**Name of Journal:** *World Journal of Gastrointestinal Oncology*

**Manuscript NO:** 76548

**Manuscript Type:** MINIREVIEWS

**Angiogenesis in gastrointestinal stromal tumors: From bench to bedside**

Papadakos SP *et al*. Angiogenesis in GIST: From bench to bedside

Stavros P Papadakos, Christos Tsagkaris, Marios Papadakis, Andreas S Papazoglou, Dimitrios V Moysidis, Constantinos G Zografos, Stamatios Theocharis

**Stavros P Papadakos,** First Department of Pathology, School of Medicine, National and Kapodistrian University of Athens, Athens 10679, Greece

**Christos Tsagkaris,** Faculty of Medicine, University of Crete, Heraklion 71003, Greece

**Marios Papadakis,** University Hospital Witten-Herdecke, University of Witten-Herdecke, Wuppertal 42283, Germany

**Andreas S Papazoglou, Dimitrios V Moysidis,** First Department of Cardiology, AHEPA University Hospital, Aristotle University of Thessaloniki, Thessaloniki 54636, Greece

**Constantinos G Zografos,** First Department of Surgery, Athens Medical School, National and Kapodistrian University of Athens, Laikon General Hospital, Athens 11527, Greece

**Stamatios Theocharis,** First Department of Pathology, Medical School, University of Athens, Athens 11527, Greece

**Author contributions:** Papadakos SP and Tsagkaris C contributed equally; Papadakos SP and Tsagkaris C contributed to the onceptualization and study design; Papadakos SP wrote the first draft; Moysidis DV, Papazoglou AS, Papadakis M and Zografos CG wrote the second draft; Tsagkaris C, Papadakis M and Theocharis S contributed to the critical revision; Papadakis M and Theocharis S contributed to the supervision

**Corresponding author: Christos Tsagkaris, MD, Academic Fellow,** Faculty of Medicine, University of Crete, Voutes Camp, Andrea Kalokairinou, Heraklion 71003, Greece. chriss20x@gmail.com

**Received:** March 20, 2022

**Revised:** May 15, 2022

**Accepted:** **July 18, 2022**

**Published online:**

**Abstract**

Gastrointestinal stromal tumors (GISTs) are rare neoplasms with an estimated incidence from 0.78 to 1-1.5 patients *per* 100000. They most commonly occur in the elderly during the eighth decade of life affecting predominantly the stomach, but also the small intestine, the omentum, mesentery and rectosigmoid. The available treatments for GIST are associated with a significant rate of recurrent disease and adverse events. Thorough understanding of GIST’s pathophysiology and translation of this knowledge into novel regimens or drug repurposing is essential to counter this challenge. The present review summarizes the existing evidence about the role of angiogenesis in GIST’s development and progression and discusses its clinical underpinnings.

**Key Words:** Gastrointestinal stromal tumor; Cancer; Oncology; Angiogenesis; Gastrointestinal oncology; Stromal tumors

Papadakos SP, Tsagkaris C, Papadakis M, Papazoglou AS, Moysidis DV, Zografos CG, Theocharis S. Angiogenesis in gastrointestinal stromal tumors: From bench to bedside. *World J Gastrointest Oncol* 2022; In press

**Core Tip:** Thorough understanding of gastrointestinal stromal tumors (GISTs)’s pathophysiology and translation of this knowledge into novel regimens or drug repurposing is essential to counter this challenge. The present review summarizes the existing evidence about the role of angiogenesis in GIST’s development and progression and discusses its clinical underpinnings.

**INTRODUCTION**

Gastrointestinal stromal tumors (GISTs) are rare neoplasms with an estimated incidence from 0.78 patients to 2 patients *per* 100000[1,2]. Their highest prevalence is noted during the eight decade of age, when they affect up to 3.06 individuals *per* 100000[3]. GIST typically present as subepithelial masses mainly in stomach (60%) and small intestine (20%-30%) with omentum, mesentery and rectosigmoid areas being less-frequently involved areas[4]. According to their primary location, GISTs could clinically present as gastrointestinal hemorrhage, anemia, dyspepsia or vomiting when the upper gastrointestinal tract is involved and as bowel obstruction, frequent urination or diarrhea in implication of the lower gastrointestinal tract[5]. The metastatic disease principally concerns the liver, omentum, and peritoneum presenting as abdominal pain or constipation while extra-intestinal metastases to lymph nodes (LN) and lungs are infrequent[4]. The pathological diagnosis relies on the tissue’s morphological and molecular characteristics. Based on their morphology, GISTs are classified into three groups according to the predominant cell type: Spindle cell type (70%), epithelioid cell type (20%) and a mixed type (10%). CD117 comprises a transmembrane protein which is the end-product of the *c-kit* expression[6]. The KIT (CD117) positivity in immunohistochemistry (IHC) in tissues which are morphologically consistent with GIST establishes the diagnosis in the 95% of the cases. In KIT negative cases, the discovered on GIST 1 (DOG1) and CD34, which is an antigen of the myeloid progenitor cells, staining or the documentation of KIT or *platelet-derived growth factor receptor* (*PDGFRA)* genemutations are sufficient to institute a diagnosis. Seldom in pediatric and young populations, GIST formation arises in the context of succinate dehydrogenase-deficiency in conjunction with paragangliomas and pulmonary chondromas[7,8].

The pharmacologic targeting of angiogenesis in cancer therapeutics was introduced as a groundbreaking approach. Nevertheless, the anti-vascular endothelial growth factor (VEGF) targeting alone or in conjunction with chemotherapy displayed only modest benefit in overall survival in solid tumors indicating the complexity of the mechanisms that regulate tumor angiogenesis[9]. Τhus, the necessity arose to develop a broad spectrum of anti-angiogenic treatments such as: Direct VEGFR2 antagonists (ramucirumab), VEGF-Traps (aflibercept), several receptor tyrosine kinases inhibitors targeting the PDGF-R, CD117 (c-KIT), fibroblast growth factor receptors (FGFR), epidermal growth factor receptor, RET, RAF kinases and the repurposing of drugs like the mammalian target of rapamycin inhibitors and lenalidomide[9,10]. In fact, anti-angiogenetic therapy has gained ground in the management of advanced, unresectable disease. Imatinib, an abl, c-KIT and PDGF-R tyrosine kinase inhibitor (TKI), constitutes the empiric treatment when the mutational status of the disease remains unknown and the first line of treatment in KIT and PDGFRA positive metastatic, inoperable GISTs. The D842V mutation in *PDGFRA* comprises a therapeutic exception and is being treated with avapritinib while *KIT* and *PDGFRA* wild type tumors are treated with sunitinib or regorafinib[11].

