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Insight into TLR4 receptor inhibitory activity via QSAR for the treatment of *Mycoplasma pneumonia* disease†

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Mycoplasma pneumoniae (MP) is one of the most common pathogenic organisms causing upper and lower respiratory tract infections, lung injury, and even death in young children. Toll-like receptors (TLRs) play an important role in innate immunity by allowing the host to recognize pathogens invading the body. Previous studies demonstrated that TLR4 is a potential therapeutic target for the treatment of MP pneumonia. Therefore, the present study aimed to screen biologically active ingredients that target the TLR4 receptor pathway. We first used molecular docking to screen out the active compounds inhibiting the TLR4 pathway, and then used regression and classification machine learning algorithms to establish a quantitative structure–activity relationship (QSAR) model to predict the biological activity of the screened compounds. A total of 78 molecules were used in QSAR modelling, which were retrieved from the ChEMBL database. The QSAR models had acceptable correlation coefficients of R^2 on the training and testing dataset in the range of 0.96 to 0.91 and 0.93 to 0.76, respectively. The multiclass classification models showed accuracy on training and testing data within ranges of 1.0 to 0.70, 0.96 to 0.63, and log loss ranges from 0.27 to 8.63, respectively. In addition, molecular descriptors and fingerprints have been studied as structural elements involved in increased and decreased inhibitory activities. These results provide a quantitative analysis of QSAR and classification models applicable for high-throughput screening, as well as insights into the mechanisms of inhibition of TLR4 antagonists.

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1. Introduction

Mycoplasmas are highly pleomorphic microorganisms, contain a very small genome ranging from 0.58–2.20 Mb and cannot flourish without cholesterol.¹ As a human pathogen, *Mycoplasma pneumoniae* (MP) lack a cell wall and has a specialized tip organelle responsible for its cytoadherence.² MP causes asthma, bronchitis, pneumonia, and pharyngitis in humans. Young children and young adults are most likely to be affected by the MP infection.³ There were outbreaks of MP infections worldwide from 2010 to 2012 in different regions of the World.^{4,5} The pathophysiological changes that take place because of MP infection are attributed to several factors, including membrane

lipoproteins, polysaccharides, and viral nucleases.⁶ Previous studies demonstrated that pattern recognition receptors (PRRs) on immune cells detect pathogen-associated molecular patterns (PAMPs) on the surface of pathogens when they invade the host.⁷ The toll-like receptor family (TLRs) is a family of type I trans-membrane transport receptors that are primarily expressed on the surface of epithelial cells and immune cells. They play an important role in the early recognition and inflammatory response of the host against pathogens.⁸ Among the TLRs, TLR4 is the first in the TLR family. It can recognize pathogenic microorganisms, triggering the production of cytokines by cells, chemokines, adhesion factors, and acute phase proteins that regulate inflammatory responses,⁹ which plays an important role in the occurrence of inflammation, especially the activation of the immune system.¹⁰ A previous study had demonstrated that exposure of mouse macrophages to MP lipids activated the TLR4 signaling pathway, resulting in the expression of tumor necrosis factor (TNF)- α and interleukin-1 β mediated by autophagy and nuclear factor kappa B (NF- κ B).¹¹ Accordingly, these findings suggest that TLR4 is not only involved in the body's ability to resist pathogen invasion, but that it may also play an important role in the development and occurrence of inflammation-mediated injuries. Thus, the inhibition of TLR4 in MP infections needs to be further explored. Meanwhile, anti-inflammatory

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compounds that target the TLR4 signaling pathway have received much attention, and it has become a research hotspot in this field, which can help in discovering new therapeutic targets.

Recently, more and more active compounds in natural products have been found to inhibit the TLR4 pathway. Studies revealed that natural products are rich in molecules that possess the potential to inhibit the TLR4 protein and have attracted the attention of researchers.^{12–15} Furthermore, TLR4 inhibition mediated by small molecules led to an array of research focusing on the molecular mechanism of action of TLR4 inhibitors.¹⁶ However, future studies are needed to confirm these findings. Besides, the emergence of antibiotic resistance represents another challenge in the context of the treatment of MP infections.¹⁷ It is therefore crucial to find alternatives to antibiotics to prevent the emergence of resistance against anti-mycoplasma drugs. Hence, active ingredients of natural products could be used to modulate the host immune inflammatory response. The active ingredients of natural products require more experimental data on its toxicology and pharmacology before its use in clinical trials, which is a lengthy process. To screen TLR4 inhibitors in a limited period of time, it is necessary to use fast, accurate, and reliable screening methods based on detailed study of TLR4 inhibition and regulation. In this context, modern machine learning technologies make better use of information obtained from several sources to predict the bioactivities of drugs for several diseases, thus facilitating the discovery of new drugs more efficiently. In recent years, computer-aided drug screening is advancing towards practicality and is emerging as a core technology for innovative drug research. Several drug libraries

were screened in a very short time-period, leading to the discovery of many active compounds in traditional Chinese medicines and the successful repurposing of several approved drugs.^{18,19} Based on previous research, virtual screening paved the way for the future development of improved chemical analogs for use in treating a wide variety of human and animal diseases through medicinal chemistry structure–activity relationships and drug screenings.²⁰ Molecular docking and quantitative structure activity relationship (QSAR) models provide structural information and insight into TLR4 inhibitors that can be used to guide more effective drug development, including screening and rational drug discovery of TLR4 inhibitors. Therefore, the objective of the present study was to identify lead compounds that can inhibit the TLR4 protein for the treatment of MP pneumonia. The regression and classification QSAR models were developed from a set of known chemical TLR4 inhibitors. These QSAR models will be used to predict and classify the bioactive compounds based on their predicted bioactivity (pIC_{50}) values and provide theoretical foundations to enable the development of potent drugs from natural products for the prevention and treatment of MP-pneumonia. The flow chart for the experimental process is shown in Fig. 1.

2. Materials and methods

2.1. Molecular docking

The set of 50.2 K Discovery Diversity Set (DDS) compounds were used for docking to the crystal structure of human TLR4 (PDB ID: 4G8A), which is taken from the RCSB PDB database. The

