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Gaussian field-based 3D-QSAR and molecular simulation studies to design potent pyrimidinesulfonamide hybrids as selective BRAF^{V600E} inhibitors†

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The "RAS-RAF-MEK-ERK" pathway is an important signaling pathway in melanoma. BRAF^{V600E} (70–90%) is the most common mutation in this pathway. BRAF inhibitors have four types of conformers: type I (α C-IN/ DFG-IN), type II (aC-IN/DFG-OUT), type I1/2 (aC-OUT/DFG-IN), and type I/II (aC-OUT/DFG-OUT). Firstand second-generation BRAF inhibitors show resistance to BRAF^{V600E} and are ineffective against malignancies induced by dimer BRAF mutants causing 'paradoxical' activation. In the present study, we performed molecular modeling of pyrimidine-sulfonamide hybrids inhibitors using 3D-QSAR, molecular docking, and molecular dynamics simulations. Previous reports reveal the importance of pyrimidine and sulfonamide mojeties in the development of BRAF^{v600E} inhibitors. Analysis of 3D-QSAR models provided novel pyrimidine sulfonamide hybrid BRAF^{V600E} inhibitors. The designed compounds share similarities with several structural moieties present in first- and second-generation BRAF inhibitors. A total library of 88 designed compounds was generated and molecular docking studies were performed with them. Four molecules (T109, T183, T160, and T126) were identified as hits and selected for detailed studies. Molecular dynamics simulations were performed at 900 ns and binding was calculated. Based on molecular docking and simulation studies, it was found that the designed compounds have better interactions with the core active site [the nucleotide (ADP or ATP) binding site, DFG motif, and the phospho-acceptor site (activation segment) of BRAF^{V600E} protein than previous inhibitors. Similar to the FDA-approved BRAF^{V600E} inhibitors the developed compounds have [aC-OUT/DFG-IN] conformation. Compounds T126, T160 and T183 interacted with DIF (Leu505), making them potentially useful against BRAF^{V600E} resistance and malignancies induced by dimer BRAF mutants. The synthesis and biological evaluation of the designed molecules is in progress, which may lead to some potent BRAF^{V600E} selective inhibitors.

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1. Introduction

Melanomas arise from uncontrolled proliferation of melanocytes, which are found throughout the body and determine the color of the skin, eye, inner ear, iris, rectum, leptomeninges, and basal layer of the hair bulb matrix.^{1,2} For all sight of skin cancer melanoma account for only 4% but have death rate higher than 50%.³ In 2020, 19 292 789 new cases were estimated worldwide for all cancers, including 324 635 (1.7%) melanoma cases. The total number of cancer deaths was 9 958 133, of which 57 043 (0.6%) were due to melanoma.⁴

The "RAS-RAF-MEK-ERK" pathway is the main component of a signal transduction pathway that regulates gene expression and supports mammalian cells' proliferation and survival.⁵ The RAS-RAF-MEK-ERK pathway is responsible for the activation of BRAF and regulates cell proliferation and differentiation.⁶ Activation of the small G protein Ras leads to activation of BRAF.⁷ BRAF activates the MAP kinase/ERK signaling pathway for normal cell control and proliferation.⁸ Normal and oncogenic BRAF signaling pathway is depicted in Fig. 1a. BRAF is the most frequently mutated protein kinase in human melanomas.⁹ The prevalence of BRAF mutations in human malignancies varies, with melanoma having between 70–90% of all mutations,^{10,11} 5–20% in colon cancer,¹² 30–50% in thyroid cancer, ~10% in papillary thyroid cancer,¹³ 5–20% in colorectal cancer, and 5–30% in ovarian cancer.¹⁴ Among BRAF gene mutations, the BRAF^{V600E} gene is the most common mutation in melanoma, accounting for 70–90%.^{15,16} It occurs at position 600 with a single point mutation (Val600 \rightarrow Glu), in which the hydrophobic valine is replaced by polar, hydrophilic glutamic acid.¹⁷ BRAF^{V600WT} is the second most common mutation of BRAF in this BRAF^{V600K}, (10–30%).¹⁸ The rarer codon mutations BRAF^{V600R,V600D} (6–3%), BRAF^{V600E2,600-MandV600G} (<1%)¹⁹ occur in a small percentage of patients and are considered 8% to 10% of all BRAF mutations.²⁰

1.1 BRAF kinase conformations

The BRAF kinase domain contains two lobes: small N and large C-terminal lobe, which are connected by a catalytic cleft or flexible hinge region. The catalytic site is located in the cleft



Fig. 1 (a) Normal and oncogenic BRAF signaling pathway notes: normal activation of RAS-RAF pathway shows normal cell proliferation and survival. Oncogenic BRAF signaling causes an increase in cell proliferation and survival from benign nevus to metastatic melanoma (b) V600E mutation in BRAF protein structure (c) BRAF inhibitor binding mode with conformations (α C-helix and DFG motif).

Table 1 Important amino acid sequence in BRAF protein

BRAF protein region	Amino acid	sequence	Different binding pocket and C-lobe	Amino acid sequence	
CR1	t1 150–290 RBD 155–227 Glycine-rich ATP-phosphate-binding CRD 234–280 loop (P-loop) and ribose pocket		463-471		
CR2	360-375		αC-Helix	-	492-504
CR3 (protein kinase domain)	457-717		RAF selective pocket ^a	DIF	504-511
2			-	DFG motif	594-596
			Gatekeeper residue		T529
			Hinge region (adenine region)		530-535
			Catalytic loop	- ,	574-581
			Activation segment		599-623

between two lobes and contains the ATP-binding site. This region is divided into a front cleft (containing the ATP-binding site), a gate area, and a back cleft. The small N-terminal domain lobe has an α C-helix, predominantly antiparallel β -sheet anchors, a glycine-rich ATP-phosphate-binding loop (P-loop), and ATP orients. The large C-terminal lobe consists of α -helices.

At the interface of the N-and C-terminal lobes, the nucleotide (ADP or ATP) binding site, the magnesium binding site (DFG [Asp594–Phe595–Gly596] motif), and the phospho-acceptor site [activation segment (AS)] are located near the DFG motif, which are three major active sites.^{8,21}

The ATP pocket consists of the following main regions. Adenine region (1) is similar to the hinge region. Sugar region (2) is where ATP ribose is located, while other three regions include hydrophobic (3), type-I (4) and type-II (5) regions. The solvent-accessible region is also part of the ATP region, but generally, ATP is not located in this region.^{22,23} The structural domain of the BRAF between the N-terminal and C-terminal lobes contains three conserved regions (CR): CR1, CR2, and CR3.²⁴ In humans, the BRAF protein kinase contains 766 amino acids, with key amino acid residues listed in Table 1.^{25,26} BRAF protein structure and different binding conformations (α C-helix and DFG) motif with inhibitors are depicted in Fig. 1b and c.