All the above mentioned drugs achieve, at least partially, their cytotoxicity disrupting signaling pathways which are implicated in angiogenesis, as it would be further analyzed below. This suggests that angiogenesis might be of paramount importance for the carcinogenesis process in GISTs and an attempt to summarize all the pre-clinical and clinical data would be of great value.

**The role of angiogenesis in GIST’s development and progression**

***The molecular mechanisms of angiogenesis in GISTs–preclinical data***

The regulation of angiogenesis is necessary for cancer cells initially to cope with their increased metabolic needs and in the process to promote their metastatic potential. Its significance was firstly recognized by Folkman[12], which stated that the magnified rate of neovascularization compared with wound healing and inflammation as a result of an interplay between tumor cells and endothelial cells was a prerequisite in order to achieve tumor growth[12]. Presently, it is widely known that the angiogenic process is being coordinated by the balance of several angiogenesis inducers and inhibitors in tumor’s microenvironment. The dominance of the pro-angiogenetic factors, a phenomenon called “angiogenic switch”[13], triggers the angiogenesis and could result either as result of the consequent hypoxia from the increased tumor proliferation or by the immune cell infiltration[14]. The primary induction phase with the undeveloped vessels paves the way for the remodeling phase when the blood vessel generation is sustained[15]. Several models of angiogenesis have been described explaining partially the poor outcomes of the selective angiogenic blockage as certain tumors can utilize alternative modes of angiogenesis[14]. Their analytical presentation has been done elsewhere[14,16,17] and goes beyond the scope of this review but a brief presentation in Table 1 would be helpful.

Xenograft studies in mice constitute an invaluable source of evidence about the angiogenetic mechanisms in GISTs. Our fundamental conceptualization about the orchestration of the angiogenetic process descended from Giner *et al*[18]. They utilized an intensely CD117, DOG1 and CD34-positive GIST with continual Ki-67 expression in about 15% of the tumor’s mass. The neovascularization experiments demonstrated the propagation of the induction phase during the first 96 h after implantation which proceeded by the remodeling phase. The induction phase was guided by the *VEGF*, *VEGFC*, *PDGFA*, *PDGFB* gene expression in conformity with their receptors. In more detail, the IHC data indicate that the VEGF ligand and the VEGFR2, VEGFR3 were positive at day 4 after the xenografting. As regards the chemokine expression, CXCL9, CXCL10, GRO and their receptors CXCR3, CXCR2 were stained in tumor cells and stroma soon after the implantation with a slight staining predominance of the chemokine receptors. These effects are possibly orchestrated by hypoxia-inducible factor (HIF)1α and the CXCL12/CXCR4 axis, which are constantly expressed[18].

The angiogenetic process in GIST has been further delineated and several regulatory molecules have been identified. CCL2 represents a chemokine expressed by the tumor cells to attract CCR2-expressing endothelial progenitor cells from the circulation as documented in HER-2/neu-driven breast cancer[19]. On the other hand, the VEGF-induced nuclear factor kappa B (NF-kB) upregulation is frequently utilized to attract inflammatory cell into tumor to stimulate the angiogenesis[20]. The bromodomain and extraterminal domain family mediates immunity regulating several signaling pathways[21]. In GISTs, the BRD4 upregulation enhanced the migratory and invasion processes regulating angiogenesis through the NF-kB/CCL2 signaling pathway. The BRD4-expressing cells attract tumor-associated macrophages *via* the expression of CCL2 potentiating the tumor’s microvessel density and secrete various pro-angiogenic molecules such as VEGFA, LOX and MMP9[22,23]. Towards the same direction, mutations of the protein phosphatase 2, regulatory subunit A, alpha (PPP2R1A) affect the carcinogenesis process[24,25]. In GISTs, mutations in *PPP2R1A* gene are found in nearly 20% of the cases and correlate with a more aggressive tumor phenotype. They result in increased growth rate *via* enhancing phosphorylation of c-kit, Akt1/2, ERK1/2 and WNK1. The latter seems to mediate the regulation of the angiogenetic process[26,27]. A further analysis of the specific mechanisms would be of great value and it should be applied.

Furthermore, while the contribution of epigenetic mechanisms in the GIST progression is well established, its impact in the angiogenetic mechanisms could be further delineated. Several gaps in our understanding that remain unaddressed by the subdivisions according to the driver gene mutation status could be further elucidated by the tumor’s epigenetic landscape. The alterations in the tumor’s methylation profile are associated with a more aggressive phenotype[28] and the methylation status of the CD133 could reshape the management of the disease and it would be presented below in more depth[29]. The KDM4 family members (KDM4A-D) reshaping the structure of chromatin are implicated in the pathogenesis of a wide variety of cancers[30]. In GIST, the upregulation of KDM4D potentiates the angiogenesis in vivo, as indicated by the overexpression of CD31 in IHC. These effects are mediated by the HIF1β/VEGFA pathway in the presence of demethylation in the promoters of the *H3K9me3* and *H3K36me3* genes[31].

Finally, it is worth mentioning that several multi-TKIs exert their anti-tumor efficacy at least partially by the inhibition of angiogenesis. Cabozantinib exerts it’s activity inhibiting the receptor tyrosine kinases MET, VEGFR2, Flt-3, c-Kit and RET[32,33] while sorafenib inhibits the signaling of VEGFR-1, VEGFR-2, VEGFR-3, PDGFR-β, Flt-3, c-Kit and the Raf serine/threonine kinases[34]. Both diminish the tumor’s microvascular density as evidenced by CD31 IHC[35,36].

