

## ORIGINAL RESEARCH ARTICLE

## Covalent docking-based virtual screening and molecular dynamics simulations identify C02b as a potential *KRAS*(G12C) inhibitor

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### Abstract

Decades of efforts to target the “undruggable” *Kirsten rat sarcoma viral oncogene homolog* (*KRAS*) oncoprotein yielded promising results in 2021 with the approval of Sotorasib, a *KRAS*(G12C) inhibitor for non-small cell lung cancer patients with the *KRAS*(G12C) variant. Before Sotorasib’s approval, developing *KRAS*(G12C) covalent inhibitors faced challenges, particularly their inability to preserve the *KRAS* protein in the GDP-bound state, which hindered clinical trial progression. Considering the importance of developing inhibitors targeting *KRAS*(G12C), we aimed to identify compounds analogous to Sotorasib, resulting in the discovery of a total of 174 Sotorasib scaffold-compounds. We then performed covalent docking-based virtual screening to examine the binding affinity of Sotorasib-like compounds to *KRAS*(G12C). Four compounds showed comparable binding energies according to Glide score to Sotorasib for targeting *KRAS*(G12C). Subsequently, molecular dynamics (MD) simulations were conducted for these four compounds, spanning 100 ns, 300 ns, and 500 ns durations, to identify the most stable complex with the lowest root-mean-square deviation (RMSD), similar to the *KRAS*(G12C)-Sotorasib reference complex. Additional dynamic cross-correlation matrix and PCA were performed as post-MD analyses to investigate the movements of two switches and the flexible regions of *KRAS*(G12C)-Sotorasib and -C02b complexes. As a result, among these four compounds, *KRAS*(G12C)-C02b was found as the optimal candidate. Further investigations beyond this study may provide more insight into C02b’s inhibitory effect on *KRAS*(G12C), offering a deeper understanding of its potential as a therapeutic agent.

**Keywords:** *KRAS*(G12C); Sotorasib-like compounds; Molecular dynamics simulations

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### 1. Introduction

Cancer is a complex disease characterized by the number of key hallmarks, including sustained proliferative signaling, evasion of growth suppressors, resistance to cell death, enabling replicative immortality, inducing angiogenesis, activation of invasion and metastasis, deregulation of cellular energetics, avoidance of immune destruction, promotion of inflammation, genome instability and mutation, and alteration of

the tumor microenvironment.<sup>1,2</sup> These characteristics collectively contribute to the development and progression of cancer, making it a difficult and complex disease to treat. Understanding these characteristics is critical for developing cancer therapies that target specific aspects of the disease's biology, as cancer is greatly influenced by evolutionary principles, particularly through tumor evolution.<sup>3-6</sup> Genetic mutations and natural selection drive the development of diverse cell populations within a tumor, resulting in genetic diversity and heterogeneity.<sup>7</sup> This diversity enables cancer cells to adapt to various environmental challenges, such as treatments or changes in the microenvironment, resulting in the survival of drug-resistant and more aggressive cell clones.<sup>8-11</sup>

*Kirsten rat sarcoma viral oncogene homolog* (*KRAS*) signaling regulates important cellular functions through a network of intracellular signaling pathways.<sup>12</sup> *KRAS* functions normally as a molecular switch, cycling between an inactive (GDP-bound) and active (GTP-bound) form. When activated, GTP-bound *KRAS* drives downstream signaling pathways that regulate cell growth, survival, and proliferation, such as the MAPK/ERK and PI3K/AKT pathways.<sup>13</sup> *KRAS* mutations, such as the common G12C mutation, maintain *KRAS* in an active state, resulting in uncontrolled signaling and cellular responses, ultimately driving cancer development and progression by promoting cell growth, inhibiting apoptosis, enhancing metastasis, and modulating the tumor microenvironment.<sup>14</sup>

*KRAS* gene mutations, particularly the G12C variant, are prevalent in a variety of cancers, including non-small cell lung cancer (NSCLC) and colorectal cancer.<sup>15,16</sup> *KRAS*(G12C) inhibitors are an emerging class of targeted cancer therapy that targets a specific mutation (G12C) in the *KRAS* gene.<sup>17</sup> *KRAS*(G12C) inhibitors aim to address a long-standing problem in cancer treatment, as *KRAS* mutations were previously considered “undruggable”<sup>18</sup> Sotorasib (AMG 510) was the first *KRAS*(G12C) inhibitor to receive accelerated FDA approval in May 2021 for the treatment of adult patients with NSCLC harboring the *KRAS*(G12C) mutation.<sup>19</sup> This marked a significant milestone in cancer therapy by introducing the first drug approved to target *KRAS* mutations. Sotorasib functions by irreversibly binding to the mutant *KRAS*(G12C), inhibiting its activity.<sup>20</sup> Specifically, Sotorasib forms a covalent bond with cysteine 12 in the switch II pocket of *KRAS*(G12C),<sup>21</sup> preventing its ability to activate downstream signaling pathways that promote cell growth and proliferation.<sup>13</sup>

Although Sotorasib has been approved as a drug for targeting *KRAS*(G12C),<sup>19</sup> there are still a limited number of virtual screening approaches to identify potential inhibitors for the *KRAS*(G12C).<sup>22-24</sup> Given this urging

need to identify novel pharmacophores targeting the *KRAS*(G12C), we performed a covalent docking-based virtual screening in conjunction with molecular dynamics (MD) simulations to investigate the stability of *compounds* binding to the target *protein*. Based on scaffold-hopping approach with Sotorasib as a template, we identified 174 molecules exhibiting structural similarities to Sotorasib among billions of compounds. Our covalent docking-based virtual screening revealed potential putative binders of *KRAS*(G12C), namely, C01, C02a, C02b, and C03, with notable Glide score similarities to Sotorasib. MD simulations further demonstrated that the *KRAS*(G12C)-C02b complex exhibited stable binding interactions with a low root-mean-square deviation (RMSD) value, comparable to the reference FDA-approved drug Sotorasib. Moreover, the calculations of binding free energy indicated that these favorable interactions were highly preserved from the last 50 nanoseconds (ns) up to 500 ns of simulation time. To further investigate the dynamics of the *KRAS*(G12C)-Sotorasib and *KRAS*(G12C)-C02b complexes, we conducted post-MD analyses, including dynamic cross-correlation matrix (DCCM) and principal component analysis (PCA). These analyses aimed to elucidate the movements of the two switches and identify the flexible regions within the complexes. These findings suggest that C02b may serve as a promising inhibitor candidate for the *KRAS*(G12C) mutation, and further advanced studies are needed to validate its potential.

## 2. Materials and methods

### 2.1. Similarity search

Sotorasib served as a template to identify similar compounds within various chemical spaces including eXplore, Freedom Space, GalaXi, and KnowledgeSpace. The scaffold-hopping search was conducted using the *infiniSee* software (version 5.0.1; BioSolveIT GmbH, Sankt Augustin, Germany, 2023, [www.biosolveit.de/infiniSee](http://www.biosolveit.de/infiniSee)). The chemical spaces encompassing both building blocks and reactions allowed for the creation of a maximum number of virtual product molecules. Scaffold Hopper was executed using the *infiniSee* software.<sup>25</sup> The acrylamide warhead was selected from Sotorasib<sup>22</sup> (Figure A1 in Appendix). The target similarity was set at 1.00 to ensure a precise match, while a minimum similarity threshold was established at 90% to guarantee a significant level of similarity. Furthermore, a total diversity score of 1.00 was ensured to uncover compounds with a high degree of similarity with Sotorasib. As a result, 174 compounds containing the acrylamide warhead and showing structural similarities to Sotorasib were identified and saved in the structure-data file (SDF) format.