1.2 Different binding modes of BRAF protein with inhibitors

The DFG motif and the α C-helix represent an important structural element of kinases. In physiological kinase activation, the DFG motif plays a fundamental role as it shifts from the DFG-OUT (inactive or closed state occupying part of the ATPbinding site) to the DFG-IN (active or open state), where it moves towards the base of the α C-helix, which occupies part of the allosteric binding site and influences the catalytic process.

When the DFG motif is in the IN position, the Asp594 residue faces the active site of the kinase. In this condition, the glycinerich loop and the activation segment of the BRAF kinase are separated from one another, the hydrophobic connection between these two areas is disrupted. As a result, the catalytic cleft accessible to ATP. In this conformation, the inhibitors form interactions with the hinge region (ATP-binding site) by forming \sim 1–3 hydrogen bonds and hydrophobic interaction around the adenine regions of the ATP-binding site of the protein. The inhibitors also interact with other different regions of ATP, including hydrophobic regions adjacent to the adenine region, the ribose region, and solvent accessible region.

When the DFG motif is in the OUT position the aspartate residue (Asp594) displaced from the active site, therefore hydrophobic connection between the glycine rich loop and activation section of the BRAF kinase domain, which is brought near to one another. This contact renders ATP unavailable to the catalytic cleft of BRAF. DFG-OUT conformation, Asp594 residue supports allosteric movement extending outward from the ATP-binding site, and inhibitors bind to allosteric site that is close to the ATP binding pocket. The α C-helix near the ATP site is called the α C-IN position, the α C-helix away from the ATP site is called the α C-OUT position.²⁷⁻³³ Based on α C-helix and DFG motif, four structural types of inhibitors are currently being developed. (1) Type I, (α C-IN/DFG-IN). (2) Type II, (α C-OUT/DFG-OUT). (3) Type I_{1/2}, (α C-OUT/DFG-IN). (4) Type I/II, (α C-OUT/DFG-OUT).³⁴⁻³⁷

1.3 Pyrimidine-sulfonamide hybrids BRAF inhibitors

Dabrafenib mesylate type $I_{1/2}$, (α C-OUT/DFG-IN) inhibitor is an FDA-approved pyrimidine-thiazole based diphenyl sulfonamide derivative for the treatment of metastatic melanoma in patients with BRAF^{V600E} mutation.^{38,39} Dabrafenib was first developed by GlaxoSmithKline (GSK) under the name GSK2118436. It is an ATP-competitive inhibitor of BRAF kinases. Dabrafenib inhibits BRAF^{V600E} most potently but is less effective against BRAF^{V600WT}.40,41 In 2011, vemurafenib, a pyrolo-pyrimidine based diphenyl sulfonamide derivative⁴² of type $I_{1/2}$, (α C-OUT/ DFG-IN)43 received FDA approval for first oral BRAF^{V600E} inhibitor.44,45 Encorafenib (LGX-818), a pyrazolo-pyrimidine based phenyl sulfonamide derivative,³⁹ was approved by the FDA in June 2018 for the treatment of BRAF^{V600E/K} mutated cancer. Patients with metastatic melanoma can take a combination of encorafenib and binimetinib (inhibitors of BRAF and MEK).46,47 The FDA-approved first- and second-generation adenosine triphosphate (ATP) competitive BRAF inhibitors are



shown resistance against monomeric BRAF^{V600} species mutations and also ineffective against malignancies driven by dimeric BRAF mutants causing 'paradoxical' activation.⁴⁸ Preclinical studies of BRAF inhibitors in melanoma shown that the development of resistance is more complex compared to single mutations.⁴⁹ BRAF inhibitor resistance depends on epigenetic and transcriptomic mechanisms of MAPK/ERK or PI3K/Akt activation of receptor tyrosine kinases.⁵⁰

In 2019 Abdel-Maksoud et al. synthesized BRAF inhibitors possessing 5-(pyrimidin-4-yl)imidazo[2,1-b]thiazole sulfonamide containing scaffold,⁵¹ and in 2021, Ali et al. synthesized novel (imidazole-5-yl)pyrimidine-sulfonamide hybrids derivatives.52 In 2020 Ammar et al. synthesized imidazothiazole moiety containing pyrimidine-sulfonamide derivatives. The synthesized compounds showed good BRAF^{V600E} kinase inhibition values in in vitro experiments.53 Abdel-Maksoud MS, et al. synthesized a new series of 4-(1H-benzo[d]imidazole-1-yl) pyrimidin-2-amine linked sulfonamide derivatives in 2021. The IC50 of the synthesized compounds were evaluated against BRAF^{V600E}, wild-type BRAF, and CRAF and potential inhibitory activity was found.⁵⁴ In 2012, Li et al. synthesized a new series of ATP-competitive BRAF inhibitors based on aryl/heteroaryl and o-substitutions on the pyrazolo [1,5-a] pyrimidine scaffold sulfonamide derivative with highly cellular potency and BRAF selectivity.55 To date, the FDA has approved four drugs for the treatment of melanoma, three of which contain a sulfonamide and pyrimidine moiety respectively (Fig. 2). BRAF inhibitors having pyrimidine and sulfonamide moiety (1-51) are also depicted in Fig. 2.

2. Computational approach for designing of new pyrimidine– sulfonamide hybrid containing compounds as potent BRAF^{V600E} inhibitors

Literature reports indicate that pyrimidine and sulfonamide moieties play a very important role in BRAF mutant inhibitor development. The aim of this work is to investigate the structural requirements of pyrimidine–sulfonamide hybrid derivatives as inhibitors of BRAF^{V600E} mutant. Typically, 3D-QSAR investigations are employed for statistical analysis of SAR for a group of compounds with similar structures and to look into the characteristics of an interaction between a ligand and its target protein.⁵⁶

Molecular docking is used to understanding the structural relationship between a ligand and its target and predicting the best binding conformation.⁵⁷ The flexibility, changes in the structural conformation of the target binding site, and stability of a ligand-receptor complex suggested by molecular docking was studied in details by the use of molecular dynamics (MD) simulations.⁵⁸ As a result, it is thought that combining 3D-QSAR, molecular docking, and MD simulations provides an efficient way to examine how the ligand and receptor attach to one another. By utilizing 3D-QSAR, molecular docking, and MD simulations in the current study, we have successfully proven molecular modelling of inhibitors of pyrimidine–sulfonamide hybrids. To the best of our knowledge, no 3D-QSAR studies have been reported for these types of compounds.⁵⁹

In this study, we built more selective 3D-QSAR models based on Gaussian techniques and SAR of these was summarized. The results of the 3D-QSAR investigation were validated using molecular docking, and the interactions between proteins and ligands were examined. We applied MD simulations to understand the flexibility, structural conformational changes and to estimate the free energy of binding of mutant BRAF kinases with the designed compounds.

