Molecular and functional characterization of a new 3’ end KIT juxtamembrane deletion in a duodenal GIST treated with neoadjuvant Imatinib

Vittorio Perfetti1,*, Erik Laurini2,*, Suzana Aulić2, Maurizio Fermeglia2, Roberta Riboni3, Marco Lucioni3, Elena Dallera3, Sara Delfanti4, Luigi Pugliese5, Francesco Saverio Latteri6, Andrea Pietrabissa5 and Sabrina Pricl2

1Internal Medicine, Ospedale SS Annunziata-ASST Pavia and Department of Molecular Medicine University of Pavia, 27100 Pavia, Italy
2Molecular Simulation Engineering (MOSE) Laboratory, Pharmaceutical and Molecular Biology Division, DEA, University of Trieste, 34127 Trieste, Italy
3Department of Molecular Medicine and Anatomic Pathology Section, Fondazione IRCCS Policlinico San Matteo, 27100 Pavia, Italy
4Department of Oncology and Hematology, Fondazione IRCCS Policlinico San Matteo, 27100 Pavia, Italy
5Department of Surgery, General Surgery II, University of Pavia and Fondazione IRCCS Policlinico San Matteo, 27100 Pavia, Italy
6General Surgery, Ospedale Cannizzaro, 95126 Catania, Italy
*These authors contributed equally to this work and are First co-authors

Correspondence to: Sabrina Pricl, email: sabrina.pricl@dia.units.it

Keywords: KIT, GIST, novel deletion mutation, Imatinib, neoadjuvant treatment

ABSTRACT

Gastrointestinal stromal tumors (GISTs) are the most common mesenchymal tumors of the gastrointestinal tract. GISTs express the receptor tyrosine kinase KIT, and the majority of GISTs present KIT gain-of-function mutations that cluster in the 5’ end of the receptor juxtamembrane domain. On the other hand, little information is known about GISTs carrying mutations in the 3’ end of the KIT juxtamembrane domain. Here we report and discuss a clinical case of localized duodenal GIST whose molecular characterization revealed the presence of a new 21 nucleotide/7 amino acid deletion in the 3’ end of the KIT juxtamembrane domain (Δ574–580). The patient was treated with Imatinib at standard regimen dose (400 mg/day), and responded well as the original tumor mass reduced, ultimately allowing conservative surgery. In line with these clinical evidences computer simulations, biophysical techniques and in vitro experiments demonstrated that the receptor tyrosine kinase KIT carrying the Δ574–580 mutation displays constitutive phosphorylation, which can be switched-off upon Imatinib treatment. In addition, results from this study showed that a clinical useful procedure, neoadjuvant treatment, can occasionally be of value for the understanding of the molecular pathogenesis of GIST.

INTRODUCTION

Gastrointestinal stromal tumors (GISTs) are a relatively rare entity, accounting for less than 1% of GI tumors; nonetheless, they represent the most common mesenchymal tumors of the GI tract. GISTs arise from the muscle layer and are usually found in the stomach (60–70%), the proximal small intestine (25–30%), but can occur anywhere along the GI tract, exceptionally in the esophagus [1, 2]. In nearly 80–90% of GISTs, the oncogenic driver is a gain-of-function genetic alteration (mutations) in the receptor tyrosine kinase (RTK) KIT and it is now widely believed that GISTs arise from KIT-expressing interstitial cells of Cajal or their precursors [3].
The vast majority of KIT mutations (60–70%) are in-frame deletions (other include missense mutations (20–30%) and internal tandem duplications) clustered in the 5′ end of KIT juxtamembrane (KIT-JM) domain (exon 11) between Q550 and E561. Alterations in the 3′ end distal part of KIT-JM are rarely reported, and these include missense point mutations in codon L576, in-frame deletions, and rare internal tandem duplications of 1 up to more than 20 codons that are more often observed in gastric GISTs and associated with a favorable outcome [4]. Functional and molecular characterization of these rare 3-end variants is lacking. Other alternative mutational “hotspots” in KIT extracellular (exon 9, 18% of cases) and kinase (exon 13 and 17, less than 2%) domains have been identified in the GISTs that are negative for the exon 11 mutation [5, 6]. In the absence of KIT mutations, GISTs can harbor mutations of the PDGFRA gene (5–15% of cases), mainly located at exons 12 and 18, which are homologous with KIT exons 11 and 17 [7, 8]. The remaining cases (12–15%) lack KIT and PDGFRA mutations (KIT and PDGFRA wild-type GISTs), but these may include BRAF mutations (3%), loss of function of succinate dehydrogenase (SDH) complex (3%), and NF-1 mutations [9].

Seventy five % of GISTs are less than 4 cm of size, and it is recommended that tumors greater than 2 cm of size should undergo surgical resection [10]. In particular, approximately 20–25% of gastric and 40 to 50% of small intestinal GISTs are clinically malignant, metastases commonly develop in the abdominal cavity and liver, and may develop 10 to 15 years after primary surgery necessitating long-term clinical follow-up [2].

Most GISTs are localized and are managed by surgery alone; however, locally advanced or metastatic cases, as well as high-risk operated GISTs, require systemic therapy with Imatinib, a small molecule tyrosine kinase inhibitor. Response to Imatinib correlates with the presence and type of RTK KIT mutations. GISTs with the most common 5′ end KIT exon 11 mutations show the highest response rates, whereas the responsiveness of exon 9 KIT mutations appears to be sensitive to increased drug concentrations [11]. On the other hand, information on Imatinib activity in case of the rare 3′ end KIT-JM mutations is lacking. Neoadjuvant Imatinib is now considered a valuable option for treating KIT-mutated GISTs [12]; it can render a locally advanced GIST resectable, allow to perform less invasive procedures or to promote preservation of function, especially if the tumor is located in an anatomically difficult position, as in case of the rare subset of GISTs that arise in the duodenum (2–5% of cases) [13]. Furthermore, neoadjuvant treatment has the potential to be of value for tumor biology, providing a way to gain in vivo information on Imatinib sensitivity in case of new mutations of undetermined functionality.

