Imatinib as preoperative therapy in Chinese patients with recurrent or metastatic GISTs

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Objective: Imatinib has dramatically altered the options for management of patients with gastrointestinal stromal tumours. However, it has become clear that secondary resistance to the drug develops during long-term therapy. The purpose of our study was to retrospectively analyze safety and long-term outcomes in Chinese patients with recurrent or metastatic GISTs treated with imatinib preoperatively.

Methods: Between June 2003 and June 2011, 22 patients underwent surgery for recurrent or metastatic GISTs after preoperative treatment with imatinib.

Results: Complete resection was accomplished in 8 of the 10 responsive disease (RD) patients (80%), and in 3 of the 12 patients (25%) who had progression disease (PD). The amount of blood loss during the operation in PD patients was higher than in RD patients. There was 1 hospital death in PD group related to surgery, while the other patients recovered with conservative therapy because complications were mild. The difference in median PFS between patients with RD and those with PD was significant (24.8 vs. 2.81 months, P<0.001). The difference in 2-year OS rate between patients with RD and those with PD was not significant (100% vs. 87.5%, P>0.05).

Conclusions: Our study indicates that surgical intervention can improve the PFS of Chinese patients with recurrent or metastatic GISTs responsive to imatinib, but does not prolong OS as well as in patients who develop imatinib resistance. Surgical resection following imatinib treatment is feasible and can be considered for patients with advanced GISTs responsive to imatinib.

Key Words: Gastrointestinal stromal tumor; imatinib; targeted therapy; surgical intervention; pathology


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Introduction

Gastrointestinal stromal tumors (GISTs) are the most common mesenchymal tumors of the gastrointestinal system and are thought to originate from intestinal cells of Caja (1). Before the specific tyrosine kinase inhibitor imatinib was investigated for GISTs treatment in 2000, GISTs proved refractory to any treatment other than surgery, including conventional chemotherapy and radiation therapy, making observation alone the standard of care after surgical resection (2,3). Imatinib is a success story of molecular biology that has dramatically altered the management of patients with gastrointestinal stromal tumours. However, as patient follow up has continued, secondary resistance has appeared because responses to imatinib are of limited duration. Indeed, the median time to progression during imatinib mesylate treatment is 2 years (4). The current clinical practice at The University of Texas M.D. Anderson Cancer Center for patients with either locally advanced, recurrent, or metastatic GISTs is to use imatinib as first-line treatment and then to consider surgical resection when the tumor has demonstrated adequate response to allow for
complete resection of any residual disease (5). However, the indications, timing, and role of surgery for these patients are not well established. The effect of preoperative imatinib on surgical resection rates and postoperative outcome in patients with advanced GISTs is unknown. In the present study, we aimed to define the role of surgery and its long-term outcome in patients with advanced GISTs treated with imatinib preoperatively.

**Patients and methods**

**Patient characteristics**

We reviewed the medical records of 113 patients with unresectable recurrent or metastatic GISTs who had been treated with imatinib at Fudan University Cancer Center, Shanghai, China, between June 2003 and June 2011. Twenty-two patients who underwent surgery after imatinib treatment were enrolled into the study.

**Imatinib treatment**

Front-line imatinib treatment consisted of 400 mg once daily. Patients who experienced disease progression before surgery were treated with 600 or 800 mg imatinib daily. Tumor response to imatinib was evaluated in all patients with serial CT imaging. According to the Response Evaluation Criteria in Solid Tumors criteria (6), an experienced radiologist retrospectively re-reviewed all radiographic data for the 22 patients in our study and categorized the response to imatinib based on changes in tumor size, degree and extent of enhancement, and the presence or absence of solid nodules within the tumor. Consistent with Southwest Oncology Group (SWOG) criteria, an increase in the sum of the longest diameters of the target lesions alone was not regarded as disease progression if accompanied by definite cystic change in the tumor suggesting necrosis (7). A resistant lesion was confirmed by radiographic imaging, such as computed tomography (CT), magnetic resonance imaging (MRI). Radiographic evaluations were carried out every 3 months during imatinib treatment.