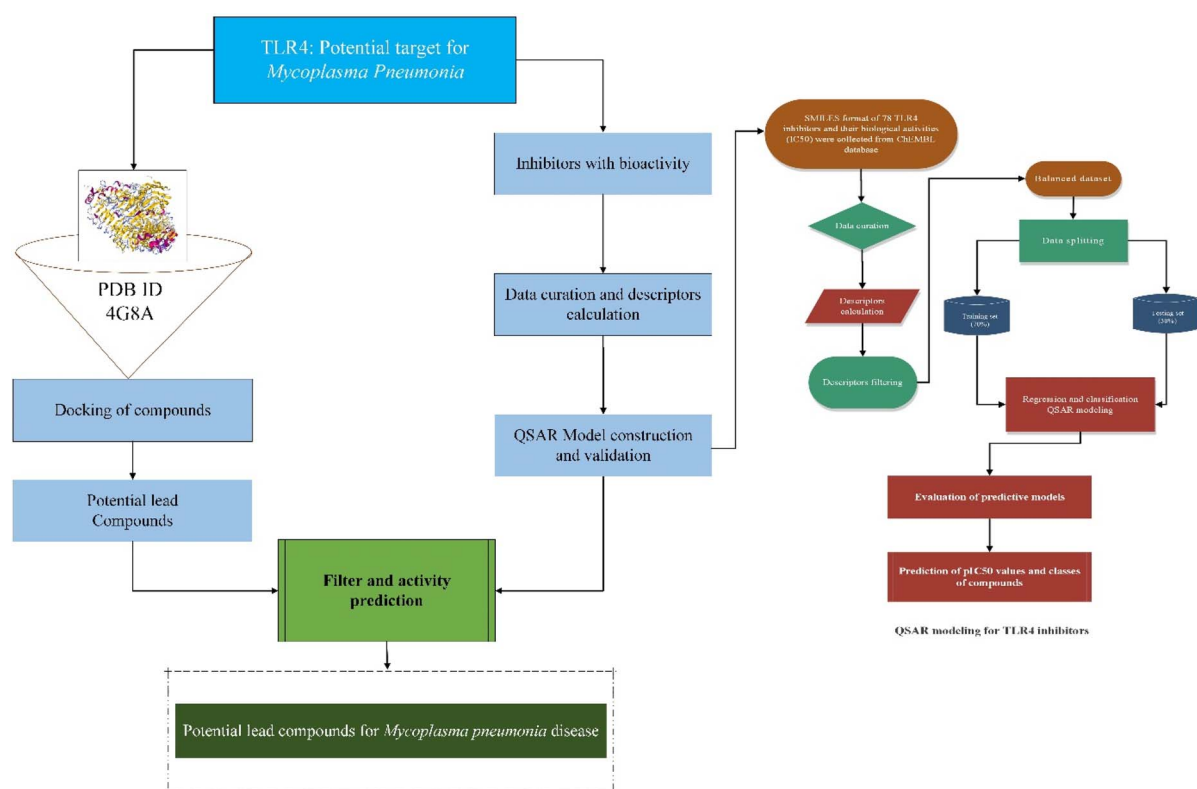


Fig. 1 Schematic diagram represents the workflow of experimental process.



structure of each ligand as well as the structure of the protein were both preprocessed prior to docking. The docking process was visualized with PyMol (version 11.4) using Schrödinger Maestro (version 11.4). Schrödinger's Protein Preparation Wizard tool was used to prepare the protein structure.²¹ The water molecules have been deleted, and the hydrogens, residues and side chains have been added. Using the force field (OPLS2005, RMSD is 0.30 Å), the energy optimization was carried out in order to minimize the protein structure.^{22,23} Maestro (version 11.4) was used to generate the receptor grid, selecting amino acid residues with hydrogen and ionic bonding Glu42/Asp60/Arg87/Glu135/Ser183/Arg234/Arg264/Asn265/Glu266/Arg289 based on the complex formed by the TLR4 with the extracellular adaptor MD-2 (a glycoprotein) and the hydrophobic portion of lipid polysaccharide (LPS), and the grid size was set to 20 Å × 20 Å × 20 Å so that new compounds could be attached to the same binding site and co-crystallized ligands could be excluded.²⁴ In Schrödinger Maestro, the LigPrep module was used to prepare the ligand structure. After the ligands were loaded and optimized, and the 3D structure was obtained.²⁵ For docking, the prepared molecules were imported into Glide. In the generated grid of the target protein, the ligand molecules are docked through geometric and energy matching. In the Glide module, standard precision (SP) was utilized with the high-throughput screening (HTVS) setting, and the top 10% of compounds were selected. Thereafter, extra precision (XP) screenings were performed in the Glide module and the top 10% of small molecules were selected for further screening.^{26–28} Best docking pose of each ligand was kept for the further analysis.²⁹

2.2. Dataset

From the ChEMBL database (version 29),³⁰ we compiled a list of 78 TLR4 inhibitors along with their IC₅₀ values. ChEMBL is a searchable database of more than 2 million compounds, molecule bioactivities, and information on bioactive molecules exhibiting drug-like properties. In ESI S1† (Excel sheet), the list of TLR4 inhibitory molecules used in the QSAR model, simplified molecular input line entry system (SMILES) notation, pIC₅₀ and ChEMBL ID as well as Lipinski's descriptors are presented. For a more uniform distribution of the data, the IC₅₀ values were converted to the pIC₅₀ by taking the negative logarithm of the IC₅₀ values by using the equation $\text{pIC}_{50} = -\log_{10}(\text{IC}_{50})$. Here, pIC₅₀ is the negative logarithm of IC₅₀. As this study is primarily concerned with developing regression and classification models of biological activity. Thus, the IC₅₀ values were divided into three classes (high (<1 μM), moderate (≤3 μM and >1 μM) and low (>3 μM)) for a clear distinction between the potency of these compounds. The geometry is optimized using MOPAC (Molecular Orbital Package) using the AM1 method, the 3D coordinates are preserved, and the energy is minimized in Merck Molecular Force Field (MMFF94) for all ligands before descriptors and molecular fingerprints are calculated. Molecular descriptors were calculated, and the data was curated based on the calculated descriptors. Model building relies on qualitative and/or quantitative chemical information that can be obtained from molecular fingerprints. The PaDel molecular fingerprints and descriptors were calculated

using ChemDes web-based platform.³¹ To clean the initial data collected from the ChEMBL database, data preprocessing has been performed as described previously.^{32–35} As part of the data curation process, some key steps are performed including (i) structural cleaning and conversion, (ii) removal of duplicates, (iii) removal of mixtures and inorganics, (iv) normalization of specific chemotypes, and (v) manual verification of the data. All other inhibitory targets except for TLR4, as well as the pIC₅₀ values for each, were removed from the data. Molecules with missing values for the pIC₅₀ and SMILES notation, as well as duplicate value entries, were removed. The feature selection process is also known as variable selection or attribute selection. Predictive modeling involves automatic selection of attributes most relevant to the problem. As a first step, all the descriptors and molecular fingerprints were checked manually for any missing values, and all the columns containing zero values were removed from the files using a Python script. The fingerprints were then combined into a single csv (Comma Separated Values) file. A variation selection method was initially applied in order to remove the redundant features from the list of combined features.

