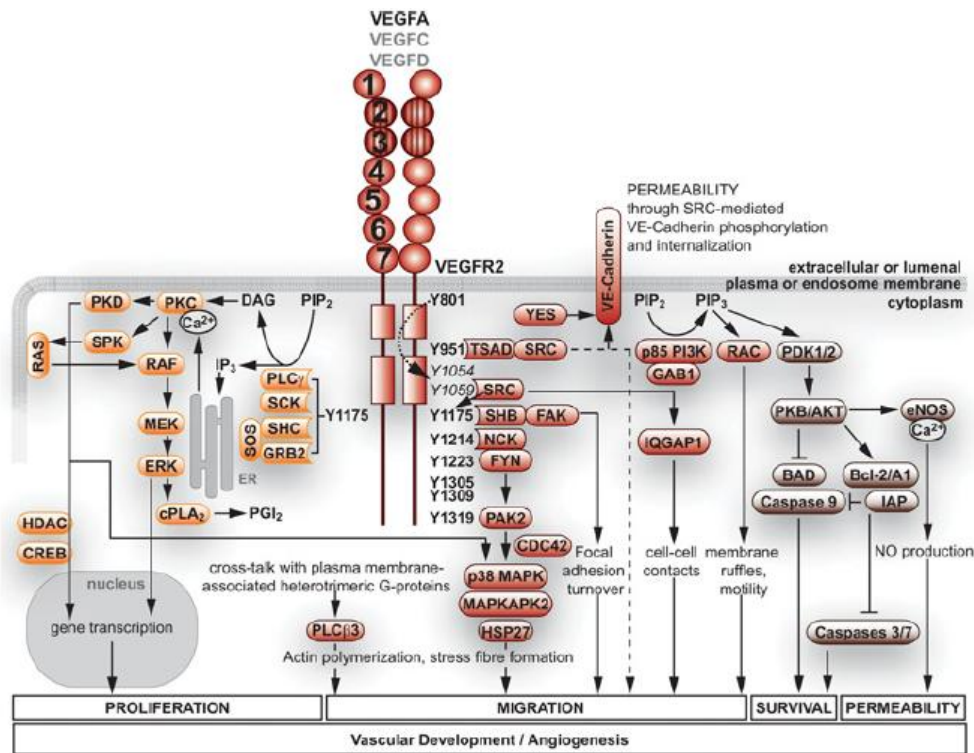


Figure 6. Signaling pathways of VEGFR2 (110).

Overexpression, mutation of the receptors and elevation of VEGF have also been related to enhanced angiogenesis and tumor development (111-113).

3.2.1.2 PDGFR family

The PDGF family consists of four ligand chains, PDGF-A, PDGF-B, PDGF-C and PDGF-D, which create five disulphide bonded dimeric isomorphs, PDGF-AA, PDGF-

BB, PDGF-AB, PDGF-CC and PDGF-DD (114). The ligands can bind to three membrane bound dimeric receptors, PDGFR $\alpha\alpha$, PDGFR $\beta\beta$, and PDGFR $\alpha\beta$ (115). Moreover, low density lipoprotein receptor-related protein and NRP1 seem to mediate PDGFR signaling as co-receptors (116,117).

PDGF synthesis is enhanced in response to low oxygen tension (118), thrombin (119), or other growth factors and cytokines (120). A number of cells secrete PDGFs, ie. cells associated with reproductive processes (121,122), vascular functions (123,124), cells of the nervous system (125-127), immune cells (128) and some other cell types (129,130).

PDGF isoforms are synthesized as precursor molecules and after dimerization they are proteolytically processed to the active forms that bind to their receptors. PDGF-AA binds only to PDGFR $\alpha\alpha$, while PDGF-BB can bind to all three dimeric receptors. PDGF-AB and PDGF-CC can activate dimeric receptors containing at least one PDGFR α monomer, while PDGF-DD those, containing at least one PDGFR β monomer. By binding to PDGF receptors, PDGFs target a broad spectrum of mesoderm-derived cells, like fibroblasts, smooth muscle cells, pericytes, mesangial- and glia cells (131-135). Soluble form of both PDGFR α and β are detected and compete with cell-associated PDGF receptors for ligand binding, antagonizing the effects of their membrane bound counterparts (136,137).

The extracellular parts of the PDGF receptors contain 5 Ig-like domains, of which domain 2 and 3 are responsible for ligand binding, while domain 4 stabilizes the dimer by a direct receptor-receptor interaction. The intracellular parts contain split tyrosine kinase domains (138). Ligand-induced dimerization of the receptors is followed by autophosphorylation, which activates their kinase activity and creates docking sites for SH2-domain-containing signaling molecules (139).

The PDGFs play crucial roles during development, and they are essential in wound healing and inflammation (140,141). Increased PDGF activity has been linked with several disorders and pathological conditions, such as atherosclerosis, fibrosis and malignant diseases (142,143). Most solid tumors secrete PDGF and express their receptors on tumor-, endothelial- or perivascular cells (144,145). Both overexpression, and different mutations of either the ligands or the receptors is associated with tumorigenesis (146-148). Moreover, all PDGF ligands are documented to be involved in the process of angiogenesis by activating signaling pathways triggering endothelial cell

proliferation (24-26). On the other hand, as pericytes and VSMCs express PDGFR β , processes driven by this receptor, such as mural cell recruitment to capillaries stabilize the vasculature, thus inhibiting tumor cell extravasation and metastatization (149).

Due to the high structural similarities with VEGFRs - all have a split kinase domain - they induce similar signaling cascades. PI3K and PLC mobilizes Ca²⁺ from intracellular stores, regulating vascular permeability (150,151). The RAS/RAF/MEK pathway is implicated in the stimulation of cell proliferation, migration, and differentiation (152). By enhancing the serine/threonine kinase AKT/PKB pathway, it mediates antiapoptotic effects (153). PDGFRs are also involved in the activation of the JNK/SAPK pathway, thus regulating cell migration (154). Binding of PDGF isoforms to PDGF receptors, with the subsequent signaling is depicted in **Figure 7**.

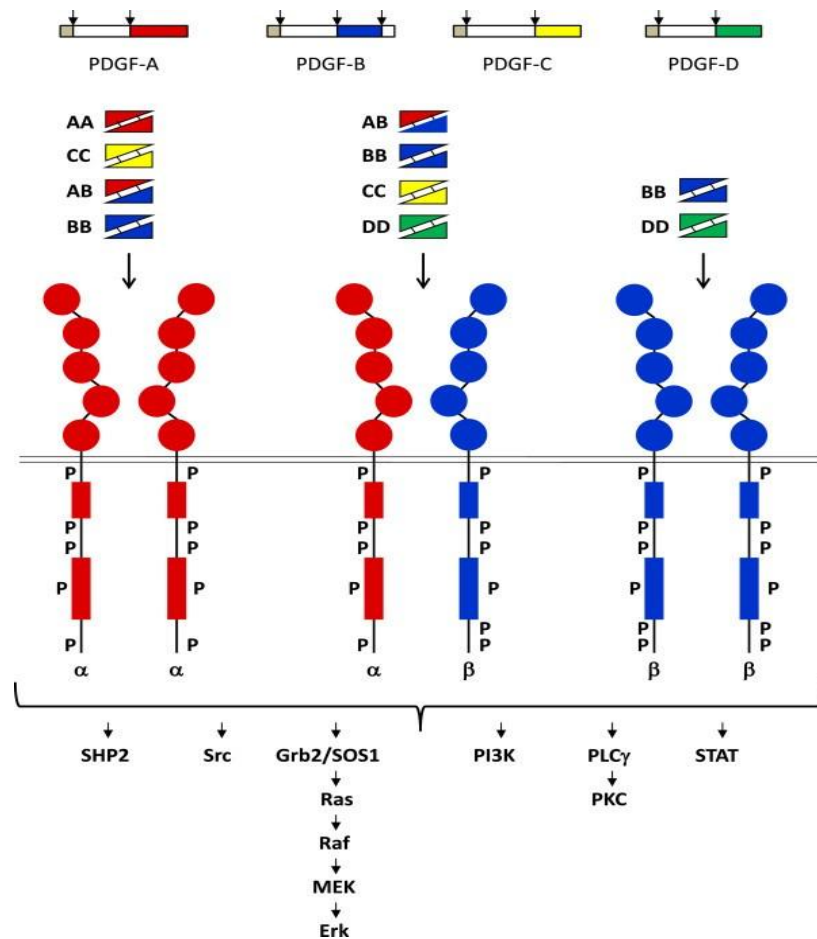


Figure 7. Binding of PDGF isoforms to PDGF receptors, and subsequent signaling (155).

3.2.1.3 FGFR family

The FGF family consists of 23 ligands, of which 18 can be secreted and bound to 4 high-affinity cell surface receptors, FGFR1-4 in mammals (156). Four FGFs do not bind

to FGF receptors, while one ligand is only expressed in mice (157). Despite most ligands act in an autocrine or paracrine manner, endocrine members of the family have also been identified (158). FGFs share a central core of 140 amino acids that is highly homologous between all family members (159,160). The ligands have strong affinity for the glycosaminoglycan side-chains of not only cell surface proteoglycans, but heparin sulphate proteoglycans and heparin-like glycosaminoglycans as well (161,162). They may protect ligands from degradation and stabilise the FGF ligand–receptor complex. A specific FGF ligand can bind to more FGFRs, although it may have a preference for a particular one (163). Ligands are produced in either epithelial or mesenchymal cells and usually activate receptors of the opposite tissue specificity.

The extracellular ligand-binding part of the receptors contains three immunoglobulin-like domains, which are important in receptor dimerisation. Different isoforms of FGFR1–3 are generated by alternative expression of the IgIII domain. Upon the presence of exon 8 or 9 they either express the "b or c" splice variant of domain III. Different isoforms are expressed in different tissues and have distinct binding specificities. FGFR4 has only one possible form (164). Truncated receptor splice variants coded by IgXa (X refers to the Ig domain being shortened by alternative splicing) are also secreted and may function as an inhibitor for FGFs (165). Structure and splicing of FGFRs are shown in **Figure 8**.

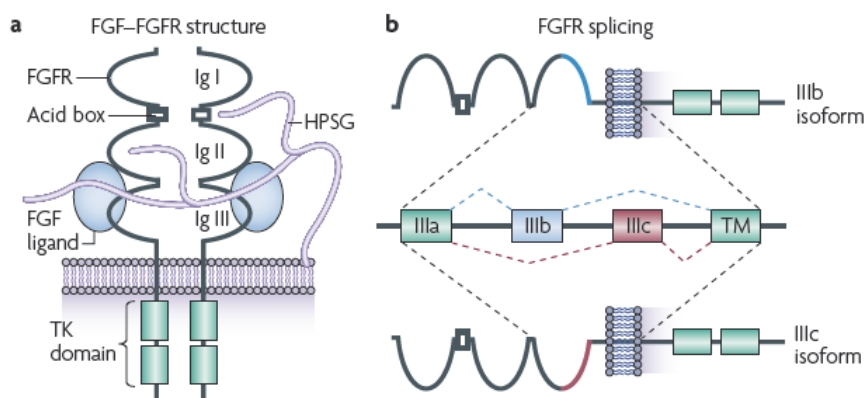


Figure 8. Structure (A.) and splicing (B.) of FGFR (166). The extracellular ligand-binding part of the receptors contain three immunoglobulin-like domains, which are important in receptor dimerisation. Different isoforms of FGFR1–3 are generated by alternative expression of the IgIII domain. Upon the presence of exon 8 or 9 they either express the "b or c" splice variant of domain III.

The FGFR signaling pathway plays a key role in embryonic development, wound healing and angiogenesis. Deregulated FGFR signalling contributes to pathological

conditions, including different malignancies. Both the receptors and the ligands are expressed by different tumor cell types (167-170). Beside translocation of FGFRs in hematological malignancies, amplification, overexpression, activating mutation or SNP of the receptors can also lead to tumorigenesis (156). Moreover, abnormalities of different FGFs are also linked to tumor progression (171,172).

Binding of bFGF (FGF2) to FGFR1IIIc is considered to be the major regulator of FGF induced angiogenesis, being the most intensively expressed both on ECs and smooth muscle cells (173). Thus it has also a diverse role, by triggering EC proliferation and stabilizing vessel walls.

Ligand binding leads to the activation of signaling cascades similar to the ones of VEGFRs and PDGFRs because of the resemblance of their kinase domain. Receptor dimerization results in the recruitment of adaptor proteins to activate RAS/RAF/MEK pathways, that regulate gene expression and cell proliferation (174). Recruitment of PI3K activates an AKT and mTOR dependent antiapoptotic pathway (175). Hydrolysis of phosphatidylinositol 4,5-bisphosphate (PIP2) by PLC γ triggers the release of intracellular Ca²⁺ (176), enhancing vascular permeability. The p38/MAPK cascade is responsible for the translocation of FGF1 to the nucleus (177), while activation of the JAK/STAT pathway, or the p70 S6 kinase also plays important roles in the regulation of FGFR signaling by mediating immunity, proliferation and apoptosis (178,179).

3.2.1.4 TIE receptor family

The family consists of two tyrosine kinase receptors, the TIE receptor 1 and 2 which can bind four ligands, Angiopoietin 1-4 (Ang1-4). The ligands are secreted glycoproteins, composed of an N-terminal superclustering domain, responsible for the creation of the higher order multimers of the ligands; a coiled-coil domain, responsible for ligand homo-oligomerization, which is essential for receptor activation and a C-terminal fibrinogen-like domain, responsible for receptor binding. The later are separated by the linker region (180,181). The extracellular part of TIE1 and TIE2 consists of three Ig-like domains that are flanked by three EGF-like cysteine repeats followed by three fibronectin type III domains. The intracellular part has a split kinase domain. Ligands bind to the Ig- and EGF-like domains of the TIE receptors (182). The structure and binding of Angs and TIE receptors are shown in **Figure 9**.

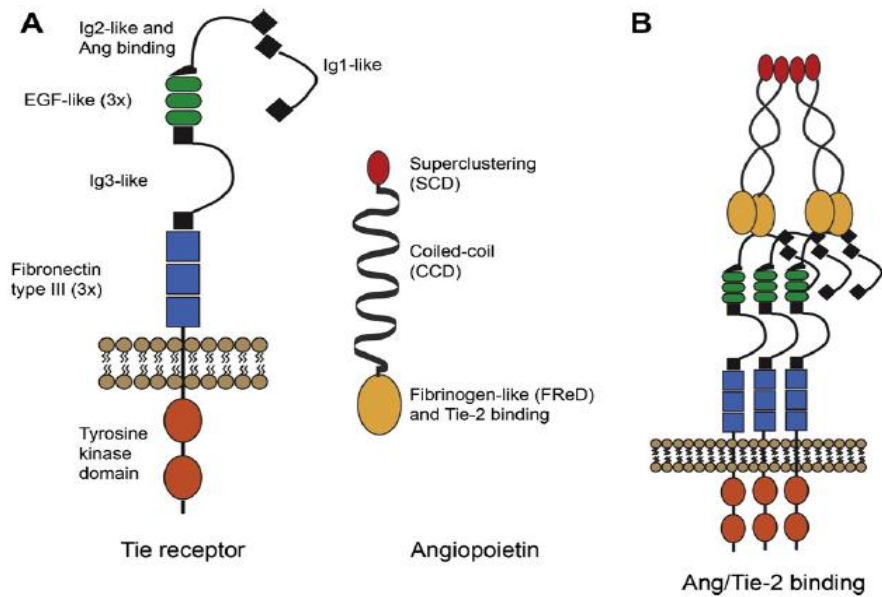


Figure 9. Structure (A.) and binding (B.) of TIE receptors and angiopoietins (183).

Ang1 is primarily secreted by mesenchymal cells, such as perivascular cells and fibroblasts, and acts in a paracrine manner on the endothelium (184,185). Ang2 is expressed by ECs (186) and retinal neurons (187). Following cytokine activation of the endothelium, it is rapidly released and acts in an autocrine manner on ECs (188). Ang1 and Ang2 bind to TIE2 with similar affinity (182). Ang1 acts as an agonist of the TIE2 receptor, activating EC survival, integrity and vessel maturation (189), whereas Ang2 is thought to be an antagonist, displacing the more active ligand Ang1. However, Ang2 has also been reported to induce receptor phosphorylation in a context-dependent manner (190). The molecular basis for these contradictory functions has not been unraveled. However, a number of factors have been implicated in controlling agonistic versus antagonistic functions (191,192). When functioning as a TIE2 antagonist, Ang2 enhances pericyte dissociation from the vessels and increases vascular permeability (193). In the absence of VEGF these unstable vessels die, but in the presence of VEGF they migrate or proliferate, depending on whether they are located at the sprouting tip or the stalk part of a newly forming blood vessel (194). Ang3 and -4 are counterparts of the same gene locus found in mouse and human, respectively, but with divergent functions. Moreover, while Ang3 is secreted by a number of mouse tissues, Ang4 is expressed to a high level only in the human lung (195).

The full length or proteolytically cleaved orphan receptor, TIE1 heterodimerize with TIE2 (196), and thus act as a co-receptor. Both TIE1 and -2 are expressed by

(lymph)endothelial cells (197,198), but TIE2 is present on hematopoietic cells and endothelial precursor cells as well (199,200). In addition to TIE receptors, angiopoietins have been found to bind integrins as well (201). Following context dependent binding of the ligands, the receptors are autophosphorylated, and intracellular signaling pathways are activated, mediating endothelial survival (202), migration, and permeability (203). Ang1 and TIE2 are shown to be essential in the recruitment of pericytes and their interaction with endothelial cells (204).

The Ang-TIE system plays a key role during vessel maturation, stabilization and remodeling, thus abnormalities of either member of the family results in diseases manifested in vascular malformations. Both Ang1 and Ang2 can be expressed by tumor cells and tumor endothelial cells (205). Despite pericyte coverage of the tumor vasculature is massively increased and thereby stabilized in response to Ang1 overexpression (206), the role of Ang1 in tumor-associated angiogenesis and metastasis remains controversial (207). Angiogenesis enhancing functions of Ang2 during VEGF or FGF induced angiogenesis has been demonstrated (208). On the other hand, an agonistic effect of Ang2 on TIE2 has also been documented (190). Furthermore, both TIE1 and -2 are present on both tumor cells and tumor ECs (209), and their expression is upregulated during angiogenesis (210,211). These findings suggest that tumor growth promoting or inhibiting functions of angiopoietins are possibly dependent on the tumor cell type and the balance between both angiopoietins and TIE1/TIE2 (190).

3.2.1.5 TGF β R family

The TGF β R family includes TGF β ligands: TGF- β 1-3, bone morphogenic proteins (BMPs), growth and differentiation factors (GDFs), activins, inhibins, leftys, nodal and the mullerian inhibitory substance (MIS). Ligands can be secreted by tumor or a number of stromal cells (212). The receptors of the superfamily include the Type I receptors: Alk (activin receptor-like kinase) receptors, and Type II receptors: TGF β RII, BMP RII, Act RII (activin type II receptor) and RIIB, and MIS RII. Co-receptors of the family include TGF β RIII/betaglycan, endoglin, BAMBI (BMP and activin membrane-bound inhibitor homologue) Crypto and Cryptic (213).

TGF β and its receptor, TGF β R are key molecules in cancer development and progression. On one hand, TGF β can inhibit tumorigenesis by suppressing cell cycle

progression and stimulating apoptosis in early stages of cancer (214). On the other hand, it can also promote cancer development and spreading by the suppression of anti-tumor immunity (215), modulation of cell invasion via modification of the microenvironment (216), induction of epithelial-mesenchymal transition (217) and degradation of the extracellular matrix (ECM) (218).

ECM and basement membrane (BM) degradation is a critical step in tumor invasion and metastasis. TGF β plays an important role in ECM degradation via upregulating MMP2 expression in different tumor types (219), and facilitates tumor cell infiltration by degrading basement membrane components via the regulation of protease expression (220). TGF β also plays a crucial role in angiogenesis by promoting endothelial cell proliferation and migration through signalling via ALK1 and by causing vessel maturation through ALK5 (41). Highly expressed ligands of the other family members, such as endoglin, can antagonize the inhibitory effects of TGF β and contribute to proliferation, migration, and capillary formation of endothelial cells (218).

3.2.1.6 Notch receptor family

Four receptors (Notch1-4), and five ligands, Jagged 1-2 and delta-like ligand (DLL) 1, 3-4 of the Notch family have been described in mammals. The receptors and the ligands are both type I transmembrane proteins, and activation requires contact of the sender and the receiver cell. The activation of Notch signaling involves three proteolytic events, resulting in the release of Notch intracellular domain (NICD) into the cytoplasm and from there it translocates to the nucleus (221). With the binding of NICD, the CSL family is converted into a transcriptional activator by displacing corepressors and by recruiting coactivators to induce transcription of Notch target genes (222).

Notch signaling plays an important role in cell proliferation, survival, apoptosis and differentiation, as their targets include proteins and factors involved in the control of cell cycle and survival (p21, cyclin D3), transcription factors (c-MYC, NF- κ B), growth factor receptors (HER/ErbB, IGF1-R) and regulators of apoptosis (survivin) (223-229). Recently, non-canonical Notch signaling that does not involve CSL or require cleavage has also been identified. This induces development via interaction with other molecules, such as members of the the IL6/JAK/STAT-, the Wnt pathway, the uPA/uPAR axis or BCR-ABL (230-233). Activation of target genes via Notch is shown in **Figure 10**.

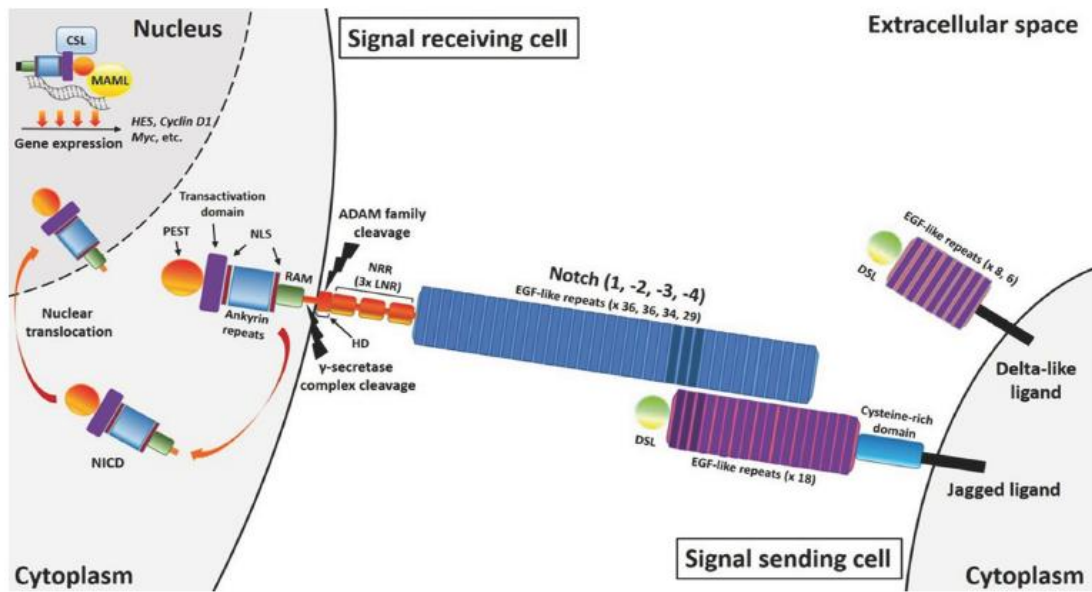


Figure 10. Activation of the Notch signaling pathway (234). Three proteolytic events result in the release of NICD into the cytoplasm and from there it translocates to the nucleus. With the binding of NICD, the CSL family is converted into a transcriptional activator to facilitate transcription of Notch target genes.

Although classically known for its role in embryonic development, the Notch pathway have been shown to mediate tumorigenesis (235) by regulating the formation of cancer stem cells (236), and epithelial- or endothelial-mesenchymal transition (237,238). Aberrant activation of Notch signaling is implicated in various neoplastic processes. Somatic gain-of-function mutations in Notch receptors have been identified first in T-cell acute lymphoblastic leukemia and later in other malignancies as well (239).

Functional studies have demonstrated the key role of Notch signaling in vascular development and postnatal angiogenesis (240), which is emphasized by the involvement of both ligands and receptors in vascular diseases. The cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL) and the Alagille syndrome can be traced back to mutations in Notch3 and Jagged1, respectively (241,242). In the endothelium, as well as in VSMCs, both Notch receptors and ligands are highly expressed (243-250). DLL4 is generally considered as a marker of arterial and tip cell phenotype (244), moreover, it is more robustly expressed in tumor ECs, compared to the neighboring normal vessels (251). Thus it is the main ligand expressed in tumor neoangiogenesis (252).

The “tip” cell is located on the leading edge of an angiogenic sprout. It is highly migratory because of the presence of filopodia and expresses high levels of VEGFR2

and -3 (253,254). Stalk cells are located adjacent to the tip cell, extend fewer filopodia but form lumen, and proliferate to support sprout elongation. The sprouting process is maintained until proangiogenic signals decrease, and quiescence is reestablished (255). It has been proposed that in response to VEGF, DLL4 is induced on tip cells. High DLL4 expression on tip cells lead and guide new sprouts, and is thought to activate Notch and suppress the tip phenotype on adjacent (stalk) ECs. As a result, Notch activity is high in stalk cells but low in tip cells, and a VEGF feedback loop is activated, whereby Notch activation causes reduction of VEGFR2 and VEGFR3 and induction of VEGFR1 on stalk cells. The absence of Notch signaling in the tip cells results in high VEGFR2 and VEGFR3 expression, causing enhanced sensitivity to external VEGF, while decreased VEGF receptor expression on the stalk cells limit sprouting and ensures the proper number of tip cells at the angiogenic front (254,256-258). Inactivation of DLL4-Notch leads to the formation of a highly branched and dense vascular network with excessive filopodia. Moreover, these vessels are often not fully lumenized, thus are unable to deliver the necessary amount of oxygen and nutrients to the tissue (252). In contrast, upregulation of DLL4 inhibits VEGF-induced endothelial cell proliferation (259) and downregulates VEGFR2 expression (260).

3.2.2 Mechanisms of tumor-induced angiogenesis

3.2.2.1 Sprouting angiogenesis

The first identified form of angiogenesis is endothelial sprouting. It was originally described by Ausprunk and Folkmann in 1977. According to their hypothesis, in response to the NO mediated VEGF signal, the postcapillary venules become dilatated, cell-cell interactions get lost, the basement membrane (BM) degrades, and thus the vessel becomes fenestrated. Endothelial cells lose their polarity and migrate to the connective tissue. Then a tube is formed from the ECs, which is followed by lumen formation, synthetization of a new BM and recruitment of pericytes (261). This hypothesis has been developed further by Paku and Paweletz in 1991. According to their model, in response to Ang2, tumor cells secrete MMPs and plasminogen activators, which digest the pericytes on the vessel wall. Consequently, the electron density of the mother vessel is changed. A gel-sol transition occurs in the BM, this is maintained by MMP2, which binds $\alpha_v\beta_3$ integrin. Growth factors are liberated from the

BM, such as bFGF and VEGF, thus inducing migration of ECs with a maintained polarity, lumen and except from tip cells, a maintained BM. As the sprouts grow the BM on the tip is continuously synthesized, pericytes are recruited in response to bFGF and PDGF and the interaction between ECs and mural cells is stabilized by TGF- β 1 and Ang1 (262). This hypothesis provides explanations missing from the former model of Ausprunk and Folkmann. These include the loosening and then regaining ECs polarity, the fact that there is de- and redifferentiation in the same process and that the lumen is formed before the synthesis of the BM, although formation of BM is known to be the facilitator for lumen formation.

3.2.2.2 Vessel incorporation or co-option

In 1987 WD Thompson raised the possibility that tumors acquire their vasculature by vessel incorporation, instead of vessel ingrowth (263). This theory was proven by Josephine Holash in 1999 (264,265). Angiogenesis by vessel incorporation usually occurs, when tumors grow or metastasize into well vascularized tissues, such as lung, liver or skin. Tumor cells grow along the preexisting, well developed vessels of the host tissue, thus annexing its vasculature. The process is faster than sprouting angiogenesis, as it does not require EC proliferation. Moreover, on the periphery of the tumor these vessels provide surface for sprouting angiogenesis. Meanwhile, in response to the locally predominant antiangiogenic factors, ECs can undergo apoptosis in the centre. This in one hand causes necrosis of the tumor mass, but on the other hand it also triggers the extravasation and metastatization of tumor cells. Maintenance of incorporated vessels is secured by the interaction of Ang1-TIE2.