Disease status at the time of surgery was classified into two clinical categories. The first category included patients who achieved partial response (PR), stable disease (SD) or complete response (CR) to imatinib. These patients were classified as having responsive disease (RD). The second category included patients who experienced secondary resistance and primary resistance to imatinib during the treatment. Primary resistance was defined as progression occurring in patients who had never shown any response to imatinib or progression encountered within 3 months after initiation of imatinib treatment. Secondary resistance was defined as progression occurring in patients who initially had a response to imatinib treatment or a progression free survival interval exceeding 3 months (8). These patients were classified as having progression disease (PD).

**Surgical intervention**

The standard approach is to plan resection at the point of maximal response when the disease has stabilized but not begun to progress. In patients with RD, a reasonable time to consider surgery is at 12 months after starting imatinib treatment (4,9). All surgical procedures in this study were elective. In RD patients with no evidence of change in tumor size or enhancement throughout treatment for at least two months, the resection was considered by the surgeon based on a review of computed tomography (CT) imaging by experienced radiologists. However, in PD patients, surgery was done as soon as progression occurred.

**Mutation analysis**

Eight pretreatment formalin-fixed and paraffin-embedded surgical specimens from primary tumors were available in our institution, but most specimens were from other medical centers. Tumor specimens from the second resections done after imatinib therapy were all from our center. The fresh samples were flash frozen in liquid nitrogen at the time of surgical resection and maintained at −80 °C. Genomic DNA was extracted using a standard phenol-chloroform organic extraction protocol. Adequate DNA for mutation analysis was obtained, which was then analysed for mutation of KIT exon 9, 11, 13, 17, and PDGFRA exon 12 and 18 as described previously in the literature (10).

**Statistical analysis**

Progression-free survival (PFS) was measured from the date of surgery to the date of documented disease progression or death from any cause. Overall survival (OS) was measured from the date of surgery to death from any cause. Survival curves were calculated according to the Kaplan-Meier method. We used the Statistical Package for the Social Sciences, version 13.0 (SPSS, Chicago, IL). P<0.05 was
considered statistically significant.

**Results**

**Patient characteristics**

A total of 22 patients (13 men and 9 women) underwent surgery for GIST after preoperative treatment with imatinib. The median age of the patients was 49.3 years (range from 33 to 72 years) at surgery after imatinib treatment. All patients received operations in our institution, including 4 patients who experienced more than 2 excisions. Primary sites of GIST were the stomach in 7 patients (31.8%), small intestine in 7 patients (31.8%), colon and rectum in 4 patients (18.2%), extra-gastrointestinal in 4 patients (18.2%). At the time of surgery, 15 patients had locally recurrent GISTs, and 7 patients had metastatic GISTs, including 3 tumors which metastasized to the liver, 2 tumors which metastasized to the peritoneum, and 2 tumors which metastasized to extragastrointestinal sites. At the first surgery in 2 patients, the surgeon found that the whole abdominal cavity and pelvic cavity were covered with hundreds of the GISTs (Table 1).

**Preoperative imatinib treatment**

Prior to surgery, the median duration of imatinib treatment for all patients was 14 months (range, 2-51 months), 18.5 months (range, 5-51 months) for patients with RD, and 8 months (range, 2-30 months) for patients with PD. Except for one patient who showed primary resistance during treatment, 21 of 22 patients had an initial response to imatinib. Of the 22 patients treated, 1 (4.5%) patient had a complete response, 4 (18.2%) patients had a partial response, 5 (22.7%) had stable disease, and 12 (54.6%) had progressive disease (Table 2). Of the 12 patients with progressive disease, one exhibited primary resistance to imatinib, while 11 developed secondary resistance to imatinib. In RD group, treatment was interrupted in 4 patients after a range of 2 to 12 months, mostly for financial reasons or drug adverse effects. After imatinib treatment was resumed in these 4 patients, all patients were still sensitive to imatinib and disease control continued. Prior to surgical resection, 8 (36.4%) patients were administered 400 mg daily, 7 (31.8%) patients, including 3 RD cases and 4 with secondary resistance, were escalated to 600 mg a day, and 7 (31.8%) patients were escalated to 800 mg per day due to different tolerance. After imatinib treatment was resumed in these 4 patients, all patients were still sensitive to imatinib and disease control continued. Prior to surgical resection, 8 (36.4%) patients were administered 400 mg daily, 7 (31.8%) patients, including 3 RD cases and 4 with secondary resistance, were escalated to 600 mg a day, and 7 (31.8%) patients were escalated to 800 mg per day due to different tolerance. In the 14 patients who were administered higher doses, 10 (71.4%) patients who had shown secondary resistance continued to experience disease progression and 4 (28.6%) patients had partial remission, which classified them as RD patients. Of the 11 patients with PD, except one hospital death and three who continued imatinib treatment due to experiencing local disease progression, imatinib therapy was discontinued and postsurgical treatment was with Chinese medicine or sunitinib.

