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

SUPPLEMENTARY MATERIALS

Computational details

The optimized structure of KIT was taken from our previous work (Tamborini et al., 2006a, 2006b; McAuliffe et al., 2008; Negri et al., 2009; Woodman et al., 2009; Conca et al., 2013). Mutations ∆559, T670I and ∆574-580 were introduced into the wild-type structure of KIT/imatinib complex by swapping the mutant residue into the specific site, according to a dedicated and well-validated procedure (Tamborini et al., 2006a, 2006b; McAuliffe et al., 2008; Negri et al., 2009; Woodman et al., 2009; Conca et al., 2013). Each protein/ligand system was relaxed in a box of TIP3P water molecules (Jorgensen et al., 1983). To achieve electroneutrality, a suitable number of neutralizing ions were added; further, the solution ionic strength was adjusted to the physiological value of 0.15 M by adding the required amounts of Na⁺ and Cl⁻ ions. Each hydrated complex system was then gradually heated to 25°C and then equilibrated for 5 ns via molecular dynamics (MD) simulations under isobaric/isothermal conditions (i.e., NPT ensemble). Finally, each MD equilibrated system was subjected to 50 ns of NPT MD data collection runs to estimate the corresponding drug/protein free energy of binding (Pierotti et al., 2010; Pierotti et al., 2011).

All simulations were carried out using the Pmemd modules of Amber 16 (Case et al., 2016).

All energetic analysis was carried out for only a single MD trajectory of each KIT/imatinib complex considered, with unbound protein and substrate snapshots taken from the snapshots of that trajectory.

The affinity of the mutant KIT isoforms for imatinib and ATP, expressed as the value of the corresponding free energy of binding \( \Delta G_{\text{bind}} \), was estimated using the Molecular Mechanics/Poisson-Boltzmann Surface Area (MM/PBSA) method (Kollman et al., 2000, Puerotti et al., 2011). Accordingly, the binding free energy between two biological entities (e.g., a drug and its protein target) in a solvent was obtained as the sum of the interaction energy between the receptor and the ligand \( \Delta E_{\text{MM}} \), the solvation free energy \( \Delta G_{\text{solv}} \), and the conformational entropy contribution \( -T\Delta S \), averaged over a series of snapshots from the corresponding MD trajectories:

\[
\Delta G_{\text{bind}} = \Delta E_{\text{MM}} + \Delta G_{\text{solv}} - T\Delta S
\]  

(S1)

The \( \Delta E_{\text{MM}} \) term in Eq. (1) can be obtained directly from the molecular mechanics interaction energies as:

\[
\Delta E_{\text{MM}} = \Delta E_{\text{int}} + \Delta E_{\text{vdW}} + \Delta E_{\text{ele}}
\]  

(S2)

where \( \Delta E_{\text{int}}, \Delta E_{\text{vdW}}, \) and \( \Delta E_{\text{ele}} \) are the internal, van der Waals and electrostatic components of the nonbonded interaction energy, respectively. Since in this work we adopted the “single trajectory protocol”, then \( \Delta E_{\text{int}} = 0 \) in Eq. (S2).

The second term Eq. (S1), the solvation energy \( \Delta G_{\text{solv}} \) can also be partitioned into two different contributions:

\[
\Delta G_{\text{solv}} = \Delta G_{\text{PB}} + \Delta G_{\text{NP}}
\]  

(S3)

The polar term of \( \Delta G_{\text{solv}}, \Delta G_{\text{PB}} \) was estimated by solving the Poisson-Boltzmann equations numerically and calculating the electrostatic energy according to the electrostatic potential. In these calculations, the interior and exterior dielectric constant values were set equal to 1 and 80, respectively. A grid spacing of 0.5 Å, extending 20% beyond the dimensions of the solute, was employed. The value of nonpolar component of \( \Delta G_{\text{solv}}, \Delta G_{\text{NP}} \) was calculated using the following relationship (Sitkoff et al., 1994):

\[
\Delta G_{\text{NP}} = \gamma \times S + \beta
\]  

(S4)

in which \( \gamma = 0.00542 \text{ kcal (mol Å²)}^{-1}, \beta = 0.92 \text{ kcal/mol}, \) and \( S \) is the molecular surface area estimated by means of the MSMS software (Sanner et al., 1993).

