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Antitumor Agents 252. Application of Validated QSAR Models to Database Mining: Discovery of Novel Tylophorine Derivatives as Potential Anticancer Agents

Shuxing Zhang^{†,§}, Linyi Wei^{†,≠}, Ken Bastow[≠], Weifan Zheng[§], Arnold Brossi[≠], Kuo-Hsiung Lee^{≠,*}, and Alexander Tropsha^{§,*}

[§]Laboratory for Molecular Modeling and Carolina Exploratory Center for Cheminformatics Research, Division of Medicinal Chemistry and Natural Products, School of Pharmacy, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina 27599

[#]Natural Products Laboratory, Division of Medicinal Chemistry and Natural Products, School of Pharmacy, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina 27599

Abstract

A combined approach of validated QSAR modeling and virtual screening was successfully applied to the discovery of novel tylophrine derivatives as anticancer agents. QSAR models have been initially developed for 52 chemically diverse phenanthrine-based tylophrine derivatives (PBTs) with known experimental EC_{50} using chemical topological descriptors (calculated with the MolConnZ program) and variable selection k nearest neighbor (kNN) method. Several validation protocols have been applied to achieve robust QSAR models. The original dataset was divided into multiple training and test sets, and the models were considered acceptable only if the leave-one-out cross-validated R^2 (q²) values were greater than 0.5 for the training sets and the correlation coefficient R^2 values were greater than 0.6 for the test sets. Furthermore, the q^2 values for the actual dataset were shown to be significantly higher than those obtained for the same dataset with randomized target properties (Y-randomization test), indicating that models were statistically significant. Ten best models were then employed to mine a commercially available ChemDiv Database (ca. 500K compounds) resulting in 34 consensus hits with moderate to high predicted activities. Ten structurally diverse hits were experimentally tested and eight were confirmed active with the highest experimental EC_{50} of $1.8\mu M$ implying an exceptionally high hit rate (80%). The same ten models were further applied to predict EC50 for four new PBTs, and the correlation coefficient (R^2) between the experimental and predicted EC_{50} for these compounds plus eight active consensus hits was shown to be as high as 0.57. Our studies suggest that the approach combining validated QSAR modeling and virtual screening could be successfully used as a general tool for the discovery of novel biologically active compounds.

Introduction

Natural products have been the major source of anticancer drugs. According to a recent review¹ on New Chemical Entities (NCE), from 1981 to 2002, approximately 74% of anticancer drugs were either natural products, or natural product-based synthetic compounds, or their mimetics. (+)-(*S*)-Tylophorine (**1**) and its analogues are phenanthroindolizidine alkaloids, commonly referred to as tylophora alkaloids. They are a small group of alkaloids known for their profound cytotoxic activities^{2–5}. Evaluation of these compounds in the

^{*}A. Tropsha, alex_tropsha@unc.edu. *K.H. Lee, khlee@unc.edu.

[†]Both authors have contributed equally to this paper

antitumor screening at the National Cancer Institute (NCI) showed a uniform and potent inhibitory effect on the cell growth (GI₅₀, $\cong 10^{-8}$ M) in all 60 cell lines, with notable selectivity toward several refractory cell lines, including melanoma and lung tumor cell lines⁶. Earlier studies in 1960s' demonstrated that their mechanism of antitumor activity was due to irreversible inhibition of the protein synthesis at the elongation stage of the translation cycle^{7–10}. In 1990s', several key metabolic enzymes were reported as biological targets of tylophorine alkaloids including thymidylate synthase (TS)¹¹ and dihydrofolate reductase (DHFR)¹². In addition, these agents were found to induce cell apoptosis as well¹³. Most recently, Gao et al. demonstrated that tylophorine analogues had significant inhibitory effect on NF-*k*B mediated transcription¹⁴. These discoveries exemplified the great potential of developing tylophorine derivatives as a new class of antitumor drugs.