2.3. Constructing regression and classification models

In QSAR modeling, Random Forest (RF) regression and classification algorithms are frequently used since they provide good predictive performance and high model interpretability. The RF algorithm is an ensemble of regression and classification trees.³⁶ It is a robust algorithm with higher performance in the presence of very high-dimensional parameters, spaces, and outliers than other machine learning algorithms.³⁷ In addition, random sampling reduces overfitting, and the RF algorithm recognizes important features that affect the QSAR model, allowing for further evaluation of the model and improved prediction accuracy. The dataset is divided into two main subsets, the training subset (70%) and the test subset (30%). The training subset of the data is used to train the model, whereas the test subset is used to cross-validate the model. First, the model was evaluated using 5-fold cross-validation, providing a more general model;³⁸ then, the model's hyperparameters were tuned using the Optune framework (Optune-A hyperparameter optimization framework).³⁹ For the training, testing, and cross-validation sets of compounds, the performance of regression models was assessed using root mean squared error (RMSE), mean absolute error (MAE), and squared correlation coefficient (R^2). The predictability of the model was also assessed by calculating the residual error between the predicted and experimental pIC₅₀ values. The performance of the classification model was evaluated in terms of classification accuracy, RMSE, MAE, and Log loss over the training, testing, and validation sets of compounds. Additionally, the RF model was compared with Decision Tree Regression (DTR), AdaBoost Regression (ABR), Gradient Boosting Regression (GBR) and Extra Trees Regression (ETR) models trained on the same data. RF classification model was also compared with KNeighbors classifier, Support Vector Classifier (SVC), Decision Tree Classifier, AdaBoost Classifier, Gradient Boosting Classifier, Linear Discriminant Analysis and Quadratic Discriminant Analysis, on both training and testing data. The R^2 , accuracy, RMSE, MAE, Log



Loss, and the performance of these models are analyzed and compared. For these compounds, a set of descriptors and fingerprints was selected in the same manner as for the fingerprints that were used to build the QSAR model. Based on these features, pIC_{50} values were predicted for DDS set of compounds.

2.4. Statistical analysis for assessing predictive models

For the training, testing, and cross-validation sets of compounds, the performance of regression models was evaluated using the root mean squared error (RMSE), mean absolute error (MAE), and squared correlation coefficient (R^2). The predictive performance of the model was further examined by calculating the residual error between the predicted and experimental pIC_{50} values. While the performance of the classification model for the training, testing, and validation set of compounds was evaluated in terms of classification accuracy, RMSE, MAE, and log loss.

2.5. Prediction of small molecule compounds

Based on 50.2 K DDSs of compounds screened by docking, the top 30 compounds were selected. ChemDes online website was used to compute chemical descriptors and molecular fingerprints for all these compounds using canonical smiles format. A set of fingerprints and descriptors were selected for these compounds, like those on which the QSAR model was based. Using these fingerprints, the pIC_{50} values and potencies of DDS compounds were predicted.

3. Results and discussion

3.1. Molecular docking

The docking technique is a computational method for predicting the orientation of a ligand or protein when bound to an enzyme or receptor protein.⁴⁰ In the field of drug discovery and design, docking is extremely useful as it provides

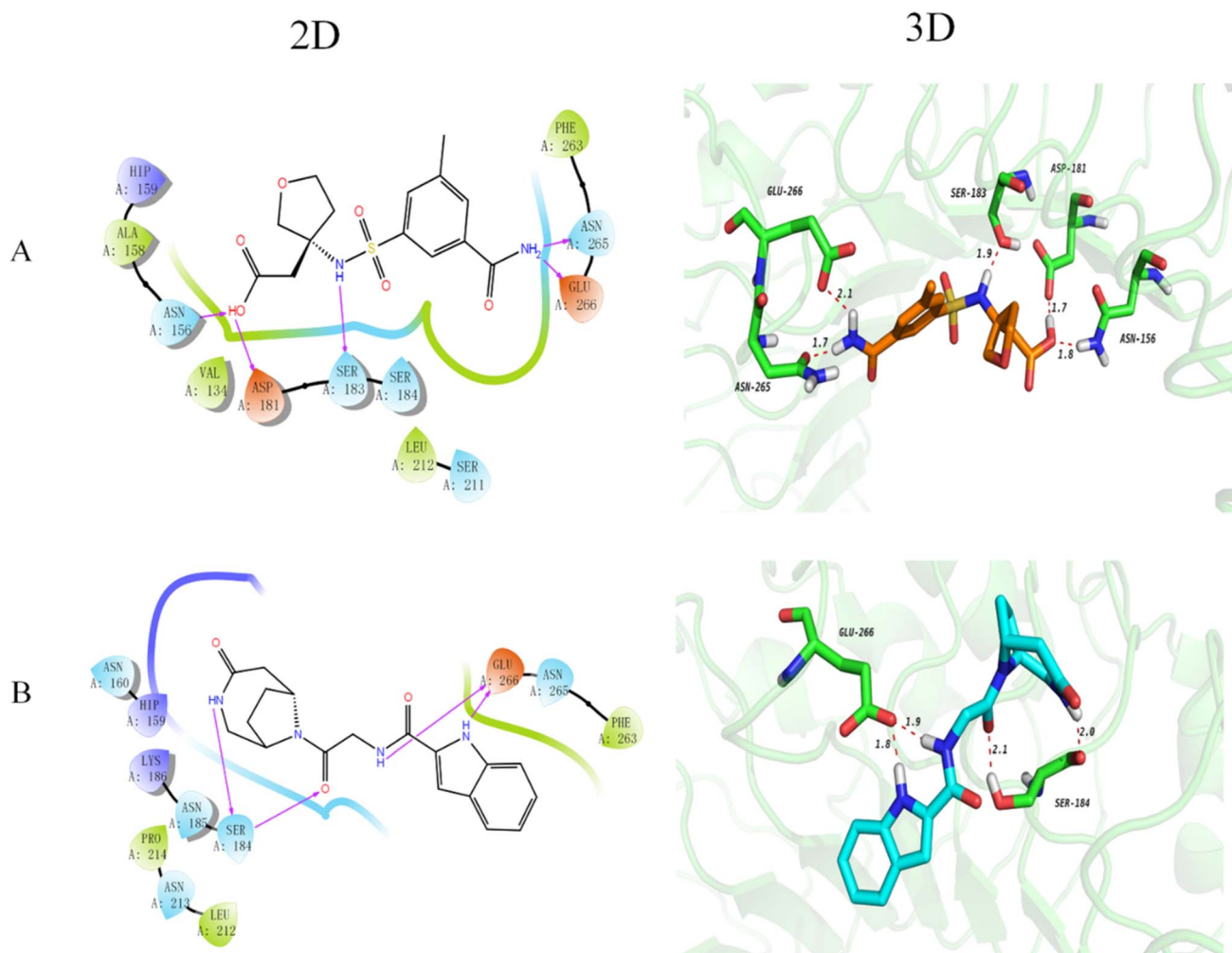


Fig. 2 2D and 3D representation of the docked compounds. Panel (A) and (B) represents top 2 top 2 compounds (*(R)*-2-(3-(3-carbamoyl-5-methylphenyl)sulfonamido) tetrahydrofuran-3-yl)acetic acid and *N*-(2-oxo-2-((6*R*)-4-oxo-3,9-diazabicyclo[4.2.1]nonan-9-yl)ethyl)-1*H*-indole-2-carboxamide, respectively. The hydrogen bonds and their lengths (Å) are displayed in these cartoons.