## 2.2. Protein and ligand preparation

The 174 compounds containing the acrylamide warhead were prepared and optimized for molecular docking using LigPrep.<sup>26,27</sup> Compounds were obtained using Epik at pH  $7.0 \pm 1.0$ .<sup>28</sup> The protein structure of the inactive GDP-bound state of human KRAS(G12C) (PDB ID: 6OIM) was retrieved from the Protein Data Bank, with a resolution of 1.65 Å.<sup>29</sup> Protein preparation was performed using the protein preparation wizard in Maestro,<sup>30</sup> which fixed all the missing residues and atoms of the protein structure. Several crucial steps in the preparation included introducing hydrogen atoms, refining hydrogen bond networks, and addressing missing side chains through the utilization of Prime. Subsequently, water molecules were eliminated, and a restrained minimization employing the OPLS3e force field was carried out.<sup>31</sup> The resulting optimized protein structure was then utilized for the subsequent docking process.

## 2.3. Covalent dockingbased virtual screening

A set of 174 pharmacophores similar to Sotorasib was employed in covalent docking-based virtual screening (CovDock-VS)<sup>32</sup> against the KRAS(G12C) target (PDB ID: 6OIM). The reactive residue Cys12 was specifically selected, and grid generation for docking was created as a cubic grid box centered at the coordinates (x, y, and z) of Sotorasib, with a length of 12 Å. The acrylamide electrophilic warhead group, identical to the template drug Sotorasib, was selected for the Michael addition – a reaction responsible for the covalent bonding of ligands featuring an acrylamide functional group with the side chain of nucleophilic protein residues. Subsequently, the ligands were selected and prioritized based on their Glide scores for the binding modes of pre-reactive complexes.<sup>33,34</sup>

## 2.4. MD simulations

MD simulations were conducted on the protein-ligand complexes derived from the covalent docking-based virtual screening using GRONingen MACHine for Chemical Simulations (GROMACS) 2023.1.<sup>35</sup> The CHARMM-GUI (Chemistry at HARvard Macromolecular Mechanics Graphical User Interface) server was used for solution building. The CHARMM36 force field<sup>36</sup> was applied to define atom types and assign atomic partial charges for both the ligand and protein. Furthermore, the CHARMM-GUI server was employed to generate the topology for both the KRAS(G12C) receptor and the promising inhibitors. For the setup of MD simulations, the protein-ligand complexes were centered in an octahedral box and solvated with the TIP3P water model. Neutralization of the complexes charge was achieved by adding 150 mM ions. The minimization process concluded when the maximum force was reduced

to <10.0 kJ/mol, with a maximum duration of 100 picoseconds (ps) for the minimization steps. Subsequently, a 100 ps equilibration phase with position restraints on the protein and ligand molecules was conducted using NVT (Number of Volume and Temperature) and NPT (Number of Pressure and Temperature) ensembles. During this period, the systems were heated to 303.15 K using a modified Berendsen thermostat (V-rescale),<sup>37,38</sup> and pressure was maintained using C-rescale pressure coupling set to a reference pressure of 1 bar.<sup>39</sup> For energy minimization, NVT, and NPT relaxation simulations, short-range interactions utilized a smooth force switch with a cut-off of 1.2 nm, and long-range electrostatics were computed using the Particle-Mesh-Ewald (PME) algorithm.<sup>40</sup> In addition, hydrogen bonds were constrained by employing the Linear Constraint Solver algorithm.<sup>41</sup> Finally, MD simulations were conducted for durations of 100 ns, 300 ns, and 500 ns (Table 1) without applying restraints. A 2 femtoseconds integration time step was employed, and trajectory snapshots were captured at 1 ps intervals.

MDS analyses were performed using GROMACS, a Linux-x86\_64-multicore CUDA-enabled program on an NVIDIA GeForce GTX 1060 GPU and an 8-core Intel Core i5-7000 central processing infrastructure. The computation of a single complex required approximately 100 ns of processing time per simulation day. GROMACS modules were used to analyze MD trajectories, focusing on parameters such as root RMSD/mean-square deviation fluctuation (RMSF) and hydrogen bond (H-bond) analysis. Graphs were generated using XMGrace<sup>42</sup> and visual molecular dynamics (VMD).<sup>43</sup>

The calculation of binding free energy was performed using the gmx\_MMPBSA tool, which is based on Assisted Model Building with Energy Refinement MM/PB(GB)SA approach. This tool is designed for conducting end-state free energy calculations using GROMACS trajectory files. Specifically, the final 50 ns of the simulation trajectory were considered for the molecular mechanics with Generalized Born surface area (MM/GBSA) calculation.

**Table 1. KRAS(G12C) complexes feature with their atom numbers for duration 500 ns**

Atoms	KRAS(G12C)-Sotorasib	KRAS(G12C)-C02b
Protein	2,704	2,704
Ligand	72	87
Na <sup>+</sup>	14	13
Cl <sup>-</sup>	7	7
Water (TIP3P)	7,914	7,923

Abbreviations: TIP3P: Transferable intermolecular potential three-point; KRAS: Kirsten rat sarcoma viral oncogene homolog.

## 2.5. Post-MD simulations analyses

The analysis employed Bio3D,<sup>44</sup> an R library-based tool, to generate time-correlated DCCM for various ligand-bound forms of KRAS(G12C). This approach aimed to discern the localization and nature of fluctuations in C $\alpha$  atoms within residues crucial to KRAS(G12C) function. Given the limitations of DCCM, which may overlook correlated but perpendicular motions, the analysis was complemented with PCA. In PCA, simulation frames are organized based on their principal components (eigenvectors), which represent the key directions capturing the most significant variability in the data. This approach allows for effective separation of structures by emphasizing dominant patterns and variations within the dataset. For both analyses, the final 400 ns of all MD simulations were utilized at 20 ps intervals per Bio3D file size constraints. This comprehensive approach provides a nuanced understanding of the dynamic behavior of KRAS(G12C) ligand-bound forms, integrating both normal mode analysis and PCA techniques to capture a broad spectrum of structural fluctuations.

## 3. Results

### 3.1. Covalent docking-based virtual screening

It is widely accepted that molecules with comparable chemical structures will have similar pharmacological properties. In ligand-based studies, Sotorasib, an FDA-approved drug for treating KRAS(G12C) protein in NSCLC, was reported to be a reliable template. To evaluate ligand interactions, we used the InfiniSee, a platform capable of examining billions of compounds across chemical spaces, including eXplore, Freedom Space, GalaXi, CHEMriya, and REAL, to identify potential chemical scaffolds similar to the template. This platform also provides access to KnowledgeSpace, a literature-based virtual chemical space with a strong emphasis on synthetic accessibility. These technological advancements offer great potential in the fields of drug discovery and chemical research due to its capacity to provide a greater spectrum of molecules, allowing the design of more effective and innovative therapeutic solutions. As a result, 174 molecules exhibiting structural similarities to Sotorasib were found from billions of compounds in the chemical spaces (Table A1 in Appendix). Subsequently, the selected compounds were further evaluated using a covalent docking-based virtual screening approach.

### 3.2. Covalent docking-based virtual screening identifies four compounds

For the covalent-based virtual screening approach, the X-ray structure of the KRAS(G12C) (PDB ID: 6OIM), forming a complex with Sotorasib, was selected.<sup>29</sup> The

