Supplementary Information

Discovery of Novel SOS1 Inhibitors Using Machine Learning

Lihui Duo,^a Yi Chen,^{c,e} Qiupei Liu,^{a,c} Zhangyi Ma,^a Amin Farjudian,^g Wan Yong Ho,^d Sze Shin Low,^a Jian Ding,^{c,e} Jianfeng Ren,^{a,*} Jonathan D. Hirst,^{b,*} Hua Xie,^{c,e,f*} Bencan Tang^{a,*}

- Nottingham Ningbo China Beacons of Excellence Research and Innovation Institute, Key Laboratory for Carbonaceous Waste Processing and Process Intensification Research of Zhejiang Province, Department of Chemical and Environmental Engineering, The University of Nottingham Ningbo China, 199 Taikang East Road, Ningbo 315100 (P. R. China); shxld1@nottingham.edu.cn (L.D.); biyzm3@nottingham.edu.cn (Z.M); sze-shin.low@nottingham.edu.cn (S.S.L)
- ² School of Chemistry, University of Nottingham, University Park, Nottingham NG7 2RD (UK);
- ³ Division of Antitumor Pharmacology, State Key Laboratory of Drug Research, Shanghai Institute of Materia Medica, Chinese Academy of Sciences, 555 Zuchongzhi Road, 201203 Shanghai, China School of Chemistry, s19-chenyi@simm.ac.cn (Y.C); qiupei.liu@nottingham.edu.cn (Q.L);
- Faculty of Medicine and Health Sciences, University of Nottingham (Malaysia Campus), Semenyih 43500, Malaysia; WanYong.Ho@nottingham.edu.my
- ⁵ University of Chinese Academy of Sciences, No.19A Yuquan Road, Beijing 100049, China; jding@simm.ac.cn
- ⁶ Zhongshan Institute for Drug Discovery, Shanghai Institute of Materia Medica, Chinese Academy of Sciences, Zhongshan Tsuihang New District, Zhongshan, 528400, China;
- School of Mathematics, Watson Building, University of Birmingham, Edgbaston, Birmingham, B15 2TT UK;
- # L.D., Y.C., and Q.L. contributed equally.
- * Correspondence: Jianfeng.Ren@nottingham.edu.cn (J.R), jonathan.hirst@nottingham.ac.uk (JD. H), hxie@simm.ac.cn (H,X), Bencan.Tang@nottingham.edu.cn (B,T).

Supplementary Methods

1 Molecular dataset

The SMILES representations of all SOS1-related molecules from ChEMBL have been documented in distinct Excel files. Active and inactive molecules were distinguished with pChEMBL value 7 as the threshold, and their molecular structure information could be found in the folder named image on GitHub.

2 Algorithm introduction

K-nearest-neighbor Regressor is a versatile non-parametric algorithm extensively employed in data analysis and prediction tasks, and it relies on the concept of feature similarity, operating under the assumption that data points with similar features tend to exhibit similar outcomes [1]. In practice, this algorithm identifies the nearest neighbors of a given data point within the training dataset based on their feature resemblance and employs a local interpolation technique to predict results [1].

Ridge Regressor is a commonly used technique when dealing with multicollinearity, where independent variables exhibit high correlation [2]. By building upon linear regression, this algorithm introduces L2 regularization to add a penalty term to the regression coefficients, thus mitigating multicollinearity, and enhancing model stability [2].

Lasso Regressor, akin to ridge regression, combats multicollinearity by incorporating the L1 regularization to penalize the absolute value of regression coefficients, reduce variability and improve the accuracy of linear regression models [3].

Elastic Net Regressor is a flexible algorithm that blends the best of Ridge and Lasso Regression methods, and it is well suited for handling the challenges of high dimensionality, primarily focusing on feature selection and regression [4]. Elastic Net combines the L1 (Lasso) and L2 (Ridge) regularization, which controls feature selection, prevents overfitting, and enhances the model robustness in the presence of

multicollinearity. By fine-tuning the mixing ratio between these two regularization terms, Elastic Net adapts to various modeling requirements, making it a potent tool for regression tasks in diverse scenarios [4].

Decision-tree Regressor is a tree-structured representation of decision-making processes that may categorize or predict continuous data, which splits the training data from the root node to the decision nodes [5]. Providing data type flexibility and legibility of resulting models, the Decision-tree Regressor could tackle the multi-class classification problems, but it suffers from potential noise and overfitting.

