

Proteins in harmony: Tuning selectivity in early drug discovery Burggraaff, L.

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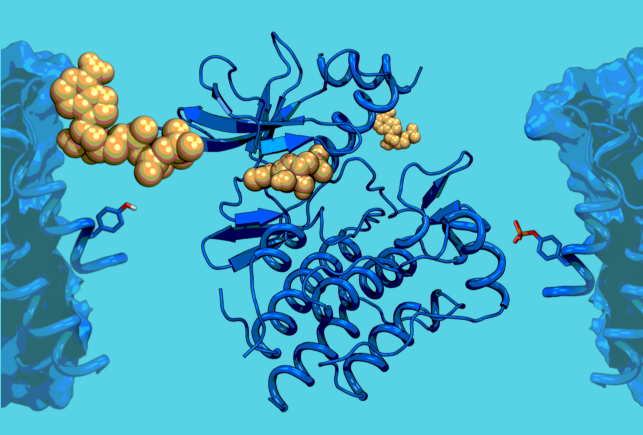
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Chapter five

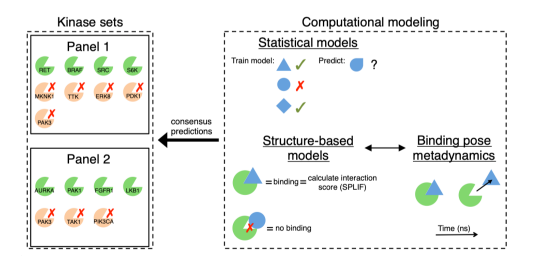
Successive Statistical and Structure-Based Modeling to Identify Chemically Novel Kinase Inhibitors



Lindsey Burggraaff, Eelke B. Lenselink, Willem Jespers, Jesper van Engelen, Brandon J. Bongers, Marina Gorostiola González, Rongfang Liu, Holger H. Hoos, Herman W.T. van Vlijmen, Adriaan P. IJzerman, and Gerard J.P. van Westen. Journal of Chemical Information and Modeling. 2020. doi: 10.1021/acs.jcim.9b01204.

Abstract

Kinases are frequently studied in the context of anti-cancer drugs. Their involvement in cell responses such as proliferation, differentiation, and apoptosis, makes them interesting subjects in multi-target drug design. In this study a workflow is presented that models the bioactivity spectra for two panels of kinases: 1) inhibition of RET, BRAF, SRC, and S6K, while avoiding inhibition of MKNK1, TTK, ERK8, PDK1, and PAK3; 2) inhibition of AURKA, PAK1, FGFR1, and LKB1, while avoiding inhibition of PAK3, TAK1, and PIK3CA. Both statistical and structurebased models were included, which were thoroughly benchmarked and optimized. A virtual screening was performed to test the workflow for one of the main targets, RET kinase. This resulted in 5 novel and chemically dissimilar RET inhibitors with remaining RET activity of < 60% (at a concentration of 10 µM) and similarities with known RET inhibitors from 0.18 to 0.29 (Tanimoto, ECFP6). The four more potent inhibitors were assessed in a concentration range, and proved to be modestly active with a pIC₅₀ value of 5.1 for the most active compound. The experimental validation of inhibitors for RET strongly indicates that the multi-target workflow is able to detect novel inhibitors for kinases, and hence this workflow can potentially be applied in polypharmacology modeling. We conclude that this approach can identify new chemical matter for existing targets. Moreover, this workflow can easily be applied to other targets as well.





Introduction

Compound promiscuity can be leveraged to develop multi-target drugs. Such multi-target drugs can replace existing multi-drug treatments, while maintaining the therapeutic effect.¹ One of the advantages of a multi-target drug, or single-drug treatment is that no drug-drug interactions occur, making the treatment less risky and harmful for the patient.² However, since multi-target drugs are designed to bind to multiple proteins, they may tend to be more promiscuous as well. Therefore, when developing multi-target compounds, off-target binding and pathways should also be considered.

In the light of the "Multi-Targeting Drug" DREAM challenge,³ bioactivities were computationally modeled for two panels of kinases. The first panel was based on treatment of medullary thyroid carcinoma, where kinases RET, BRAF, SRC, and S6K, should be inhibited and MKNK1, TTK, ERK8, PDK1, and PAK3, should not be affected.^{4,5} RET was considered the main on-target kinase in this panel and thus was prioritized over other kinases in the panel. The second panel was based on tauopathies in neurogenerative disease:

Box 5.1 The Dialogue for Reverse Engineering Assessments and Methods (DREAM) challenge is a contest organized by a community of researchers and is open to participants worldwide. Starting from 2006, every year multiple challenges are organized aimed at a different subject within biological science and human health. A common motive throughout the challenges is that it is promoted to find a solution using computational models and techniques. The "multi-targeting drug DREAM challenge" was part of the 12th year edition of DREAM challenges.

compounds should inhibit AURKA, PAK1, FGFR1, and LKB1, and not bind to PAK3, TAK1, and PIK3CA.³ Since the main on-targets AURKA and PAK1, and additional on-targets FGFR1 and LKB1, are targeted in the central nervous system, compounds for panel 2 kinases should additionally be able to pass the blood-brain barrier.

This study describes a rigorous workflow to model the bioactivity spectra of compounds in kinases (**Figure 5.1**) and identify novel inhibitors. Every step in the workflow was extensively benchmarked, and each model was validated prior to virtual screening. A consensus scoring approach was used to rank virtual screening compounds, and only compounds with a Tanimoto similarity (ECFP6 (extended-connectivity bit string, diameter 6))⁶ < 0.4 were considered to make sure that existing active molecules would not be 'rediscovered'. This consensus approach encompassed statistical modeling techniques such as quantitative structure-activity relationship (QSAR) models and proteochemometric (PCM) modeling.^{7–9} Moreover, structure-based docking and pose metadynamics were applied.¹⁰ Next to compound ranking, this approach was also used to exclude inactive-predicted compounds from virtual screening along the way. Fast machine learning models were applied to discard compounds early in the workflow. Subsequently, slower, but information rich, structure-based models were applied that consequently only had to score a smaller fraction of compounds compared to the entire initial virtual screening set. In this way, millions of compounds were screened rapidly and scored accurately.

In addition to the extensive benchmark validations, the workflow was also validated in vitro for one of the main on-target kinases. Forty-six compounds were experimentally validated for RET, of which 15 were selected based on the consensus approach, based on statistical models only, and 25 compounds based on structure-based modeling. This resulted in a total of 5 inhibitors causing RET activity < 60% at a concentration of 10 μ M. The four most potent compounds were further inspected and their IC₅₀ were determined. Although the most potent RET inhibitor

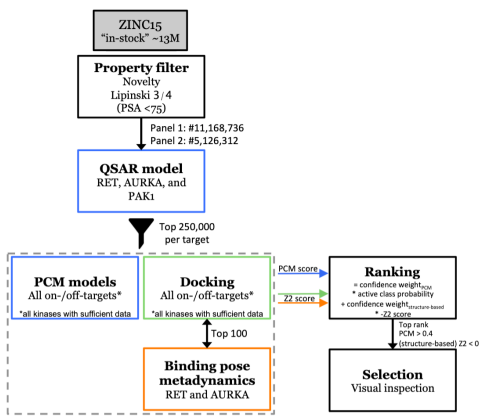


Figure 5.1. Virtual screening workflow for the two panels of kinases. Statistical models (blue), structure-based models (green), and molecular dynamics (orange) were applied to rank the virtual screening compounds.

was modestly active with a pIC_{50} of 5.1 ± 0.1 , all compounds were chemically distinct from known RET inhibitors with Tanimoto (ECFP6) similarities smaller than 0.29. Therefore, the identified inhibitors in this study provide a new starting point for hit/lead optimization based on novel scaffolds.

Results

Data curation and filtering

Compound bioactivity data was derived from different sources to increase data availability for the benchmark sets in model training and validation. Kinase compound data was retrieved from the following sources: ChEMBL database (version 23)¹¹, publicly available sets from Eidogen¹², and ExCAPE-DB¹³. The data was curated by standardizing chemical structures and bioactivity values (see Methods section for details). In this way a dataset was constituted that contained compound bioactivity information for 512 kinases. This dataset was used for training and testing of statistical models. Since validation was performed using cross-validation, the dataset was divided into subsets; five subsets were created based on chemical clustering of the compounds per target (see Methods section for details). The biggest subset was used in training of every



dataset available at doi: 10.4121/uuid:6af1d9de-281f-4221-b7e1-e7c01b90dfe0.

model, whereas the remaining four subsets were used rotationally for testing. The subsets that were not used in testing in the specific iteration were added to the training set.

Additionally, separate active/inactive/decoy datasets were constructed for validation of structure-based models. These benchmark sets were constructed for each panel kinase separately. Data for these benchmark sets were derived from ChEMBL (version 23), which was curated and filtered to construct a benchmark set for each kinase containing inactive and chemically diverse active compounds: only compounds with measured pChEMBL¹⁴ values were included in the benchmark sets and a restrain was set for the number of active compounds. Up to a maximum of 100 actives per kinase were included, whereas the number of inactive compounds was not limited. The 100 active compounds were selected based on chemical diversity by clustering the compounds with pChEMBL value > 6.5 into 100 clusters and selecting only the cluster centers. Additionally, DUD-E¹⁵ decoys were added to each benchmark set. These decoys have similar physicochemical properties as active ligands, but differ in chemical structure. The structure-based benchmark sets were smaller than the training and test sets of statistical models, but big enough to allow for validation of the models. The smaller size of the structure-based benchmark sets allowed for quicker model evaluation, resulting in validation of many protein structures.