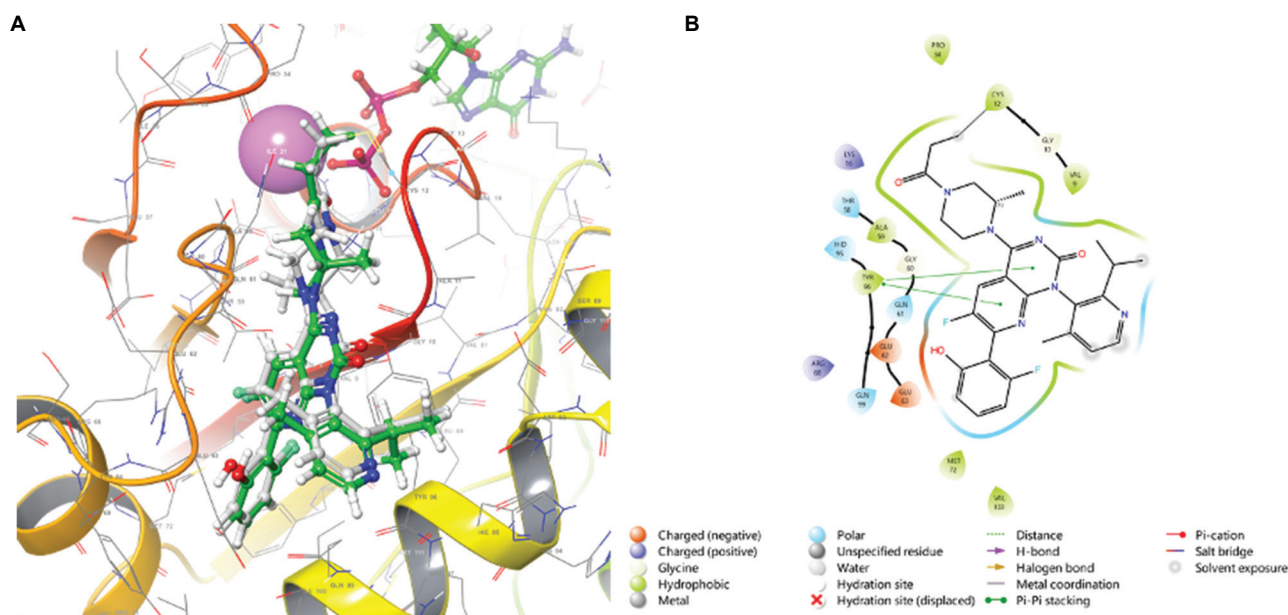
COVDock protocol was utilized to investigate the redocking capabilities as well as the binding of the co-crystallized Sotorasib to the KRAS(G12C) with GDP. As a result, we observed a similar binding mechanism to that observed in the crystallized structure, validating the covalent docking protocol (Figure 1A). Next, the binding modes of these four compounds were compared with the co-crystallized Sotorasib according to their 2-D interaction diagram (Figure 1B). The precision of the COVDock protocol in reproducing the binding mechanism of the co-crystallized ligand with the KRAS(G12C) protein has been demonstrated by prior researchers.<sup>22,45</sup>

The set of 174 selected compounds was then subjected to a covalent docking-based virtual screening, with their binding energies ranked according to their Glide scores. While none of the ranked compounds displayed a higher estimated binding energy compared to Sotorasib, which possesses a value of  $-8.1$  kcal/mol, four compounds namely C01, C02a, C02b, and C03 exhibited closer and notable Glide scores of  $-7.8$  kcal/mol,  $-7.5$  kcal/mol,  $-7.3$  kcal/mol, and  $-7.0$  kcal/mol, respectively (Table 2 and Figure A1).

The comparison of binding modes for the four compounds with co-crystallized Sotorasib exhibited Pi-Pi stacking interactions with the residue TYR96. Similarly, compounds C01, C02a, C02b, and C03 exhibited similar interactions (Figure 2). Furthermore, these compounds formed hydrogen bonds with additional residues (LYS16, ALA59, and GLN61) in the binding pocket of KRAS(G12C) (Figure 2). Taken together, the four candidates show promise as covalent binders of KRAS(G12C) in its inactive GDP-bound configuration.

### 3.3. MD simulations of C01, C02a, C02b, C03, and Sotorasib with KRAS(G12C) protein

The root RMSD analysis allows us to monitor the fluctuation in the three-dimensional structure over time, offering valuable insight into the mobility of binding pocket residues during the MD simulation. RMSD is a measure used in computer simulations to determine how far a molecule or part of it has moved from its initial position. To assess possible fluctuations in the 3D structure over time, we ran simulations on various KRAS(G12C) protein complexes, including KRAS-C01, KRAS-C02a, KRAS(G12C)-C02b, KRAS-C03, and KRAS(G12C)-Sotorasib. Initially, we ran simulations for all complexes for 100 ns. Following that, we increased the simulation time for KRAS-C01, KRAS(G12C)-C02b, and KRAS(G12C)-Sotorasib to 300 ns. Finally, we ran simulations for KRAS(G12C)-C02b and KRAS(G12C)-Sotorasib for 500 ns. This enabled us to compare the structural stability of potential therapeutic



**Figure 1.** (A) The binding modes of the X-ray-bound Sotorasib and the re-docked Sotorasib were superimposed. (B) Amino acid interactions of Sotorasib with *Kirsten rat sarcoma viral oncogene homolog*(G12C) and compound prioritization based on molecular dynamics simulations and binding free energy calculations.

**Table 2. The binding affinities and physicochemical properties of KRAS-C01, KRAS-C02a, KRAS(G12C)-C02b, and KRAS-C03 resembling the chemical scaffold of Sotorasib. TPSA: vdW polar surface area, LogP: Predicted octanol/water partition coefficient**

Compound number	Compound name	Similarity	Glide score (kcal/mol)	Prime energy	Molecular weight	TPSA	LogP
	Sotorasib	1.00	-8.157	-19.6	560.6	102	4
C01	Shen_92298609_93443091_95741143	0.917	-7.801	-78.3	605.770	95.660	7.178
C02a	Shen_92298609_44473994_95741143	0.916	-7.596	-16.7	600.731	108.550	6.512
C02b	Shen_92298609_36250864_95741143	0.916	-7.347	-22.2	600.731	108.550	6.512
C03	Shen_92298609_50382533_8869305	0.916	-7.074	-5.6	611.675	130.480	4.944

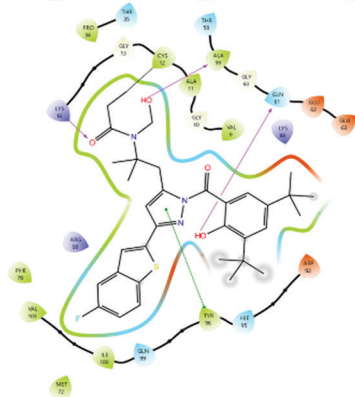
Abbreviations: TPSA: Topological polar surface area.

candidates with the clinically approved drug Sotorasib using metrics such as RMSD and MM/GBSA to assess their binding strength. The 100 ns MD simulations revealed that, apart from KRAS-C02a and KRAS-C03, all complexes maintained an average RMSD value ranging from 0.2 to 0.4 nm, indicating a relatively stable simulation completion that was closely matched with their initial conformations (Figure 3). Furthermore, KRAS-C01 was unable to maintain its stability in subsequent 300 ns MD simulations (Figure 3). Complexes such as KRAS(G12C)-C02b and KRAS(G12C)-Sotorasib, on the other hand, successfully completed the simulation with RMSD values ranging from 0.2 to 0.4 nm (Figure 3). Continuing the 500 ns MD simulation, KRAS(G12C)-C02b maintained its stability until around 450 ns when its RMSD ranged from 0.3 to 0.6 nm near the end of the simulation at 500 ns (Figure 3). Of note, KRAS(G12C)-Sotorasib maintained an RMSD

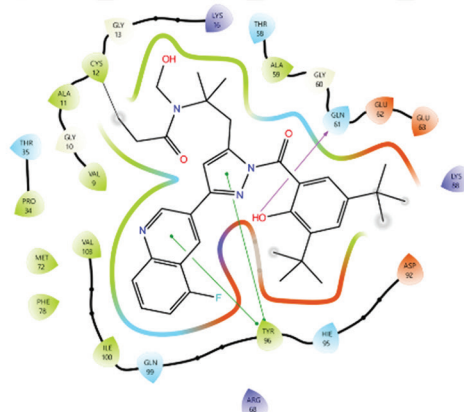
value ranging from 0.2 to 0.4 nm throughout the entire simulation. Thus, based on the RMSD of the ligand after least square fitting to the protein backbone, KRAS(G12C)-C02b exhibits a therapeutic candidate attribute comparable to the reference complex, KRAS(G12C)-Sotorasib, in that it maintains binding stability and shows no evidence of dissociation from the binding site.

