Targeting gastrointestinal stromal tumors: the role of regorafenib

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Abstract: Gastrointestinal stromal tumor (GIST) is a devastating disease in the metastatic setting, but its natural history has been dramatically altered by the development of small molecule tyrosine kinase inhibitors, most notably imatinib. Although patients with advanced GIST live much longer today than they did in the past, imatinib-refractory disease remains a tremendous problem. For disease that is refractory to imatinib and sunitinib, regorafenib is an excellent option. In this review, we discuss the biology and clinical work establishing regorafenib as the standard of care for advanced GIST refractory to both imatinib and sunitinib.

Keywords: regorafenib, GIST, refractory, imatinib

Introduction to gastrointestinal stromal tumors

Gastrointestinal stromal tumor (GIST) is one of the most common soft tissue sarcoma subtypes; each year ~3,300–6,000 new cases are diagnosed.1,2 They occur largely in middle-aged and elderly persons, with ~60% located in the stomach, 30% in the small intestine, and 10% in other regions of the gastrointestinal (GI) tract.3 Size, mitotic rate, and location of the primary lesion are the most important prognostic factors.4

The initial presentation of patients with GIST often reflects the underlying biology of the tumor. GISTs are often highly vascularized and may be relatively large at presentation. GI bleeding occurs in more than half of the patients; a palpable mass is present in one-third of the patients. Pain, in some cases due to obstruction, occurs in nearly one-quarter of the patients.5 Nonspecific presenting symptoms are common, such as early satiety or bloating.

Because of the initial symptoms related to abdominal pathology, computed tomography (CT) is a frequent initial imaging modality in patients with GIST. Triple-phase CT (intravenous and oral contrast with venous and arterial phases) allows observation of the extent and vascularity of these tumors, and assesses for hepatic involvement.5 GISTs typically metastasize to the liver and GI tract, although dissemination to other sites, notably the lungs, is also possible. CT imaging of these regions can provide staging information in these areas.

While clinical and radiographic findings may be suggestive, the definitive diagnosis of GIST relies on tissue diagnosis. Histologically, GIST can have a variable appearance, but has been classified in the past as frequently displaying features of smooth muscle tumors. A key observation in GIST diagnosis and therapy was the central role of mitogen receptor and GIST marker, c-Kit; gain-of-function of this receptor is crucial for tumor growth in most cases of GISTs.6 Overexpression of c-Kit protein is detected immunohistochemically in the majority of GISTs, which is a key diagnostic finding.
GIST can also occur from mutations in the platelet-derived growth factor receptor-α (PDGFRα) gene. So-called wild-type GISTs lack mutations in both KIT, and PDGFRα.7 The GIST morphology ranges from predominantly spindle-shaped (70%), epithelioid (20%), to a mixed phenotype (10%). A minor subset (5%) of GISTs are c-Kit negative, and these often occur in the stomach with an epithelioid cell morphology and a usually less aggressive course.10 DOG1 (ANO1) is expressed in GISTs regardless of KIT expression and supports the diagnosis.

In patients with primary and localized GIST, the standard therapy is surgical resection with the intention to cure. In the pre-imatinibera, approximately one-half of the patients diagnosed with localized GIST would relapse and patients with recurrent, unresectable, and/or metastatic GIST would die in their first year following diagnosis. Today, complete resection for primary GIST is associated with a 5-year recurrence-free survival rate of 70%. Image-guided core needle biopsy is also useful in diagnostic evaluation and should be done first if GIST is suspected in the case of a large tumor and where neoadjuvant imatinib is being considered.4

**Tyrosine kinase inhibitors have revolutionized the treatment of GIST**

In 1998, investigators made a critical discovery showing that gain-of-function mutations in the proto-oncogene for the tyrosine kinase receptor c-Kit drive GISTs.12 These mutations are present in over 85% of GISTs and are able to induce tumor growth in vitro and in vivo.13,14 Exon 11 encodes the protein’s juxtamembrane domain and is the most common location for mutations. Mutations of exon 11 permit ligand-independent receptor activation and independent tumor growth.15,16 The prognostic significance associated with different KIT mutations can be profound; exon 11 KIT mutations yield ~89% 5-year survival versus 40% for other mutation types.13,16,17 Most KIT mutations are found in exons 9, 11, 13, or 17.