2.1 Materials and methods

2.1.1 Gaussian based 3D QSAR

2.1.1.1 Dataset collection. Structurally different pyrimidinesulfonamide hybrid derivatives (1-51) were selected from the literature, exhibiting a broad spectrum of BRAF^{V600E} inhibitory activities (IC₅₀ 1.20–1910 nM).^{51–54} The 2D structures of the pyrimidine–sulfonamide hybrid derivatives (1-51) were prepared using the Chem 2D program. The geometry of all these molecules was converted to a 3D structure using Schrodinger's Maestro 12.9. The following options were chosen: (i) the force field used was OPLS 2005, (ii) one low energy ring conformation per ligand was generated, (iii) all possible ionization states at biological pH (iv) depending on the number of chiral centers, probable tautomers and stereospecificity were considered when processing the 3D structure of the molecules with the LigPrep module.^{59,60}

2.1.1.2 Training and test set selection. The IC_{50} values of the data set were changed to negative logarithmic forms [pIC₅₀ = $-\log_{10}(IC_{50})$] to linearize the data. pIC₅₀ values between 5.719 and 8.921 were used in this study as dependent variable for the development of 3D-QSAR models. The dataset was divided into training and test (validation) set (70 : 30) (Table 2) using random selection tool present in 3D-Gaussian based QSAR of Maestro 12.9.^{59,60}

2.1.1.3 Alignment procedure. Correct alignment of optimized conformers from selected dataset in a fixed lattice is the most crucial need for successful generation of models. It depends on the correct relative placement of the molecules from the dataset. In this study, the molecules were aligned using the structure alignment tools (flexible shape-based alignment, common scaffold alignment, maximum common substructure, smarts) of Maestro 12.9 tools.^{60,61}

2.1.1.4 Generation of 3D QSAR model. Models were generated using a grid spacing of 0.1 Å and extending the grid by 3.0 Å beyond the limits of the training set using Maestro 12.9. The force field values were set as ignore force field within 2.0 Å. The truncated steric and electrostatic force field was preset to 30.0 kcal mol⁻¹. Variables were removed with a standard deviation (SD) <0.01 and the number of ligands to exclude from the crossvalidation was set as 1. Five features including steric, electrostatic, H-bond acceptor, H-bond donor, and hydrophobic field was considered for model buildup.^{60,62}

2.1.1.5 Partial least square (PLS) analysis. The partial least squares (PLS) were used to establish the 3D-QSAR model. In this various statistical parameters such as standard deviation (SD), non-cross-validated correlation coefficient (r^2), cross-correlation validation, ($r_{\rm CV}^2$), r^2 scramble, Fischer's test (*F*-test), variance

ratio (*p* value), root mean squared error (RMSE), Pearson-*r* values, leave-one-out (LOO) cross-validated correlation coefficient (q^2) and Pearson-*r* were considered for validation of the developed models. The quality of the developed models was examined by SD of regression < SD of actual activities (1), $1 > r^2 > 0.5$, $r_{\rm CV}^2 > 0.6$, $r_{\rm scramble}^2 < 0.6$, Fischer's test (*F*-test) value (as large as possible), *p* value (as less as possible), $q^2 > 0.5$, Pearson-*r* values, and RMSE close to 1. To evaluate the predictive ability of the QSAR models developed, pIC₅₀ values as dependent variables and Gaussian intensities as descriptors (independent variables) were used.^{60,63}

2.1.2 Molecular docking. The 3D structure of human BRAF kinase (hBRk) with V600E mutation was taken from the research collaboratory for structural bioinformatics (RCSB) protein database,⁶⁴ PDB ID 4XV2.⁶⁵ Dabrafenib (*N*-{3-[5-(2-aminopyrimidin-4-yl)-2-*tert*-butyl-1,3-thiazol-4-yl]-2-fluorophenyl}-

2,6-difluorobenzenesulfonamide) is bound in the catalytic pocket. Only the a chain of the kinase was retained, and dabrafenib and water molecules were removed. The hBRk was prepared for molecular docking with Chimera 1.14.66,67 All nonpolar hydrogens were merged, gasteiger charges were added to each atom, atom types were determined, and the final structure of the prepared receptor was saved in pdbqt file format. The center of the grid box was set at the position of the F39 atom of dabrafenib, with cartesian coordinates -0.4, -4.9, -21.6. The size of the box was 20 \times 20 \times 20 \mathring{A}^3 and both exhaustiveness and the number of modes were set to 100. The 3D structure of the ligands was determined by converting SMILES to 3D geometry and then optimizing the geometry in RDKit.68 The Auto Dock Tools 4 Python script prepare_ligand4.py⁶⁹ was used to prepare the ligands. A threshold of 4 kcal mol⁻¹ relative to the highest score was introduced to save docked conformations. All results were visually inspected, and the conformation with the lowest binding energy was retained. AutoDock Vina69 was used for molecular docking simulations. Dabrafenib was docked to the hBRk to validate our docking procedure.

2.1.3 Molecular dynamics simulations. The initial structures of the ligand-receptor complexes for the MD simulation were obtained by molecular docking experiment. The ligands were parametrized in the AMBER 20 Antechamber module⁷⁰ using the GAFF force field.⁷¹ For the kinase, AMBER ff19SB⁷² force field was employed. The PDB2PQR web-server73 was accessed to determine the protonation state of the side chains of the residues at physiological pH. The ligand-receptor complex was impregnated with pre-equilibrated TIP3P water molecules in a truncated octahedral periodic box. The minimum distance between the edges of the water box and the nearest atom of the complex was set at 12 Å. The system was neutralized by six Cl⁻ anions, followed by the addition of Na⁺ and Cl⁻ ions according to the recommendations of Machado and Pantano74 to achieve a neutral environment with a salt concentration of 0.15 M.