**Surgical intervention**

Complete resection was accomplished in 8 of the 10 RD patients (80%). Three of the 12 PD patients (25%), including 1 patient who showed primary resistance to imatinib, achieved complete resection. The other 9 (75%) PD patients had incomplete resections, including 2 patients who underwent palliative resection because of hundreds of
tumors covering the entire abdominal cavity. The amount of blood loss during the operation was below 1,000 mL in 6 of 10 RD patients, and none lost over 5,000 mL. On the contrary, in the PD patients, except for 3 (25%) patients who lost less than 1,000 mL, 3 (25%) patients lost more than 5,000 mL, and 1 patient lost over 10,000 mL. There was one hospital death related to surgery, and 2 patients experienced postoperative complications. One patient died 2 days after operation due to disseminated intravascular coagulation. One PD patient lost over 5,000 mL of blood during surgery, was in shock during the operation, and recovered after extensive effort treatment. One patient experienced postoperative acute renal failure, but recovered with conservative therapy because the complication was comparatively mild.

**KIT/PDGFRA genotyping before and after imatinib treatment**

For all primary tumors, activating mutations in KIT were seen in 17 (77.3%) patients, including 12 mutations in exon 11, 4 mutations in exon 9 and 1 mutation in exon 13. PDGFRA mutations were seen in 4 (18.2%) patients, while KIT and PDGFRA wild type gene was found in the one (4.5%) patient who showed primary imatinib resistance. After surgery following imatinib treatment, mutation analysis was performed. In addition to the original mutations, 3 of 12 patients with secondary imatinib resistance harbored secondary mutations in KIT exon 17 in the progressive lesion, while the other lesions showed only the original mutations (Table 3).

**Progression-free survival and overall survival analysis**

Except for one patients who died while in hospital, all other patients were alive at last followup. After a median follow-up of 53 months after surgery (range, 8-84 months) for the surviving patients, PFS for RD and PD patients was 24.8 and 2.81 months, respectively. The difference in PFS between patients with RD and those with PD was significant (P<0.001). All PD patients experienced disease progression within 11 months after surgery. To evaluate the role of surgery in the setting of recurrent or metastatic GISTs, we reanalyzed the data of 3 patients with recurrent or metastatic GISTs who had RD during imatinib treatment. The median PFS for these 3 patients was 20 months. The difference in PFS between patients with recurrent or metastatic GISTs exhibiting RD and patients with PD was still significant (Figure 1A, P<0.001). In the 21 patients who were still alive, 9 patients remain alive with unresectable GISTs, while the other 13 patients are disease-free. The OS for the 10 patients exhibiting RD on imatinib and for the 11 patients who experienced PD has not yet reached the median. For RD patients, the estimated OS was 100% at 2 years, especially one patient was alive for more than 7 years. For PD patients, OS was 87.5% at 2 years. The difference in OS between RD and PD patients was not

| Table 2 Clinical data of preoperative imatinib treatment for GISTs |
|-------------------|--------------------------|------------------|
| **Disease status at surgery** | **RD (n=10)** | **PD (n=12)** |
| **Duration of Imatinib (months)** | 18.5 | 8 |
| **Range** | 5-84 | 2-30 |
| **Dose of Imatinib** | 400 mg | 6 |
| **Range** | 400 mg | 2-84 |
| **Dose increment of Imatinib** | 600 mg | 3 |
| **Range** | 600 mg | 7 |
| **Response** | 800 mg | 1 |
| **Complete response** | 1 | 0 |
| **Partial response** | 4 | 0 |
| **Stable disease** | 5 | 0 |
| **Progressive disease** | 12 | 12 |
| **Primary resistance** | 0 | 1 |
| **Secondary resistance** | 0 | 11 |
| **Imatinib continue** | 6 | 0 |
| **Imatinib interrupt** | 9 | 3 |