Random Forest Regressor is an ensemble approach to integrating multiple decision trees to tackle the problems of plagued bias and variance in decision trees, in order to improve the prediction performance and the model robustness [6]. By training each decision tree on a distinct data subset and introducing random feature selection for each split, Random Forest Regressor could effectively reduce correlations among individual trees, and leverage the predictions from multiple decision trees to enhance the ensemble's stability and accuracy [6]. Furthermore, its insight into feature importance aids feature selection and model interpretation, making Random Forest Regressor a widely adopted and potent tool, especially valuable for tackling high-dimensional complex data across various applications [6].

Extra-Tree Regressor is a powerful ensemble learning technique that builds a multitude of randomized decision trees through a meta estimator [7]. This approach introduces randomness by selecting random subsets of data and features during the tree-building process, which results in a more robust model with reduced overfitting and a higher prediction accuracy through ensemble-based averaging [7].

Adaboost Regressor first fits a regressor on the original dataset and then fits subsequent copies of the regressor on the same dataset with the instance weights being changed in accordance with the error of the most recent prediction, to concentrate on challenging instances [8].

Gradient Boosting Regressor is a powerful ensemble learning algorithm for regression that combines a collection of weak regression models such as decision trees in a sequential manner to progressively enhance the predictive performance by

minimizing the loss function. This approach is particularly effective at capturing intricate nonlinear relationships while maintaining robustness, although it may require additional tuning and training time compared to alternative algorithms [9].

Support Vector Regressor (SVR) finds a regression plane with the closest possible proximity to a subset of data points called support vectors, while allowing a controlled degree of deviation from these points [10]. SVR is able to capture the underlying patterns and relationships within the data while maintaining a balance between the model complexity and the predictive accuracy, in order to regress a more accurate prediction [10].

3 External validation dataset

The dataset has been documented in distinct Excel files on GitHub (https://github.com/cristinaduo/ML-based-SOS1-VS.git).

4 Similarity calculation

Similarity between molecules have been calculated from the Morgan fingerprints.

Supplementary Tables

Table S1. The optimal hyperparameters of 10 constructed models.

ML models	The optimal hyperparameters	Search space
		'criterion' in [squared_error,
Decision Tree	'criterion': friedman_mse,	friedman_mse];
Decision Tree	'min_samples_split': 7	'min_samples_split' ranges
		from 2 to 9.
		'bootstrap' in [True, False];
Extra Tree	-	'min_samples_split' ranges
		from 2 to 9.
Didaa		'alpha' in [0.001, 0.01, 0.1,
Ridge	-	1, 10];
		'learning_rate' in [0.001,
A 1-D4	'learning_rate': 1, 'loss':	0.01, 0.1, 1];
AdaBoost	expomential	'loss' in [linear, square,
		'loss' in [linear, square, expomential].
Gradient	'min samples split': 6	'min_samples_split' ranges
Boosting	mm_samples_spirt.0	from 2 to 9.
		'C' in [0.001, 0.01, 0.1, 1,
SVR	'C': 10, 'gamma': auto	10, 20, 50, 100];
		'gamma' in [scale, auto].
		'n_neighbors' ranges from 2
	'alaguithus': hall tuga	to 10;
K-Neighbors	'algorithm': ball_tree,	'algorithm' in [auto,
	'n_neighbors': 9, 'p': 1	ball_tree, kd_tree, brute];
		'p' in [1, 2].
T	(-1-1-2-0-01-(-1-4'-2-1-1-4'-2-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-	'alpha' in [0.001, 0.01, 0.1,
Lasso	'alpha': 0.01, 'selection': random	1, 10];

		'selection' in [cyclic, random].
Elastic Net	alpha': 0.01, '11_ratio': 0.7.	'alpha' in [0.001, 0.01, 0.1, 1, 10]; '11_ratio' in [0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8].
Random Forest	'bootstrap': False, 'max_features': sqrt, 'min_samples_split': 9	'bootstrap' in [True, False]; 'max_features' in [auto, log2, sqrt]; 'min_samples_split' ranges from 2 to 9.

Table S2. Algorithms mean performance for model validation using 90% data for model refitting, and 10% reserved data for model verification: a measure of overfitting and external data validation.