To understand the fluctuations at the per residue level upon ligand binding, root mean-square fluctuation (RMSF) calculations were performed. Consistent with previously published findings,<sup>26,31,46</sup> it was observed that, in contrast to the entire protein, fluctuations were predominantly concentrated around switch-I (residues 30 – 38) and switch-II (residues 60 – 76). In Figure 4A, high peaks were noted in the RMSF graph within the region encompassing switch-I and switch-II loops in both complexes. Furthermore, hydrogen bond occupancy,

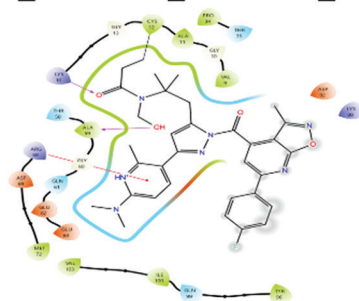
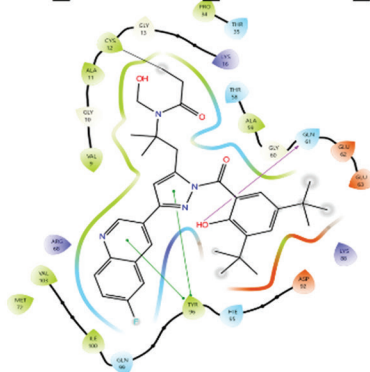
C01 (Shen\_92298609\_93443091\_95741143)



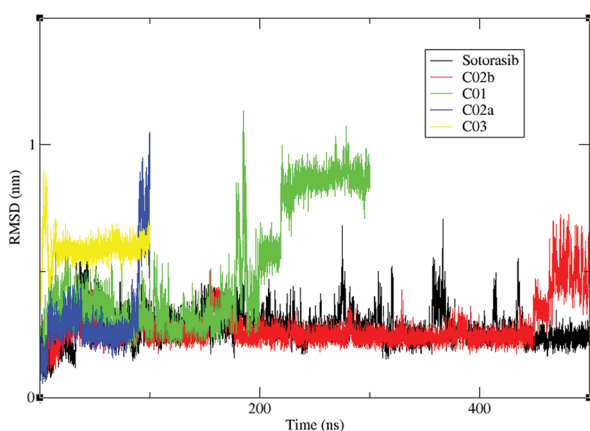
C02a (Shen\_92298609\_44473994\_95741143)



C02b (Shen\_92298609\_36250864\_95741143) C03 (Shen\_92298609\_50382533\_8869305)



**Figure 2.** Amino acid interactions of the top-ranking compounds with *Kirsten rat sarcoma viral oncogene homolog*(G12C).



**Figure 3.** Lig-fit-prot RMSD graph spectrums of Sotorasib, KRAS-C01, KRAS-C02a, KRAS(G12C)-C02b, KRAS-C03, and KRAS(G12C)-Sotorasib in complex with KRAS(G12C).

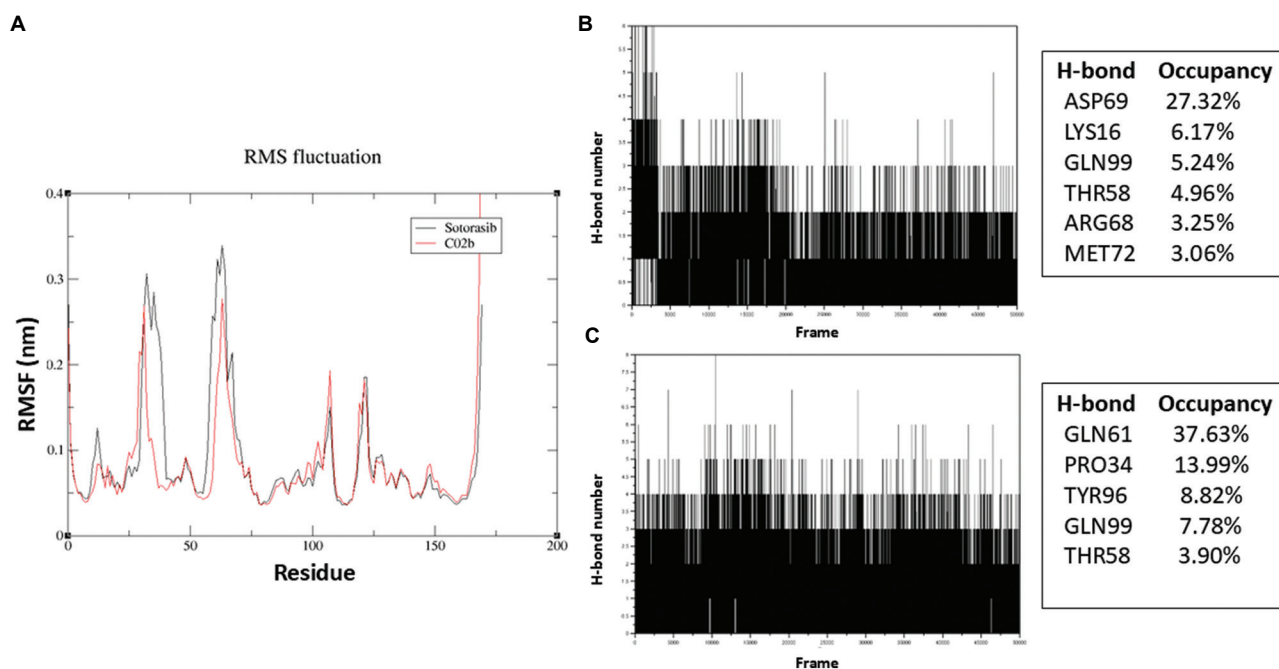
Abbreviations: RMSD: Root-mean-square deviation; KRAS: *Kirsten rat sarcoma viral oncogene homolog*.

quantified as the average number of hydrogen bonds per time frame, was also computed for these key residues. The percentages of hydrogen bond occupancies are detailed

in Figures 4B and C. The elevated rate of hydrogen bond formation between C02b and GLN61, as well as Sotorasib and ASP69, signifies a robust and dependable hydrogen-bonding profile for these interactions.

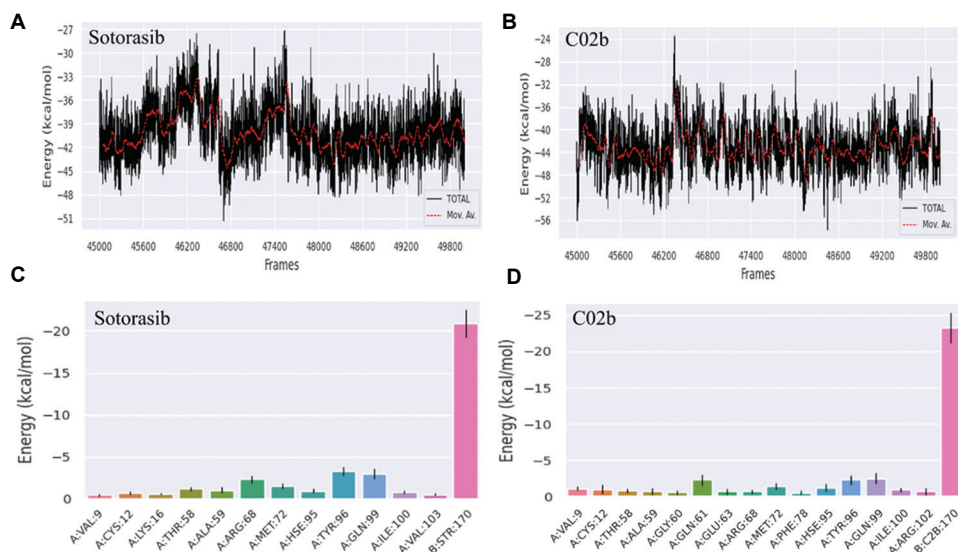
### 3.4. MM/GBSA calculation for evaluating binding free energy (dG)

Afterward, binding free energy (dG) was calculated using gmx\_MMPBSA, and the MM/GBSA dG distribution graph was generated for the last 50 ns of the 500 ns MD simulation. For KRAS(G12C)-Sotorasib, the average dG was  $-41$  kcal/mol (Figure 5A), whereas for KRAS(G12C)-C02b, this value was  $-44$  kcal/mol (Figure 5B). In addition, the Delta TDC (Total DeComposition) values, illustrating the residues contributing to the binding free energy, were investigated. During the interaction with KRAS G12C-Sotorasib, TYR96, and GLN99 were found to contribute significantly to the binding energy, followed by ARG68 and MET72 (Figure 5C). Examination of the Delta TDC values for KRAS(G12C)-C02b revealed that, like in the reference complex, TYR96 and GLN99 contributed most significantly to the binding free energy, followed by GLN61 and MET72,



**Figure 4.** (A) RMSF profiles of KRAS(G12C) in complex with reference and hit compounds. H-bonding interactions of (B) KRAS(G12C)-Sotorasib and (C) KRAS(G12C)-C02b.