***The association between imaging data and angiogenesis in GISTs***

There have been several classification systems to stratify the malignant potential of GISTs such as: The National Institutes of Healthv consensus criteria (Fletcher's criteria), the Armed Forces Institute of Pathology criteria (Miettinen's criteria) or the International Union against Cancer TNM classification. Their main drawback constitute the inability to validate the tumor’s aggressiveness without surgical resection and detailed pathologic examination of the entire tumor to estimate the mitotic count[37,38] . Although taking into consideration the current therapeutic trends, the management of the advanced, unresectable disease is unequivocal, there are margins for improvement in the management of primary localized disease, especially in small-sized tumors. It could not be emphasized enough that even small GISTs could develop malignant behavior. Τhereat, it could provide us a wealth of valuable predictive and prognostic information an attempt to incorporate imaging data about the vascularization of the tumor such as the vessels’ irregularity or the blood perfusion[39].

The above mentioned gap was attempted to be filled by a landmark study by Iannicelli *et al*[40], the computed tomography (CT) constitutes the fundamental imaging modality in patients presenting with the clinical manifestations of GIST. Reviewing past literature, several studies have documented that aim to associate certain imaging features with pathologic parameters[41,42]. Iannicelli *et al*[40] presented that GISTs with irregular margins tended to have superior mitotic rate than tumor with regular margins. Furthermore, a heterogonous pattern of contrast enhancement (CE), the angiogenesis and necrosis correlated with an increased tumor size and a more aggressive clinical behavior. It worth mentioning that the intensity of CE although it represents a novel mark of biologic activity, was not correlated with neither the number of mitoses nor the tumor’s risk stratification[40]. The above comprise an indirect link between tumor’s margins and mitotic rate, which is essential in order to stratify before surgery the clinical behavior of the tumor and highlight the importance of angiogenesis in disease progression. The latter could also be deduced by dynamic positron emission tomography analysis. Strauss *et al*[43] reported an association between the rate in which the F-18-fluorodeoxyglucose diffused into the tumor with the expression of VEGF-A[43]. The main limitation of CT comprises it’s low sensitivity as regards the imaging of vascularity in small sized tumors[39]. This divergence could be addressed by the endoscopic ultrasound (EUS) technology.

The utilization of EUS has emerged during the last decades. Its ability to evade the intervention of the abdominal fat and gastrointestinal gas in conjunction with the capability of FNA biopsy render it a useful tool towards a more personalized approach in the management of GIST. In EUS the GISTs are visualized as hypoechoic masses arising from the muscularis propria or the muscularis mucosae. The presence of irregular margins, cystic areas or malignant LN herald bad prognosis[44]. The usage of contrast media enhances further the diagnostic capacity of the EUS and promotes the tumor’s vascularity as a valuable prognostic biomarker. The role of CE-EUS in the management has been extensively reviewed elsewhere[45] and we intend to delineate the fundamentals. Sakamoto *et al*[39] classified the tumor’s vascularity into two subgroups according to the pattern of perfusion (homogenous or heterogeneous) and vessel appearance (regular or irregular). The homogenous perfusion with regular vessels were considered as signs of mild clinical behavior. Furthermore, they compared the diagnostic sensitivity of contrast-enhanced harmonic US, Power-Doppler EUS and CE-multidetector CT to visualize tumor vessels. In GISTs larger than 3 cm their sensitivities were 100%, 75% and 42% respectively. The differences became more emphatic in tumors less than 3 cm: 100%, 25% and 0%, respectively. It was noteworthy that every malignant lesion less than 3 cm in the cohort had been detected by the CEH-EUS before surgery[39]. The above indicate that CE-US comprises a powerful tool to visualize vascularity. Taking a step further, Yamashita *et al*[46] demonstrated an association between the imaging findings on CE-US and the pathologic risk stratification. In more depth, the large vessels lacked elastic tissue, indicating that neovascularization constitutes the underlying pathogenetic mechanism, and expressed VEGF[46].

It becomes evident that the imaging findings of vascularity might be sensational and practice changing in a subset of patients with small sized tumors (< 3 cm) and aggressive phenotype. A more substantial body of evidence should be collected in order to address properly those dilemmas.

***Angiogenesis mediators as biomarkers in GIST–clinical data***

The development of biomarkers comprises an essential step towards the individualization of medical practice. Liquid biopsy provides a cutting-edge, non-invasive technology to access predictive information to guide the therapeutic management in a wide variety of diseases[47-51]. It’s application in GIST treatment has been started to emerge[52,53]. Reviewing subsequent and more recent literature, an extensive number of studies has been found associating molecules implicated in angiogenesis with pathologic features. Although there are several limitations in the above mentioned research, the importance of angiogenesis in GIST’s malignant progression is delineated. In Table 2 are summarized the most significant data.

**CONCLUSION**

As highlighted above, angiogenesis mediates an extensive proportion of GIST’s malignant dynamics. Several signaling pathways are implicated in the regulation of angiogenesis such as: The VEGF, the fibroblast growth factor-2 (FGF2), the PDGF, the angiopoietins, the Eph/ephrin signaling, the Apelin/APLNR pathway, the HIFs and several chemokines[14]. The VEGF signaling comprises the most well-studied pathway in GIST angiogenesis.

The FGF2/R2 signaling has been extensively studied in GIST as a drug resistance mechanism. Sergei *et al*[54] and Boichuk *et al*[55] demonstrated that the blockage of FGFR2 signaling could enhance the responsiveness to DNA-Topoisomerase II inhibitors[54] while the downregulation of FGF2 signaling might stimulate the response to imatinib[55]. It’s contribution in GIST progression has been reviewed[56] but data about potential effects in GIST vascularization process are missing. Towards the same direction, the Eph/ephrin system has been investigated in carcinogenesis[57,58]. It would be of paramount importance an attempt to outline its contribution in GIST angiogenesis.