Under this perspective, in this work we report and discuss a clinical case of localized duodenal GIST whose molecular characterization revealed the presence of a new 21 nucleotide/7 amino acid deletion in the 3′ end of KIT-JM domain (∆574–580). The patient was treated with Imatinib at standard regimen dose (400 mg/day), and responded well as the original tumor mass reduced, ultimately allowing conservative surgery. In line with these clinical evidences computer simulations, biophysical techniques and in vitro experiments demonstrated that the TKR KIT carrying the ∆574–580 mutation displays constitutive phosphorylation, which can be switched-off upon Imatinib treatment.

RESULTS

Patient clinical history

On March 2015 a 33-year-old man with silent medical history presented at the emergency unit with acute GI tract hemorrhagic anemia. No other clinical signs or symptoms were present. A CT scan showed a lesion in the II duodenal tract and the patient was admitted. Endoscopy showed persistent bleeding from the submucosal ulcerated duodenal mass. Contrast-enhanced abdomen NMR detected an oval mass (3.4 × 2.2 × 3.1 cm) with regular margins, in the medial aspect of the duodenal wall, immediately distal to the papilla and extending to nearby pancreatic head, displaying contact with the inferior cava vein (Figure 1, top panel, left).

Cytology obtained via echoendoscopy was highly suggestive for GIST, spindle cell type. Molecular typing was not readily possible, clinical conditions prompted intervention and the patient was eager to start treatment. The mass was technically resectable but required pancreatic-duodenectomy, with consequent significant morbidity. Bleeding indicated significant growth. 18F-FDG-PET/CT scan showed intense metabolic activity (SUV 6.3) (Figure 1, top panel, right), as expected for most GISTs, providing a way for testing Imatinib activity after a short drug trial in the context of a neoadjuvant approach (despite limited information in duodenal GISTs, Imatinib-sensitive mutations involving exon 11 are not uncommon) [14]. Indeed, a multidisciplinary evaluation proposed neoadjuvant treatment at the standard 400 mg/day Imatinib dose, which was started on early April 2015. Concomitantly, the patient underwent also a complete cardiac evaluation. Routine liver and renal function tests were within normal range.

Tumor histology

The duodenal lesion after Imatinib neoadjuvant treatment showed size reduction (2.5 × 1.5 × 1.5 cm); the lesion was of solid, white-yellowish appearance, entirely constituted by proliferation of spindle cells, with mild cellular atypia, and a storiform growth pattern (Supplementary Figure 1A). Immunophenotypic profile was: CD117/c-KIT +/− and DOG1 + (Supplementary
Figure 1B and 1C), CD34 −/+ , smooth muscle actin −, desmin −, H-caldesmon −/+ , S100 −, CAM5.2 −. Focal necrosis (less than 5%) and plurifocal hemorrhagic areas were observed. Signs of tumor regression included fibrosis and hydropic degeneration and blood vessel hyalinization. Mitotic index was 1/50 HPF, Ki67 < 3%. The lesion was localized within the muscular layer with focal extension into the submucosa, and was partially surrounded by a fibrotic rim. Resection margins were clear of tumor tissue. Risk assessment was not feasible because neoadjuvant treatment alters the two critical parameters, size and mitotic index.

KIT and PDGFRA molecular analysis

DNA sequencing of the duodenal GIST identified a new 21 codon in frame deletion c1718:1739del21 (p.T574_H580delTQLPYDH) encompassing the internal part of the juxtamembrane zipper region of KIT (JM-Z, 10 amino acids, residues 572–581), an area adjacent to the first trans-phosphorylation sites (Tyr 568 and Tyr 570) (see Supplementary Figure 2). As result of the deletion, the JM-Z consisted of just the 3 amino acids DPK. Interestingly, mutations described in this area include point mutations and internal tandem duplications [4]. No mutations were found in the KIT exons 9, 13 and 17, as well as in PDGFRA exons 12, 14 and 18 hot spots (Supplementary Figure 3).

Patient evaluation after Imatinib treatment

After prolonged discussion with the patient and informed consent obtained, Imatinib was started at the usual daily dose of 400 mg. Patient was monitored for hemoglobin concentration and occult fecal blood. Counts were stable and at day +30 of treatment, metabolic activity was not detected at a second FDG-PET-CT (Figure 1, middle panel, right), attesting optimal Imatinib sensitivity. The treatment was protracted till conservative surgery could be planned (approximately 8 months later). At that time, CE-NMR documented a mass of 1.3 cm size (Figure 1, middle panel, left) that allowed conservative surgery. Segmental duodenectomy with end-to-end duodenal reconstruction [14] was performed with optimal outcome, tumor resection was radical. Imatinib was started again as soon as clinical conditions allowed, and this because tumor histology after neoadjuvant treatment cannot reliably predict risk category. The patient was free of recurrence at last contrast-enhanced CT scan of the abdomen (approximately 1.5 years after surgery, Figure 1, bottom panel).

