A Case of Hereditary Gastrointestinal Stromal Tumor (KIT Mutation p.Asp820Tyr) in a Portuguese Family and a Good Response to Imatinib – An Update

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ABSTRACT

Gastrointestinal stromal tumors (GIST) are the most common mesenchymal tumors of the human gastrointestinal tract. They are derived from transformed neoplastic precursors of the interstitial cells of Cajal. They account for 0.1% to 3% of all gastrointestinal cancers. Up to 20% of cancers of the small bowel are GISTs. Less than 5% of cases are associated with hereditary predisposition, like Neurofibromatosis type I, Carney syndrome, and Familial GIST syndrome. The study objectives were: to update the response to imatinib of the family of our institution and to review the families cases published in the literature with this syndrome. Review of the cases of Familial GIST syndrome published in literature and update of the only Portuguese family by consultation of clinical processes.

In the literature, we found 35 cases of unrelated families with this syndrome. This report is also an update on the only Portuguese family with this syndrome and a good response to imatinib. The role of imatinib was not established in cases of Familial GIST syndrome. In our opinion, it seems wise to use 400 mg/day for an indefinite period. The objective is disease control and hindering the development of additional lesions.

Key words: germline GIST gene mutation, gastrointestinal cancer, surgery, imatinib, stroma tumours

INTRODUCTION

Gastrointestinal stromal tumors (GIST) were first recognized as a distinct clinicopathologic syndrome in 1983 by Mazur and Clark (1) and may be defined morphologically as spindle cell, epithelioid, or occasionally pleomorphic mesenchymal tumors of the gastrointestinal tract that usually express the KIT protein (2). They are the most common mesenchymal tumors of the human...
gastro-intestinal tract and are derived from transformed neoplastic precursors of the interstitial cells of Cajal (1). They develop mainly within gastric and small bowel muscular layers (3-4).

GISTs only account for 0.1% to 3% of all gastrointestinal cancers, but up they comprise 20% of cancers of the small bowel. They are found most frequently in the stomach (60%), duodenum and small bowel (35%), and less frequently in colon, rectum, esophagus, omentum and mesentery (<5%). They occur mostly in the fifth to seventh decades of life and affect equally men and women (4).

This entity has a wide spectrum of clinical behaviors, spanning from indolent and curable disorders to highly malignant disease that metastasize and become lethal (1).

Annual incidence is between 10 to 20 cases per million (2,4), and prevalence is 129 per million per year (4). No specific epidemiological risk factors for GIST have been described (5). More than 95% of GISTs are sporadic and less than 5% are associated with hereditary predisposition (3).

Approximately 95% of sporadic GISTs have over-expression of the KIT protein (CD117), a trans-membrane tyrosine kinase receptor. Activating somatic mutations of the proto-oncogenes KIT or platelet-derived growth factor receptor alpha (PDGFRA) are mutually exclusive. These mutations occur in 85–90% of the cases, and are considered the primary genetic events in GIST pathogenesis. These genes code proteins with similar structure and function. KIT somatic mutations are present in 80–85% of the cases and occur predominantly in the juxtamembrane domain (exon 11; 60–70%), followed by the extracellular domain (exon 9; 10%), or, rarely, in the Kinase I and II domains (exons 13 and 17, respectively; ~2%). Mutations in PDGFRA are detected in 5–10% of the sporadic cases and involve exon 12 (homologous to KIT exon 11; 1%), exon 14 (homologous to KIT exon 13; <1%), or exon 18 (homologous to KIT exon 17; 6%) (3,6). Both of genes (KIT and PDGFRA) are located within human chromosome 4q12 (4).

TNM classification has several limitations in GIST staging. Therefore its use is not recommended in this disease.

The standard approach to GIST tumors ≥ 2 cm in size is biopsy/excision, because they are associated with a higher risk of aggressive behavior. If there is an abdominal nodule not amenable to endoscopic assessment, laparoscopic/laparotomic excision is the standard approach (7).

Prognosis depends on mitotic rate, tumor size and location (gastric GISTs have a better prognosis than small bowel or rectal GISTs) (7).

**CASE REPORT**

Update of the only portuguese family with Familiar GIST syndrome, be consultation of clinical processes; and a comprehensive literature review was carried out, searching for of the cases of Familial GIST syndrome published in literature.