**REFERENCES**

1 **van der Graaf WTA**, Tielen R, Bonenkamp JJ, Lemmens V, Verhoeven RHA, de Wilt JHW. Nationwide trends in the incidence and outcome of patients with gastrointestinal stromal tumour in the imatinib era. *Br J Surg* 2018; **105**: 1020-1027 [PMID: 29664995 DOI: 10.1002/bjs.10809]

2 **Casali PG**, Blay JY, Abecassis N, Bajpai J, Bauer S, Biagini R, Bielack S, Bonvalot S, Boukovinas I, Bovee JVMG, Boye K, Brodowicz T, Buonadonna A, De Álava E, Dei Tos AP, Del Muro XG, Dufresne A, Eriksson M, Fedenko A, Ferraresi V, Ferrari A, Frezza AM, Gasperoni S, Gelderblom H, Gouin F, Grignani G, Haas R, Hassan AB, Hindi N, Hohenberger P, Joensuu H, Jones RL, Jungels C, Jutte P, Kasper B, Kawai A, Kopeckova K, Krákorová DA, Le Cesne A, Le Grange F, Legius E, Leithner A, Lopez-Pousa A, Martin-Broto J, Merimsky O, Messiou C, Miah AB, Mir O, Montemurro M, Morosi C, Palmerini E, Pantaleo MA, Piana R, Piperno-Neumann S, Reichardt P, Rutkowski P, Safwat AA, Sangalli C, Sbaraglia M, Scheipl S, Schöffski P, Sleijfer S, Strauss D, Strauss SJ, Hall KS, Trama A, Unk M, van de Sande MAJ, van der Graaf WTA, van Houdt WJ, Frebourg T, Gronchi A, Stacchiotti S; ESMO Guidelines Committee, EURACAN and GENTURIS. Electronic address: clinicalguidelines@esmo.org. Gastrointestinal stromal tumours: ESMO-EURACAN-GENTURIS Clinical Practice Guidelines for diagnosis, treatment and follow-up. *Ann Oncol* 2022; **33**: 20-33 [PMID: 34560242 DOI: 10.1016/j.annonc.2021.09.005]

3 **Ma GL**, Murphy JD, Martinez ME, Sicklick JK. Epidemiology of gastrointestinal stromal tumors in the era of histology codes: results of a population-based study. *Cancer Epidemiol Biomarkers Prev* 2015; **24**: 298-302 [PMID: 25277795 DOI: 10.1158/1055-9965.EPI-14-1002]

4 **Nishida T**, Blay JY, Hirota S, Kitagawa Y, Kang YK. The standard diagnosis, treatment, and follow-up of gastrointestinal stromal tumors based on guidelines. *Gastric Cancer* 2016; **19**: 3-14 [PMID: 26276366 DOI: 10.1007/s10120-015-0526-8]

5 **Caterino S**, Lorenzon L, Petrucciani N, Iannicelli E, Pilozzi E, Romiti A, Cavallini M, Ziparo V. Gastrointestinal stromal tumors: correlation between symptoms at presentation, tumor location and prognostic factors in 47 consecutive patients. *World J Surg Oncol* 2011; **9**: 13 [PMID: 21284869 DOI: 10.1186/1477-7819-9-13]

6 **Sarlomo-Rikala M**, Kovatich AJ, Barusevicius A, Miettinen M. CD117: a sensitive marker for gastrointestinal stromal tumors that is more specific than CD34. *Mod Pathol* 1998; **11**: 728-734 [PMID: 9720500]

7 **Joensuu H**, Hohenberger P, Corless CL. Gastrointestinal stromal tumour. *Lancet* 2013; **382**: 973-983 [PMID: 23623056 DOI: 10.1016/S0140-6736(13)60106-3]

8 **Miettinen M**, Wang ZF, Sarlomo-Rikala M, Osuch C, Rutkowski P, Lasota J. Succinate dehydrogenase-deficient GISTs: a clinicopathologic, immunohistochemical, and molecular genetic study of 66 gastric GISTs with predilection to young age. *Am J Surg Pathol* 2011; **35**: 1712-1721 [PMID: 21997692 DOI: 10.1097/PAS.0b013e3182260752]

9 **Jain RK**. Antiangiogenesis strategies revisited: from starving tumors to alleviating hypoxia. *Cancer Cell* 2014; **26**: 605-622 [PMID: 25517747 DOI: 10.1016/j.ccell.2014.10.006]

10 **Teng LS**, Jin KT, He KF, Zhang J, Wang HH, Cao J. Clinical applications of VEGF-trap (aflibercept) in cancer treatment. *J Chin Med Assoc* 2010; **73**: 449-456 [PMID: 20875616 DOI: 10.1016/S1726-4901(10)70097-6]

11 **Klug LR**, Khosroyani HM, Kent JD, Heinrich MC. New treatment strategies for advanced-stage gastrointestinal stromal tumours. *Nat Rev Clin Oncol* 2022; **19**: 328-341 [PMID: 35217782 DOI: 10.1038/s41571-022-00606-4]

12 **Folkman J**. Tumor angiogenesis: therapeutic implications. *N Engl J Med* 1971; **285**: 1182-1186 [PMID: 4938153 DOI: 10.1056/NEJM197111182852108]

13 **Hanahan D**, Folkman J. Patterns and emerging mechanisms of the angiogenic switch during tumorigenesis. *Cell* 1996; **86**: 353-364 [PMID: 8756718 DOI: 10.1016/s0092-8674(00)80108-7]

14 **Lugano R**, Ramachandran M, Dimberg A. Tumor angiogenesis: causes, consequences, challenges and opportunities. *Cell Mol Life Sci* 2020; **77**: 1745-1770 [PMID: 31690961 DOI: 10.1007/s00018-019-03351-7]

15 **Llombart-Bosch A**, López-Guerrero JA, Carda Batalla C, Ruíz Suarí A, Peydró-Olaya A. Structural basis of tumoral angiogenesis. *Adv Exp Med Biol* 2003; **532**: 69-89 [PMID: 12908551 DOI: 10.1007/978-1-4615-0081-0\_8]

16 **Carmeliet P**, Jain RK. Molecular mechanisms and clinical applications of angiogenesis. *Nature* 2011; **473**: 298-307 [PMID: 21593862 DOI: 10.1038/nature10144]

17 **Kuczynski EA**, Vermeulen PB, Pezzella F, Kerbel RS, Reynolds AR. Vessel co-option in cancer. *Nat Rev Clin Oncol* 2019; **16**: 469-493 [PMID: 30816337 DOI: 10.1038/s41571-019-0181-9]