Figure 1: MRI and CT scan of patient at time of diagnosis and after Imatinib treatment. Top panel: Contrast-enhanced nuclear magnetic resonance of the abdomen (left) and whole body 18F-FDG PET/CT of the patient with a duodenal GIST (arrow) at time of diagnosis (right). The mass displayed contact with various structures (left) and manifested intense metabolic activity (right, SUV 6.3). Middle panel: Contrast-enhanced nuclear magnetic resonance (left) of the patient abdomen at time of post-neoadjuvant surgery. The mass was reduced in size allowing to perform segmental duodenectomy safely. Whole body 18F-FDG PET/CT of the patient (right) that after a single month neoadjuvant Imatinib showed absent metabolic activity, attesting optimal drug sensitivity. Bottom panel: patient history timeline.
Computational results

The calculated free energy of binding values listed in upper part of Table 1 show that the newly reported Δ574–580 mutant receptor is endowed with an affinity toward Imatinib (ΔGbind = −8.58 kcal/mol) slightly lower than that of the activating but Imatinib-responsive Δ559 isoform (ΔGbind = −9.15 kcal/mol; ΔΔGbind = −0.57 kcal/mol). Contextually, this deletion mutant seems to be provided with better affinity to the inhibitor compared to the WT receptor (ΔGbind value is −8.19 kcal/mol, [15]). From the structural point of view, no significant perturbation of the overall 3D structure of the Imatinib/Δ574–580 complex is detected compared to the Imatinib responsive KIT Δ559 isoform (Figure 2). On the other hand, the T670I is found very resistant to the TK inhibitor, as expected (ΔGbind = −6.38 kcal/mol).

In order to gain further insight into the binding mode of Imatinib to Δ574–580 KIT mutant receptor, the energetic contribution (ΔGbind,RES) for those residues which afford a substantial contribution to the binding was calculated. As shown in the left panel of Figure 3, the major contribution to Δ574–580 KIT/Imatinib binding stems from the combined action of four hydrogen bonds involving the side chains of residues E640, T670, C673, and D810 and further stabilizing interactions provided by two clustered hydrophobic regions (HRI and HRII) of the receptor including residues: A621, Y672, and L799 (HRI) and V643, L783, and H790 (HRII), respectively. The strength of the hydrogen bonds network is ensued by the persistence and optimal average dynamics length (ADL) of these interactions, monitored during all the equilibrated MD simulation. Indeed, the ADL values for each residue involved fall in the typical range of the permanent hydrogen bond interactions (ADL_{E640} = 1.93 ± 0.01 Å; ADL_{T670} = 1.91 ± 0.03 Å; ADL_{C673} = 2.15 ± 0.04 Å; ADL_{D810} = 2.05 ± 0.01 Å). Such efficient intermolecular interaction scheme accounts for the most substantial favorable energetic term to the total drug binding free energy (ΔGbind = −8.58 kcal/mol, Table 1), as these residues per se afford a stabilizing contribution of -8.23 kcal/mol (Figure 3, right panel).

In the case of the Imatinib/KIT Δ559 complex, no meaningful differences are detected at the individual residue binding level (Figure 3, right panel), and the identified interactions are in agreement with those described in previous works for these prototypical Imatinib-responsive KIT mutant [15–20]. Conversely, in the case of the Imatinib/KIT T670I assembly, the overall reduction of drug affinity cannot be attributed either to a single residue or to a particular cluster of binding site residues. Instead, a general decrease of the binding energy for all residues involved is predicted. This analysis confirms that the conformation adopted by the newly reported Δ574–580 KIT mutant and, in particular, of its inhibitor binding site, is comparable to the one characterizing the Imatinib-sensitive Δ559 KIT receptor; concomitantly, the quantitative examination of the essential molecular determinants for Imatinib binding provides a sensible explanation of the efficiency of this inhibitor against this new KIT juxtamembrane mutant.

The lower half of Table 1 lists the ΔGbind value obtained in silico for the Δ574–580 KIT mutant. The calculated ΔGbind values for the reference KIT variants Δ559 and T670I are also reported for comparison. As seen from these numbers, and from the related ΔΔGbind values in the last row of the top half of Table 1, the Δ574–580 KIT isoform shows an affinity for ATP (ΔGbind = −24.02 kcal/mol) utterly comparable to that of the Δ559 (ΔGbind = −24.20 kcal/mol; ΔΔGbind = −0.18 kcal/mol) and the T670I (ΔGbind = −25.14 kcal/mol; ΔΔGbind = +0.94 kcal/mol) mutants, supporting the activating character of these mutations [15–17, 19].

Isothermal titration calorimetry (ITC)

Isothermal titration calorimetry (ITC) experiments were to validate in silico prediction of KIT mutants affinity for Imatinib via determination of the drug binding thermodynamics (i.e., ΔG_{bind,ITC} and its major enthalpic (ΔH_{bind,ITC}) and entropic (-TΔS_{bind,ITC}) components, the

Figure 2: In silico analysis of Δ574–580 new KIT mutation. (Left) The solvated 3D model of the Imatinib/Δ574–580 complex. The secondary structure of the protein is in ribbon representation (red, α-helices; magenta, β-sheets; gray, turns and coil, while Imatinib is portrayed via its van der Waals surface. Na+ and Cl− ions and counterions are shown as purple and green spheres, respectively. Water molecules are depicted as transparent, light blue spheres. Hydrogen atoms are omitted for clarity. Superposition of MD equilibrated snapshots of Δ559 (green) and the Δ574–580 (blue) KIT in complex with Imatinib (center) and details of the corresponding Imatinib binding site (right).
dissociation constant $K_{\text{ITC}}$, and the binding stoichiometry $n$). Figure 4 shows the titration curves for Imatinib binding to ∆574–580, ∆559 and T670I KIT isoforms, while the relevant numerical results obtained from data fitting with a binding model that assumes one set of identical binding sites are displayed in Table 2.