3.2.2.3 Intussusceptive microvascular growth

Intussusceptive microvascular growth was first described by Sybill Patan in 1996 in a human colon adenocarcinoma model (266). This type of vessel growth is characterized by the incorporation of peritumoral capillaries, which are then separated by connective tissue pillars. The original model of Caduff has been redrawn by Paku and his colleagues in 2011. After the formation of an intraluminal endothelial bridge the BM is locally degraded, thus the EC can attach to a collagene bundle from the underlying collagene layer. The actin cytoskeleton of the EC exerts pulling force to the collagene bundle,

which in turn is transported through the vessel lumen. Finally, connective tissue cells are immigrated to the pillars and new collagenous connective tissue is deposited (267), thus the process results in lots of vessels with big lumens, providing surface for EC sprouting. In the absence of EC proliferation, the process is faster than sprouting and does not involve permeability of the vessel wall. This suggests and some experimental data also supports, that intussusception does not rely on VEGF signalization, but instead is regulated by physiologic stimuli and cytokines, that lead to vessel maturation, such as bFGF (268), PDGF-BB (269), Ang1-TIE2 complex (270), ephrin-ephrin receptor (271). However, the role of VEGF in the process is still contested (269,272,273). Moreover, it is also shown, that in response to some antiangiogenic treatment, such as vatalanib or the mTOR inhibitor sirolimus, experimental tumors switch from endothelial sprouting to intussusceptive angiogenesis (274,275).

3.2.2.4 Glomeruloid angiogenesis

In 1992 Hauro Ohtani described coiled vascular structures in human gastrointestinal carcinoma. As they resemble renal glomeruli, they were called glomeruloid structures (276). These are characterized by many, tightly associated capillary loops with different thickness of BMs. The molecular mechanism behind the process is still not clear. On one hand, Sundberg et al. suggests that VEGF is a key mediator of inducing glomeruloid body formation and maintaining these vessels (277). On the other hand, it was also shown that the glomeruloid structure is created by proliferating and migrating tumor cells, which pull the capillaries and their branching points into the tumor cell nests. Thus, this type of vascular growth cannot be termed as true angiogenesis, but rather a reorganization of tumor blood vessels, which does not require EC proliferation (278). Moreover, these vessels seem to be able to provide enough oxygen and nutrients to the cells, as no necrosis was observed in tumors developing them. As a result, glomeruloid bodies are not only diagnostic markers of glioblastoma, but also poor prognostic factors in many cancer types (279).

3.2.2.5 Postnatal vasculogenesis

Postnatal vasculogenesis refers to the process, in which bone marrow derived VEGFR2+, TIE2+, AC133+, CD34+ endothelial progenitor cells (EPCs) incorporate

into the EC layer of the tumor capillary network in response to tumor derived VEGF (280) and other proangiogenic factors. After incorporation into the EC layer, EPCs are differentiated to mature ECs. Thereafter, by producing pro-angiogenic factors, such as VEGF and PlGF, they mediate the attraction of additional EPCs to the tumor vasculature. VEGF mobilizes these cells from the bone marrow via the stromal derived factor (SDF) and its receptor, CXCR4. The process of postnatal vasculogenesis was described by Takayuki Asahara and his group both in physiologic and pathologic conditions at the end of the 1990s (281).

3.2.2.6 Vessel-like structures, formed by tumor cells

It has been shown, that not only ECs, but aggressive tumor cells can also form vessel-like structures, which facilitate tumor perfusion.

In 1941 Béla Kellner described tumor sinusoids in soft tissue sarcomas (282). These are lumens, which are covered exclusively by tumor cells, and are responsible for the transport of blood cells within the tissue.

It is also possible that tumor cells form a lumen together with ECs, without expressing endothelial or embryonal markers. These structures are called mosaic vessels and were also first described in the 1940s (283). The genesis of these types of vessels is still not well understood. They are considered to be formed either by the apoptosis of incorporated ECs, which is followed by the occupancy of the lumen or invasion of the vessel by tumor cells. The process is thus thought to be mediated via Ang2 signalling.

Vasculogenic mimicry refers to the process, when tumor cells express differential markers to completely resemble BM covered ECs. The process was first described by the group of Mary J Hendrix in uveal melanoma, where highly aggressive melanoma cells formed vessel-like channels and upregulated endothelial genes and genes involved in microvascular channel formation. Meanwhile, these cells downregulate classical melanoma markers. None of the main angiogenic cytokines, such as VEGF, PDGF, bFGF, TGF- β , Ang, Notch, TNF- α seem to induce the formation of these channels (284).

Different types of vascularization mechanisms in cancer are shown in **Figure 11**.

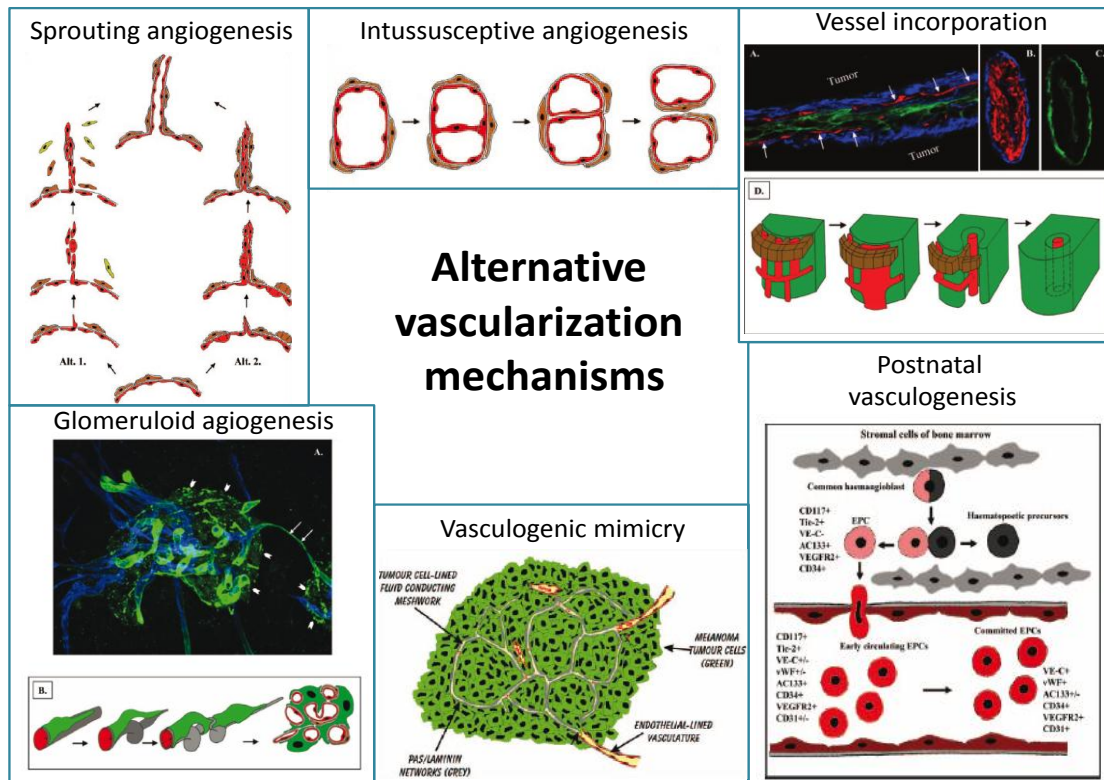


Figure 11. Alternative vascularization mechanisms in cancer (285).

3.2.3 Characteristics of tumor blood vessels

Because of the dominance of proangiogenic factors in malignant tissues, tumor blood vessels differ from the normal ones both in function and structure. In a healthy adult, ECs have a long half-life with a well differentiated BM (286), which is stabilized by pericytes and VSMCs, to influence permeability and contractility of the vessel wall. (287). Moreover, they secrete a minor amount of survival factors for the ECs, such as VEGF and Ang1. In case of Ang2 and MMP effect pericytes and VSMCs get detached from the BM and after their apoptosis, ECs also get prepared for cell death (288). In a healthy adult the endothelial tube is connected with cell-cell junctions. Maintenance of the vessel is supported by autocrin factors and signals launched by the oxygen sensors, such as the PHD2, which secures the appropriate blood flow (289).

In contrast, tumor blood vessels usually have deficient pericyte and VSMC coverage, which are not tightly attached to the ECs. They are often immature, less contractile and their shape is abnormal (290,291). The BM is often degraded and loosely attached to the EC, or it is thick and thus hinder contraction and perfusion (292). The morphology of ECs are changed and cell-cell connections are often lost (293). As a result, vessels get

dilated negatively impacting the ratio of vessel surface, which supplies the tumor tissue with oxygen and nutrients. Tumor cells can press in the instable vessel wall, thus vessel diameter becomes uneven (294). Because of this and the altered EC morphology, vessels get curved and form serpentine-like structures, and may also create high amount of anastomosis within the vessel wall (291). This is further supported by the impaired cell junctions, resulting in the loss of the signals of oxygen sensors, eventually leading to abnormal flow in the tumor vasculature (295). Because of the degraded BM, ECs lose their polarity, thus get detached from the endothelial layer and may form a plug in the lumen resulting in thrombosis and further injury of the vessel wall. This leads to enhanced metastatization capacity and the creation of mosaic blood vessels (296). Moreover, degraded mural cells and BM leads to fenestrated vessel walls, and thus not only metastatizing tumor cells may intravasate to the lumen (297), but vessels get hyperpermeabilized as well, causing an increase in the interstitial fluid pressure, which cannot be restored because of the decreased lymphatic function (298,299). Thus, the oxygen supply of the tumor decreases, resulting in an increased glycolitic activity to provide energy and the necessary building material for the growing tumor (300). The altered acidic microenvironment and abnormal morphology of tumor vessels increase the invasion and metastatization ability of malignus cells, creating a vicious circle in tumor progression. This microenvironment selects for hypoxia resistant, aggressive tumor cells (301). Because of the high intratumoral pressure, interstitial fluid, often containing tumor cells may infiltrate to lymphatic vessels, thus enhancing lymphatic tumor spread (302). Furthermore, some immune cells also promote invasion in these circumstances (303). Unlike in tumors, in case of physiological angiogenesis, these processes are balanced by the generation of antiangiogenic factors. Moreover, as a result of the inadequate blood flow, the resulting hypoxia and/or the tightened BM, efficacy of conventional anticancer therapies are inadequate due to limited penetration.

These abnormalities of tumor blood vessels and the resulting aggressiveness of cancers, accompanied by decreased efficacy of conventional treatment led Rakes Jain to the elaboration of the vessel normalization theory (304). Accordingly, the inhibition of the signal of proangiogenic factors and the subsequent normalization of tumor blood vessels has become a critical step in cancer therapy. Proposed structures of normal, tumorous, normalized and inadequate blood vessels are shown in **Figure 12**.

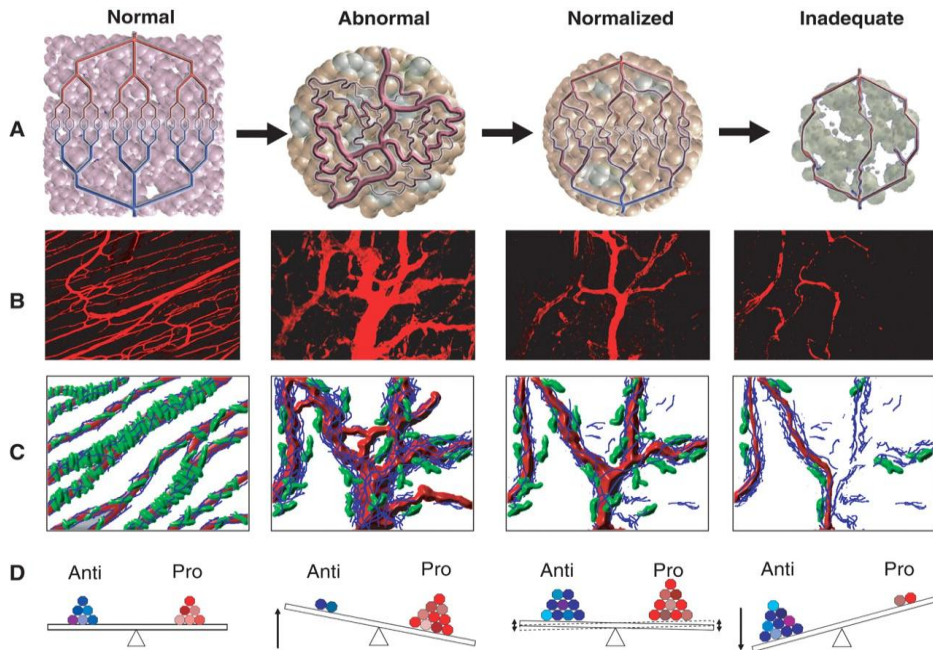


Figure 12. Proposed structures of normal, tumorous, normalized and inadequate blood vessels. (A.) Schematic structures of the vascular system. (B.) Two-photon images of normal blood vessels in skeletal muscle and subsequent images showing human colon carcinoma vasculature in mice at day 0, day 3, and day 5 after administration of VEGFR2-specific antibody. (C.) Diagrams depicting the changes in pericyte (green) and basement membrane (blue) coverage. (D.) Changes in the balance of pro-, and antiangiogenic factors in the tissue (305).

3.2.4 Inhibition of the tumor vasculature

In the complex processes outlined above, several differently acting agents are capable of blocking the blood supply of a tumor. It can be suppressed in distinct cells and also in diverse levels and modes.

3.2.4.1 Conventional chemotherapeutic agents

Conventional therapeutic agents have been found to have antiangiogenic functions as "side effect". These include microtubule targeting agents, such as Vinca-alkaloids or Taxanes, that block endothelial cell proliferation (306). Beside other drugs, thalidomide and lenalidomide also have complex antiangiogenic functions (307,308).

3.2.4.2 Vascular disrupting agents

Vascular disrupting agents (VDAs) selectively target tumor vessels, causing fast and dynamic effects (309). They destroy rapidly dividing endothelial cells in the tumor

tissue by targeting the colchicine binding site of tubulins (tubulin-binding agents) (310) or induce vascular collapse through TNF- α (flavonoid-type VDAs) (311). As a result, vascular supply shuts down, causing necrotization in response to the insufficient oxygen and nutrient delivery (312,313). By this strategy both preexisting and newly formed vessels can be targeted, with the inhibition of the metastatic potential as well. VDAs mostly act on advanced tumors, which are resistant for conventional therapy. However, their effect is very short, and can cause serious cardiac toxicities (314). Moreover, vessel density is higher at the edge of tumors, so targeting it with VDAs is not effective enough. Consequently, often there is no visible tumor shrinkage following VDA therapy. Furthermore, as tumors can also be fed by diffusion from nearby tissues, the monotherapy can leave a surviving rim at the edge of the tumor, which allows rapid tumor regrowth (315). Combretastatin A-4 Phosphate (CA4P), 5,6 dimethylxanthenone-4-acetic acid (DMXAA) and NPI-2358 are the most investigated VDAs.

3.2.4.3 Vasoactive agents

Vasoactive agents also combinatorially block existing vessels and suppress the formation of new ones. Moreover, they do not target preferentially the larger vessels in the tumor center, but also the small ones in the periphery, causing hyperabnormalization and hyperpermeability of the vessels. This allows chemotherapeutic agents better access to the tumor. However, these drugs are highly toxic when administered systemically, thus local, small dose and metronomic application is preferable.

The most often used vasoactive agents are inflammatory modulators. Among them the most important are the followings:

IL-2, a cytokine, that induces T cells, augments natural killer cell activity and demonstrates vasopermeability activity (316).

TNF- α , an inflammatory cytokine, that is principally produced by activated macrophages and monocytes and has direct effects on tumor cells. After exposure to low-dose TNF- α , hyperpermeability, hemorrhagic necrosis, extravasation of erythrocytes, edema and vessel congestion were observed (317), leading to an increased intratumoral chemotherapeutic drug concentration or effect of radiation (318). In the clinic combination therapy of melphalan and TNF- α is a popular approach to treat patients with unresectable advanced sarcoma and advanced melanoma (319).

Histamine is an inflammatory modulator that causes edema in small vessels by locally increasing the lymph flow into the extracellular space and by promoting hyperpermeability of the endothelium (320).

3.2.4.4 Angiogenesis inhibitors

Angiogenesis inhibitors (AIs) block the formation of new vessels from preexisting ones, but do not affect already established vasculature. Despite the fact that they are used in advanced tumors in the clinic, they are thought to mainly act on small vessels of the tumor edge, mostly at the early stage of tumorigenesis or metastatization.

As discussed above, endothelial sprouting is activated in physiological angiogenesis in response to hypoxia, thus, suppression of HIF1 to bind to the HRE of proangiogenic molecules is one of the main options to block the process. In the absence of selective HIF inhibitors (321) and the presence of a number of factors that also regulate angiogenic growth factor expression in tumors, the common way is to target molecules downstream of HIF by the blockade of receptor - ligand communication.

Growth factors can be targeted with either monoclonal antibodies (mABs) or soluble „trap/decoy” receptors. mABs are produced from a common germ cell and have affinity for a specific antigen, thus inhibiting their binding to the corresponding receptors (322). Although having just one target, their effectiveness may be broad, as angiogenic growth factors usually bind to a number of isoforms of their target receptors. mABs are usually given intravenously, and because of their high molecular weight, their half-life is long (weeks). The resulting more prolonged inhibition allows less frequent dosing. The high molecular weight of mABs reduces diffusional capacity, thus renal filtration is limited, but their penetration into the brain is also impeded. Antiangiogenic antibodies as single agents exhibit only limited clinical activity, thus they are usually applied in combination that significantly improves their therapeutic effect.

Decoy receptors are soluble proteins that compete with their membrane bound counterparts with high affinity for their ligands. In the absence of transmembrane and intracellular domains, they fail to mediate signal, thus function as a trap for angiogenic growth factors (323). These traps can either be physiologically present, as a result of proteolytic cleavage of their transmembrane counterparts (eg. soluble VEGFR1), or designed and added as external therapy (324,325).

Receptors on ECs can be targeted at their extracellular part by mABs, thus blocking ligand binding, or by RTKIs at the tyrosine kinase domain, which serve as a docking site for molecules mediating the angiogenic signal (326,327). Anti-receptor mABs can usually target only one of the receptors, thus do not mediate widespread effect.

RTKIs can be divided to five subgroups:

- Type I kinase inhibitors recognize the active conformation of the receptor. They typically consist of a heterocyclic ring system that occupies the purine binding site of the enzyme, and form one to three hydrogen bonds by that part of the inhibitor, which mimic the purine ring of the adenine moiety, thus actively competing with ATP. Extra interactions may also be formed at hydrophobic regions adjacent to the hinge region. The hydrophilic region of the enzyme can be exploited for maximizing the solubility of the compounds. Since the targeted ATP pocket is conserved through the kinome, Type I inhibitors usually have low kinase selectivity, thereby enhancing the potential for off-target side effects. Examples of type I tyrosine kinase inhibitors targeting the VEGF pathway are sunitinib and pazopanib.
- Type II inhibitors recognize the inactive, unphosphorylated conformation of the kinase and indirectly compete with ATP by occupying the hydrophobic pocket, which is created by the DFG-out conformation of the activation loop. It is also known as the allosteric site, thus type II inhibitors can modulate kinase activity in an allosteric way. The DFG-out conformation is unique to all receptors, thus the hydrophobic interactions with the DFG pocket confer a high degree of selectivity. Some type II inhibitors are able to form a hydrogen bond directly to the ATP-binding site, but this is not necessary for their functionality. Sorafenib is a type II kinase inhibitor (328).
- Type III or allosteric inhibitors bind outside the catalytic domain of the kinase, in regions that are involved in the regulatory activity of the enzyme. They block the binding of ATP by modulating the conformation of the receptor. As they exploit the binding sites and regulatory mechanisms that are unique to the target, a high degree of kinase selectivity is exhibited. Additionally, allosteric modulators can provide delicate regulation of kinases, which is not easily performed with ATP-competitors.
- Type IV kinase inhibitors also act allosterically by forming a reversible interaction outside the ATP binding pocket, in the kinase substrate binding site, thus are not

competing with ATP. Since this area is unique for the substrate, it also ensures high degree of selectivity.

- The fifth class of kinase inhibitors are known as ‘covalent’ inhibitors. These bind covalently to cysteines proximal to the ATP binding site. Sulfur, an electron-rich atom is present in the cysteine residue, and reacts with the electrophilic groups of the inhibitor. As a result, by sharing electrons, they bind irreversibly, thus allowing the inhibitor to prevent the binding of ATP. The cysteine residue, and thus the binding site of the inhibitor can be variably located in the kinase domain. Examples of covalent tyrosine kinase inhibitors are quinazoline-based inhibitors (329) such as vandetanib.

Most small-molecule kinase inhibitors developed to date compete with ATP, thus target (nearby) the ATP-binding site. This region is common to all RTKs, thus selectivity is ensured by the region of the inhibitor, which is not similar to the structure of ATP.

In contrast to the mABs, RTKIs are much smaller, which on one hand may result in penetration to the brain, but also in decreased stability. The subsequent shorter half-life (hours) makes daily dosing necessary. Fortunately oral application is possible, which allows more comfort for the patient. However, absorption is often influenced by food and concomitant medications. Interactions in absorption and broad metabolism of RTKIs may be responsible for their limited activity or increased toxicity.

The toxic effects of antiangiogenic RTKIs can in part be attributed to their lack of selectivity. However, selective inhibitors may also induce toxicities, because their target kinases are not exclusively expressed by endothelial cells. As discussed above, although normal vasculature remains quiescent during adulthood, growth factor signaling in normal endothelial cells is still important for their survival and the maintenance of vascular integrity. Moreover, as specific kinases are involved in the normal physiology of organs like kidneys and the thyroid gland, specific toxicities, such as nephrotic syndrome might be related to the interference of RTKIs with the normal function of these organs (330). Furthermore, bleeding and wound healing abnormalities may be caused by the disturbance of the close interaction of PDGFR, FGFR and fibroblasts (331). In line with that, most common toxicities of antiangiogenic tyrosine kinase inhibitor therapy include hypertension, bleeding, fatigue, diarrhea, nausea, vomiting, hand-foot syndrome, and myelosuppression (332).

In order to enhance their effect, antiangiogenic agents can be combined, resulting in either vertical or horizontal blockade. Horizontal combinations have targets at the same level of the pathway, like different growth factors or different RTKs. Vertical combination, for example blocking the ligand and its receptor simultaneously leads to more effective suppression of a given pathway (333).

Because of the structural similarities of the main angiogenic RTKs (VEGFRs, PDGFRs and FGFRs), they activate overlapping signaling cascades. As a result, most antiangiogenic RTKIs block more than one isoforms of these receptors, (albeit with different affinity), thus horizontal blockade can be achieved by using them alone (multi-target RTKIs). This makes tumors less prone to switch from one driver angiogenic molecule to another, and may result in enhanced tumor growth inhibition (334). Moreover, the target receptors of antiangiogenic RTKIs are often expressed not only by endothelial-, but tumor cells as well, exhibiting direct antitumor properties beside the blockade of the vasculature. On the other hand, inhibition of PDGFR and FGFR signaling and the subsequent loss of mural cell function can also result in vascular destabilization and enhanced tumor leakiness, which may support metastatization (335). Furthermore, simultaneous receptor blockade may lead to increased toxicity (336). Antiangiogenic agents approved for the treatment of cancer are shown in **Table 1**.

Table 1. Antiangiogenic agents approved for the treatment of cancer. Target status defined as half maximal inhibitory concentration (IC50)<1000 nM. Approval of FDA, if not, labelled.

Name	Target receptor/Type	Developer	Indication
mAB			
Bevacizumab (Avastin)	Recombinant humanized anti-VEGF mAB (337)	Genentech	Metastatic colorectal cancer (CRC); metastatic non-small cell lung cancer (NSCLC); progressive glioblastoma; metastatic renal cell carcinoma (RCC); metastatic cervical cancer; platinum resistant ovarian, fallopian tube, or

			primary peritoneal cancer
Ramucirumab (Cyramza, IMC-1121B)	Full human Anti- VEGFR2 mAB (338)	ImClone Systems Inc.	Metastatic CRC; metastatic NSCLC; advanced gastric or gastroesophageal junction adenocarcinoma
RTKI			
Axitinib (Inlyta, AG013736)	VEGFR1-3, PDGFR α , - β , c-KIT, FLT3, CSF-1R (339)	Pfizer	Advanced RCC
Cabozantinib (Cabometyx, XL184, BMS907351)	VEGFR1-3, PDGFR β , TIE2, c- MET, KIT, FLT3, AXL, RON (340)	Exelixis	Advanced RCC
Nintedanib (Vargatef, BIBF1120)	VEGFR1-3, PDGFR α , - β , FGFR1-4, IGF1R, insuline receptor, FLT3, LCK, Src, LYN (341)	Boehringer Ingelheim	Advanced lung adenocarcinoma (EMEA)
Pazopanib (Votrient, GW786034B)	VEGFR1-3, PDGFR α , - β , FGFR1, -3-4, c-KIT, c-FMS, LCK, ITK, FAK (342)	GlaxoSmith Kline	Advanced soft tissue sarcoma; advanced RCC
Regorafenib (Stivarga, BAY 73- 4506)	VEGFR1-3, PDGFR β , FGFR1, TIE2, BRAF, BRAF (V600E), RAF-1, RET, KIT (343)	Bayer	Advanced gastrointestinal stromal tumor (GIST); metastatic CRC
Sorafenib (Nexavar, BAY 43-	VEGFR2-3, PDGFR β , FGFR1, EGFR, HER2,	Bayer	Advanced RCC; unresectable hepatocellular carcinoma; progressive differentiated

9006)	BRAF, BRAF (V599E), RAF-1, IGF1R, FLT3, C-KIT, ERK1, MEK1 (344)		thyroid carcinoma
Sunitinib (Sutent, SU11248)	VEGFR2, PDGFR β , FGFR1, EGFR, FLT3, Kit (345)	Pfizer	progressive well differentiated pancreatic neuroendocrine tumors (pNET); metastatic RCC; GIST after disease progression, or intolerance to imatinib mesylate
Vandetanib (Caprelsa, ZD6474)	VEGFR2-3, EGFR (346)	AstraZeneca	Advanced thyroid cancer
Trap receptor			
(Ziv-) Aflibercept (Zaltrap, AVE0005)	Recombinant fusion protein of the extracellular domains of human VEGFR1 - 2 (347)	Sanofi and Regeneron	Metastatic CRC

The role of VEGF and its receptor as an important factor in the vascularization, metastatization and proliferation of human colorectal cancer was already postulated in the mid-1990s (348). The first drug inhibiting the VEGF-VEGFR axis in colorectal cancer (CRC), bevacizumab was approved in 2004 (349). The other antiangiogenic mAB, ramucirumab was just approved in 2015 (350). Ziv-Aflibercept was approved by the FDA in 2012 (351). Beside the spread of antiangiogenic RTKIs in the clinic, regorafenib, a dual VEGFR2-TIE2 blocking drug is the only antivascular RTKI, being used for the treatment of CRC since 2012 (352). All other classical VEGFR inhibitors have failed to demonstrate unequivocal benefit in this patient population.

3.2.5 Resistance to antiangiogenic tyrosine kinase inhibitor therapy

Initially, no resistance to antiangiogenic tyrosine kinase inhibitors was expected, because they target genetically stable ECs and therefore unlikely to develop mutations. In spite of that, drug resistance in patients treated with antiangiogenic therapies is an important clinical problem. Both primary (no initial response is shown to therapy) and secondary (after a short regression period, the tumor recovers) resistances have been documented. The following mechanisms are considered to explain the phenomenon:

Although VEGF signaling is the predominant stimulator of angiogenesis, as seen above, several other pathways are involved in the promotion of neovascularization. Thus, inhibition of VEGFR-mediated pathways may not be sufficient to completely inhibit vessel growth. Activation of alternative angiogenic pathways may circumvent inhibition by antiangiogenic tyrosine kinase inhibitors (353).

In line with that, as discussed above, VEGF signalization does not drive all forms of angiogenesis, and thus by activating alternative vascularization mechanisms, vascular development can be achieved in spite of effective VEGFR inhibition.