The virtual screening set that was screened using both statistical- and structure-based models was derived from the ZINC15¹⁶ database. All "in stock" compounds were filtered on drug-like properties by discarding compounds that did not adhere to 3 of 4 Lipinski rules.¹⁷ Furthermore, compounds were filtered on novelty: compounds with Tanimoto similarity (ECFP6) > 0.4 compared to existing actives (pChEMBL > 5) on the kinases in the respective panel were excluded from the virtual screening set. The virtual screening set was additionally filtered for panel 2 kinases by including the likelihood of compound passing the central nervous system by only keeping compounds with polar surface area < 75 Å². This resulted in a virtual screening set for panel 1 of 11,168,736 compounds, and for panel 2 of 5,126,312 compounds.

Statistical models

Separate quantitative structure-activity relationship (QSAR) models were constructed for the main on-targets RET, AURKA, and PAK1, as a first filter for bioactive compounds. The models were validated with 4-fold cross-validation using standardized benchmark sets (see Methods for details). The benchmark sets were constructed per target and were extracted from the main statistical benchmark set containing 512 kinases. Chemical descriptors were calculated for every compound: FCFP4 (feature-connectivity bit string, diameter 4) fingerprints and physicochemical descriptors (listed in table S5.1). These descriptors describe the compounds and were used in training the models. The RET, AURKA, and PAK1 QSAR models were predictive with ROC (receiver operating characteristic) scores higher than random (Table 1). The ROC of the QSAR models was comparable between targets (ROC 0.76 ±0.01), whereas the Matthews correlation coefficient (MCC) varied slightly more (MCC 0.30 ±0.08).

The performances of the QSAR models were sufficient as a first filter for bioactive compounds, discarding the least active compounds and steering clear of the decision boundary. Virtual screening set 1 was screened using the RET QSAR, and virtual screening set 2 was screened using the AURKA and PAK1 QSAR models separately. Using the active class probability score, the most promising compounds (250,000 compounds per RET/AURKA/PAK1) were selected

to be further processed in the structure-based approaches. This pre-screening of compounds with a simple, but fast model decreased the number of compounds effectively. As a result, subsequent screening and scoring steps proceeded quicker, since fewer compounds had to be screened. The subsequent steps, proteochemometrics (PCM)⁹ and structure-based modeling, were carried out in parallel. Additionally, QSAR models were constructed for the remaining kinases simultaneously. These QSAR models were compared to the more advanced PCM models to select the best approach for scoring. The performances of these QSAR models are included in **Table 1**.

The PCM models that were created were applied solely for the purpose of scoring and not as a filtering step. PIK3CA was excluded from modeling, as insufficient data was available to build and validate the models. The PCM models were constructed for each kinase separately and were based on the particular kinase and its L4 level family members as annotated in ChEMBL.¹⁸ In addition to the compound descriptors used in QSAR modeling (FCFP4 fingerprints and physicochemical properties), the PCM models included protein descriptors that were based on a full kinome sequence alignment (see Methods for details). Initial PCM models were trained using default random forest settings: 300 trees, log2(m) features at every node in the tree, no maximum tree depth, a minimum of 2 samples to consider a node for splitting, and bootstrap enabled. This resulted in an overall performance of: MCC 0.22 and ROC 0.69 (average over all panel kinases). Since the PCM models were intended for scoring of compounds, the models were optimized to enhance predictive performance. Optimal settings were explored by tuning hyperparameters with random search, a basic approach to automated machine learning. 19 Approximately 500 random forest models were trained during optimization, resulting in the best model with performance MCC 0.25 and ROC 0.74, and following settings: 300 trees (fixed) with 43 features at every node in the tree, a maximum tree depth of 99, a minimum of 12 samples to consider a node for splitting, and bootstrapping disabled. The performance of the QSAR and PCM models per kinase are shown in Table 5.1.

Predictions for most kinases were comparable between QSAR and PCM modeling. However, for target PAK3, PCM clearly outperformed QSAR with MCC difference 0.18 and ROC difference 0.20. MCC could not be calculated for ERK8 due to a small dataset and consequently a lack of a predicted class (total number of compounds for ERK8 is 332). Using QSAR, the MCC displays a negative correlation for ERK8 (-0.12), which is also reflected by the ROC score that is worse than random (ROC < 0.5). PDK1 has high early enrichment (BEDROC (Boltzmann-Enhanced Discrimination of the Receiver Operating Characteristic)) with both QSAR and PCM. Although overall enrichment (ROC) for PDK1 is lower than the early enrichments, it is still good with ROC 0.71 (QSAR) and 0.72 (PCM). The average over all targets shows that PCM predicts slightly better than QSAR, with a similar MCC score of 0.25, but higher (BED)ROC scores: differences BEDROC = 0.06, and ROC = 0.06. Moreover, the nature of PCM models allows for extrapolation of bioactivities from related kinases to the target of interest. Therefore, it was hypothesized that the PCM models will perform better than QSAR models when applied to a more diverse chemical dataset such as the virtual screening sets. The performances of the PCM models of the main on-targets AURKA and PAK1 were higher than the average performance over all targets. However, main on-target RET had a PCM MCC value (0.15) that was lower than average (0.25). Nevertheless, (BED)ROC scores were higher than the average: RET BEDROC 0.64, RET ROC 0.76, average BEDROC 0.58, and average ROC 0.74. Moreover, all RET scores



Table 5.1. PCM performance per target	(mean over 4-fold cross-validation).
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		PCM			QSAR		Number of	compounds
Target	MCC	BEDROC $(\alpha = 20)$	ROC	MCC	BEDROC (α = 20)	ROC	Active (pChEM- BL ≥ 6.5)	Inactive (pChEM- BL < 6.5)
RET	0.15	0.64	0.76	0.23	0.63	0.75	1492	416
BRAF	0.18	0.74	0.56	0.20	0.75	0.54	1119	1359
SRC	0.28	0.47	0.72	0.26	0.47	0.72	4642	2238
S6K	0.38	0.85	0.79	0.45	0.85	0.78	1662	685
MKNK1	0.09*	0.42	0.61	0.01	0.32	0.50	549	51
TTK1	0.22	0.45	0.78	0.26	0.44	0.75	663	276
ERK8	****	0.05	0.48	-0.12	0.02	0.35	302	30
PDK1	0.27*	0.85	0.72	0.31	0.86	0.71	579	536
PAK3	0.25	0.72	0.91	0.07	0.27	0.71	1204	53
AURKA	0.37	0.65	0.78	0.38	0.47	0.77	3165	1674
PAK1	0.32	0.74	0.86	0.28	0.66	0.77	712	114
FGFR1	0.41	0.70	0.85	0.77	0.71	0.82	2477	928
LKB1	0.57**	0.53	0.76	0.26*	0.45	0.63	429	47
TAK1	0.15***	0.27	0.68	0.12*	0.33	0.69	1204	53
Average	0.25	0.58	0.74	0.25	0.52	0.68	295	56

Asterisks indicate that no value could be determined due to lack of predicted (positive/negative) classes: * = 1 cross-validation failed, *** = 2 cross-validations failed, *** = 3 cross-validations failed, and **** = 4 cross-validations failed. Indicated in bold in each column is the best performing model for that given parameter.

were better than random indicating the predictive power of the model.

The settings that corresponded to the best performing model in 4-fold cross-validation were applied to train PCM models on the full dataset per kinase (including test set in training). The models were applied to score all the virtual screening compounds that passed the filtering step using QSAR models.

Structure-based models

Structure-based scoring was performed in addition to scoring of compounds using the PCM models. For many kinases,multiple crystal structures have been deposited in the PDB, and it is often not obvious which crystal structure should be used prospectively. Therefore we performed a rigorous benchmark to determine the best enriching crystal structure. The number of validated crystal structures for all targets ranged from 1 to 135, excluding ERK8 and PAK3 for which no crystal structures were available at the time (December 2017). The crystal structures that were deemed suitable for virtual screening were X-ray protein structures that contained a co-crystalized orthosteric ligand. A total of 499 crystal structures were benchmarked using corresponding compound benchmark sets that were composed for each target separately. The benchmark sets containing active compounds, inactive compounds, and decoys, were docked into the orthosteric binding pockets, from which a docking score was derived for each compound. The decoys were considered as inactive compounds when calculating actives enrichment for each crystal structure. Enrichment was calculated based on best docking score per compound. It was observed that actives enrichment varied greatly between different structures of the same kinase: ROC ranging

from 0.58 (PDB: 2IVS) to 0.77 (2IVU) for RET, 0.47 (5OSF) to 0.80 (2XNG and 4O0W) for AURKA, and 0.65 (3Q4Z and 4O0T) to 0.95 (5IME) for PAK1. The models per target were ranked based on the sum of the overall (ROC) and early enrichment (BEDROC, $\alpha = 160.9$). Models with a score ROC+BEDROC ≥ 1 (e.g. ROC 0.40 + BEDROC 0.60) were considered sufficient for prediction of active compounds.

