**Gastrointestinal Stromal Tumor – An Overview**

**Abstract**
Gastrointestinal stromal tumors (GISTs) are rare tumors but are most common mesenchymal tumors of the digestive tract. They are commonly seen in the stomach (60%) and small intestine (30%). GISTs are likely derived from the interstitial cells of Cajal or their stem cell precursors. They are best characterized by computerized tomography and have a specific staining pattern on immunohistochemistry, i.e., C-Kit and DOG-1. The treatment of GIST is based on the risk assessment for relapse, and patients with localized GIST require resection with or without adjuvant imatinib mesylate (IM). Advanced unresectable tumors are usually treated with IM, with a number of further options available for patients post progression on IM. There is an increasing emphasis on identifying C-Kit and platelet-derived growth factor receptor alpha mutations in all patients with GIST, as these are driver mutations with current and future therapeutic implications.

**Keywords:** Gastrointestinal stromal tumor, Indian data, overview

**Introduction**
Gastrointestinal stromal tumors (GISTs), in general, are rare tumors but they are the most common mesenchymal tumors of the digestive tract and rarely can arise from the intra-abdominal soft tissues.[1] They are commonly seen in the stomach (60%) and small intestine (30%) but are also seen in the rectum and colon.[2,3] The discovery of the c-KIT gene (cellular homolog of the oncogene v-KIT) set the ball rolling in elucidation of the pathogenesis of GIST as well as its classification as a separate disease entity.[4,5]

**Global Incidence, Epidemiology – Global and India**
The global incidence of GIST is unknown due to the rarity of the disease, but available data suggest variances across geographical regions. Reported incidence from Northern Norway, Hong Kong, and Korea is approximately 9–22 cases per million inhabitants, while incidences are lower from North America, Slovakia, etc., (4.3–6.8 cases per million).[6] Although GISTs can arise at any age, they are most commonly seen beyond the age of 50 years (median age – 63 years). There is no large scale data from India with regard to the incidence or clinical presentation. Small single-institution studies have shown a median age range at a diagnosis of 50–58 years, with a greater incidence of presentation with advanced/metastatic disease, though this is possibly due to underreporting of early cases.[7,8]

**Clinical Presentation – Global and India**
Approximately 18%–25% of patients have been diagnosed with GIST based on imaging or while being investigated for other illnesses. The most common site is the stomach (60%), but GISTs can be found throughout the gastrointestinal (GI) tract including the jejunum and ileum (30%), duodenum (5%), colon/rectum (4%), and esophagus or appendix (<1%). Occasionally, they may present with emergent complications such as hemorrhage, tumor rupture, bowel perforation, or obstruction. The increased awareness of GIST as a different disease with improved diagnostic criteria and routine use of adjuvant imatinib have resulted in the pickup of smaller tumors at the diagnosis of GIST. This has resulted in a new subgroup of GISTs called mini-GISTs (measuring between 1 and 2 cm) and micro-GISTs (measuring < 1 cm). GISTs are rare in the pediatric age group, and most of them are observed in the second decade with a female predisposition.

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They almost exclusively arise in the stomach with frequent nodal involvement in this age group.\textsuperscript{[9]}

A snapshot of Indian data with regard to epidemiology and presentation is presented in Table 1.\textsuperscript{[7-13]}

**Molecular Basis of Gastrointestinal Stromal Tumor**

The central role in the pathogenesis of GIST is occupied by the KIT and platelet-derived growth factor receptor alpha (PDGFRA) genes. These genes encode tyrosine kinase receptors comprising an extracellular ligand-binding region, a transmembrane sequence, a juxtamembrane domain, and two cytoplasmic kinase domains.\textsuperscript{[14]} Mutations in KIT and PDGFRA result in constitutive activation and signaling (in the absence of the endogenous ligand) causing a chain of cellular events, leading to the advent of GIST. A majority of mutations are seen in the KIT gene (approximately 80%, predominantly juxtamembrane) and found in exon 11 (70%) and exon 9 (10%).\textsuperscript{[1,15]} Mutations in exon 11 themselves are a heterogeneous group, with mutations in codon 557–558 of exon 11 having a different biological behavior compared to other exon 11 mutants.\textsuperscript{[16-18]} PDGFRA mutations are seen in <10% of GISTs, primarily either exon 18 or 14. GISTs with PDGFRA mutations are considered less aggressive compared to KIT-mutant GISTs. Approximately 10% of all GISTs are considered “wild type” and are characterized by the absence of KIT and PDGFRA mutations. However, targeted exome sequencing analyses in this “wild type” cohort have identified germ line mutations involving succinate dehydrogenase (SDH), resulting in a complete loss or reduction in SDH protein, thereby causing GIST.\textsuperscript{[19]}

