BINARY QSAR: A NEW METHOD FOR THE DETERMINATION OF QUANTITATIVE STRUCTURE ACTIVITY RELATIONSHIPS

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A new method (particularly suited to the analysis of High Throughput Screening data) is presented for the determination of quantitative structure activity relationships. The method, termed "Binary QSAR," accepts binary activity measurements (e.g., pass/fail or active/inactive) and molecular descriptor vectors as input. A Bayesian inference technique is used to predict whether or not a new compound will be active or inactive. Experiments were conducted on a data set of 1947 molecules. The results show that the method exhibits high accuracy and is robust to measurement errors.

1 Introduction

The automation of physical experiments through robotics to effectively perform hundreds of thousands or millions of experiments in a short time has opened the door to a large-scale brute-force approach to drug discovery. This approach is generally called High Throughput Screening (HTS). The motivation behind this approach is to reduce, and possibly eliminate, time-consuming and costly manual interventions by physically synthesizing and testing a very large number of compounds. This HTS brute-force ideal can, perhaps, be realized when a few million compounds need to be tested; however, two factors will likely interfere with the HTS ideal:

- The number of possible ligands. The number of stable drug candidates is not known. Estimates vary widely; however, the lowest estimate is that there are at least ten trillion reasonable candidates. Even if one million candidates can be tested per day an exhaustive test of all candidates would require ten million days, or more than 27,300 years. A throughput rate of one billion measurements per day would require over 27 years.
- The economics of HTS. The cost per HTS measurement is not negligible. The average cost of raw materials and overhead results in a rate of approximately \$5 per measurement^a. This is certainly a substantial improvement over manual measurement; however, a one-million-test-per-day rate results in a daily expenditure of \$5 million (sufficiently high to warrant an attempt at cost reduction).

^a This figure is based upon communications with High Throughput Screening system manufacturers and screening facility operators.

These two factors strongly suggest that "Brute Force HTS" will have to become "Smart HTS" rather quickly. In other words, to reduce the total number of experiments, an experiment/analysis cycle will have to be developed so that, for example, the results of an HTS run on 100,000 compounds are analyzed and used to determine the next 100,000 compounds to be tested.

It is generally accepted that the structure, composition, or physical properties of a ligand directly affect its biological activity against a target. The attempt to transform this qualitative belief into a quantitative method of activity assessment is known as the determination of *Quantitative Structure Activity Relationships* (QSAR) which began with the work of Hansch^b and further developed by others^{c,d}. Determining a QSAR generally proceeds as follows:

- 1. Define a quantitative measure of activity (e.g., the amount of ligand needed to produce an interference with the functioning of the target).
- 2. Express the ligand in some quantitative manner; that is, select a collection of numbers that characterize the ligand. These numbers are called *molecular descriptors* or, *descriptors*.
- 3. Determine a functional relationship between activity and the selected descriptors; that is, search for a mathematical function, f, that has the property that "activity = f (descriptors)" to a suitably high level of accuracy.
- 4. Use the determined activity measure, molecular descriptors and determined functional relationship to predict the activity of new candidate ligands.

Currently, QSAR techniques are applied to relatively small data sets consisting of several tens, or perhaps several hundreds, of molecules for which activity measurements are available. These activity measurements are performed manually in the laboratory and produce relatively accurate measurements (e.g., IC_{50} numbers: the concentration of ligand required to attain 50% inhibition). The most widely used method of determining the functional relationship is the statistical technique of regression or least squares.

It is natural and tempting to assume that all one needs to do is apply current QSAR methodology to the large scale data sets of HTS and provide the necessary analysis portion of the proposed HTS experiment/analysis cycle. Unfortunately,

^b Hansch,C., Fujita,T. ρ - σ - π Analysis. A Method for the Correlation of Biological Activity and Chemical Structure. *J.Am.Chem.Soc.* 1964.

^c Cramer,R.D., Patterson,D.E., Bunce,J.D. Comparative Molecular Field Analysis (CoMFA). 1. Effect of Shape on Binding of Steroids to Carrier Proteins. *J.Am.Chem.Soc.*, 1988, *110*, 5959-5967.