18 **Giner F**, Machado I, Lopez-Guerrero JA, Mayordomo-Aranda E, Llombart-Bosch A. High-risk gastrointestinal stromal tumour (GIST) and synovial sarcoma display similar angiogenic profiles: a nude mice xenograft study. *Ecancermedicalscience* 2017; **11**: 726 [PMID: 28386296 DOI: 10.3332/ecancer.2017.726]

19 **Chen X**, Wang Y, Nelson D, Tian S, Mulvey E, Patel B, Conti I, Jaen J, Rollins BJ. CCL2/CCR2 Regulates the Tumor Microenvironment in HER-2/neu-Driven Mammary Carcinomas in Mice. *PLoS One* 2016; **11**: e0165595 [PMID: 27820834 DOI: 10.1371/journal.pone.0165595]

20 **Jiang BH**, Liu LZ. PI3K/PTEN signaling in angiogenesis and tumorigenesis. *Adv Cancer Res* 2009; **102**: 19-65 [PMID: 19595306 DOI: 10.1016/S0065-230X(09)02002-8]

21 **Wang N**, Wu R, Tang D, Kang R. The BET family in immunity and disease. *Signal Transduct Target Ther* 2021; **6**: 23 [PMID: 33462181 DOI: 10.1038/s41392-020-00384-4]

22 **Mu J**, Sun P, Ma Z, Sun P. BRD4 promotes tumor progression and NF-κB/CCL2-dependent tumor-associated macrophage recruitment in GIST. *Cell Death Dis* 2019; **10**: 935 [PMID: 31819043 DOI: 10.1038/s41419-019-2170-4]

23 **Liu N**, Ling R, Tang X, Yu Y, Zhou Y, Chen D. Post-Translational Modifications of BRD4: Therapeutic Targets for Tumor. *Front Oncol* 2022; **12**: 847701 [PMID: 35402244 DOI: 10.3389/fonc.2022.847701]

24 **Calin GA**, di Iasio MG, Caprini E, Vorechovsky I, Natali PG, Sozzi G, Croce CM, Barbanti-Brodano G, Russo G, Negrini M. Low frequency of alterations of the alpha (PPP2R1A) and beta (PPP2R1B) isoforms of the subunit A of the serine-threonine phosphatase 2A in human neoplasms. *Oncogene* 2000; **19**: 1191-1195 [PMID: 10713707 DOI: 10.1038/sj.onc.1203389]

25 **Janssens V**, Goris J. Protein phosphatase 2A: a highly regulated family of serine/threonine phosphatases implicated in cell growth and signalling. *Biochem J* 2001; **353**: 417-439 [PMID: 11171037 DOI: 10.1042/0264-6021:3530417]

26 **Xie J**, Yoon J, Yang SS, Lin SH, Huang CL. WNK1 protein kinase regulates embryonic cardiovascular development through the OSR1 signaling cascade. *J Biol Chem* 2013; **288**: 8566-8574 [PMID: 23386621 DOI: 10.1074/jbc.M113.451575]

27 **Lai JG**, Tsai SM, Tu HC, Chen WC, Kou FJ, Lu JW, Wang HD, Huang CL, Yuh CH. Zebrafish WNK lysine deficient protein kinase 1 (wnk1) affects angiogenesis associated with VEGF signaling. *PLoS One* 2014; **9**: e106129 [PMID: 25171174 DOI: 10.1371/journal.pone.0106129]

28 **Okamoto Y**, Sawaki A, Ito S, Nishida T, Takahashi T, Toyota M, Suzuki H, Shinomura Y, Takeuchi I, Shinjo K, An B, Ito H, Yamao K, Fujii M, Murakami H, Osada H, Kataoka H, Joh T, Sekido Y, Kondo Y. Aberrant DNA methylation associated with aggressiveness of gastrointestinal stromal tumour. *Gut* 2012; **61**: 392-401 [PMID: 21708825 DOI: 10.1136/gut.2011.241034]

29 **Geddert H**, Braun A, Kayser C, Dimmler A, Faller G, Agaimy A, Haller F, Moskalev EA. Epigenetic Regulation of CD133 in Gastrointestinal Stromal Tumors. *Am J Clin Pathol* 2017; **147**: 515-524 [PMID: 28398518 DOI: 10.1093/ajcp/aqx028]

30 **Lee DH**, Kim GW, Jeon YH, Yoo J, Lee SW, Kwon SH. Advances in histone demethylase KDM4 as cancer therapeutic targets. *FASEB J* 2020; **34**: 3461-3484 [PMID: 31961018 DOI: 10.1096/fj.201902584R]

31 **Hu F**, Li H, Liu L, Xu F, Lai S, Luo X, Hu J, Yang X. Histone demethylase KDM4D promotes gastrointestinal stromal tumor progression through HIF1β/VEGFA signalling. *Mol Cancer* 2018; **17**: 107 [PMID: 30060750 DOI: 10.1186/s12943-018-0861-6]

32 **Yakes FM**, Chen J, Tan J, Yamaguchi K, Shi Y, Yu P, Qian F, Chu F, Bentzien F, Cancilla B, Orf J, You A, Laird AD, Engst S, Lee L, Lesch J, Chou YC, Joly AH. Cabozantinib (XL184), a novel MET and VEGFR2 inhibitor, simultaneously suppresses metastasis, angiogenesis, and tumor growth. *Mol Cancer Ther* 2011; **10**: 2298-2308 [PMID: 21926191 DOI: 10.1158/1535-7163.MCT-11-0264]

33 **Grüllich C**. Cabozantinib: a MET, RET, and VEGFR2 tyrosine kinase inhibitor. *Recent Results Cancer Res* 2014; **201**: 207-214 [PMID: 24756794 DOI: 10.1007/978-3-642-54490-3\_12]

34 **Wilhelm SM**, Adnane L, Newell P, Villanueva A, Llovet JM, Lynch M. Preclinical overview of sorafenib, a multikinase inhibitor that targets both Raf and VEGF and PDGF receptor tyrosine kinase signaling. *Mol Cancer Ther* 2008; **7**: 3129-3140 [PMID: 18852116 DOI: 10.1158/1535-7163.MCT-08-0013]