**Evaluation and Workup**

Although GISTs may often be diagnosed incidentally, once suspected, a complete diagnostic evaluation is essential.

a. Endoscopy and endoscopic ultrasound (EUS) – Since small GISTs are often evaluated initially as submucosal tumors, the initial investigation for such tumors would be an EUS-fine-needle aspiration (FNA).\textsuperscript{[20]} EUS-FNA is indicated for lesions measuring >1 cm (usually between 1 and 2 cm) that are suspected to be potentially malignant\textsuperscript{[21]}. For larger tumors of the stomach (potential candidates for resection), an endoscopy is an ideal investigation for description of the tumor as well as taking adequate biopsies.

b. Radiology and Nuclear imaging – The imaging modality of choice for accurate detection and staging of a GIST in the current era is a contrast-enhanced computerized tomographic (CECT) scan. Since GISTs normally arise

<table>
<thead>
<tr>
<th>Centre</th>
<th>Number of patients</th>
<th>Median age</th>
<th>Common sites</th>
<th>Presentation</th>
<th>Highlights</th>
</tr>
</thead>
<tbody>
<tr>
<td>NIMS, Hyderabad\textsuperscript{[7]}</td>
<td>50</td>
<td>50</td>
<td>Stomach, Jejunum, Colon, Small intestine, Stomach, colorectal, Stomach, Small bowel, Duodenum</td>
<td>60% curative resection</td>
<td>Differences in outcomes between intermediate and high risk</td>
</tr>
<tr>
<td>CMC, Vellore\textsuperscript{[8]}</td>
<td>92</td>
<td>4th-5th decade</td>
<td>Stomach, Jejunum, Colon, Small intestine, Stomach, colorectal, Stomach, Small bowel, Duodenum</td>
<td>24% curative resection</td>
<td>70.4% high risk</td>
</tr>
<tr>
<td>GB Pant, New Delhi\textsuperscript{[9]}</td>
<td>92</td>
<td>50</td>
<td>Stomach, Small intestine, Duodenum</td>
<td>80% R0 resection</td>
<td>Nuclear pleomorphism as predictor of recurrence</td>
</tr>
<tr>
<td>The Cancer Institute (WIA), Chennai\textsuperscript{[10]}</td>
<td>24</td>
<td>56</td>
<td>Stomach, Small intestine, Rectum</td>
<td>All metastatic</td>
<td>Common sites of metastases - liver</td>
</tr>
<tr>
<td>Kidwai Memorial, Bengaluru\textsuperscript{[11]}</td>
<td>44</td>
<td>56</td>
<td>Stomach, Small intestine, Stomach, Duodenum</td>
<td>52% localized at presentation</td>
<td>65% high risk</td>
</tr>
<tr>
<td>TMH, Mumbai (nonmetastatic cohort)\textsuperscript{[12]}</td>
<td>103</td>
<td>54</td>
<td>Stomach, Small intestine, Stomach, Duodenum, Jejunum</td>
<td>100% localized</td>
<td>59% high risk, 18% intermediate risk, 21% low risk, 2% very low risk</td>
</tr>
<tr>
<td>TMH, Mumbai (metastatic cohort)\textsuperscript{[13]}</td>
<td>83</td>
<td>54</td>
<td>Stomach, Small intestine, Duodenum, Rectal</td>
<td>All metastatic</td>
<td>Commonest mutation - exon 11 c-kit</td>
</tr>
</tbody>
</table>

NIMS - Nizam’s Institute of Medical Sciences; CMC - Christian Medical College; GB Pant - Govind Ballabh Pant Institute of Post Graduate Medical Education and Research; TMH - Tata Memorial Hospital

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from the outer muscular layers, their exophytic nature is well captured on a CECT.\textsuperscript{[23]} In specific situations such as in the case of anorectal GISTs, a magnetic resonance imaging may provide additional information beyond a CECT.