^d Rogers, D., Hopfinger, A.J., Application of Genetic Function Approximation to Quantitative Structure-Activity Relationships and Quantitative Structure-Property Relationships, *J. Chem. Info. Comp. Sci.*, 1994, 34.

^e Hogg,R.V, Tanis,E.A. Probability and Statistical Inference. MacMillan Publishing Company, New York, 1993.

two critical factors render the current QSAR technology practically useless for HTS:

- Precision Loss. HTS has given rise to the following trade-off: higher throughput reduces the precision of the activity measurement. Many HTS technologies report a binary condition: a candidate ligand is either "active" or "inactive." Some HTS technologies report a discrete measure, e.g., activity on a scale from 1 to 10. In either case, current QSAR technology requires a continuous activity measurement; e.g., accurate to 2 or 3 decimal places.
- 2. Potentially Significant Error Rate. Many HTS techniques have the unfortunate property that the activity measurement is error prone. The error rate is significant enough to warrant special attention since current QSAR technology is very sensitive to outliers and errors. A significant error rate will neutralize the predictive capabilities of current QSAR technology.

Consider a simple example. Suppose that activity y is linearly related to a single descriptor x; that is, y = mx + b. A conventional data set would consist of n observations (y_i,x_i) . Without loss of generality we may assume that m > 0, the x_i have mean 0 and variance 1 and that activity is indicated by the condition that y < 0 (i.e., when x < -b/m). The linear regression estimates for m and b are

$$\hat{m} = \frac{1}{n} \sum_{i=1}^{n} y_i x_i, \quad \hat{b} = \overline{y}, \quad \overline{y} = \frac{1}{n} \sum_{i=1}^{n} y_i$$
 (1)

When presented with binary measurements (1 is active, 0 is inactive) representing the condition that $y_i < 0$ the linear regression estimates become

$$\hat{m} = \frac{1}{n} \sum_{x_i < -b/m} x_i, \quad \hat{b} = \frac{a}{n}$$
 (2)

where a is the number of active compounds. These estimates are completely different than those obtained from non-binary input (e.g., the b estimate is always in [0,1] for binary data). For example, the descriptor value at the boundary between active and inactive is

$$x = \frac{-1}{\sum_{x_i < -b/m} x_i / a} \tag{3}$$

i.e., inversely proportional to the mean active descriptor value. Contrast this with -b/m, the true descriptor value at the boundary. Fundamentally, the trouble stems from the fact that the assumptions of linear regression are not satisfied with binary HTS data.

The problems do not lie with the concepts of QSAR itself but with the underlying mathematical techniques used to determine the functional relationship between structure and biological activity. Indeed, the fundamentals of QSAR are a promising avenue for HTS data analysis.

In this article, we present a new method for the determination of a QSAR from binary activity data. We will outline the method and give the results of experiments conducted to validate the method and assess its sensitivity to measurement error.

2 Methods

Let $\{(y_i, L_i)\}$ be m results of a High Throughput Screening (HTS) experiment on a common target in which y_i is either 0 or 1 (either "inactive" or "active") and L_i is a molecule. In other words, the output of the HTS for each molecule is *binary*. Note that continuous activity data can be made binary using a threshold criterion. We will assume that there is a mapping, D, from molecules to real n-vectors that corresponds to the calculation of a set of n molecular descriptors; let $x_i = D(L_i)$. In the text to follow Y denotes a random variable with value 0 or 1 and $X = (X_1, ..., X_n)$ denotes a random variable over n-vectors (a random molecule).