Although the phenanthroindolizidine alkaloid tylocrebrine (2) previously failed in clinical trials due to its CNS toxicity¹⁵, the very profound cytotoxicity of these alkaloids¹⁶, particularly against multidrug resistant cancer cells¹⁶, sparked our interest in additional studies of this class of potential anticancer agents. Recently, we have advanced a novel series of polar, watersoluble phenanthrene-based tylophorine derivatives (PBTs) (6, 18) with $EC_{50} \approx 10^{-7}$ M against the A549 human lung cancer cell line¹⁷. These compounds could possibly have lower or no CNS toxicity because their increased polarity should prevent them from penetrating the bloodbrain barrier. Although the biological target of PBTs is unknown, several structure-activity trends have been observed¹⁷: 1) A planar phenanthrene system is required, but not sufficient for cytotoxic activity. 2) An N-hydrophilic substituent at the C-9 position is essential for the enhanced cytotoxicity and should be linked through a methylene rather than a carbonyl group. 3) The C-9 N-hydrophilic substituent is ideal for the introduction of a polar moiety. Analogs containing terminal carboxylic acid or hydroxymethyl groups are more favorable than those with methyl esters. 4) On the phenanthrene skeleton, a methoxyl substituent best fits both the steric and electronic requirements at the C-6 position and is preferred over benzyloxyl and hydroxyl groups. 5) Adding a methylenedioxyl ring at the 2, 3 positions of the planar phenanthrene system dramatically enhances the cytotoxic activity and leads to the most potent derivatives. The new PBT derivatives possess a novel structure and show remarkable EC_{50} values in the sub-micromolar range¹⁷, comparable to those of the front-line antineoplastic drugs, and suggesting that this new class of compounds may have a great potential as antitumor agents. The availability of experimental data on PBT derivatives afforded us an opportunity to apply advanced computational drug discovery approaches, in particular QSAR modeling, towards knowledge based accelerated discovery of novel anticancer agents.

Many different QSAR approaches have been developed during the past few decades^{18–21}. Modern methods are characterized by the use of multiple descriptors of chemical structure combined with the application of both linear and non-linear optimization approaches, and a strong emphasis on rigorous model validation to afford robust and predictive OSAR models (see recent reviews^{22;23}) The most important recent developments in the field have concurred with a substantial increase in the size of experimental datasets available for the analysis and an increased application of QSAR models as virtual screening tools to discover biologically active molecules in chemical databases and/or virtual chemical libraries^{23;24}. The latter focus differs substantially from the traditional emphasis on developing so called explanatory OSAR models characterized by high statistical significance but only as applied to training sets of molecules with known chemical structure and biological activity. Our laboratory recently developed a robust computational QSAR modeling framework that combines various algorithms for model development (such as k nearest neighbor (kNN) QSAR approach with variable selection²⁵), model validation, and model implementation for virtual screening^{24;26}. This strategy was recently applied to several experimental datasets leading to the successful discovery of novel anticonvulsant agents²⁶ and D₁ dopaminergic antagonists²⁷.

In this paper, we discuss the application of the *k*NN QSAR method to a dataset of 52 PBTs with known EC₅₀ values (Table 1, Table 2 & Table 3). The structures were characterized with MolConnZ descriptors²⁸. The models developed for the PBT dataset have been extensively validated using several criteria of robustness and accuracy²⁹. Several validated models with the high predictive power were used to mine the commercially available ChemDiv³⁰ database resulting in 34 consensus hits with the moderate to high predicted activities. Ten structurally diverse hits were experimentally tested and eight compounds were confirmed active, with the most potent compound having EC₅₀ of 1.8μ M. The predictive power of these models were further confirmed by the high correlation coefficient between the predicted and actual cytotoxicity for an external set comprised of four new PBTs and the eight active ChemDiv hits, which were not included in the original 52 PBTs dataset. The correlation coefficient (R²) was as high as 0.57. The results of this study suggest that rigorously validated QSAR models could be successfully used as virtual screening tools for prioritizing untested compounds for experimental biological evaluation.

Materials and Methods

Chemistry and Biological Activity Data

All PBTs used in this study were synthesized and evaluated (Table 1, 2, 3 & 5) in one of our laboratories. The general synthetic procedure, biological activity, physical and spectral data have been reported previously¹⁷. The hit compounds identified by the means of database mining were purchased from ChemDiv, Inc. The human A549 lung cancer cell line was used for the cytotoxicity screening of both PBTs synthesized earlier as well as novel computational hits, employing a cell-based sulforhodamine B (SRB) microtitre plate assay³¹. The screening method was reported in detail elsewhere¹⁷.

Generation of Molecular Descriptors

All chemical structures were generated using SYBYL 7.0³². Molecular descriptors were calculated for each compound with the MolConnZ software version 4.05^{28;33}. MolConnZ produced more than 400 descriptors; however, in our study, only 244 significant descriptors were used after removing those with zero variance. The descriptors were range-scaled prior to model generation because the absolute scales of different descriptors differed in some cases by orders of magnitude. Range scaling helps to avoid disproportional weightings of descriptors upon the Euclidean distance calculations in multidimensional descriptor space.