Kinases LKB1 and MKNK1 only had a few crystal structures available (1 and 2 structures, respectively) of which performance was low (ROC+BEDROC: 0.50 and 0.72 respectively). Moreover, kinase S6K of which 18 structures were available, only reached maximum performance of ROC+BEDROC 0.80. Therefore, additional protein structures were created for these kinases by application of induced-fit docking. In contrast to docking, induced-fit docking accommodates the ligand and additionally re-orientates the side chains of binding pocket residues. This allows the residues to change conformation in order for the ligand to fit into the binding pocket. Five ligands for each LKB1, MKNK1, and S6K, were docked with induced-fit docking into the respective binding pocket. Subsequently, the ligands were removed from the protein, keeping only the altered protein structures. This resulted in 95 additional protein structures for LKB1, 44 for MKNK1, and 74 for S6K. These additional structures were validated using the same benchmark sets as used for the initial structures. The protein structures that were created using induced-fit docking varied in performance from ROC+BEDROC 0.27 to 1.07. For all three kinases, an induced-fit structure was generated that outperformed the initial crystal structure. The best performance for LKB1 was ROC+BEDROC 0.99, for MKNK1 it was 1.07, and for S6K it was 0.97.

Based on the docking performances of all models, five protein structures per target were selected for structural protein-ligand interaction fingerprints (SPLIF)²⁰ calculations. With SPLIF, interactions between the binding pocket residues and (docked) ligand are calculated, resulting in a SPLIF score that indicates the similarity between the interactions of the (docked) ligand compared to a reference compound. In this case, the reference compound consists of the cocrystalized ligand in the corresponding structure. The structures for SPLIF calculations were selected based on diversity of the co-crystalized ligands per kinase. The diversity of these ligands was assessed by clustering the ligands using affinity propagation clustering. 21 The cluster centers of the structures with highest ROC+BEDROC in docking were selected as reference. Consequently, the corresponding protein structures and docked benchmark set poses were thus used in SPLIF calculations. The SPLIF similarity scores for each benchmark compound were used to calculate actives enrichment based on SPLIFs. Especially for MKNK1 actives enrichment increased significantly compared to enrichment based on docking scores: ROC+BEDROC SPLIF 1.57 versus ROC+BEDROC docking 1.07. Based on early enrichment, BEDROC (a = 160.9), the performance using SPLIF scores increased for 9 of 13 kinases (Figure 5.2). However, for targets BRAF, PDK1, PAK1, and FGFR1 docking scores enriched (BEDROC (a = 160.9)) better than SPLIF scores.

An ensemble score was constituted, which combined docking and SPLIF scores. This ensemble score, the Z2 score, ²² was based on only docking scores, only SPLIF scores, or a combination of docking and SPLIF scores (see Methods for details). Prior to calculation of Z2-scores, the docking and SPLIF scores were normalized to Z-scores (more negative Z-score means better

The protein structures resulting from induced-fit docking are available at doi: 10.4121/uuid:9e61b6a6-88e5-4a18-ba19-dbf1bbdd656a



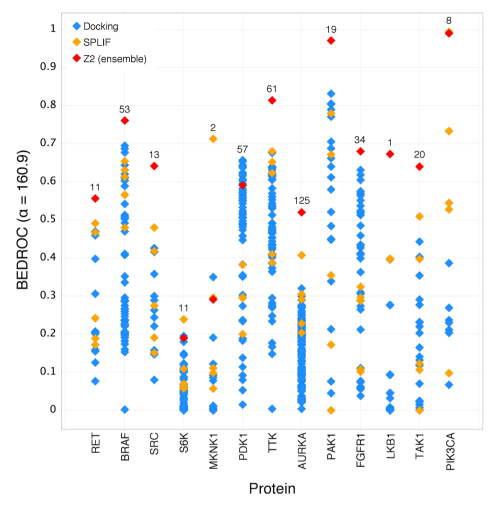


Figure 5.2. Early enrichment of actives, BEDROC ($\alpha = 160.9$), per crystal structure for each target. Enrichment reached by docking scores (blue), SPLIF scores (orange), and Z2-scores (red), are shown for all 13 kinases that had a crystal structure available. Numbers on top indicate the number of crystal structures for each kinase.

binding). This normalization was done with respect to the actives from the benchmark set for a given kinase. The early enrichments of active compounds based on Z2-scoring increased performance for 10 of the 13 targets (**Figure 5.2**). The best (ensemble of) models per target are listed in **Table S5.2**. The best performances of the main on-targets RET, AURKA, and PAK1 were reached by Z2-scoring. The performance of PAK1 was very good with BEDROC ($\alpha = 160.9$) 0.97. However, it should be noted that the number of compounds in this benchmark set was low due to lack of data (63 active compounds, 10 inactive compounds and 3886 decoys). Therefore, this model may not be representative when applied to a virtual screening set. The best BEDROC performances of RET and AURKA were 0.56 and 0.52 respectively. Furthermore, the overall performance for these targets were good with ROC 0.85 for RET and ROC 0.82 for AURKA.

Prospective structure-based docking

The virtual screening compounds resulting from QSAR filtering were docked into the protein structures that resulted in the best performances in benchmarking. Not the entire virtual screening set could be docked on all kinases because of time constraints. Therefore the top 250,000 compounds for RET, AURKA, and PAK1, with highest active-class probabilities, were selected for docking. This included all active-predicted compounds for RET and AURKA (167,828 for RET and 315 for AURKA), and additional, consecutively ranked compounds that did not reach the QSAR activity threshold (active class probability > 0.5). For PAK1 the top 250,000 compounds were selected from a total of 298,163 active-predicted compounds. Subsequently, the compounds were assigned a structure-based Z2-score, with the exception of S6K, MKNK1, and PDK1 for which SPLIF or docking score gave the best BEDROC ($\alpha = 160.9$) performance and thus Z2-scores were replaced with the respective score. The best scoring method for S6K underperformed with a BEDROC of 0.24. Although this model was not discarded immediately, the poor performance was taken into account later when reliability weights were assigned to the corresponding method and kinase.

Binding pose metadynamics

Binding pose metadynamics^{23,24} was performed to re-score the docked poses and scores of compounds. Since binding pose metadynamics is a time-consuming modeling technique, only the main on-targets (RET, AURKA, and PAK1) were subjected to this method. Binding pose metadynamics measures the persistence of ligand-protein interactions and the movements of the ligand's heavy atoms, which are sampled during variation of the complex's free energy states throughout the simulation. The result of binding pose metadynamics is a metadynamicscomposition score. This composition score was calculated for the top 100 compounds (based on docking score) from the benchmark set of each included kinase. The docking scores from which these top 100 compounds were chosen were derived from a single protein structure per target to allow for easy and direct comparison. The selected protein structures had the best actives enrichment based on docking and included 2IVU for RET, 2BMC for AURKA, and 4EQC for PAK1. The composition score was added to the docking score, resulting in a combined score. The actives enrichment for the targets was re-evaluated using this new score. It was observed that performance (based on the top 100 compounds) of PAK1 did not increase (ROC+BEDROC difference 0.01). However, actives enrichment for RET and AURKA increased with performances (ROC+BEDROC) 1.27 for RET and 1.76 for AURKA, compared to initial (docking) performance of 1.26 for RET and 0.77 for AURKA. Therefore, for RET and AURKA the docking scores of virtual screening compounds were re-scored with metadynamics-composition scores, resulting in an indirect re-ranking of the top 100 virtual screening compounds, which was based on Z2-scores.

Polypharmacology ranking of compounds

The virtual screening compounds were scored with both a statistical model score (PCM score) and structure-based score. A final compound rank per target was constituted by adding a weight to the predictions made by statistical models and structure-based models (**Table 5.2**). The ranking order of compounds for off-targets, i.e. kinases compounds should not interact with, was reversed: ranking compounds with high predicted activity as lowest, and compounds with worst predicted activity as highest. The weights of the statistical PCM model were based on the



ROC score corrected for the size of the training dataset per target. The structure-based models were considered to be better suited to select novel chemistry and were therefore assigned higher weights than the PCM model weights: structure-based models were attributed more weight compared to equally performing PCM models (same ROC). The weights of the structure-based models were calculated by taking the sum of BEDROC+ROC. These weights were reduced by penalizing the models when induced-fit structures were used and when the numbers of compounds in the benchmark sets were insufficient (for details see Methods). The weights were subsequently used to rank the compounds per target: Structure-based Z2-scores were multiplied by the structure-based weight, PCM predicted class probability was multiplied by the statistical model weight, and as final step the derived scores were summed up to retrieve a final rank per target.

Based on overall target weights (**Table 5.2**), predictions for kinases LKB1, ERK8, PAK3, TAK1, and PIK3CA, were not very reliable. However, predictions for the more important main ontargets RET, AURKA, and PAK1 were considered to be reliable. The workflow was evaluated by the selection and experimental validation of compounds for one of the main on-targets: RET. The consensus approach was used to select 15 highly ranked virtual screening compounds for RET (all compounds had PCM score > 0.4 and structure-based score Z2 < 0), which were validated in vitro. Moreover, the performance of the different approaches was compared on the RET models by additionally selecting compounds based on the predictions of only statistical models, and only structure-based models.

Table 5.2. Weights assigned to each target per modeling technique.

Model		On-ta	argets				off-targets		
Panel 1 kinases	RET	BRAF	SRC	S6K	MKNK1	TTK	ERK8	PDK1	PAK3
PCM	0.54	0.14	0.90	0.67	0.13	0.41	-0.02	0.35	0.70
Structure Based	1.44	1.66	1.56	0.48	1.42	1.79	n.a.	1.45	n.a.
Total	1.98	1.80	2.46	1.15	1.55	2.20	-0.02	1.80	0.70
Panel 2 kinases	AURKA	PAK1	FGFR1	LKB1	TAK1	PIK3CA	PAK3		
PCM	0.93	0.50	0.98	0.28	0.16	n.a.	0.70		
Structure Based	1.34	1.23	1.62	0.04	0.46	0.97	n.a.		
Total	2.27	1.73	2.60	0.32	0.62	0.97	0.70		

Experimental validation

The models were evaluated by experimental validation of predicted actives for RET kinase. A total selection of 46 compounds was purchased and validated for RET inhibition. These compounds were selected by different criteria: 6 compounds based on predicted activity by QSAR and PCM modeling, 25 compounds based on structure-based docking (including rescoring by binding pose metadynamics) and SPLIF scoring, and 15 compounds based on consensus scoring of statistical and structure-based models.