We present a case of a Caucasian 48 year-old woman, school worker, complaining of bloating, sporadic abdominal pain and heartburn for one year. She had a past medical history of depression and had suffered an unilateral oophorectomy years before (ovarian benign cyst).

There was a history of cancer in her family: Her mother had been operated 18 months before for metastatic GIST (segmental bowel resection and exciscional biopsy of gastric exophytic tumor). The examination of the resected specimen revealed a mutation in 17 exon of c-kit gene [c.2458G>T (p.Asp820Tyr)], in more than 20% of tumor cells), and she had been started on Imatinib.

Due to the discovery of this mutation, our patient and some members of her family were submitted to genetic testing by another team, in another Hospital, and they published a paper about the mutation (3). However, the fraction of family we follow had not been included in this study (fig. 1). Our patient did carry this mutation.

We have found only 35 reports of unrelated families in world, with this syndrome (see table 1). In Portugal there is only one such family, living the Oporto area (3).

When we first saw our patient, she had already been submitted to upper GI endoscopy and computerized tomography (CT) scan of the abdomen. Endoscopy revealed two polypoid tumors (of submucosa), in the gastric fundus, 10 mm each. The biopsy of those lesions did not reveal malignancy.

CT scan (fig. 2) revealed two polypoid tumors in the gastric fundus (about 14 mm) and cyst inclusion on the site of a previous oophorectomy.

To complete the study, an endoscopic ultrasound was performed, which revealed two subepithelial lesions, both in the dependence of the muscularis mucosa, with 18 and 13 mm largest diameter, in the anterior wall of the body and proximal small curvature of the gastric cardia.

After discussing with the patient, an atypical gastrectomy was decided, to remove the lesions. During laparotomy, some millimetric lesions were
Vítor Devezas et al found all over the gastric wall (fig. 3), and we decided to resect just the two major lesions.

After surgery, GIST was confirmed in the two lesions removed. After a consultation group with other specialties, we decided to proceed to imatinib treatment and imaging surveillance.

After 24 months with Imatinib and no evidence of disease progression, we performed a second-look surgery with total gastrectomy. Pathological examination of the stomach found no residual tumor, confirming the effectiveness of imatinib therapy. Since this was a case of familial GIST syndrome, our group decided to keep the treatment with Imatinib going.

![Figure 1 – Family tree. The arrow identifies our patient. The individuals fill the black indicate achievement by GIST disease. The individuals with a plus (+) was submit to genetic study and was positive for mutation of c-kit (c.2458G>T (p.Asp820Tyr)). Individuals with a minus (-) was negative in genetic study for mutation. The squares indicate the men and the circles the women.](image1)

![Figure 2 – Images of CT scan with pediculated tumor in gastric fundus (arrow)](image2)
### Table 1 – Families in world with Hereditary GIST syndrome with mutation, country, organs affected and accompanying signs and symptoms