Table 1 The results of docking and the partial prediction of pIC₅₀ values of the Top 30 DDS compounds

S. No.	Drug names	MW	Docking score	RFR model	ETR model	DTR model	ABR model	GBR model
1	(R)-2-(3-(3-Carbamoyl-5-methylphenyl)sulfonamido) tetrahydrofuran-3-yl)acetic acid	342.37	−5.228	5.430829277	6.314821411	6.958607315	4.813787229	5.136554754
2	N-(2-Oxo-2-((6R)-4-oxo-3,9-diazabicyclo[4.2.1]nonan-9-yl)ethyl)-1H-indole-2-carboxamide	340.38	−5.127	4.815372966	6.16129079	4.978810701	4.644380954	4.864479656
3	4-(N-(2-Carbamoylphenyl)sulfamoyl)-3-fluorobenzoic acid	338.31	−5.014	5.131434015	5.942968037	4.779891912	4.644380954	4.848622926
4	(R)-3-(5-Oxo-2,5-dihydro-1H-1,2,4-triazol-3-yl)-N-((3-(trifluoromethyl)-1H-1,2,4-triazol-5-yl)methyl)piperidine-1-carboxamide	360.30	−4.888	5.335803885	4.311301872	5	5.128070842	5.154233598
5	1-(((1S,2R)-2-Hydroxy-1,2,3,4-tetrahydronaphthalen-1-yl)carbamoyl)cyclopent-3-enecarboxylic acid	301.34	−4.857	5.162769848	5.498993506	5.229147988	5.853871964	5.136223419
6	3-(3-Amino-5-methylisoxazole-4-sulfonamido)-4-methoxybenzoic acid	327.31	−4.841	5.42557839	5.515008442	7.096910013	6.198970004	4.768100468
7	5-(2-(2-Aminoethyl)amino)thiazol-4-yl)-2-hydroxybenzamide	278.33	−4.788	5.062946566	4.451147788	6.795880017	4.676057904	4.715750754
8	(R)-3-((4-Amino-6,7-dimethoxyquinazolin-2-yl)amino)-2-hydroxy-2-methylpropanoic acid	322.32	−4.765	5.384378737	4.723592771	5	5.551578992	5.318842099
9	4-((3-(1-(3-Methylbutanoyl)piperidin-4-yl)ureido)methyl)benzoic acid	361.44	−4.729	5.352768581	4.604814799	6.958607315	5.128070842	4.930009052
10	(S)-3-(5-Fluoropyridine-3-sulfonamido)-2-hydroxypropanoic acid	264.23	−4.626	5.469392305	5.131277132	6.958607315	5.857242359	5.598004671
11	3,4-Difluoro-N-(2-((2R,4R)-4-hydroxy-2-(hydroxymethyl)pyrrolidin-1-yl)-2-oxoethyl)benzamide	314.29	−4.559	5.303713126	4.866537918	6.958607315	5.121293699	4.907629008
12	(S)-3-(2,3-Dichlorophenylsulfonamido)-2-hydroxypropanoic acid	314.14	−4.530	5.448314838	4.792741186	6.958607315	5.121293699	4.849140284
13	2-(((3R,4R)-3-Methyltetrahydro-2H-pyran-4-yl)amino)-5-sulfamoylbenzoic acid	314.36	−4.524	5.737739993	4.597185429	6.795880017	4.644380954	5.211471563
14	(R)-2-(3-Methyl-1H-1,2,4-triazol-5-yl)-N-(4-oxochroman-3-yl)acetamide	286.29	−4.508	5.198586102	4.835972975	6.795880017	5.121293699	4.841100257
15	4-(N-(2-Amino-2-oxoethyl)-N-benzylsulfamoyl)-1H-pyrrole-2-carboxylic acid	337.35	−4.507	4.93587296	5.078963707	4.256568635	4.644380954	5.278315448
16	(R)-4-Isopropoxy-3-(1-(tetrahydrofuran-3-yl)-1H-pyrazole-4-sulfonamido)benzoic acid	395.43	−4.504	5.659485927	5.673346573	5	4.676057904	5.283553443
17	3-(N-(5-Cyano-2-(methylamino)phenyl)sulfamoyl)-5-fluorobenzoic acid	349.34	−4.489	5.2393042	6.270021811	4.586700236	4.676057904	5.202654077
18	1-((2-(5-Fluoro-1H-indol-3-yl)ethyl)carbamoyl)azetidine-3-carboxylic acid	305.30	−4.463	5.138107357	5.712863539	6.958607315	5.121293699	4.859899139
19	4-(3-Ethylphenylsulfonamido)-3-hydroxybenzoic acid	321.35	−4.456	5.326559803	5.193816275	5	5.266000713	5.426620204
20	(R)-3-(3-(5,6-Dimethyl-4-oxo-1,4-dihydrothieno[2,3-d]pyrimidin-2-yl)propanamido)-2-hydroxy-2-methylpropanoic acid	353.39	−4.450	5.372020827	5.783184637	5	5.121293699	5.442711298
21	(2R,3S,4R)-1-(<i>tert</i> -butoxycarbonyl)-3,4-dihydroxypyrrolidine-2-carboxylic acid	247.25	−4.443	5.469071944	5.613893366	5	5.121293699	4.953810874
22	2,4-Difluoro-N-(2-((2R,4R)-4-hydroxy-2-(hydroxymethyl)pyrrolidin-1-yl)-2-oxoethyl)benzamide	314.29	−4.411	5.278958496	4.61805273	6.958607315	4.813787229	4.931341817
23	(3R,5R)-1-(6-(((3-Cyclopropyl-1H-pyrazol-5-yl)methyl)amino)pyrimidin-4-yl)-5-((dimethylamino)methyl)pyrrolidin-3-ol	357.45	−4.399	5.284046701	4.564151855	6.602059991	5.121293699	5.125098228

Table 1 (Contd.)

S. No.	Drug names	MW	Docking score	RFR model	ETR model	DTR model	ABR model	GBR model
24	(R)-2-(1-Oxo-1,2-dihydroisoquinoline-3-carboxamido)-3-phenylpropanoic acid	336.34	-4.386	5.163010804	5.302824295	5.853871964	5.595633098	5.484515313
25	(R)-4-(N-(2-Oxo-2-((tetrahydro-2H-pyran-3-yl)amino)ethyl)sulfamoyl)benzoic acid	342.37	-4.380	5.820985738	5.221450444	6.958607315	5.035830554	5.35223672
26	(R)-2-(6,7-Dihydro-5H-pyrrolo[1,2-d]imidazole-3-sulfonamido)-2-(3-methoxyphenyl)acetic acid	351.38	-4.365	5.275202795	5.291271987	5	4.676057904	5.012575341
27	(R)-N-(1-Amino-3-methoxy-1-oxopropan-2-yl)-7-methyl-1H-indole-2-carboxamide	275.30	-4.342	5.130831094	5.946522112	6.795880017	5.121293699	4.992894312
28	N-((2R,3R)-4-Hydroxy-3-(methylthio)butan-2-yl)-2-oxo-2,3-dihydrobenzo[d]oxazole-6-sulfonamide	332.40	-4.339	5.36250487	5.963766686	6.958607315	4.813787229	4.866545583
29	1-(2-Morpholino-2-oxoethyl)-3-(pyridin-3-yl)urea	264.28	-4.278	5.323176653	4.938421827	6.795880017	5.121293699	4.845185655
30	(3R,4R)-1-(3-(Carbamoylphenethyl)carbamoyl)-4-methylpiperidine-3-carboxylic acid	333.38	-4.260	5.316126193	4.52083702	6.476253533	5.121293699	4.632121561