As seen in Table 2, the stoichiometry of Imatinib binding to all KIT variants is approximately 1, confirming binding of one inhibitor molecule per RTK. The derived $K_{\text{ITC}}$ confirm strong Imatinib binding to both deletion mutants (359 nM and 127 nM for ∆574–580 and ∆559 KIT, respectively). The ineffectiveness of Imatinib is obvious for T670I, with a $K_{\text{ITC}}$ of approximately 30 μM. This analysis clearly demonstrate that Imatinib binds to the inactivated forms of the newly reported deletion mutant ∆574–580 with affinity comparable to that of the activating yet responding ∆559 isoform, and the loss of binding affinity of the inhibitor for the notoriously resistant T670I KIT variant. Finally, a comparison of the experimentally derived $\Delta G_{\text{bind,ITC}}$ values (Table 2) with the computer-predicted ones ($\Delta G_{\text{bind}}$, Table 1) reveals that the two data sets are in excellent agreement, ultimately yielding a direct validation of the in silico results presented above.

### Table 1: Binding free energies ($\Delta G_{\text{bind}}$) and binding free energy differences ($\Delta \Delta G_{\text{bind}}$) for the ∆559, ∆574–580, and T670I KIT receptors in complex with Imatinib and ATP

|                      | ∆574–580 | ∆559 | T670I |
|----------------------|----------|------|-------|
| **Imatinib binding** |          |      |       |
| $\Delta G_{\text{bind}}$ (kcal/mol) | -8.58 ± 0.08 | -9.15 ± 0.12 | -6.38 ± 0.11 |
| $\Delta \Delta G_{\text{bind}}$ (kcal/mol) | -0.57 | | -2.77 |
| **ATP binding**      |          |      |       |
| $\Delta G_{\text{bind}}$ (kcal/mol) | -24.02 ± 0.48 | -24.20 ± 0.82 | -25.14 ± 0.94 |
| $\Delta \Delta G_{\text{bind}}$ (kcal/mol) | -0.18 | | +0.94 |

According to its definition ($\Delta \Delta G_{\text{bind}} = \Delta G_{\text{bind,∆559}} - \Delta G_{\text{bind,mut}}$), negative values calculated for $\Delta \Delta G_{\text{bind}}$ indicate that the considered amino acid substitution at a given position of KIT is unfavorable in terms of imatinib/ATP affinity, whereas positive $\Delta \Delta G_{\text{bind}}$ values indicate that the considered mutation is favorable.

### Biological effects of the ∆574–580 mutation on KIT receptor activity

Next, the effect of Imatinib on the activation of the ∆574–580 KIT mutant was examined. Again, the same study was conducted in parallel on the reference ∆559 and T670I KIT variants for comparison. Accordingly, the HEK293T cells previously transfected with the mutated KIT expression constructs were cultured in absence and in presence of 1 and 5 μM Imatinib, respectively. As shown in Figure 5A, all proteins are expressed and phosphorylated, and both the mature (145 kDa) and partially glycosylated (125 kDa) forms of the receptor could be detected by immunoblotting of the lysates. The new ∆574–580 KIT mutant reveals constitutive phosphorylation of both 145 and 125 kDa forms of the receptor by the Western blot with anti-phospho-KIT antibodies. This behavior matches that observed for the well-defined ∆559 and T670I KIT variants, thereby confirming the constitutive KIT receptor-activating role of the ∆574–580 deletion, in agreement with the computational predictions reported in Table 1.

Also, again in accordance with in silico calculations/ITC determinations the newly reported KIT deletion...
variant is sensitive to Imatinib treatment at both concentrations (Figure 5A). Moreover, it exhibits a profile similar to that of the prototypical Imatinib-sensitive Δ559 KIT, in that the relative KIT phosphorylation levels are strongly reduced compared to the untreated cells (about 75%, Figure 5B). On the other hand, the T670I KIT mutant does not show any reduction of the KIT phosphorylation level in the presence of Imatinib, in line with its intrinsic Imatinib-resistant profile.

Furthermore, the effects of the new reported deletion on the typical signaling pathway triggered by KIT were investigated by Western blotting in absence and in presence of Imatinib. As shown in Figure 5C, phosphorylation of ERK1/2 and AKT is detectable in untreated cell expressing Δ559 or Δ574–580, thereby confirming the constitutive activation character of the new deletion KIT mutant. Interestingly, at both Imatinib concentrations (1 µM and 5 µM), the inhibition promoted by the inhibitor switches off the Δ574–580 KIT pathway, as both AKT and ERK1/2 phosphorylation is reduced (Figures 5D and 5E). As a similar effect is seen for the well-known Imatinib-sensitive Δ559 KIT isoform, these evidences support the Imatinib-sensitive behavior of this new KIT juxtamembrane variant, as anticipated by computer-assisted predictions and confirmed by isothermal titration calorimetry experiments.