Dataset Division into Training and Test Sets

It is commonly accepted that the internal validation of QSAR models built from training sets is sufficient to confirm their predictive power^{34–38}. However, previous studies in this as well as several other laboratories demonstrated that no correlation exists between leave-one-out (LOO) cross-validated R^2 (q^2) for the training set and the correlation coefficient R^2 between the predicted and observed activities for the test set^{29;39}. These findings indicated that in order to obtain QSAR models with high predictive ability, external validation was critical. Thus, a dataset of 52 compounds was divided into multiple chemically diverse training and test sets with a rational approach implemented in our group⁴⁰ based on the Sphere Exclusion (SE) algorithm⁴¹. SE is a general procedure that is typically applied to molecules characterized by multiple descriptors of their chemical structures. The entire dataset can then be treated as a collection of points (each point corresponding to an individual compound) in the MolConnZ descriptor space. The goal of the SE method is to divide a dataset into two subsets (training and test sets) using a diversity sampling procedure⁴⁰.

The SE algorithm used in this study included the following steps.⁴⁰ The algorithm starts with the calculation of the distance matrix D between points representing compounds in the

multidimensional descriptor space. Let D_{min} and D_{max} be the minimum and maximum elements of *D*, respectively. *N* probe sphere radii are defined by the following formulas: $R_{min}=R_1=D_{min}$, $R_{max}=R_N=D_{max}/4$, $R_i=R_1+(i-1)*(R_N-R_1)/(N-1)$, where i=2, ..., N-1. Each probe sphere radius corresponds to one division into the training and test sets. Once the sphere size is defined the subsequent calculations include the following steps: (i) Select randomly a point in the MolConnZ descriptor space. (ii) Include it in the training set. (iii) Construct a probe sphere around this point. (iv) Select points from this sphere and include them alternatively into test and training sets. (v) Exclude all points within this sphere from further consideration. (vi) If no more compounds left, stop. Otherwise let *m* be the number of probe spheres constructed and *n* be the number of remaining points. Let d_{ij} (i=1,...,m; j=1,...,n) be the distances between the remaining points and probe sphere centers. Select a point corresponding to the lowest d_{ij} value and go to step (ii). The training sets were used to build models and the test sets were used for model validation.

kNN QSAR Method

Our implementation of this method²⁵ employs the *k*NN pattern recognition principle⁴² and a variable selection procedure. Briefly, a subset of *nvar* descriptors (number of selected variables) is selected randomly. A model is built using this random descriptor selection with LOO cross-validation, where each compound is eliminated from the training set and its biological activity is predicted as the average activity of its *k* most similar molecules (usually k = 1-5). The value *k* is optimized during the model building process to give the best prediction for the training set. The similarity is characterized by the Euclidean distance between compounds in multidimensional space of selected descriptors. We used weighted molecular similarity²⁷ to calculate the estimated activities \hat{y}_i of compounds excluded by LOO procedure using the following formula

$$\widehat{y}_{i} = \frac{\sum_{j=1}^{k} a_{j} w_{ij}}{\sum_{j=1}^{k} w_{ij}},$$
(1)

where a_i was the observed activity of the *j*th compound, and weights w_{ij} are defined as

$$w_{ij} = \left(1 - \frac{d_{ij}}{\sum_{j=1}^{k} d_{ij}}\right),$$
(2)

and d_{ij} was the distance between compound *i* and its *j*th nearest neighbor. After each run, cross-validated q² is calculated

$$q^{2} = 1 - \frac{\sum (y_{i} - \widehat{y}_{i})^{2}}{\sum (y_{i} - \overline{y})^{2}},$$
(3)

where y_i , and \bar{y} are the actual and average values of activity. The summation in (3) is performed over all compounds. A method of simulated annealing with the Metropolis-like acceptance

In summary, the *k*NN QSAR algorithm generates both an optimal *k* value and an optimal *nvar* subset of descriptors that afford a QSAR model with the highest value of q^2 . Figure 1 shows both the overall flowchart of the current implementation of the *k*NN method (1a) and the predictive QSAR modeling workflow (1b).