Positron-emission tomography (PET) provides functional information that may help in staging, especially when combined with morphological information provided by computerized tomography (CT). A PET/CT also helps in differentiating necrotic tissue from viable tissue, recurrent tumor from scar tissue, specifically when assessing response in tumors post therapy.\textsuperscript{[23]} The Choi criteria, a combination of tumor density (15\% change) and modified tumor size (>10\%), is an excellent criteria for early response evaluation and has prognostic value but is yet to be taken as the standard criteria for response assessment in clinical trials in GIST.\textsuperscript{[24]}

c. Pathology – The key to an accurate diagnosis of GIST is the pathological evaluation of a biopsy specimen.

- Gross pathology – GISTs are usually well-circumscribed or multinodular tumor that may develop in any portion of the gut wall, but most GISTs are centered in the submucosa or the muscularis propria. Some tumors are predominantly extramural and extremely large tumors may even extend to or infiltrate into the adjacent organs. On cut sections, the tumors are gray–white and solid with a fleshy appearance and often show hemorrhage, necrosis, or cystic change\textsuperscript{[25]}

- Microscopic features – GI stromal tumors are very cellular lesions, with 70\% of cases composed of spindle cells, 20\% of epithelioid cells, and the remainder having a mixed cellular composition. Depending on the cytological features including cellularity, nuclear atypia, mitosis, and necrosis, they can be further categorized into potentially benign and malignant GIST. However, in GIST, tumor size and mitotic count per 50 HPF are regarded as the best predictor of malignant behavior. No other histologic parameter has correlated as strongly with metastatic risk or survival\textsuperscript{[26]}

- Immunohistochemistry (IHC) – On IHC, KIT (CD117) expression is a sensitive and specific marker for GIST, with about 95\% of GISTs showing a strong and diffuse cytoplasmic staining for KIT, although some tumors, particularly those with an epithelioid morphology, can show membranous staining.\textsuperscript{[27]} CD117 is a highly sensitive marker of GIST which can be consistently expressed in seminomas, mastocytomas, and granulocytic sarcomas, and rarely, expressed in angiosarcoma, metastatic melanoma, clear cell sarcoma, and the Ewing sarcoma/primitive neuroectodermal tumor family of tumors.\textsuperscript{[28]} DOG1 (discovered on GIST1) is also a synchronously used IHC marker for the diagnosis of GIST.\textsuperscript{[29]} The sensitivity of KIT and DOG1, together, approximates 95\% for the diagnosis of GIST, with <3\% of GISTs being KIT and DOG1 negative. Importantly, DOG1 stains about one-third of KIT-negative GIST, thereby making it a useful adjunct in the diagnosis of GIST.\textsuperscript{[30]} Other IHC markers which stain positively in GIST include CD34, caldesmon, smooth muscle actin, desmin, S-100 protein, and keratin

- The differential diagnosis for GIST includes smooth muscle tumors, nerve sheath tumors, fibromatoses, inflammatory fibroid polyps, inflammatory myofibroblastic tumors, follicular dendritic cell tumors, and other types of sarcomas.

**Principles of Management (Overview)**

Once the diagnosis of GIST is established with adequate pathology and radiology, management is based on expected tumor prognosis and staging. A number of prognostic classification systems have developed over the past two decades, including NIH criteria developed by Fletcher, Goh’s modified Armed Forces Institute of Pathology risk criteria, and the Joensuu’s modified NIH criteria.\textsuperscript{[30-33]} Table 2 offers a snapshot of the widely used system as proposed by Miettinen.\textsuperscript{[26,24]} A deficit in these scores is their nonrecognition of KIT mutational status in their risk assessment. It is important to note that these classifications are based on a pathological evaluation of completely resected specimens in patients with nonmetastatic GIST.