**DISCUSSION**

Resistance to tyrosine kinase inhibitors is a major problem in cancer targeted therapy. Gastrointestinal stromal tumors are not exception to this, in that the majority (nearly 85%) of GISTS patients present primary activating mutations in the gene encoding the mast/stem cell growth factor receptor KIT, a type III RTK that plays an essential role in the regulation of cell survival and proliferation, hematopoiesis, stem cell maintenance, gametogenesis, mast cell development, migration and function, and in melanogenesis. Most of the reported KIT mutations in GISTS (missense, deletion and deletion/insertion variants) cluster in the receptor juxtamembrane domain (residues 550–586), encoded by the exon 11 of the related KIT gene. According to the COSMIC database, the most often reported KIT-JM mutations are point substitutions at positions 557 (e.g., W557R/G/S/C, V559D/A/G, and V560D/A/G), thereby affecting the 5′ end of the KIT-JM domain. At the same time, the efficacy of the tyrosine kinase inhibitor Imatinib in GISTS has been linked to the capacity of this small molecule to inhibit these aberrant receptor-activating mutant KIT proteins. Alterations at the other extreme of the KIT-JM (3′ end) are much less frequent, and these include missense point mutations in codon L576, in-frame deletions, and rare internal tandem duplications of 1 up to more than 20 codons. The latter KIT variants are more often observed in gastric GISTS, and are associated with a favorable outcome [4].

In this work we presented and discussed a newly discovered 3′ end KIT-JM 7-residues deletion mutant Δ574–5870 (i.e., delTQLPYDG), found in a 33-year-old male patient with a rare duodenal GIST that was difficult to resect at time of presentation. Neoadjuvant Imatinib at

### Table 2: Thermodynamic parameters extracted from the calorimetric evaluation of unphosphorylated Δ574–580, Δ559, and T670I mutant KITs titrated Imatinib

| KIT mutant | ΔGbind,ITC (kcal/mol) | ΔHbind,ITC (kcal/mol) | -TΔSbind,ITC (kcal/mol) | Kd,ITC (nM) | n   |
|------------|-----------------------|-----------------------|------------------------|-------------|-----|
| Δ574–580   | −8.80                 | −10.73                | 1.93                   | 359         | 1.12|
| Δ559       | −9.41                 | −11.67                | 2.26                   | 127         | 1.09|
| T670I      | −6.18                 | −7.25                 | 1.07                   | 29450       | 0.96|

Experiments were performed in triplicate at 25°C. Standard deviation values are within 1–3% for ΔHbind,ITC, 3–5% for Kd,ITC, and 1–2% for n.

Figure 4: Isothermal titration calorimetry of KIT mutants with Imatinib. ITC integrated heat curves for Imatinib binding to Δ574–580 (left), Δ559 (center), and T670I (right) KIT mutants. The inset shows ITC raw data for the tiration of Δ574–580 KIT mutant with Imatinib as an example. Solid curves represent data fitting with a 1:1 binding model.
standard regimen (400 mg/day), reduced the tumor mass to an extent that conservative surgery could be practiced. Molecular analysis revealed that no other mutations were present either in the KIT exons 9, 13 and 17, or in the PDGFRA exons 12, 14 and 18.

In line with the in vivo observations, computer-based simulations predicted that the new Δ574–580 KIT variant was indeed Imatinib-responsive, with an inhibitor affinity comparable to that of the known activating and Imatinib-responding Δ559 KIT isoform. In fact, the calculated ΔG_{bind} for this mutant is −8.58 ± 0.08 kcal/mol, corresponding to K_d or IC_{50} values (obtained from the fundamental relationship ΔG_{bind} = RTln K_d = RT ln IC_{50}) of 516 nM, while the corresponding values of these quantities for the Δ559 KIT mutants are −9.15 ± 0.12 kcal/mol and 198 nM, respectively (Table 1). At the same time, a validation test performed on the Imatinib refractory T670I KIT mutant yielded a ΔG_{bind} value of −6.38 ± 0.11 kcal/mol, corresponding to a K_d > 21 µM (Table 1).

Isothermal titration calorimetry (ITC) experiments (Figure 4) performed on purified, unphosphorylated KIT constructs fully supported the in silico predictions, leading to ΔG_{bind,ITC} and K_{d,ITC} values −8.80 kcal/mol and 359 nM for the Δ574–580 KIT and −9.41 kcal/mol and 127 nM for the Δ559 KIT, respectively (Table 2). The experimental values of ΔG_{bind,ITC} and K_{d,ITC} for the T670I KIT mutant (−6.18 kcal/mol and > 29 µM, Table 2) was also found in agreement with the corresponding computed values (Table 1), thereby ultimately validating the computational strategy and results adopted in the present study.

Cell-based assays next confirmed the constitutive activation of the KIT receptor bearing the Δ574–580 mutation; concomitantly, the capacity of Imatinib to inhibit this mutant KIT phosphorylation was also verified. Importantly, the effect of Δ574–580 KIT inhibition by Imatinib shared similar effects to those observed in cells transfected with the Δ559 KIT mutant, for which a strong phosphorylation of both AKT and ERK1/2 was detected (Figure 5).

In summary, in the present study we reported and characterized a new 7-residue deletion mutation Δ574–580 in the rarely involved 3′end of tyrosine kinase receptor