A treatment breakthrough occurred in 2001, when the first case report was published describing a patient with GIST who had successfully been treated with the tyrosine kinase inhibitor (TKI), imatinib. This patient had rapidly progressive, metastatic GIST. The patient progressed despite multiple prior systemic therapies (dacarbazine, doxorubicin, ifosfamide, interferon-α, and thalidomide). Imatinib led to a complete metabolic response within 1 month of treatment initiation.18 The same year, researchers assembled 36 patients with GIST with advanced disease to evaluate the safety and efficacy of imatinib.19 Minor responses were elicited in 17% (six of 36) of patients, while partial remissions were achieved in 53% (19 of 36) of patients.

In 2002, 147 patients with advanced GIST were enrolled in a multicenter Phase II trial in which they were randomized to receive either 400 or 600 mg of imatinib daily. The patients had undergone a number of previous treatment modalities: 15% received radiotherapy, 51% received chemotherapy, and 98% had a previous operation. No patient achieved a complete response, but 54% (79 patients) had partial response and 28% (41 patients) had stable disease (treatment benefit lasting >6 months). Neither dose was superior, although the study was underpowered to detect a difference at 52-month follow-up.20 Subsequent studies have continued to document the efficacy of imatinib in both the metastatic and adjuvant settings.19–23

Imatinib has become the gold standard for GIST therapy, as it provides a stable response, typically for 18–36 months. Prior to the introduction of imatinib, a median survival time of 10–20 months was generally seen. Today, this has improved to 51–57 months.24 Fortunately, imatinib has a manageable toxicity profile. Common side effects include diarrhea, nausea, headache, edema, myalgia, rash, and some abdominal discomfort.

Despite the vast improvement in GIST therapy with the advent of imatinib, the drug is not curative for patients with metastatic disease, and resistance ultimately emerges. Investigation of planned salvage therapies has been undertaken. For example, the multitargeted TKI sunitinib has been approved for treatment of patients with GIST after progression on imatinib therapy. Sunitinib, improves progression-free survival (PFS) in imatinib-refractory patients with GIST, particularly so in patients with exon 9 KIT mutations.25 Sunitinib can also be given on the continuous daily dosing schedule without apparent loss of efficacy.25

**Regorafenib is a small molecule TKI that also inhibits angiogenesis**

GIST biology has proven to be more complex than originally believed, as evidenced by the molecular heterogeneity found in all GISTs and the numerous subgroups identified.26 These tumors are distinct from nerve sheath tumors or GI smooth muscle tumors, and may originate in the interstitial cells of Cajal.27 Activating mutations of c-KIT are not sufficient to cause GIST; additional genomic alterations are present, but their exact biological contribution remain largely unknown.28–30

The success of imatinib and sunitinib raised interest in a wide variety of multitargeted small molecule TKIs that affect c-Kit to varying degrees, such as dasatinib and sorafenib.31,32
During this period of TKI investigation, regorafenib’s potent inhibitory activity was recognized.\textsuperscript{33–35} While orally available and structurally similar to sorafenib, regorafenib is a TKI against multiple targets important for tumor angiogenesis, oncogenesis, and overall maintenance of the tumor microenvironment (Figure 1).\textsuperscript{33} Vascular endothelial growth factor receptor 2 (VEGFR-2) and other tyrosine kinases with immunoglobulin and epidermal growth factor homology domain 2 are critical to the biology of both normal and tumor vasculatures, but have been targeted successfully in cancer. While angiogenesis is a well-recognized component of tumor development and has been demonstrated to be an important therapeutic target in many malignancies,\textsuperscript{36,37} its role in GIST was largely underappreciated despite the frequent vascularity of these tumors. Similar to tumor growth, angiogenesis is governed by interconnected tyrosine kinase-driven signaling pathways, which are affected by other small molecule TKI GIST therapies, including sunitinib and sorafenib.