The minimization-heating-equilibration-production protocol was used. First, the system was subjected to geometry optimization with periodic boundary conditions in all directions. In 10 000 optimization cycles (4000 steepest descent +

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 Table 2
 Experimental and predicted BRAF^{V600E} inhibitory IC₅₀ values of pyrimidine-sulfonamide hybrid derivatives used in current study

S. No.	R ₁	х	n	R_2	$\mathrm{IC}_{50}\left(\mathrm{nM} ight)$	pIC ₅₀	Predicted pIC ₅₀ values (Gaussian)	Ref.
1	\square	F	1	ОН	5.620	8.250	7.802	52
2	Br	F	1	ОН	26.300	7.580	7.901	52
3	F	F	1	ОН	30.300	7.519	7.748	52
4	H ₃ C	F	1	ОН	8.680	8.061	7.858	52
5	но	F	1	ОН	2.490	8.604	8.436	52
6		Cl	1	ОН	4.250	8.372	7.824	52
7	Br	Cl	1	ОН	21.800	7.662	7.923	52
8	F	Cl	1	ОН	14.800	7.830	7.770	52
9	но	Cl	1	ОН	7.810	8.107	8.464	52
10	F F	_	2	F	22.300	7.652	7.440	51
11		_	3	F	89.000	7.051	7.376	51
12	F F	_	3	F	26.000	7.585	7.575	51
13	o	_	3	F	25.100	7.600	7.773	51

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S. No.	R ₁	x	п	R ₂	$IC_{50}(nM)$	pIC ₅₀	Predicted pIC ₅₀ values (Gaussian)	Ref.
14	CI	_	3	F	16.200	7.790	7.778	51
15	Br	_	3	F	9.300	8.032	7.778	51
16	H ₂ N	_	2	F	28.300	7.543	7.615	53
17	H ₂ N	_	2	CH ₃	59.100	7.228	7.403	53
18	H ₂ N	_	2	CF ₃	141.000	6.851	7.323	53
19	H ₂ N	_	2	OCH3	23.400	7.631	7.375	53
20	H ₂ N	_	2	CI	16.700	7.777	7.611	53
21	H ₂ N	_	2	Br	73.700	7.133	7.598	53
22	H ₂ N	_	2	F	12.700	7.896	8.112	53
23	H ₂ N	_	2		152.000	6.818	6.651	53
24	H ₂ N	_	3	F	13.500	7.870	8.067	53
25	H ₂ N	_	3	CF ₃	11.400	7.943	7.721	53

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Table 2 (Contd.)

S. No.	R ₁	Х	n	R ₂	IC_{50} (nM)	pIC ₅₀	Predicted pIC ₅₀ values (Gaussian)	Ref.
26	H ₂ N	_	3		20.400	7.690	8.326	53
27	H ₂ N	_	3		4.310	8.366	8.332	53
28	H ₂ N	_	3	F	6.210	8.207	8.286	53
29	H ₂ N	_	3		24.500	7.611	7.758	53
30	H ₂ N	_	3	OCH ₃	1.200	8.921	8.331	53
31	H ₂ N	_	3	Br	4.310	8.366	8.332	53
32	H ₂ N	_	3	F	6.210	8.207	8.286	53
33	F	_	3	OCH3	25.100	7.600	7.507	53
34	F	_	3	Br	9.300	8.032	7.741	53
35	O ₂ N	_	3	OCH3	131.0	6.883	7.454	53
36	O ₂ N	_	3	Br	20.0	7.699	7.453	53
37	O ₂ N	_	3	F	35.0	7.456	7.407	53

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S. No.	R ₁	Х	n	R_2	IC_{50} (nM)	pIC ₅₀	Predicted pIC_{50} values (Gaussian)	Ref.
38		_	2	_	1910.0	5.719	5.987	54
39	F	_	2	_	740.0	6.131	6.055	54
40	CI	_	2	_	820.0	6.086	6.066	54
41	Br	_	2	_	910.0	6.041	6.066	54
42	CF ₃	_	2	_	620.0	6.208	6.183	54
43	OCH3	_	2	_	980.0	6.009	6.043	54
44	CH ₃	_	2	_	790.0	6.102	5.993	54
45		_	3	_	1120.0	5.951	6.045	54
46	F	_	3	_	530.0	6.276	6.062	54
47	CI	_	3	_	790.0	6.102	6.056	54
48	Br	_	3	_	810.0	6.092	6.043	54
49	CF ₃	—	3	_	490.0	6.310	6.259	54

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Table 2 (Contd.)

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S. No.	R ₁	X	n	R_2	IC_{50} (nM)	pIC ₅₀	Predicted pIC ₅₀ values (Gaussian)	Ref.
50	OCH3	_	3	_	820.0	6.086	6.185	54
51		_	3	_	1280.0	5.893	6.409	54

6000 conjugate gradient), both the protein and the ligand were constrained with harmonic potential $k = 10.0 \text{ mol}^{-1} \text{ Å}^{-2}$. The system was heated stepwise from 0 K to 310 K in 500 ps without any constraints. After an equilibration period of 500 ps, a productive unconstrained molecular dynamics (MD) simulation of 1 µs duration was started, except for the drug dabrafenib, for which the total simulation time was 300 ns. A time step of 2 fs at constant pressure (1 atm) and temperature (310 K) was used. A langevine thermostat with a collision frequency of 1 ps^{-1} was used for temperature control. In all simulation protocols, hydrogen atoms were constrained by the SHAKE algorithm.75 The cut-off distance for non-bonded interaction was 11 Å, while for long-range electrostatic interactions, the particle mesh Ewald method⁷⁶ was used. Periodic boundary conditions were employed in all directions. MD simulations were performed using the molecular dynamics package Amber.77 MD simulations were performed on the Isabella cluster of the University Computing Center, University of Zagreb, Croatia.

2.1.3.1 Binding free energy calculation. The free energies of binding (ΔG_{bind}) between a series of potential drugs (ligands) and the hBRk were calculated according to well-established molecular mechanics/generalized Born surface area (MM/ GBSA) protocol.⁷⁸ The formula

$$\Delta G_{\text{bind}} = \Delta H - T\Delta S \approx \Delta E_{\text{MM}} + \Delta G_{\text{sol}} - T\Delta S \tag{1}$$

$$\Delta E_{\rm MM} = \Delta E_{\rm internal} + \Delta E_{\rm electrostatic} + \Delta E_{\rm vdW} \tag{2}$$

$$\Delta G_{\rm sol} = \Delta G_{\rm GB} + \Delta G_{\rm SA} \tag{3}$$

within single-trajectory approach is implemented in the MMPSBA.py script of the AmberTools package. The sum of the bond, angle and dihedral energy ($\Delta E_{internal}$), electrostatic ($\Delta E_{electrostatic}$) and van der Waals (ΔE_{vdW}) energies is ΔE_{MM} , the change in MM energy contribution in the gas phase. ΔG_{sol} is a change in solvation free energy, with a polar component (ΔG_{GB} , electrostatic solvation energy) and a non-polar, non-electrostatic solvation contribution (ΔG_{SA}). $T\Delta S$ is the conformational entropy at binding. The production phase trajectory