understanding of the best binding affinity of the potent ligand.⁴¹ Here, docking of the DDS compounds to TLR4 receptor protein provides insights and highlights important protein-ligand interactions. The 2D and 3D docking poses of the representative top 2 compounds are represented in Fig. 2. It has been noted that longer bond results in weaker hydrogen bonding and *vice versa*. Compound 1 (Fig. 2(A)) formed five hydrogen bonds with TLR4 protein, showing that the carboxyl group forms two hydrogen bonds with ASP181 and ASN156 as hydrogen bond donor and acceptor, respectively, with distances of 1.7 Å and 1.8 Å, respectively. Further, the NH of the sulfonamide forms a hydrogen bond with SER183 at a distance of 1.9 Å and the NH₂ group forms two hydrogen bonds with ASN265 and GLU266 with distances of 1.7 Å and 2.1 Å, respectively. Compound 2 (Fig. 2(D)) forms 4 hydrogen bonds with the TLR4 protein that is the NH on the bridged ring acts as a hydrogen bond donor to form a hydrogen bond with SER184 with a distance of 2.0 Å and the carbonyl group acts as a hydrogen bond acceptor to form a hydrogen bond with SER184 with a distance of 2.1 Å. Besides, the NH form a hydrogen bond with GLU266 with a distance of 1.9 Å and the NH of indole form a hydrogen bond with GLU266 with a distance of 1.8 Å. Furthermore, the docking scores of the top 30 compounds of DDS set of compounds along with their prediction results are shown in Table 1. It is worthy to mention here that docking results needs further verification by biological testing to further confirm and modify the experimental process for these potent ligands for the treatment of MP pneumonia.

3.2. Exploring the chemical space of TLR4 inhibitory compounds

The QSAR paradigm is based on the link between the molecular structure of compounds and their respective biological activities. Molecular descriptors play a critical role in providing analytical descriptions of the physicochemical properties of molecules. In order to adequately account for these structural features, it is of utmost importance to select appropriate descriptors for a QSAR analysis. Todeschini *et al.*⁴² have compiled a handbook providing comprehensive coverage of molecular descriptors. This study selected a subset of molecular descriptors and fingerprints representative of the general characteristics of molecules (*i.e.*, flexibility, molecular size, solubility, polarity, electronic properties, and charge, as well as chemical reactivity) that are mostly correlated with bioactivities. Using such descriptors, we can explore the chemical space of inhibitors as mentioned earlier.⁴³ Lipinski's rule-of-five (Ro5) descriptors are used to examine the general chemical space of the investigated data set. According to previous studies have suggested that chemical space analysis provides a mechanistic explanation of the correlation between MW, ALog *P*, number of hydrogen bond donors and number of hydrogen bond acceptors of a compound and its degree of potency and pIC₅₀ values.⁴⁴ In order to visualize the relative distribution of the bioactivity classes and Ro5, scatter and box plots were created, as shown in Fig. 3(A–G). The results showed that 60% of compounds have molecular weight of less than 500 Da (Fig. 3(G)). Whereas the



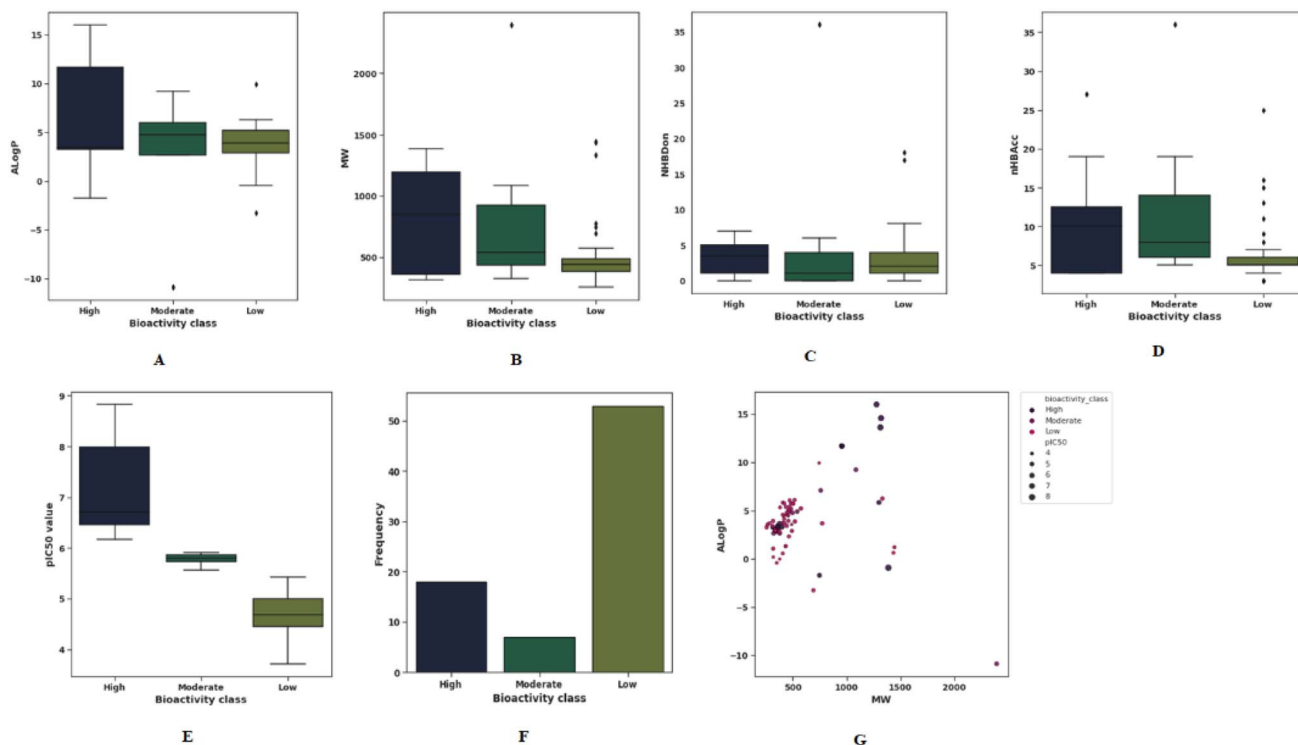


Fig. 3 Panels (A–F) show exploratory TLR4 inhibitors data analysis and panel (G) shows chemical space analysis. The scatter plot showed the diversity of ALog *P* versus MW of TLR4 inhibitory compounds.