35 **Gebreyohannes YK**, Schöffski P, Van Looy T, Wellens J, Vreys L, Cornillie J, Vanleeuw U, Aftab DT, Debiec-Rychter M, Sciot R, Wozniak A. Cabozantinib Is Active against Human Gastrointestinal Stromal Tumor Xenografts Carrying Different KIT Mutations. *Mol Cancer Ther* 2016; **15**: 2845-2852 [PMID: 27777285 DOI: 10.1158/1535-7163.MCT-16-0224]

36 **Huynh H**, Lee JW, Chow PK, Ngo VC, Lew GB, Lam IW, Ong HS, Chung A, Soo KC. Sorafenib induces growth suppression in mouse models of gastrointestinal stromal tumor. *Mol Cancer Ther* 2009; **8**: 152-159 [PMID: 19139124 DOI: 10.1158/1535-7163.MCT-08-0553]

37 **Agaimy A**. Gastrointestinal stromal tumors (GIST) from risk stratification systems to the new TNM proposal: more questions than answers? A review emphasizing the need for a standardized GIST reporting. *Int J Clin Exp Pathol* 2010; **3**: 461-471 [PMID: 20606727]

38 **Takahashi T**, Nakajima K, Nishitani A, Souma Y, Hirota S, Sawa Y, Nishida T. An enhanced risk-group stratification system for more practical prognostication of clinically malignant gastrointestinal stromal tumors. *Int J Clin Oncol* 2007; **12**: 369-374 [PMID: 17929119 DOI: 10.1007/s10147-007-0705-7]

39 **Sakamoto H**, Kitano M, Matsui S, Kamata K, Komaki T, Imai H, Dote K, Kudo M. Estimation of malignant potential of GI stromal tumors by contrast-enhanced harmonic EUS (with videos). *Gastrointest Endosc* 2011; **73**: 227-237 [PMID: 21295636 DOI: 10.1016/j.gie.2010.10.011]

40 **Iannicelli E**, Carbonetti F, Federici GF, Martini I, Caterino S, Pilozzi E, Panzuto F, Briani C, David V. Evaluation of the Relationships Between Computed Tomography Features, Pathological Findings, and Prognostic Risk Assessment in Gastrointestinal Stromal Tumors. *J Comput Assist Tomogr* 2017; **41**: 271-278 [PMID: 27753723 DOI: 10.1097/RCT.0000000000000499]

41 **Horton KM**, Juluru K, Montogomery E, Fishman EK. Computed tomography imaging of gastrointestinal stromal tumors with pathology correlation. *J Comput Assist Tomogr* 2004; **28**: 811-817 [PMID: 15538156 DOI: 10.1097/00004728-200411000-00014]

42 **Baheti AD**, Shinagare AB, O'Neill AC, Krajewski KM, Hornick JL, George S, Ramaiya NH, Tirumani SH. MDCT and clinicopathological features of small bowel gastrointestinal stromal tumours in 102 patients: a single institute experience. *Br J Radiol* 2015; **88**: 20150085 [PMID: 26111069 DOI: 10.1259/bjr.20150085]

43 **Strauss LG**, Dimitrakopoulou-Strauss A, Koczan D, Pan L, Hohenberger P. Correlation of dynamic PET and gene array data in patients with gastrointestinal stromal tumors. *ScientificWorldJournal* 2012; **2012**: 721313 [PMID: 22701369 DOI: 10.1100/2012/721313]

44 **Palazzo L**, Landi B, Cellier C, Cuillerier E, Roseau G, Barbier JP. Endosonographic features predictive of benign and malignant gastrointestinal stromal cell tumours. *Gut* 2000; **46**: 88-92 [PMID: 10601061 DOI: 10.1136/gut.46.1.88]

45 **Chhoda A**, Jain D, Surabhi VR, Singhal S. Contrast Enhanced Harmonic Endoscopic Ultrasound: A Novel Approach for Diagnosis and Management of Gastrointestinal Stromal Tumors. *Clin Endosc* 2018; **51**: 215-221 [PMID: 29874903 DOI: 10.5946/ce.2017.170]

46 **Yamashita Y**, Kato J, Ueda K, Nakamura Y, Abe H, Tamura T, Itonaga M, Yoshida T, Maeda H, Moribata K, Niwa T, Maekita T, Iguchi M, Tamai H, Ichinose M. Contrast-enhanced endoscopic ultrasonography can predict a higher malignant potential of gastrointestinal stromal tumors by visualizing large newly formed vessels. *J Clin Ultrasound* 2015; **43**: 89-97 [PMID: 25043900 DOI: 10.1002/jcu.22195]

47 **Hadjimichael AC**, Pergaris A, Kaspiris A, Foukas AF, Theocharis SE. Liquid Biopsy: A New Translational Diagnostic and Monitoring Tool for Musculoskeletal Tumors. *Int J Mol Sci* 2021; **22** [PMID: 34768955 DOI: 10.3390/ijms222111526]

48 **Masaoutis C**, Korkolopoulou P, Theocharis S. Exosomes in sarcomas: Tiny messengers with broad implications in diagnosis, surveillance, prognosis and treatment. *Cancer Lett* 2019; **449**: 172-177 [PMID: 30779943 DOI: 10.1016/j.canlet.2019.02.025]

49 **Lone SN**, Nisar S, Masoodi T, Singh M, Rizwan A, Hashem S, El-Rifai W, Bedognetti D, Batra SK, Haris M, Bhat AA, Macha MA. Liquid biopsy: a step closer to transform diagnosis, prognosis and future of cancer treatments. *Mol Cancer* 2022; **21**: 79 [PMID: 35303879 DOI: 10.1186/s12943-022-01543-7]

50 **Heydari R**, Abdollahpour-Alitappeh M, Shekari F, Meyfour A. Emerging Role of Extracellular Vesicles in Biomarking the Gastrointestinal Diseases. *Expert Rev Mol Diagn* 2021; **21**: 939-962 [PMID: 34308738 DOI: 10.1080/14737159.2021.1954909]

51 **Koulouris A**, Tsagkaris C, Messaritakis I, Gouvas N, Sfakianaki M, Trypaki M, Spyrou V, Christodoulakis M, Athanasakis E, Xynos E, Tzardi M, Mavroudis D, Souglakos J. Resectable Colorectal Cancer: Current Perceptions on the Correlation of Recurrence Risk, Microbiota and Detection of Genetic Mutations in Liquid Biopsies. *Cancers (Basel)* 2021; **13** [PMID: 34298740 DOI: 10.3390/cancers13143522]