Robustness and Predictive Power of QSAR Models

The robustness of the models was examined by comparing them to those obtained when using randomized activity of the training set (this procedure is commonly referred to as Y-randomization test)⁴⁴. Briefly, the QSAR calculations were repeated with the randomized activities of the training sets. Then the q^2 values for actual and random activities of training sets were compared to see whether there was a significant difference as expected for robust models. This test was applied to all data divisions considered in this study and it was repeated five times for each division.

To estimate the predictive power of a QSAR model, the following parameters were used²⁹: (i) correlation coefficient R^2 between the predicted and observed activities; (ii) coefficients of

determination (predicted versus observed activities R_0^2 , and observed versus predicted activities

 $R_{0}^{(2)}$; (iii) slopes *k* and *k*' of regression lines (predicted versus observed activities, and observed versus predicted activities) through the origin. We concluded that a QSAR model had an acceptable predictive capability if the following conditions were satisfied²⁹:

$$q^2 > 0.5;$$
 (4)

$$R^2 > 0.6;$$
 (5)

$$|R_0^2 - R_0^2| < 0.3; \tag{6}$$

$$0.85 \le k \le 1.15 \text{ or } 0.85 \le k' \le 1.15.$$
 (7)

Applicability Domain of kNN QSAR Models

Following the procedures developed earlier,^{44;45} the distances (similarity) of compounds in our training set were compiled to produce an applicability domain threshold (i.e., similarity threshold), D_T , calculated as follows:

$$D_{T} = \langle d \rangle + Z\sigma \tag{8}$$

where $\langle d \rangle$ is the average of Euclidean distances between *k* nearest neighbors of all compounds of the training set used in model derivation, σ is the standard deviation of these distances, and *Z* is the empirical parameter to control the significance level. The default value of *Z* was set to 0.5, which formally placed the boundary for the compounds to be predicted at one-half of the standard deviation (assuming a normal distribution of distances between *k* nearest neighbor compounds in the training set). Thus, if the distance of an external compound from at least one of its nearest neighbors in the training set exceeded this threshold, the prediction was considered unreliable.

Database Mining

A commercially available chemical database, Chemical Diversity (ChemDiv)³⁰, containing ca. 500K compounds, was used for virtual screening. MolConnZ descriptors were generated for each compound in the databases and linearly normalized based on the maximum and minimum values of each descriptor in the training set²⁷.

Ten best *k*NN models were used to predict activities of the database compounds that were within the applicability domain of each individual model. The results for each individual prediction exercise were then combined and the mean predicted activity was calculated for each compound that was within the applicability domain of multiple models. The number of models that predicted each compound in the database and the standard deviation of those predictions for each compound were also recorded. We selected a subset of compounds as hits that were predicted by at least 50% of the models and exhibited a small standard deviation across all models. We also performed an additional analysis as to whether the hits resulting from database mining possessed the features of PBTs essential for their activities. This additional consideration was considered important because it helped us identify novel scaffolds expected to have anticancer activities.

Results and Discussions

QSAR Models and Their Robustness

In the *k*NN QSAR method, *nvar* can be set to any value that is less than the total number of descriptors. Since the optimal number for *nvar* is not known *a priori*, multiple models have to be generated to examine the relationship between q^2 and *nvar*. As previously discussed, Y-randomization is a widely used technique to ensure the robustness of a QSAR model⁴⁶. It is expected that the resulting QSAR models from the randomized sets should generally have low training set fitness q^2 and low test set R^2 . Sometimes, though infrequently, high training set q^2 may be obtained due to a chance correlation or structural redundancy of the training set q^2 and test set R^2 values, it implies that an acceptable QSAR model cannot be built for the given dataset by the current modeling method.

To compare results from the actual data set with those from data sets with randomized activity values, models with 10, 20, 30, 40 and 50 descriptors were generated. Figure 2 shows a plot of q^2 vs. *nvar* for the actual and random data sets obtained with *k*NN calculations. Every q^2 value was the average of 10 independent computations. Overall, we have obtained consistently much higher q^2 values for the actual data set compared to those from Y-randomization. The q^2 values for the real data set were in the range 0.60 to 0.7 while they were from -0.01 to 0.05 for the random data sets. This demonstrated that the high q^2 values of the models for the real data sets were not due to chance correlations.