**Surgical Management**

**General principles**

Surgery is the primary treatment of choice for patients with localized or potentially resectable GIST lesions. Preoperative histological diagnosis is not mandatory if the diagnosis of GIST is strongly suspected and if it appears to be resectable. Biopsy is necessary to confirm the diagnosis if neoadjuvant imatinib is considered prior to attempted resection in a patient who has a large/locally advanced lesion clinicoradiologically suspected to be a GIST.

The surgical procedure should aim to resect the tumor with histologically negative margins. Segmental or wedge resection to obtain negative margins is often appropriate. On laparotomy/laparoscopy, the abdomen should be thoroughly explored to identify and remove any previously undetected peritoneal metastatic deposits. Although primary GISTs may demonstrate inflammatory adhesions to the surrounding organs, a true invasion is not frequent. GISTs are fragile and should be handled with care to avoid tumor rupture. The goal is to achieve complete gross resection of the tumor with an intact pseudocapsule. Violation of the tumor pseudocapsule causes risk for subsequent tumor seeding into the peritoneum.\textsuperscript{[32]}

Lymphadenectomy is not indicated unless enlarged or pathologic nodes are seen on imaging or intraoperatively, as GISTs rarely spread to regional nodes. A macroscopically complete resection with negative or positive microscopic
Margins (R0 or R1 resection, respectively) is associated with a better prognosis than a macroscopically incomplete excision (R2 excision). Re-resection is generally not indicated for microscopically positive margins on final pathology. The presence of microscopically positive margins (R1) after macroscopic total resection may not confer a worse prognosis. In a review of data from more than 800 patients enrolled in two large North American multi-institutional trials, there was no difference in recurrence-free survival in those who had R1 versus R0 resection. Optimal management after R1 resection is still not well defined and may include re-resection, watchful waiting, and/or systemic therapy.

**Extent of surgical resection**

Complete resection with at least a 1-to 2-cm gross margin should be the goal of surgical treatment.

Gastric tumors, the most common location for GIST, typically require only a partial gastrectomy or even gastric wedge resection to achieve these margins. Partial gastrectomy confers the same progression-free survival (PFS) as total gastrectomy but spares the perioperative and postoperative morbidity of the more extensive surgery. GISTs arising from the small intestine and larger bowel (including rectum) often require segmental resection of the involved gut.

**Minimally invasive approaches**

Minimally invasive resection of gastric GISTs is comparable to open techniques. A meta-analysis of 19 studies (n = 1060 GIST cases) revealed no difference in long-term outcomes of GIST resections using laparotomy and laparoscopy.

Laparoscopic resections of GISTs are associated with decreased blood loss and shorter hospital stays when compared with open surgery. Studies have shown the feasibility of resecting larger gastric tumors with excellent oncologic outcomes. Current guidelines do not generally recommend laparoscopic resection for tumors >5 cm in diameter.

**Management of small gastrointestinal stromal tumors**

All GISTs ≥2 cm in size should be resected. However, there is no or limited consensus on the management of smaller GISTs (<2 cm) and guidelines vary in their recommendations. The natural history of small GISTs, their growth rate, and metastatic potential is unknown. Endoscopic mucosal resection techniques may not be able to provide R0 resections, as GISTs originate in the submucosa. Submucosal dissection technique can be occasionally employed for few small lesions at critical locations or in patients unfit or unwilling for surgery. Gastric GISTs behave less aggressively than the small bowel, colorectal, or GISTs in other locations. Gastric GISTs that are <2 cm in diameter and asymptomatic are currently referred to as very small, mini- (1–2 cm) or micro-(<1 cm) GISTs. Many of these are found only incidentally on endoscopy, in pathologic specimens after gastric resection, or on autopsy. They generally demonstrate benign clinical behavior. Only those lesions that are found to have high-risk features on EUS (irregular borders, cystic spaces, ulceration, echogenic foci, and internal heterogeneity) are considered to be at risk for progression and are considered for surgery.