Resistance to kinase inhibitors can also result from a mutation in the target site of the receptor even on endothelial cells, although resistance is less likely to arise if multiple receptors are being targeted at the same time. Moreover, multi-target inhibitors mostly hit the ATP binding site, thus they are less precisely linked to their target receptor and for that reason, are less sensitive for dislodging due to a mutation of the target kinase. In spite of that, several studies reported mutations in target kinases recently that correlate with resistance to antiangiogenic RTKIs (354,355).

Decreased vessel number is accompanied by hypoxia as a result of antiangiogenic therapy, but hypoxia may select for more resistant tumor cells, thus increasing tumor aggressiveness and invasion (356).

The metastatization potential might also be enhanced by the loss of tumor vessel integrity, because of thrombotic events (357) or the decreased number of pericytes as a result of PDGFR and FGFR inhibiting functions of multi-target RTKIs (358).

Beside the above mechanisms, probably the most important cause of resistance is suboptimal pharmacokinetics and/or the localization of the drug out of the target site. Multidrug resistance proteins (MDRs) may also be involved in the removal of antiangiogenic RTKIs from the tumor tissue (359).

3.3 TKI imaging

Mass spectrometry (MS) is an analytical chemistry technique, which generates ions from molecules by excitation energy, and helps to identify the structure and relative intensity of molecules from their mass to charge ratio (m/z). By virtue of its speed and sensitivity, MS has become a key technique in medical research and drug development.

In the last decades MS technology has been dramatically improved and had an impact on the research field of angiogenesis as well. The first proteomic study of HUVECs was published in 2003, when 53 proteins were identified using Time of flight (TOF) MS (360). Since then, the number of detected proteins has increased, and also those being differentially expressed upon pro/antiangiogenic stimulation have been identified (361). Moreover, stable isotope labeling with amino acids (SILAC) of ECs enables accurate quantitative proteomic analysis of cell cultures (362).

MS technology can be used in several fields of angiogenesis research. For example, the proteome of subcellular compartments can be studied (363). Of that area, the cellular secretome is of particular interest, because it allows investigation of the communication between ECs and surrounding cells (364). Cellular regulatory mechanisms such as protein trafficking, posttranslational modifications or phosphorylation of molecules can also be examined by MS (365). It allows the characterization of the cellular and molecular cascades that regulate developmental angiogenesis (366). MS analysis can be further extended to mice treated with (anti)angiogenic therapeutics to detect molecular changes in the vasculature that are critical for tumour progression. The combination of mass spectrometry and separation techniques, such as liquid-chromatography and electrophoresis has already identified numerous disease biomarkers from various body fluids (367). Moreover, using laser capture microdissection, the highest regional selectivity could be applied for down to single cell analyses (368), which helps to build a picture about spatial distribution of the analytes. The need for this spatial information and the time consuming, arduous nature of laser capture microdissection promoted the spread of mass spectrometry imaging techniques. This ensures visualization of molecules on complex surfaces to identify and localize elements, lipids, peptides, proteins, pharmaceuticals and metabolites in biological tissues (369-373).

Although the last half century has witnessed dramatic advances in the field of medical imaging, there is still an urgent need for the development of more advanced techniques

in the drug discovery process. This is particularly important in the narrowing of the selection of potential hits and leads as candidates for further development. One of the reasons this has been difficult to accomplish in the past is that until recently, the only avenue for visualizing the in vivo distribution of drugs in targeted tissues was the use of labels, commonly radioactive and such, a safety risk. Methods, like positron emission tomography and autoradiography can provide information on the distribution of a radio-labelled compound even at cellular level (374). However, both of these methods rely on quantitative data based upon the relative strength of the label rather than the relative concentration of the drug. If a drug is metabolized, the label can follow the altered structure, that is neither active, nor the precursor of an active form, and the readout of distribution may have little to do with the mode of action or the actual efficacy of the drug (374). For these reasons, unlabeled i.e. “cold compound” would provide evidence that relate only to the drug structure and not to the chemistry of labeling material in a modified drug molecule. Other methods rely on the use of isotopes with relatively short half-lives or fluorescent tags which makes long-term pharmacological analysis impossible or alters the chemical structure of the drug and thus, the binding affinity and/or avidity to its target molecule (375). From this point of view, it is particularly important that the applied methods can be used to investigate the characteristics of the unaltered native compound (i.e. the same agent that is being administered to patients). Meanwhile the pharmacological properties of novel drug candidates are routinely characterized during the preclinical phase of drug development, in-depth and routine determination of the adsorption, distribution, metabolism and elimination (ADME) of compounds became the focus of research only in the last decades. Although ADME can fundamentally influence the therapeutic benefit of different drugs, until recently, extensive ADME studies were conducted rather late in the process of drug development, mainly in phase I clinical studies. This may be one of the key factors behind the low, 11%, overall first-in-man to registration rate of novel drug candidates in the 1990s. This proportion was especially poor (5%) among drugs in the field of oncology (376). Mass spectrometry is a powerful technique, enabling the parallel determination of label-free drugs and their metabolites from different tissue compartments, that gives researchers the opportunity to analyse the adsorption, distribution and elimination of the native drug and its active/toxic metabolites as well.

One of the best techniques for such complex experiments is matrix-assisted laser desorption ionization (MALDI) MS (377). Briefly, a suitable matrix material is added to the sample surface, which extracts the analytes, helps the formation of analyte-doped crystals on the surface, and absorbs the laser energy for soft-ionization of sample molecules (378). Thus, sample preparation is crucial for successful detection of the desired molecule (379). Then, a pulsed laser irradiates the sample, triggering ablation and desorption of the sample-matrix mixture. Finally, the analyte molecules are ionized by being protonated or deprotonated in the hot plume of ablated gases, and then are accelerated into the mass spectrometer linked to analyse them. The use of MALDI mass spectrometry in pharmacological studies dates back to the mid 1990s when in vitro metabolites were characterized by this technique (380).

The development of MALDI mass spectrometry imaging (MSI) sources for high mass resolution and mass accuracy analysers allows for the separation of ions with the same nominal mass and confident assignment of elemental formulas. Furthermore, various MS/MS techniques are available to link with FT-MS systems. This, by fragmenting the precursor molecule and yielding a structure specific fragment ion map, further ensures the identification of the desired molecule. This technology has recently been employed to determine the exact tissue compartment localization of small drug molecules (381).

Although localization of antitumor molecules in their target site is considered to be important to show their efficacy, studies published to date focus mainly on the measurements of compounds from the blood, urine and occasionally from tissue homogenates (382). Consequently, the lack of greater clinical success of anticancer agents is, at least in part, due to our limited knowledge of their pharmacokinetic profile, bioavailability and distribution at the tumor tissue. Antiangiogenic drug imaging is also still in its infancy, as the only study reported so far on the distribution of an antiangiogenic antibody, detected by MALDI-MSI was published in 2014 (383). Our group was the first to show imaging data on the distribution of antiangiogenic RTKIs.

However, considering that the inhibition of the tumour vessel network may influence the dispersal of the antiangiogenic drug itself, such spatial distribution data could greatly help researchers to better understand their mode of action and to identify the best potential therapeutic schedules of these agents.

4. AIMS

The clinical experiences with antiangiogenic RTKIs are controversial, despite their predicted inhibitory and normalizing effects on vessel growth. No biomarker of tumor response has yet been linked to the effect of these agents. Taken into consideration that ADME can fundamentally influence the therapeutic benefit of different drugs, the following specific aims have been defined:

1. to develop a method for the detection of antiangiogenic RTKIs and their metabolites in different tissues.
2. to analyse the intratumoral distribution and levels of different antiangiogenic receptor tyrosine kinase inhibitors (RTKIs) of mice bearing subcutaneously growing murine tumors by using MALDI-MSI.
3. to evaluate the potential associations between intratumoral antiangiogenic drug levels and distributions and **3.1./** tumor growth inhibition; **3.2./** blood vessel density and area; **3.3./** blood vessel integrity (basement membrane, pericyte and α -smooth muscle actin (SMA) coverage **3.4./** the size and localization of hypoxic areas of the tumor tissue; **3.5./** the expression and distribution of receptors targeted by the RTKIs.

5. METHODS

5.1 In vivo tumor models and treatments

5.1.1 Tumor models

Two different mouse colon adenocarcinoma models, C26 and C38 were used for our experiments. The C26 cell line was cultured in RPMI 1640 medium with 10% fetal bovine serum and 1% penicillin/streptomycin (all from Sigma Aldrich, Steinheim, Germany) in a humidified atmosphere at 37°C, 5% CO₂. Groups of six 8-week-old female Balb/C mice from the colony of the National Oncology Institute, Budapest were inoculated subcutaneously (sc.) with 2x10⁶ C26 cells. The C38 tumors were maintained by serial s.c. transplantations in 8-week-old female C57Bl/6 mice. Tumors were cut into cubes measuring 5×5×5 mm. Animals were anesthetized and one piece of tumor tissue was transplanted into the back of each mouse.

All animal-model protocols were developed and conducted in accordance with the ARRIVE guidelines (384) and the animal welfare regulations of the Department of Experimental Pharmacology, National Institute of Oncology, Budapest, Hungary (permission number: 22.1/722/3/2010). Mice were kept on a daily 12-h light/12-h dark cycle and held in conventional animal house in microisolator cages with water and laboratory chow ad libitum.

5.1.2 Drugs

For drug treatment of C26 bearing mice, 5 antiangiogenic RTKIs either approved by both the FDA and the EMEA (pazopanib, sorafenib, sunitinib) or investigated in Phase III trials (motesanib, vatalanib) were selected. In the C38 model a vatalanib-treated and a control group was generated.

Drugs were purchased from LC Laboratories, (Woburn, MA, USA, CAS. No. for motesanib: 453562-69-1; pazopanib: 444731-52-6; sorafenib: 284461-73-0; sunitinib: 557795-19-4 and vatalanib: 212141-54-3) at >99% purity and suspended in 2% carboxymethylcellulose with 2 mg/mL methyl-4-hydroxybenzoate (both from Sigma Aldrich). Control animals received the suspending medium only.

These RTKIs are known to have significant IC₅₀ values of the main angiogenic receptor, VEGFR2, and may also inhibit PDGF- and FGF receptors (342,344,385-387).

The IC50 values (nM) of the applied compounds against the main antiangiogenic receptors as measured in a cell-free assay are shown in **Table 2**.

Table 2. IC50 values (nM) of the main antiangiogenic RTKIs against human angiogenic receptors measured in a cell free assay. IC50 values of the corresponding mouse receptors are shown in brackets. ND: not defined. *not defined which FGFR is tested.

Target receptor	Motesanib	Pazopanib	Sorafenib	Sunitinib	Vatalanib
VEGFR1	2	10	ND	ND	77
VEGFR2	3 (6)	30	90 (15)	9	37 (270)
VEGFR3	6	47	(20)	ND	660
PDGFR α	ND	71	ND	ND	ND
PDGFR β	84	84	(57)	8	580
FGFR1	>2800*	140	580	830	ND
FGFR2	ND	ND	ND	ND	ND
FGFR3	ND	130	ND	ND	ND
FGFR4	ND	800	ND	ND	ND
TIE1	ND	ND	ND	ND	ND
TIE2	ND	4520	ND	ND	ND
Ref.	(385)	(342)	(344)	(386)	(387)

Pazopanib, sunitinib, and vatalanib are Type I inhibitors, while motesanib and sorafenib belong to the group of Type II inhibitors (327,388). Out of this five compounds pazopanib, sorafenib and sunitinib are already used for cancer therapy. All three are approved for advanced RCC. Besides RCC, pazopanib is effective in advanced soft tissue sarcoma (389). Sorafenib is used for the treatment of unresectable hepatocellular carcinoma and locally recurrent or metastatic, progressive differentiated thyroid carcinoma, refractory to radioactive iodine treatment (41). Sunitinib therapy is approved in progressive well-differentiated pNET and GIST after disease progression or intolerance to imatinib mesylate (390). Several completed or still ongoing clinical trials have assessed the effects of these approved antiangiogenic RTKIs in colorectal cancer with controversial results (352,391-395). Motesanib and vatalanib have been investigated in clinical trials in different tumor types. In colorectal cancer, the only study completed to date with motesanib in first- or second line treated CRC patients demonstrated modest efficacy (396), while vatalanib is showing promising results in CRC patients only with activated vessel density (high phospho-VEGFR expressing vessels) or low lactate dehydrogenase levels (397,398).

5.1.3 In vivo treatment

The administration and dose level for each compound was chosen to be tolerable and effective according to the literature (385,399-402). Moreover, to mimic the clinical situation, RTKIs were administered orally.

Accordingly, in case of the C26 model, treatment began 14 days after tumor cell injection and was performed per os with a feeding tube once daily, at a dose of 100 mg/kg, 5 times a week for two weeks. In the C38 model, treatment began 9 days after tumor implantation and was performed as in the C26 model.

During the growth of tumors animals were weighed three times per week and monitored daily for tumor related symptoms. To evaluate the in vivo effects of the different antiangiogenic RTKIs, two diameters of the tumors were measured three times a week, and tumor volume was calculated with the formula $\text{width}^2 \times \text{length} \times \pi/6$.

To assess intratumoral hypoxia, a bolus of i.p. pimonidazole (60 mg/kg; Hypoxyprobe Inc., MA, USA) was administered 1 hour before the mice were sacrificed. Pimonidazole is activated by reduction in hypoxic cells and forms adducts with thiol groups in proteins, peptides and amino acids.

Two hours after the last oral treatment, peripheral blood was drawn from the canthus and mice were sacrificed. Tumors and normal organs were removed, tumor weight was measured and samples were snap frozen by submerging the tissues into dry ice cooled isopentane. Frozen tissues were stored at -80 °C until utilization.

5.2 Analysis of vascular parameters and target receptors

For the analysis of vascular parameters and target receptor expressions, 10 serial frozen sections were cut from each tumor. Sections #5 and #7 were used to analyse the distribution and levels of the given RTKI by MALDI-MSI and for subsequent haematoxylin&eosin (HE) staining. Sections #1-4 were labeled with either of the following primary antibodies: anti-FGFR1, anti-PDGFR α , anti-PDGFR β and anti-VEGFR2. For hypoxia detection (section #6), we used the Hypoxyprobe-1 Plus Kit. Sections #8-10 were labeled with either of the following primary antibodies: anti-laminin (for endothelial basement membrane labeling), anti-desmin (for pericyte labeling) and anti- α -smooth muscle actin (anti- α SMA). All of the above primary antibodies were developed with an appropriate fluorescent secondary antibody. For

intratumoral microvessel density (MVD) and microvessel area measurements, sections #1-4, #6 and #8-10 were co-stained with anti-mouse CD31 antibody, followed by a counterstain with Hoechst 33342 before mounting under glass coverslips in ProlongGold Antifade Reagent (Invitrogen, Carlsbad, CA, USA; Catalog number: P36930). Primary and secondary antibodies and dyes used for immunohistochemical labeling are listed in **Table 3**.

Slides were scanned by TissueFAXS using a 20x objective and analysed by TissueQuest 4.0.0140 (both from TissueGnostics GmbH, Vienna, Austria) and ImageJ software packages.

For microvessel density and microvessel area measurements, images of ten different viable intratumoral regions were assessed separately for each section. Microvessel areas were calculated by measuring the number of CD31-positive pixels, while MVD was estimated by counting the number of vessels in all ten fields of view, and calculating the average of them. For quantification of VEGFR2 expression, both signal area and density were calculated similarly to CD31. In case of VEGFR2, positively labelled vessels and tumor cells were also counted for data analysis. The percentages of microvessels that were positive for laminin, desmin or α SMA were also calculated. For quantification of hypoxia, PDGFR α , PDGFR β and FGFR1 expressions, the percentages of the areas of positively labeled cells were determined across the entire section.

Table 3. Specifications of different antibodies and the nucleic acid stain used in our experiments.

Antibody	Species	Manufacturer	Catalog No.	Dilution
Primary				
Anti-CD31	rat monoclonal	BD Pharmingen, BD Biosciences, Franklin Lakes, NJ, USA	550274	1:50
Hypoxyprobe-1 Plus Kit (FITC-Mab1)	mouse monoclonal	Hypoxyprobe Inc., Middlesex Turnpike Burlington, MA, USA	HP1-100Kit	1:100
Anti-Laminin	rabbit polyclonal	DAKO, Glostrup Denmark	Z0097	1:200

Anti-Desmin	rabbit polyclonal	Abcam, Cambridge, UK	Ab32362	1:200
Anti- α SMA	mouse monoclonal	DAKO, Glostrup Denmark	M0851	1:200
Anti-FGFR1	rabbit polyclonal	Cell Signaling Technology, Danvers, MA, USA	9740	1:50
Anti-PDGFR α	rabbit polyclonal	Cell Signaling Technology, Danvers, MA, USA	3174	1:50
Anti-PDGFR β	rabbit polyclonal	Cell Signaling Technology, Danvers, MA, USA	4564	1:50
Anti-VEGFR2	rabbit polyclonal	Cell Signaling Technology, Danvers, MA, USA	2479	1:50
Secondary				
Alexa 488	Anti-mouse goat F(ab') ₂	Cell Signaling Technology, Danvers, MA, USA	4408	1:1000
Alexa 488	Anti-rabbit goat F(ab') ₂	Cell Signaling Technology, Danvers, MA, USA	4412	1:1000
Alexa 555	Anti-rat goat F(ab') ₂	Cell Signaling Technology, Danvers, MA, USA	4417	1:1000
Nucleic acid stain				
Hoechst 33342		Molecular probes, Eugene, OR, USA	H3570	1:10000

5.3 Detection of the compounds and analysis of drug distribution and intensity

5.3.1 Ionization technique and instrumentation

MALDI-MS was used for the detection of RTKIs. MALDI-MS is a soft ionization technique, thus it is mostly used to detect and analyse molecules which tend to be fragile and fragment when ionized by conventional ionization methods.

As a matrix, molecules with mobile electron system are suitable, which are showing light absorption in the wavelength of the laser, thus transmit the laser energy to the analyte molecule. This process facilitates the ionization of the sample. Besides, the matrix is able to block the aggregation of analyte molecules.

For the analysis of small molecules, the most commonly used matrices are 3,5-dimethoxy-4-hydroxycinnamic acid (sinapinic acid), α -cyano-4-hydroxycinnamic acid (CHCA) and 2,5-dihydroxybenzoic acid (DHB). A solution of one of these matrices is usually made of a mixture of highly purified water and an organic solvent such as acetonitrile (ACN), acetone, chloroform, methanol or ethanol. The most suitable matrix is variable among different molecular structure analytes, thus more matrices should be tested to reach maximal quality of the mass spectrum and highest signal intensity of the analyte. A counter ion source such as trifluoroacetic acid (TFA) is usually added to generate the $[M+H]^+$ ions (quasimolecular ions). The analyte molecules are mixed with the matrix solution either on a target plate or on tissue surface. The mixture of a hydrophilic (water) and a hydrophobic (organic solvent) component of the matrix solution allows both hydrophobic and hydrophilic molecules to dissolve into the solution. The solvents vaporize in a few seconds, while matrix molecules co-crystallize with the analyte molecules embedded into matrix crystals.

MALDI (**Figure 13.**) makes it possible to detect molecules of interest with high accuracy and sensitivity from small amounts of complex samples, as one of the main advantages of this ionization technique is that it has great tolerance towards salts and buffers, being present in physiological samples (403). However, matrix application can result in the generation of high background signals below 500 Da mass range (404).

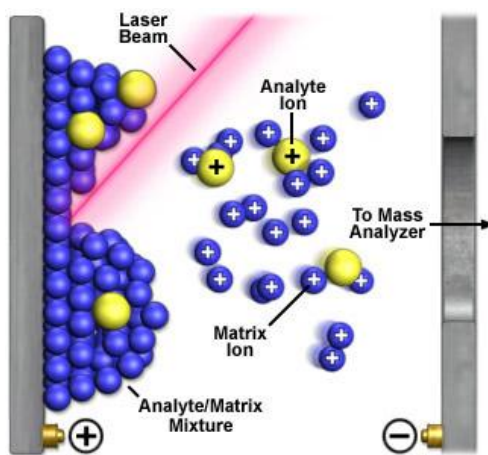


Figure 13. Ionization of the analyte molecule (yellow) with matrix (blue) using MALDI (405). Solution of the matrix is applied to the surface of the sample. In turn, analyte molecules dissolve into the solution. The solvents vaporize, while matrix molecules co-crystallize with the analyte molecules embedded into matrix crystals, thus blocking their aggregation. The matrix (a molecule with mobile electron system) absorbs light in the wavelength of the laser and transmits the laser energy to the analyte molecule. This process facilitates the ionization of the sample.

Both for compound characterization and drug detection in blood on a MALDI target plate, as well as for tissue imaging a MALDI LTQ Orbitrap XL mass spectrometer (**Figure 14.**, Thermo Fisher Scientific, Bremen, Germany) was used.

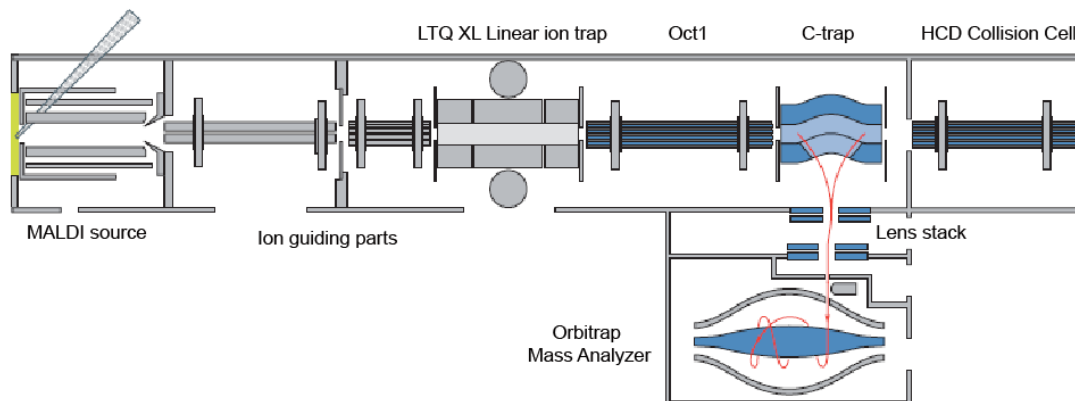


Figure 14. Structure of the MALDI LTQ Orbitrap XL mass spectrometer (406).

After the insertion of the plate, the sample is targeted by a nitrogen laser at the UV range. By absorbing light in the wavelength of the laser, the matrix gets excited, ionized and desorbed from the surface of the sample. Finally matrix molecules transfer/remove protons to/from the analyte molecules, which also get into gas phase by desorption from the sample surface. The analyte molecules in turn get singly or multiply charged depending on the matrix, the laser intensity and the applied voltage. One of the main advantages of the MALDI technique is that the laser is operated in a pulsed mode, thus generation of ions is a discrete process, and if the analysis is linked in time with ion generation, hardly any sample is wasted. This is one reason of the high sensitivity of the MALDI technique, as even fmol (10^{-15}) of analyte molecules can be detected (407). On the other hand, pulsed laser is the reason, why MALDI ionization cannot be linked with some specific mass analysers. After the ion beam is formed, ions are accelerated and sent to the mass analyser part of the instrument.

The mass analyser can either be responsible for fragmentation of the gas phase analyte molecules, or separation of both precursor molecules and their fragment ions based on their m/z ratio, depending on the scan type used. Moreover, they transfer ions to the detector. The mean free path in the analyser (the average distance traveled by an ion between collisions) has to be larger than the length of the instrument. That means that collision with the wall of the instrument is more likely than it is with other ions. This is

of major importance, because collisions may modify the direction or energy of ions, which mainly impact data outcome. Therefore, analysers are operated in high vacuum.

The instrument used in our experiments has a hybrid linear quadrupole ion trap (LTQ)-orbitrap mass analyser, and both of the mass analysers function as mass detectors as well. Compared to other hybrid instruments, LTQ-Orbitrap is characterized by high ion transmission (30%–50%) (406).

The LTQ (**Figure 15.**) part consists of 4 parallel metal rods, which conduct electricity. Direct current (DC) and alternate current (AC) voltages are both switched to the rods in such manner, that the opposite rods have the same potential, while the neighbouring rods have potentials with opposite signs. Ions leave the ion source and enter the quadrupole in response to accelerating voltages. Here the positive potential rods toss the positive ions, while attract the negative ions. The negative rods do the opposite. As a result of the AC voltage the relative charge of the rods is changing from time to time, thus, the ions oscillate while passing through the rods. In a quadrupole mass analyser, the correct magnitude of the radio frequency and DC voltages applied to the rods allow ions of specific m/z to maintain stable trajectories from the ion source to the detector, whereas ions with different m/z values are unable to maintain stable trajectories.

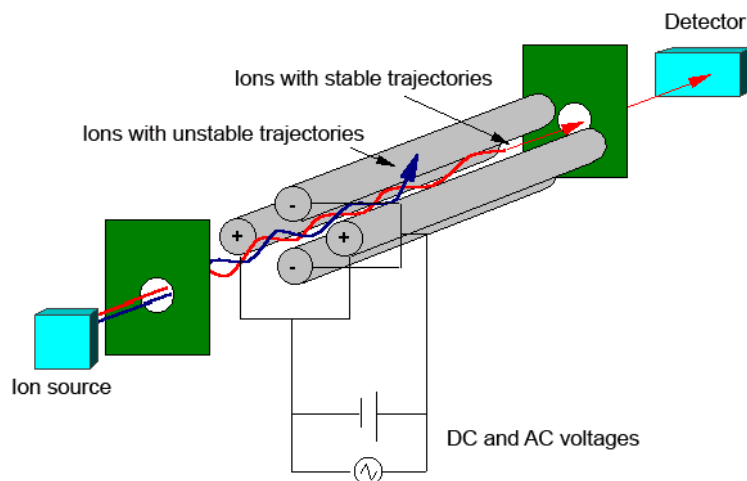


Figure 15. Structure of a quadrupole mass analyser (408). The 4 parallel rods conduct electricity. DC and AC voltages are switched to the rods in an alternating manner. Two opposite rods have a certain applied potential, while the other two rods have the same potential with negative sign. As a result of the AC voltage the relative charge of the rods is changing from time to time, thus the ions oscillate as determined by their charge while passing through the rods with a speed defined by their mass. For given DC and AC voltages, only ions of a certain mass-to-charge ratio pass through the quadrupole and all other ions are thrown out of their original path.

The linear ion trap part of the instrument is an independent MS detector, thus able not just to store and isolate, but fragment and then send ions through one of the three exit slots either to the Orbitrap for further analysis or to a secondary electron multiplier (SEM) detector. Fragmentation occurs in the LTQ by collision-induced dissociation (CID). The precursor ions are accelerated by electric potential to high kinetic energy and then collide with helium. The kinetic energy is partly converted into internal energy which results in breakage of bonds and the molecular ion falls apart into smaller fragments. These fragment ions can then be analysed by a mass spectrometer.

Mass analysis (**Figure 16.**) is achieved by making ion trajectories unstable in a mass-selective manner. The main radio frequency voltage is ramped and simultaneously AC voltage is applied to the exit rods to facilitate ejection. As the main radio frequency voltage is increased, ions of greater and greater m/z values become unstable and are ejected through the exit slots.

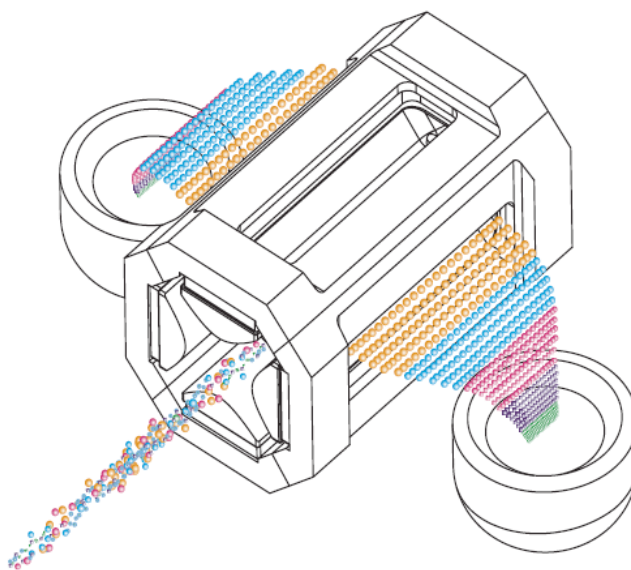


Figure 16. Operation of LTQ. Ions of different m/z values (represented by colored spheres) enter the linear ion trap. The ions are ejected from the trap in order of increasing m/z values. After exiting the trap through slots in the exit rods, the ions strike the conversion dynodes on each side of the trap (409).