was divided into 20 segments of 50 ns length. Except for the drug dabrafenib (DAB), where the 300 ns trajectory was divided into 6 segments of 50 ns length. From each segment, 100 geometries were sampled at regular time steps and ΔG_{bind} was calculated. The final ΔG_{bind} was expressed as the mean \pm standard deviation for all 20 segments. The calculated MM/ GBSA free energies of binding were further decomposed into the specific contributions for each residue. In this way, the contributions of each amino acid side chain to ΔG_{bind} were determined and the nature of the energy change in terms of interaction and solvation energies, or entropic contributions were identified.⁷⁹ Since we are comparing the energies of the same receptor and of ligands with approximately the same size and number of rotatable bonds, entropic contributions ($-T\Delta S$) were neglected.

2.1.3.2 Cluster analysis. The structures of each complex were clustered based on the RMSD of the C α atoms of each residue using the *k*-means algorithm. The maximum number of iterations was set to 1000, with randomized initial set of points and sieving set was equal to 10. The frames closest to the centroids of the clusters were identified and considered as representative structures of the conformations. The CPPTRAJ module⁸⁰ was used to perform the cluster analysis.

3. Results and discussion

3.1 Gaussian-based QSAR model generation

Model was generated on structurally aligned pyrimidine– sulfonamide hybrid derivatives (1-51) with BRAF^{V600E} inhibitory activity. There was a moderate disparity between the actual and predicted values of the training and test molecules, which is known as residual activity, and it was within the Gaussian models' allowed ranges (<1), this indicates strong linear correlation.

The developed QSAR models were evaluated using the following parameters: squared correlation coefficient (r^2), cross-validated correlation coefficient (q^2 training), r^2 scramble, stability, Fisher's test (the variance ratio indicating the statistical significance of the model), *p*-value (magnitude of the

variance ratio, where small *p*-values usually indicate a higher degree of significance), SD (SD of the regression), RMSE test (root-mean-square error of the test set), q^2 test (value for the predicted activities of the test set) and Pearson-*r* (correlation coefficient between observed and predicted activities for the test set).⁸¹

3.1.1 Gaussian-based 3D-QSAR model. Model was generated using PLS with five factors, including steric, electrostatic, hydrophobic, HBD, and HBA fields of the training set.⁵⁹ The $r_{\rm CV}^2$ value of 0.7873 was derived from the cross-validation leave one out (LOO) method, while non-cross-validation analysis yielded an r^2 value of 0.9603. SD of the regression is 0.1855, Fischer's ratio value is 140.5, and stability of the model is 0.87. The statistical summary of the model is listed in Table 3.

The results for this model showed that the steric and electrostatic contributions were, respectively, 0.416 and 0.114. This implies that for protein–ligand interactions, the steric field is more significant than the electrostatic groups. Steric and electrostatic field contributions are given in (Table 3 and Fig. 3). The pIC₅₀ values were predicted using 15 test set inhibitors for model validation. The predictive q^2 for these models was 0.6384, which shows that they have a respectable level of predictive power. The contributions of the hydrophobic, HBD, and HBA fields were 0.169, 0.184, and 0.118, respectively (Table 3). The field contributions of the steric and hydrophobic intensities were more than those of the electrostatic, HBA, and HBD fields, indicating that the importance of the steric and hydrophobic fields for protein–ligand interactions is greater.

3.1.2 Different Gaussian contour maps. The contours of each of the five PLS components of the training set are described and shown in Fig. 3. In the Gaussian steric contour map (Fig. 3b), the phenyl ring (green contours) and the imidazo [2,1-*b*] thiazole moiety (red contours) are shown as favourable and unfavorable sites for steric interactions respectively.

The imidazo[2,1-*b*] thiazole moiety (green contours) with electropositive groups and the phenyl ring (red contours) with electronegative groups are active in the Gaussian electrostatic contour map (Fig. 3c). The green contour maps over the phenyl ring (Fig. 3d) of the Gaussian hydrophobic contour map show that they are hydrophobic, whereas the red contour maps over the imidazo[2,1-*b*] thiazole moiety show that they are not hydrophobic and may reduce BRAF^{V600E} activity. The hydrogen bond acceptor (HBA) groups in the phenyl ring (red contour) are unfavorable, whereas the HBA groups in the sulfonamide moiety (green contour) are favorable for activity, in the Gaussian HBA contour map (Fig. 3e). The green contour with the phenyl ring has a favorable hydrogen bond donor (HBD) group in the Gaussian HBD contour map (Fig. 3f), and substituting HBD groups at this position may boost BRAFV600E activity.

3.1.3 Validation of Gaussian-based 3D-QSAR model. Predicted *versus* experimental activity of the training and test sets was plotted to demonstrate the model's validity see Fig. 3g–i. Comparison of the experimentally observed and predicted pIC₅₀ values of pyrimidine–sulfonamide hybrid inhibitors, shows that the Gaussian model performs well in predicting the activities for both training and test molecules. After analyzing the statistical parameters, we came to the conclusion that the models had good predictability and offered precise information about the chemical properties of the ligands, which would contribute to antagonistic activity against BRAF^{V600E}.

Fig. 4 shows the structural requirements of the designed pyrimidine–sulfonamide hybrid derivatives based on 3D-QSAR. In designed compound Fig. 4, the alkyl chain of potent compound Fig. 3a is replaced by phenyl ring due to its higher hydrophobicity and imidazothiazole moiety was eliminated. Structure activity relationship (SAR) of designed compound on the basis of different parameters like hydrogen bond acceptor (HBA), (hydrogen bond donor) HBD, electrostatic hydrophobic

Statistical parame	Statistical parameters of generated 3D OSAR model						
PLS data	Gaussian	Acceptable value					
SD	0.1855	SD of regression \leq SD of actual activities (1)					
r^2	0.9603	$1 > r^2 > 0.5$					
$r_{\rm CV}^{2}$	0.7873	>0.6					
r^2 scramble	0.6879	<0.6					
Stability	0.87	>0.6 or = 1					
F	140.5	Larger values of F					
р	2.10×10^{-19}	Smaller values of p, greater degree of confidence					
RMSE	0.53	Close to 1					
q^2	0.6384	>0.5, and $(R^2 - Q^2)$ 0.3					
Pearson-r	0.806	Close to 1, lesser scattered predictive activities in the plot of observed vs. predicted activities					
Field contribution	. (%)						

Gaussian steric Gaussian electrostatic Gaussian hydrophobic Gaussian H-bond acceptor Gaussian H-bond donor

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0.416

0.114

0.189

0.118

0.184



Fig. 3 Gaussian based 3D-QSAR model. (a) Potent BRAF^{V600E} inhibitor $[IC_{50} = 1.200 \text{ nM}; (S. No. 30)]$ (b) steric contour map (c) electrostatic contour map (d) hydrophobic contour map (e) H-bond acceptor contour map (f) H-bond donor contour map. Relation between experimental and predicted BRAF^{V600E} inhibitory activity values of (g) all set (pIC₅₀) (h) test set (pIC₅₀) (i) training set (pIC₅₀).

and steric requirements at specific position are also depicted in Fig. 4.

