A Model-Based Ensembling Approach for Developing QSARs

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Abstract
Ensemble methods have become popular for QSAR modeling, but most studies have assumed balanced data consisting of approximately equal numbers of active and inactive compounds. Cheminformatics data is often far from being balanced. We extend the application of ensemble methods to include cases of imbalance of class membership and to more adequately assess model output. Based on the extension, we propose an ensemble method called MBEnsemble that automatically determines the appropriate tuning parameters to provide reliable predictions and maximize the F-measure. Results from multiple datasets demonstrate that the proposed ensemble technique works well on imbalanced data.

Keywords
Ensemble; Imbalanced Data; F-measure; Majority Vote; Probability Averaging and Threshold

Introduction
Compounds with similar chemical structure often have similar biology activity1. The goal of quantitative structure-activity relationship (QSAR) modeling is to determine whether chemical structures are quantitatively correlated with biology activities. A QSAR study incorporates statistical and mathematical approaches and uses computer-based tools to implement those approaches. Over the last decade, many QSAR modeling tools have been developed. Some popular examples include Decision Tree (DT)2, K-Nearest Neighbors (KNN)3, Support Vector Machines (SVM)4, Neural Networks (NNet)4,5 and Random Forest (RF)6,7. All of these methods are reputed for their application in QSAR modeling. However, none of them fully address all “practical” features required by QSAR modeling, including the ability to effectively handle imbalanced data and multiple mechanisms.

Typically, binary designations are used to indicate presence of the studied biological activity. Each compound is assigned a value of one or zero, with one indicating desired activity and zero indicating no or little activity. When compound collections have unequal numbers of active and inactive compounds, the resulting data on activity and structural descriptors is said to be imbalanced. In the real world, biologically active compounds tend to be rare when compared with inactive compounds, so the chemical data set is usually highly imbalanced. High-throughput screening data submitted to PubChem8 by the Molecular Libraries Screening Centers Network9 tend to have activity rates much less than 0.1%.

QSAR modeling is complicated by the imbalanced feature of the data. It is more difficult to predict active than to predict inactive no matter which QSAR model is utilized. For extremely
imbalanced data, the activity rate is so low that some methods may predict inactive status for the entire collection. Although the resulting accuracy rate is high, we miss all compounds that are truly active. Since one of the major goals of a QSAR study is to identify chemical structures that lead to active chemical reaction, QSAR modeling needs the ability to handle imbalanced data.

Significantly different chemical structures may cause the same biological activity. In other words, multiple mechanisms can lead to the same biological response, and therefore correct prediction depends on the ability to distinguish multiple regions of activity amidst an overwhelming excess of inactive structures. It is difficult to detect all the mechanisms and most conventional methods may only recognize some of them. When using a multiple linear regression model, for example, we may detect at most one mechanism. In general, the number of mechanisms is unknown and many popular QSAR methods fail because of this uncertainty. DT, for example, may ignore some active structures due to using a single descriptor as the splitting variable. With these complexities induced by multiple mechanisms, the ability to detect multiple mechanisms is crucial for QSAR modeling.

No single modeling approach has been shown to be optimal for all QSAR studies. Moreover, some modeling approaches have been shown to be highly sensitive to small perturbations in their training data. For this reason, the method of ensembling has gained popularity in recent years. The method of ensembling aggregates results from several individual models in an attempt to achieve substantial improvement over all individual models. Ensemble models can be designed in many different ways. Dietterich points out that the performance of ensembles depends critically on three factors.

In this paper, we focus on the “family ensemble”. Our choice of the word family indicates that all individual models used as input of the ensemble come from a common ancestor, i.e., a common data-mining algorithm. There are three factors for implementing a family ensemble: (a) the base learner, which is the data-mining algorithm used to create all individual models for the ensemble – examples are DT and KNN; (b) the selection of training datasets – the goal is to create an ensemble whose individual input models are as diverse as possible; and (c) the strategy for combining results from all individual input models, including specifying weights on the results of all base learners, e.g., the strategy of majority voting assigns equal weights to each learner. RF belongs to the class of family ensembles. It uses DT as the base learner, bootstrapping to select the training set, and makes ensemble prediction by majority vote.

A recent study by Bruce et al. compared many family ensembles for their effectiveness on balanced biological activity data. They concluded that several family ensembles are more accurate than individual methods, but a single decision tree remains competitive. Because imbalance is a common feature of QSAR datasets, we believe it is important to study the performance of family ensembles on imbalanced datasets. Moreover, we also believe that well-constructed family ensembles, which use an “unstable” base learner (able to obtain very different results from slightly different subsets of the training dataset) and employ a flexible method to aggregate the results, can outperform the individual method on imbalanced data. Our goal in this paper is to propose a model-based ensembling method that provides good prediction on imbalanced data. Additionally, we motivate use of the F-measure as an appropriate assessment criterion for QSAR studies.