52 **Ko TK**, Lee E, Ng CC, Yang VS, Farid M, Teh BT, Chan JY, Somasundaram N. Circulating Tumor DNA Mutations in Progressive Gastrointestinal Stromal Tumors Identify Biomarkers of Treatment Resistance and Uncover Potential Therapeutic Strategies. *Front Oncol* 2022; **12**: 840843 [PMID: 35273917 DOI: 10.3389/fonc.2022.840843]

53 **Li J**, Guo S, Sun Z, Fu Y. Noncoding RNAs in Drug Resistance of Gastrointestinal Stromal Tumor. *Front Cell Dev Biol* 2022; **10**: 808591 [PMID: 35174150 DOI: 10.3389/fcell.2022.808591]

54 **Sergei B**, Pavel D, Aigul G, Firyuza B, Ilmira N, Ilshat M, Aida A, Refat K, Natalia A, Elena S, Vera G. Inhibition of FGFR2-Signaling Attenuates a Homology-Mediated DNA Repair in GIST and Sensitizes Them to DNA-Topoisomerase II Inhibitors. *Int J Mol Sci* 2020; **21** [PMID: 31948066 DOI: 10.3390/ijms21010352]

55 **Boichuk S**, Galembikova A, Mikheeva E, Bikinieva F, Aukhadieva A, Dunaev P, Khalikov D, Petrov S, Kurtasanov R, Valeeva E, Kireev I, Dugina V, Lushnikova A, Novikova M, Kopnin P. Inhibition of FGF2-Mediated Signaling in GIST-Promising Approach for Overcoming Resistance to Imatinib. *Cancers (Basel)* 2020; **12** [PMID: 32599808 DOI: 10.3390/cancers12061674]

56 **Napolitano A**, Ostler AE, Jones RL, Huang PH. Fibroblast Growth Factor Receptor (FGFR) Signaling in GIST and Soft Tissue Sarcomas. *Cells* 2021; **10** [PMID: 34204560 DOI: 10.3390/cells10061533]

57 **Pergaris A**, Danas E, Goutas D, Sykaras AG, Soranidis A, Theocharis S. The Clinical Impact of the EPH/Ephrin System in Cancer: Unwinding the Thread. *Int J Mol Sci* 2021; **22** [PMID: 34445116 DOI: 10.3390/ijms22168412]

58 **Papadakos SP**, Petrogiannopoulos L, Pergaris A, Theocharis S. The EPH/Ephrin System in Colorectal Cancer. *Int J Mol Sci* 2022; **23** [PMID: 35269901 DOI: 10.3390/ijms23052761]

59 **Zhao Y**, Wang Q, Deng X, Zhao Y. Altered angiogenesis gene expression in gastrointestinal stromal tumors: potential use in diagnosis, outcome prediction, and treatment. *Neoplasma* 2012; **59**: 384-392 [PMID: 22489693 DOI: 10.4149/neo\_2012\_050]

60 **Kang BW**, Kim JG, Chae YS, Bae HI, Kwon O, Chung HY, Yu W, Song HS, Kang YN, Ryu SW, Lee KH, Bae YK, Choi JH, Kim SW, Ryoo HM, Cho CH, Chae HD, Park KW, Gu MJ, Bae BJ. Clinical significance of vascular endothelial growth factor and vascular endothelial growth factor receptor-2 gene polymorphisms in patients with gastrointestinal stromal tumors. *Asia Pac J Clin Oncol* 2014; **10**: e40-e45 [PMID: 23551429 DOI: 10.1111/ajco.12068]

61 **Toda-Ishii M**, Akaike K, Suehara Y, Mukaihara K, Kubota D, Kohsaka S, Okubo T, Mitani K, Mogushi K, Takagi T, Kaneko K, Yao T, Saito T. Clinicopathological effects of protein phosphatase 2, regulatory subunit A, alpha mutations in gastrointestinal stromal tumors. *Mod Pathol* 2016; **29**: 1424-1432 [PMID: 27469332 DOI: 10.1038/modpathol.2016.138]

62 **Liu N**, Huang J, Sun S, Zhou Z, Zhang J, Gao F, Sun Q. Expression of matrix metalloproteinase-9, cyclooxygenase-2 and vascular endothelial growth factor are increased in gastrointestinal stromal tumors. *Int J Clin Exp Med* 2015; **8**: 6495-6501 [PMID: 26131278]

63 **Takahashi R**, Tanaka S, Kitadai Y, Sumii M, Yoshihara M, Haruma K, Chayama K. Expression of vascular endothelial growth factor and angiogenesis in gastrointestinal stromal tumor of the stomach. *Oncology* 2003; **64**: 266-274 [PMID: 12697968 DOI: 10.1159/000069316]

64 **Verboom MC**, Kloth JSL, Swen JJ, van der Straaten T, Bovée JVMG, Sleijfer S, Reyners AKL, Mathijssen RHJ, Guchelaar HJ, Steeghs N, Gelderblom H. Genetic polymorphisms in angiogenesis-related genes are associated with worse progression-free survival of patients with advanced gastrointestinal stromal tumours treated with imatinib. *Eur J Cancer* 2017; **86**: 226-232 [PMID: 29054076 DOI: 10.1016/j.ejca.2017.09.025]

65 **Chen WT**, Huang CJ, Wu MT, Yang SF, Su YC, Chai CY. Hypoxia-inducible factor-1alpha is associated with risk of aggressive behavior and tumor angiogenesis in gastrointestinal stromal tumor. *Jpn J Clin Oncol* 2005; **35**: 207-213 [PMID: 15845570 DOI: 10.1093/jjco/hyi067]

66 **Basilio-de-Oliveira RP**, Pannain VL. Prognostic angiogenic markers (endoglin, VEGF, CD31) and tumor cell proliferation (Ki67) for gastrointestinal stromal tumors. *World J Gastroenterol* 2015; **21**: 6924-6930 [PMID: 26078569 DOI: 10.3748/wjg.v21.i22.6924]