ALog *P* for the majority of compounds varies between 2 and 7. Molecular lipophilicity can be measured by ALog *P* where a high ALog *P* value indicates high lipophilicity whereas a low value suggests low lipophilicity. The ALog *P* is a computational estimator of the logarithm of the partition coefficient between water and octanol, which has been indispensable in determining molecular hydrophobicity. The boxplots indicate distribution and frequency of high-, moderate- and Low-class compounds over the Ro5 (Fig. 3 (A–D)). The Ro5 index may be used to distinguish compounds based on their pharmacological effects based on their molecular properties, namely the octanol–water partition coefficient ($\log P < 5$), the molecular weight (< 500), the number of hydrogen bond donors (> 5), and the number of hydrogen bond acceptors (< 10). The Ro5 were found to be of limited use in contributing to our understanding of the targets–ligands relationship (*i.e.*, their affinity towards the target) as they were based solely on general ligand properties. Oprea *et al.* demonstrated that the Ro5 criteria are not effective in discriminating between drugs and non-drugs based on the

availability of more than 90% of the chemical reagents listed in the Available Chemical Directory that meets Ro5 criteria.⁴⁵ The Ro5 criteria, however, do not eliminate the possibility that they may be used to narrow the pharmacokinetic space for therapeutically relevant compounds. In addition, Benet *et al.* have demonstrated that a QSAR model developed using the Ro5 criteria can effectively predict drug disposition characteristics for drugs that meet or do not meet the Ro5.⁴⁶

3.3. Regression QSAR models of TLR4 inhibitors

The development of QSAR models was conducted using curated data sets containing 78 structurally diverse compounds spanning several scaffolds to predict TLR4 inhibitory activity. The molecular features of compounds are described using several 1D and 2D descriptors and fingerprint types. Several rounds of data splitting were used to test the generalization and interpolation abilities of QSAR model. This model was designed using a data split ratio of 70/30, in which 70% of the data set was used as the internal set, while 30% of the data set was used as

Table 2 Performance summary of different regression models

Models	R2 (train)	RMSE (train)	MAE (train)	R2 (test)	RMSE (test)	MAE (test)	R2 (CV)	RMSE (CV)	MAE (CV)
RFR	0.91	0.36	0.25	0.89	0.39	0.3	0.71	0.65	0.68
ETR	0.96	0.23	0.06	0.76	0.53	0.41	0.76	0.56	0.41
DTR	0.96	0.22	0.04	0.82	0.53	0.42	0.63	0.69	0.51
ABR	0.91	0.36	0.26	0.89	0.39	0.29	0.74	0.59	0.42
GBR	0.96	0.23	0.06	0.93	0.29	0.21	0.79	0.51	0.36



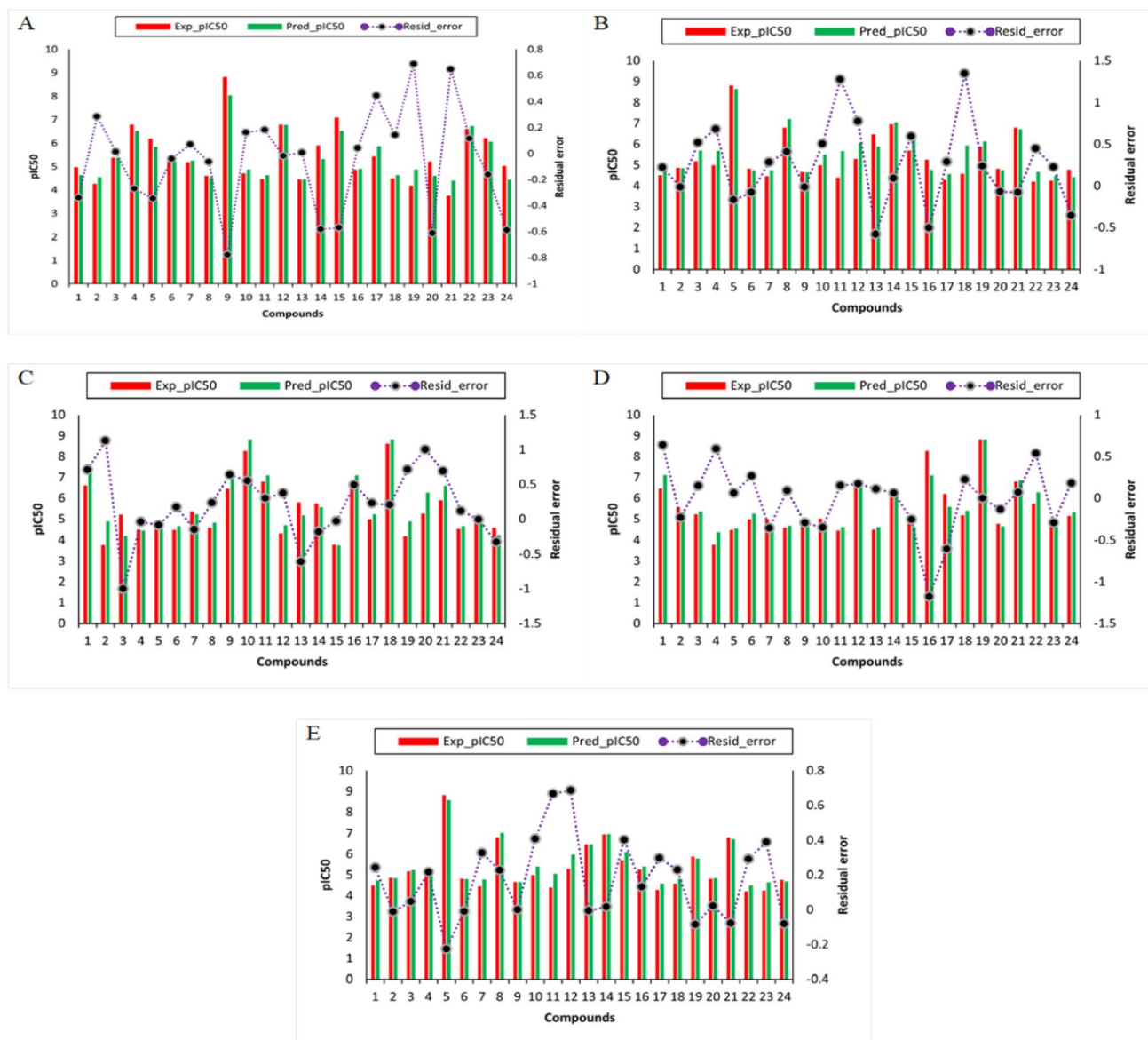


Fig. 4 Plot of experimental *versus* predicted pIC_{50} values of TLR4 inhibition for QSAR regression models generated by RF (A), ETR (B), DTR (C), ABR (D) and GBR (E) for the testing set (24 compounds). The X-axis shows compounds. The Y-axis on the left represent the pIC_{50} values and the Y-axis on the right shows residual error.

the external set. In the first subset of 70% of the data, the model was internally validated by using it both as the training set as well as the cross-validation set, and its performance was evaluated according to R^2 , RMSE, and MAE. This second set of data containing 30% of the bioactivity data was used for external validation, and the performance of the models was based on R^2 , RMSE, and MAE. Table 2 summarizes the results of the models

constructed using the RF algorithm. The descriptors and fingerprints including feature importance score on which the final RF model was trained and tested is provided in ESI S2.† The feature importance score was calculated from the RF algorithm based on its built-in function for determining feature importance. The descriptors and fingerprints which have a score greater than 0.00011 were included in the training of the

Table 3 Performance summary of RF classification model on train, test, and validation data