The change in solute entropy upon association \( -T\Delta S \) was estimated using the Nmode module of Amber 16. The normal-node analysis was performed for the minimized structures of the complexes, KIT, and ligands using a distance-dependent dielectric constant \( \varepsilon = 4r \). In the first step of this calculation, an 8-Å sphere around the ligand was cut out from an MD snapshot for each ligand-protein complex. This value was shown to be large enough to yield converged mean changes in solute entropy. On the basis of the size-reduced snapshots of the complex, we generated structures of the uncomplexed reactants by removing the atoms of the protein and ligand, respectively. Each of those structures was minimized, using a distance-dependent dielectric constant \( \varepsilon = 4r \), to account for solvent screening, and its entropy was calculated using classical statistical formulas and normal mode-analysis. To minimize the effects due to different
conformations adopted by individual snapshots we averaged the estimation of entropy over 50 snapshots.

A per-residue binding free energy decomposition (PRBFED) was performed exploiting the MD trajectory of each complex. This analysis was carried out using the MM/GBSA approach (Tsui and Case, 2000), and was based on the same snapshots used in the binding free energy calculation.

**Cloning and 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. (2009). Accordingly, a construct encoding an N-terminal 6xHis tag + r3c protease site + KIT residues 544-976 was cloned into the baculovirus expression vector pVL1393 (Invitrogen). Proteins were expressed in Sf9 cells for 48 h; cells were then harvested and frozen (−80°C). Cellular lysates were obtained by treatment with 25 mM Tris (pH 7.4), 250 mM NaCl, 0.25 mM TCEP, and 20 mM imidazole. After binding to a Ni-NTA column, each protein was step-eluted using the same buffer employed for pellet lysing with 250 mM imidazole. Each peak eluted fraction was subjected to dialysis at 4°C for 12 h. Next, the His tag was cleaved with the r3C protease and removed by reloading the proteins on the Ni-NTA column. Each Flow-through was collected from the column and concentrated. Next, each protein was feeded to a Superdex-200 column equilibrated in 25 mM Tris (pH 7.4), 250 mM NaCl, 0.5 mM TCEP, and 1mM EDTA. The peak fractions were pooled, concentrated (up to 6 mg/ml) and quickly frozen.

**Isothermal titration calorimetry (ITC) experiments**

A solution of each purified KIT mutant protein (10 µM, sample cell) was titrated with 36 step-by-step injections (spacing 150 s) of 2 µL volume of Imatinib (250 µM, syringe). Solutions and buffer were degassed for 30 min at room temperature under stirring at 750 rpm prior to each experiment.

**SUPPLEMENTARY REFERENCES**

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Supplementary Figure 1: Tumor histology: (A) Histopathologic features (HE, 200×): the tumor mainly consisted of spindle cells exhibiting a fascicular growth pattern. (B) Immunostaining for c-KIT/CD117 (400×): most tumor cells showed membranous and/or cytoplasmic positivity for CD117. (C) Immunostaining for DOG1 (400×): most tumor cells showed strong immunoreactivity for DOG1.
Supplementary Figure 2: Sequencing of exon 11 of WT and mutated KIT. Top panel: comparison between exon 11 of WT and patient KIT. Bottom panel: Patient KIT sequence (partial in length) comprising the region encoding for the newly reported mutation Δ574-580, sequence showing the in frame deletion mutation c1718:1739del21.
Supplementary Figure 3: Sanger sequencing of KIT exons 9, 13 and 17 hotspots (first three panels from top), and of PDGFRA exons 12, 14 and 18 hotspots (last three panels).