**Resectable gastrointestinal stromal tumor with a higher risk of perioperative morbidity**

If imaging findings which suggest a complex surgical procedure is required (like total gastrectomy with adjacent organs removal), then a multidisciplinary consultation regarding the use of preoperative imatinib as a neoadjuvant therapy to downsize the tumor and to avoid a multivisceral/morbid resection is recommended. Furthermore, large abdominal tumors felt to be at a significant risk of tumor rupture during surgery and can be treated with preoperative imatinib.

The duration of preoperative medical therapy varies from 3 months to 1 year. However, it is generally 6–12 months, which corresponds to the time interval when the maximum degree of tumor shrinkage is achieved.

**Unresectable, recurrent, or metastatic gastrointestinal stromal tumor**

Imatinib and further TKIs are the primary therapies for metastatic GIST. Surgery is indicated when:

- R0 resection at primary and all metastatic sites is possible
- Limited disease progression refractory to imatinib
- Locally advanced or previously unresectable tumors

<table>
<thead>
<tr>
<th>Tumor size (cm)</th>
<th>Mitotic rate (HPFs)</th>
<th>Gastric</th>
<th>Jejunum/ileum</th>
<th>Duodenum</th>
<th>Rectum</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤2</td>
<td>≤5/50</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>2-5</td>
<td>≤5/50</td>
<td>98.1</td>
<td>95.7</td>
<td>91.7</td>
<td>91.5</td>
</tr>
<tr>
<td>5-10</td>
<td>≤5/50</td>
<td>96.4</td>
<td>76</td>
<td>66</td>
<td>43</td>
</tr>
<tr>
<td>&gt;10</td>
<td>≤5/50</td>
<td>88</td>
<td>48</td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤2</td>
<td>&gt;5/50</td>
<td>100</td>
<td>50</td>
<td>-</td>
<td>46</td>
</tr>
<tr>
<td>2-5</td>
<td>&gt;5/50</td>
<td>84</td>
<td>27</td>
<td>50</td>
<td>48</td>
</tr>
<tr>
<td>5-10</td>
<td>&gt;5/50</td>
<td>45</td>
<td>15</td>
<td>14</td>
<td>29</td>
</tr>
<tr>
<td>10</td>
<td>&gt;5/50</td>
<td>14</td>
<td>10</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2: Rates of progression-free survival for gastrointestinal stromal tumors of the stomach, small intestine, and rectum as per Miettinen’s Classification
after a favorable response to preoperative imatinib

• Management of symptomatic bleeding, obstruction, or similar local tumor-related symptoms.

Medical Management

Brief history

The medical management of GIST revolves around the role of imatinib mesylate (IM, a selective tyrosine-kinase inhibitor of KIT, PDGFRA, and ABL), which was initially used for the treatment of chronic myeloid leukemia with great success. Based on the dramatic response of a heavily pretreated patient with GIST to IM which was published as a brief report in 2001, IM is now almost the universal first line of management for patients who are candidates for systemic treatment.\(^{[43]}\)

Adjuvant treatment post resection of gastrointestinal stromal tumor

The treatment of GIST post resection is heavily dependent on the risk of recurrence as assessed by the risk recurrence and prognostic scores previously discussed. Patients classified as very low risk or low risk need no further treatment post resection and should be kept under surveillance/observation.

For patients with intermediate-risk GIST (recurrence risk 10%–24%), there are differing opinions on the need of adjuvant IM as well as the duration of IM.\(^{[42,44,45]}\) Currently, there is a trend toward treating these patients with adjuvant IM for 3 years, though an individualized approach is recommended.