These ions are focused toward the ion detection system where they are detected. If the linear ion trap functions as a mass detector as well, the ions strike the conversion dynodes in the off-axis ion detection systems upon ion ejection. This is located in both sides of the trap. Upon striking the surface of the conversion dynode by an ion, secondary particles are produced. These secondary particles are focused by the curved

surface of the conversion dynode and are accelerated by a voltage gradient into the electron multiplier. The current that leaves the electron multiplier via the anode is converted to a voltage and recorded by the data system.

If the detector is the orbitrap, ions move through the gas-free radio frequency octapole (Oct 1) into the gas-filled curved linear trap (C-Trap) on their way from the linear trap to the Orbitrap. Ions in the C-Trap are returned by a trap electrode. Upon their passage, using nitrogen collision gas the ions lose kinetic energy and cool down, thus preventing them from leaving the C-Trap through the gate. From the C-trap, ion packets are injected tangentially into the field of the Orbitrap.

The orbitrap (**Figure 17.**) is both a mass analyser and a detector, consisting of a spindle-shaped central electrode surrounded by a pair of bell-shaped outer electrodes (410).

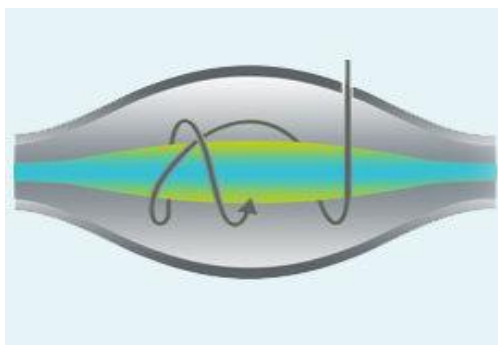


Figure 17. Orbitrap mass analyser (411), consisting of an outer bell-like electrode and an inner spindle-like electrode. Ions are trapped in an orbital motion around the spindle based on their m/z value. Image currents from the trapped ions are detected and converted into mass spectrum using Fourier transformation of the frequency signal.

Upon ions leaving the C-trap, the electric field in the orbitrap is increased by elevating voltage on the inner electrode. Thus, ions get squeezed towards the central electrode until they reach the desired orbit inside the trap. At that moment ramping of the voltage is stopped, the field becomes static and detection starts. Ions in the orbitrap move with different rotational frequencies based on their m/z ratio, but with the same axial frequency along the central electrode. The outer electrode is split in two symmetrical pick-up sensors connected to a differential amplifier. Axial oscillations of ion rings are detected by their image current induced on the outer electrode. This signal is converted into mass spectrum by Fourier-transformation (406).

From the linear ion trap ions are either sent to the orbitrap or to the high energy collisional dissociation (HCD) part of the instrument through the C-trap. The offset

between the C-Trap and HCD is used to accelerate the precursors into the gas-filled collision cell. It is supplied with a nitrogen collision gas providing increased gas pressure inside the multipole. HCD is specific to the orbitrap, in which fragmentation takes place outside the trap. After dissociation ions return to the C-trap before injection into the orbitrap for mass analysis. Despite the name, the collision energy of HCD is typically in the low regimen, thus fragmentation is not as efficient as in the linear ion trap, but as fragment ions of approx. 5%–10% m/z of the precursor ion mass can be observed, it may be used, when fragments are in a low m/z range (412).

Analysers are characterized by the following features:

Linear dynamic range is the concentration range in which signal intensity is in a linear correlation with the concentration.

Mass resolution defines how accurately two neighbouring m/z values can be distinguished. The border between the low and high mass accuracy instruments is 10^4 . This means, that ions of a molecule of hundred mass numbers can be differentiated by 0.01 m/z . ($R_s=100/0.01=10^4$).

The mass range is the range between the m/z values the instrument can minimally and maximally detect.

The detection limit shows the minimal amount of the analyte molecule, the instrument can detect. This can be highly influenced by the ion transmission efficiency. If the distance between the ion source and the detector is long enough, the efficiency of ion transmission is high enough.

Appropriate temperature of the instrument is essential, as the sample must be kept in gas phase during the ionization process.

The speed of the mass analyser is expressed in the amount of spectrums generated in a certain amount of time. This is important so that the mass spectrum reflects the real mass intensity dispersion of each sampled spot.

The LTQ Orbitrap XL has the following measuring properties:

Resolution: 60 000 (Full Width Half Maximum) @ m/z 400 with a scan repetition rate of 1 second.

Cycle Time: 1 scan at 60 000 resolution @ m/z 400 per second.

Mass Range: m/z 50–2 000; m/z 200–4 000.

Mass Accuracy: <3 ppm root mean square for 2 h period with external calibration using defined conditions, <2 ppm root mean square with internal calibration.

Dynamic Range: >10 000 between mass spectra, >4 000 between highest and lowest detectable mass in one spectrum.

Three scan modes are defined in MS. The mass spectrometry scan mode is a single stage mass analysis ($n = 1$). In that scan mode no fragmentation of the precursor ion occurs. The MS/MS scan is a two stage mass analysis ($n = 2$). In an MS/MS scan, precursor ions are fragmented into product ions. An MS^n scan usually involves three to ten stages of mass analysis ($n = 3-10$), in which the precursor ion is first fragmented, and afterwards fragment ions are selected as precursor ions and fragmented again. This process can be repeated 10 times. The instrument has MS/MS and MS^n scan functions.

5.3.2 Compound characterization

Drugs were dissolved in 50% methanol (Sigma-Aldrich, Steinheim, Germany) at HPLC grade (99.8+%) at 0.5 mg/mL concentration. After the selection of the appropriate matrix molecule, 7.5 mg/mL CHCA (Sigma Aldrich, Steinheim, Germany) was dissolved in 50% ACN (Merck, Darmstadt, Germany) and 0.1% TFA (Sigma-Aldrich, Steinheim, Germany) was added. 1 μ L of the compound solution was applied with 1 μ L matrix solution to the MALDI plate.

Full scan was performed, to determine the m/z value of the RTKIs and their fragment ions. Full scan presents a full mass spectrum of the analyte. In that case, ions are scanned from the first mass to the last without interruption. To determine the precursor ion peak, single-stage full scan was done. In that analysis, ions formed in the ion source are stored in the mass analyser and afterwards they are sequentially scanned to produce a full mass spectrum. Fragment ions were determined in a two-stage full scan experiment. In the first stage of mass analysis, the ions formed in the ion source are stored in the mass analyser. Then, ions of one specific mass-to-charge ratio (the precursor ions) are selected and all other ones are ejected from the mass analyser. The precursor ions are excited and collide with background gas. This facilitates their fragmentation to create one or more product ions. In the second stage of mass analysis the product ions are first stored and then sequentially scanned out of the mass analyser to produce a full product ion mass spectrum.

Full mass spectra were obtained at 60,000 resolution by using positive polarity. The spots were sampled in survey mode (accidentally choosing sampled spots) collecting 20 experiments for a single run. The nitrogen laser was set to 10 μJ . The detected precursor ions were isolated with m/z 2.0 width isolation window, and were fragmented by using 40% normalized collision energy (NCE) during a 30 ms activation time, while activation Q of 0.25 was applied. MS/MS spectra were collected at normal scan rate in centroid format.

5.3.3 Compound detection in the blood

Blood was removed from the canthus just before sacrificing the animals. After centrifugation plasma samples were stored in $-80\text{ }^{\circ}\text{C}$ until utilization. Acetonitrile precipitation was performed as the following: 20 μL of the plasma sample was removed and mixed with 40 μL ice cold 100% ACN to precipitate blood components. After vortexing and centrifugation at room temperature for 15000 rpm and 15 minutes, the supernatant was removed and dried with speed vac. Then the sample was diluted in 20 μL 0.1% TFA solvent. Pierce C18 Tips (Thermo Fisher Scientific, Rockford, IL, USA) were used to concentrate RTKIs from the precipitated plasma samples, following the manufacturer's instructions. 2 μL concentrate was eluted from the column, and 1 μL of this was applied on the MALDI plate with 1 μL matrix solution using the same instrument settings as used for compound characterization.

5.3.4 Tissue imaging of antiangiogenic RTKIs

10- μm frozen sections were cut using a cryotome and placed on glass slides. After drying of the tissue, 0.5 mL matrix solution was applied stepwise to avoid wetting of the sections by using an airbrush, while its position was kept constant. Full mass spectra were collected by performing a single stage full scan using the Orbitrap mass analyser at 60000 resolution (at m/z 400), in positive mode with a 150–800 Da mass range and 100 μm raster size. The nitrogen laser was operated at 10.0 μJ . For ensuring a known number of charges in the linear trap, in particular, in order to avoid overfilling the ion trap, automatic gain control (AGC) was used. AGC is a pre-scan event, which is performed before an analytical scan, for which the pre-scan serves as a prediction of the

number of charges, and the injection time. Depending on the result of the pre-scan event, the parameters of the live scan can be changed.

For obtaining MS/MS data two-stage full scan was performed. The observed peaks of the precursor drugs were isolated with m/z 2.0 width isolation window and fragmented, using 40% NCE, 30 ms activation time and 0.25 activation Q . For MS/MS spectra generation the minimal signal required by the linear ion trap was 500 counts. The fragment ions were analysed in the linear ion trap at normal scan rate.

5.3.5 Quantification of the compounds

For tissue quantification of intratumoral drug concentration, calibration curves of each compound were established on untreated control C26 and C38 tumor tissue sections. After determining the detection limit of the instrument, drugs were dissolved and diluted in 50% methanol (concentration range: 0.001–0.5 $\mu\text{mol}\cdot\text{mL}^{-1}$), and 0.5 μL from each concentration was applied on a tissue section. Spraying and detection conditions were the same as those during the tissue sample analysis of the in vivo treated tumors. Average signal intensities of the applied concentrations were measured and normalized to total ion current (TIC) by Xcalibur v 2.0.7 and ImageQuest™ software packages.

Calibration curves were created, which were then used to estimate the tissue drug concentrations of in vivo treated tumor sections. Because of the possibility of generating nonspecific precursor ion peaks from the tissue itself, the average signal intensities and the corresponding concentrations obtained from RTKI-treated tumors were compared with that of non-treated control tumors.

Throughout our MS experiments drugs were considered to be detected if the precursor molecule and at least one fragment ion was discovered in the spectra.

Evaluation of all MS spectra was performed with Xcalibur v 2.0.7 software, while the visualization of the drugs and fragment ions in/on tissue was implemented with the ImageQuest™ software (both from Thermo Fisher Scientific, San José, CA).

5.4 Statistical analysis

Differences in parametric and non-parametric variables between multiple groups were analysed using one-way ANOVA and Dunnett's posthoc test or using the Kruskal–Wallis test followed by a posthoc Dunn's multiple comparison test, respectively. For

comparing two groups, unpaired t-tests were applied to analyse parametric and Mann-Whitney U tests were used to analyse non-parametric data. Differences were considered statistically significant when $p < 0.05$. All Statistical analyses were carried out using GraphPad Prism 5.0 software (GraphPad Inc., San Diego, CA, USA).

Figure 18. shows the flow chart of the experiments.

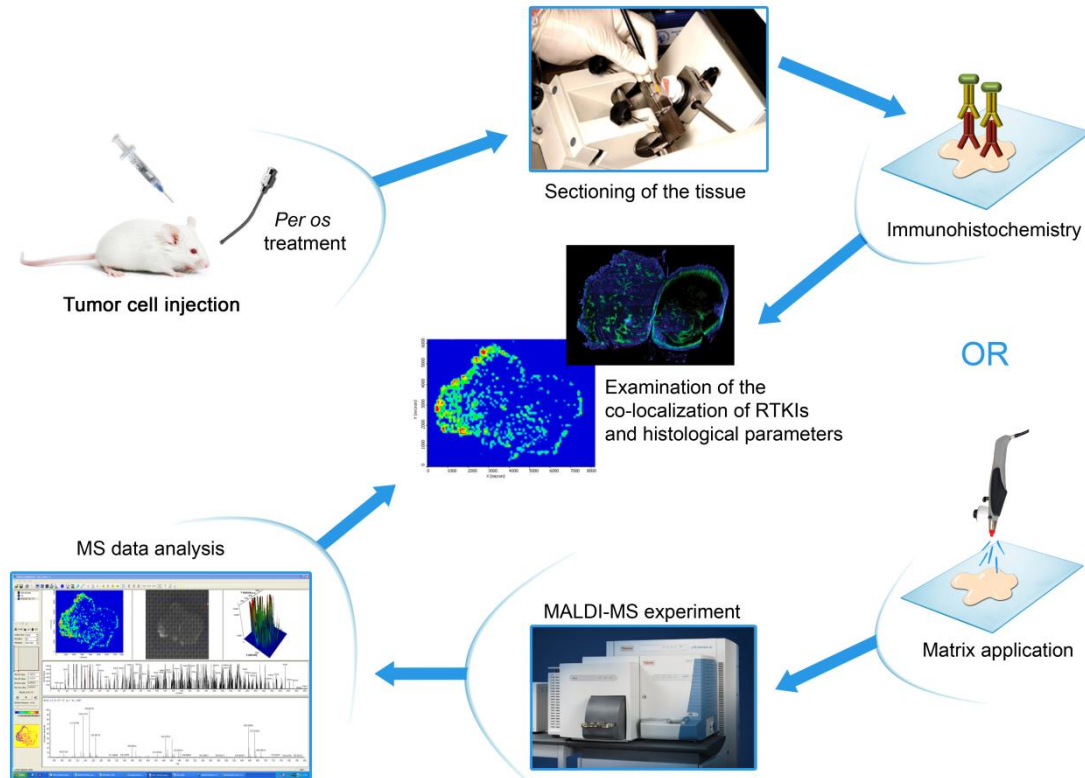


Figure 18. Flow chart of the study procedure. Nine serial frozen sections were cut from C26 or C38 mouse tumors treated with different antiangiogenic RTKIs. Sections were then used to analyse drug dispersal by MALDI-MSI, for subsequent HE staining and for immunolabeling with antibodies against CD31, laminin, desmin, α SMA and the target receptors of RTKIs. An additional slide was also used for hypoxia detection. After scanning the tissue sections, the antitumor and antivascular properties of RTKIs were correlated with their tumor tissue distribution data obtained by MALDI-MSI.

6. RESULTS

6.1 Tumor growth inhibition

A significant ($p=0.0194$) relative tumor growth inhibitor effect was shown after two weeks of RTKI treatment, when comparing the % change in tumor volume in the C26 model. Sunitinib was the only drug affecting tumor growth in this experimental setting (**Figure 19A.**).

In the C38 model two weeks of vatalanib treatment significantly reduced ($p=0.0173$) tumor burden (**Figure 19B.**).

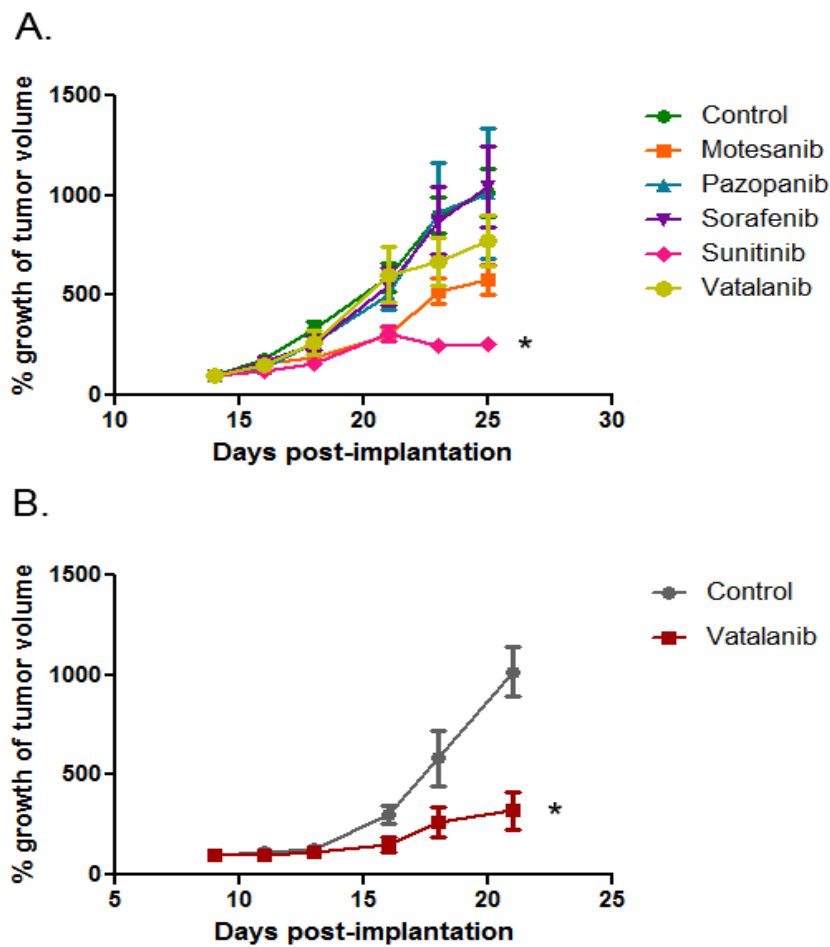


Figure 19. In vivo growth inhibition of different RTKIs in C26 and C38 tumors. (**A.**) Out of the five different antiangiogenic RTKIs (motesanib, pazopanib, sorafenib, sunitinib and vatalanib), only sunitinib reduced significantly the in vivo growth of C26 mouse colon adenocarcinoma cells in Balb/C mice, as shown by the one-way Anova test ($p=0.0194$) (**B.**) In contrast to the C26 model, vatalanib demonstrated a significant growth inhibitory effect in C57Bl/6 mice bearing C38 tumors, as shown by unpaired t-test ($p=0.0173$). Growth curves are means for six mice per group; bars, SEM.

In both models, control tumors weighted more than two times more than the ones treated with the effective compound (mean tumor weights 0.427g vs. 1.046g for sunitinib treated and control tumors, respectively in the C26 model and 0.811g vs. 2.146g for vatalanib treated and control mice, respectively in the C36 model, data not shown).

6.2 Immunohistochemical analysis

The growth of s.c. tumors in mice is known to be angiogenesis-dependent (413). Thus, in the next step of the experiment we tested, if the differences in tumor growth are in line with the effects of drugs on vascular parameters and the expression of target angiogenic receptors of the RTKIs. Therefore, we stained tissue sections for these parameters.

The expression level of antiangiogenic RTKs may significantly influence treatment response, and in turn, successful therapy can also regulate target receptor localization and function. In the C26 model expression of PDGFR α , - β and FGFR1 was observed not only on mural cells, but also on tumor cells. Expression patterns of PDGFR α , - β and FGFR-1 did not change in response to treatment with any of the compounds (p=0.8265, 0.1261 and 0.2983, respectively) in the C26 model (**Figure 20-22.**).

The receptor distribution of the C38 model showed a different pattern compared to the C26 tumors. C38 tumor cells did not express the aforementioned receptors, but a definite cell population expressing PDGFR α , - β and FGFR1 was detectable on the mural cells. Similarly to the C26 model, no change in the expression of PDGFR α , - β and FGFR1 (p=0.7601, 0.7497 and 0.7178, respectively) was seen (**Figure 20-22.**).

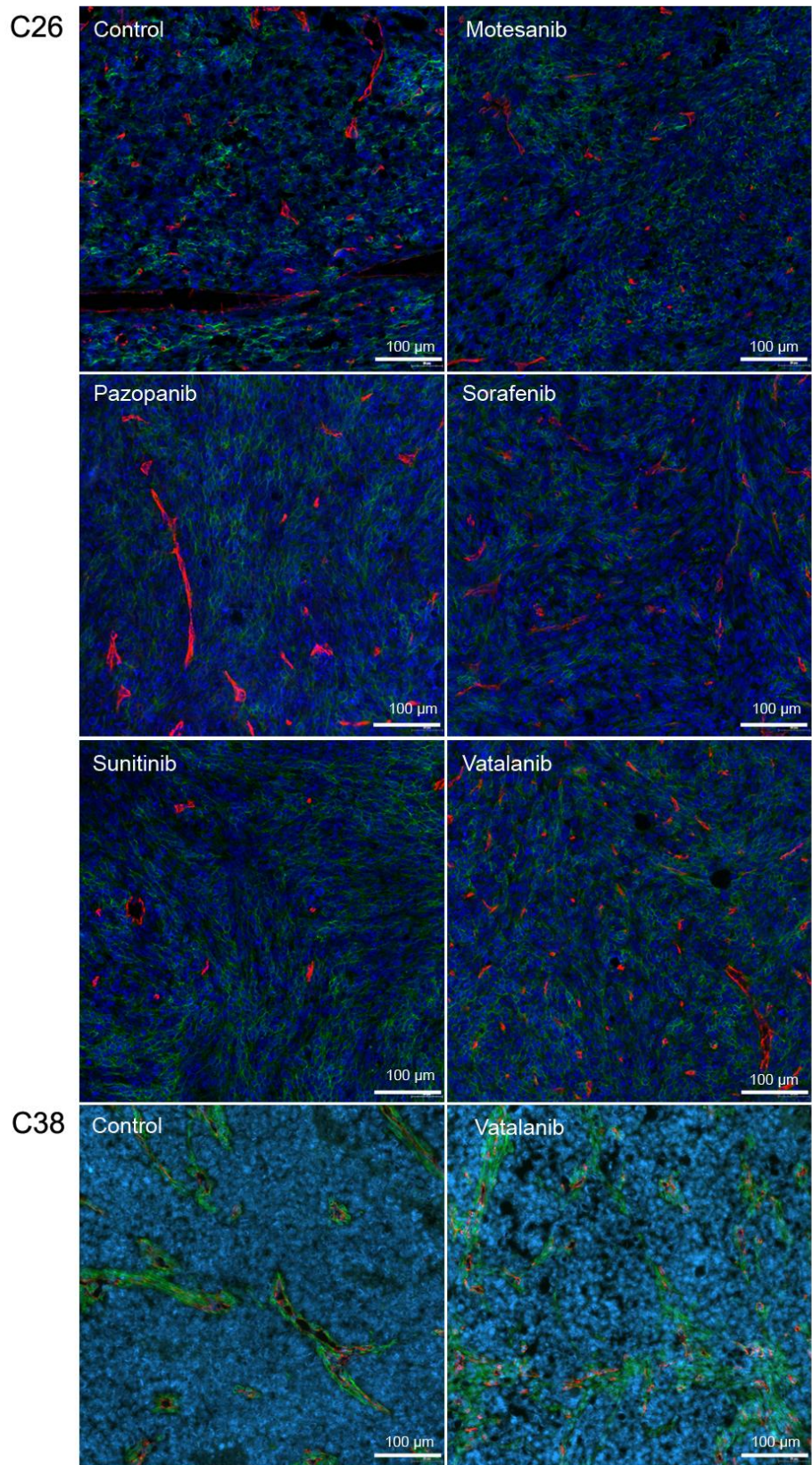


Figure 20. Low power views of C26 and C38 tumor sections stained for PDGFR α (green). Microvessels are labeled with anti-CD31 (red). Nuclei are stained with Hoechst 33342 (blue).

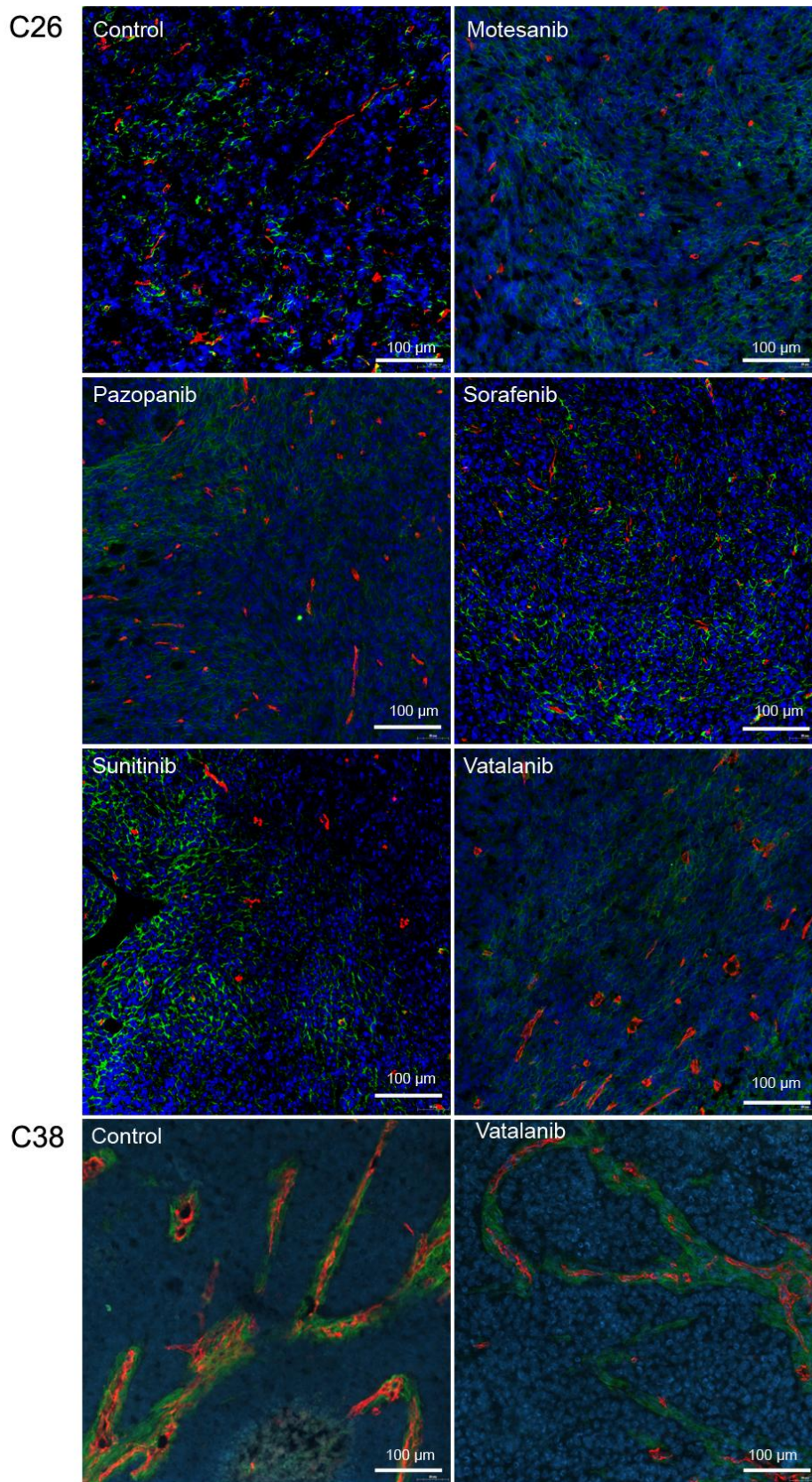


Figure 21. Low power views of C26 and C38 tumor sections stained for PDGFR β (green). Microvessels are labeled with anti-CD31 (red). Nuclei are stained with Hoechst 33342 (blue).