3.2 Rationality of designed compound compared with USFDA approved BRAF inhibitors

Based on a Gaussian-based 3D-QSAR model, the developed pyrimidine–sulfonamide hybrid derivative's structural moieties

were assessed with the FDA-approved BRAF inhibitors: firstgeneration – sorafenib and second generation – dabrafenib, vemurafenib, and encorafenib for rationality.

Pyrimidine and sulfonamide (SO_2NH_2) moieties were selected as a core moiety for development of new BRAF^{V600E} inhibitors. The pyrimidine moiety is similar to the most potent drugs on the market, dabrafenib and encorafenib. The



Fig. 4 Structural requirements of pyrimidine–sulfonamide hybrid derivatives for BRAF^{V600E} activity and selectivity based on the analysis of Gaussian based contour maps.



Fig. 5 Structure of new designed pyrimidine-sulfonamide hybrid compounds

sulfonamide moiety is similar to all 2nd generation drugs. The *N*-phenyl ring moiety has similarity with sorafenib (1st generation), dabrafenib, vemurafenib, and encorafenib (2nd generation). The rational of the designed compound is depicted in Fig. S1.†

3.2.1 Different derivatives of designed pyrimidine–sulfonamide hybrid moieties. Importance of different function groups, such as OH (HBA/HBD), NO₂, Cl, Br, F, and I (electronwithdrawing), OCH₃, OH (electron-donating group), different heterocyclic moieties, such as quinoline, pyrrole, thiophene, and naphthol (steric requirements), were chosen based on the SAR of the designed compound (Fig. 4). On the basis of above findings, a library of 88 newly designed compounds was created and is shown in Fig. 5 and Table S1.[†]

3.3 Molecular docking and molecular dynamics simulations

Being aware of the potential problems with molecular docking experiments,82 we docked 92 compounds (88 designed compounds and 4 FDA approved BRAF inhibitor drugs) to the kinase receptor. The docking procedure was validated by redocking dabrafenib to the hBRk binding site. The root mean squared deviation (RMSD) between the experimental and docked structure is small (0.47 Å). Additionally, the rotation of the dabrafenib's methyl group contributes the most to the overall RMSD, so we have demonstrated the correctness of our approach. The superimposed experimental and docked dabrafenib structures in the enzyme pocket are displayed in Fig. S2.† Among all docked compounds, dabrafenib performed best with a docking score of -12.7 kcal mol⁻¹. Four molecules, T109, T183, T160 and T126 had scores below -12 kcal mol⁻¹, so they were selected for detailed studies. Binding poses of T109 (red), T126 (orange), T160 (sea green), T183 (sky blue), and the reference drug dabrafenib (magenta) as determined by molecular docking to the BRAF kinase protein pocket are depicted in Fig. S3.[†] It is interesting that none of the docked hit molecules nor dabrafenib (docked or experimentally determined structure) are in contact with Glu600. Fig. S4[†] shows docked T160 (sea green) and dabrafenib (magenta) to BRAF. CA atom of Glu600 (black) is more than 14 Å apart from the nearest nonhydrogen atom of T160/dabrafenib ligands. All docking scores are collected in Table S2.†

After visual inspection of the best docked poses of the top 4 compounds forming our group of hit molecules, they served as initial geometries for MD simulations. Before conclusions were drawn, the conformational dynamics and stability of each complex were examined using a 1 μ s long molecular dynamics

simulation. The RMSD and radius of gyration (RoG) of the backbone α C atoms of the receptor relative to the initial structures were calculated and the stability of the complexes was monitored. Flexibility of the residues' side chain was evaluated using RMSF. The compounds were ranked based on their ability to bind to hBRk, and their interaction pattern was investigated.

3.3.1 Molecular dynamics simulation study of compound T 160. In the first 900 ns of the simulation, the RMSD fluctuated around a mean value of 2.44 Å. Then an increase in RMSD was observed, and the mean value for the last 100 ns was 3.05 Å. These changes prompted us to perform a cluster analysis to identify relevant changes in the complex structure. We chose to use the k-means algorithm, where k ranges from 2 to 5. The results were analyzed using the Davies-Bouldin index (DBI), the pseudo-F statistic (pSF), and the ratio between the sum of squares of the regression and the sum of squares of error (SSR/ SST) (Table S3[†]). The analysis confirmed the presence of three relevant conformations with selected parameters collected in Table S4.[†] For each conformation, a representative structure was identified as a frame closest to the cluster centroid. The distribution of the conformations of the T160:BRAF kinase (T160:hBRk) complex along the trajectory and their RMSD relative to the initial frame are shown in Fig. 6. Along with the increase in the mean RMSD values after the 900 ns mark, a slight decrease in the radius of gyration (RoG) from 19.24 Å to 19.15 Å was observed.

In all three conformations, the secondary structure was generally well preserved. However, there were some minor changes in the secondary structure that could affect the geometry of the catalytic pocket Fig. 7. The first region includes residues that are in direct contact with the ligand - Arg462, Ile463, Gly464, and Ser465, while these residues do not have a defined secondary structure in the dominant conformation A, they form a β-sheet in conformations B and C. The second region with considerable changes is the region between Ile592 and Phe 635, which is almost completely without defined secondary structure. In the crystal structure (Fig. S5[†])⁸³ and in conformation A, only Pro622 to Ile625 from this region form an H4 α-helix. In conformation B, the H4 helix is absent but two new helices are formed (Glu611-Leu613 and Ile617-Trp619). The H4 helix and the Ile617-Trp619 helix are present in the C conformation.