The evidence for the adjuvant treatment of resected GIST with IM was initially evaluated in tumors considered high risk for recurrence with the ACSOSG Z9000 (Alliance) intergroup phase 2 trial. Their definition of “high risk” tumors, i.e., tumor diameter >10 cm, intraperitoneal tumor rupture, or up to four peritoneal implants (more important from a historic standpoint than currently appropriate or standard), received 1 year of adjuvant IM and showed an improved 1-, 3-, and 5-year overall survival (OS) rate which was 99, 97, and 83%, as compared to historical controls.\(^{[46]}\) The current standard of 3 years of adjuvant IM for resected GIST is based on the SSG XVIII Phase III trial, which evaluated 3 years of adjuvant IM versus 1 year.\(^{[47]}\) The definition of high risk used in this study was tumors with the longest tumor diameter >10.0 cm, mitotic count >10 mitoses/50 hpf, tumor diameter >5.0 cm, and mitotic count over 5 per hpf or tumor rupture before surgery or at surgery. The study showed an improved 5-year recurrence free survival (65.6% vs. 47.9%; \(P < 0.001\)) as well as 5-year OS (92.0% vs. 81.7%; \(P = 0.02\)) with 3 years of IM as opposed to 1 year. Interestingly, the longer duration of IM benefitted exon 11 KIT mutants as opposed to no significant improvement for subsets of patients whose GIST harbored KIT exon 9 mutation or PDGFRA mutations, underlying the importance of mutation testing in GIST.\(^{[48]}\) Even in the subset of exon 11 KIT mutants, deletions that affected exon 11 codons 557 and/or 558 benefitted the most with 3 years of IM as opposed other subsets of exon 11 KIT mutants. While 3 years of adjuvant IM should be considered the current standard for resected IM, studies evaluating 5 years of adjuvant IM have been completed showed the feasibility of such an approach, with head-to-head comparison results still awaited.\(^{[44,49]}\)

There are two specific scenarios where the standard dosing of IM 400 mg as adjuvant needs a mutation-specific personalized approach. First, GISTs harboring the PDGFRA exon 18/D842V mutation (comprising 60%–70% of PDGFRA mutants) are considered relatively resistant to IM and current recommendations are for no adjuvant treatment for these mutants.\(^{[45,50,51]}\) These mutants are considered to be resistant to other TKIs such as sunitinib and regorafenib as well. Second, KIT exon 9 mutants have better outcomes with IM 800 mg/day dosing in the advanced/metastatic setting and it may be worthwhile considering an increased dose for these patients in the adjuvant setting as well, though there is no firm evidence for the same.\(^{[45,52]}\)

Neoadjuvant Imatinib in gastrointestinal stromal tumor

Although neoadjuvant IM has been used since 2003, data regarding the feasibility and efficacy have started emerging only recently. Indications for the use of neoadjuvant IM are yet to standardized, but common scenarios where a neoadjuvant approach with IM is considered are as follows:

• Distal anorectal sphincter complex GIST – for organ and/or sphincter preservation
• Bulky duodenal–pancreatic region GISTs to minimize the extent of multivisceral resections and avoid intraoperative tumor rupture
• Difficult initial location of tumors – gastroesophageal junction, rectum, duodenum
• Marginally resectable GIST to facilitate R0 resection.

The single largest published experience with the use of neoadjuvant IM comes from the pooled data of the European Organisation for Research and Treatment of Cancer (EORTC) Soft Tissue and Bone Sarcoma Group (STBSG) sarcoma group, which evaluated 161 patients with locally advanced nonmetastatic GISTs. Almost 83.2% of the patients underwent R0 resection with such an approach, with only two patients progressing on neoadjuvant IM. Five-year disease-free survival rates were 65%, with a median OS of 104 months. Patients had received IM for a period of 4–12 months prior to surgical evaluation in this study.\(^{[53]}\) The feasible and efficacious use of neoadjuvant IM has also been documented in the Indian setup with large retrospective studies from Tata Memorial Hospital.\(^{[13,54]}\)

Prospective single-arm studies have also validated the use of neoadjuvant IM. The Apollon phase II study prospectively evaluated 41 patients (locoregionally...
advanced, nonmetastatic) GIST with a median tumor size of 10.8 cm with 6 months of neoadjuvant IM. R0 resections were performed in 30 (n = 34) patients, with two patients having metastatic disease at resection. The PFS rate at 3 years was 85.2%, which was considered promising in view of the fact that patients were not planned for adjuvant IM.[55] A more recent single-arm phase II study in the Asian population evaluated 56 patients with high-risk gastric GIST and found response rates of 63% with an R0 resection of 91% using 6–9 months of IM. Although the median follow-up was only 32 months, 2-year OS and PFS rates were encouraging 98% and 89%, respectively.[56]

While the use of neoadjuvant IM has clinical implications and appeal in select cases of GIST, multiple questions remain to be answered.