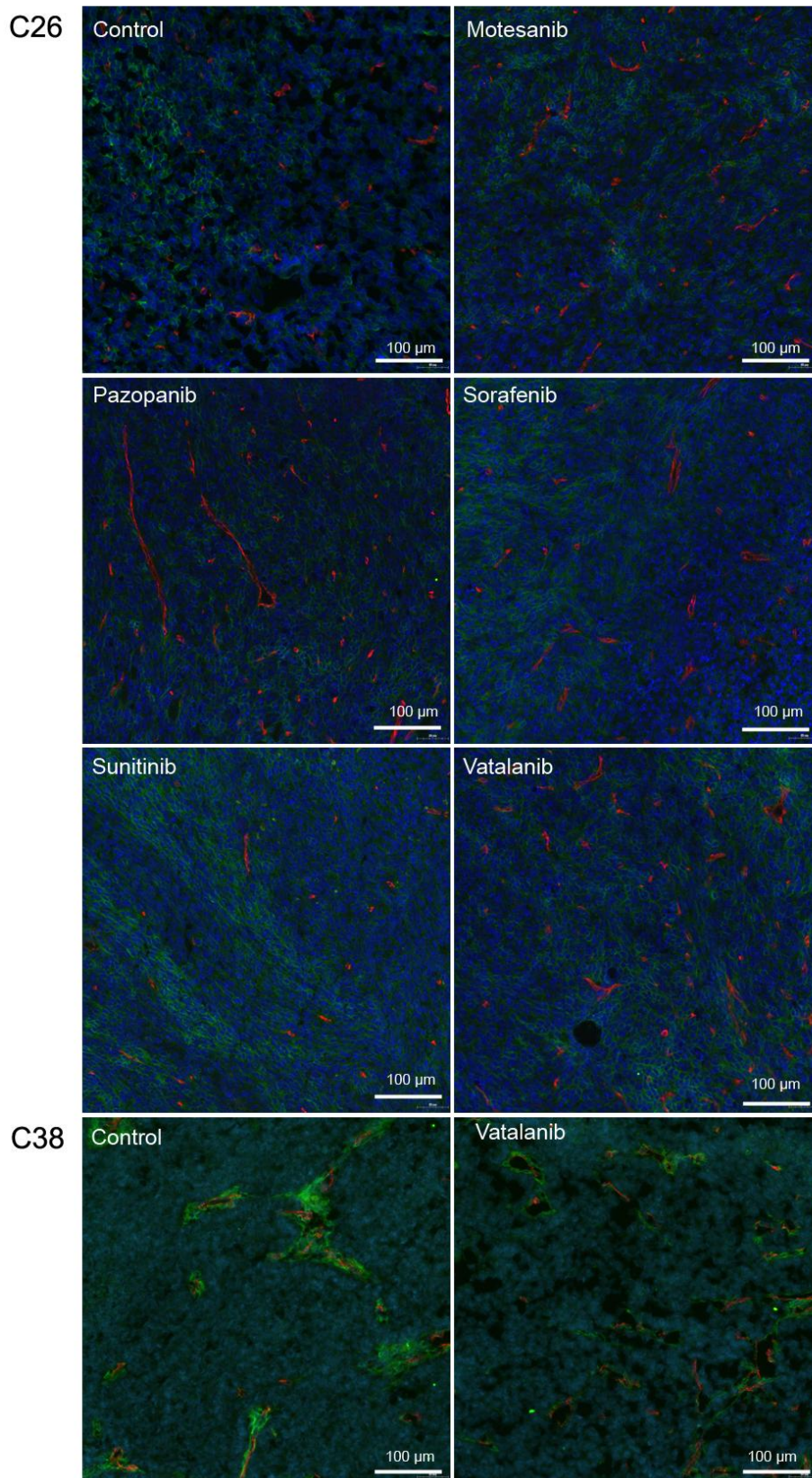


Figure 22. Low power views of C26 and C38 tumor sections stained for FGFR1 (green). Microvessels are labeled with anti-CD31 (red). Nuclei are stained with Hoechst 33342 (blue).

VEGFR2 expression was detected both on tumor and endothelial cells in the C26 model. Significant differences were shown both when counting the VEGFR2 signal and when measuring the area of VEGFR2+ cells in C26 tumors ($p=0.0296$ and 0.022 respectively; **Figure 23, 25**). Post-hoc test showed that VEGFR2 expression was altered only in the sunitinib treated group.

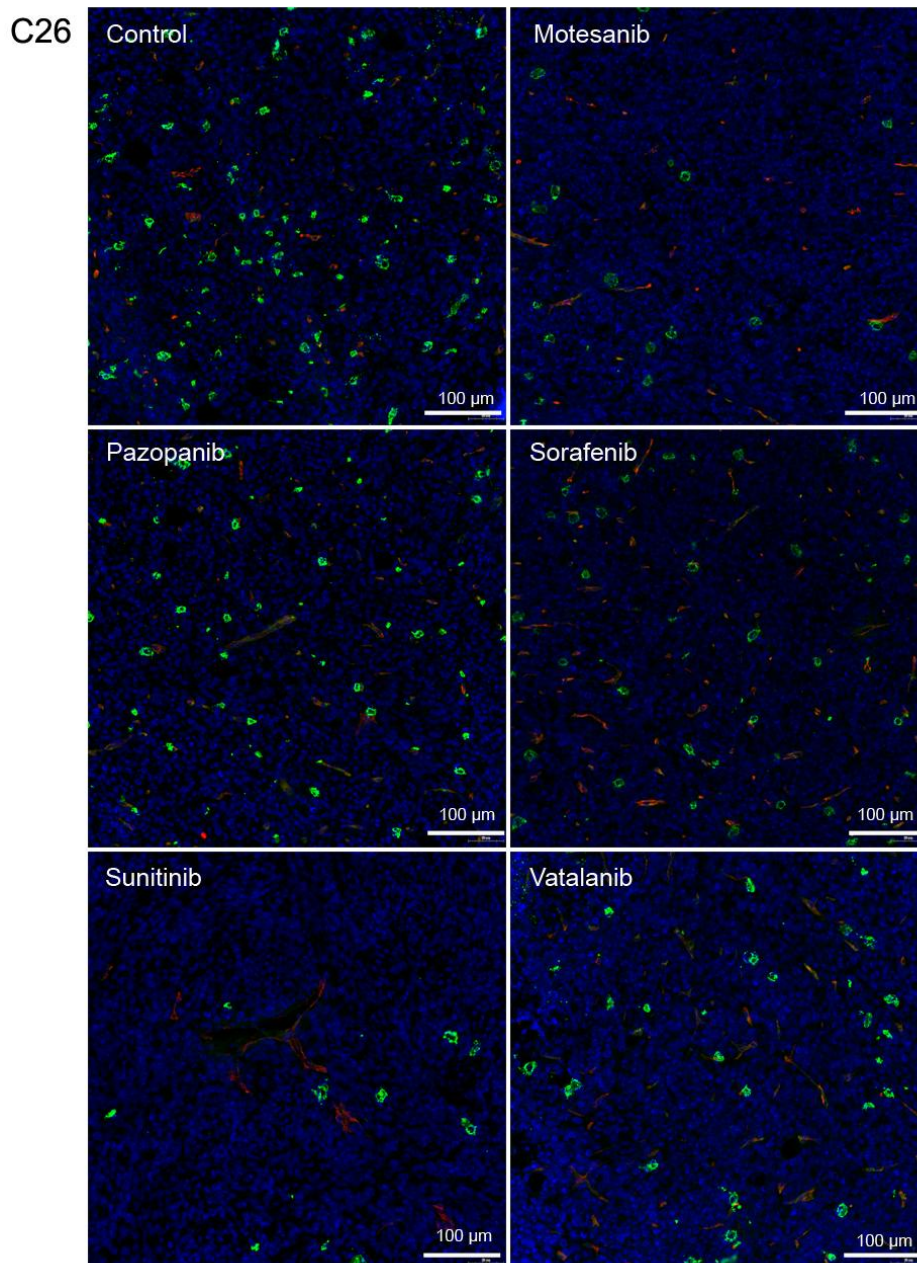


Figure 23. Low power views of C26 tumors stained for VEGFR2 (green). Microvessels are labeled with anti-CD31 (red). Nuclei are stained with Hoechst 33342 (blue).

C38 tumors were characterized by a weak VEGFR2 expression of the CD31+ endothelial layer, but no signal of the receptor on tumor cells was observed. No difference in the expression of VEGFR2 in the treated vs. non treated groups ($p=0.6857$ and 0.4857 for VEGFR2 density and area respectively) was detected (**Figure 24-25.**).

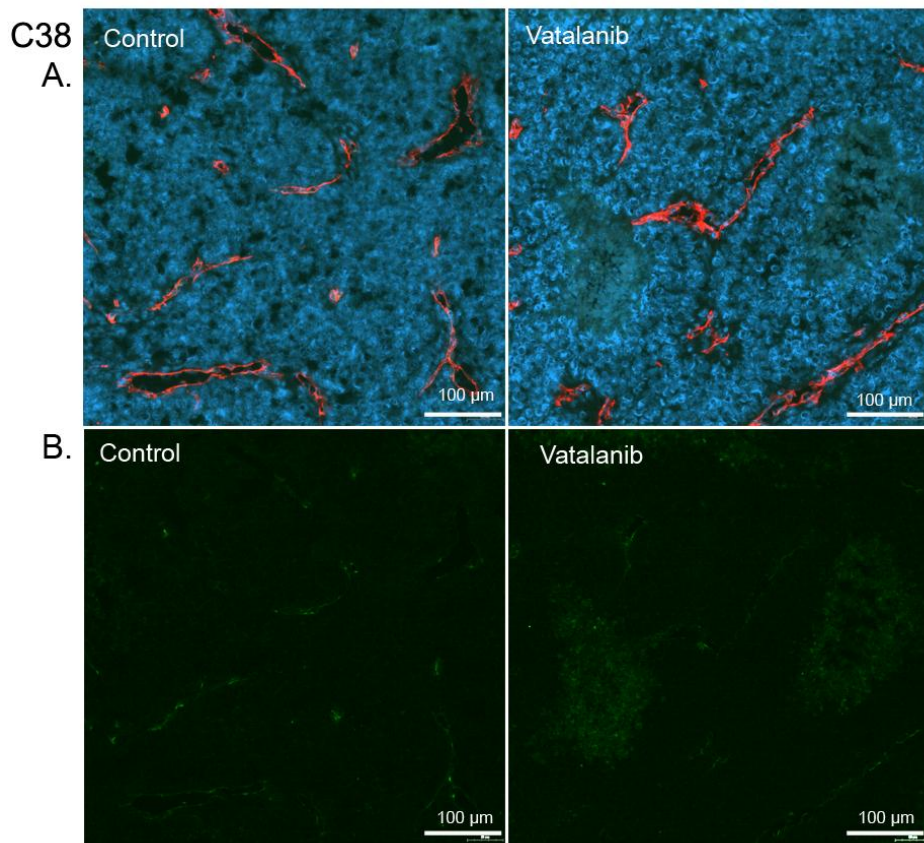


Figure 24. (A.) Low power views of C38 tumors stained for VEGFR2 (green). Microvessels are labeled with anti-CD31 (red). Nuclei are stained with Hoechst 33342 (blue). (B.) The same images without counterstain with anti-CD31 and Hoechst 33342.

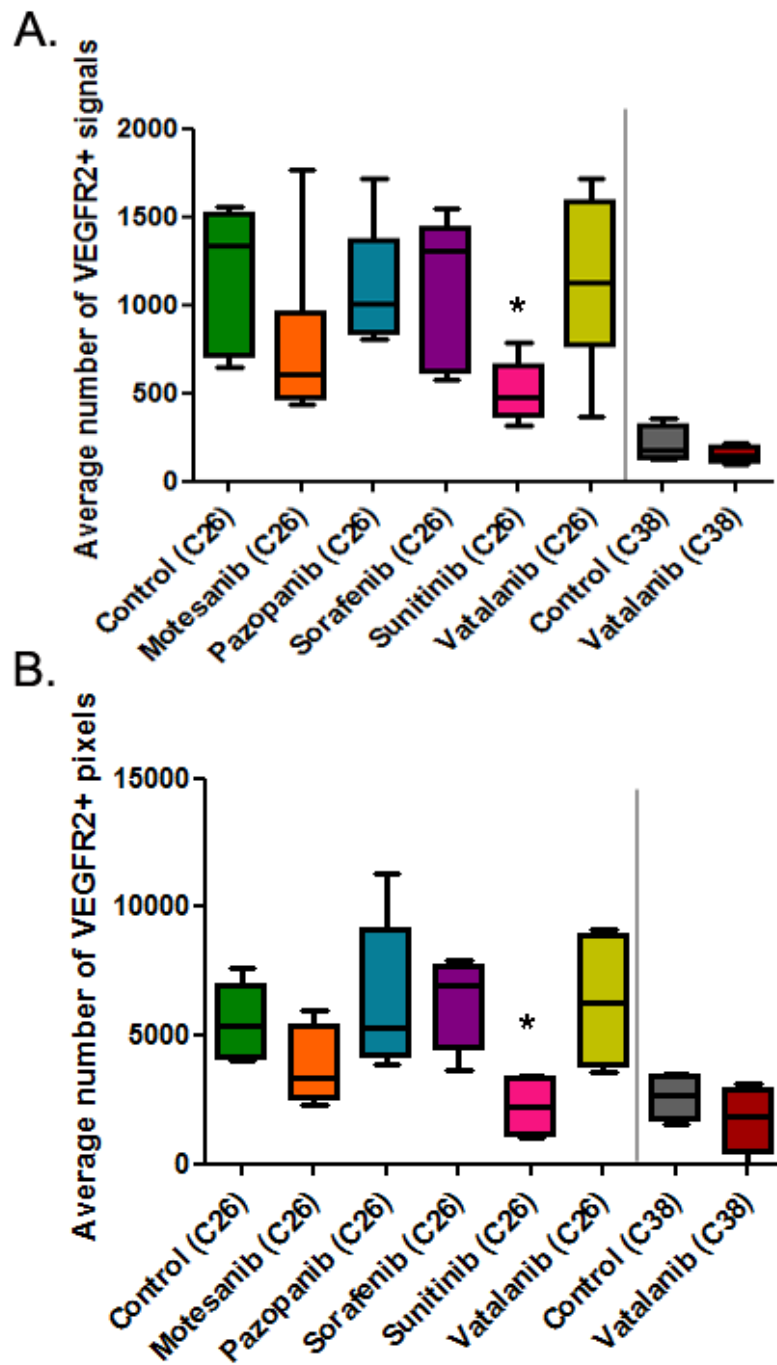


Figure 25. VEGFR2 (A.) densities and (B.) areas of C26 and C38 tumors. Data are shown as box (first and third quartiles) and whisker (maximum to minimum) plots with the mean (horizontal bar) from 6 animals per group. VEGFR2 densities were counted in ten viable intratumoral regions. VEGFR2 areas were calculated by counting the number of VEGFR2-positive pixels in ten viable intratumoral regions. $p=0.0296$ and 0.022 for VEGFR2 density and VEGFR2 area respectively in the C26 model as shown by the Kruskal-Wallis test, $p=0.6857$ and 0.4857 for VEGFR2 density and VEGFR2 area respectively in the C38 model as shown by the Mann Whitney U test.

We found a significant difference between the MVD of the C26 groups, $p < 0.0001$. The post-hoc test showed a suppressed MVD by sunitinib, motesanib and minimally vatalanib in that model. Microvessel area was also decreased in the sunitinib, motesanib and less intensively in the sorafenib treated group, $p < 0.0001$ (**Figure 26-27**).

In the C38 model however, no difference in the vessel density ($p = 0.235$), but in vessel area was detected, $p = 0.0341$ (**Figure 26-27**). It is also important to mention, that major differences in the vasculature of the two groups were observed. While C26 tumors had lots of small vessels, C38 tumors were characterized by only a few, but large and complex vascular structures.

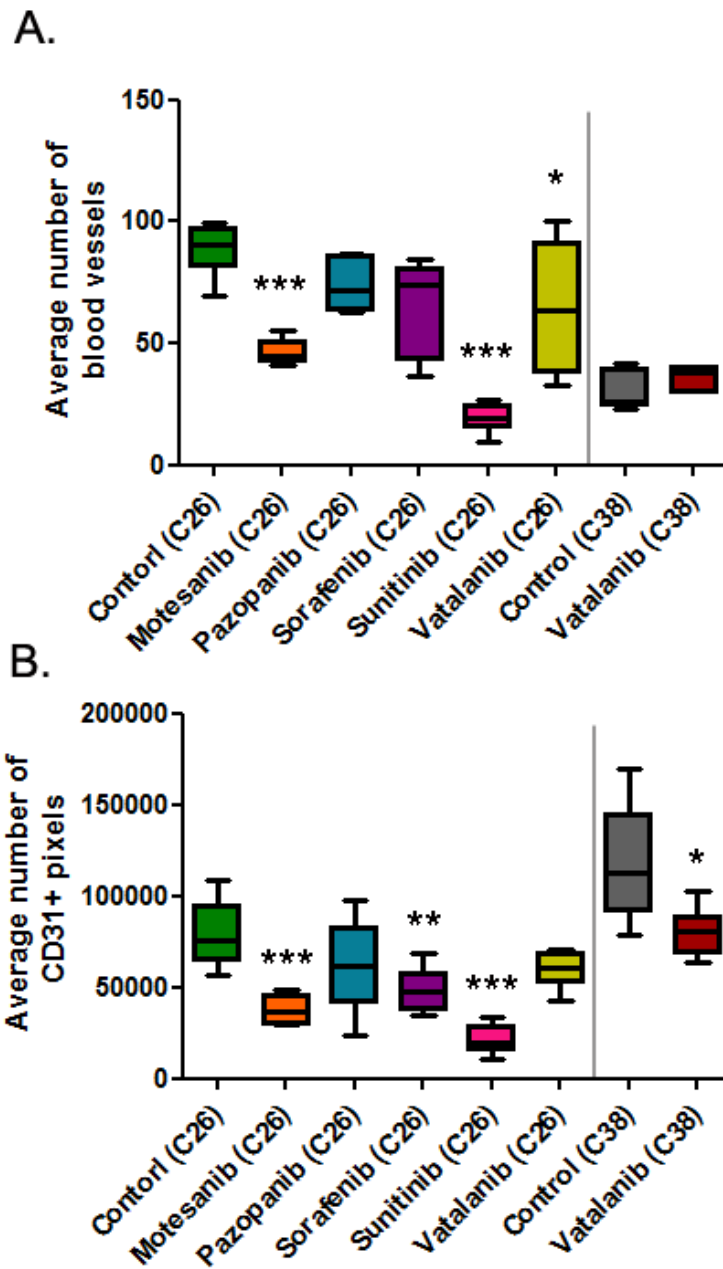


Figure 26. Microvessel density (A.) and microvessel area (B.) data of C26 and C38 tumors. Data are shown as box (first and third quartiles) and whisker (maximum to minimum) plots with the mean (horizontal bar) from 6 animals per group. Microvessel densities were counted in ten viable intratumoral regions. Microvessel areas were calculated by counting the number of CD31-positive pixels in ten viable intratumoral regions. $p < 0.0001$ both for MVD and vessel area in the C26 model as shown by the one-way Anova test, $p = 0.235$ and 0.0341 for vessel density and area respectively as shown by unpaired t-test in the C38 model.

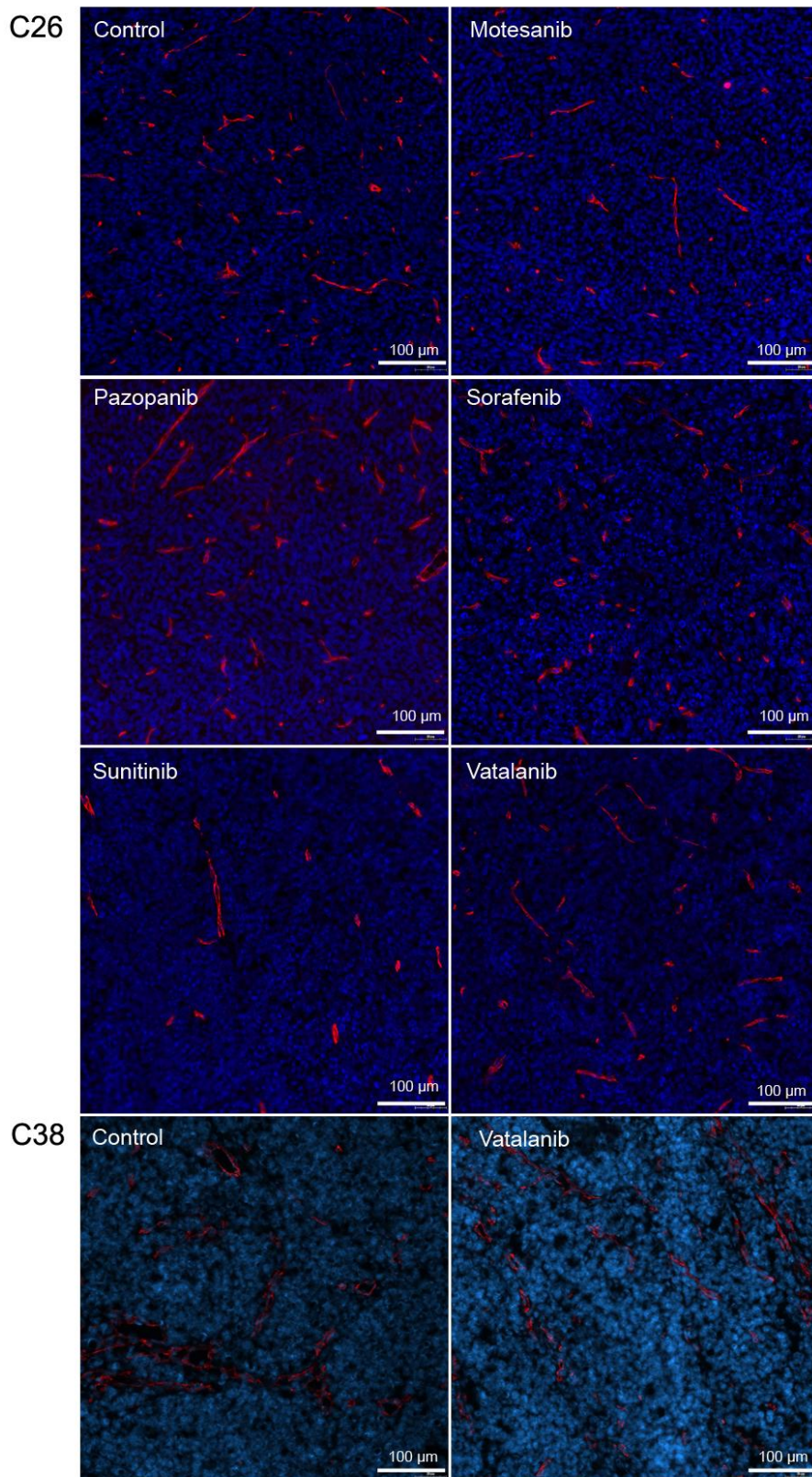


Figure 27. Microvessel density and area of C26 and C38 tumors. Tumors were labeled for the endothelial cell marker CD31 (red) and nuclei are stained with Hoechst 33342 (blue).

MVD clearly correlated with tumor oxygenation. Hypoxia was located in the less vascularized areas of the tumor. Accordingly, a significant increase ($p=0.0152$) in the intratumoral hypoxic areas was observed in the sunitinib treated group (**Figure 28-29**). The ratio of hypoxic areas did not differ ($p=0.9143$) in the C38 model (**Figure 28-29**).

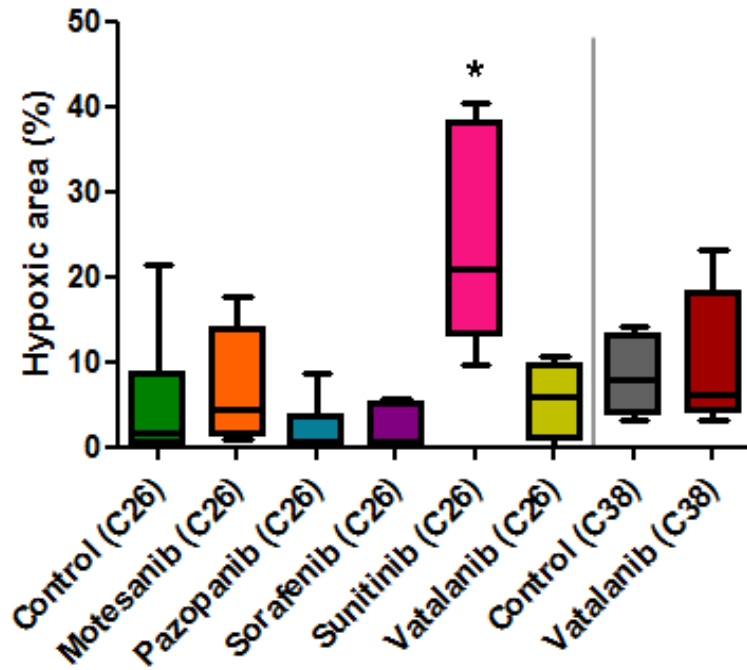


Figure 28. Hypoxic areas of C26 and C38 tumors. Data are shown as box (first and third quartiles) and whisker (maximum to minimum) plots with the mean (horizontal bar) from 6 animals per group. Hypoxic areas are shown in the percentage of the total tumor sections. $p=0.0152$ as shown by the Kruskal-Wallis test in the C26 model and 0.9143 as shown by the Mann-Whitney U test in the C38 model.

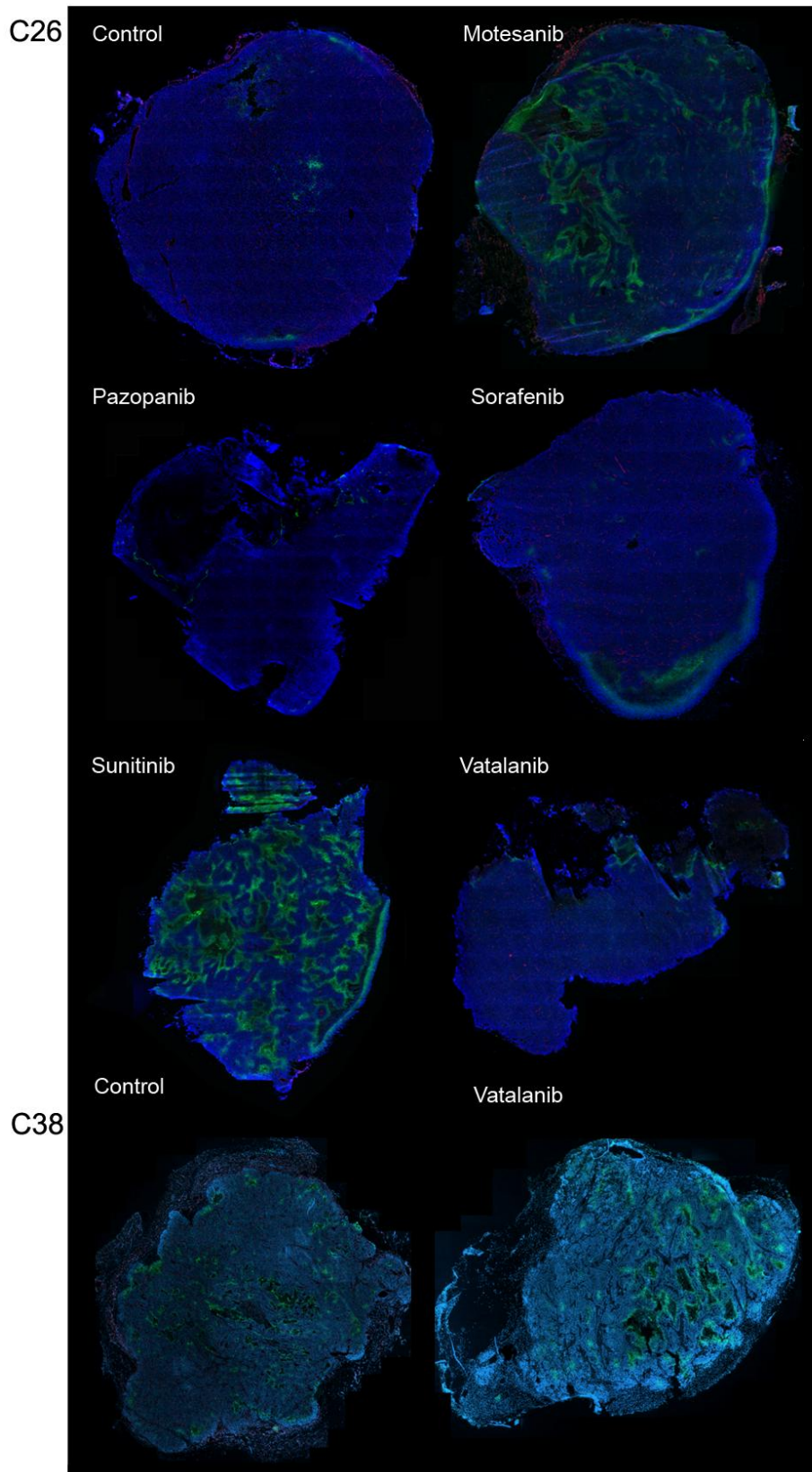


Figure 29. Representative images of hypoxic areas in the C26 and C38 tumors. Green: anti-pimonidazole staining for hypoxia; blue: nuclear staining with Hoechst 33342.

Beside the inhibition of endothelial cell proliferation, multi-target antiangiogenic RTKIs also influence PDGFR and FGFR positive pericyte and VSMC recruitment to tumor blood vessels. Therefore, the inhibition of these receptors may result in not only decreased MVD and consequently lower blood flow rate of tumors, but could also facilitate cancer cell metastatization. To observe the structural changes of the vasculature in response to treatment, we examined the expression of laminin, desmin and α SMA of tumor sections. While all vessels remained underlaid with a definite layer of laminin and covered with α SMA, desmin expression has decreased in response to sunitinib and motesanib treatment in the C26 model; $p=0.0135$ (**Figure 30-32.**).