Algorithm	Train R ²	Test R ²	Train RMSE	Test RMSE
Decision Tree	0.986(0.0019)	0.833(0.0549)	0.185(0.0133)	0.623(0.0998)
Extra Tree	0.996(0.0011)	0.833(0.0566)	0.099(0.0147)	0.622(0.0942)
Ridge	0.992(0.0014)	0.894(0.0187)	0.142(0.0118)	0.500(0.0462)
AdaBoost	0.940(0.0041)	0.898(0.0166)	0.389(0.0132)	0.491(0.0459)
Gradient	0.000(0.0010)	0.000(0.0242)	0.222(0.0102)	0.494(0.0697)
Boosting	0.980(0.0019)	0.900(0.0242)	0.222(0.0103)	0.484(0.0687)
SVR	0.992(0.0015)	0.902(0.0182)	0.144(0.0131)	0.482(0.0495)
K-Neighbors	0.996(0.0011)	0.906(0.0184)	0.099(0.0147)	0.470(0.0461)
Lasso	0.946(0.0032)	0.910(0.0200)	0.367(0.0102)	0.462(0.0571)
Elastic Net	0.955(0.0024)	0.912(0.0189)	0.335(0.0085)	0.456(0.0535)
Random	0.094(0.0015)	0.016(0.0145)	0.202/0.0099\	0.445(0.0427)
Forest	0.984(0.0015)	0.916(0.0145)	0.203(0.0088)	0.445(0.0437)

Table S3. The information of 10% data for model validation.

Series	Compound ID	Structure	Predicted pChEMBL	Actual pChEMB L
1	CHEMBL4529467		8.40	8.70
2	CHEMBL4469357		8.38	8.70
3	CHEMBL4451252		8.38	7.60
4	CHEMBL4435672		8.37	8.52
5	CHEMBL4572922		8.32	7.46
6	CHEMBL4572076	F NH O	8.27	8.30
7	CHEMBL4539190		8.26	8.52
8	CHEMBL4554249		8.25	8.70
9	CHEMBL4540213		8.24	8.70
10	CHEMBL4441820		8.20	8.40

11	CHEMBL4546387		8.17	8.10
12	CHEMBL4551748	N NH ₂	8.14	8.30
13	CHEMBL4459389		8.14	8.30
14	CHEMBL4513254		7.93	8.52
15	CHEMBL4533487		7.81	7.57
16	CHEMBL4464090		7.77	8.10
17	CHEMBL4448274		7.70	7.47
18	CHEMBL4524954	OH HN	7.68	7.85
19	CHEMBL4463184		7.64	7.44
20	CHEMBL4570224		7.63	7.52
21	CHEMBL4443395	NH N	7.63	7.54
22	CHEMBL4515122		7.58	7.46
23	CHEMBL4453639		7.55	7.40

24 CHEMBL4583745	NH N	7.52	7.37
25 CHEMBL4473214		7.52	7.44
26 CHEMBL4561965	NH CH	7.51	7.66
27 CHEMBL4455684	No.	7.46	7.55

Table S4. Inhibition rate for selected carboxylic acid compounds at various concentrations in activity confirmation assays. Quantified data represents the mean \pm SD from two independent biological replicates.

Inhibition rate (%)						
Concentration (µg/mL)	50	40	25	10	0.5	
CL01545355	61.6 ± 4.0	53.3 ± 0.5	38.0 ± 3.3	17.9 ± 3.3	-1.6 ± 0.2	
CL01545365	72.3 ± 2.7	66.6 ± 0.1	53.8 ± 1.6	31.9 ± 2.8	0.8 ± 1.1	
CL01545444	49.9 ± 0.5	45.5 ± 0.0	31.1 ± 1.4	14.5 ± 3.3	1.5 ± 1.1	
CL01545464	60.5 ± 4.0	55.3 ± 3.2	42.9 ± 3.4	24.2 ± 1.4	3.4 ± 1.8	

Table S5. Drug-likeness prediction of the molecule (CL01545365).

Properties		Value	Optimal Range	Proper	rties Value	Optimal Range
Physicochemi	Molecular Weight	389.5	100~600	Metabolism CYP1	A2	
cal				inhibit	or	
Property	nHA	7	0~12	CYP1A	A2 +	
				substra	ite	
	nHD	2	0~7	CYP20	C19	
				inhibit	or	
	TPSA	99.600	0~140	CYP20	C19	
				substra	ite	
	logS	-3.675	-4~0.5	CYP20	C9 inhibitor++	
	logP	4.128	0~3	CYP20	C9	
				substra	ıte	
	logD	1.210	1~3	CYP2I	D6 ++	
				inhibit	or	
Absorption	Caco-2	-5.673	>-5.15	CYP2I	D6	
	Permeability			substra	ite	

	MDCK	1.1	2-20 × 10-6		CYP3A4	<u>-</u>	
	Permeability	10-5			inhibitor		
	Pgp-inhibitor				CYP3A4	++	
					substrate		
	Pgp-substrate			Excretion	CL	0.857	5~15
	HIA				T1/2	0.103	3
	F20%			Toxicity	LD50	1228.80	>500
						2	
	F30%				hERG Blockers		
Distribution	PPB	96.990%	<90%		H-HT	+	
	VD	0.245	0.04~20		DILI	+++	
	BBB Penetration				AMES		
	Fu	1.630%			SkinSen		

Molecular weight (MW) contains hydrogen atoms.

nHA:Number of hydrogen bonds acceptors.

nHD:Number of hydrogen bonds donors.