Our fundamental method uses the conditional distribution Pr(Y|X) in order to determine the probability that a new molecule L is active with Pr(Y=1|X=D(L)). Let a denote the prior probability Pr(Y=1) and define f(x,y) = Pr(X=x|Y=y). We can now use Bayes' Theorem^f and write

$$p(x) = \Pr(Y = 1 \mid X = x) = \frac{f(x,1)a}{f(x,1)a + f(x,0)(1-a)}$$
(4)

and upon rearranging we have

$$p(x) = \left[1 + \frac{f(x,0)}{f(x,1)} \frac{1-a}{a} \right]^{-1}.$$
 (5)

Without loss of generality we can assume that each descriptor has mean 0 and variance 1. Suppose that the individual molecular descriptors $\{X_i\}$ are mutually independent. In this case we have

$$p(x) = \left[1 + \frac{1 - a}{a} \prod_{j=1}^{n} \frac{f_j(x_j, 0)}{f_j(x_j, 1)}\right]^{-1}.$$
 (6)

where $f_j(x,y) = \Pr(X_j = x | Y = y)$. Thus, we must estimate distributions of the form $\Pr(X_j = x | Y = y)$ and the prior probability a.

The random variable, Y, takes on the values 0 or 1. Now, the maximum likelihood estimate for a is S/m where $S = y_1 + ... + y_m$. This is an unbiased estimate with the smallest possible variance over all unbiased estimators; however, for small sample sizes, or samples in which a single value is observed it is possible to have an estimate of a = 0. This can be catastrophic for the above formulae which require likelihood ratios. For this reason we use the biased Bayes estimate under a uniform

^f Feller, W. An Introduction to Probability Theory and its Applications, Vol.1, Wiley & Sons Inc., New York, 1950.

prior which is a = (S+1)/(m+2) which is always in (0,1) and is well defined when m is zero. Let m_1 be the number of active molecules in the data set and $m_0 = m - m_1$. We now have

$$p(x) \approx \left[1 + \frac{m_0 + 1}{m_1 + 1} \prod_{j=1}^{n} \frac{f_j(x_j, 0)}{f_j(x_j, 1)} \right]^{-1}.$$
 (7)

Let Z be a random variable over the reals and let f(z) be the probability density for Z. We can estimate f by accumulating a histogram of observed sample values on a set of B bins $(b_0,b_1],...,(b_{B-1},b_B]$ defined by B+1 numbers $b_k < b_{k+1}$, b_0 is minus infinity and b_B is infinity. The usual procedure for counting the number of observations among m samples in bin k>0 is

$$B_k = \sum_{i=1}^m \delta(z_i \in (b_{k-1}, b_k]) = \sum_{i=1}^m \int_{b_{k-1}}^{b_k} \delta(x - z_i) dx$$
 (8)

which has unfortunate sensitivity to the selection of bin boundaries since observations close to the boundary between two bins are treated as if they were in the middle of one of the bins. To reduce the sensitivity to the bin boundaries we replace the delta function observation density with a Gaussian with variance σ^2 (this can be interpreted as an observation error as well as a smoothing parameter). We now have

$$B_{k} = \sum_{i=1}^{m} \int_{b_{k-1}}^{b_{k}} \frac{1}{\sigma\sqrt{2\pi}} \exp\left[-\frac{1}{2} \frac{(x-z_{i})^{2}}{\sigma^{2}}\right] dx$$

$$= \frac{1}{2} \sum_{i=1}^{m} \left[\operatorname{erf}\left(\frac{b_{k}-z_{i}}{\sigma\sqrt{2}}\right) - \operatorname{erf}\left(\frac{b_{k-1}-z_{i}}{\sigma\sqrt{2}}\right) \right]$$
(9)

which is more efficiently calculated with

$$B_k = E_k - E_{k-1}, \quad E_k = \frac{1}{2} \sum_{i=1}^m \text{erf} \left(\frac{b_k - z_i}{\sigma \sqrt{2}} \right).$$
 (10)

Even with this smoothing, it may happen that some bins are essentially 0. We therefore construct the final distributions by adding a constant to each bin before normalizing (similar to the Bayes estimate of a). Thus, the discrete density for Z is estimated with

$$\Pr(z \in (b_{k-1}, b_k]) \approx \frac{B_k + 1/c}{c + B/c}, \quad c = \sum_{l=1}^{B} B_l$$
 (11)