Both laminin and α SMA expression were definite and did not change in response to treatment in the C38 model (**Figure 30-31.**), but unlike in C26, no difference in desmin expression was observed either $p=0.9143$ (**Figure 32-33.**).

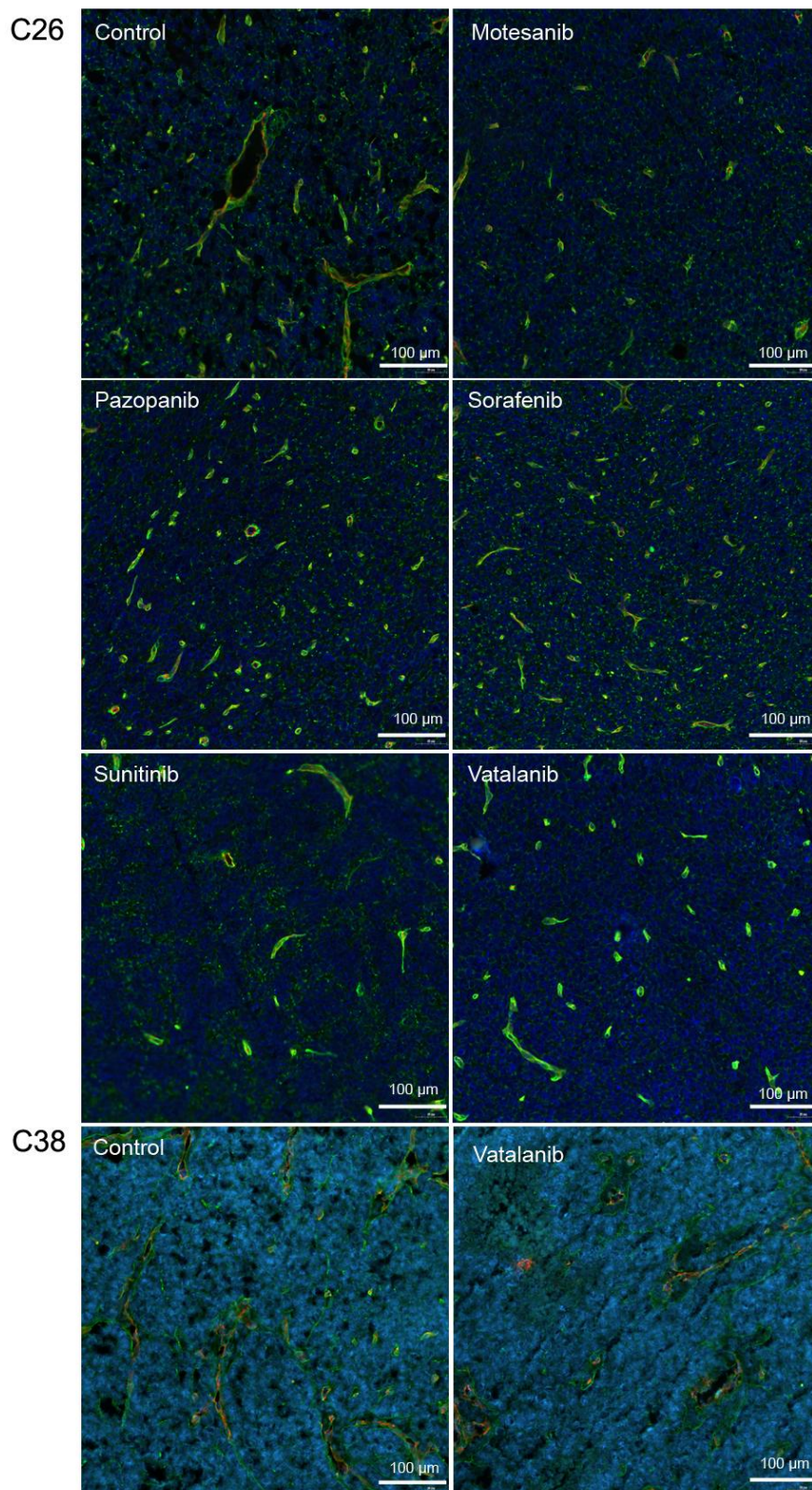


Figure 30. Low power views of C26 and C38 tumor sections stained for the capillary basement membrane component laminin (green) and CD31 (red). Nuclei are stained with Hoechst 33342 (blue). Note that tumor cells are also weakly positive for laminin.

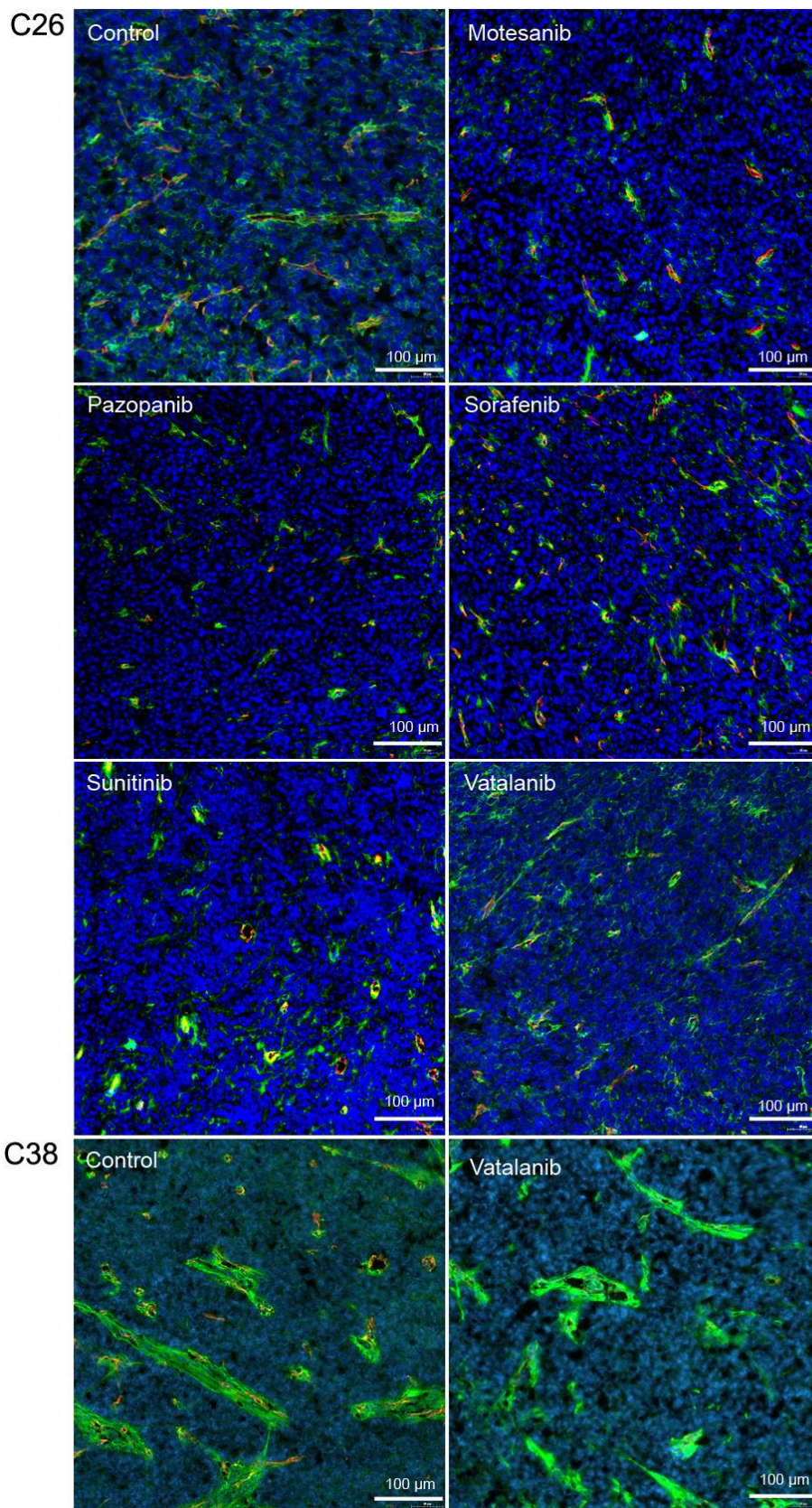


Figure 31. Low power views of C26 and C38 tumor sections stained for α SMA (green) and CD31 (red). Nuclei are stained with Hoechst 33342 (blue).

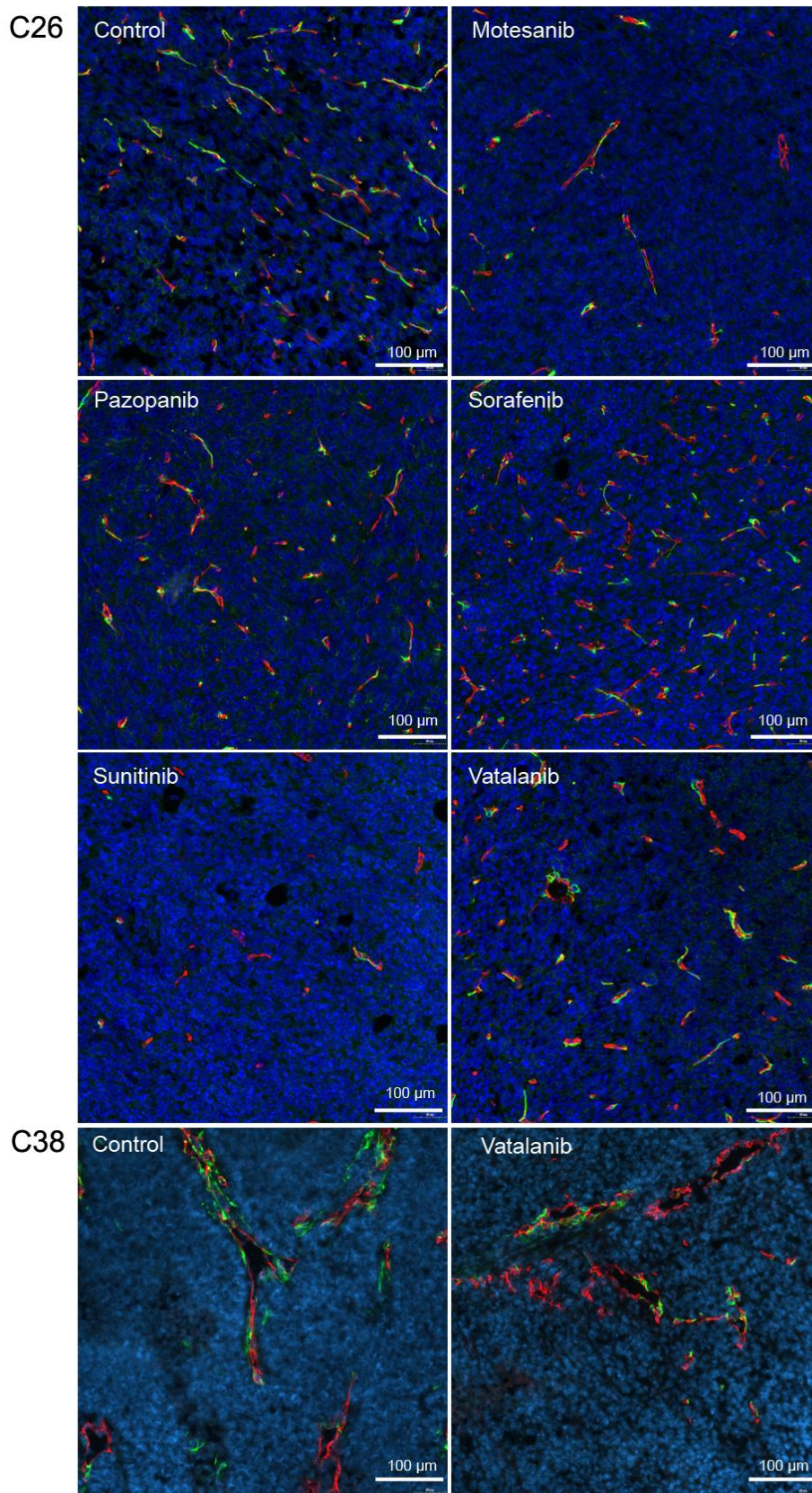


Figure 32. Desmin expression in C26 and C38 tumors. Tumor sections are immunolabeled for pericyte desmin (green) and CD31 (red). Nuclei are counterstained with Hoechst 33342 (blue).

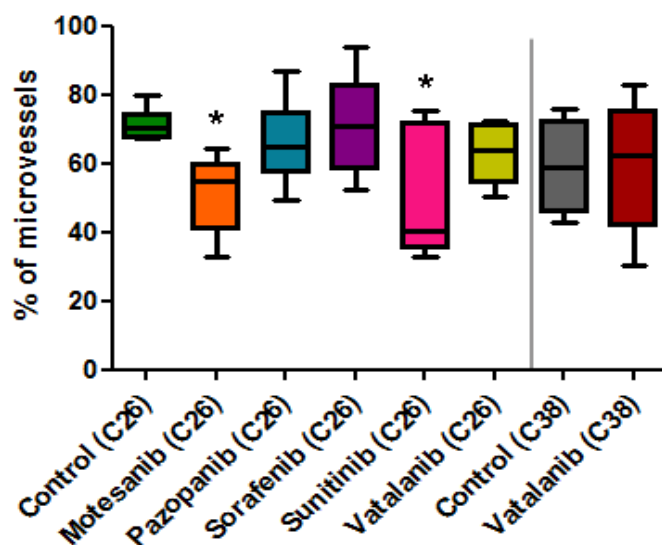


Figure 33. Desmin expression of C26 and C38 tumors. Data are shown as box (first and third quartiles) and whisker (maximum to minimum) plots with the mean (horizontal bar) from 6 animals per group. Desmin expression is expressed in the % of the microvessels covered by desmin expressing pericytes. $p=0.0135$ for the C26 model and 0.9143 for the C38 model as shown by the one-way Anova and the Mann-Whitney U tests respectively.

6.3 Mass spectrometric analysis

6.3.1 Compound characterization

To determine if these differences in the biological effects of the compounds arise from the differences in the intratumoral drug distribution, we have developed a method to characterize the antiangiogenic RTKIs in tissue samples.

First, the m/z value of the quasimolecular ion and fragmentation pattern of each drug compound were defined on a MALDI stainless steel target plate.

The quasimolecular ion of motesanib was detected at m/z 374.199. Fragmentation of the molecule resulted in ions at m/z 212.1, which corresponds to the split at the amide bond of nicotinamide. The fragment ions corresponding to the indoline formamide moiety were identified at m/z 189.1. Cleavage of the pyridine moiety and charge retention resulted fragment ions at m/z 163.1.

Pazopanib was detected with a quasimolecular ion at m/z 438.17. Subsequent MS/MS fragmentation of the precursor ions led to the loss of the amidogen group, generating fragment ions at m/z 421.1. Further loss of the sulfur dioxide eventuated fragment ions

at m/z 357.1, while the presence of fragment ions at m/z 342.1 indicated the loss of an additional methyl group.

The quasimolecular ion of sorafenib was found at m/z 465.093. Dehydroxylation of the formamide moiety and subsequent bond retention resulted in fragment ions at m/z 447.1, while cleavage of the pyridine ring eventuated fragment ions at m/z 425.1. Fragmentation of the molecule also led to detection of ions at m/z 270.2, corresponding to the loss of the chloro-trifluoromethyl-phenylamine group. Presence of fragment ions at m/z 252.2 indicated the cleavage of the chloro-trifluoromethyl-phenyl ring and the scission of the carboxamide group.

Sunitinib was identified with a quasimolecular ion at m/z 399.218. Subsequent MS/MS fragmentation of the precursor ions led to the loss of the terminal diethylamino group, generating fragment ions at m/z 326.1, while the presence of fragment ions at m/z 283.1 indicated a cleavage at the amide group.

The quasimolecular ion of vatalanib was seen at m/z 347.105. Scission of the benzene ring resulted in the generation of ions at m/z 320.2. Loss of the chloride eventuated fragment ions at m/z 311.2. Decomposition of the phthalazin-amine ring led to the detection of ions at m/z 294.2. Loss of the pyridine moiety resulted in fragment ions at m/z 268.1, while cleavage of the methylpyridine group indicated the detection of ions at m/z 254.1. Loss of the chlorophenyl-amine group resulted in fragment ions at m/z 220.2.

When applied to the tissue surface, drug molecules showed similar ionization and fragmentation properties to those generated on the MALDI plate. The same quasimolecular ions and fragment ions detected on both surfaces are shown in **Table 4**.

Table 4. Chemical properties of the studied drugs

	Motesanib	Pazopanib	Sorafenib	Sunitinib	Vatalanib
Chemical formula	C ₂₂ H ₂₃ N ₅ O	C ₂₁ H ₂₃ N ₇ O ₂ S	C ₂₁ H ₁₆ ClF ₃ N ₄ O ₃	C ₂₂ H ₂₇ FN ₄ O ₂	C ₂₀ H ₁₅ ClN ₄
Molecular weight (g/mol)	373.45	437.52	464.82	398.47	346.81
Quasimolecular ion [M+H]⁺ (m/z)	374.199	438.170	465.093	399.218	347.105
Fragment ions (m/z)	212.1;189.1; 163.1	421.1; 357.1; 342.1	447.1; 425.1; 270.2; 252.2	326.1; 283.1	320.2; 311.2; 294.2; 268.1; 254.1; 220.2

6.3.2 Precursor compound and metabolite detection in the blood

Adsorption of the drugs was examined in the peripheral blood, drawn just before sacrificing the animals. In both models all applied drugs absorbed successfully with notable signal intensities being observed in the peripheral blood.

Moreover, all so far identified metabolites of motesanib (414) could also be characterized in blood samples, however, 2-amino nicotinamide metabolite (*m/z* 283.157), the lactam form of this metabolite (*m/z* 297.137), the carbinolamine metabolite (*m/z* 372.184), and the oxindole metabolite (*m/z* 388.179) were found with high signal intensities, reaching 5-90% of the signal intensity of the precursor compound (**Figure 34**).

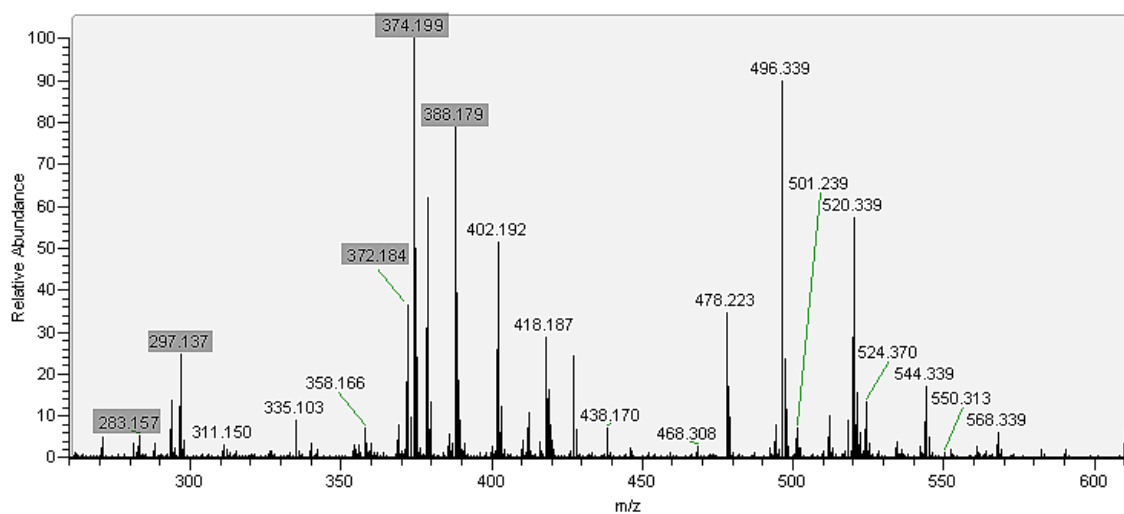


Figure 34. A representative mass spectrum of a blood sample taken from a motesanib-treated mouse. Marked are peaks of motesanib and its main metabolites.

Metabolization of pazopanib is less remarkable. Indeed, although all so far detected (415) metabolites of pazopanib were traceable, but none of them reached 5% of the signal intensity of the precursor pazopanib (**Figure 35.**).

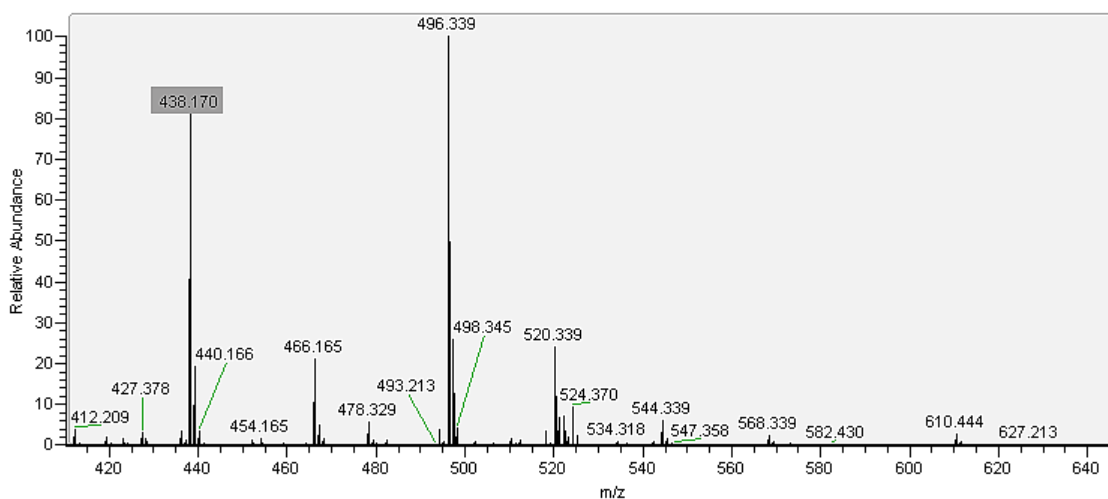


Figure 35. A representative mass spectrum of a blood sample taken from a pazopanib-treated mouse. Peak of pazopanib is marked.

Sorafenib was present with the lowest signal intensity in the blood samples. Moreover, metabolization of sorafenib is even less known than that of pazopanib. Neither the N-oxide, nor the glucuronide metabolite of sorafenib was reliably detected, but the desmethylated metabolite (m/z 451.078) was traceable (**Figure 36.**).

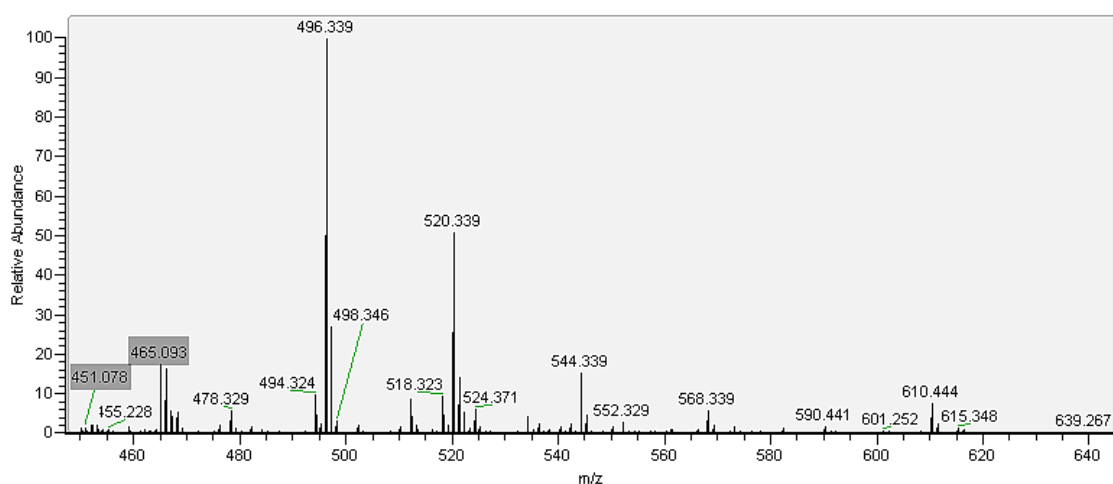


Figure 36. A representative mass spectrum of a blood sample taken from a sorafenib-treated mouse. Marked are peaks of sorafenib and its main metabolite.

Sunitinib was measured in all plasma samples, moreover, all metabolites of the precursor compound were also traceable and could be characterized. Presumed structures and MS/MS spectra of the precursor compound and its metabolites in blood plasma are presented in **Figure 37**.

The previously described bis-desethylated metabolite (M1) of sunitinib (349), with the quasimolecular ion at m/z 343.000 could be detected only in a few blood samples performing full mass scans. However, isolating and fragmenting the proposed peak of that metabolite resulted in fragment ions at m/z 326.2 and 283.1 in all samples. Stepwise elevation of the collision energy proved that the detected fragment ions are formed by the fragmentation of M1. The missing precursor ion in full mass spectra may be explained by the low concentration of M1 that appeared to be below the detection limit of the FT analyser compared to the linear ion trap.

The signal generated at m/z 358.126 of M2 indicates the loss of the terminal diethylamine group, with the oxidation of the molecule. This resulted in fragment ions at m/z 283.1 but not at m/z 326.1. The presence of fragment ions at m/z 340.2 refers to the terminal dehydroxilation of the molecule.

M3, an active metabolite of sunitinib (SU012662) (416) was formed by the mono-desethylation of the molecule, resulting a quasimolecular ion at m/z 371.188 and the same fragment ions as sunitinib.

Two mono-hydroxylated variations of the active metabolite were detected at m/z 387.182. M4 was modified at the indolylidene-dimethylpyrrole moiety, resulting fragment ions at m/z 342.2 and 299.1. M5 was hydroxylated at the carbon next to the amide nitrogen, which generated fragment ions at m/z 283.1. The detected fragment ion peak at m/z 369.2 could be derived from both molecules by dehydroxylation.

Loss of two hydrogen atoms of the terminal ethyl group of sunitinib eventuated in a metabolite (M6) at m/z 397.203. Fragmentation of the molecule generated ions at m/z 326.1 and 283.1.

Fragment ions of a previously described metabolite with the quasimolecular ion at m/z 397.224 (M7) could also be detected by MS/MS (349). Signals of fragments were generated at m/z 324.2 and 281.2, suggesting defluorination and subsequent dehydroxylation of the molecule. M7 was not traceable by full MS, probably because of the signal suppression of M6 at m/z 397.203.