kNN QSAR Model Validation

Generally, we accept models with q^2 values for the training set greater than 0.5 and R^2 values for predicted vs. actual activities of the test set compounds greater than 0.6.³⁴ Table 4 presents 10 best models obtained from multiple *k*NN analyses. In order to estimate the statistical significance of the models, the original dataset of 52 compounds was divided into 50 training and test sets. Multiple *k*NN models with the high q^2 values (greater than 0.5) were collected. However, similar to our previous observations²⁹, no correlation was found between q^2 and R^2 (Figure 3) demonstrating that q^2 alone does not serve as an estimate of the predictive power of *k*NN models. On the basis of our criteria, acceptable models with both high statistical significance ($q^2 > 0.5$) and predictive power ($R^2 > 0.6$) represented only a fraction of all models with $q^2 > 0.5$ (Figure 3). So the aforementioned conditions (Eq. 4, 5, 6, and 7) are indeed very important. Based on all of these criteria, the best models were obtained for the test sets including 14 and 18 compounds, with the optimal number of descriptors of 15 and 20, respectively (Table 4).

Figure 4 shows the correlation coefficient between actual and calculated activity for the training and test sets with $q^2 = 0.59$ and $R^2 = 0.81$ respectively. Two outlier points (compound **20** and **54**) were poorly predicted in the training set. The one in black circle represents antofine (Figure 4), a positional isomer of tylophorine isolated from *Asclepiadaceae* by Dr. T. S. Wu in Taiwan⁴⁷. It was used as a reference compound when we screened PBTs. A possible explanation for this observation is that antofine lacks the chemical structure descriptors that are most meaningful and statistically significant in terms of correlation with biological activity: a *N*-hydrophilic substituent and free rotated C-9 methylene bond, which were proven to be required for the PBTs analogs activity in our previous study¹⁷. Studies on the mechanisms of action for antofine and PBTs are still ongoing, and it remains possible that they have different mechanisms of action. In regards to compound **20** (in grey circle, Figure 4), the terminal -Cl group may cause some solubility or cell membrane transportation problem that would result in diminished activity in our cell-based assay. After excluding these two outlying points, the q^2 for training set increased to 0.69. Given the high predictive R^2 (0.81), this model was considered acceptable for virtual screening.

Interpreting Predictive QSAR Models

Upon the analysis of our QSAR models, a number of MolConnZ descriptors were found in most of the accepted models, suggesting that they played critical roles in defining antitumor activity of PBTs. These included molecular connectivity descriptors such as Chi indices, hydrogen bond counts, E-state descriptors, shape indices, etc. The results were consistent with our preliminary SAR observations described in the **Introduction** Section.

Based on the MolConnZ manual and personal communication with Dr. Lowell Hall, one of the principal developers of MolConnZ software, these selected descriptors were grouped into the following classes: 1) The high frequency of Chi indices, including dXvp7, dXvp10, nXp6, Xvp7, etc., suggested the importance of those structure features such as size, branching, cyclicity and so on. Our previous SAR study showed that the para-positioned functional groups and three intra-atomic distances were very important for the antitumor activity. These features defined the relative spatial dispositions of three significant atoms: the oxygen of C-9 chain terminal, the nitrogen atom, and a methoxyl oxygen of the phenanthrene ring. 2) Hydrogen bond donors/acceptors counts and E-states descriptors, such as nHBint9, SHBd, Hmax, and Hmin, indicated the importance of polar hydrogen atoms and hydrogen bond interactions during binding. This observation agreed with the finding that the presence of a hydrogen bond acceptor/donor group at C-9 chain terminus appeared essential for the cytotoxic activity, and analogs containing terminal carboxylic acid or hydroxymethyl groups were more favorable than those with methyl esters. 3) The appearance of atom type counts and E-states descriptors

(SssCH2, nsssCH, and SsOH) demonstrated the high importance of electron accessibility for those atoms. Our studies indeed found that an *N*-hydrophilic substituent at the C-9 position was essential for enhanced cytotoxicity and should be linked through a methylene rather than a carbonyl group. 4) Several steric crowding related descriptors (n2Pag12 and Tm) occurred many times in our models. This point was reflected in our finding that, for example, a methoxyl substituent best fitted both the steric and electronic requirements at the C-6 position. 5) Several Kappa and complexity indices (ka1, ka2, tets1, tets2, graph complexity, etc.) were found in many models, suggesting the shape related features were important for the cytotoxicity of our PBTs. It was noted that a planar phenanthrene system was required, although not sufficient, for the cytotoxic activity. Also adding a methylenedioxyl ring at the 2, 3 positions of the planar phenanthrene system dramatically enhanced the cytotoxic activity and led to the most potent derivatives. 6) Last, but not the least, descriptors such as nCl and SsCl indicated the importance of a -Cl group to the activity. Inclusion of this group in structures led to the significant change of their activities (e.g. compound **20, 64** and **68**).