| Table 3 Mutation Analysis of KIT and PDGFRA in Patients before and after imatinib treatment |
|-------------------|--------------------------|------------------|
| **Disease status at surgery** | **RD (n=10)** | **PD (n=12)** |
| **Before imatinib treatment** | **KIT exon 9 mutation dupc502+c503** | 1 |
| **KIT exon 11 mutation V559D** | 4 |
| **KIT exon 11 mutation del552-553** | 3 |
| **KIT exon 11 mutation del553-558** | 1 |
| **KIT exon 13 mutation K642E** | 0 |
| **PDGFRA exon18 mutation D842V** | 1 |
| **Wild type** | 0 |
| **After imatinib treatment** | **Original mutation** | 10 |
| **Secondary mutation KIT exon 17 Y823D** | 9 | 3 |
significant \((\text{Figure 1B, } P>0.05)\).

**Discussion**

Imatinib has shown considerable clinical efficacy in treating advanced GISTs, and has been shown to prolong OS of patients with recurrent or metastatic disease. Preoperative imatinib therapy for primary or recurrent unresectable GISTs commonly results in tumor downstaging and enables curative surgical resection (11), or allows large tumors to be resected with less blood loss. SYM and his group found that earlier surgical intervention in patients with advanced, imatinib-responsive GIST may be optimal in preventing tumor progression (12). In our investigation, blood loss during surgery and hospital deaths for PD patients were much higher than for RD patients. 75% of PD patients lost more than 5,000 mL of blood, while 3 (25%) PD patients lost over 10,000 mL. One PD patient died of disseminated intravascular coagulation 2 days after surgery, due to much blood loss. Another PD patient went into shock during the operation because of blood loss exceeding 8,000 mL, only to recover after doctors’ effective treatment. In our study, 10 patients developed disease progression even after dose escalation of imatinib. Of these patients, the tumor in one patient grew so quickly that it occupied the entire pelvis \((\text{Figure 2A,B})\), which induced dysuresia and dyschezia. In another patient who show secondary imatinib resistance, after escalating the dose from 400 to 600 mg
once daily; tumor necrosis occurred which led to high body temperature for several days before surgery. Since sunitinib at that time had not been approved in China, surgical intervention was the only treatment modality available, and both these patients underwent palliative surgery to reduce tumor size (Figure 2C,D). Over several months of follow up, we found that these palliative operations improved the two patients’ quality of life greatly. Our observation suggests that careful patient selection is required for the performance of surgery in patients with recurrent or multiple metastatic disease after imatinib therapy, because the probability of vascular friability in these patients is greatly elevated, leading to greater blood loss during surgery. Palliative nonresectional surgery should be considered only for specific symptoms just like difficult defecation due to tumor oppression or palliative care, either by focal debulking of symptomatic deposits or using bypass procedures.

In the present study, imatinib treatment was stopped one week before surgical procedures in accordance with the standard of care. In 6 of 10 RD patients, imatinib was reintroduced at the preoperative dose as soon as solid food was started. The median time of imatinib recommencement was 9 days (range, 2-20 days) after surgery. We found that postoperative complications were likely unrelated to imatinib treatment. Some series have shown that continuing imatinib treatment until 1 day before surgery does not increase postoperative complications and that imatinib can be safely restarted within 2 weeks after surgery (13). Four of 22 patients interrupted imatinib treatment for 2-12 months, mostly for financial reasons or drug adverse effects. Imatinib treatment was resumed after these patients developed disease progression, and all 4 patients were still sensitive to imatinib when treatment was resumed. These findings are in agreement with a Korean study, which found that in patients with advanced GISTs controlled with imatinib, imatinib interruption resulted in a high risk of PD within 1 year. However, the majority of disease was controlled with imatinib resumption after PD, and there were no statistically significant differences in PFS and OS between patients who stopped and restarted imatinib and patients who continued treatment.