Finally, to evaluate the density f at a particular point, z, we calculate

$$f(z) \approx \hat{f}(z) = \sum_{k=1}^{B} \frac{B_k + 1/c}{c + B/c} \int_{b_k - 1}^{b_k} \frac{1}{\sigma\sqrt{2\pi}} \exp\left\{-\frac{(x - z)^2}{2\sigma^2}\right\} dx$$
 (12)

which simplifies to

$$\hat{f}(z) = \frac{1}{2} \sum_{k=1}^{B} \frac{B_k + 1/c}{c + B/c} \left[\operatorname{erf} \left(\frac{b_k - z}{\sigma \sqrt{2}} \right) - \operatorname{erf} \left(\frac{b_{k-1} - z}{\sigma \sqrt{2}} \right) \right]$$
(13)

We can thus model each of the descriptor distributions $f_j(x,y)$ for y in $\{0,1\}$ and j in $\{1,...,n\}$; that is, for each descriptor, we estimate two distributions: one for the active molecules in the training set and one for the inactive molecules. We thus arrive at the estimate

$$p(x) \approx \left[1 + \frac{m_0 + 1}{m_1 + 1} \prod_{j=1}^{n} \frac{\hat{f}_j(x_j, 0)}{\hat{f}_j(x_j, 1)} \right]^{-1}.$$
 (14)

The above considerations suggest the following computational procedure:

- 1. For each molecule in the experimental data set, compute $d_i = D(L_i)$.
- 2. Perform a principal component analysis^g to produce a matrix Q and a vector u such that the covariance matrix of $\{x_i = Q(d_i u)\}$ is the identity matrix.
- 3. Estimate the parameters of the probability model, p, from $\{(y_i, x_i)\}$.
- 4. The probability that a new molecule L is active is then estimated as p(Q(D(L)-u)).

3 Results and Discussion

A set of 1,947 small molecules with molar refractivity data were chosen to test the correctness and robustness to measurement error of the method. The range of molecular weights in the data set was (28,1609) with a mean weight of 312 and a standard deviation of 131. The molar refractivity values were in the range (0.3,30) with a mean of 8.3 and a standard deviation of 3.27.

An inactivity criterion of $y_i > y_0$ was used to create a binary experimental value; that is, if the molar refractivity was greater than a pre-set threshold then the molecule was considered inactive. Several values of y_0 were chosen which resulted in different divisions of the data set into active and inactive (between 51% and 97.5%). Table 1 gives the threshold values used in the experiments and the percentage of molecules in the data set that were, as a result, classified as inactive.

^g Glen, W.D., Dunn, W.J., Scott, R.D. Principal Components Analysis and Partial Least Squares Regression. *Tetrahedron Comput. Methodol.*, 1989, 2, 349-376.

y_0	$% > y_0$	y_0	$% > y_0$
2.650	97.48	5.575	80.28
2.975	97.02	5.900	77.04
3.300	96.46	6.225	73.70
3.625	95.07	6.550	70.31
3.950	93.07	6.875	67.13
4.275	90.24	7.200	63.79
4.600	87.78	7.525	60.25
4.925	85.82	7.850	56.09
5.250	83.05	8.175	51.52

Table 1. Proportion of inactive compounds as a function of the threshold.

Four molecular descriptors were chosen to represent the molecules: two zero'th order and two first order connectivity indices^h defined as follows:

$$\chi_0 = \sum_i d_i^{-1/2} \qquad \chi_1 = \sum_{ij} (d_i d_j)^{-1/2}
\chi_0^{\nu} = \sum_i s_i^{-1/2} \qquad \chi_1^{\nu} = \sum_{ij} (s_i s_j)^{-1/2}$$
(15)

where d is the number of heavy atoms connected to atom i and s = (v-h) / (Z-v-1), where Z is the atomic number, h is the number of attached hydrogens, and v is the number of valence electrons. The sums over ij denote summation over all heavy atom bonds i-j.

The descriptors were calculated using the 1998.03 version of the MOEⁱ software from Chemical Computing Group Inc (the descriptor codes used were chi0, chi1, chi0v, chi1v). The prediction method was implemented using the SVL programming language built into MOE.