**Treatment of advanced/metastatic gastrointestinal stromal tumor**

Patients with advanced/metastatic GIST have good median OS. These patients should have a KIT mutation testing done and if required, PDGFRA analysis should also be conducted while planning treatment.

The initial treatment for advanced/metastatic GIST is IM, with a dose of 400 mg/day recommended for KIT exon 11 mutants and 800 mg/day for KIT exon 9 mutants. This is based on a PFS benefit seen with 800 mg/day dosing in KIT exon 9 mutants.[57] It is prudent to consider more frequent clinical and radiological tumor assessments in the early course of treatment for advanced GIST, either with CT or PET-CT scans. The median time to response to IM is 3 months, though a slower rate of responses is also known. Although the Choi criteria is appealing, using response evaluation criteria in solid tumors also provides a fair estimation of response/progression.[58]

Treatment with IM should be considered until progression or there are tolerance issues even if a complete response is achieved or in the less common scenario of macroscopic resection of residual disease sites.[59] The median progression-free survival with first-line IM in older series is approximately 18–24 months, though this figure has increased to approximately 30–36 months in later reports, albeit with smaller sample sizes.[8,19,53,60,61] These differences can be explained by upfront treatment with imatinib in the current era (as opposed to the prior use of ineffective chemotherapy and radiotherapy) as well as reducing tumor size at diagnosis across time periods. Beyond median values, there is a cohort of patients who will remain progression free for 6–10 years on IM alone.[62]

An important aspect of the management of advanced/metastatic GIST is the requirement for continued treatment with IM as opposed to cessation post a specified time duration. The concept of IM cessation after 1 year and 3 years was studied in the BFR14 studies. The results showed that IM interruption after 1/3 years in patients responding to IM resulted in a high risk of rapid progression in patients with advanced GIST.[63,64]

Although treatment responses with IM are remarkable, approximately 80% of patients will progress after a median of 2–3 years due to the development of secondary KIT mutations.[65] Secondary mutations typically occur in the ATP-binding pocket of KIT encoded by exons 13 and 14, and the activation loop (A-loop) encoded by exons 17 and 18.[66] Although there is no robust evidence to suggest KIT mutation-based individualized treatment post progression with IM, an option of repeating a biopsy in such a setting to evaluate mutational status is gaining ground. There is some evidence to suggest that sunitinib has preferentially better action against secondary KIT exon 13/14 mutant GISTs, while regorafenib has superior action against exon 17 mutants.[67,68]

Options post progression with IM have emerged with trials confirming the efficacy of drugs such as sunitinib and regorafenib in this scenario. Sunitinib (an oral multikinase receptor tyrosine kinase inhibitor targeting KIT, PDGFRs, VEGFR-1, VEGFR-2, VEGFR-3, FLT3, and RET) has been shown to improve median time to progression as compared to placebo in the second-line setting (27.3 weeks vs. 6.4 weeks; P < 0.0001) in a randomized setting and is currently recommended for the same.[69] Careful monitoring of the side effects with sunitinib is required, considering the increased incidence of fatigue, diarrhea, skin discoloration, and nausea seen with sunitinib as opposed to the usually well-tolerated imatinib. In patients who have progressed on both imatinib and sunitinib, the recommended option for further treatment is regorafenib (oral multikinase inhibitor inhibiting VEGFR1-3, TEK, KIT, RET, RAF1, BRAF, BRAFV600E, PDGFR, and FGFR). The GRID study evaluated regorafenib versus placebo in a randomized control Phase III trial and showed an improvement in PFS which was statistically significant (4.8 months vs. 0.9 months; P < 0.0001). The most common regorafenib-related adverse events of grade 3 or higher seen in the study were hypertension, hand–foot skin reaction, and diarrhea.