TPSA:Topological Polar Surface Area.

logS:log of the aqueous solubility.

logP:log of the octanol/water partition coefficient.

logD:logP at physiological PH 7.4.

Caco-2 Permeability:apparent Caco-2 cell permeability in log unit.

MDCK Permeability:apparent MDCK cell permeability in cm/s.

Pgp-inhibitor:possibility of being Pgp-inhibitor.

Pgp-substrate:possibility of being Pgp-substrate.

HIA:Human Intestinal Absorption.

F20%:20% bioavailability.

F30%:30% bioavailability.

PPB:Plasma Protein Binding.

VD: Volume Distribution.

BBB Penetration:Blood-Brain Barrier Penetration.

Fu:the fraction unbound in plasms.

CYP1A2 inhibitor:possibility of being inhibitor.

CYP1A2 substrate:possibility of being substrate.

CYP2C19 inhibitor:possibility of being inhibitor.

CYP2C19 substrate:possibility of being substrate.

CYP2C9 inhibitor: possibility of being inhibitor.

CYP2C9 substrate: possibility of being substrate.

CYP2D6 inhibitor: the possibility of being inhibitor.

CYP2D6 substrate: the possibility of being substrate.

CYP3A4 inhibitor: the possibility of being inhibitor.

CYP3A4 substrate: the possibility of being substrate.

CL: Clearance.

T1/2: half-life.

LD50: the dose amount of a tested molecule to kill 50% of the treated animals within a given period (mg/kg).

hERG Blockers: the probability of being active.

H-HT: Human Hepatotoxicity.

DILI: Drug-Induced Liver Injury.

AMES: Ames Mutagenicity; SkinSen: Skin sensitization.

Supplementary Figures

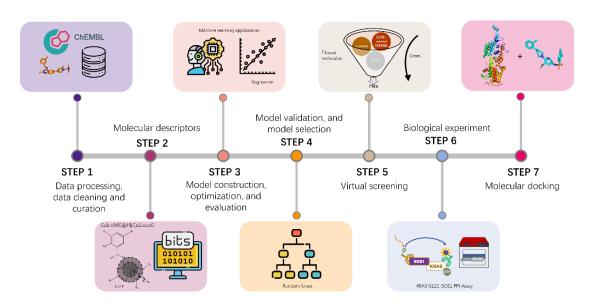


Figure. S1. Overview of the proposed pipeline for virtual screening small molecules as SOS1 inhibitors using machine-learning techniques with molecular docking. The framework consists of seven steps: 1) Raw data collection from the ChEMBL, data processing including curation, cleaning, and deduplicate. 2) Molecular representation generation: converting data into molecular fingerprints. 3) Model construction, optimization, and evaluation: implementing 10 different machine learning techniques, evaluating and adjusting the model parameters. 4) Model validation: comparison of model performance capabilities and best predictive model selection. 5) Virtual screening: molecules are searched from in-house libraries based on ML-based LBVS and hits are identified and ranked. 6) Biological experiment: KRAS G12C/SOS1 PPI Assay. 7) Molecular docking: hits and receptor interactions study.

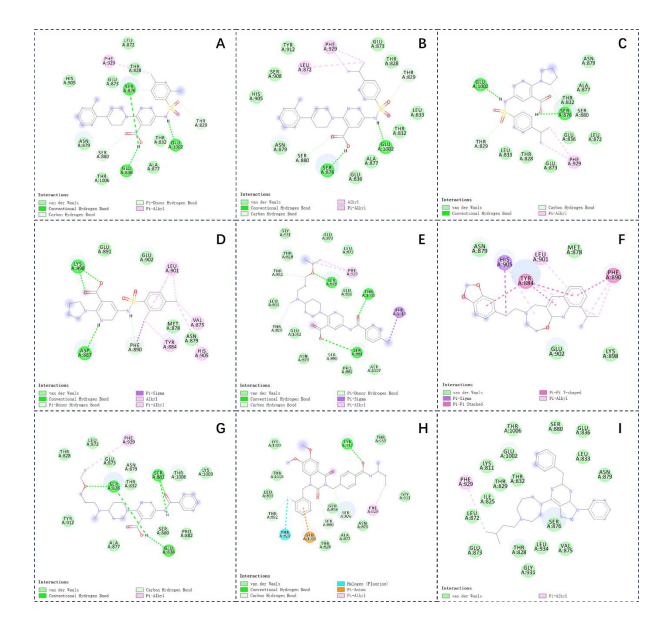


Figure. S2. 2D interaction mode of nine hit compounds with SOS1 protein (PDB:6CSM). (A) CL01545444; (B) CL01545464; (C) CL01545365; (D) CL01545355; (E) CL00838284; (F) CL01132463; (G) CL00838287; (H) CL00817024; (I) CL01027021. The receptor-ligand interaction was visualized using the BIOVIA Discovery Studio Visualizer (Version 2023, San Diego, Systèmes).