Accuracy (train)	RMSE (train)	MAE (train)	Accuracy (test)	RMSE (test)	MAE (test)	Accuracy (CV)	RMSE (CV)	MAE (CV)	Log loss
1	0	0	0.96	0.2	0.04	0.82	0.46	0.19	0.27



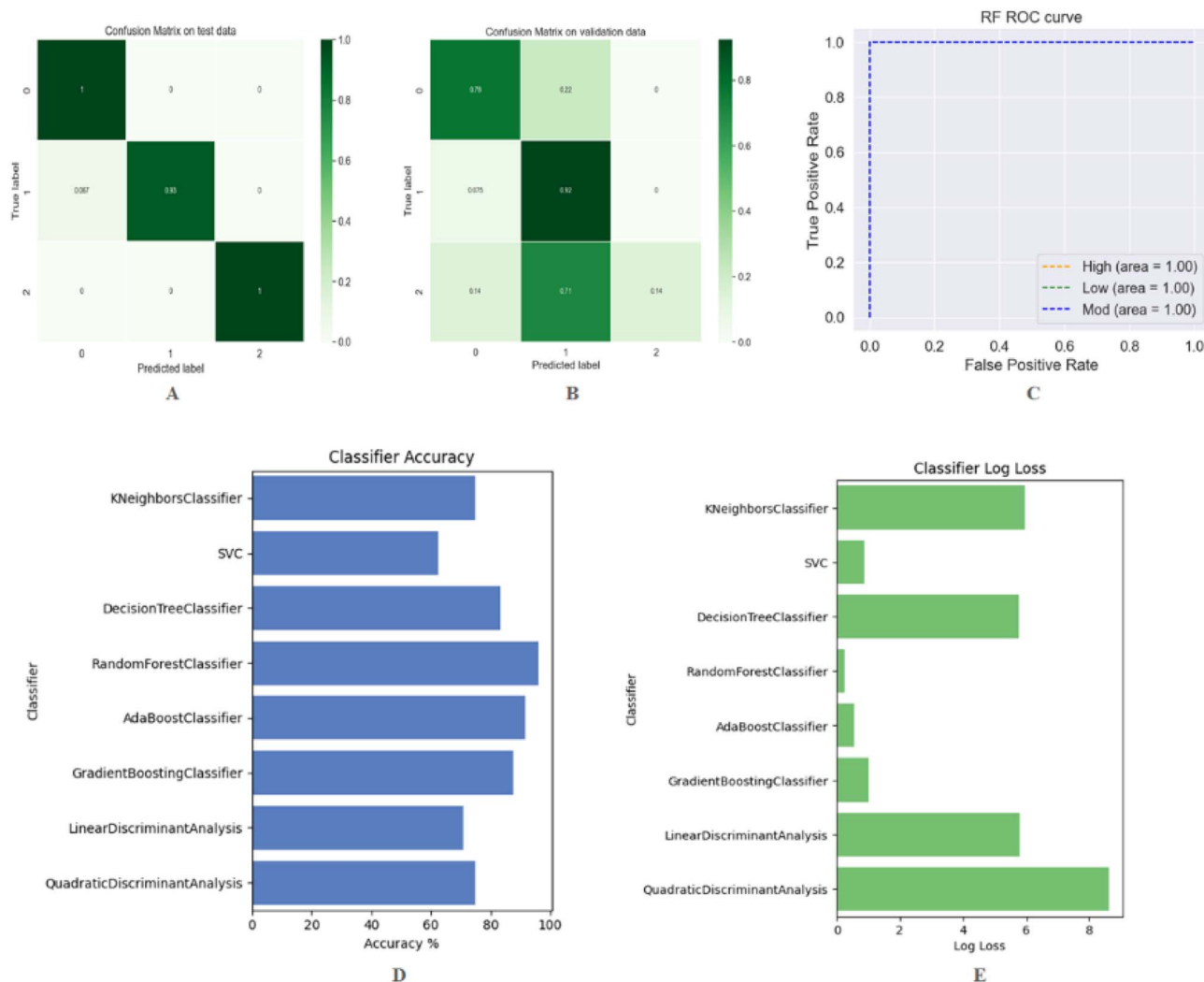


Fig. 5 (A and B) Depicts the confusion matrix related to the test and 5-fold cross validation. (C) Shows the multiclass ROC curves for the RF classifier. The performance accuracy and log loss of each classifier are illustrated in (D) and (E) respectively.

model. While the descriptors and fingerprints whose score is below 0.00011 were discarded because these descriptors and fingerprints do not affect the model performance significantly based on our pre-experiments. The predictive power of the model was assessed according to the thresholds suggested by previous studies,^{47,48} in which acceptable models possess R^2 (train) > 0.6 and R^2 (test) > 0.5. Meanwhile, it is necessary to develop a generalized QSAR model that is capable of automatically identifying informative features from a large pool of chemical descriptors that will result in a better comprehension of the mechanism of chemical compounds. The model has to be simple, user-friendly, and consistent with acceptable prediction results. Thus, the RF algorithm was utilized since it is easily able to calculate the relative importance of molecular fingerprints used during model construction. Further, the RF method was employed in order to select informative molecular descriptors because of its built-in function for measuring the importance of descriptors. Additionally, an insight into the applicability domain of the TLR4 QSAR model is established in the current study for the prediction of pIC_{50} values for DDS set of

compounds. In the present study, it was taken into account the robustness of the model, goodness-of-fit and predictability were used to determine how well the model was performing. In this study, the model was further evaluated against several other algorithms and developed QSAR models for the prediction of potent compounds. The performance summary of the RF regression model and its comparison with other models (extra-trees regression (ETR), decision tree regression (DTR), adaboost regression (ABR), gradient boosting regression (GBR)) is shown in Table 2. While the difference (residual error) between the predicted and experimental bioactivity is represented in Fig. 4. From these results, it has been suggested that these models performed better on training, testing and validation data. The comparison results showed that the proposed models are quite promising and hold potential for the prediction of TLR4 inhibitory compounds. To the best of our knowledge, this is the first study that proposed QSAR models for TLR4 receptor as a target for MP pneumonia. In a previous study, novel antagonists of Toll-Like Receptor 4 (TLR4) were identified using structure and ligand-based virtual screening. They noted that



Table 4 Classification report of RF classification model on the test and cross validation data

	Precision	Recall	F1-score	Support
Classification report of RF model on test data				
0	0.888889	1	0.941176	8
1	1	0.933333	0.965517	15
2	1	1	1	1
Accuracy	0.958333	0.958333	0.958333	0.958333
Macro avg.	0.962963	0.977778	0.968898	24
Weighted avg.	0.962963	0.958333	0.95884	24
Classification report of RF model on cross validation data				
0	0.736842	0.777778	0.756757	18
1	0.844828	0.924528	0.882883	53
2	1	0.142857	0.25	7
Accuracy	0.820513	0.820513	0.820513	0.820513
Macro avg.	0.860557	0.615054	0.62988	78
Weighted avg.	0.833834	0.820513	0.79698	78

three *in silico* hits exhibited promising anti-TLR4 activities with micromolar IC₅₀ values, and significantly reduced the production of TNF- α .⁴⁹ Incredibly, several TLR4 antagonists are already being investigated as potential anti-sepsis drugs. The most advanced of these compounds, TAK-242 and eritoran, have shown promising results in preclinical studies. These two strategies failed to meet their primary objective which was to reduce the mortality rate in patients with sepsis.^{50,51} However, a critical need remains for the identification of new TLR4 antagonists that can serve as novel therapeutics.