Compounds that were selected using only statistical models satisfied the set active-class threshold criteria of both QSAR (active probability > 0.6) and PCM (active probability > 0.5). The structure-based thresholds were docking score < -8 and SPLIF score > 0.25. Additionally, structure-based compounds with a docking score < -10 were selected that did not necessarily adhere to the SPLIF score criteria. Furthermore, compounds that were selected based on consensus scoring fitted the following thresholds: statistical predictions PCM > 0.4, and structure-based score Z2 < 0. The compounds that were selected based on structure-based predictions or with consensus scoring were additionally inspected visually by checking the 3D docking pose and interaction with the hinge region.

The 46 compounds were first tested using single point measurements at two concentrations: 10 μ M and 0.1 μ M (**Table S5.3**). The top ten compounds showing RET inhibition (RET activity < 80%, concentration 10 μ M), were based on either consensus scoring or structure-based scoring. None of these compounds was from statistical scoring only. Five compounds showed inhibitory activity (RET activity < 60%) for RET at a concentration of 10 μ M, of which one compound also showed slight RET inhibition at concentration 0.1 μ M (RET activity < 100%). The activities of the four most potent hits were assessed more accurately by a potency determination in triplicate, yielding pIC₅₀ values. The inhibitors were modestly active. ZINC33008650 was the best inhibitor with a pIC₅₀ of 5.1 \pm 0.1, followed by ZINC72312837 (pIC₅₀ 4.6 \pm 0.2), ZINC12324934 (pIC₅₀

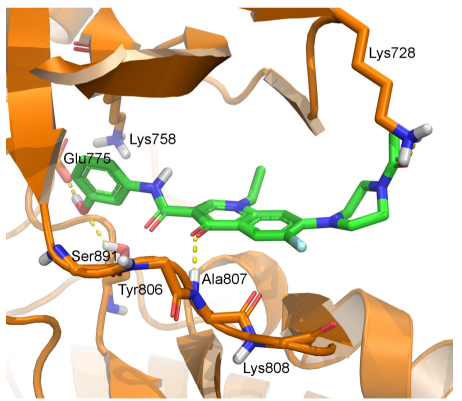


Figure 5.3. ZINC12324934 (green) docked into the orthosteric binding pocket of RET (orange) (PDB:2IVU). Hydrogen bonds are displayed as yellow dotted lines.



4.6 \pm 0.2), and ZINC9518200 (pIC₅₀ 4.0 \pm 0.2). The docked pose of ZINC12324934 in RET suggests that the compound is an orthosteric binder with potential to bind to the hinge region, as evidenced by a hydrogen bond interaction with the backbone of Ala807 (**Figure 5.3**). Additional hydrogen bonds are observed between the ligand and Glu775, Ser891, and Lys728. Although the inhibitors showed modest activity, their chemical diversity compared to known RET inhibitors was high since the compounds were pre-filtered on novelty by selecting on Tanimoto similarity < 0.4 (compared to inhibitors with pChEMBL value \geq 5).

Bioactivity spectra prediction

The bioactivity profile of the five most active inhibitors for RET was predicted for all kinases in the panel. The bioactivity spectra for both on- and off-targets are shown in **Table 5.3**, where positive values indicate binding and values below and including zero indicate no binding. Based on the entire predicted bioactivity spectra, the most potent RET inhibitors comply better with inactivity on off-targets than activity on on-targets. However, the predictions are not equally reliable for each kinase (based on the previously assessed weights shown in **Table 5.2**). Nevertheless, assuming that the predictions of the kinases that scored equally well or better compared to RET (weight 1.98) are the most accurate, conclusions can be drawn for on-target SRC and off-target TTK (weights 2.46 and 2.20, respectively). Although the RET inhibitors were not predicted active for SRC, inactivity was predicted for off-target TTK. The compounds were selected based on bioactivity for RET, and as a result may not show optimal bioactivity spectra. However, the novelty filter to which the compounds were subjected prior to virtual screening included all of the kinases in the panel (as opposed to only RET). Therefore, all predicted interactions might indicate novel starting points for future research.

Compound	On-targets				Off-targets				
	RET	BRAF	SRC	S6K	MKNK1	TTK	ERK8*	PDK1	PAK3*
ZINC33008650	1	0	0	0	-2	-2	0	-2	0
ZINC12324934§	0	0	0	0	0	0	0	0	0
ZINC9518200	2	0	0	0	-1	-1	0	-1	0
ZINC72312837	2	0	0	0	-1	-1	0	-2	0
ZINC65184824	1	-1	0	0	-2	-1	0	-2	0

Table 5.3. Predicted bioactivity spectra for the panel 1 kinases of the five most potent RET inhibitors.

Manual inspection of hits

The four most potent hit compounds resulting from virtual screening and experimental validation were inspected based on their novelty and patentability. Novelty was re-evaluated by similarity searching (Tanimoto ECFP6) on a more strict threshold for RET compounds (pChEMBL ≥ 4) in the most recent version of ChEMBL (version 25) and patentability was checked in SureChEMBL²⁵ (similarity > 0.9). The most similar RET actives were: CHEMBL1979093 for ZINC33008650 (similarity 0.24, RET pChEMBL 7.2), CHEMBL1983715 for ZINC12324934 (similarity 0.18, RET pChEMBL 6.9), CHEMBL1977134 for ZINC9518200 (similarity 0.20,

^{*}Predictions based on limited structure-based data (no crystal structure available).

Sompound was selected on RET docking score only, therefore no Z2-score was available for this compound and no structure-based weight could be assigned.

Table 5.4. Novel RET inhibitors and their most similar RET actives in ChEMBL based on Tanimoto (ECFP6).

Compound	Closest similar (pChEMBL ≥ 4)	Similarity (Tanimoto
F	ì	ECFP6) 0.24
——————————————————————————————————————	NH ₂	
ZINC33008650	CHEMBL1979093	
pIC ₅₀ 5.1 ±0.1	RET pChEMBL 7.2	
Ô	Ö	0.29
ифин		
	NH ₂	
ZINC72312837 pIC ₅₀ 4.6 ±0.2	CHEMBL1965570 RET pChEMBL 7.6	
	0	0.18
L L L L L L L L L L L L L L L L L L L		
ZINC12324934 pIC ₅₀ 4.6 ±0.2	CHEMBL1983715 RET pChEMBL 6.9	
	5.	0.20
H ₂ N NH NH		
ZINC9518200 pIC ₅₀ 4.0 ±0.2	CHEMBL1977134 RET pChEMBL 8.4	



RET pChEMBL 8.4), and CHEMBL1965570 for ZINC72312837 (similarity 0.29, RET pChEMBL 7.6) (**Table 5.4**). None of the compounds was listed as patented in SureChEMBL.

Additionally, novelty of the chemical scaffolds was checked, which were compared based on Bemis-Murcko²⁶ scaffold trees with known active compounds for RET in ChEMBL. None of the four hit compounds was clustered in the same scaffold cluster as a known active, supporting the novelty of the inhibitors. Although the hit compounds were identified as modestly active binders, further SAR around these inhibitors may yield more potent derivatives. The novel scaffolds may be explored along different vectors to enhance affinity of the ligands and may reveal a new chemical space for RET inhibitors.

Discussion

An elaborate virtual screening process was constituted in which both statistical and structure-based models were applied that were all carefully tuned and validated. Statistical models were validated using as many compound data as possible, with attention paid to chemical diversity of the subsets in cross-validation. Structure-based models were slower than the statistical models and therefore smaller benchmark sets were used for validation of these models. These benchmark sets were composed of diverse actives to cover a large chemical space, irrespective of the smaller size of the dataset. By application of statistical QSAR models and structure-based docking successively, computational time was used efficiently in virtual screening: compounds that had the least predicted activity probability for the main on-targets based on the QSAR models were excluded from docking. The consensus scoring approach, in which statistical PCM scores were combined with structure-based predictions, resulted in sets of active-predicted compounds per kinase. However, compounds were excluded from bioactivity modeling if the respective kinase had a poor reliability weight. This reliability weight was based on the performances per model per target derived from the benchmarking steps.

It was observed that the lack of compound data resulted in lower performance of statistical models in particular, as seen for kinase ERK8. Although structure-based models also performed worse for targets with less compound data and less crystal structures, this could partially be resolved by creation of additional protein structures using induced-fit docking and alternative scoring with SPLIF and Z2. However, alternate structures may also have been created using more elaborate ligand-protein sampling methods such as molecular dynamics.²⁷ Interestingly, performance between different structures of the same kinase varied greatly, rendering the benchmarking of multiple protein structures necessary.

Based on the final reliability scores, kinases RET, BRAF, SRC, MKNK1, TTK, PDK1, AURKA, PAK1, and FGFR1, are best used in bioactivity modeling, whereas predictions for kinases S6K, ERK8, PAK3, LKB1, TAK1, and PIK3CA, may not be accurate enough and thus should be excluded for now. Although the models of the first kinases are considered reliable, identification of compounds that fit the desired bioactivity profiles on these targets remains challenging. Since hit rates are rarely 100%, compounds with a well-predicted profile would need experimental validation. The hit rate for the experimental validation for RET in this study was 11% over 46 compounds, using pIC $_{50} \ge 4$ (or RET activity < 60% at a concentration of 10 μ M) as a threshold for active compounds. On the hypothesis that similar hit rates will be achieved for the other

kinases, the probability of a compound being active on a number of kinases thus decreases with every additional target. Therefore, the number of experimentally validated compounds should be expanded, to increase the chance of finding a compound that fully fits the desired bioactivity profile.