The construction for model-based ensembling is described in the Methods section. We investigate the proposed ensemble method by studying the datasets used by Bruce et al. plus two additional datasets obtained from the Molecular Libraries Screening Center Network through PubChem. Results for these datasets and discussions comparing performance are
presented in the Results and Discussion section. Finally, in the Summary section we review
the advantages and limitations of the model-based ensembling method.

Methods

The proposed method, model-based ensembling (MBEnsemble), is developed for QSAR
classification problems that make predictions for binary designations of activity. MBEnsemble
is able to automatically adjust the decision rule determining prediction that a compound is
active according to the performance of base learners on training datasets. As will be
demonstrated later in this paper, this feature is especially suitable for imbalanced data. All
feature of MBEnsemble will be further discussed in this and subsequent subsections. To
construct a family ensemble, it is necessary to carefully decide three factors: (a) the base learner
for the ensemble; (b) the manipulation of the dataset for each learner; and (c) the scheme to
combine the results from all learners. Before introducing MBEnsemble, we first discuss some
basic concepts used to develop MBEnsemble.

Probability Averaging

There are various schemes to aggregate multiple learners. Majority vote (MV) is a common
choice, e.g. RF uses MV for classification problems. It jointly uses the learners by counting a
vote from each learner and the class that receives the largest number of votes is selected as the
final decision. Suppose there are \( m \) independent learners in the ensemble and each learner has
accuracy rate \( \theta \). The accuracy rate of the ensemble using MV is:

\[
\theta_{E,MV} = \begin{cases} 
\frac{\sum_{k=0}^{m/2} \binom{m}{k} \theta^k (1-\theta)^{m-k}}{2m} & \text{if } m \text{ is odd} \\
\frac{\sum_{k=m/2}^{m} \binom{m}{k} \theta^k (1-\theta)^{m-k} + \frac{1}{m} \left( \frac{m}{m/2} \right) \theta^{m/2} (1-\theta)^{m/2}}{2m} & \text{if } m \text{ is even}
\end{cases}
\]

While MV has received much attention, other schemes can be more effective. Using a dataset
on hand-written digit recognition, Kittler et al.\(^{13}\) compared six combination schemes: sum rule,
min rule, max rule, product rule, median rule, and MV. They concluded that the median rule
outperformed the other five combination schemes. For data with binary designations of activity,
probability averaging (PA), which averages over the predicted probabilities of being active
obtained from all \( m \) learners, is a competitive alternative to the median rule. As the mean of
probabilities, PA has benefits over the median rule and results in the following approximation
of ensemble accuracy rate:

\[
\theta_{E,PA} \approx \Pr(Z \leq z_\theta \sqrt{m}),
\]

where \( Z \) represents a random variable that follows the standard Gaussian distribution, \( \Pr(Z \leq z) \)
the cumulative distribution function and \( \Pr(Z \leq z_\theta) = \theta \).

Figure 1 displays the gain in accuracy due to PA relative to the accuracy due to MV, namely
\((\theta_{E,PA} - \theta_{E,MV})/\theta_{E,MV}\). PA can get more than 6% improvement in accuracy over MV if the
base learner is more accurate than a learner that performs random selection, i.e. \( \theta > 0.5 \). The
goal of ensembling is to achieve the best possible performance and the base learner used in the
ensemble is usually better than random guessing. Therefore, we choose PA to make decisions
in MBEnsemble.
Threshold

The above accuracy rates, \( \theta_{E, MV} \) and \( \theta_{E, PA} \), assume that the base learners are independent of each other. In practice, it is difficult to ensure independence among all base learners. To relax the restriction of independence and have a further understanding of ensemble prediction based on PA, we generalize the assumptions as follows: (1) \( V_{yi} \), the estimated probability of being active from the \( i \)th learner for the compound with true response \( Y \) (0 \( \equiv \) inactive, 1 \( \equiv \) active), follows a distribution with mean \( \mu_Y \) and standard deviation \( \sigma_Y \); (2) the correlation between \( V_{yi} \) and \( V_{yj} \) for \( i \neq j \) is \( \rho_Y \); (3) \( V_{1i} \) and \( V_{0i} \) are independent of each other; and (4) the truly activity rate is known to be \( p \). PA prediction is based on a preset threshold \( \delta \) and

\[
\bar{V} = \frac{1}{m} \sum_{i=1}^{m} (V_{1i}I(Y=1) + V_{0i}I(Y=0))
\]

where \( I(\cdot) \) is the indicator function. If \( \bar{V} > \delta \), the compound is predicted to be active; otherwise, the compound is predicted to be inactive.