Database Mining with Predictive QSAR Models

The ChemDiv database of over 500K compounds was screened with our ten best QSAR models within a defined applicability domain (i.e., similarity threshold). Formally, a QSAR model can predict the target property of any compound for which chemical descriptors have been calculated. Since the training set models are developed with the *k*NN QSAR approach by interpolating activities of the nearest neighbor compounds, the applicability domain should help avoid making predictions for compounds that differ substantially from the training set molecules⁴⁴. We hypothesized that the higher the number of models with a stringent applicability domain that predict a compound's activity, the more likely the compound actually has the predicted activity. This postulate may also apply to the standard deviation of the predictions made for a single compound. The smaller the prediction variance across all models, the more confidence we have that the predicted biological activity for that compound is accurate. The predicted activities for each compound from those selected models were averaged to yield consensus values.

Thirty-four consensus hits were identified with moderate to high predicted activities. Among them, some compounds shared very similar core chemical structures while the others were quite different. Eventually, ten structurally diverse hits (**59–68**, Table 5.) with moderate to high predicted activity were purchased and screened using the same cell line and assay method as used for the PBTs screening. The chemical structure and experimental biological data are shown in Table 5. Eight (**59**, **61–62**, **64–68**) of ten compounds were confirmed to be active against A549 lung cancer cell line, and compound **68** (ChemDiv #: K915-0700) showed the best activity with EC₅₀ of 1.8μ M, comparable to the active PBTs. This hit (**68**) has a novel core structure which is different from either known PBTs or other anticancer drugs, and our screening established the anticancer activity for this structure for the first time.

Prediction of Anticancer Activity for an External Data Set

The accurate prediction of the bioactivity is a more challenging task compared to the relative ranking in virtual screening. A reliable and truly predictive QSAR model should be able to accurately predict activities of new compounds in external sets. To this end, the *k*NN QSAR models validated with the test sets were used to predict the activity of four new compounds (Table 5), which were not available prior to our QSAR studies of the 52 PBT derivatives dataset. Concurrently, the eight active hits (Table 6) from ChemDiv, which had moderate to high predicted activities, were also used to evaluate the accurate activity prediction capability of our models in a quantitative manner.

As mentioned above, all of the external compounds displayed moderate to high predicted activity that ranged between 0.15 and 72.4µM (Table 5 & 6), while the original training set compounds (Table 1, 2 & 3) featured EC_{50} activity values that ranged from 0.08 to 80µM. Table 5 & 6 lists the average predicted activity values for the external data set obtained from the best kNN models. We intentionally selected a series of compounds that had a wide range of predicted activity, paralleling those used in the training and test sets during the model building. This hit selection strategy helped us confirm the predictive power of our models in a wide applicability domain. The resulting correlation coefficient R^2 was as high as 0.57 (Figure 5). It was interesting to analyze the performance of QSAR models on the congeneric (with similar core structure) compounds and novel (core structure different) compounds. With four congeneric PBTs, the difference between the predicted and experimental activity $(-\log EC_{50})$ was about 0.2. For the eight structurally diverse hits (Table 6), the result was not as good as that for the congeneric compounds. Two reasons could be suggested and discussed: 1) The congeneric and the novel hit compounds had high structural dissimilarity. The models were based on the structure-activity of the training set (congeneric), so the selected descriptors were not sufficient to reliably and accurately predict the activity of external diverse structures (novel). This problem always existed in statistical modeling and the final results could be improved by using additional descriptors. 2) Since compounds were tested in whole cell based assay, activity could deviate from the predicted values due to different mechanisms of inhibition, transportation, metabolism, etc. Although the prediction accuracy of the activity for screening hits was not as good as that for the original congeneric (core structure similar) compounds, the high experimental hit rate (eight out of ten hits were active) and the capability of detecting novel active structures from a large chemical database confirmed that this method was a very useful and powerful tool for lead identification. The overall high correlation coefficient (R²=0.57) demonstrated that our QSAR models were very robust and predictive for most of the compound structures (both congeneric and novel) and could be used to diversify the chemical repertoire of anticancer agents.