While sequencing of treatment as mentioned above is preferable, options beyond this paradigm also need exploration. A commonly used treatment modality is increased dosing of IM from 400 mg to 800 mg per day. Long-term results of the EORTC-STBSG/AIGTG phase III trial have shown that 17.4% of patients who had progressed on 400 mg dosing and were administered dose escalated IM, 800 mg per day, remained progression free for >1 year. Whether such a benefit is limited to KIT exon 9 mutants or can be generalized across subgroups remains to be seen, but such a strategy can be used as an interim measure in resource-constrained settings.[70] Another strategy is the evaluation of metastasectomy or resection in patients with limited progression. While resection is generally considered in advanced disease in patients responding to
Radiation management
The role of radiotherapy in the management of GIST is limited, as GIST is traditionally considered a radioresistant tumor.[77] A few scenarios where radiotherapy may be attempted include as follows:[78]

- Radiotherapy of metastases for palliation of local symptoms
- Radiotherapy of focally progressing lesions, with the aim of overcoming emergent resistant clones
- Definitive radiotherapy alternative to surgery in localized GIST in elderly patients with comorbidities or in case of unresectable tumors.

Special Situations in Gastrointestinal Stromal Tumor
Syndromes associated with gastrointestinal stromal tumor
GIST is a predominantly sporadic disease, but since 1998, germ line mutations with familial predisposition to GIST have been identified.[79,80]

- Neurofibromatosis type 1 (von Recklinghausen’s disease)
- Carney–Stratakis Syndrome
- Carney’s triad

Available data suggest that GISTs associated with the above-mentioned hereditary syndromes are generally less sensitive to treatment with IM.[80]

Treatment of advanced gastrointestinal stromal tumor with platelet-derived growth factor receptor alpha mutations
Patients with PDGFRA mutations are seen in 5%–15% of localized GISTs and 1%–3% of advanced GISTs, likely because of the better prognosis of these GISTs when presenting with localized disease.[81,82] While patients with resected disease and harboring PDGFRA D842V mutations are recommended not to receive adjuvant IM, there are no such recommendations for patients with advanced disease.

Two large retrospective studies have been conducted solely focusing on the outcomes in PDGFRA-mutant tumors. The first study comprised data collated from investigators attached to members of the EORTC-STBSG, the French Sarcoma Group (GSF-GETO), the Italian Sarcoma Group, and the Spanish Sarcoma Groups (GEIS).[83] Eighty-eight patients were evaluated, of whom 32 patients (55%) had PDGFRA-D842V substitutions, whereas 17 (29%) had mutations affecting other codons of exon 18 and nine patients (16%) had mutations in other exons (exon 12 and exon 4). The study clearly showed that none of the 31 evaluable patients with D842V mutations had a response to IM, while responses were seen in the non-D842V subgroup. This also translated into markedly different PFS and OS between the two mutant cohorts. The second study comprising 71 patients from the Netherlands and United States suggested that a small cohort of patients with D842V mutations respond to IM, though such patients would need frequent monitoring if started on IM.[84]

Based on the available evidence, a majority of patients with PDGFRA non-D842V respond to and should be treated with IM, while for patients with D842V mutations, the appropriate line of management at this juncture is still to be elucidated. Treatment with second-line TKIs may be attempted. Avapritinib, a potent and selective kinase inhibitor with broad activity against oncogenic KIT/ PDGFRA mutants, including PDGFRA, has recently shown activity in pretreated unresectable PDGFRA D842V and KIT-mutated GISTs and is likely to become an option for treatment in this subset of patients.[85]

Stage-wise Prognosis with Current Management
The survival of patients with GIST has improved markedly over the past two decades as understanding of the disease biology has grown along with improvement in surgical techniques and the increasing benefits with imatinib and other TKIs. The expected survival for patients receiving multimodality management in the current era is detailed in Table 3.[86]

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Conflicts of interest
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