The flexibility of the residues of the hBRk protein in the complex was examined by calculating root mean squared fluctuations (RMSF) for each residue (Fig. 8). High RMSF values indicated greater fluctuations and flexibility. The averaged RMSF value was 1.55 Å. Almost all of the first 50 residues were



having higher than average RMSF values, indicating high flexibility of the N-terminus of the kinase. The highest RMSF value was having Met627, a residue whose side chain pointed toward the solvent, far from the binding site of the ligand. The previously mentioned unstructured region was having two fragments with RMSF values above 3 Å. These two fragments, Ser607–Ser614 and Ile625–Asn631, were not in direct contact with the ligand. It is important to note that the catalytically relevant residues Asp594, Phe595 and Gly596 have RMSF values below 1 Å. Hydrogen bond analysis revealed that on average 2.11 hydrogen bonds were established between T160 and the kinase. This is on average only 0.02 less than in the complex with T183 and the hBRk. The predominant hydrogen bond formed in almost 75% of the simulation time was between the hydroxyl group of Thr529 (hydrogen acceptor) and the N4 atom of T160 with an average bond length of 1.89 ± 0.08 Å. Changes in the 3D structure of the protein were reflected in the hydrogen bonding patterns in different conformations. For example, in A, T160 formed four hydrogen bonds with three residues (two with



Fig. 7 Structural differences between conformations A (blue), B (red) and C (green). Superposition of three structures of the T160:hBRk complex representing three conformations.



Fig. 8 Root mean square fluctuations per residue of the T160:hBRk complex.

Ser465 and one with Thr529 and Asp594). In addition, 15 residues had at least one heavy atom in a 4.0 Å zone away from the non-hydrogen atoms of T160. This illustrated the large number of stabilizing van der Waals interactions. In conformation B, only two hydrogen bonds were formed (with Lys483 and Thr529). Finally, conformation C exhibited three hydrogen bonds. The detailed interactions are shown in Fig. 9.

Although the MD simulation for DAB was 300 ns long, some conclusions can be drawn. The flexibility of residues changes slightly when T160 is replaced in the catalytic pocket by dabrafenib, the drug approved for the treatment of melanoma with V600E or V600K mutation (Fig. S6†), with a slight increase in the flexibility of residues Ala544–Lys548 and a decrease in the flexibility of Ile625–Asn631. In addition, dabrafenib has on average one more hydrogen bond than T160, being predominantly (90% of the simulation time) bound to the Gly593 oxygen atom with an average hydrogen bond length of 1.84 Å, which is shorter than the hydrogen bond between Thr529 and T160. Numerous van der Waals interactions together with a higher number of hydrogen bonds could be responsible for the higher binding affinity to hBRk than T160.

3.3.2 Calculation of the free energy of binding. The binding affinity of the hit molecules and the reference drug dabrafenib was estimated using the MM/GBSA approach, in which the free energies of solvation were determined by solving

the generalized Born equation. For the complex T160:hBRk, the free energy of binding without entropy contribution was -59.1 \pm 2.6 kcal mol⁻¹. T160 had the highest binding potential to hBRk of all the compounds studied. The MM/GBSA binding free energy decomposition was used to identify key residues with dominant contribution to protein-ligand binding. In the study of the SARS-CoV-2 virus, its main protease⁸⁴ and the NS3 protease of Kyasanur forest disease virus,⁸⁵ a threshold of -1.5kcal mol⁻¹ was set for the free energy of binding of a single residue to classify it as a residue with dominant contribution, and the same criteria was applied in the present study. Table 4 lists the residues with dominant contributions for T109, T126, T160, and T183 and dabrafenib. Fig. 10 shows the main interactions based on the decomposition of the binding free energy. In addition to the polar and uncharged Thr529 and the positively charged Lys483, four hydrophobic residues (Val471, Leu505, Leu514, Trp531, and Phe583) contributed significantly to the binding free energy of T160. This finding confirmed the results of previous analyzes on the importance of van der Waals interactions.

As expected, the Gly593 residue, which forms the hydrogen bond, has the largest contribution to the binding of dabrafenib (DAB). Other stabilizing interactions are established with the electron-rich residues Phe583 and Trp531, the hydrophobic Val471, Leu514, and Ile527, and the polar Thr529 and Cys532. The free energy of binding for the DAB:hBRk complex was estimated to be -61.7 ± 1.0 kcal mol⁻¹, slightly better than for T160:hBRk.

The results obtained by the analysis of Gaussian-based contour maps suggest that the introduction of the HBA group at the phenyl moiety should increase the activity of the pyrimidine–sulfonamide hybrid derivatives. Comparing compounds T160 and T126, which differ only in the presence of a nitro group at the *para* position of the phenyl moiety in T160, one can observe a drastic change in the binding free energy for T160 is -59.1 ± 2.6 kcal mol⁻¹, it is only -39.7 ± 2.2 kcal mol⁻¹ for T126 (*vide infra*). The MD simulations revealed that the nitro group sits in a hydrophobic pocket and is surrounded by hydrogen atoms. To gain a deeper insight into the nature of why some structural modifications predicted by Gaussian-based contour maps contribute to higher selectivity, additional MD simulations are needed, including non-substituted pyrimidine–sulfonamide



Fig. 9 Hydrogen bonds patterns in three T160:hBRk conformations.

Table 4	Contributions of the most importa	nt amino acid residues	for the binding of T10	9, T126, T160,	T183 and dabrafenib to hBRk
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T109		T126		T160	T160		T183		Dabrafenib (DAB)	
Residue	$\Delta G_{ m bind}$									
Lys483	-6.86	Lys483	-2.68	Thr529	-3.63	Lys483	-3.09	Gly593	-3.45	
Asp594	-3.93	Phe583	-2.27	Lys483	-2.74	Asp594	-2.38	Trp531	-2.63	
Ile527	-2.21	Leu514	-2.19	Leu514	-2.43	Val471	-2.08	Phe583	-2.36	
Val471	-2.07	Val471	-2.16	Val471	-2.05	Gly593	-2.07	Thr529	-2.23	
Phe583	-1.77	Thr529	-1.80	Leu505	-1.85	Ile463	-2.02	Leu514	-1.93	
Leu514	-1.51	Ile463	-1.73	Phe583	-1.77	Leu514	-1.97	Val471	-1.89	
		Leu505	-1.58	Trp531	-1.62	Ile527	-1.92	Ile527	-1.81	
						Phe583	-1.91	Cys532	-1.70	
						Leu505	-1.56	•		



Fig. 10 Insight into the catalytic site neighborhood of T160 in hBRk, with residues with the most favorable single residue binding free enerav

scaffolds, but this is beyond the scope of the present manuscript.