Similar to other small molecule TKIs, regorafenib has direct inhibitory effects on tumor growth through inhibition of KIT, as well as Ras and B-Rapidly Accelerated Fibrosarcoma (BRAF) downstream of Platelet Derived Growth Factor Receptor Beta (PDGFR\textbeta).\textsuperscript{38} For patients with some imatinib resistance mediated by secondary KIT mutations, direct inhibition of c-Kit by regorafenib may play a particularly important role.\textsuperscript{39}

Interestingly, in colorectal cancer (CRC) cell lines, regorafenib has been demonstrated as an important agonist of the tumor suppressor Src Homology region 2 domain containing Phosphatase-1 (SHP-1).\textsuperscript{40} Regorafenib may also alter mitogen activated protein (MAP) kinases, Extracellular signal Regulated Kinase (ERK) pathway, and c-Jun N-terminal Kinase (JNK/Jun) in complex ways that are incompletely characterized.\textsuperscript{41–43} In other systems, increased cytosolic Ca\textsuperscript{++} may play a role.\textsuperscript{44} Regorafenib directly impairs mitochondrial function by uncoupling oxidative phosphorylation, which may play a role in rare reports of hepatic toxicity.\textsuperscript{45}

Unlike imatinib, and to a greater degree than other small molecule TKIs, regorafenib inhibits endothelial cells based on biochemical and cellular kinase proliferation assays by targeting VEGFR-2 and TIE-2.\textsuperscript{33} This additional activity allows regorafenib to maintain activity even in xenograft models of GIST that are highly resistant to c-Kit targeting.\textsuperscript{39} Similar explanations can be invoked for the use of regorafenib in GIST.

VEGFR targeting is mechanistically important, as regorafenib improves overall survival in patients with metastatic, treatment-refractory CRCs.\textsuperscript{47,48} Given its similarity to pazopanib, which improves PFS in soft tissue sarcoma patients,\textsuperscript{49,50} regorafenib is now being broadly tested in soft tissue sarcomas.\textsuperscript{51}

**Regorafenib has a toxicity profile acceptable for most patients with refractory metastatic GIST**

In the first reported Phase I trial of regorafenib, the drug was used in patients with advanced solid tumors refractory to...
standard chemotherapy. A total of 53 patients were recruited for this study to assess the drug’s safety, pharmacokinetics, pharmacodynamics, and efficacy profile.

Dosing on the Phase I study ranged from 10 to 220 mg/day on a 21 of 28-day oral cycle. While five of 12 patients treated at the 220 mg/day dosing experienced toxicities, none were life-threatening. The 160 mg/day dosing schedule was deemed superior based on its toxicity profile (two of 12 patients at this dose had dose-limiting toxicity) and was selected for further investigation.

Surprisingly, given a highly refractory and diverse patient population, 66% of these patients experienced disease control (ie, partial response or stable disease). An expanded Phase I cohort analyzed the safety in metastatic CRC.

The safety of regorafenib has been tested extensively, and its toxicity profile is well established and consistent (Table 1). Like other VEGFR-targeted TKIs, regorafenib can induce hypertension, requiring additional antihypertensive therapy. Another common toxicity of regorafenib in some patients is hand–foot skin reactions (HFSRs). HFSR is not as frequent in patients on sunitinib, but grade 3 or higher HFSR has been observed in all the randomized controlled trials of regorafenib. Maculopapular rashes and generalized desquamation can also occur, but are rarely severe. Diarrhea occurs in <10% of patients treated with regorafenib. It is generally not severe and can be responsive to Lomotil (diphenoxylate and atropine). Although rare, regorafenib has been associated with fatal (grade 5) toxicities, including cardiac arrest and hepatic failure.