For each threshold value y_0 a predictive model was estimated (with Gaussian smoothing parameter $\sigma = 0.25$) and evaluated against the data set. Estimation of model parameters required approximately 2 seconds of CPU time for each threshold value.

Performance was measured as follows. Let m_0 be the number of inactives in the data set and m_1 the number of actives. Let c_0 be the number of inactives correctly labeled by the model and let c_1 be the number of actives correctly labeled by the model. Performance is measured with the following three percentages: $a = 100(c_0+c_1)/(m_0+m_1)$ the percentage of correctly predicted activity values over all of the data, $a_0=100c_0/m_0$ the percentage of correctly predicted activity values over the inactive molecules only, and $a_1=100c_1/m_1$ the percentage of correctly predicted activity values over the active molecules only. Figure 1 summarizes these results.

^h Bicerano, J., *Prediction of Polymer Properties*, Marcel Decker Publishing, New York, 1996.

ⁱ Consult http://www.chemcomp.com for more information.

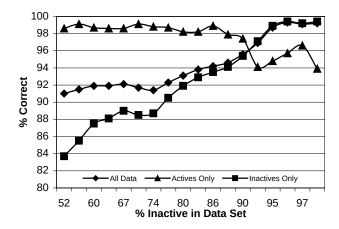


Figure 1. Proportion of inactive compounds as a function of the threshold.

At all threshold values y_i , accuracy is extremely significant (statistically) and well above what is expected from random assignments using the percentage of actives in the data set. Overall accuracy was between 91% and 99.5%. More importantly, the accuracy on the active subset was between 93% and 99.5%. Accuracy in the inactive subset was between 83.5% and 99.5%. It should be noted that the drop in accuracy in the active subset (in the right-most portion of the graph) was probably due to the fact that very few molecules were used to estimate the active distributions. This small sample fall-off would likely be eliminated with a larger data set. When the number of actives in the data set was below 10% the model performed very well with all accuracy measures above 90% correct.

To measure the sensitivity of the method to errors in the measured activity values, the following procedure was used. To each refractivity value y_i a uniform random error was introduced in the range [-ws,ws] where s is the standard deviation of the refractivity values and w is a pre-set error scale factor. This resulted in an activity criterion of $y_i + e_i > y_0$ where e_i is uniform in [-ws,ws]. The data set was trained on the modified data and accuracy measured against the actual criterion $y_i > y_0$. Three values of w were used: 0.1, 0.5 and 1.0. Each value of w was used 10 times for each value of y_0 and the accuracy results averaged. Figure 2 presents the accuracy results for the entire data set.

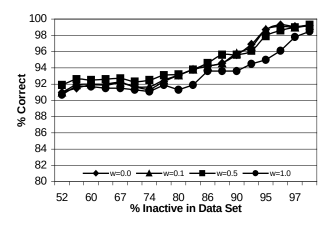


Figure 2. Accuracy results for entire data set with uniform errors.

At all error widths of w overall accuracy is maintained between 90% and 99.5%. Figure 3 presents the accuracy results on the active subset of the data for each of the w values.

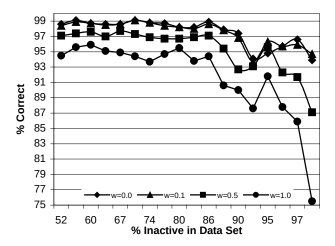


Figure 3. Accuracy results for actives only with uniform errors.

High statistical significance was maintained at all error rates and reasonable accuracy was maintained. The active-only subset showed some accuracy fall-off at very large error widths in the small sample region of the plot (lower right). This was probably due to the small number of actives that were used to estimate the

probability distributions. Despite this fall-off, it is clear that the method is capable of withstanding very high levels of measurement error in the data set.

Another form of error generation was tested. Each activity measurement was randomly inverted with probability p; that is, active and inactive was reversed for randomly selected data points. Error rate values of 5% and 10% were used. Each error rate was used 10 times and the accuracy results averaged. Figure 4 presents the overall accuracy rates for each of the error rates.