| Gene | Exon | Mutation     | Country | Organs affected | Pigmentation anomalies | Dysphagia | Urticaria pigmentosa | Mastocytosis | Reference |
|------|------|--------------|---------|----------------|------------------------|---------|---------------------|-------------|-----------|
| KIT  | Exon 8 | p.Asp419del | German | Small intestine | No | Yes | No | Yes | [10] |
| KIT  | Exon 11 | p.Tyr553Cys | Japan | Small intestine/Stomach/Colon | No | No | No | No | [13] |
| KIT  | Exon 11 | p.W557R | Italy | Small intestine | Yes | Yes | No | No | [14] |
| KIT  | Exon 11 | p.W557R | Canada | Small intestine | Yes | No | No | No | [15] |
| KIT  | Exon 11 | p.V559A | Argentine | Small intestine/Stomach | Yes | Yes | No | No | [4] |
| KIT  | Exon 11 | p.V559A | USA | Small intestine | Yes | No | Yes | No | [1] |
| KIT  | Exon 11 | p.V559A | Korea | Small intestine | No | No | No | No | [17] |
| KIT  | Exon 11 | p.V559A | Italy | Small intestine/Stomach | Yes | No | Yes | Yes | [18] |
| KIT  | Exon 11 | p.V559A | Japan | Small intestine/Stomach | Yes | No | No | No | [19] |
| KIT  | Exon 11 | p.V559del | Japan | nd | Yes | No | No | No | [9] |
| KIT  | Exon 11 | p.V560G | Korea | Small intestine | Yes | No | No | No | [20] |
| KIT  | Exon 11 | p.V560del | Japan | Small intestine/Stomach | No | No | No | No | [8] |
| KIT  | Exon 13 | p.575s_577delinsA | Belgium | Rectum | No | No | No | No | [21] |
| KIT  | Exon 11 | p.L576_P577insG | Spain | Small intestine/Stomach | Yes | No | No | No | [22] |
| KIT  | Exon 11 | p.L576P | Germany | Small intestine | Yes | Yes | No | No | [23] |
| KIT  | Exon 11 | p.d579del | USA | Small intestine | No | No | No | No | [24] |
| KIT  | Exon 11 | p.d579del | USA | Stomach | No | No | No | No | [25] |
| KIT  | Exon 11 | p.d579del | USA | Small intestine/Stomach | Yes | No | No | No | [26] |
| KIT  | Exon 11 | p.d579del | USA | Small intestine/Stomach | No | No | No | No | [27] |
| KIT  | Exon 11 | p.d579del | USA | Small intestine/Stomach | No | No | No | No | [27] |
| KIT  | Exon 13 | p.K642E | France | Small intestine | No | No | No | No | [28] |
| KIT  | Exon 13 | p.K642E | France | nd | Yes | Yes | No | No | [29] |
| KIT  | Exon 13 | p.K642E | France | nd | Yes | Yes | No | No | [29] |
| KIT  | Exon 13 | p.K642E | France | nd | Yes | Yes | No | No | [29] |
| KIT  | Exon 13 | p.K642E | UK | Esophagus/Stomach/Small intestine/Rectum | No | Yes | No | No | [30] |
| KIT  | Exon 13 | p.K642E | Spain | Stomach/Small intestine/Rectum | No | No | No | No | [31] |
| KIT  | Exon 13 | p.K642T | Japan | Small intestine/Stomach | No | Yes | No | No | [32] |
| KIT  | Exon 17 | p.D820Y | Japan | Small intestine/Stomach | No | Yes | No | No | [15-16] |
| KIT  | Exon 17 | p.D820Y | Irlanda | Small intestine/Stomach | No | Yes | No | No | [33] |
| KIT  | Exon 17 | p.D820Y | Portugal | Stomach/Small intestine/Rectum | No | No | No | No | [3] |
| KIT  | Exon 17 | p.D822Y | Germany | Small intestine/Stomach | nd | nd | nd | nd | [34] |
| PDGFRA | Exon 12 | p.Y555C | Belgium | Small intestine | No | No | No | No | [35] |
| PDGFRA | Exon 12 | p.V561D | USA | Small intestine/Stomach | No | No | No | No | [36] |
| PDGFRA | Exon 14 | p.P653L | Italy | Stomach | No | No | No | No | [37] |
| PDGFRA | Exon 18 | p.D846Y | France | Stomach | No | No | No | No | [38] |
DISCUSSION

Hereditary predisposition to GIST is related with Neurofibromatosis Type I, Carney syndrome, and Familial GIST syndrome. The latter is a rare autosomal dominant genetic disorder originated by germline gain-of-function mutations of KIT or PDGFRA (3,8). It was described for the first time in 1998 by Nishida and his group (5), with identification the mutation in exon 11 of KIT (9).

Most often, patients affected with Familial GIST develop multiple tumors at an early age, in contrast with sporadic cases that are characterized most often by solitary tumors that appear at a latter age. Other less common clinical features reported in patients with Familial GIST syndrome are skin hyperpigmentation, dysphagia, urticaria pigmentosa, and mastocytosis (3,7,10-12). None of these were to be found in this family.

Among patients with familial GIST syndrome, the average age of index cases was 47 years (earlier than in sporadic cases). There is a documented case of a 15 year-old (11). There is no predominance of sex. Worldwide, there are four families with germline mutation on PDGFRA gene and the remaining 31 families with KIT gene mutations (one in exon 9, 19 in exon 13, seven in exon 13 and four in exon 17).