There is also an association between regorafenib therapy and an increased incidence of myocardial ischemia, potentially linked to the inhibition of VEGF pertaining to its role in cardiovascular function. Patients with preexisting cardiovascular comorbidities, including venous thrombus, embolic events such as stroke or transient ischemic attacks, pulmonary embolus, cardiac arrhythmias requiring antiarrhythmic therapy, uncontrolled hypertension despite optical medical management, or severe heart failure within 6 months before the start of treatment were excluded from the trial. Although a Phase I trial assessing the cardiovascular safety of regorafenib in solid tumors showed the drug to have modest effects on QT/QTc and left ventricular ejection fraction. In 72% of patients treated with regorafenib, the drug dosage had to be modified due to adverse events. However, the incidence of adverse events leading to permanent drug discontinuation was 6.1% in the regorafenib arm, similar to the 7.6% in the placebo-treated patients.

Despite some potentially serious toxicities, regorafenib is well tolerated in the vast majority of patients. Patients participating in a randomized Phase III trial of regorafenib were given a quality-of-life survey. There was no significant difference in the quality-of-life between patients treated with regorafenib and those receiving best supportive care, though quality-of-life worsened after disease progression.

**Regorafenib improves PFS for patients with refractory metastatic GIST**

The preliminary efficacy from the Phase I study was further evaluated in a multicenter, Phase II study of regorafenib in 33 patients. These patients had previously been treated with both imatinib and sunitinib, yet they progressed, or they were unable to tolerate the treatments. Twenty-six patients

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<th>Toxicity</th>
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<th>GRID Grade 4 (%)</th>
<th>GRID Grade 5 (%)</th>
<th>CORRECT* Grade 3 (%)</th>
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Note: *The CORRECT trial used regorafenib in the treatment of colorectal cancer, not gastrointestinal tumors.

Abbreviations: GRID, GIST-Regorafenib In progressive Disease; HFSR, hand–foot skin reaction; CORRECT, regorafenib monotherapy for previously treated metastatic colorectal cancer.
(75%) derived clinical benefit from regorafenib, as evidenced by either a partial or complete tumor response, or stable disease for a minimum of 16 weeks. Most patients (22 of 26) had demonstrated stable disease, but four exhibited partial responses by response evaluation criteria in solid tumors (RECIST). The lack of substantial tumor remission in a majority of treated patients, as has been seen with other TKIs in the treatment of GIST, might suggest that alternative response assessment criteria should be used.

PFS for the entire cohort on the Phase II study was 10 months (95% confidence interval, 8.3–14.9 months). Patients with exon 11 KIT mutations experienced longer PFS versus those with exon 9 mutations. However, this was an exploratory analysis based on a small sample and may be secondary to the relative aggressiveness of the two mutational domains, rather than any increased associated response to therapy. Clinical responses were seen despite dose reductions in a majority of patients for grade 3 toxicities affecting their quality-of-life.

These results led to the GIST-Regorafenib In progressive Disease (GRID) trial, an international, multicenter, randomized, placebo-controlled, Phase III trial with 199 patients who had either metastatic or unresectable disease. Furthermore, these patients previously had unsuccessful treatment with imatinib (understood as progression or intolerance to treatment), and were unsuccessfully treated with sunitinib (identified by disease progression). Patients were randomized using 2-to-1 to treatment with either regorafenib or placebo. All patients also received best supportive care (defined as any method to preserve the comfort and dignity of the patient, excluding disease-specific antineoplastic therapy). Oral regorafenib treatment was given at the same dose and schedule as the Phase II trial, with 160 mg/day taken for 21 days of each 28-day cycle.

The primary endpoint of the study was PFS, which was 4.8 months for those who received regorafenib, versus 0.9 months for those who received placebo (hazard ratio =0.27; P<0.00001). No patient in either group demonstrated a complete response. The disease control rate (defined as rate of complete or partial response plus stable disease lasting at least 12 weeks) was 52.6% (70 of 133) for regorafenib and 9.1% (six of 66) for placebo (95% confidence interval, 54.72–32.49; P<0.00001).