---

**Figure 5: Biological effects of the Δ574–580 mutation on KIT receptor activity.** (A) Western blot analysis of HEK293 cells transfected with the indicated KIT mutant constructs, untreated (0 µM) or treated with 1 or 5 µM Imatinib. (B) Quantitative analysis of relative KIT phosphorylation (bottom) expressed as % ratio between phosphorylated and total KIT levels and referred to untreated cells. Data represent the average of three independent experiments ± the standard deviation. (C) Effect of Imatinib on AKT and ERK1/2 phosphorylation induced by expression of Δ559 and Δ574–580 KIT deleted mutants in transiently transfected HEK293 cells. Arrows indicate the 145 and 125 kDa KIT receptor forms. (D and E) Quantitative analysis of relative AKT (D) and ERK1/2 phosphorylation (E) expressed as % ratio between phosphorylated and total protein levels and referred to untreated cells. Data represent the average of three independent experiments ± the standard deviation.
KIT juxtamembrane region, found in a 33-year-old male patient diagnosed with localized duodenal GIST who successfully underwent neoadjuvant imatinib treatment at standard dose (400 mg/day). In line with the in vivo observation of imatinib optimal sensitivity as attested by early complete metabolic response at FDG-PET, the results achieved by a combination of experimental and computational techniques indicate that the new Δ574–580 KIT mutant does not influence the overall sensitivity of KIT mutant toward Imatinib. Molecular simulations reveal that the deletion of 7 amino acids in the juxtamembrane domain of the receptor does not perturb the overall protein structure as well the binding region in which Imatinib is encased (Figures 2 and 3). Furthermore, the predicted affinity values of this KIT variant for ATP (Table 1) are in line with those calculated for two renown activating KIT mutations, i.e., the Δ559 and T670I KIT receptor [15–17, 19]. The molecular rationale yielded by the computational procedure is validated through ITC-based receptor/drug binding experiments and immunoblotting analysis of the effect of this new deletion on the intrinsic KIT activity and its response to Imatinib. Indeed, phosphorylation of the Δ574–580 KIT mutant is inhibited by Imatinib in a way similar to that observed for the Imatinib-sensitive Δ559 KIT control, whereas the kinase activity and thus receptor phosphorylation of the T670I, the prototypical Imatinib resistant KIT variant, is not inhibited at drug concentrations between 1 and 5 μM. The positive response to the kinase inhibitor of the Δ574–580 KIT form is further confirmed by the analysis of its signaling pathway in presence and in absence of Imatinib, according to which both AKT and ERK1/2 phosphorylation is indeed impaired at both Imatinib concentrations considered (Figure 5).

In addition, results from this study showed that a clinical useful procedure, neoadjuvant treatment, can occasionally be of value for the understanding of the molecular pathogenesis of GIST.

MATERIALS AND METHODS

KIT and PDGFRA molecular analysis

Mutation analysis of KIT exons 9, 11, 13, and 17, as well as PDGFRA exons 12, 14, and 18, was performed using direct sequencing of PCR products as described previously [21, 22]. Tumor cells comprised 98% of the cells in the sample used for gene sequencing. Three independent PCR experiments were performed to confirm sequencing results.

Computational Details

The 3D Δ574–580 (delTQLPYDH) KIT mutant model was obtained starting from our optimized WT KIT model in complex with Imatinib [15–20], following a consolidated methodology [23–27]. The validated structures of the Δ559 and T670I KIT mutants were also considered as reference for Imatinib responsive and resistant TK isoforms, respectively [15–20]. Following the same procedure, the complexes of all proteins with ATP were obtained from the corresponding optimized WT/KIT ensemble [15–17].

Each protein/ligand system was solvated, gradually heated to 25°C, equilibrated and subjected molecular dynamics (MD) data collection runs to perform drug/protein free energy of binding analysis [28–29].

All simulations were carried out using the Pmemd modules of Amber 16 [30], running on our Mose25 CPU/GPU calculation cluster (see the Supporting Information appendix for full computational details).

Cloning an KIT mutants purification for experimental Imatinib binding studies

Δ559, T670I, and Δ574–580 mutant KIT constructs were produced according to the established methodology described by Gajiwala et al. [31]. All mutant plasmids were sequenced to verify the success of mutagenesis experiments. The purified c-KIT protein was not phosphorylated as judged by routine mass spectrometry (data not shown). Full details are provided in the Supporting information appendix.

Isothermal titration calorimetry (ITC) experiments

Calorimetric titrations were carried out on a MicroCal PEAQ-ITC calorimeter (Malvern, UK) at 25°C. Before titrations, all KIT mutant proteins used were buffer-exchanged into an identical lot of HBS buffer (10 mM Hepes pH7.5, 150 mM NaCl). Gel filtration was used to control buffer heat dilution effects. Each protein sample was thoroughly degassed before ITC experiments, and these were run in triplicate (see Supporting information appendix for details).

Construction and transfection of mutated KIT

Δ559, T670I and the new variant Δ574–580 mutant KIT receptors were obtained as following a methodology reported in our previous work [15–17, 19]. Briefly, an expression vector carrying wild-type (WT) human complementary DNA (cDNA) for KIT (kind gift of Professor Y. Yarden, Weizmann Institute, Rehovot, Israel) was used to generate all mutated forms of KIT via site-directed mutagenesis using the commercial QuickChange Site-Directed Mutagenesis kit (Promega, Madison, WI), following manufacturer’s instruction. We then constructed mutant Δ559, in which amino acid 559 is removed from the juxtamembrane region by deleting nucleotides 1696 – 1698 from the portion of the exon 11-derived WT
For the T670I mutant, the second base of the T670 triplet codon ACA (i.e., cytosine 2030), was mutated to a thymine. Finally, the ∆574–580 mutant was obtained by removing nucleotides 1741–1761 from the corresponding portion of the WT cDNA. All plasmid inserts were sequenced after mutagenesis to verify their identity.

Cell cultures and transfections

HEK293T cells were maintained in Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented by 10% fetal calf serum, in 5% CO₂ humidified atmosphere, and transfected by Lipofectamine 2000 (Invitrogen), according to the manufacturer’s instructions. Cells were harvested 48 h after transfection following overnight serum starvation. Cell lysates were produced in RIPA modified buffer, and Western blot analysis was performed as described in our previous work [15–17, 19]. Anti-KIT (clone H300) antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA); antiphospho KIT (Y719, ref. 3391), AKT and phospho AKT (Ser473, ref. 9271) antibodies were obtained from Cell Signal Technology (Beverly, MA, USA) while ERK1/2 and phospho ERK1/2 (T202/Y204, ref. SAB4301578) antibodies were acquired from Sigma-Aldrich (St. Louis, MO, USA).