3.3.3 Compound T109, T126 and T183 molecular dynamics simulations study. The same protocol used for the analysis of the T160:hBRk complex was applied to the analysis of the T109:hBRk, T126:hBRk, and T183:hBRk complexes. The stability of the complexes was confirmed by tracking RMSD and RoG along the trajectory (Fig. S7[†]).

All complexes remained stable but had higher averaged RMSD values compared to the T160:hBRk complex. The RoG values were very similar and range from 18.93 Å to 19.29 Å. The existence of multiple conformations was confirmed by cluster analysis. The cluster analysis data are summarized in Tables S5-S10.† Three complexes shared the presence of an initial shortlived conformation that converts to another conformation within 100 ns. The relative ratios between the populations of conformations A and B are approximately 0.6 to 0.3.

The primary geometric difference between the conformations of T109:hBRk and T126:hBRk, just as with T160:hBRk, was in unstructured fragments. However, in T183:hBRk, conformations A and B showed differences in secondary structure motifs compared to the representative structure of the initial conformation C (Fig. 11). For example, a shift of the H6 α -helix axis was observed. Slightly less pronounced was the shift of the H1 α -helix axis, but it played a larger role in the geometry of the

binding site. While the C-terminus of the H1 α -helix had van der Waals contacts with the catalytic Phe595 residue, the Nterminus is tilted toward the unstructured loop region, reducing the volume of the cleft.

hBRk residues had higher average RMSF values, 1.81 Å and 1.80 Å, when T183 and T126 were bound in the active pocket, respectively, compared with T109 (1.55 Å). The RMSF diagram (Fig. S8[†]) shows similar flexibility patterns. Again, the Nterminus and unstructured regions exhibited above average flexibility, while the conserved triad, which was important for catalysis, exhibited low flexibility. T126 had the lowest average number of hydrogen bonds (0.95) between ligand and the receptor. The hydrogen bond connecting the hydroxyl oxygen of Thr529 and the NH group of T126 was present only during 23% of the simulation time, with an average length of 1.91 Å. For comparison, the analogous bond in T160:hBRk was present for about 75% of the simulation time. In T183:hBRk, Gly593 acted as a hydrogen bond acceptor, and the 1.83 \pm 0.10 Å long bond was present during 82% of the trajectory. In addition to hydrogen bonding, van der Waals interactions also contributed to free energy of binding. While the free energy of binding of



Fig. 11 Superposition of three structures of the T183:hBRk complex representing three conformations with circled regions of major structural rearrangements. Conformations A (blue), B (red) and C (green).



Fig. 12 FDA inhibitors and designed compound interaction with different region of BRAF^{V600E} and their binding conformation on the basis of DFG motif and α C-helix.

T109 and T183 was almost within 1 kcal mol⁻¹ (-55.5 ± 3.1 kcal mol⁻¹ and -56.6 ± 3.0 kcal mol⁻¹, respectively), the value for T126 is only 39.7 \pm 2.2 kcal mol⁻¹. T109, T160, and T183 can serve as excellent starting points for introducing chemical modifications to optimize their potential to inhibit BRAF kinase.

4. Binding study of BRAF inhibitors and designed compounds with DFG motif and α C-helix of BRAF^{V600E} protein and their conformation

BRAF inhibitors are having 4 different types binding conformations with DFG motif and α C-helix.⁸⁶ Employing this notion, by using molecular docking and simulation studies, we succeeded in locating our designed molecule's binding conformation with the DFG motif and α C-helix (Fig. 12).

The Asp594 residue of the DFG motif was facing the active site of the kinase in the designed compounds (T109, T126, T160 and T183). Compounds T109, T126, T160 and T183 have interactions with the hinge region (ATP-binding site) by forming \sim 1-3 hydrogen bonds and hydrophobic interaction around the adenine regions of the ATP binding site of the protein. The designed compounds also has interactions with other various regions of ATP, including hydrophobic regions adjacent to the adenine region, the ribose region, and the solvent assessable region (DFG-IN). The αC-helix away from the ATP site is referred to as the aC-OUT position. Thus, designed compounds interacted with the [aC-OUT/DFG-IN] confirmation, just like dabrafenib. encorafenib and vemurafenib.23,87 Additionally, compounds T126, T160, and T183 interact with the Leu505 amino acid sequence at the dimerization interface (DIF). BRAF dimerization contributes to the pathogenic function of diseaseassociated mutant Raf proteins and exhibits activity similar to the constitutively active BRAF^{V600E} mutant once this activated state has been attained. It has also been found to alter

therapeutic responses and disease progression in patients treated with BRAF inhibitors.⁸⁸ Oncogenic BRAF dimers shown resistance against BRAF inhibitors and causes paradoxical activation.⁸⁹

5. Conclusion

The pyrimidine-sulfonamide hybrid derivatives were designed based on the different structural moieties of the first and second-generation BRAF inhibitors. 3D-QSAR and molecular docking studies were performed for the designed compounds to determine their binding affinity. Molecular dynamics simulations were performed to understand the flexibility, structural conformation changes and interaction pattern of the mutant BRAF kinase with hit compounds. Analysis of the Gaussianbased 3D-QSAR models showed that the contribution of steric and hydrophobic fields was higher than the other field intensities. These models provided considerable insight into the key properties of the inhibitors, their structural features, and their inhibitory potential. The developed compounds showed good interactions with the core active site [the nucleotide (ADP or ATP) binding site, DFG motif, the phospho-acceptor site (activation segment), adjacent to the DFG motif and the aC-helix] of the BRAF^{V600E} protein. Similar to the FDA-approved BRAF^{V600E} inhibitors the developed compounds have $[\alpha C-OUT/DFG-IN]$ conformation and compound T126, T160, and T183 interacted with dimerization interface and may effective against malignancies driven by dimer interface. The synthesis of these designed molecules is in progress. The synthesized molecules will be tested further to support the results of in silico studies.

Author contributions

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Abbreviations

ADP	Adenosine diphosphate
ATP	Adenosine triphosphate
CR	Conserved regions
CRD	Cysteine-rich domain
DAB	Dabrafenib
DIF	Dimerization interface
FDA	Food and drug administration
3D-	Three-dimensional quantitative structure-activity
QSAR	relationship
hBRk	Human BRAF kinase
LOO	Leave one out
MAPK	Mitogen-activated protein kinase
MD	Molecular dynamics
MM/	Molecular mechanics/generalized Born surface area
GBSA	
PLS	Partial least square
RMSD	Root mean squire deviation
RMSE	Root mean squire error
RMSF	Root mean squared fluctuations
RoG	Radius of gyration

Conflicts of interest

The authors declare no competing interest.

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