The five hits that proved active on RET kinase were found with either consensus scoring or structure-based modeling. Although the number of compounds validated using statistical models is smaller, it is plausible that statistical modeling by itself is not a strong enough predictor to identify really novel compounds.²⁸ Since the virtual screening compounds were pre-filtered on novelty (Tanimoto ECFP6 < 0.4) with known actives, the models were challenged with identification of chemical dissimilar actives.²⁹ This is a difficult task for statistical models as they rely on chemical patterns of known actives, and therefore dissimilar compounds may lie outside the applicability domain of these models.²⁹ To resolve this issue, the chemical space can be expanded by addition of chemical diversity.8 Although this requires biological experiments to validate (in)activity of new compounds, an iterative screening approach may be applied to expand the chemical space efficiently and cost-effectively.³⁰ Nevertheless, identification of novel chemistry without the need for addition of much experimental data may be achieved through structure-based modeling, as these models are not particularly biased towards known chemistry. However, bias in structure-based models may indirectly be present, as the best performing protein structures were based on enrichment of known actives. Nevertheless, this is a minor issue compared to the limited scope and bias of chemical space of statistical models.

The applied workflow in this study employs a filtering step through which compounds are discarded that were categorized as inactive by statistical QSAR models. Based on the previous statement on applicability domain, it is assumed good inhibitors can be wrongly categorized as "inactive" and neglected by the statistical model. One might reason that docking of all compounds may be a more effective method to identify novel inhibitors, as docking of 170 million compounds in proteins AmpC β-lactamase and the D4 dopamine receptor resulted in novel chemical scaffolds.³¹ However, docking of millions of compounds on multiple proteins is a very time consuming task (~170,000 compounds per day for 1 RET crystal structure (24 cores)), which is unfeasible without the required computational resources. Thus, considering the scope of the task, the possibility that the QSAR models discard 'good' compounds is accepted, as the QSAR models decrease the runtime of the workflow and make the task comprehensive. As a final remark, it should be mentioned that the workflow applied to kinases did not implement orthosteric or allosteric binding of compounds. Therefore, inhibitors were not tuned to bind to the DFG-in or DFG-out kinase conformation, a conformational change that influences inhibitor binding greatly.³² Although structure-based docking was focused on the ATP-binding pocket, the most optimal crystal structures were selected based on benchmark sets that were not filtered for DFG-out and allosteric binders. As a consequence, crystal structures may have been selected that enriched DFG-out binders better than DFG-in binders. Moreover, since the statistical models were trained on sets that were not filtered for DFG-out binders, these models were also not able to distinguish DFG-in from DFG-out binders. As a result, it is undetermined whether the five hits from the screening workflow bind to the DFG-in conformation of RET. The docked poses of the hits do not constitute optimal hinge-binding, suggesting that it is plausible that they may be DFG-out binders. Moreover, two of the five most potent hit compounds contained a ureamotif, a motif that is often associated with DFG-out binders. 32,33 To capture the binding type of



compounds, machine learning models could have been used to predict the type of compound as an additional score, something that will be considered for future work.^{34,35}

Conclusion

An extensive workflow was designed to predict compound activity in kinases and to model the compounds' bioactivity spectra in kinases. The workflow can easily be expanded to other targets as well. By combining statistical and structure-based modeling, processing speed was optimized, while the accuracy of predictions was preserved. Every single kinase target was validated separately, which enabled reliability weights to be assigned to the predictions for every target. The workflow was experimentally validated by testing predictions made for RET kinase, a target with a good reliability score. A selection of 46 compounds was tested in vitro, of which 5 compounds showed RET inhibition (activity < 60%) at a concentration of 10 μ M. The four most potent inhibitors had pIC₅₀ values ranging from 4.0 to 5.1. The Tanimoto similarities (ECFP6) of these inhibitors with known RET actives was \leq 0.29. Moreover, the compounds contained unique chemical scaffolds, underscoring the true novelty of these inhibitors.

Methods

Dataset statistical models

Training and validation sets for statistical models were constituted from compound information derived from ChEMBL (version 23)11, publicly available sets from Eidogen12, and ExCAPE-DB13. The compounds with experimental bioactivity for any kinase were filtered on molecular weight < 700, duplicates were removed, and compounds were standardized using BIOVIA Pipeline Pilot 2016³⁶: salts were removed, largest fragment was kept, stereochemistry and π -systems were standardized, and charges were neutralized. The resulting set contained 512 kinases and 123,246 bioactivities. For training and validation of the classification models the threshold for 'active' compounds was set at pChEMBL/pK₁/pIC₅₀/pEC₅₀ \geq 6.5 as we did previously.³⁷ Compounds that did not reach this threshold were termed 'inactive'. All compounds per target were divided into five subsets using clustering with the Cluster Molecules component (FCFP4 clustering) in Pipeline Pilot 2016. Compounds were clustered into five clusters per each kinase family (same L2 level in ChEMBL). First, compounds were separated based on activity (active when pChEMBL > 6.5) and then clustered into five clusters, after which active and inactive compounds were combined again. To ensure that every panel kinase was represented in each cluster, clustering into five clusters was done in two steps: the panel kinase was clustered first, followed by coclustering of compounds from related kinases into the same cluster. The biggest cluster, or subset, was used as fixed training set, while the other four subsets were rotationally used as test or training set in 4-fold cross-validation. Datasets used in QSAR modeling only contained information of the respective kinase. In PCM modeling, additionally data of related kinases was included based on L4 classification in ChEMBL.

Dataset structure-based docking

For each kinase target, with the exception of PAK3 and ERK8 because of lack of a crystal structure, a benchmark set was created to validate structure-based docking for each target. The benchmarking set for structure-based docking included active compounds, inactive compounds, and decoy compounds, which were derived from ChEMBL (version 23) and the

DUD-e webserver¹⁵. Compounds with reported pChEMBL value for the challenge kinases, with confidence score 9, assay type B, and molecular weight < 550, were standardized using BIOVIA Pipeline Pilot 2016³⁶ by removing salts and keeping the largest fragment. An activity gap was realized to better distinguish active and inactive compounds: compounds with a pChEMBL value > 6.5 were assigned the label 'active', and compounds with a pChEMBL value > 4 and < 5.5 were labeled 'inactive'. Compounds with pChEMBL value ≤ 4 were excluded as inactive compounds to limit the number of inactives. Exceptions in the thresholds were made for activity labeling of compounds for targets RET and MKNK1: for RET compounds were labeled as inactive when pChEMBL value > 4.5 and < 6.5, and for MKNK1 compounds were defined as active when pChEMBL value > 6 to increase the fraction of active compounds. The thresholds used for RET were not intended to alter the number of (in)active compounds; RET was used as exploratory case and therefore for this kinase 'initial' thresholds were used. However, the 'general' threshold for all kinases was adapted since this yielded increased performance.

Scaffold trees were generated for the compounds based on Bemis-Murcko scaffolds²⁶ and the compound with the highest pChEMBL value per scaffold class, per activity class, per kinase, was kept. Compounds labeled as active were clustered into 100 clusters using the Cluster Molecules component (k-means) in BIOVIA Pipeline Pilot 2016. From the resulting clusters, only the cluster centers were kept. Kinases that had less than 100 active compounds available were excluded from this clustering step. The resulting active compound sets of maximum 100 compounds per kinase were used to generate decoys for each target kinase using the DUD-e webserver. For target PIK3CA decoys were generated based on only 80% of the active molecules as DUD-e decoy generation failed for the remaining fraction. The decoys were considered as inactive compounds when used in model validation. The collected actives, inactives, and decoys were prepared for docking using LigPrep from Schrödinger²³.

Screening dataset

Compounds for virtual screening were extracted from the ZINC15 compound library¹6. These compounds were selected based on "in stock" status and were filtered on a maximum of one Lipinski's rule of five violation. Additionally, the screening set was filtered on novelty by only including compounds with Tanimoto similarity (ECFP6) < 0.4 compared to known 'active' compounds (pChEMBL value ≥ 5) in ChEMBL (version 23) for the kinases of the respective panel. For panel 2, covering AURKA, PAK1, FGFR1, and LKB1, PAK3, TAK1, and PIK3CA, the screening set was additionally filtered on PSA < 75 to allow permeability of the blood-brain barrier. The compounds in the screening dataset were additionally prepared for structure-based docking by using LigPrep.

Statistical modeling - QSAR

Single target QSAR models were constructed for the main on-target kinases RET, AURKA, and PAK1. QSARs were built using BIOVIA Pipeline Pilot 2016 (version 18.0.1.1604). Models were trained on compound descriptors FCFP4 (3000 most frequent bits) and 86 physicochemical descriptors (**S1**). The following settings were applied in training categorical Random Forest QSAR models: 1000 trees, log2(m) number of descriptors, equalized class sizes and seed 12345. The classification threshold for active compounds was set at pChEMBL > 6.5.