Abbreviations

CT, computed tomography; ¹⁸F-FDG-PET/CT, 2-deoxy-2-[fluorine-18]fluoro- D-glucose integrated with computed tomography; GIST, gastrointestinal stromal tumors; JM, juxtamembrane; NMR, nuclear magnetic resonance; RTK, receptor tyrosine kinase; SUV , standardized uptake value; 3D, three-dimensional.

Authors’ contributions

Conception and design: V. Perfetti, S. Pricl; Development of methodology: M. Lucioni, E. Dallera, S. Delfanti, R. Riboni, L. Pugliese, F.S. Latteri, A. Pietrabissa, E. Laurini, S. Aulić, M. Fermeglia; Acquisition of data (acquired and managed patients, computational studies, biophysical and molecular biology characterization, etc.): V. Perfetti, E. Laurini, S. Aulić; Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): V. Perfetti, E. Laurini, S. Aulić, M. Fermeglia, S. Pricl; Writing, review, and/or revision of the manuscript: V. Perfetti, M. Lucioni, E. Dallera, L. Pugliese, F.S. Latteri, A. Pietrabissa, E. Laurini, S. Aulić, M. Fermeglia, S. Pricl; Study supervision: V. Perfetti, S. Pricl.

CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

FINANCIAL SUPPORT

This work has received financial support from the Italian Association for Cancer Research (AIRC IG 2015 Id.17413 to SP), ED is recipient of a fellowship from Anatomic Pathology, IRCCS Fondazione Policlinico San Matteo, Pavia, Italy. RR is a recipient of a grant from the Italian Ministry of Health (5 per Mille) of IRCCS Fondazione Policlinico San Matteo, Pavia, Italy.

REFERENCES

1. Nilsson B, Bömминg P, Meis-Kindblom JM, Odén A, Dortok A, Gustavsson B, Sablinska K, Kindblom LG. Gastrointestinal stromal tumors: the incidence, prevalence, clinical course, and prognostication in the preimatinib mesylate era—a population-based study in western Sweden. Cancer. 2005; 103:821–829.
2. Miettinen M, Lasota J. Gastrointestinal stromal tumors: review on morphology, molecular pathology, prognosis, and differential diagnosis. Arch Pathol Lab Med. 2006; 130:1466–1478.
3. Kindblom LG, Remotti HE, Aldenborg F, Meis-Kindblom JM. Gastrointestinal pacemaker cell tumor (GIPACT): gastrointestinal stromal tumors show phenotypic characteristics of the interstitial cells of Cajal. Am J Pathol. 1998; 152:1259–1269.
4. Lasota J, Dansonka-Mieszkowska A, Stachura T, Schneider-Stock K, Kallajoki M, Steigen, SE, Sarlomo-Rikala M, Boltze C, Kordek R, Roessner A, Stachura J, Miettinen M. Gastrointestinal stromal tumors with internal tandem duplications in 3’ end of KIT juxtamembrane domain occur predominantly in stomach and generally seem to have a favorable course. Mod Pathol. 2003; 16:1257–1264.
5. Hirota S, Isozaki K, Moriyama Y, Hashimoto K, Nishida T, Ishiguro S, Kawano K, Hanada M, Kurata A, Takeda M, Muhammad Tunio G, Matsuzawa Y, Kanakura Y, et al. Gain-of-function mutations of c-kit in human gastrointestinal stromal tumors. Science. 1998; 279:77–80.
6. Miettinen M, Lasota J. Gastrointestinal stromal tumors. Gastroenterol Clin North Am. 2013; 42:399–415.
7. Heinrich MC, Corless CL, Duensing A, McGreevey L, Chen, CJ, Joseph N, Singer S, Griffith DJ, Haley A, Town A, Demetri GD, Fletcher CD, Fletcher JA. PDGFRA activating mutations in gastrointestinal stromal tumors. Science. 2003; 299:708–710.
8. Corless CL, Schroeder A, Griffith D, Town A, McGreevey L, Harrell P, Shiraga S, Bainbridge T, Morich J, Heinrich MC. PDGFRA mutations in gastrointestinal stromal tumors: frequency, spectrum and in vitro sensitivity to Imatinib. J Clin Oncol. 2005; 23:5357–5364.
9. Patil DT, Rubin BP. Genetics of gastrointestinal stromal tumors: a heterogeneous family of tumors? Surg Pathol Clin. 2015; 8:515–524.
10. Demetri GD, von Mehren M, Antonescu CR, DeMatteo RP, Ganjoo KN, Maki RG, Pisters PW, Raut CP, Riedel RF, Schuetze S, Sundar HM, Trent JC, Wayne JD. NCCN Task Force report: update on the management of patients with gastrointestinal stromal tumors. J Natl Compr Canc Netw. 2010; 8:S1–S41.

11. Gastrointestinal Stromal Tumor Meta-Analysis Group (MetaGIST). Comparison of two doses of imatinib for the treatment of unresectable or metastatic gastrointestinal stromal tumors: a meta-analysis of 1,640 patients. J Clin Oncol. 2010; 28:1247–1253.

12. Ford SJ, Gronchi A. Indications for surgery in advanced/metastatic GIST. Eur J Cancer. 2016; 63:154–67.

13. Buchs NC, Bucher P, Gervaz P, Ostermann S, Pugin F, Morel P. Segmental duodenectomy for gastrointestinal stromal tumor of the duodenum. World J Gastroenterol. 2010; 16:2788–2792.