Conclusions

As part of our ongoing studies on plant-derived antitumor agents, we recently initiated the design and synthesis of new tylophorine analogs because of their profound anti-cancer activity. The structures of this class of compounds (PBTs) are still being optimized and their mechanisms of action are under investigation. Herein, we report the use of validated QSAR modeling analysis and database mining in advancing the discovery of novel antitumor agents. Using the QSAR modeling workflow we have developed robust models for a series of PBT derivatives with high internal and external prediction accuracy. These models were further exploited in database mining for new lead identification and computational hits have been tested experimentally yielding an exceptionally high <u>confirmed</u> hit rate of 80%. Those hits (such as compound **68**) lacking a phenanthrine ring but having high activity (e.g., $EC_{50}=1.8\mu$ M) may in fact lead to a novel structural class of anticancer agents.

In principle, virtual screening with QSAR models could be regarded as a sophisticated form of similarity searching. Our results demonstrate that the use of multiple descriptors of chemical structures such as molecular topological indices in validated QSAR models could indeed afford the identification of novel compounds. These results are in agreement with earlier observations of Brown and Martin⁴⁸ on the efficiency of 2D molecular descriptors in molecular similarity calculations leading to the identification of biologically active molecules. We place particular emphasis on the rigorous validation of QSAR models as well as conservative extrapolation limited to the applicability domain as two major factors that allow us to achieve the highest possible accuracy in predicted biological activity of compounds external to the training set. Furthermore, we select only consensus hits obtained with multiple validated QSAR models as opposed to the predictions based on a single best model. We believe that this approach should

facilitate the design of new natural product analogs as well as the search for new structures with anti-cancer activity from large chemical databases. We further suggest that our recent successes in identifying novel active compounds using combined application of rigorous QSAR modeling and database mining for several classes of compounds such as anticonvulsants²⁶, D1 antagonists²⁷ and now, anti-tumor agents positions the QSAR – virtual screening (QSAR-VS) as a general methodology for computer aided drug discovery.

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Figure 2.

Plots of q^2 vs. the number of descriptors selected for the best kNN QSAR models for 52 PBTs. The results for both actual and random (with shuffled activity values) data sets are shown. Every q^2 value is the average of 10 independent calculations. The black circles represent the actual data set, and the grey triangles represent the random data set.





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Figure 4.

Plot of actual vs. predicted activity for one of the best 10 models. This model has 18 compounds in the test set (grey triangles), and correspondingly 34 in the training set (black dots). Twenty descriptors were selected for the calculation. $q^2=0.59$ and $R^2=0.81$. The circled are two outliers, Antofine **54** (black circle) and Compound **20** (grey circle), respectively.

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Figure 5.

Plot of actual vs. predicted activity for the four designed compounds and eight computational hits. Compound 60 and 63 were shown to be inactive during experimental test, so the consensus EC_{50} values for only 12 out of 14 compounds were predicted based on the 10 best models with $R^2 = 0.57$.

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(-)-R-Tylophorine (1)



Tylocrebrine (2)



Chart 1. Structures of tylophorine, tylocrebrine, and PBTs 6 & 18





Compound	R ₁	R ₂
3	-CH ₃	-CONH(CH ₂) ₄ COOH
4	-CH ₃	-CH ₂ NH(CH ₂) ₄ COOMe
5	-CH ₃	-CH ₂ NH(CH ₂) ₄ COOH



	Compound	R ₁	R ₂
\bullet -CH ₂ NH(CH ₂) ₄ CH ₂ OH	6	-CH ₃	-CH ₂ NH(CH ₂) ₄ CH ₂ OH
7 -CH ₃ -CONH(CH ₂) ₅ COOH	7	-CH ₃	-CONH(CH ₂) ₅ COOH
8 -CH ₃ - CH ₂ NH(CH ₂) ₅ COOMe	8	-CH ₃	- CH ₂ NH(CH ₂) ₅ COOMe
9 -CH ₃ - CH ₂ NH(CH ₂) ₅ COOH	9	-CH ₃	- CH ₂ NH(CH ₂) ₅ COOH

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Compound	R ₁	R ₂
20	-CH ₃	
21	-CH ₃	- CH ₂ NH(CH ₂) ₁₀ COOH
22	-CH ₃	- CH ₂ NH(CH ₂) ₁₀ CH ₂ OH