67 **Imamura M**, Yamamoto H, Nakamura N, Oda Y, Yao T, Kakeji Y, Baba H, Maehara Y, Tsuneyoshi M. Prognostic significance of angiogenesis in gastrointestinal stromal tumor. *Mod Pathol* 2007; **20**: 529-537 [PMID: 17334345 DOI: 10.1038/modpathol.3800767]

68 **Wang TB**, Qiu WS, Wei B, Deng MH, Wei HB, Dong WG. Serum vascular endothelial growth factor and angiogenesis are related to the prognosis of patients with gastrointestinal stromal tumors. *Ir J Med Sci* 2009; **178**: 315-320 [PMID: 19367428 DOI: 10.1007/s11845-009-0315-7]

**Footnotes**

**Conflict-of-interest statement:** All theauthors report no relevant conflicts of interest for this article.

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**Provenance and peer review:** Invited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Peer-review started:** March 20, 2022

**First decision:** April 25, 2022

**Article in press:**

**Specialty type:** Oncology

**Country/Territory of origin:** Greece

**Peer-review report’s scientific quality classification**

Grade A (Excellent): 0

Grade B (Very good): B, B

Grade C (Good): C

Grade D (Fair): 0

Grade E (Poor): 0

**P-Reviewer:** Lewitowicz P, Poland; Messias LHD **S-Editor:** Fan JR **L-Editor:** A **P-Editor:** Fan JR

**Table 1 The basic mechanisms of angiogenesis**

|  |  |  |
| --- | --- | --- |
| **Angiogenetic mechanism** | **Function** | **Implicated signaling/ pathways** |
| Sprouting angiogenesis | Vessel formation from a parental one as a sprout outgrowth | VEGF, Dll4/notch pathways and neuropilins |
| Intussusceptive Angiogenesis | Splitting of a parental vessel into two newly formed | VEGF, PDGF pathways and erythropoietin |
| Vasculogenesis/Endothelial progenitor cells | Vessel formation from endothelial progenitor cells differentiating into mature endothelial cells | VEGF pathway, chemokines |
| Vasculogenic mimicry | Vessel-like formations without endothelial cells | HGFR |
| Trans-differentiation of CSCs | CSC give rise to endothelial cells | Tie-2, TGF-β, CXCL12/CXCR4 |

PDGF-R: Platelet-derived growth factor receptor; VEGF: Vascular endothelial growth factor; HGFR: Hepatocyte growth factor receptor; TGF-β: Transforming growth factor-β; CSCs: Cancer stem cells, CXCL12: C-x-c motif chemokine ligand 12; CXCR4: C-x-c motif chemokine receptor 4.

**Table 2 A brief presentation of several angiogenetic molecules in disease progression**

|  |  |  |  |
| --- | --- | --- | --- |
| **Ref.** | **Sample size** | **Molecule/methods** | **Outcomes** |
| Zhao *et al*[59] | 124 patients-62, 50% in stomach, 22.6% in small intestine | HIF-1α/IHC | Association with disease-free survival (*P* = 0.03) |
| VEGF/IHC | Association with disease-free survival (*P* = 0.002) |
| MVD/IHC | Association with disease-free survival (*P* < 0.001) |
| Kang *et al*[60] | 213 patients-63% in stomach, 25.3% in small intestine | 634G/C | Superior OS than 634 G/G (*P* = 0.054) |
| Superior RFS than 634 G/G (*P* = 0.082) |
| Mu *et al*[22] | 20 patients | BRD4/mRNA, IHC | Increased BRD4 expression compared with normal tissue |
| BRD4/IHC | Associated with poor OS (*P* < 0.01) |
| Associated with poor DFS (*P* < 0.01) |
| Toda-Ishii *et al*[61] | 94 patients–mean follow-up period 65 mo | PPP2R1A mutations/PCR | Lower OS (*P* < 0.05) |
| Lower DFS (*P* < 0.05) |
| Liu *et al*[62] | 52 patients–27 malignant cases–11 borderline–14 benign | MMP-9, COX-2, VEGF/IHC | Enhance metastasis (*P* = 0.014, *P* = 0.010, *P* = 0.032 respectively) |
| Higher mitotic count (*P* = 0.021, *P* = 0.027, *P* = 0.009 respectively) |
| Higher incidence of central necrosis (*P* < 0.01) |
| Takahashi *et al*[63] | 53 patients: 21 cases < 30 mm-9 cases with liver metastasis | VEGF/IHC | Association with liver metastasis (*P* < 0.01) |
| VEGF/IHC | Poor 10-yr OS (*P* < 0.05) |
| MVD/IHC | Association with liver metastasis (*P* < 0.05) |
| Verboom *et al*[64] | 227 patients-36 *SNPs-18* genes, median PFS 39 mo–median OS 86.5 mo | rs1570360 polymorphism in *VEGFA* gene | Association with poorer PFS (*P* = 0.015) |
| rs1870377 polymorphism in *VEGFR2* gene | Association with lower PFS (*P* = 0.037) |
| Chen *et al*[65] | 62 patients: 31 high risk–31 low risk | HIF-1α/IHC | Association with high risk disease (*P* < 0.0001) |
| Association with GIST recurrence or metastasis (*P* = 0.009) |
| Basilio-de-Oliveira and Pannain[66] | 54 patients | VEGF/IHC | Association with survival (*P* < 0.001) |
| CD105/IHC | Association with prognosis (*P* < 0.001) |
| Imamura *et al*[67] | 95 patients: 64 cases in stomach–31 in small intestine | MVD/IHC | Association with tumor grade (*P* = 0.036) |
| Association with VEGF expression (*P* < 0.0001) |
| Association with DFS after surgery (*P* = 0.0028) |
| Wang *et al*[68] | 68 patients: 20 low risk cases–48 high risk cases | Soluble VEGF | Association with lower DSS (*P* < 0.05) |
| VEGF/IHC |
| MVD/IHC |

OS: Overall survival; DSS: Disease-specific survival; DFS: Disease-free survival; PFS: Progression-free survival; VEGF: Vascular endothelial growth factor; IHC: Immunohistochemistry; MVD: Microvascular density; HIF: Hypoxia-inducible factor.