14. Beham A, Schaefer IM, Cameron S, von Hammerstein K, Füzesi L, Ramadori G, Ghadimi MB. Duodenal GIST: a single center experience. Int J Colorectal Dis. 2013; 28:581–590.

15. Conca E, Miranda C, Dal Col V, Fumagalli E, Pelosi G, Mazzoni M, Fumagalli M, Laurini E, Pierotti MA, Piloti S, Greco A, Pricl S, Tamborini E. Are two better than one? A novel double-mutant KIT in GIST that responds to Imatinib. Mol Oncol. 2013; 7:756–762.

16. Tamborini E, Negri T, Miselli F, Lagonigro MS, Pricl S, Piloti S. Response of a KIT-positive extra-abdominal fibromatosis to Imatinib mesylate and KIT genetic analysis. J Natl Cancer Inst. 2006; 98:1583–1584.

17. Tamborini E, Pricl S, Negri T, Lagonigro MS, Miselli F, Greco A, Gronchi A, Casali PG, Ferrone M, Fumagalli M, Carbone A, Pierotti MA, Piloti S. Functional analyses and molecular modeling of two c-KIT mutations responsible for Imatinib secondary resistance in GIST patients. Oncogene. 2006; 25:6140–6146.

18. McAuliffe JC, Wang WL, Pavan GM, Pricl S, Yang D, Chen SS, Lazar AJ, Pollock RE, Trent JC. Unlucky number 13? Differential effects of KIT exon 13 mutation in gastrointestinal stromal tumors. Mol Oncol. 2008; 2:161–163.

19. Negri T, Pavan GM, Virdis E, Greco A, Fumagalli M, Sandri M, Pricl S, Pierotti MA, Piloti S, Tamborini E. T670X KIT mutations in gastrointestinal stromal tumors: making sense of missense. J Natl Cancer Inst. 2009; 101:194–204.

20. Woodman SE, Trent JC, Stemke-Hale K, Lazar AJ, Pricl S, Pavan GM, Fumagalli M, Gopal YN, Yang D, Podoloff DA, Ivan D, Kim KB, Papadopoulos N. Activity of dasatinib against L576P KIT mutant melanoma: molecular, cellular, and clinical correlates. Mol Cancer Ther. 2009; 8:2079–2085.

21. Haller F, Gunawan B, von Heydebreck A, Schwager S, Schulten HJ, Wolf-Salgó J, Langer C, Ramadori G, Sültmann H, Füzesi L. Prognostic role of E2F1 and members of the CDKN2A network in gastrointestinal stromal tumors. Clin Cancer Res. 2005; 11:6589–6597.

22. Battochio A, Mohammed S, Winthrop D, Lefresne S, Mulder K, Chu Q, O’Hara C, Lai R. Detection of c-KIT and PDGFRA gene mutations in gastrointestinal stromal tumors: comparison of DHPLC and DNA sequencing methods using a single population-based cohort. Am J Clin Pathol. 2010; 133:149–155.

23. Brambilla L, Genini D, Laurini E, Merulla J, Perez L, Fumagalli M, Carbone GM, Pricl S, Catapano CV. Hitting the right spot: mechanism of action of OPB-31121, a novel and potent inhibitor of the Signal Transducer and Activator of Transcription 3 (STAT3). Mol Oncol. 2015; 9:1194–1206.

24. Brune S, Schepmann D, Klempnauer KH, Marson D, Dal Col V, Laurini E, Fumagalli M, Wünsch B, Pricl S. The sigma enigma: in vitro/in silico site-directed mutagenesis studies unveil σ1 receptor ligand binding. Biochemistry. 2014; 53:2993–3003.

25. Bozzi F, Conca E, Laurini E, Posocco P, Lo Sardo A, Jocollè G, Sanfilippo R, Gronchi A, Perrone F, Tamborini E, Pelosi G, Pierotti MA, Maestro R, et al. In vitro and in silico studies of MDM2/MDMX isoforms predict Nutlin-3A sensitivity in well/de-differentiated liposarcomas. Lab Invest. 2013; 93:1232–1240.

26. Dileo P, Pricl S, Tamborini E, Negri T, Stacchiotti S, Gronchi A, Posocco P, Laurini E, Coco P, Fumagalli E, Casali PG, Piloti S. Imatinib response in two GIST patients carrying two hitherto functionally uncharacterized PDGFRA mutations: an imaging, biochemical and molecular modeling study. Int J Cancer. 2011; 128:983–990.

27. Laurini E, Col WD, Mamolo MG, Zampieri D, Posocco P, Fumagalli M, Vio L, Pricl S. Homology model and docking-based virtual screening for ligands of the σ1 receptor. ACS Med-Chem Lett. 2011; 2:834–9.

28. Pierotti MA, Negri T, Tamborini E, Perrone F, Pricl S, Piloti S. Targeted therapies: the rare cancer paradigm. Mol Oncol. 2010; 4:19–37.

29. Pierotti MA, Tamborini E, Negri T, Pricl S, Piloti S. Targeted therapy in GIST: in silico modeling for prediction of resistance. Nat Rev Clin Oncol. 2011; 8:161–170.

30. Case DA, Betz RM, Cerutti DS, Cheatham TE 3rd, Darden TA, Duke RE, Giese TJ, Gohlke H, Goetz AW, Homeyer N, Izadi S, Janowski P, Kaus J, et al. AMBER 2016, University of California, San Francisco (CA, USA), 2016.