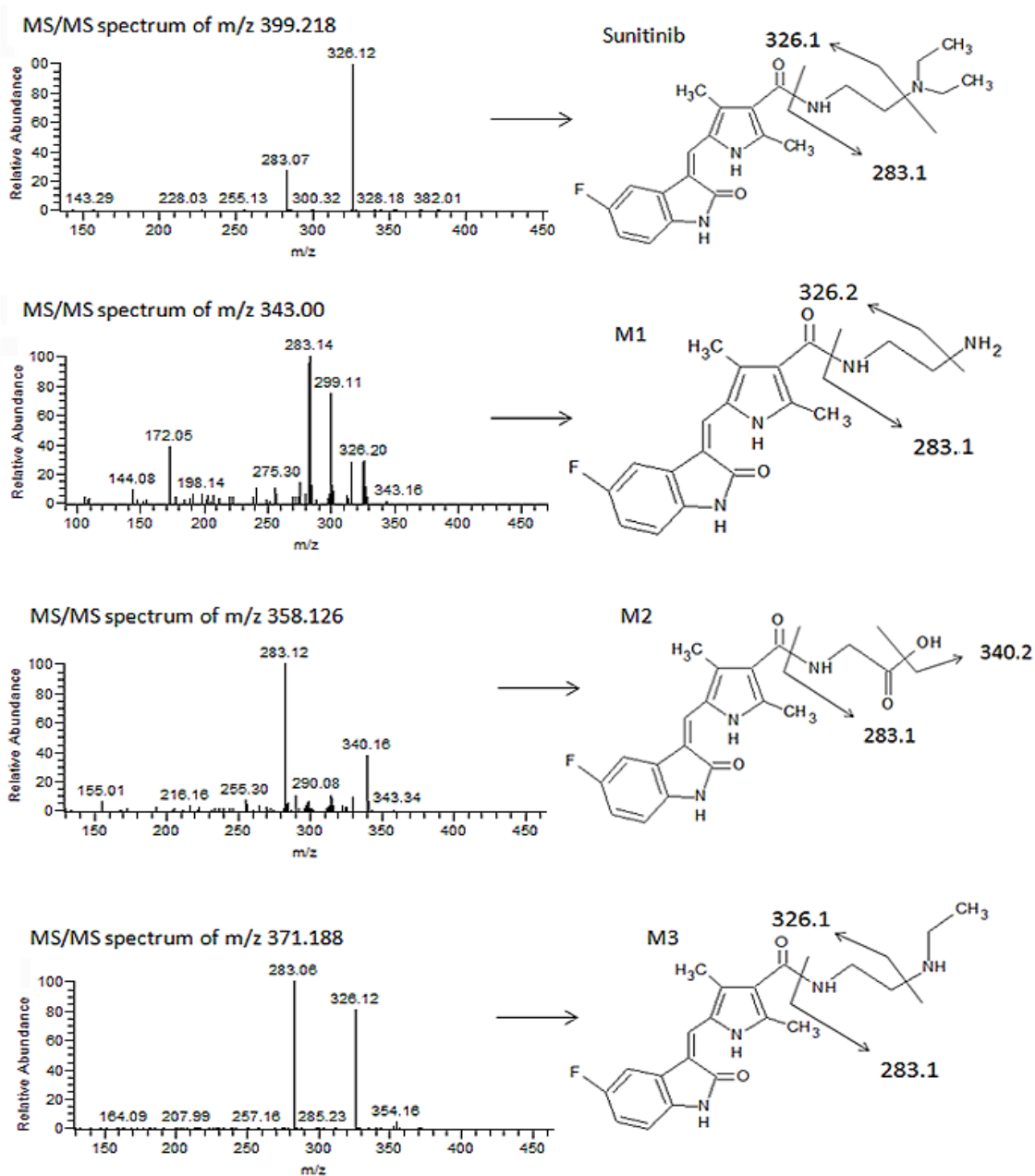
Similarly to M1, the saturated metabolite of sunitinib, M8, was detected by Speed et al. at m/z 401.00 in rat and monkey feces (349). This could only rarely be measured in our mouse model by full MS. However, when isolating the presumed metabolite peak, the detected fragment ions at m/z 285.1 and 328.2 indicated the presence of the molecule, and that the saturation occurred at the indolylidene-dimethylpyrrole moiety.

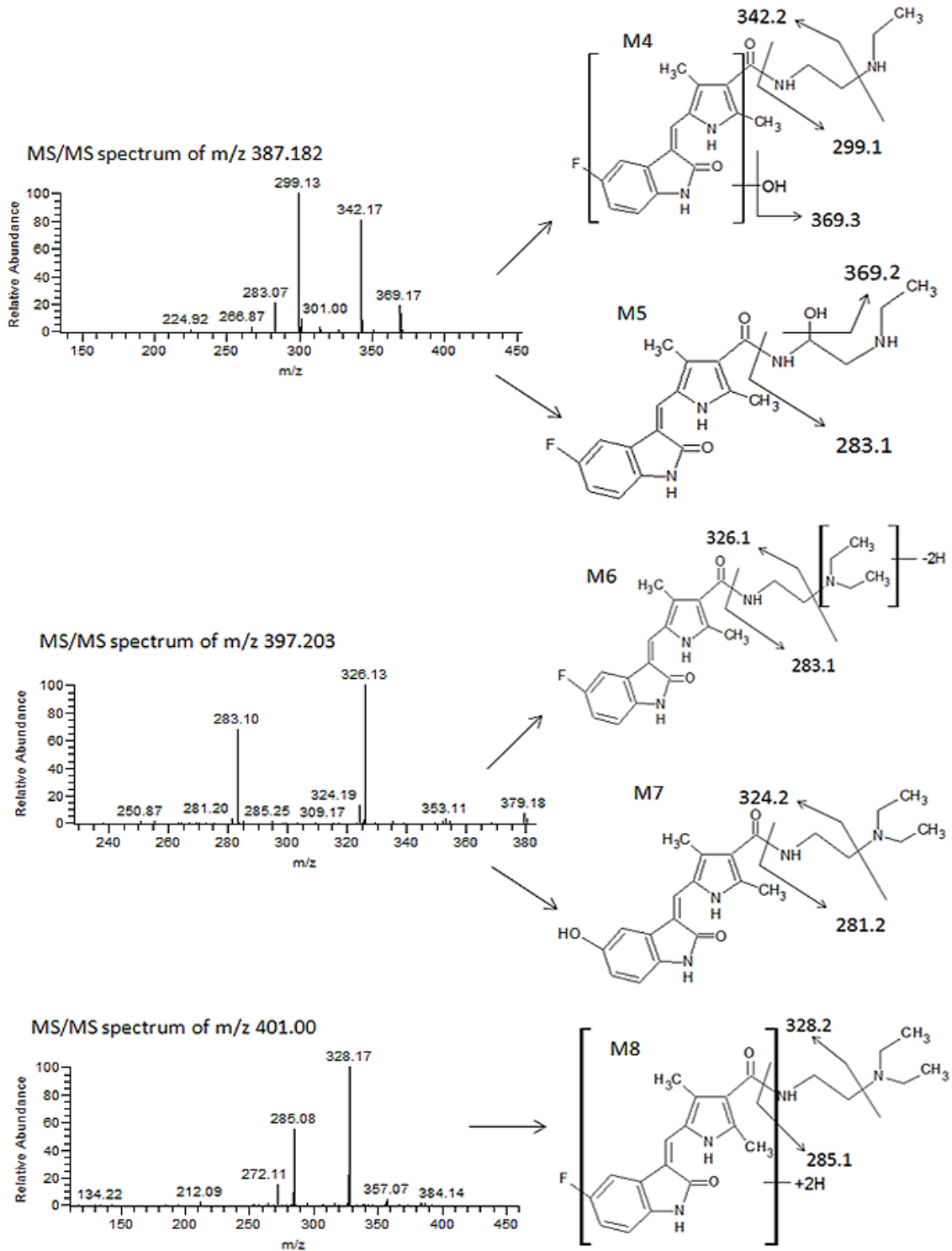
Mono-hydroxylated metabolites of sunitinib were also measured at m/z 415.214. Fragmentation of the molecule indicated the oxidation on the indolylidene-dimethylpyrrole group (M9) with 16 Da higher fragments than the corresponding ions of sunitinib at m/z 342.2 and 299.2. Moreover, upon fragmentation of the detected metabolite peak, ions at m/z 326.1 and 283.1 were also formed, indicating that the oxidation occurred either at one of the terminal carbons of the diethylamine group (M10) or at the amine moiety (M11). M11 was previously synthesized as SU012487 (349). Dehydroxylation of any of the mono-hydroxylated metabolites could result in fragment ions at m/z 397.1.

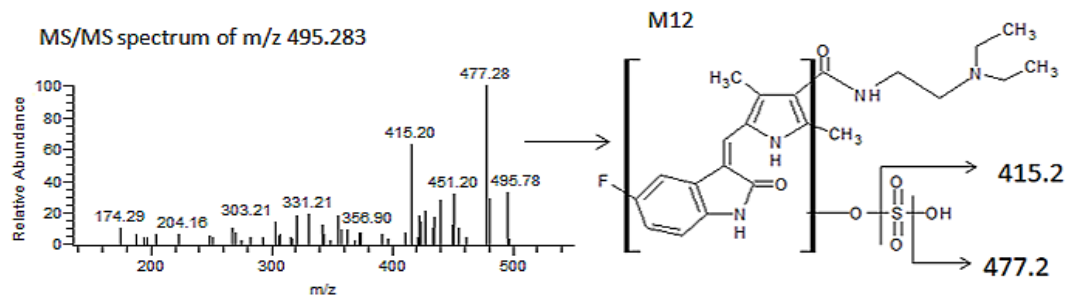
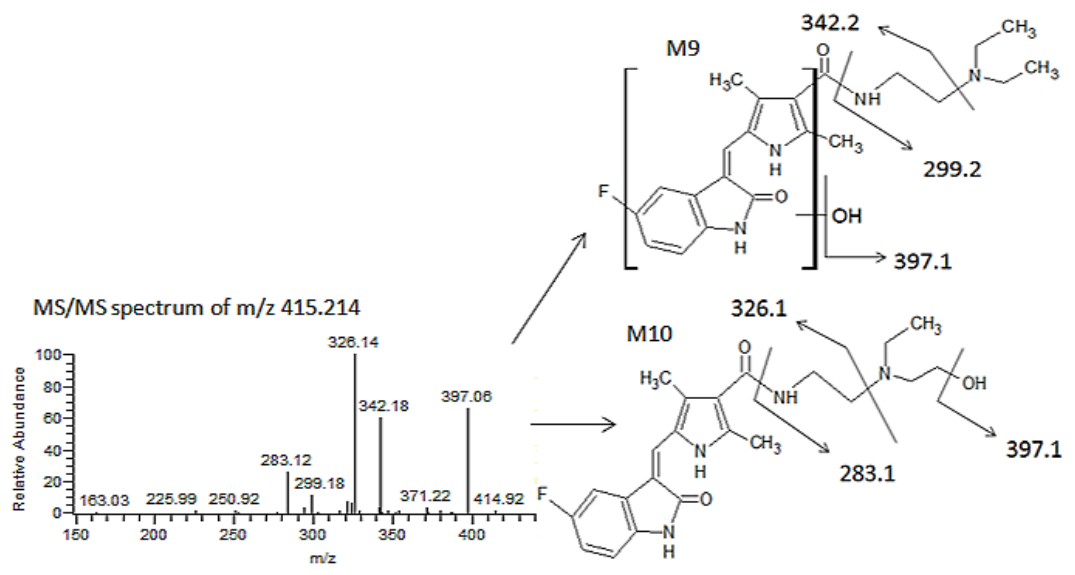
M12 at m/z 495.283 was identified as a sulphate conjugate of M9. Desulphuration of the molecule eventuated in fragment ions at m/z 415.2, while dehydroxylation resulted in fragment ions at m/z 477.2.

The glucuronide metabolite, M13, was detected at m/z 575.252. The cleavage at the amide group and the loss of the terminal diethylamino moiety resulted in fragment ions at m/z 459.2 and 502.2, respectively.

The metabolite at m/z 591.243 (M14) was generated by both the oxidation and the glucuronidation of sunitinib. When the molecule fragmented as the unmodified compound, ions at m/z 518.2 and 475.1 were generated. Dehydroxilation eventuated in a signal at m/z 573.2, while fragment ions at m/z 415.2 were formed by the loss of the dehydrated glucuronic acid. Deglucuronidation and dehydroxilation of the molecule resulted in ions at m/z 342.2.







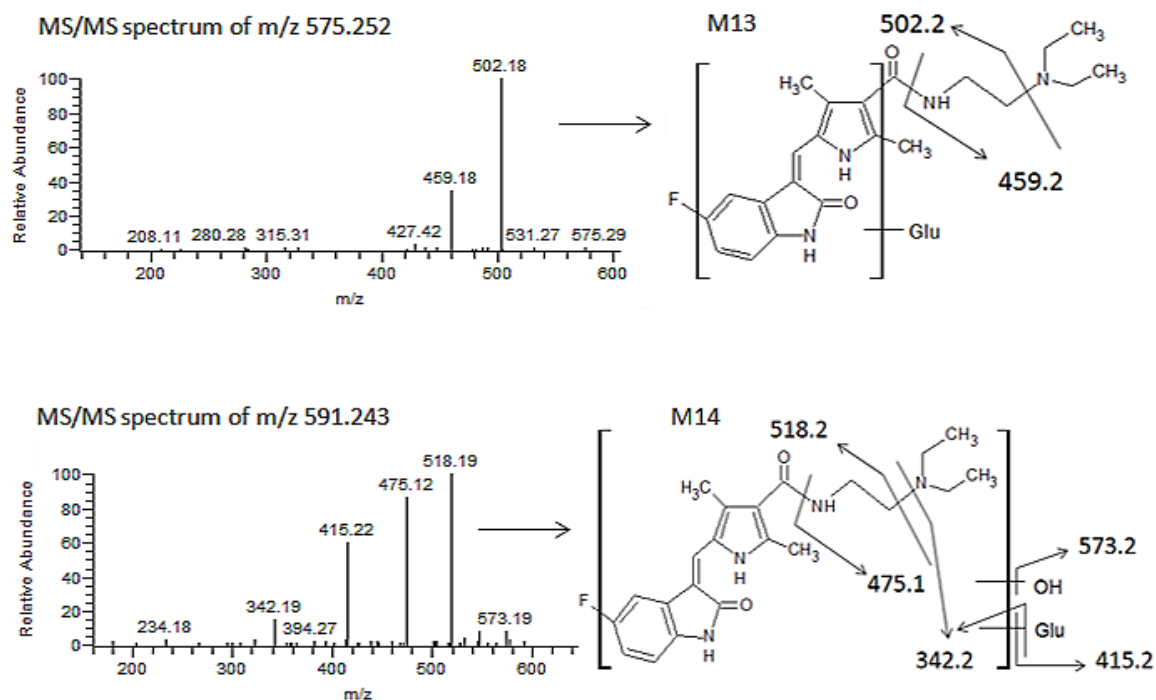


Figure 37. Detection of sunitinib and its metabolites in blood samples. MS/MS spectra of sunitinib and its metabolites with the proposed structure and fragmentation properties.

M3, the active metabolite generated 2-3-fold less intensive signal than the precursor molecule in blood samples. All the other metabolites were only traceable, with less than 5% of the signal intensity of the unmodified compound (data not shown).

Vatalanib also highly metabolized as observed in the blood samples, however, metabolites were mainly traceable, and the signal intensity of only the main oxydative metabolite (m/z 363.1) reached the 30% of the precursor compound. No difference in the signal intensities and metabolization pattern of vatalanib in the blood samples taken from the Balb/C and the C57black/6 mice was detected (**Figure 38**).

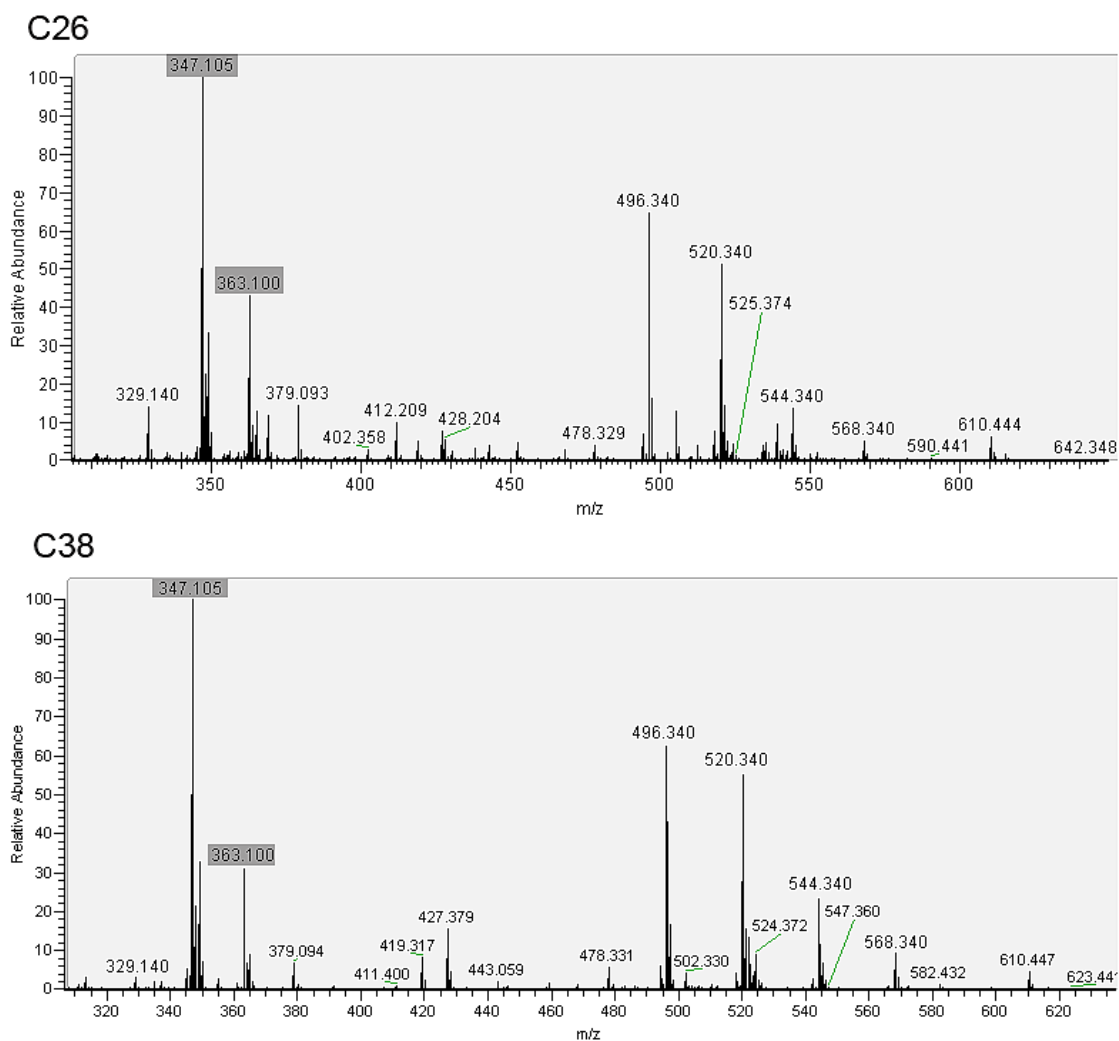


Figure 38. Representative mass spectra of blood samples of mice bearing C26 or C38 tumors, and treated with vatalanib. Marked are peaks of vatalanib and its main metabolite.

6.3.3 Tissue imaging of antiangiogenic RTKIs

Calibration of the drug molecules resulted in linear correlation between concentration and normalized average signal intensity for all compounds in the examined concentration range (**Figure 39**).

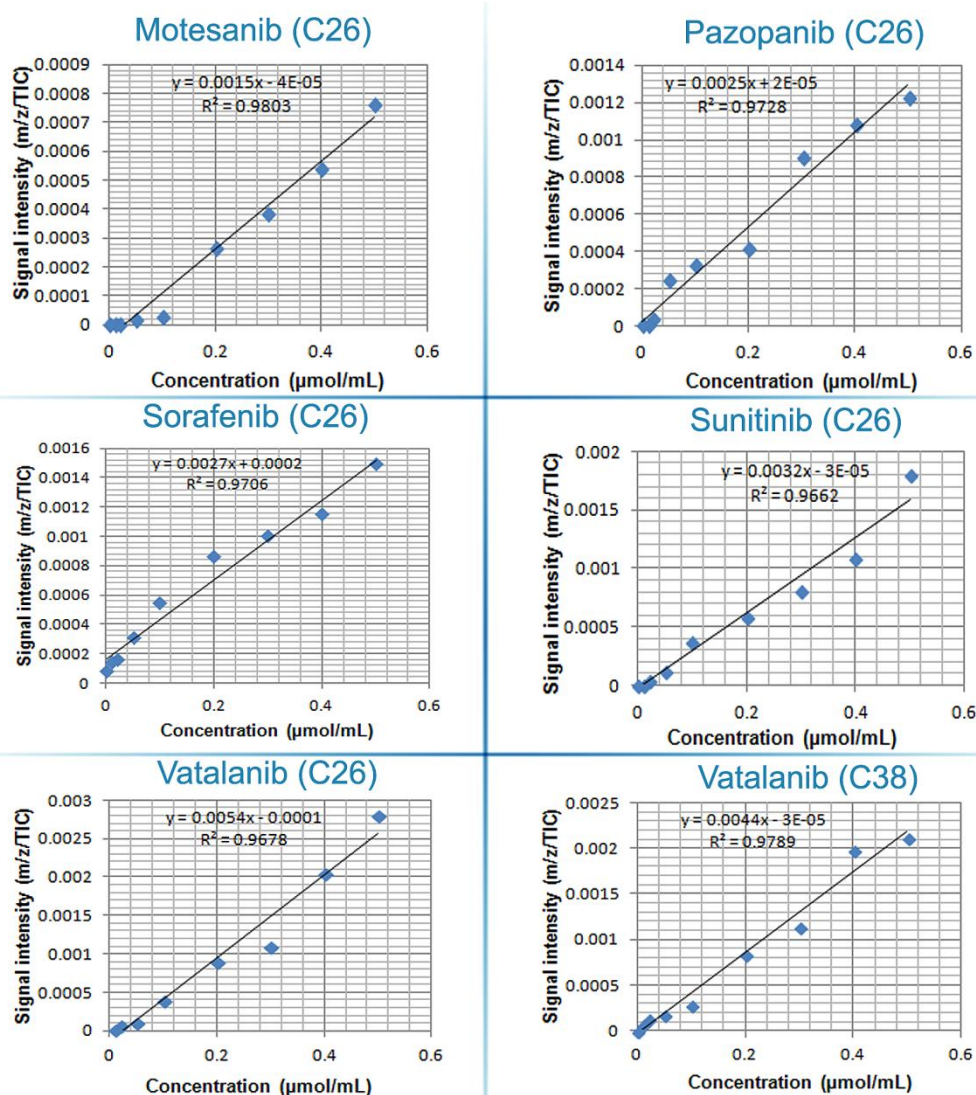


Figure 39. Calibration curves of antiangiogenic RTKIs. Drugs were dissolved and diluted in 50% methanol in the concentration range of 0.001–0.5 $\mu\text{mol}\cdot\text{mL}^{-1}$. One microliters of the compound solutions were applied on control tumor tissue surfaces. Spraying and detection conditions were the same as those during the analysis of in vivo-treated tumors. Average signal intensities of the applied concentrations were measured and normalized to TIC by using Xcalibur v 2.0.7. and ImageQuest™ softwares.

Based on the calibration curves, average signal intensities were translated into drug concentration ($\mu\text{mol}/\text{mL}$) data of C26 and C38 tumors. While intratumoral sorafenib and vatalanib levels did not differ between drug-treated and control C26 tumors ($p=1$), the concentrations of motesanib, pazopanib and sunitinib were significantly elevated (vs. control), with the highest values detected in the sunitinib-treated animals (0.0083, 0.148, 0.2372 $\mu\text{mol}/\text{mL}$, respectively; **Figure 40**).

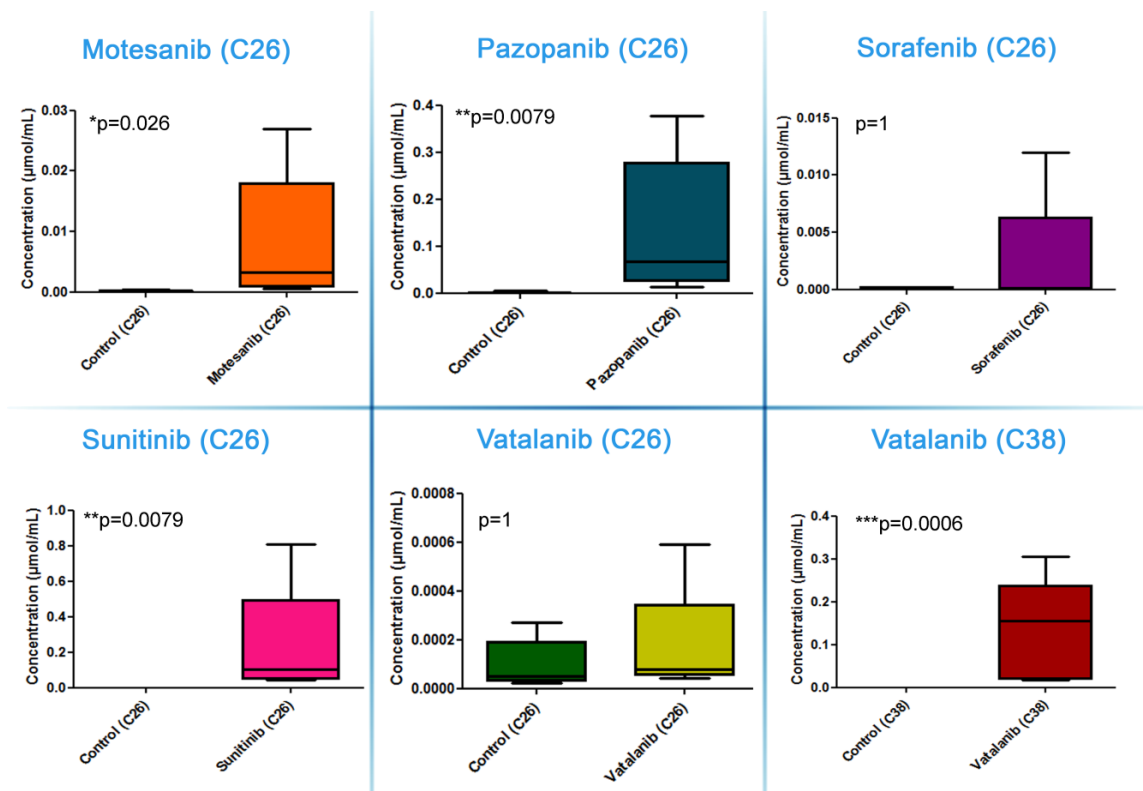


Figure 40. Tumor tissue concentrations of antiangiogenic RTKIs. Signal intensities (normalized to TIC) of the appropriate RTKIs in treated tumors and the same non-specific normalized m/z values measured in control tumors were used to calculate intratumoral drug concentrations. Data are shown as box (first and third quartiles) and whisker (maximum to minimum) plots with the mean (horizontal bar) from 6 animals per group.

Importantly, the above described drug concentrations refer to the entire tumor section and striking differences in the drug distribution were observable within the in vivo-treated C26 tumors. As for sunitinib, the drug was quite homogeneously distributed within the viable C26 tumor areas and apoptotic regions showed notably lower signal intensities (**Figures 41 and 42.**). In contrast, motesanib was seen only in one third of the C26 tumors at relatively high levels in connected areas and the intratumoral distributions of this RTKI and pazopanib (both of which were also present at relatively high average tumor tissue levels; **Figure 40.**) were inhomogeneous with the highest signal intensities observed in non-viable areas (**Figure 41.**). Only traces of sorafenib and vatalanib were detected in the C26 model. Representative images of intratumoral drug distributions are shown in **Figure 41.**

In a previously published study, we found significantly decreased C38 tumor burdens in C57Bl/6 mice treated with vatalanib (267). Accordingly, in order to determine why

mice bearing C26 tumors respond notably poorer to vatalanib than those with C38 tumors, we also utilized MALDI-MSI of C38 tumors and addressed whether there are animal model-specific variations in the tumor tissue penetration and distribution of antiangiogenic RTKIs. In contrast to the C26 model, vatalanib was well-distributed with notable signal intensities in the C38 tumors (**Figure 41.**). In line with this, in vatalanib-treated mice bearing C38 tumors, the intratumoral drug concentration was significantly higher than that in the group of untreated controls ($p=0.0006$, **Figure 40.**). It is also important to mention that we found significantly higher vatalanib concentrations in C38 than in C26 tumors ($0.142 \mu\text{mol/mL}$ vs 0.174 nmol/mL , $p=0.0025$, **Figures 40 and 41.**). No correlation between drug signal intensities in the blood and in the corresponding tumor tissue was detected.