Statistical modeling - PCM

PCM modeling was performed on all 512 kinases in the statistical modeling dataset. A multi-target PCM model was constructed using a random forest classifier in scikit-learn³⁸. Compound descriptors were the same as used in QSAR modeling. Protein descriptors were calculated based on full sequence alignment derived from kinase.com (with a total of 1567 alignment positions including gaps).^{39,40} The residues in the alignment were converted to three z-scales and an additional mean value for each of the three z-scales was added per sequence, resulting in a total of 4704 protein descriptors per kinase. Gaps were included in these descriptors, and were assigned a value of "0" for all three z-scales as was done previously.^{41,42}

The random forest model was of high complexity due to the high dimensionality of the data, which includes the compounds' physiochemical properties, FCFP4 descriptors, and protein descriptors. The complexity of the random forest models was reduced by imposing constraints on parameters, such as the number of trees and maximum depth of the trees. The hyperparameters of the random forest model were optimized by utilizing random search, which evaluates the performance of the algorithm using different and randomly chosen configurations with crossvalidation. Random search can be considered a simple form of Automated Machine Learning 19,43 and has shown to outperform other basic methods of hyperparameter optimization, such as grid search. 44 Approximately 500 random configurations were evaluated, resulting in an improvement of on average 7% AUROC over the default configuration (300 trees, log2(m) features at every node in the tree, no maximum tree depth, a minimum of 2 samples to consider a node for splitting, and bootstrap enabled) in 4-fold cross-validation. The settings that corresponded with the best model in random parameter optimization were applied in model training on the full data set, which was used in virtual screening. The final model consisted of 300 trees (fixed) with 43 features at every node in the tree, a maximum tree depth of 99, a minimum of 12 samples to consider a node for splitting, and bootstrapping disabled.

Structure-based docking

X-ray structures of all challenge proteins were extracted from the PDB⁴⁵ (except for ERK8 and PAK3 as no structure was available at the time). Crystal structures lacking co-crystalized orthosteric small-molecule ligands were discarded. The remaining structures were prepared for docking with the Protein Preparation tool from the Schrödinger 2017-4 suite after removing any other components than the protein, orthosteric ligand, and binding pocket ions. The 'add missing side chains' option was used, waters were removed, hydrogens were added, and disulphide bonds were created. The crystal structures were superposed per target and co-crystalized ligands were removed from the binding site. The grid for docking was determined for each target by the center of one of the co-crystalized ligands (box size xyz = 35 Å). Compounds were docked into the binding pocket using the Schrödinger 2017-4 suite²³ and the OPLS3 force field⁴⁶ with standard precision (SP) and standard settings. A maximum of ten poses per compound (per target) was generated.

Induced-fit docking

Induced-fit docking, as implemented in the Schrödinger 2017-4 suite, ²³ was applied to kinases LKB1, S6K, and MKNK1. These kinases showed poor enrichment of actives when docked into the available crystal structures (sum of ROC and BEDROC < 1). For S6K five active

compounds, for MKNK1 ten active compounds, and for LKB1 six active compounds were docked using induced-fit docking (see **Table S5.4** for list of compounds). The crystal structures selected for induced-fit docking were 2HW6 for MKNK1, 2WTK for LKB1, and 3WF5, 3A62, and 4RLO for S6K. The resulting protein conformations were used as alternative protein structures in addition to the original crystal structures.

Structural protein-ligand interaction fingerprints (SPLIFs)

Structural protein-ligand interaction fingerprints (SPLIFs)⁴⁷ were calculated for a maximum of ten poses per compound that were retrieved from structure-based docking. The co-crystalized ligands of the protein structures were used as reference in SPLIF calculations to derive Tanimoto-like SPLIF scores. Five protein structures per kinase were selected for SPLIF generation. These structures were selected based on diversity of their co-crystalized ligands and best active-enrichment based on docking scores. The diversity of the co-crystalized ligands was assessed using affinity propagation clustering based on FCFP6 similarity⁶. One protein structure was selected from every corresponding cluster based on best docking score performance until a maximum of five protein structures was reached. Subsequently, Tanimoto-like SPLIF scores were calculated for the compounds in the benchmark sets for each of the selected protein structures. For each compound (maximum 10 poses) the best SPLIF score was used to calculate actives enrichment based on SPLIF scores.

For RET and PIK3CA exceptions were made in the selection of benchmark proteins for SPLIF because only four and three clusters were generated, respectively. Therefore, similar co-crystalized ligands and their corresponding proteins were also selected to get a total of five protein structures per target. Furthermore, for RET structure 2IVV instead of the co-crystalized ligand, CHEMBL3775169 was used as a reference.

Binding pose metadynamics

A metadynamics-composition score¹⁰ was calculated with the binding pose metadynamics tool in the Schrödinger 2017-4 suite,²³ for compounds and poses derived from structure-based docking. The top 100 compounds for RET (PDB: 2IVU) and AURKA (PDB: 2BMC) were selected, based on best docking scores. The protein structures were prepared for metadynamics simulations by capping the termini and the run time of each simulation was set at 10 ns. The resulting metadynamics-composition scores were added to the existing docking scores to re-rank the top 100 compounds in the benchmark set, and to re-rank the top 100 compounds in the screening set.

Ensemble scoring in structure-based modeling

For every target, protein models for virtual screening were selected based on ensemble scoring of docking and SPLIF scores. The combination, or ensemble, of protein models that resulted in the best actives enrichment was used in virtual screening. The validated ensembles were based on docking scores of the top five enriching models and all five SPLIF models per target. Different ensembles were tested for each target including: docking scores only, SPLIF scores only, and docking and SPLIF scores combined. The performances of the ensembles were evaluated using Z2-scoring by averaging over the top two Z-scores.⁴⁷ Prior to ensembling, the Z-scores per compound were normalized towards the actives from the respective benchmark set. Docking



scores were normalized to Z-scores by subtracting the mean docking score (mean over docking scores from all actives in the benchmark set) from the docking score of the test compound, and subsequently dividing by the standard deviation of the (benchmark) actives' docking scores. The same approach was used in normalization of SPLIF scores to Z-scores. However, the SPLIF scores were first multiplied by -1 to change the vector into the same direction as the docking scores (better binder, more negative score).

Compound ranking per target

The compounds in virtual screening were ranked for ensembles of targets based on the scores derived from PCM modeling and structure-based ensemble score. The PCM predictions were given a confidence weight between 0 and 1 based on the prediction performance per target (with 0 being no confidence, and 1 being highest confidence). The confidence score for the PCM model could be calculated using the following equation:

$$confidence\ weight_{PCM} = (ROC*2-1)*\frac{\sqrt{nr\ of\ samples}}{\sqrt{\max nr\ of\ samples}}$$

The PCM confidence score was based on the number of compounds (of the corresponding target) in training and the ROC score derived from 4-fold cross-validation: ROC * 2 - 1 (to normalize that 0.5 corresponds with a weight of 0 and 1 corresponds with a weight of 1), multiplied by the weight based on the number of samples for the target in the training data (calculated as sqrt(number of samples)/sqrt(maximum number of samples).

The weights of the structure-based predictions were calculated by taking the total sum of ROC and BEDROC per target⁴⁸, consequently giving the structure-based predictions more weight than the statistical PCM model predictions. However, structure-based models were penalized (by multiplying with 0.5) when induced-fit models were created for the kinase. Moreover, kinases containing less than 100 actives in the benchmark set were penalized by multiplying the structure-based weight with the fraction of number of actives (with a fraction of 1 being 100 actives, and a fraction of 0.1 being 10 actives).

Finally, the obtained weights for PCM and structure-based models were applied in calculating the compound rank per kinase.

compound rank

$$= confidence\ weight_{PCM}* active\ class\ probability$$

$$+\ confidence\ weight_{structure-based}* - Z2\ score$$

Compound predictions were multiplied by the subtotal weights and summed up (statistical model weight * statistical PCM prediction + structure-based model weight * structure-based Z2 score), resulting in a final compound rank per target.

Protein kinase assay

A selection of 46 compounds was experimentally validated for inhibitory activity against RET. The compounds were ordered via Mcule Inc.⁴⁹, (Budapest) and purchased from ChemBridge, ChemDiv, ChemScene, Enamine, Life Chemicals, and Vitas M Chemical Limited. The assays were performed by ProQinase GmbH⁵⁰, Germany. First, the compounds were tested (n=1) at final assay concentrations 10 µM and 0.1 µM. The five resulting hits (< 60% remaining RET activity at concentration 10 μM) were re-evaluated by determination of IC₅₀ values (n=3). RET WT activity was measured using a radiometric protein kinase assay (33PanQinase® Activity Assay). All kinase assays were performed in 96-well FlashPlatesTM from PerkinElmer (Boston, MA, USA) in a 50 ml reaction volume. The reaction cocktail was pipetted in 4 steps in the following order: 1) 20 ml of assay buffer. 2) 5 ml of ATP solution (in H₂O). 3) 5 ml of test compound (in 10 % DMSO), and 4) 20 µl enzyme/substrate mix. The assay for all protein kinases contained 70 mM HEPES-NaOH pH 7.5, 3 mM MgCl₂, 3 mM MnCl₃, 3 mM Na-orthovanadate, 1.2 mM DTT, 50 μg/ml PEG20000, 1 μM ATP, [γ -³³P]-ATP (approx. 1.91 x 1005 cpm per well), 40 ng/50 μl protein kinase, and 0.125 μg/50μl poly(Glu, Tyr)4:1 substrate. The reaction cocktails were incubated at 30°C for 60 minutes. The reaction was stopped with 50 ml of 2 % (v/v) H₂PO₄, plates were aspirated and washed two times with 200 ml 0.9 % (w/v) NaCl. Incorporation of ³³Pi was determined with a microplate scintillation counter (Microbeta, Wallac).



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Supporting Information

Table S5.1. List of 86 physicochemical descriptors.