Abbreviations: RMSF: Root-mean-square fluctuation; H-bond: Hydrogen bond; KRAS: *Kirsten rat sarcoma viral oncogene homolog*.

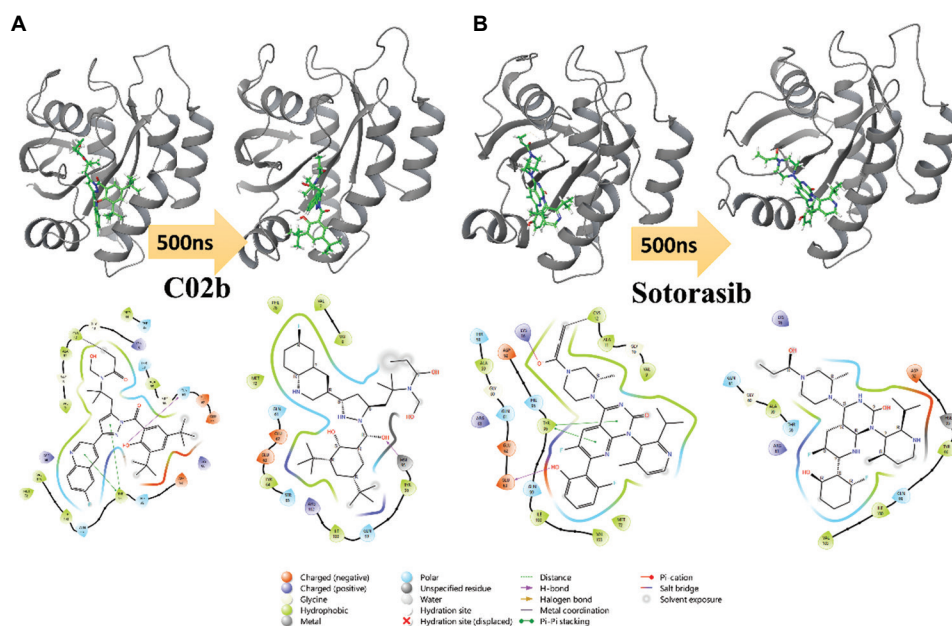


**Figure 5.** MM/GBSA dG distribution graph obtained for the last 50 ns of the 500 ns MD simulation for (A) Sotorasib, (B) C02b; and Delta total decomposition values for (C) STR: Sotorasib, (D) C2B: C02b.

Abbreviations: MM/GBSA: Molecular mechanics with generalized born surface area; MD: Molecular dynamics.

with a minor contribution from HIS95 (Figure 5D). It is worth reiterating that the interaction with GLN61 observed in the covalent docking results persisted throughout the 500 ns simulation. Both the KRAS(G12C)-C02b (Figure 6A) and KRAS(G12C)-Sotorasib (Figure 6B) complexes remained largely stable throughout the 500 ns trajectory, with minimal

deviation from the targeted binding site, maintaining their initial positions. These findings suggest that C02b exhibits binding interactions similar to the reference FDA-approved drug Sotorasib, with highly preserved interactions. Further research into C02b, which shares similarities with Sotorasib, could potentially serve as an alternative.



**Figure 6.** A schematic 2-D and 3-D representation of detailed ligand atom interactions with the protein residues at 0 ns and 500 ns. (A) C02b and (B) Sotorasib.

### 3.5. PCA and DCCM analysis

PCA was performed on the atomic backbone ( $C_{\alpha}$  position) using three configurations: First principal component (PC1), second principal component (PC2), and third principal component (PC3). The three most representative principal components cumulatively accounted for approximately 50% of all variances among the least correlated components across all 2000 frames analyzed for KRAS(G12C)-ligand complexes (Figures 7–10).

In the PCA, the PC1 dominated the overall variance, accounting for more than a third of the total variance (25.45% for Sotorasib and 20.01% for C02b). PC2 accounted 12.21% for Sotorasib and 14.1% for C02b, while PC3 exhibited the lowest variability, with 5.67% for Sotorasib and 6.28% for C02b. Together, the first three components represented 43.3% and 40.5% of the total variance for Sotorasib and C02b, respectively (Figures 7 and 9).

To further elucidate the impact of Sotorasib on the structure of KRAS(G12C), RMSF analysis was performed to compare the flexibility of the two systems. Higher fluctuation peaks were observed in the switch-I (residues 30 – 38) and switch-II (residues 60 – 76) regions of the KRAS(G12C)-Sotorasib complex (Figure 8), consistent with previous observations of dynamic structural features in those regions.<sup>46</sup> However, for KRAS(G12C)-C02b, RMSF values showed a significant decrease, especially in the switch-II region (Figure 10). The 3D structures of

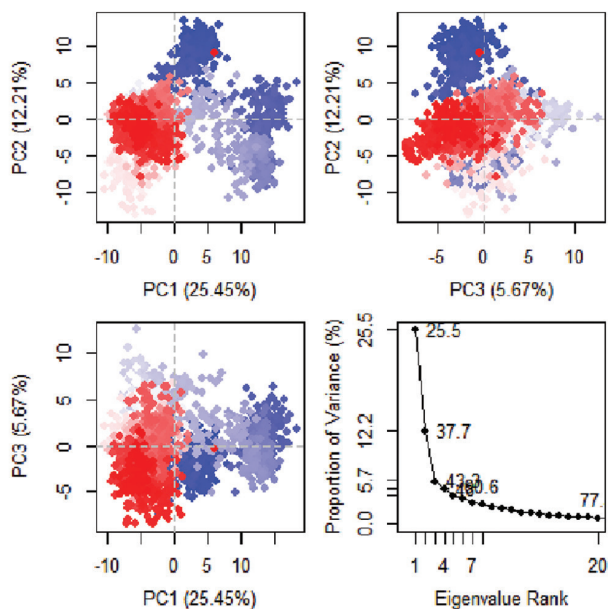
KRAS(G12C)-Sotorasib and KRAS(G12C)-C02b revealed a noticeable reduction in flexibility in the two switch regions for the structure of KRAS(G12C)-C02b compared to the reference.

The correlated conformational motions of the KRAS(G12C)-Sotorasib and KRAS(G12C)-C02b complexes were examined through DCCM analysis (Figure 11). In this analysis, regions displaying high positive values (depicted in red) indicate a strong correlation in the movement of residues in the same direction. Conversely, negative regions (depicted in blue) signify robust anti-correlated motion, where residues move in opposite directions. The color intensity along the diagonal reflects the degree of movement for the corresponding atoms.

Within the reference and C02b hit complex systems, the KRAS(G12C)-Sotorasib complex exhibits relatively stronger correlated motions compared to the KRAS(G12C)-C02b complex (Figure 11). Specifically, in the DCCM of the KRAS(G12C)-Sotorasib complex, the flexible region is observed moving in an anti-correlated manner with both switch-I (residues 30 – 38) and switch-II (residues 60 – 76), as well as displaying anti-correlated motion with the C12 residue and switch-I. This suggests that Sotorasib brings the binding pocket closer through these anti-correlated motions of the flexible regions, including switch-I, switch-II, and the C12 residue.