There was no difference in the overall survival between the two arms. The study was not intended to detect a survival benefit; crossover from the placebo group to the active treatment was allowed, with 85% of placebo-treated patients ultimately receiving regorafenib at the time of progression. Toxicity in the GRID trial (Table 1) was similar to regorafenib toxicities seen in other trials, including HFSR (56.1% vs 15.2%, regorafenib vs placebo), hypertension (48.5% vs 16.7%), diarrhea (40.9% vs 7.6%), and oral mucositis (37.9% vs 9.1%).

**Future directions**

Regorafenib is now approved for the treatment of GIST in the US and Europe and is considered by most experts to be the standard-of-care third-line agent for patients with metastatic or unresectable GIST that is refractory to imatinib and sunitinib. The impressive clinical activity in this highly refractory patient population raises the question as to whether regorafenib’s use as first- or second-line treatment would be beneficial. In light of the established benefits, including overall survival benefits, or imatinib therapy, the design of clinical trials to test this hypothesis may present some challenges.

Given the multiple kinase targets of regorafenib (Figure 1), patients treated with regorafenib as a first-line agent may be less prone to resistance; this proposition merits further investigation as it may provide further support for testing regorafenib earlier in the GIST treatment continuum. A point of interest is why some patients with KIT mutations, such as exon 17, tend to have resistance to nilotinib, yet they show response to regorafenib. An ongoing trial is evaluating the activity of regorafenib in patients with a secondary mutation in exon 17 (NCT 02606097). Trials testing the role of regorafenib in the curative setting either as an alternative to imatinib or in combination with imatinib are compelling, particularly in the neoadjuvant setting where significant questions remain regarding which patients benefit most from imatinib treatment and for how long therapy should continue.

Regorafenib-refractory disease remains a critical problem, but a number of different strategies are under investigation. New TKIs, such as ponatinib (NCT 01874665), are under investigation. Combining small molecule TKIs along with other novel agents may allow treatment to overcome resistance. The observation that PI3 kinases (PI3K) and mechanistic target of rapamycin (MTOR) are upregulated in imatinib-resistant GIST lines led to the dose-finding study of the novel PI3K inhibitor, BKM120, in combination with imatinib (NCT 01468688). For some patients, BRAF resistance may play a role in imatinib resistance and targeting this pathway may play a role in treatment. Small molecule TKIs may have increased activity when combined with each other such as imatinib and sorafenib. The ongoing SURE trial is evaluating an alternating schedule of regorafenib and sunitinib (NCT 02164240).
Monoclonal antibodies recognizing KIT are able to control imatinib-refractory GIST in xenograft models. These antibodies may directly slow GIST and enable macrophage-induced tumor clearance. These infiltrating immune and stromal cells play a critical role in the immune response to GIST, and the combination of TKI with novel immunotherapies is another source of great excitement in the field.

Understanding genetic signatures and complex signaling networks are crucial for yielding successful outcomes using targeted therapies. Since regorafenib targets multiple kinases, further investigation of the interplay between oncogenesis and angiogenesis may explain treatment successes, as well as failures. One challenge that remains is cancer resistance to drug intervention, which inevitably leads to progression and a fatal outcome. In an era of next-generation sequencing, longitudinal molecular monitoring may allow us to understand better the evolution of drug resistance against current and future regimens. Study of regorafenib-treated patients may be especially pertinent, as most regorafenib-treated patients have been previously exposed to other agents (ie, imatinib and sunitinib). Combinations with other molecularly targeted agents, such as MTOR inhibitors, may unlock crucial synergies allowing patients to maintain efficacy for much longer.

Therapy of GIST continues to be an exciting field of investigation, and regorafenib has been an important addition to our anti-GIST armamentarium. Its use has generated important questions, the answers to which may improve the use of regorafenib and shed additional light on GIST biology. This will contribute to the continuing progress against this entity.

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