	Physicochem	ical descriptors	
CMP_ATOMCOUNT_C	CMP_NUM_CHAINASSEM- BLIES	CMP_ATOMCOUNT_I	CMP_ATOMS_O_FRAC
CMP_ATOMCOUNT_C_sp3	CMP_ATOMCOUNT_META- LATOMS	CMP_Molecular_solubility	CMP_ATOMS_HETEROATOM_ FRAC
CMP_ATOMCOUNT_C_sp2	CMP_BONDS_PIBONDS	CMP_LogP	CMP_ATOMS_S_FRAC
CMP_ATOMCOUNT_C_sp	CMP_BONDS_STEREOATOMS	CMP_MOLECULAR_SUR- FACEAREA	CMP_ATOMS_P_FRAC
CMP_ATOMS_C_sp3_FRAC	CMP_BONDS_STEREOBONDS	CMP_MOLECULAR_SASA	CMP_ATOMS_HALOGEN_ FRAC
CMP_ATOMS_C_sp2_FRAC	CMP_BONDS_SINGLEBONDS	CMP_MOLECULAR_POLAR- SASA	CMP_ATOMS_OTHER_FRAC
CMP_ATOMS_C_sp_FRAC	CMP_BONDS_DOUBLEBONDS	CMP_NUM_ALIPHATICRINGS	CMP_ATOMS_STEROATOM_ FRAC
CMP_FORMALCHARGE	CMP_BONDS_TRIPLEBONDS	CMP_BONDS_HEAVYATOMS	CMP_ATOMS_POSITIVE_FRAC
CMP_ATOMCOUNT_HEAVY	CMP_NUM_ALIPHATICSIN- GLEBONDS	CMP_BONDS_HYDROGENS	CMP_ATOMS_NEGATIVE_ FRAC
CMP_BONDS_TOTAL	CMP_NUM_ALIPHATICDOU- BLEBONDS	CMP_BONDS_SINGLE_FRAC	CMP_ATOMS_H_ACCEP- TOR_FRAC
CMP_ATOMCOUNT_HYDROGENS	CMP_NUM_HYDROGENBONS	CMP_BONDS_DOUBLE_FRAC	CMP_ATOMS_H_DONOR_ FRAC
CMP_BONDS_EXPLICIT	CMP_NUM_TERMINAL- ROTOMERS	CMP_BONDS_TRIPLE_FRAC	CMP_RIGIDITY_INDEX
CMP_ATOMCOUNT_POSITIVE	CMP_ATOMCOUNT_H_AC- CEPTORS	CMP_BONDS_AROMATIC_ FRAC	CMP_MOLECULAR_WEIGHT
CMP_ATOMCOUNT_NEGA- TIVE	CMP_ATOMCOUNT_H_DO- NORS	CMP_BONDS_BRIDGE_FRAC	CMP_CLASS_LIPINSKI_PASS
CMP_BONDS_RING	CMP_ATOMCOUNT_H	CMP_BONDS_STEREO_FRAC	CMP_NUM_Ro5_Violations
CMP_BONDS_ROTATABLE	CMP_ATOMCOUNT_N	CMP_BONDS_RING_FRAC	CMP_RO3_PASS
CMP_BONDS_AROMATIC	CMP_ATOMCOUNT_O	CMP_BONDS_ALIPHATIC_ FRAC	CMP_MOLECULAR_POLAR_ SURFACEAREA
CMP_BONDS_BRIDGE	CMP_ATOMCOUNT_F	CMP_BONDS_ROTATABLE_ FRAC	CMP_MOLECULAR_POLAR_ SURFACEAREA_FRAC
CMP_NUM_RINGS	CMP_ATOMCOUNT_P	CMP_ATOMCOUNT_HALOGENS	CMP_FULL_MWT
CMP_NUM_AROMATICRINGS	CMP_ATOMCOUNT_S	CMP_ATOMS_C_FRAC	CMP_INORGANIC_FLAG
CMP_NUM_RINGASSEMBLIES	CMP_ATOMCOUNT_Cl	CMP_ATOMS_H_FRAC	
CMP_NUM_CHAINS	CMP_ATOMCOUNT_Br	CMP_ATOMS_N_FRAC	



Table S5.2. Selected structure-based models per target.

Protein	PDB	Model	B E D R O C (α=160.9)	ROC	Sum	
Challenge 1			•			
RET	2IVU	Docking+SPLIF	0.56	0.85	1.41	
	5FM2	Docking				
	2IVV*	Docking+SPLIF				
	5FM3	Docking				
	2X2K	Docking				
BRAF	5HID	Docking+SPLIF	0.75	0.9	1.66	
	1UWH	Docking	1			
	4DBN	Docking	1			
	4G9R	SPLIF				
SRC	4MXX	Docking+SPLIF	0.64	0.92	1.56	
	2BDJ	Docking				
	1YOL	Docking				
	2H8H	SPLIF	1			
S6K	3A62 IFD #74	SPLIF	0.24	0.72	0.96	
MKNK1	2HW6 IFD #25	SPLIF	0.71	0.86	1.57	
PDK1	3QD0	Docking	0.66	0.87	1.53	
TTK	4C4I	Docking	0.81	0.98	1.79	
	5N84	Docking				
	5N87	SPLIF				
	5NA0	SPLIF				
Challenge 2			•			
AURKA	2BMC	Docking	0.52	0.82	1.34	
	4B0G	SPLIF				
PAK1	4EQC	Docking+SPLIF	0.97	0.98	1.95	
	6B16	SPLIF				
FGFR1	3GQL	Docking	0.68	0.94	1.62	
	4RWL	Docking				
	3TT0	Docking				
	4NKA	Docking				
LKB1	2WTK IFD #83	Docking+SPLIF	0.67	0.81	1.48	
	2WTK IFD #34	Docking+SPLIF				
TAK1	4L53	Docking+SPLIF	0.64	0.76	1.40	
	4L3P	Docking				
	5JGB	SPLIF				
PIK3CA	4JPS	Docking	0.99	0.62	1.61	
	5DXH	SPLIF]			

Table S5.3. Bioactivities of 46 compounds validated for RET activity.

ZINC ID	Smiles	Selection method	Remaining RET a c t i v - ity with c o m - p o u n d concentration 10^{-7}	Remaining RET activity with compound concentration 10-5	IC ₅₀ (M) n1	IC ₅₀ (M) n2	IC ₅₀ (M) n3
ZINC33008650	COc1cc(NC(=O) Nc2ccc(C)c(F)c2) cc3sc(NC(=O)c4ccccc4) nc13	Con- sensus	107	16	5.16E-06	9.12E-06	7.87E-06
ZINC12324934	CCN1C=C(C(=O) Nc2cccc(O)c2) C(=O)c3cc(F)c(cc13) N4CCN(CC4)C(=O)C	Struc- ture- based	91	25	1.65E-05	4.06E-05	1.81E-05
ZINC9518200	NC1=C(NC(=O) c2ccc(F)c(F)c2)C(=O) NC(=N1)SCC(=O) Nc3ccc4OCOc4c3	Struc- ture- based	109	28	5.85E-05	7.30E-05	1.71E-04
ZINC72312837	COc1ccc(NC(=O) Nc2cccc2c3nc(n[nH]3) c4cocc4)cc1	Con- sensus	103	54	1.99E-05	1.74E-05	3.67E-05

ZINC ID	Smiles	Selection method	Remaining RET ac- tivity with compound concentra- tion 10 ⁻⁷	Remaining RET ac- tivity with compound concentra- tion 10 ⁻⁵
ZINC65184824	COc1cccc(Nc2nc(C)c(s2)C(=O)Nc3ccc4OCOc4c3) c1	Structure- based	129	55
ZINC101094323	Cc1cc(NC(=O)NC2=C(NC3CCCC3) c4ccccc4OC2=O)ccc1Br	Structure- based	97	68
ZINC72119413	Cc1ccc(NC(=O)Nc2ccc(Nc3cc(nc(C)n3)n4ccnc4) cc2)c(C)c1	Consensus	95	68
ZINC33008371	Cc1cc(NC(=O)Nc2ccc(F)cc2)cc3sc(NC(=O) c4cccc4)nc13	Consensus	146	79
ZINC49407610	COC1=CC(=O)C(=NN1c2ccc(OC)c(OC)c2)C(=O) Nc3ccc4OCOc4c3	Structure- based	120	79
ZINC65250057	Cc1cccc(c1)n2ncc3C(=C(C(=O)Nc4ccc(C)cc4F) C(=O)Nc23)O	Structure- based	134	79
ZINC33008195	COc1ccccc1NC(=O)Nc2ccc3nc(NC(=O)c4ccccc4) sc3c2	Consensus	113	84
ZINC33008629	CCOc1ccc(NC(=O)Nc2cc(OC)c3nc(NC(=O) c4ccccc4)sc3c2)cc1	Consensus	115	85
ZINC101332769	Fc1ccc(NC(=O)CN2C(=O)N(CC3CCCO3)C(=O) c4ccc(cc24)C(=O)NCCc5ccccc5)cc1Cl	Structure- based	116	85