When \( m \) is large (\( m = 100 \) used in this paper is typically considered large), \( \bar{V} \) approximately follows a mixture-of-normals distribution. The components of this mixture are: a normal component with mean \( \mu_1 \) and standard deviation \( \sigma_1 \sqrt{1+(m-1)p_1}/\sqrt{m} \) with weight \( p \), and a normal component with mean \( \mu_0 \) and standard deviation \( \sigma_0 \sqrt{1+(m-1)p_0}/\sqrt{m} \) with weight \( 1 - p \). The above assumptions imply the accuracy rate of the ensemble using PA as

\[
\theta_{PA} \approx p \Pr \left( Z > \frac{\delta - \mu_1}{\sigma_1} \sqrt{\frac{m}{1+(m-1)p_1}} \right) + (1 - p) \Pr \left( Z \leq \frac{\delta - \mu_0}{\sigma_0} \sqrt{\frac{m}{1+(m-1)p_0}} \right).
\]

Parameters \( \mu_Y \), \( \sigma_Y \) and \( \rho_Y \) depend on the base learner chosen for the ensemble, the scheme for selecting a training dataset for each base learner, and the richness of the data. On the other hand, the activity rate \( p \) depends only on the data. Once we decide the construction scheme of the ensemble for a given data set, then \( p \), \( \mu_Y \), \( \sigma_Y \) and \( \rho_Y \) become unchangeable and so prediction quality is controlled only by adjusting \( \delta \). In the Results and Discussion section, we will show the role of \( \delta \) and how distributions of \( V_{1i} \) and \( V_{0i} \) affect the selection of \( \delta \).

Assessment Using the F-Measure

In learning from imbalanced data, accuracy rate is an inappropriate measure of performance. There are many alternative measures for performance evaluation. Misclassification cost\(^{14} \), F-measure\(^{15} \) and Geometric Mean\(^{16, 17} \) are common choices to assess performance on imbalanced data. These measures are functions of the confusion matrix as shown in Table 1. Given the unit cost of a false negative \( (FN) \), \( c_1 \), the unit cost of a false positive \( (FP) \), \( c_0 \), and the total number of compounds, \( N \), the misclassification cost, F-measure and G-Mean (or geometric mean) can be defined respectively as:

- **Misclassification cost** = \( (c_1 FN + c_0 FP)/N \). All nonnegative values are possible, with zero being ideal.

- **G – Mean** = \( \sqrt{\frac{TP}{TP+FN}} \sqrt{\frac{TN}{TN+FP}} \) = \( \sqrt{\alpha^*a} \). Values range from zero to one, one being ideal.

- **F – measure** = \( \frac{(1+\alpha)TP}{(1+\alpha)TP+c\cdot FP+c\cdot FN} \) = \( (1+\alpha)/\left(\frac{TP+FP+c\cdot FN}{TP+c\cdot FP+\alpha \cdot FP}\right) \), where \( \alpha \geq 0 \) is set by the user. Values range from zero to one, one being ideal.
According to the above definitions, the ratio of $c_1$ to $c_0$, $c = c_1/c_0$, has an obvious influence on the use of misclassification cost. If $c$ is very large, most attention will be paid to reduce $FN$ and the misclassification cost will lose the control of $FP$ as well as its responsibility of assessment. Large $c$ may lead the algorithm to predict actives for all compounds if the goal of the algorithm is to minimize the misclassification cost. In reality, the ratio $c$ is large since the cost of misclassifying a rare active compound to be inactive is quite high. Although the 100% active prediction minimizes the misclassification cost, it does not provide valuable results for a QSAR study.

The geometric mean is a popular assessment measure that is typically used outside of QSAR studies\textsuperscript{16, 17}. Based on the proportion of truly active compounds that are correctly predicted ($a^+$) and the proportion of truly inactive compounds that are correctly predicted ($a^-$), the geometric mean is high when both $a^+$ and $a^-$ are high and when the difference between $a^+$ and $a^-$ is small. Consequently, the geometric mean applies equal weights to correctly identifying actives and inactives. While this strategy is preferable to only monitoring the overall accuracy rate, it still is not entirely appropriate for QSAR goals. For QSAR studies, there is very little (likely no) interest in identifying inactive compounds, hence $a^-$ is not informative and so an assessment measure based on $a^-$ is not attractive. In fact, $TN$ in Table 1 is of little use because correctly identifying inactives is not of primary value in QSAR studies. Hence we argue that because both the misclassification cost and the geometric mean directly involve $TN$, they are less desirable for assessing QSAR model effectiveness for binary outcomes in the presence of imbalanced classes.