Figure 3 Comparison of pre- and post-imatinib CT scan and tumour tissue in a recurrent GIST undergoing preoperative imatinib therapy. A. CT scan before imatinib therapy revealed a tumour located between the residual stomach and liver, adhering to the inferior caval vein, about 10 cm in diameter; B. CT scan after 6 months of preoperative imatinib therapy demonstrates a 6 cm in diameter sized residual tumour; C. gross specimen after imatinib therapy demonstrating necrosis nodule and bleeding in the tumor; D. biopsy specimen from the first resection of gastric GIST, before preoperative imatinib treatment showing been composed of affluent spindle cells, H&E 200; E. representative section of the tumour mass after imatinib therapy showing much collagen with hyaline degeneration, H&E 400; F. tumour cells before and after imatinib therapy show the same (C1676 T → A) was detected in exon 11 of the KIT gene.
who got the drug continuously (14).

All patients received evaluation for response to imatinib by CT scan based on changes in tumor size, degree, and extent of enhancement. Although imatinib therapy has dramatically improved the prognoses of patients with metastatic GISTs (15), a complete pathologic remission is rarely achieved (16). In the present study, after 6 months of imatinib treatment after recurrence, one patient with a primary gastric GIST which was radically resected but recurred in the retroperitoneum three years later achieved a complete pathologic response. Given the fact that the diameter of the recurrent nodule decreased from 10 to 6 cm, the patient was evaluated as a PR by CT scan before surgery. However, the excision sample showed no viable tumor cells and the case was finally confirmed as a CR by pathology examination (Figure 3), which suggests that a CT scan is not a definitive tool in the evaluation of GIST during imatinib therapy. This interesting case was published elsewhere.

PET is a sensitive and specific method with which to evaluate tumor response on the basis of changes in tumor metabolism (17), which can demonstrate response before CT and may indicate response or progression when the CT findings are equivocal. However, functional PET scans also may become positive again within weeks of imatinib being stopped. The role for PET in assessing the likely response of patients included in neo-adjuvant trials needs to be further investigated (18).

In our cases, GISTs with mutations of the KIT exon 11 showed a better response compared to GISTs with other mutation such as KIT exon 9, PDGFRA mutation D842V, and wild type tumors; which suggests that mutations status correlates with the grade of initial imatinib response, and that a pre-treatment mutation analysis of the biopsy specimen may facilitate the treatment decision on preoperative imatinib treatment. Resistance to imatinib after initial response, mediated by expansion of therapy resistant and potentially pre-therapeutic clones with secondary mutations has also been recognized in metastatic GISTs (19). In our study, secondary mutation of KIT exon 17 Y823D occurred in 3 of 12 PD patients who showed an initial response to therapy, which indicates that additional mutations occurring in codon 823 of KIT exon 17 may provide a mechanism for secondary imatinib resistance.

In our series, surgical intervention for patients with metastatic and unresectable GISTs who were responsive to imatinib therapy resulted in prolonged PFS. But, for patients with advanced GISTs who had disease progression after achieving SD for some period, (classified as PD patients in our study) the effect of surgical interventions was limited. Although the difference in OS between patients exhibiting RD and those with PD was not significant, the findings were different those reported by SYM et al. (12), in which OS was significantly higher in patients with RD than in those with PD. Shorter follow-up time may account for our different results. Based on our findings, complete surgical resection was achieved in 8 of 10 patients with recurrent or metastatic GISTs exhibiting response to imatinib, and we believe that the quality of life for RD patients was better than for PD patients. Moreover, these patients showed prolonged progression free survival after surgery with mostly mild surgical morbidity. Our study, along with others (20,21), suggests that surgical intervention for GISTs responsive to imatinib may be an important measure of the management strategy. Surgery can be considered a relative indication only for advanced patients in whom a few nodules show resistance to imatinib while most other lesions have not developed resistance and continue to respond, or for specific symptoms, such as gastric outlet obstruction or tumor pressure on the bladder or rectum.

In summary, we suggest that surgical resection following imatinib treatment improves progression free survival for Chinese patients with recurrent or metastatic GISTs responsive to imatinib, but does not prolong overall survival as well as in patients who develop imatinib resistance because of short follow-up time. Surgical resection following imatinib is feasible and can be considered for patients with advanced GISTs responsive to imatinib. In the near future, our findings should be further investigated by more randomized clinical trials.

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