Compound	R ₁	R ₂
23	-CH ₂ C ₆ H ₅	-CONH(CH ₂) ₅ COOMe
24	$-CH_2C_6H_5$	-CONH(CH ₂) ₅ COOH
25	$-CH_2C_6H_5$	-CH ₂ NH(CH ₂) ₅ COOH
26	$-CH_2C_6H_5$	-CH ₂ NH(CH ₂) ₅ CH ₂ OH



Compound	R ₁	\mathbf{R}_2
27	-CH ₂ C ₆ H ₅	-CH ₂ NH(CH ₂) ₄ COOMe
28	$-CH_2C_6H_5$	-CH ₂ NH(CH ₂) ₄ COOH















Compound	R ₁	R ₂
35	-CH ₂ C ₆ H ₅	-CONH(CH ₂) ₅ COOMe
36	-CH ₂ C ₆ H ₅	-CONH(CH ₂) ₅ COOH
37	$-CH_2C_6H_5$	-CH ₂ NH(CH ₂) ₅ COOH











Compound	R	EC ₅₀ (μM)
41	-NH(CH ₂) ₁₀ COOH	13.0
42	-NH(CH ₂) ₁₀ CH ₂ OH	3.6
43	-CH ₂ NH(CH ₂) ₅ COOH	9.7















Compound

R

 $EC_{50}\left(\mu M\right)$

Table 3

Structure and cytotoxic activity of PBTs compounds (47-52) used in model building.



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${f R}^2$	0.75	0.8	0.71	0.72	0.77	0.81	0.81	0.79	0.81	0.73
q²	0.52	0.53	0.72	0.72	0.56	0.51	0.58	0.54	0.59	0.55
Number of Descriptors	15	20	10	15	10	20	15	15	20	15
Training Sets	44	41	40	40	38	38	38	38	34	32
Test Sets	8	11	12	12	14	14	14	14	18	20
Models	1	2	3	4	5	9	7	8	6	10

Table 5Structure and cytotoxic activity for compounds (53–56) in the external set



55

Compound



Compound

56







R

57



Compound

58





Compound

R

Actual Actual Predic Activity Activity Activi (EC ₅₀ , μM) (-logEC ₅₀) (-logE	10.9 4.96 5.86
ChemDiv ID #	1661-1313
Structure	H Comput Aided Mol Des. Author manuscript; availablem PC 2009 September

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Table 6 acture and cytotoxic activity for the 10 computational hits (59–68) from ChemDiv database.

Predicted Activity (-logEC₅₀) 4.52 4.99 **NIH-PA Author Manuscript** Actual Activity (-logEC₅₀) N/A4.71 Actual Activity (EC₅₀, µM) 19.3 N/AChemDiv ID # 2188-3298 3253-1073 **NIH-PA Author Manuscript** HO тż Structure **NIH-PA Author Manuscript** Ω J Comput Aided Mol Des. Author manuscript; available in PMC 2009 September 30.



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Predicted Activity (-logEC₅₀) 5.76 Actual Activity (-logEC₅₀) N/AActual Activity (EC₅₀, µM) N/AChemDiv ID # 3570-0022 Structure J Comput Aided Mol Des. Author manuscript; vailable in PMC 2009 September 30.

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Predicted Activity (-logEC₅₀) 5.83 Actual Activity (-logEC₅₀) 5.10Actual Activity (EC₅₀, µM) 7.9 ChemDiv ID # 4106-0061 Structure Н thor manuscript; available in PMC 2009 September 30. tt Aided Mol Des.

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Structure

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Actual Activity (EC₅₀, µM)

> ChemDiv ID #

11.1



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Zh	nig et al.
Predicted Activity (-logEC ₅₀)	4.97
Actual Activity (-logEC ₅₀)	4.76
Actual Activity (EC _{50, μ} M)	17.3
ChemDiv DD #	C614-0150
Structure	Point Aided Mol Des. Author manuscript; available in PMC 2009 September 30.

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Predicted Activity (-logEC₅₀) 5.01Actual Activity (-logEC₅₀) 4.14 Actual Activity (EC₅₀, µM) 72.4 ChemDiv ID # K901-0689 ဂ် Structure manuscript; available in PMC 2009 September 30. Aided Mol De С

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Actual Activity (-logEC₅₀)

Actual Activity (EC₅₀, μM)

ChemDiv ID #

Structure



5.21

5.74

1.8

K915-0700