On the other hand, the DCCM of the KRAS(G12C)-C02b complex reveals the flexible regions moving in

an anti-correlated manner with switch-I and switch-II, although the dynamic movement is less pronounced

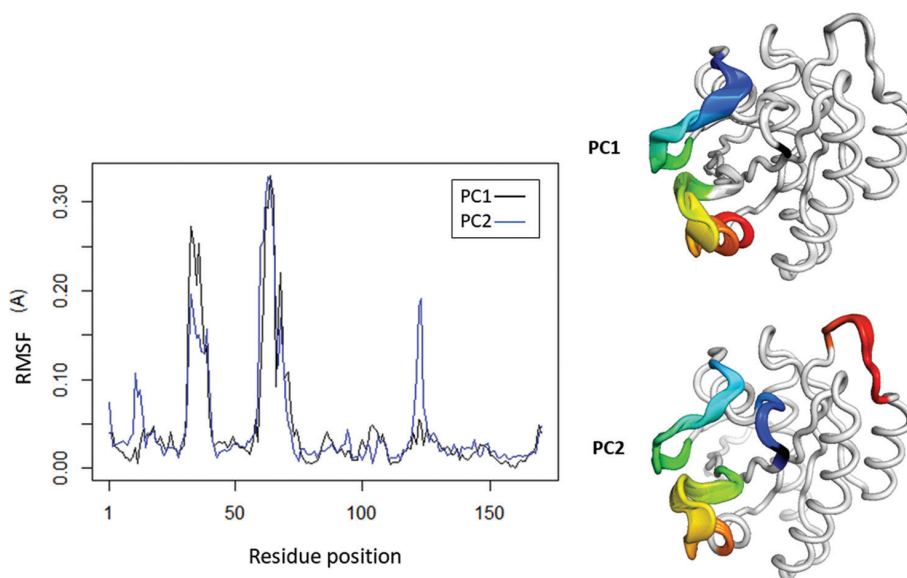


**Figure 7.** PCA dimension plots illustrating the first three principal components and displaying the cumulative percentages of variance covered by all eigenvectors for the *KRAS*(G12C)-Sotorasib structure. The color spectrum, transitioning from blue to red, delineates conformational alterations across the simulation. Blue dots signify the initial timesteps, white dots depict intermediate stages, and red dots denote the concluding timesteps. Abbreviations: PCA: Principal component analysis; *KRAS*: *Kirsten rat sarcoma viral oncogene homolog*.

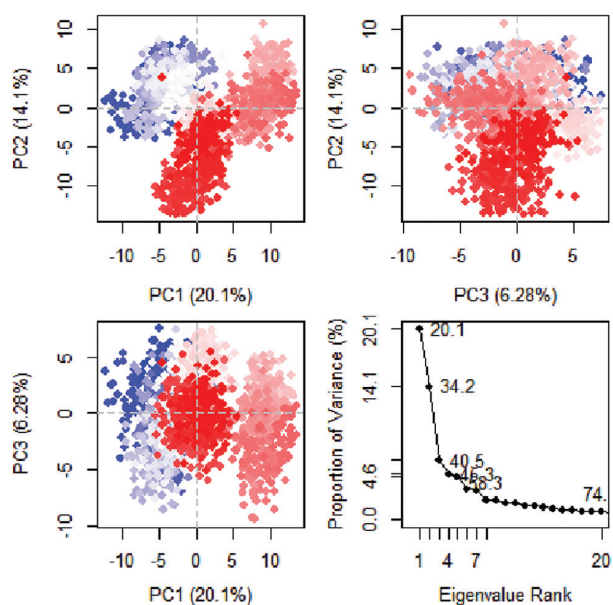
compared to the *KRAS*(G12C)-Sotorasib complex. Notably, there is no observed movement of the C12 residue in the *KRAS*(G12C)-C02b complex. These findings suggest a distinctive dynamic behavior in the correlated motions of the flexible loops and associated residues between the two complexes, highlighting the unique characteristics of Sotorasib and C02b interactions with *KRAS*(G12C).

The interaction analysis revealed notable differences between C02b and Sotorasib in their binding to the *KRAS*(G12C). C02b forms a strong and persistent interaction with GLN61 within the switch-II region, a key flexible loop implicated in *KRAS* conformational dynamics. In contrast, Sotorasib predominantly interacts with ARG68 and ASP69 in the same region. These interactions were validated through hydrogen bond occupancy analysis, which highlighted the stability and significance of the C02b-GLN61 interaction throughout the MD simulations. This distinct binding pattern of C02b suggests a potential alteration in the modulation of *KRAS*'s switch regions compared to Sotorasib, which may influence downstream signaling pathways.

Furthermore, PCA and DCCM analyses demonstrated unique dynamic behavior for the *KRAS*(G12C)-C02b complex. While Sotorasib induces a broader range of flexibility in the switch-II region, C02b exhibited reduced fluctuations in this region, potentially stabilizing the inactive GDP-bound state of *KRAS*. This stabilization is further supported by MM/GBSA binding free energy

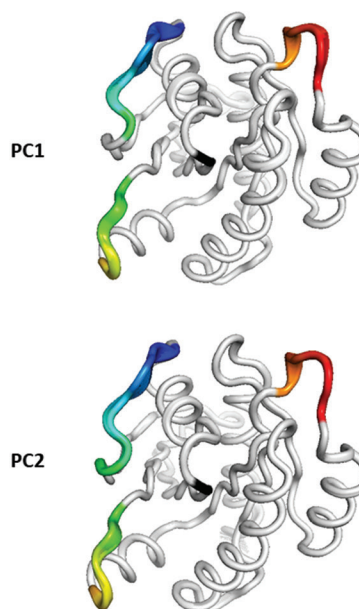
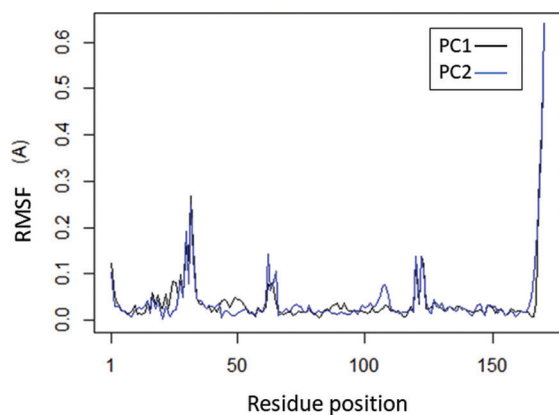


**Figure 8.** Flexibility analysis of the *KRAS*(G12C)-Sotorasib structure. The left side presents the RMSF values of the C $\alpha$  atoms within the system, while the right side displays ribbon representations of the PC1 and PC2 structures. Flexible regions are highlighted with a rainbow color scheme, providing a visual cue to the dynamic nature of these segments. Notably, the C12G residue is depicted in black, and the structures were visualized using PyMOL. Abbreviations: RMSF: Root mean-square fluctuation; *KRAS*: *Kirsten rat sarcoma viral oncogene homolog*; PC1: First principal component; PC2: Second principal component.



**Figure 9.** PCA dimension plots illustrating the first three principal components and displaying the cumulative percentages of variance covered by all eigenvectors for the *KRAS*(G12C)-C02b structure. The color spectrum, transitioning from blue to red, delineates conformational alterations across the simulation. Blue dots signify the initial timesteps, white dots depict intermediate stages, and red dots denote the concluding timesteps.

Abbreviations: PCA: Principal component analysis; *KRAS*: *Kirsten rat sarcoma viral oncogene homolog*.



**Figure 10.** Flexibility analysis of *KRAS*(G12C)-C02b structure. The left side presents the RMSF values of  $C^\alpha$  atoms within the system, while the right side displays ribbon representations of the PC1 and PC2 structures. Flexible regions are highlighted with a rainbow color scheme, providing a visual cue to the dynamic nature of these segments. Notably, the C12G residue is depicted in black, and the structures were visualized in PyMOL.