This Familial GIST syndrome had been studied and reported years before, but anyway this part of the family had not been submitted to genetic testing or put under surveillance. In fact, our patient and her mother had multiple lesions when they were submitted to surgery.

In our patient, although preoperative imaging suggested that the disease was locally contained, during laparotomy it was evident that there were multiple/metastatic GIST. Latter, imatinib treatment warranted a good response with complete remission in stomach.

In general, surgery is the standard in cases of non metastatic GISTs. The tumor must be removed en bloc with its pseudocapsule, to yield an adequate resection margin. Therefore, the goal of surgery is to accomplish an R0 excision (2,7). If R0 surgery is not feasible, or if it only can be achieved at the expense of great mutilation or loss of function, pre-treatment with tyrosine kinase inhibitors (TKI), like imatinib followed by sparing surgery (cytoreduction), is standard. Optimal timing for surgery after maximal tumor response, is 6–12 months (7).

Adjuvant therapy with imatinib 400mg/day for three years, is the standard treatment for patients with a significant risk of relapse. Adjuvant therapy should not be considered when the risk is low (7).

In patients with metastatic disease, TKI are standard treatment (2). Imatinib has at least a 10-fold more active favorable response than any other agent ever considered for treatment of GISTs (39). Treatment with TKI should be continued indefinitely, since treatment interruption is generally followed by relatively rapid tumor progression, even when lesions have been previously surgically excised (7).

In the case of confirmed progression, a dose escalation up to 800 mg/day and/or switch to a standard second-line treatment, another tyrosine kinase inhibitor – sunitinib – must be considered (7).

There is no data regarding the role of imatinib in preventing the development of GIST in Familial GIST. In some of these families, imatinib has an important role in maintaining the disease under control. We
believe it can be used similar to nonhereditary GIST.

Typically, there are mutations which display an especially strong response to imatinib, like mutations of exon 11 of KIT as p.Asp579del mutation, compared to mutations of exon 9 of KIT.

Ávila et al. (2014) treated all gastrointestinal tumor-affected family members (p.V559A) with imatinib, which has caused a progressive reduction of the melanosis and of the tumor size (4). On the other hand, Bamba et al. (2015) (8) started imatinib if tumor progression is faster than expected, with half-dose of imatinib (200mg/day) and the size of all GISTs markedly was reduced and maintained stable for one year (p.V560del).

Graham and contributors (2007) maintained their patient (p.K642E) on imatinib treatment at 300 mg/day (started at 400mg/day, but was reduced to 300mg/day because of intolerance) and disease remained stable after 19 months of treatment (until the publication of this article) (30).

Wozniak and contributors (2008), treated their patient (p.Q575_P577delinsH) with 400mg/day of imatinib and achieved either long-term partial response or stable disease after 58 months on treatment (21).

Regarding the family under study, Veiga et al. (2010) achieved partial response to imatinib and concluded that p.D820Y mutation was responsible for the weaker effect (3). We also achieved a good response in our patient (with complete remission of gastric lesions after 18 months of treatment and no relapse until now), with 400 mg/day of Imatinib. This result might be explained by different cytogenetic progression pathways described in this family (3).

**CONCLUSION**

In conclusion, this case is about a rare condition in the world and the only affected family in Portugal. Furthermore this mutation is uncommon in Familial GIST syndrome (only one of the four documented cases of exon 17 mutation of KIT gene, in a total of 35 families).

All patients carrying a germline mutation of KIT or PDGFRA need to start endoscopic screening at a young age, since the condition may develop early in life (a case in a 15 year-old patient has been reported).

TKI, like imatinib, are an important therapeutic weapon in GIST cases. The role of this drug was not established in cases of familial GIST syndrome but, in our opinion, it seems wise to use it with a dosage of 400 mg/day continued indefinitely, since mutation was present in all of cells of patient. Imatinib may serve to control disease and may prevent the development of additional lesions.

**Ethical approval**

The terms of the Helsinki Agreement have been respected.

**Author contribution**

All of author had contributed to this article. All authors approved the final manuscript.

**Conflicts of interest and source of funding**

Authors have no conflict of interest to declare.

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