3.4. Classification QSAR models of TLR4 inhibitors

The models were constructed in a similar manner to those for regression, with the exception that quantitative values were replaced by semi-quantitative labels (high, moderate and low) for bioactivity. We present a simple and general-purpose RF method for predicting the low, medium, and high-class DDS set of compounds. The analysis of the training, testing, and cross validation performance of the RF classification model is presented in Table 3. The performance of machine learning models was evaluated by evaluation metrics such as sensitivity and specificity including the classification accuracy, confusion matrixes, true positive rates for a given model and false positive rates as mentioned previously.⁵² Confusion matrixes are very popular in solving classification problems. Fig. 5(A and B)

depicts the confusion matrix related to the test and 5-fold cross validation. An important tool for evaluating a machine learning model's performance is the receiver operating characteristic curve (ROC curve). ROC curves demonstrate the correlation between true positive rates (TPR) for a given model and false positive rates (FPR). The Fig. 5(C) shows the multiclass ROC curves for the RF classifier. On a class-by-class basis, the classification report presents a summary of the principal classification metrics. Table 4 provides a classification report that includes precision, recall, and F1-scores on both test data and cross validation for each of the three classes. A deeper understanding of the classifier behavior over global accuracy may be gained by assessing the behavior of a specific class within a multiclass problem. All the evaluating parameters (classification report, performance summary, ROC curve, and confusion matrix) confirmed that the developed RF classification model performed better on training, testing, and validation data. For further verification, it is necessary to compare the proposed method with other existing methods. The developed RF model was evaluated against a variety of different classifiers including KNeighbors, support vectors, decision trees, ada-boost, gradient boosting, linear discriminant analysis, and quadratic discriminant analysis. The performance accuracy and log loss of each classifier are illustrated in Fig. 5(D) and (E) respectively. The performance summary of all these classifiers is presented in Table 5. The results clearly indicate that the RF classifier outperforms all other data-based classifiers both on the test and on the trained data. It is evident from the comparison results that the proposed RF classifier, as presented in this study, is quite promising. It has the potential to be a useful tool for discriminating low, moderate, and high potency compounds. Furthermore, the proposed RF model can be viewed as complementary to the existing method in the same area.

3.5. Predictions of DDS set of compounds

To predict the bioactivity of compounds, we applied the final selected QSAR models as prediction models. The top 30 compounds selected from DDS set on the basis of docking scores and their predicted pIC₅₀ bioactivities derived from machine learning models are summarized in Table 1, respectively. RF model has been compared with several other models, in order to reduce redundancy in the predicted results. A comparison of the results generated by the proposed models

Table 5 Performance summary of different classifiers on the train and test data

Models	Accuracy (train)	RMSE (train)	MAE (train)	Accuracy (test)	RMSE (test)	MAE (test)	Log loss
RF model	1	0	0	0.96	0.2	0.04	0.27
KNeighbors classifier	0.76	0.64	0.3	0.75	0.5	0.25	5.95
SVC	0.7	0.54	0.29	0.63	0.61	0.37	0.87
Decision-tree classifier	1	0	0.35	0.88	0.35	0.13	4.31
AdaBoost classifier	0.98	0.14	0.02	0.92	0.46	0.13	0.54
Gradient boosting classifier	1	87.5	0	0.88	0.35	0.13	0.97
Linear discriminant analysis	0.98	0.27	0.04	0.71	0.54	0.29	5.79
Quadratic discriminant analysis	1	0	0	0.75	0.5	0.25	8.63



suggests that these models have promising results and can be used to predict compounds that inhibit TLR4. This is the very first study that comes to our knowledge that proposes QSAR study to be used as a target in MP pneumonia based on a TLR4 receptor. Additionally, these results can be also used as references to find the structures of compounds that are potentially TLR4 modulating as well as saving time and money compared to the traditional biological screening procedures. In addition, previous studies have explored the importance of these QSAR models for *in silico* prediction of novel compounds that can be used to predict the activity of a target or receptor when this is a computationally intensive process that requires learning from bioactivity data.^{53–56}

4. Conclusion

The pathological process of MP pneumonia has been identified as being associated with TLR4 and has been described as one of the most difficult and threatening diseases of the human population. TLR4 can be successfully inhibited therapeutically for the prevention of MP. A widespread use of traditional antibiotics in clinical settings has led to an increase in drug resistance and other adverse effects. Thus, natural products may have the potential to be used to modulate the host immune inflammatory response in the prevention of TLR4-driven diseases. Hence, regulating the host immune inflammatory response by means of natural products might represent a new strategy for preventing MP infection. As a result, the project can serve as a theoretical base for the treatment of TLR4-related illnesses by regulating the immune response of the host species. Hence, the present research aims to screen medicinal compounds to find new candidate drugs to treat TLR4-mediated inflammatory diseases. Recently, machine learning is reshaping research in various fields.^{57–60} Furthermore, molecular docking analysis of the inhibitors and key residues in the binding pocket of the TLR4 protein also suggested the presence of several hydrogen bonds and hydrophobic interactions between inhibitors and key residues. Nevertheless, further research will be performed to test the screened and selected candidates, and in-depth molecular research will be conducted in order to understand the mechanism of action, to provide new ideas for new anti-inflammation medications targeting TLR4. This study explores the relationship between structure and activity for a library of 78 compounds from the ChEMBL database that are known TLR4 inhibitors. Using SMILES-based descriptors, we propose a simple method for constructing both regression and classification QSAR models. For the numerical description of the compounds, several 1D and 2D chemical descriptors and molecular fingerprints were employed. By analyzing the calculated molecular descriptors, the chemical space of TLR4 inhibitors was explored thus providing important insights into their molecular characteristics. Presented models have the advantage of being simple and easy to use and interpret. Key features that are correlated with pIC_{50} values, as determined by the RF algorithm. In the future, such information can be used to guide the design of novel molecular structures that act as TLR4 inhibitors. These classification and regression QSAR models

have demonstrated robust statistical results and are interpretable, which strongly supports their use for rapid screening of TLR4 inhibitors.

Author contributions

All authors contributed to the study conception and design. Data collection and formal analysis were performed by Muhammad Ishfaq, Ziaur Rahman, Akhunzada Bilawal and Sattar Hamid. Zemin Zhu, Sihong Li and Muhammad Ishfaq supervised and provided the resources and funding. Muhammad Aamir performed visualization and data curation. Syed Zahid Ali Shah helps in critical revision of the manuscript. The first draft of the manuscript was written by Muhammad Ishfaq and all authors commented on the manuscript. All authors read and approved the final version of the manuscript.

Conflicts of interest

The authors report no conflicts of interest in this work.

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