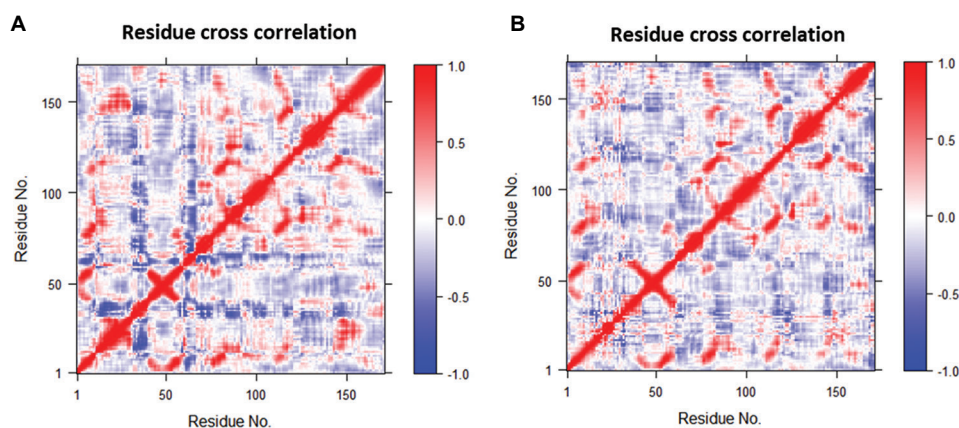
Abbreviations: RMSF: Root mean-square fluctuation; *KRAS*: *Kirsten rat sarcoma viral oncogene homolog*; PC1: First principal component; PC2: Second principal component.

calculations, which confirm the preservation of strong binding interactions for both compounds but with distinct residue-specific contributions.

#### 4. Discussion

In this study, we present potential *KRAS*(G12C) inhibitors identified through a comprehensive approach involving covalent docking-based virtual screening and MD simulations. Our strategy started with a similarity search based on the scaffold of Sotorasib, which successfully identified 174 molecules from a vast pool of compounds. Subsequently, covalent docking-based virtual screening was performed, revealing promising *KRAS*(G12C) protein binders, namely, C01, C02a, C02b, and C03. To assess the stability of the identified complexes and putative binders, MD simulations were employed. Significantly, among the candidates, C02b demonstrated exceptional stability as an inhibitor targeting the *KRAS*(G12C) during the 500 ns MD simulation, comparable to the reference.

In a previous study, it was found that Sotorasib (AMG 510) bound to the HIS95 groove and created a connected network of 25 ligand-protein interactions through van der Waals forces.<sup>29</sup> This network spanned from the backbone of helix 2 (involving HIS95 and TYR96) to the backbone of the flexible switch-II loop. According to the hydrogen bond analysis performed with VMD and decomposition analysis



**Figure 11.** The post-MD simulation time-correlated DCCM plot for the *KRAS*(G12C) (A) Sotorasib and (B) *KRAS*(G12C)-C02b complexes. Blue color indicates anti-correlation and red positive correlation.

Abbreviations: DCCM: Dynamic cross-correlation matrix; MD: Molecular dynamics; *KRAS*: *Kirsten rat sarcoma viral oncogene homolog*.

using Delta TDC values in our study, the key interaction in C02b binding was observed with the GLN61 residue located in the switch-II loop. For Sotorasib, interactions were observed with ARG68 and ASP69 residues in the same loop. In addition, it was found that the TYR96 residue was crucial for both the reference and C02b interactions.

The distinct interaction of C02b with GLN61 in the switch-II region, compared to Sotorasib's interactions with ARG68 and ASP69, likely contributes to the observed differences in conformational dynamics and flexibility of the *KRAS*(G12C) complexes. Switch-II is a critical region for nucleotide exchange and downstream effector binding, and its modulation directly influences *KRAS* signaling activity. By stabilizing GLN61, C02b may better inhibit *KRAS* activation by preventing the conformational changes required for GTP loading and effector interactions. These molecular differences may translate into unique cellular outcomes. Reduced switch-II flexibility, as observed with C02b, could lead to more robust inhibition of *KRAS*-driven downstream signaling pathways, potentially altering cellular proliferation, survival, and apoptotic processes. This distinct mechanism of action highlights C02b as a promising candidate for further preclinical investigation, offering a potential advantage in overcoming resistance mechanisms associated with existing *KRAS* inhibitors like Sotorasib. Further experimental validation will be essential to confirm these findings. Comparative studies on the effects of C02b and Sotorasib in cellular models of *KRAS*(G12C)-driven cancers could provide critical insights into their impact on downstream signaling and therapeutic efficacy.

Moreover, the structural diversity of *KRAS*(G12C) inhibitors remains narrow, leaving substantial room for the development of alternative scaffolds. Such scaffolds could offer improved pharmacokinetics, enhanced binding

interactions, and broader application to different tumor types harboring the *KRAS*(G12C) mutation. For instance, novel compounds with distinct binding modes may reduce side effects, enhance tolerability, or bypass resistance mechanisms arising in *KRAS* proteins with secondary mutations or altered conformations. These challenges highlight the urgent need for novel inhibitors that can either supplement or outperform existing treatments. Compounds like C02b, which demonstrate stable binding and distinct interaction profiles in computational analyses, hold promise for addressing these gaps. These novel compounds could expand the arsenal of therapeutic options, potentially improving outcomes when used alone or in combination with existing therapies.

While the current study primarily focuses on computational and simulation-based approaches, we have outlined the necessity of such biochemical analyses in the discussion section. Future work will include experimental assessments of compound selectivity using GTP-loaded *KRAS*(G12C) and wild-type *KRAS* proteins. These experiments will involve evaluating GDP/GTP exchange inhibition, covalent binding to Cys12, and downstream signaling modulation. We appreciate the reviewer's suggestion and recognize its importance for validating the therapeutic potential of C02b.

The catalytic domain of the RAS family maintains a relatively conserved conformation, apart from the two switch regions (switch-I and switch-II).<sup>47</sup> The conformations of these switch regions may play a crucial role in mediating interactions with other proteins.<sup>47</sup> Therefore, post-MD simulations analyses were carried out to evaluate the stability and flexibility of the two switches in the Sotorasib and C02b complexes targeting *KRAS*(G12C). The RMSD analysis determined the structural stability, while RMSF focused on flexibility, particularly in critical

functional regions. Throughout the 500 ns MD simulation, the KRAS(G12C)-C02b complex maintained low RMSD values and stability, similar to the reference complex. High peaks were observed in RMSF graph for the region containing switch-I and switch-II loops in both complexes. On the other hand, PCA identified dominant modes of motion, capturing a substantial portion of the total variance and DCCM elucidated correlated conformational motions within the complexes. The dynamic interaction of KRAS(G12C) and Sotorasib through a 10 microseconds MD simulation found that Sotorasib increased flexibility in the switch-I and Switch-II regions,<sup>46</sup> However, a previous study observed a decrease in the fluctuation of the switch-I region during an 800 ns MD simulation.<sup>48</sup> The disparity in these simulation results may be attributed to different simulation settings. In our RMSF analysis with PCA (Figures 8 and 10), through a 500 ns MD simulation of the dynamic interaction of KRAS(G12C) and Sotorasib, it was observed that Sotorasib did not stabilize KRAS switches in the complex.<sup>49</sup>

## 5. Conclusion

Taken together, our findings emphasize the distinctive stabilizing effects of C02b compared to Sotorasib on the conformational dynamics of KRAS(G12C), which may contribute to the development of potential inhibitors and highlight the importance of C02b in the development of KRAS(G12C) inhibitors in the preclinical setting.

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## Conflict of interest

The authors declare that they have no competing interests.

## Author contributions

*Conceptualization:* Safiye Merve Bostancioglu, Ahmet Acar

*Formal analysis:* Safiye Merve Bostancioglu

*Investigation:* Safiye Merve Bostancioglu, Ahmet Acar

*Methodology:* Safiye Merve Bostancioglu, Ahmet Acar

*Writing – original draft:* Safiye Merve Bostancioglu, Ahmet Acar

*Writing – review & editing:* Safiye Merve Bostancioglu, Ahmet Acar

## Ethics approval and consent to participate

Not applicable.

## Consent for publication

Not applicable.

## Availability of data

Data are available from the corresponding author upon reasonable request.