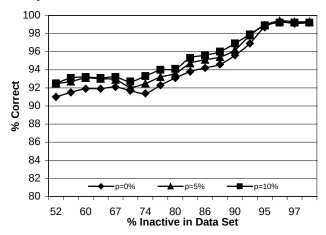


Figure 4. Accuracy results for entire data set with inversion errors.

At both error rates the accuracy on the entire data set was maintained and fell between 92% and 99.5%. It is interesting to note that the overall accuracy was better when some errors are introduced in the cases where the number of actives was below 10%. This suggests that it might be possible to improve the overall accuracy by data smoothing techniques (e.g., addition of "constructive noise") that directly model sources of error; i.e., one can take into account knowledge that there is a 5% error rate.

Figure 5 presents the accuracy results of the same experiments on the active subset.

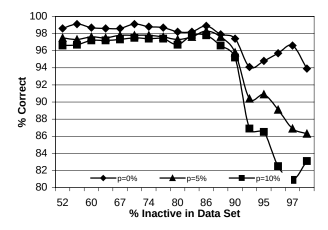


Figure 5. Accuracy results for actives only with inversion errors.

High significance is maintained at both error rates with accuracy on the active subset falling between 81% and 98%. As with the previous results, the observed accuracy fall-off was likely due to small sample effects. It must be emphasized that this fall-off is rather minor and very high statistical significance is maintained even with the small sample size.

All calculations were performed with MOE 1998.03 on a Compaq Pentium 133 with 32Mb of memory running Windows95.

4 CONCLUSIONS

A new method for QSAR analysis called Binary QSAR was presented along with results that strongly suggest the method's accuracy and robustness to measurement error. Future work will attempt to characterize the effect of the smoothing parameter upon accuracy. In the present formulation the smoothing parameter is a free parameter. As such, it can be optimized for a particular data set with a cross-validation procedure. It may be possible to eliminate the smoothing parameter by making it a function of, perhaps, the bin widths or the number of data set points.

A notable drawback of Binary QSAR is that there is no obvious way to determine the relative importance of the descriptors. In linear regression analysis, for example, each descriptor is assigned a single coefficient that can be used to estimate descriptor relevance. On the other hand, since Binary QSAR is a non-linear modeling method there is no analogous coefficient. Assessing the relative importance of the descriptors will be the subject of future work.

Binary QSAR is a fundamental shift away from the empirically fitted functional relationship methods of traditional QSAR methodology. Rather than fitting the

parameters of a model to experimental data, Binary QSAR builds predictive binary models through the use of large-scale probabilistic and statistical inference. Because data fitting is not used, the predictive capacity of Binary QSAR is not interpolative, but based on generalizations substantiated by the experimental data.

Binary QSAR has several important and immediate applications:

- Prioritization of HTS Experiments. Rather than test, for example, 5,000,000 compounds in a single run, break up the set of 5,000,000 compounds into lots of, say, 50,000 compounds. Binary QSAR could then be used to estimate the number of active compounds, or hits, in lots that have not been tested. In this way hits are found earlier and subsequent HTS experiments are more focused. Each HTS experiment proceeds from maximal diversity in the tested compounds to minimal diversity focusing on the discovered hits.
- Combinatorial Library Design. It is often the case that combinatorial
 chemistry techniques are used to create candidates for HTS experiments.
 Current combinatorial library design methods focus on maximizing the
 diversity of the resulting collection of compounds. Using Binary QSAR
 facilitates the design of more focused combinatorial libraries: the data from an
 HTS experiment is used to bias the combinatorial library towards diverse, but
 active, compounds.
- Virtual Screening and Virtual Synthesis. Once a Binary QSAR analysis is
 performed on HTS results, the resulting data model is used to search for other
 active compounds in corporate or supplier databases or even reaction pathways.

Chemical Computing Group has sought patent protection for the Binary QSAR methodology. Binary QSAR is available in Chemical Computing Group's MOE software as QuaSAR-BinaryTM and may be licensed from Chemical Computing Group Inc.