ZINC ID	Smiles	Selection method	Remaining RET ac- tivity with compound concentra- tion 10 ⁻⁷	Remaining RET ac- tivity with compound concentra- tion 10 ⁻⁵
ZINC12584529	Cc1cccc(NC(=O)CNC(=O)c2cccc2NS(=O)(=O) c3ccc4NC(=O)Nc4c3)c1C	Structure- based	114	86
ZINC33008381	COc1cc(NC(=O)Nc2cc(C)c3nc(NC(=O)c4ccccc4) sc3c2)cc(OC)c1	Consensus	123	88
ZINC6864637	O=C1Nc2ccc(NC=C3C(=O)NC(=O)N(C3=O) c4cccc4)cc2N1	Structure- based	107	91
ZINC67956408	O=C(NCc1cccnc1n2cnc3ccccc23)c4ccc5NC(=O) Nc5c4	Structure- based	102	91
ZINC83815577	CC(=O)Nc1cccc2CN(CCc12)C(=O)CCNC(=O) c3ccsc3	Structure- based	104	91
ZINC21806194	CCc1cccc(NC(=O)c2cc([nH]n2)c3ccc(OC)c(c3) S(=O)(=O)N)c1	Structure- based	120	93
ZINC9784888	CC(=O)c1cccc(NC(=O)Cc2csc(NC(=O) Nc3cccc(Cl)c3)n2)c1	Structure- based	116	93
ZINC33008624	COc1cc(NC(=O)Nc2cccc2Br)cc3sc(NC(=O) c4cccc4)nc13	Statistical modelling	106	93
ZINC33008616	COc1ccc(OC)c(NC(=O)Nc2cc(OC)c3nc(NC(=O) c4ccccc4)sc3c2)c1	Consensus	114	94
ZINC49002135	Cc1ccc2C(=O)C(=CN(CC(=O)NC3CCCC3)c2n1) C(=O)Nc4ccc(C)c(C)c4	Structure- based	114	96
ZINC35868136	COc1cccc(NC(=O)Nc2c(N)n(nc2c3ccc(Cl)cc3) c4ccc(F)cc4)c1	Statistical modelling	110	96
ZINC33008380	COc1ccc(NC(=O)Nc2cc(C)c3nc(NC(=O)c4ccccc4) sc3c2)c(OC)c1	Consensus	133	99
ZINC1430237	NC1=Nc2nc3ccccc3n2C(N1)c4c[nH]nc4c5cccc(O) c5	Structure- based	117	99
ZINC46009367	CN1C(=O)N(C)c2cc(Oc3ccccc3)c(NC(=O) Nc4cccc(c4)C(=O)C)cc12	Consensus	104	100
ZINC23135952	C1(N2CCN(CC2)C(=O)C2=CC=CO2) N=C2NN=CC2=C(NC2=CC=C(C=C2)C)N=1	Structure- based	130	100
ZINC12155670	COCC(=O)Nc1cc(cc2ncn(CCc3ccccc3)c12)C(=O) NCc4ccc(C)c(F)c4	Structure- based	133	100
ZINC103927245	FC(F)(F)c1cc(NC(=O)c2cocc2)ccc1NC(=O) Nc3ccc(NC(=O)c4cocc4)cc3C(F)(F)F	Statistical modelling	113	101
ZINC33008656	COc1cc(NC(=O)Nc2cc(OC)c3nc(NC(=O)c4ccccc4) sc3c2)cc(OC)c1	Consensus	112	101
ZINC33008626	CCOc1ccccc1NC(=O)Nc2cc(OC)c3nc(NC(=O) c4ccccc4)sc3c2	Consensus	118	101
ZINC2955974	CC(=O)OCc1cnc(C)c2OC(=Nc3ccc(F)cc3) C(=Cc12)C(=O)Nc4ccc5OCOc5c4	Structure- based	121	101
ZINC33008219	COc1ccc(Cl)cc1NC(=O)Nc2ccc3nc(NC(=O) c4cccc4)sc3c2	Consensus	141	103

ZINC ID	Smiles	Selection method	Remaining RET ac- tivity with compound concentra- tion 10 ⁻⁷	Remaining RET ac- tivity with compound concentra- tion 10 ⁻⁵
ZINC33008399	COC(=O)c1cccc(NC(=O)Nc2cc(C)c3nc(NC(=O) c4ccccc4)sc3c2)c1	Consensus	109	103
ZINC103832769	FC(F)(F)c1cc(NC(=O)c2occc2)ccc1NC(=O) Nc3ccc(NC(=O)c4occc4)cc3C(F)(F)F	Statistical modelling	123	103
ZINC32996009	Cc1ccc(cc1)C2=NN3C(=O)C=C(CNC(=O) Nc4ccc(C)c(Cl)c4)N=C3S2	Structure- based	116	105
ZINC33008637	COc1cc(NC(=O)Nc2ccc(Br)cc2)cc3sc(NC(=O) c4cccc4)nc13	Statistical modelling	120	106
ZINC12243694	COCC(=O)Nc1cc(cc2ncn(CCc3ccccc3)c12)C(=O) NCCc4cnn(C)c4	Structure- based	118	109
ZINC1211467	O=C(Cn1cc(C=C2C(=O)NC(=O)NC2=O) c3ccccc13)Nc4ccccc4	Structure- based	131	110
ZINC10279818	Clc1ccc(c(NC(=O)COc2ccc(Cl)cc2C(=O) Nc3ccccc3)c1)n4cncn4	Structure- based	107	111
ZINC72429138	COc1ccc(C2=Cn3cc(nc3C(=O)N2C)C(=O) N4CC(O)C4)c(F)c1	Structure- based	117	111
ZINC11966082	COCC(=O)Nc1cc(C(=O)N(C)CCc2c(C)n[nH]c2C) c3c(c1)ncn3Cc4ccc(F)cc4	Structure- based	118	117
ZINC33008631	COc1cc(NC(=O)Nc2cccc(Br)c2)cc3sc(NC(=O) c4cccc4)nc13	Statistical modelling	153	127
ZINC46009379	CN1C(=O)N(C)c2cc(Oc3ccccc3)c(NC(=O) Nc4ccc(C)c(C)c4)cc12	Consensus	112	128
ZINC62022653	C1(N2CCN(CC2)C(OCC)=O) N=C(NC2C=CC(=CC=2)C)C2=C(NN=N2)N=1	Structure- based	128	154



Table S5.4. List of compounds used to generate structures for S6K, MKNK1, and LKB1, with induced-fit docking.

Target kinase	Compound ID	Compound SMILES
S6K	CHEMBL6291	Cn1cc(C2=C(C(=O)NC2=O)c3cn(CCCSC(=N)N)c4ccccc34)c5ccccc15
S6K	CHEMBL3685220	NC(=O)c1cccc2c(NCc3cccc(c3)N4C(=O)Nc5ccccc5C4=O)ncnc12
S6K	CHEMBL3680430	COCCC(Nc1ncnc2c(cccc12)C(=O)N)c3cccc(NC(=O)c4cc(n[nH]4)C5CC5)c3
S6K	CHEMBL3685220	NC(=O)c1cccc2c(NCc3cccc(c3)N4C(=O)Nc5ccccc5C4=O)ncnc12
S6K	CHEMBL3675398	CNCC(Nc1ncnc2c(cccc12)C(=O)N)c3cccc(NC(=O)c4ccc(F)c(F)c4)c3
MKNK1	CHEMBL3355002	CC(C)Oc1cc2[nH]ncc2cc1Nc3ncnc4[nH]c5CCC(Cc5c34)C(=O)NCCCS(=O) (=O)C
MKNK1	CHEMBL3355000	COc1cc2[nH]ncc2cc1Nc3ncnc4[nH]c5CCC(Cc5c34)C(=O)NCc6ccccc6
MKNK1	CHEMBL3354998	CC(C)Oc1cc2[nH]ncc2cc1Nc3ncnc4[nH]c5CCCCc5c34
MKNK1	CHEMBL223360	Cc1ccc(F)c(NC(=O)Nc2ccc(cc2)c3cccc4[nH]nc(N)c34)c1
MKNK1	CHEMBL3659492	Cc1c(sc2ncnc(Nc3ccc(F)cc3OC4CCN(CC(=O)N5CCOCC5)CC4)c12)C(=O)
MKNK1	CHEMBL3417208	CN1C=C(Nc2ncnc3sc(C(=O)N)c(C)c23)C=CC1=O
MKNK1	CHEMBL1721885	Cc1[nH]c(\C=C\2/C(=O)Nc3ccc(F)cc23)c(C)c1C(=O)NC[C@H](O) CN4CCOCC4
MKNK1	CHEMBL226403	O=C1NCCc2[nH]c(cc12)c3ccnc(c3)c4cnc5cccc5c4
MKNK1	CHEMBL3632739	COC(=O)c1sc2ncnc(Nc3ccc(F)cc3OC(C)C)c2c1C
MKNK1	CHEMBL3805029	FC(F)(F)c1cc(NC(=O)COc2ccc(cc2)c3cccnc3)ccc1CN4CCNCC4
LKB1	CHEMBL502835	COC(=O)c1ccc2\C(=C(\Nc3ccc(cc3)N(C)C(=O)CN4CCN(C)CC4)/ c5cccc5)\C(=O)Nc2c1
LKB1	CHEMBL603469	C[C@]12O[C@H](C[C@]1(O)CO)n3c4ccccc4c5c6C(=O) NCc6c7c8ccccc8n2c7c35
LKB1	CHEMBL1908397	O=C(N1CCNCC1)c2ccc(\C=C\c3n[nH]c4ccccc34)cc2
LKB1	CHEMBL535	$CCN(CC)CCNC(=O)c1c(C)[nH]c(\C=C\2/C(=O)Nc3ccc(F)cc23)c1C$
LKB1	CHEMBL1287853	Cc1cnc(Nc2ccc(OCCN3CCCC3)cc2)nc1Nc4cccc(c4)S(=O)(=O)NC(C)(C)C
LKB1	CHEMBL388978	CN[C@@H]1C[C@H]2O[C@@](C)([C@@H]1OC) n3c4ccccc4c5c6CNC(=O)c6c7c8ccccc8n2c7c35