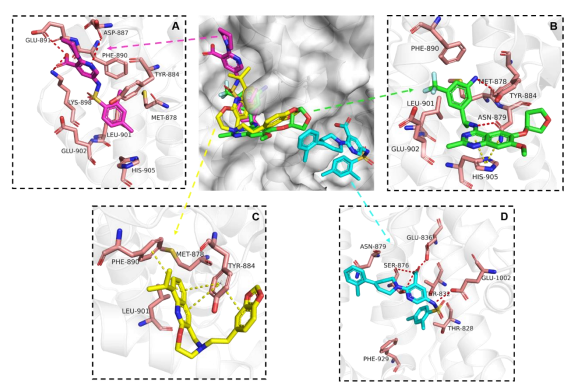


Figure. S3. Interaction mode of hit compounds with SOS1 protein (PDB:6SCM). (A) **CL01545355**; (B) **BI-3406** (C) **CL01132463**; (D) **CL01545444**; The red dashed line represents the hydrogen bond interaction, and the yellow dashed line represents *pi* stacking; The protein-ligand interactions were analyzed by PLIP (Protein-Ligand Interaction Profiler) [11].

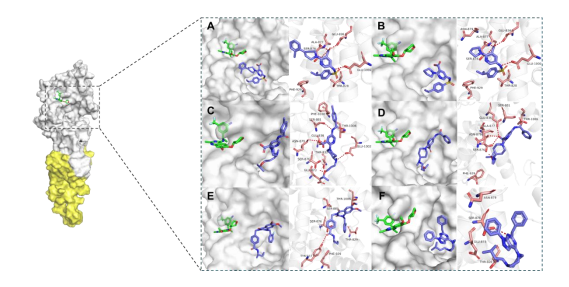


Figure. S4. Interaction mode of hit compounds with SOS1 protein (PDB:6CSM). (A) CL01545464; (B) CL01545365; (C) CL00838284; (D) CL00838287; (E) CL00817024; (F) CL01027021.

References

- [1] L. Devroye, L. Gyorfi, A. Krzyzak, G. Lugosi, On the strong universal consistency of nearest neighbor regression function estimates, The Annals of Statistics. 22 (1994) 1371–1385.
- [2] G.C. McDonald, Ridge regression, Wiley Interdisciplinary Reviews: Computational Statistics. 1 (2009) 93–100.
- [3] R. Tibshirani, Regression shrinkage and selection via the lasso, Journal of the Royal Statistical Society Series B: Statistical Methodology. 58 (1996) 267–288.
- [4] Z. Zhang, Z. Lai, Y. Xu, L. Shao, J. Wu, G.-S. Xie, Discriminative elastic-net regularized linear regression, IEEE Transactions on Image Processing. 26 (2017) 1466–1481.
- [5] J.R. Quinlan, Induction of Decision Trees, Machine Learning. 1 (1986) 81–106.
- [6] L. Breiman, Random forests, Machine Learning. 45 (2001) 5–32.
- [7] P. Geurts, D. Ernst, L. Wehenkel, Extremely randomized trees, Machine Learning. 63 (2006) 3–42.
- [8] D.P. Solomatine, D.L. Shrestha, AdaBoost. RT: a boosting algorithm for regression problems, in: 2004 IEEE International Joint Conference on Neural Networks (IEEE Cat. No. 04CH37541), IEEE, 2004: pp. 1163–1168.
- [9] J.H. Friedman, Greedy function approximation: a gradient boosting machine, Annals of Statistics. (2001) 1189–1232.
- [10] A. Singh, N. Thakur, A. Sharma, A review of supervised machine learning algorithms, in: 2016 3rd International Conference on Computing for Sustainable Global Development (INDIACom), Ieee, 2016: pp. 1310–1315.
- [11] M.F. Adasme, K.L. Linnemann, S.N. Bolz, F. Kaiser, S. Salentin, V.J. Haupt, M. Schroeder, PLIP 2021: Expanding the scope of the protein–ligand interaction profiler to DNA and RNA, Nucleic Acids Research. 49 (2021) W530–W534.