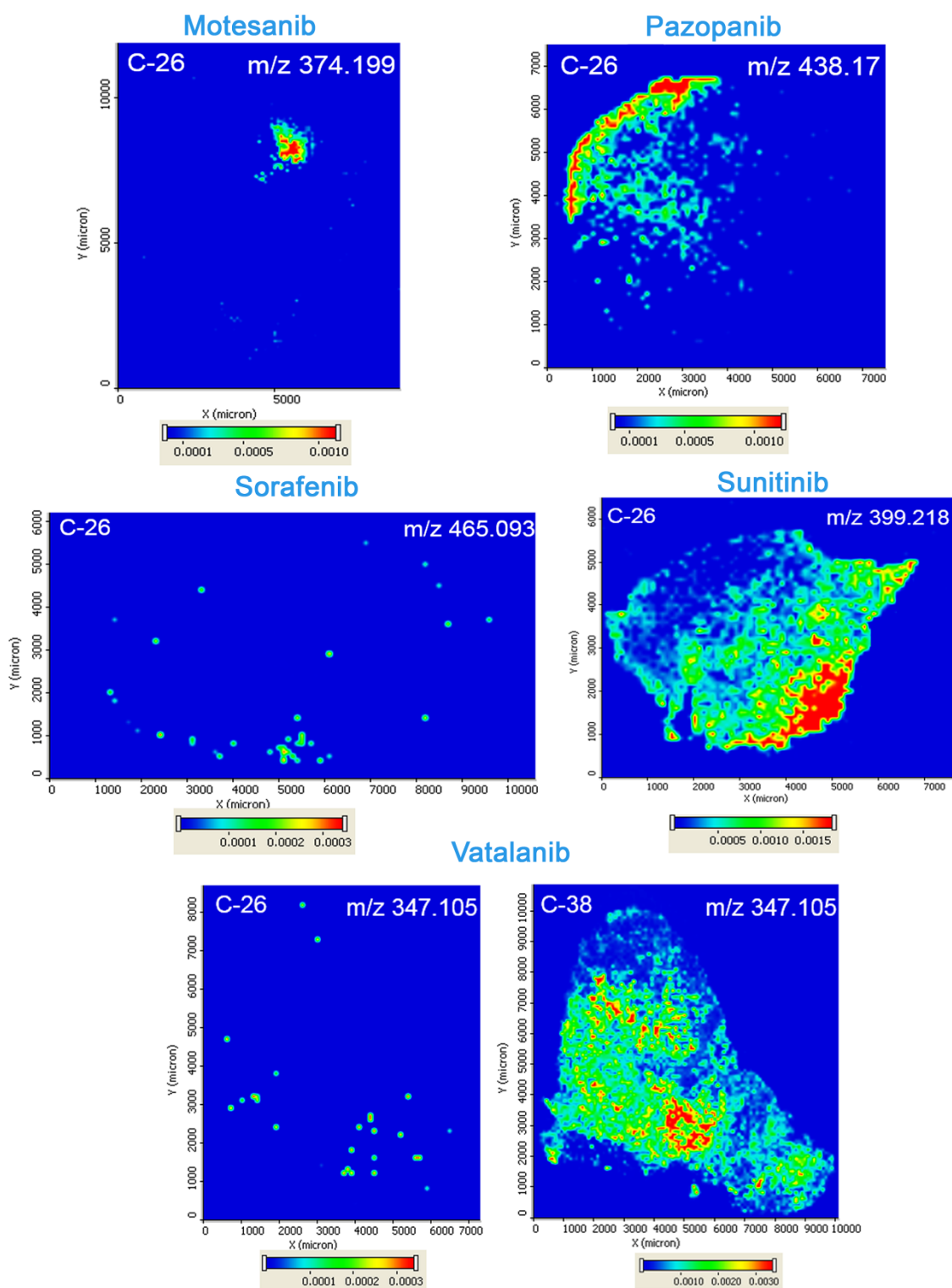


Figure 41. Representative images of drug distribution in C26 and C38 tumors after two weeks of treatment with different antiangiogenic RTKIs. Precursor ion signals of RTKIs were normalized to TIC.

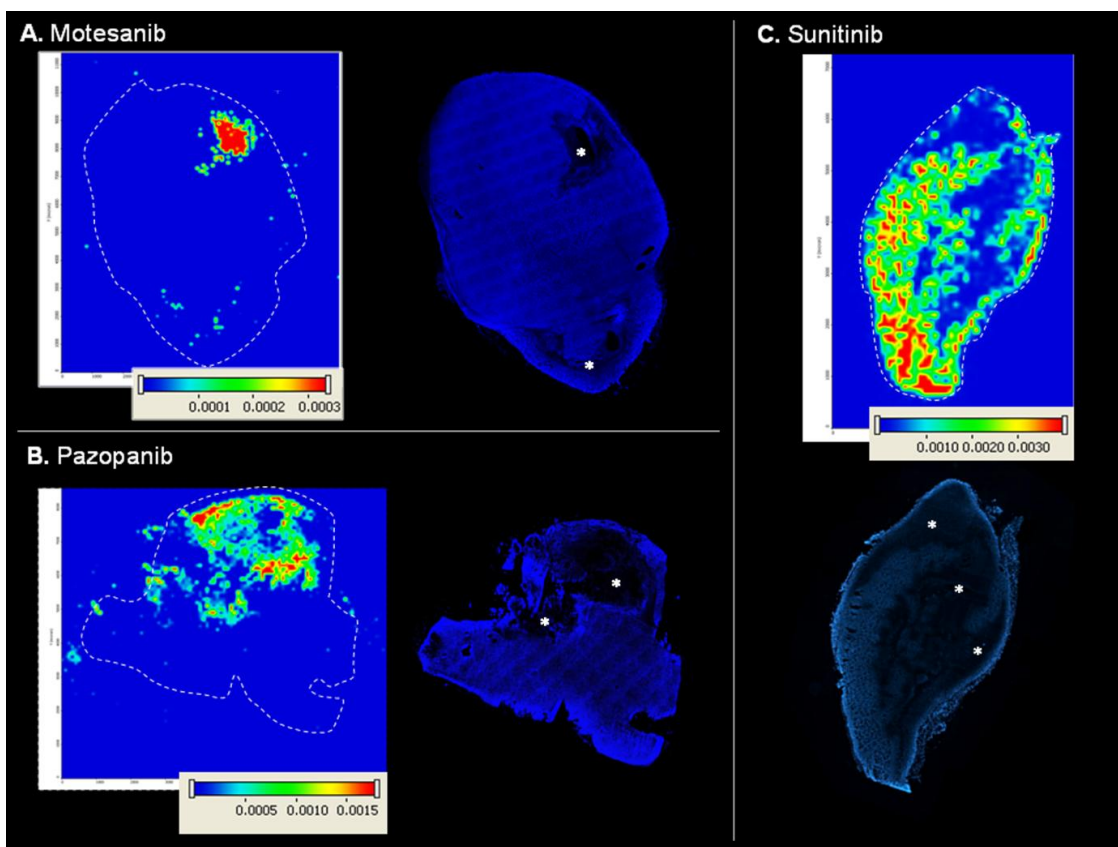


Figure 42. Co-localization of drug compounds (as visualized by MALDI-MSI) and non-viable areas in C26 tumors treated with motesanib (A.), pazopanib (B.) and sunitinib (C.). Asterisks mark non-viable intratumoral areas that appear black due to lack of nuclear counterstain (Hoechst 33342, blue). Tumor boundaries are delineated with dashed line in MALDI-MS images.

All detected RTKIs and their fragment ions showed co-localization within the tissues. This co-localization can be interpreted as a molecular fingerprint that confirms the identity of RTKIs. Representative examples showing the distribution of sunitinib and its fragment ions in tumor, liver and kidney samples are shown in **Figure 43**.

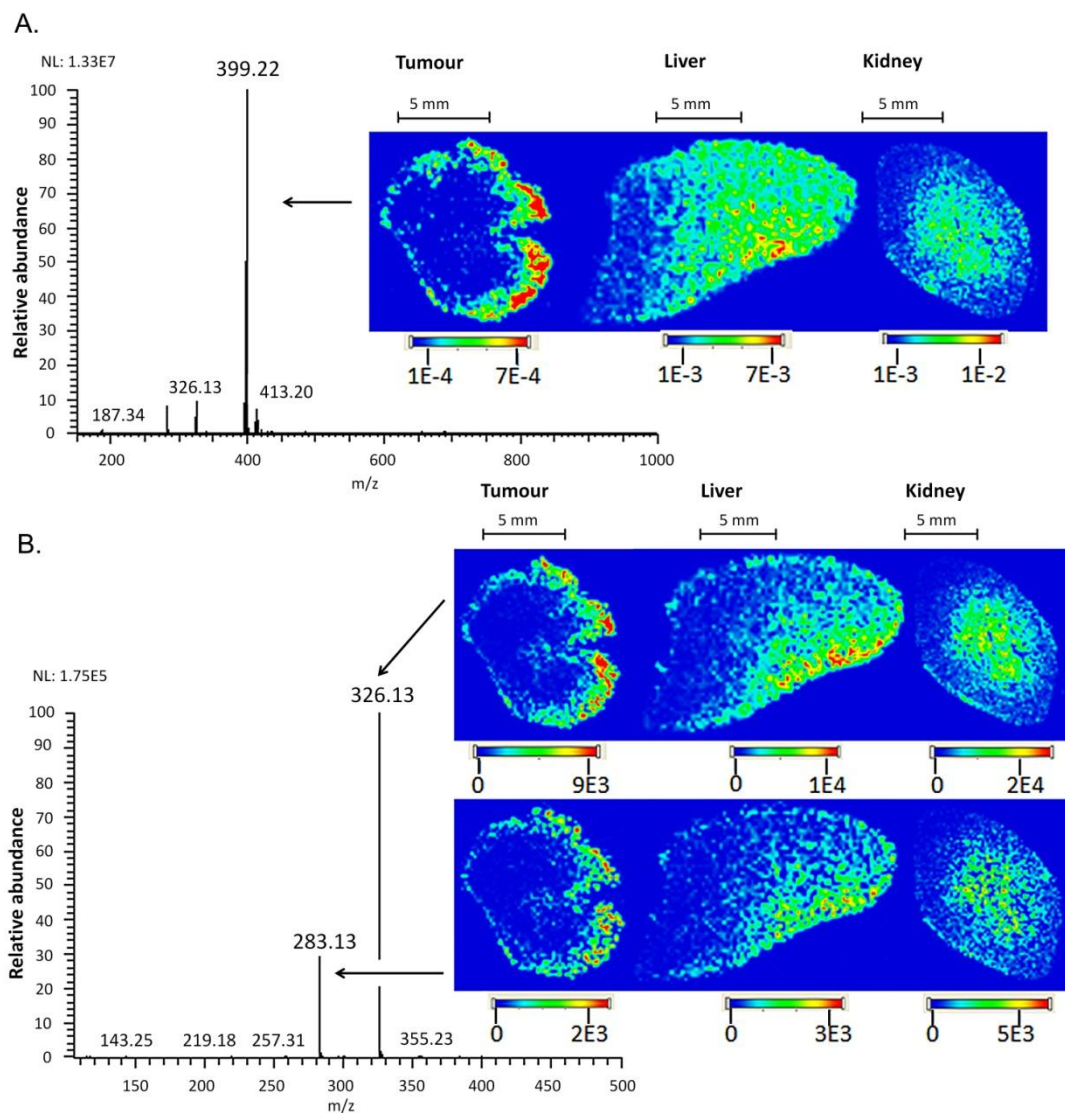


Figure 43. (A.) Full mass spectrum of sunitinib (399.218) and images of the distribution of the precursor molecule in tumor, liver and kidney tissues after 2 weeks of treatment. Signal of sunitinib is normalized to TIC. (B.) MS/MS spectrum of sunitinib and images of the distribution of the fragment ions (m/z 326.1 and 283.1) in tumor, liver and kidney tissues.

We also identified several sunitinib metabolites within the different tissues. In particular, the mono-desethylated (m/z 371.188), the desaturated (m/z 397.203), and the monohydroxylated (m/z 415.215) metabolites were observable by imaging (**Figure 44.**).

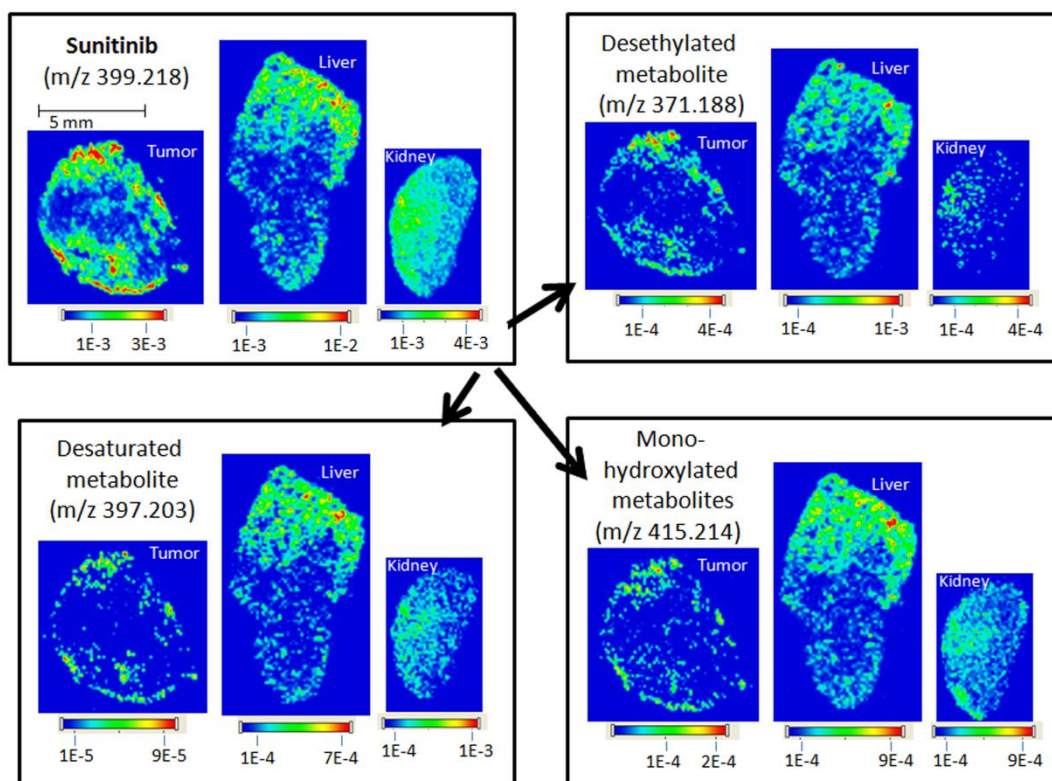


Figure 44. Distribution properties of sunitinib and its metabolites. Precursor molecule, desethylated metabolite (SU012662, M3), desaturated metabolite (M6) and mono-hydroxylated metabolites (M9, M10 and/or M11) in tumor, liver and kidney tissue sections.

Similarly, the carbinolamine (m/z 372.184) and the oxindole metabolite (m/z 388.179) of motesanib could also be shown in tissue (data not shown). Moreover, the main oxydative metabolite (m/z 363.1) of vatalanib was also identified in tissue sections, however, only in the C38 model (data not shown).

The precursor compounds and all the measured metabolites showed an overlapping tissue pattern.

7. DISCUSSION

Angiogenesis research has led to the identification of several regulators of the process, some of which represent therapeutic targets. However, results of trials with antiangiogenic agents have been both encouraging and disappointing. The most important problem in the clinical application of these drugs is assessing the tumor response that can be inadequate. Tumor shrinkages characterized by cavitation have been observed and these do not meet the usual standard radiologic criteria for response. A relevant clinical challenge is therefore to find the best techniques for monitoring the effects of antivasular drugs. Especially antiangiogenic RTKI treatment raises a lot of questions. As the main receptors being involved in the angiogenic process have high structural similarities in the kinase domain, and thus activate similar signaling cascades, a relevant attempt is to develop drugs with a broad specificity, blocking not only the VEGF pathway, but PDGF and FGF signaling as well (417). Indeed, all of the approved antiangiogenic RTKIs are multi-target inhibitors, which beside the better efficacy caused by hitting multiple targets on one hand, can lead to increased toxicities on the other hand. Moreover, by blocking mural cell recruitment, treatment can affect vessel integrity as well, emerging the metastatizing potential of the tumor. However, by destructing the tumor vasculature, it is also questionable, whether the drug can reach the place of action in an effective level. Nevertheless, although the combination of antiangiogenic agents with conventional chemotherapy is highly problematic and should be carefully designed, there had been no studies that focused directly on the exact intratumoral distribution of these agents during and after their delivery.

In our study, the combination of the tolerability of the ionization mode, the resolving power of the Orbitrap with the sensitivity of the linear ion trap made MALDI-MS an ideal technique for both drug and metabolite detection in different tissue compartments (**Figures 37 and 44.**). Besides detecting non labeled compounds, another advantage of MALDI-MSI compared to other previously used methods is that these techniques require either fluid samples (such as urine, blood or sweat) or the homogenization of the tissue (418-420). Therefore, they are not capable of analyzing the spatial tissue distribution of a compound in an organ or in a solid tumor.

The current study is the first describing the tissue distribution of unlabelled antiangiogenic RTKIs and their metabolites by MSI and provides the first direct evidence that antiangiogenic drugs given orally are transported to, taken up and metabolized within the targeted compartment, the adenocarcinoma tumor. Moreover, the presented results are the first demonstrating that MALDI-MSI is a versatile and simple method of conducting ADME studies on antiangiogenic RTKIs.

The observed overlap in the distribution pattern of the RTKIs and their fragment ions confirms the identity of the drugs (**Figure 43.**). Co-localization of the RTKIs and their metabolites (**Figure 44.**) suggests that the chemical properties responsible for drug dispersion remain similar in case of the metabolites, and accordingly, they may contribute to the tumor growth inhibitory activity of the precursor compound as well. Alternatively, the co-localization may indicate that the drug is being taken up and metabolized locally rather than being transported from other sites of metabolism, such as the liver, back to the same location as the precursor compound. Further studies are warranted to confirm or rule out these assumptions.

To the best of our knowledge, this is the first study reporting the head-to-head comparison of the intratumoral concentrations and distributions of various unlabeled antiangiogenic RTKIs by MSI. We found that oral administration of motesanib, pazopanib, sorafenib, sunitinib or vatalanib resulted in the absorption of all the five drugs with notable signal intensities being observed in the circulation. Surprisingly, only motesanib, pazopanib and sunitinib treatments resulted in significantly elevated intratumoral drug levels in the C26 model with the highest concentrations and the most homogeneous tumor tissue distributions observed in sunitinib-treated animals (**Figure 41.**). The intratumoral distributions of motesanib and pazopanib were inhomogeneous and notable signal intensities were confined to non-viable areas (**Figure 42.**). We also found, that both sorafenib and vatalanib was only traceable in the C26 tumors. In contrast, vatalanib was always detectable at homogeneously high concentrations throughout the malignant tissue in the C38 model (**Figure 41.**). Chances, therefore, are that besides their dose, schedule and direct antivasculature activity, the phenotype of the host vasculature and/or the tumor type are also likely to influence the tumor tissue levels and distribution of antiangiogenic RTKIs. The possible mechanisms linking inadequate antiangiogenic RTKI tumor concentration and endothelial- or tumor-specific

characteristics involve lysosomal degradation of RTKIs (421) and increased RTKI efflux by the tumor (422) or the endothelial (423) cells or both.

Importantly, high viable intratumoral drug concentrations were linked with tumor growth inhibition both in the C26 model (sunitinib) and C38 model (vatalanib) (**Figure 19.**). Moreover, suppressed vascular supply was also observable in these treatment groups. Although decreased MVD and/or microvessel area could also sporadically be observed in other treatment groups in the C26 model (**Figure 26-27.**). However, the low number of microvessels in case of vatalanib was accompanied by a relatively high microvessel area, while decreased microvessel area in case of sorafenib was linked with a relatively high MVD (**Figure 26.**). These opposing parameters of the vasculature could keep tumor blood flow levels, and consequently tumor burden high in these treatment groups. The suppressed MVD was linked with decreased microvessel area in case of motesanib, while both parameters were high in the pazopanib treated group (**Figure 26.**), both of which were present in the tumor in relatively high concentrations (**Figure 40.**), although in the non-viable tumor areas. A number of possible explanations of these opposing results exist, of which probably the most adequate is the difference in the efflux of the drugs by ECs and tumor cells. However, further experiments are needed to examine these parameters.

Hypoxic area ratios clearly correlated with decreased MVD (**Figures 26, 28-29.**).

We observed that VEGFR2 expression was significantly reduced only in sunitinib treated C26 tumors, while no difference in the expression pattern of the other receptors, or that of VEGFR2 in the C38 model were observed (**Figures 20-25.**). While no evidence exists that the expression profile of PDGFRs or FGFRs should change in response to receptor blockade, Domingues et al. documented, that successful therapy downregulates VEGFR2 expression (424). The lack of the decrease in VEGFR2 expression in the vatalanib treated C38 tumors may be explained by the different receptor expression profile of the two models, and probably by the fact, that vatalanib treatment induces a switch from sprouting angiogenesis to intussusception, which in contrast to sprouting, is not dependent on VEGFR signaling (267). However, it is also important to mention, that the intensity of VEGFR2 signal was not analysed.

Furthermore, one could also assume that since PDGFB is a key survival factor for the pericyte population and pericytes have a crucial role in the maintenance of vascular

stability (425), RTKIs with potent anti-PDGFR β activity may not promote normalization but, instead, might destabilize the vasculature and thus interfere with drug delivery. Our actual findings, however, do not support this assumption. In Balb/C mice bearing C26 tumors, treatment with sunitinib (the tested RTKI with the lowest IC50 value against PDGFR β , (386)) significantly decreased the pericyte coverage of tumor capillaries, as assessed by desmin expression (**Figures 32-33.**), and was also found in the highest intratumoral concentration (**Figures 40-41.**). However, how much this high concentration is the result of drug accumulation is still an open question. No difference in the desmin expression of C38 tumors was expected, as the IC50 value of vatalanib against PDGFR β is less remarkable (**Table 2.**).

In our study also no changes in the structure of the vasculature, as examined by laminin and α SMA expression was observed in any group (**Figures 30-31.**).

Although antiangiogenic drugs also have direct effects against autocrine tumor cell signaling, the main effect of antivasular agents is exerted on the tumor vasculature itself and, consequently, they influence the efficacy of their own delivery. Additionally, recent clinical data raised serious concern that bevacizumab can significantly reduce the uptake of chemotherapy by human tumors (426). Of note, this is in contrast to the "vessel normalization theory" proposed by Jain and colleagues whereby treatment with an antiangiogenic agent such as bevacizumab (427) normalizes the chaotic tumor blood vessel network thus increasing chemotherapeutic drug delivery. It is also unclear whether antiangiogenic RTKIs - which are typically used as monotherapies in the indications for which they are so far approved - after a long time of treatment, can efficiently penetrate tumor tissues. As the tumor mass grows and the given antiangiogenic RTKI exerts its antivasular effects, blood capillaries may become nonfunctional or separated by longer distances resulting in limited drug delivery to RTK expressing tumor cells located distally from functional blood capillaries. Thus, the net result of antiangiogenic RTKI treatment in solid tumors might be tightly balanced by the (potentially opposing) antivasular and direct antitumor effects of these drugs.

Our current findings demonstrate the potential of MALDI-MSI to help in optimizing the dose and schedule of RTKIs, which is of crucial importance, because there is a pressing need for biomarkers of antiangiogenic therapy in the clinics (428).

8. CONCLUSIONS

1. Our results provide the first evidence that MALDI-MSI can be used to conduct ADME studies on low molecular weight antiangiogenic drugs.
2. Limited tumor tissue drug penetration contributes to primary resistance against angiogenesis inhibitors.
3. Drug concentration detected in the viable regions of the tumor is related to the antitumor and antivasular effects of the applied compounds.
4. The effects of antiangiogenic RTKIs are dependent on the tumor model used.
5. Effective treatment in the C26 model, but not the C38 model resulted in a decreased expression of VEGFR2 and desmin, and an increase in the intratumoral hypoxia.

9. SUMMARY

Since Judah Folkman postulated that tumor growth is angiogenesis dependent, several agents have been developed to inhibit the progress of new vessel formation. Despite the considerable progress made with antiangiogenic cancer therapies, the overall survival benefit with these drugs is still frustratingly limited for most patients and the mechanisms of primary and acquired resistances are unclear.

To find the potential causes of the observed ineffectiveness, we elaborated a method for the detection of antiangiogenic RTKIs and their metabolites in different tissue compartments. We correlated the antitumor and antivascular properties, as well as target receptor expressions of five different antiangiogenic RTKIs (motesanib, pazopanib, sorafenib, sunitinib, vatalanib) with their tumor tissue drug distribution data.

In Balb/C mice bearing C26 tumors, all five RTKIs absorbed efficiently, but only sunitinib exhibited broad-spectrum antivascular and antitumor activities by significantly inhibiting both tumor growth and vascularisation, while simultaneously suppressing VEGFR2, desmin expression, and increasing intratumoral hypoxia. Importantly, the highest tumor tissue drug concentrations and the most homogeneous intratumoral drug distributions have been found in sunitinib-treated animals. Motesanib and pazopanib were also present intratumorally in relatively high concentrations in connected areas, but their expression co-localized with non-viable tissue areas. Meanwhile sorafenib and vatalanib was detected only in traces in the tumor tissue. In another animal model, where - in contrast to the C26 model - vatalanib was detectable at homogeneously high concentrations throughout the malignant tissue, the drug significantly reduced the growth and microvessel area of C38 tumors in C57Bl/6 mice.

Our results suggest, that the tumor tissue penetration and thus the antiangiogenic and antitumor potential of small molecule angiogenesis inhibitors varies among the tumor models and demonstrates the potential of MALDI-MSI to predict the efficacy of unlabelled small molecule antiangiogenic drugs in malignant tissues. Our approach is thus a major technical and preclinical advance demonstrating that primary resistance to angiogenesis inhibitors involves limited tumor tissue drug penetration. We also conclude that MALDI-MSI can significantly aid in the improvement of antivascular cancer therapies.

10. ÖSSZEFOGLALÁS

Mióta Judah Folkman felvetette, hogy a tumornövekedésben az angiogenezis fontos szerepet tölt be, számos, az új erek képződésének gátlását célzó szer került kifejlesztésre. Az antiangiogén rákterápia területén elért jelentős haladás ellenére azonban a teljes túlélés meghosszabítására tett kísérletek ezen gyógyszercsoporttal korlátozottak és az elsődleges, és szerzett rezisztencia mechanizmusai tisztázatlanok.

Ezen hatástalanság okainak felkutatására kidolgoztunk egy módszert az antiangiogén receptor tirozinkináz-inhibitorok (RTKI-k) és metabolitjaik különböző szöveti környezetben való detektálására. Összevetettük öt különböző RTKI (motesanib, pazopanib, sorafenib, sunitinib, vatalanib) tumorelles és angiogenezist gátló hatását, valamint célreceptor expresszióját a gyógyszerek intratumorális denzitásával.

C26 tumort hordozó Balb/C egereknél mind az öt RTKI sikeresen felszívódott, de egyedül a sunitinib mutatott széles körű antivaszkuláris és antitumorális hatást szignifikánsan gátolva a tumornövekedést, és csökkentve mind az erek mennyiségét, mind a VEGFR2- és a dezmin kifejeződést, valamint növelve a hipoxiát. A legmagasabb intratumorális gyógyszerkoncentráció és leghomogénebb gyógyszereloszlás a sunitinib esetén volt megfigyelhető. Ezzel szemben, a motesanib és a pazopanib szintén relatív magas koncentrációban és összefüggő területeken volt detektálható, de jelenlétük a tumor elhalt területein volt jellemző. A sorafenib és a vatalanib csak nyomokban volt megtalálható a tumorszövetben. Egy másik állatmodellben, ahol a vatalanib homogéne magas koncentrációban volt jelen a tumorszövetben, ez a gyógyszer is szignifikánsan csökkentette a tumornövekedést és az intratumorális erek területét C57Bl/6 egerekben növekvő C38 tumorokban.

Eredményeink bizonyítják, hogy a kis molekulású angiogenezis inhibitorok tumorszövetbe való eljutása és az ennek következtében kialakuló növekedésgátló és antiangiogén hatás függ az alkalmazott tumormodelltől, valamint hogy a MALDI képképző tömegspektrometria alkalmas ezen jelöletlen gyógyszercsoport hatásának előrejelzésére, így segíthet az antivaszkuláris rákterápiák hatékonyságának növelésében. Az általunk alkalmazott módszer ezért egy fontos technikai és preklinikai újítás, mely bizonyítja, hogy az antiangiogén kezeléssel szembeni primer rezisztencia egyik oka az elégtelen intratumorális penetráció.

11. REFERENCES

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12. PUBLICATIONS

Connected to the thesis:

- Connell JJ, Sugihara Y, Torok S, Dome B, Tovari J, Fehniger TE, Marko-Varga G, Vegvari A. Localization of sunitinib in in vivo animal and in vitro experimental models by MALDI mass spectrometry imaging. *ANALYTICAL AND BIOANALYTICAL CHEMISTRY* 407:(8) pp. 2245-2253. (2015)
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- Hoda MA, Rozsas A, Lang E, Klikovits T, Lohinai Z, Torok S, Berta J, Bendek M, Berger W, Hegedus B, Klepetko W, Renyi-Vamos F, Grusch M, Dome B, Laszlo V. High circulating activin A level is associated with tumor progression and predicts poor prognosis in lung adenocarcinoma. *ONCOTARGET* 7:(12) pp. 13388-13399. (2016)
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