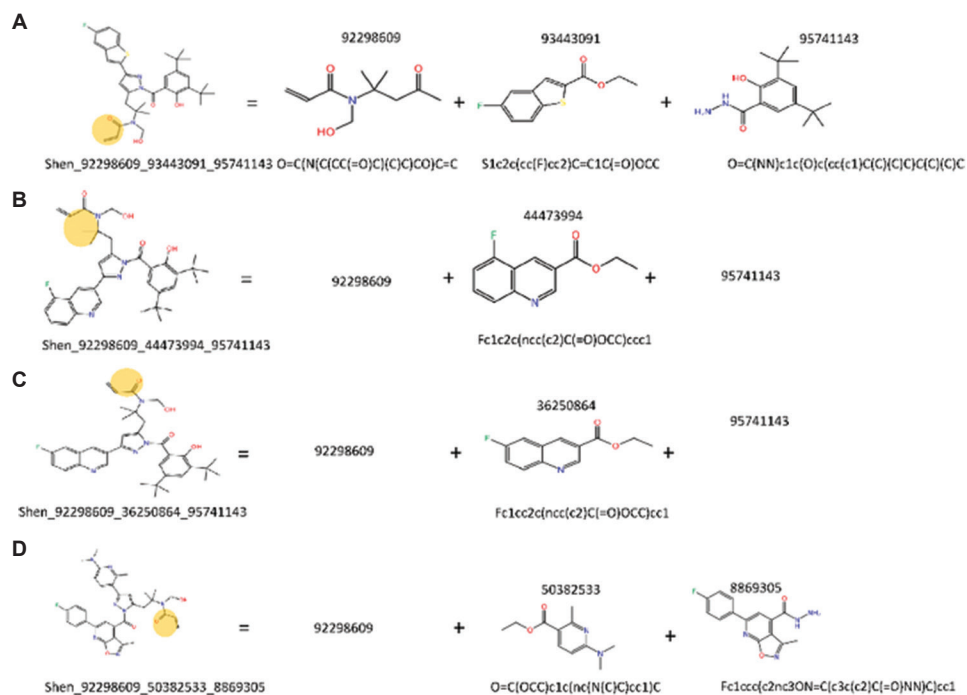
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## Appendices



**Figure A1.** The acrylamide warheads and building blocks of compounds. (A) C01, (B) C02a, (C) C02b, and (D) C03.

Table A1. 174 compounds that have structural similarities to Sotorasib

No	Name
1	rxn501b__EMOL916035__EMOL4353066__EMOL45478818
2	rxn501b__EMOL485662__EMOL4353066__EMOL45478818
3	rxn508__EMOL916035__EMOL37154489__EMOL724744
4	rxn508__EMOL916035__EMOL37154489__EMOL4854098
5	rxn508__EMOL916035__EMOL37154489__EMOL300131641
6	rxn508__EMOL916035__EMOL37154489__EMOL9779925
7	rxn508__EMOL916035__EMOL37154489__EMOL2863515
8	rxn501b__EMOL882112__EMOL4353066__EMOL45478818
9	rxn508__EMOL916035__EMOL50157604__EMOL724744
10	rxn508__EMOL916035__EMOL2862529__EMOL4854098
11	rxn508__EMOL916035__EMOL50157604__EMOL2863515
12	rxn508__EMOL916035__EMOL50157604__EMOL300131641
13	rxn508__EMOL916035__EMOL2862529__EMOL300131641
14	rxn508__EMOL916035__EMOL50157604__EMOL9779925
15	rxn508__EMOL916035__EMOL2862529__EMOL2863515
16	rxn508__EMOL916035__EMOL50157604__EMOL4854098
17	rxn508__EMOL916035__EMOL2862529__EMOL9779925
18	rxn508__EMOL916035__EMOL2862529__EMOL724744
19	rxn508__EMOL916035__EMOL37154489__EMOL30070316
20	rxn508__EMOL916035__EMOL37154489__EMOL36725275
21	rxn508__EMOL916035__EMOL50157604__EMOL36725275
22	rxn508__EMOL916035__EMOL2862529__EMOL36725275
23	rxn508__EMOL916035__EMOL50157604__EMOL30070316
24	rxn508__EMOL916035__EMOL2862529__EMOL30070316
25	rxn501b__EMOL916035__EMOL4353066__EMOL48597142
26	rxn501b__EMOL916035__EMOL4353066__EMOL45478663
27	rxn508__EMOL916035__EMOL37154489__EMOL44474549
28	rxn508__EMOL916035__EMOL50157604__EMOL44474549
29	rxn508__EMOL916035__EMOL2862529__EMOL44474549
30	rxn508__EMOL916035__EMOL37154489__EMOL43633603
31	rxn508__EMOL916035__EMOL2862529__EMOL43633603
32	rxn508__EMOL916035__EMOL50157604__EMOL43633603
33	rxn508__EMOL916035__EMOL37154489__EMOL49844674
34	rxn508__EMOL916035__EMOL37154489__EMOL1522728
35	rxn508__EMOL916035__EMOL37154489__EMOL11554196
36	rxn508__EMOL916035__EMOL37154489__EMOL1885674
37	rxn508__EMOL916035__EMOL37154489__EMOL42785649
38	rxn508__EMOL916035__EMOL37154489__EMOL11488490
39	rxn508__EMOL916035__EMOL89945076__EMOL724744

(Cont'd...)

Table A1. (Continued)

No	Name
40	rxn508__EMOL916035__EMOL89945076__EMOL9779925
41	rxn508__EMOL916035__EMOL89945076__EMOL4854098
42	rxn508__EMOL916035__EMOL89945076__EMOL2863515
43	rxn508__EMOL916035__EMOL89945076__EMOL300131641
44	rxn508__EMOL916035__EMOL37154489__EMOL904614
45	rxn508__EMOL916035__EMOL2862529__EMOL42785649
46	rxn508__EMOL916035__EMOL50157604__EMOL49844674
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53	rxn508__EMOL916035__EMOL2862529__EMOL1522728
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64	rxn508__EMOL485662__EMOL37154489__EMOL300131641
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76	Shen__92298609__48620700__44152703
77	Shen__92298609__93443091__44152703
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83	Shen_92298609_49239556_44152703
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85	Shen_92298609_44473994_44152703
86	Shen_92298609_44473403_44152703
87	Shen_92298609_44473381_44152703
88	Shen_92298609_48573899_44152703
89	Shen_92298609_48552545_107053254
90	Shen_92298609_48552545_81813113
91	Shen_92298609_32456150_44152703
92	Shen_92298609_36745965_44152703
93	Shen_92298609_830425_44152703
94	Shen_92298609_42824050_44152703
95	Shen_92298609_42817358_44152703
96	Shen_92298609_25041346_44152703
97	Shen_92298609_97662126_29552264
98	Shen_92298609_36249205_29552264
99	Shen_92298609_48552545_8869305
100	Shen_92298609_36249205_107053254
101	Shen_92298609_97662126_107053254
102	Shen_92298609_56970423_44152703
103	Shen_92298609_49823880_44152703
104	Shen_92298609_36249205_81813113
105	Shen_92298609_97662126_81813113
106	Shen_92298609_50382533_29552264
107	Gray_107373053_6886343_107509511
108	Shen_92298609_53885480_44152703
109	Shen_92298609_53885502_44152703
110	Shen_92298609_44473387_44152703
111	Shen_92298609_53885486_44152703
112	Shen_92298609_42816206_44152703
113	Shen_92298609_36828355_44152703
114	Shen_92298609_53890255_44152703
115	Shen_92298609_50382533_107053254
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Table A1. (Continued)

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124	Shen_92298609_50272909_107053254
125	Shen_92298609_106068876_44152703
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128	Shen_92298609_43045611_44152703
129	Shen_92298609_50272909_81813113
130	Shen_92298609_48620700_95741143
131	Shen_92298609_48620564_95741143
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136	Shen_92298609_97662126_8869305
137	Shen_92298609_81802090_29552264
138	Shen_92298609_76750201_29552264
139	Shen_92298609_106978508_29552264
140	Shen_92298609_95759057_29552264
141	Shen_92298609_48552545_50495850
142	Shen_92298609_93443091_95741143
143	Shen_92298609_93443091_96426646
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145	Shen_92298609_106083795_95741143
146	Shen_92298609_106083795_96426646
147	Shen_92298609_106083765_96426646
148	Shen_92298609_81802090_107053254
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150	Shen_92298609_76750201_107053254
151	Shen_92298609_49253559_44152703
152	Shen_92298609_1451473_44152703
153	Shen_92298609_49253379_44152703
154	Shen_92298609_48552545_2066545
155	Shen_92298609_95759057_107053254
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Table A1. (Continued)

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