In the spirit of misclassification cost, the F-measure uses $\alpha$ to control the numbers of $FN$ and $FP$. When $\alpha$ approaches zero, the F-measure approaches a measure that is quite popular in the text-mining literature, namely the precision, where precision is defined as $TP/(TP + FP)$. Precision is exactly equivalent to the hit rate that is more commonly known in the QSAR community. When $\alpha$ approaches infinity, the F-measure approaches another popular measure called recall, where recall is defined as $TP/(TP + FN)$. Recall is the proportion of truly active compounds that are predicted to be active. Using the notation introduced for the geometric mean, recall is exactly equivalent to $a^+$.

The F-measure is actually a weighted harmonic mean of precision and recall, and $\alpha$ is the weight for recall. Therefore, the F-measure takes values between zero (indicating the worst performance) and one (indicating the best performance). A commonly used F-measure is $F_1$ that uses $\alpha = 1$ and has equal weight on recall and precision.

A geometric mean based on precision and recall has also been proposed\textsuperscript{17}, defined as

$$\sqrt{\frac{TP}{TP+FN} \cdot \frac{TP}{TP+FP}}$$

where values range from zero to one, one being ideal. This measure enjoys many of the benefits of the F-measure but does not allow unequal weights to be applied to precision and recall. Although we use the equal-weight version of the F-measure for the remainder of this paper, we ascribe to the belief that there are studies for which unequal weights are appropriate and necessary, and hence we find value in the F-measure.

To more clearly see difference and similarities between the six assessment measures accuracy ($A$), misclassification cost ($MC$), geometric mean based on accuracy rates ($G_1$), geometric mean based on precision and recall ($G_2$), the equal-weight F-measure ($F_1$), and an unequal-weight F-measure ($F_2$), consider the following two confusion matrices (A and B), both ordered as described in Table 1 with, $c_1 = 10 c_0 = 1$:

- **A**: $TP=90$, $FN=10$, $FP=20$, $TN=80$. Then $A=0.85$, $MC=0.60$, $G_1=0.85$, $G_2=0.86$, $F_1=0.86$, $F_2=0.87$. 

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Confusion matrix B is reflective of imbalance, and most would agree that the predictive model is far less effective in this case than in matrix A. However, \( G_1 \) rates these matrices equally while \( A \) assigns only marginal penalty to matrix B. Even worse, \( MC \) rates matrix B as better than matrix A. On the other hand, \( G_2, F_1, \) and \( F_2 \) all significantly penalize matrix B, albeit in varying amounts.

To provide further assistance in calibrating numerical values of the F-measure to other assessment measures, two addition confusion matrices (C and D) are considered below. These matrices fix the total number of truly active and truly inaction compounds to match the corresponding totals in confusion matrices A and B discussed above, but instead use a “random guess” approach to arbitrarily provide correct prediction for half of the true actives and correct predictions for half of the true inactives. Assessment measures on confusion matrices C and D are shown below:

\[ \begin{align*}
\text{C: } & TP=50, FN=50, FP=50, TN=50. \text{ Then } A=0.50, MC=2.75, G_1=0.50, G_2=0.50, \\
& F_1=0.50, F_2=0.50. \\
\text{D: } & TP=50, FN=50, FP=500, TN=500. \text{ Then } A=0.50, MC=0.91, G_1=0.50, G_2=0.21, \\
& F_1=0.15, F_2=0.20. 
\end{align*} \]

Again we see that \( G_2, F_1, \) and \( F_2 \) significantly penalize matrix D but in varying amounts, and their values are much worse for these random guesses than for matrices A and B that result in more true positives and fewer false negatives and false positives.

Based on features of the misclassification cost, geometric mean and F-measure, we choose \( F_1 \) as the performance assessment measure and use it as a tool to find the appropriate threshold \( \delta \). Other options for \( \alpha \) can also be used, depending on the needs of the study. Results, as presented in the Results and Discussion section, using other values of \( \alpha \) are available upon request.

**MBEnsemble**

As mentioned in the beginning of the Methods section, there are three factors determining the construction and performance of a family ensemble. For Factor (a), DT is probably the most desired learner\(^{12}\). There are several reasons for this choice: the DT algorithm is very scalable for binary classification and hence can work for very large datasets; DT has the ability to deal with collinear descriptors; DT is interpretable; and DT can give big changes in estimation as a result of small changes in the training dataset, thus leading to less correlated learners. Because of its valuable properties, Bruce et al.\(^ {12}\), Svetnik et al.\(^ {7}\) and Dietterich\(^ {11}\) all suggest using DT as the base learner for ensembling methods. Accordingly, MBEnsemble consists of 100 decision trees.

For Factor (b), 10-fold cross-validation is used (the result by Kohavi\(^ {18}\) implies that 10-fold cross-validation with decision tree works well) and only 70% of the descriptors are randomly selected for each training dataset in each fold. When it comes to selection of base learners for an ensemble, there are two major factors that greatly impact performance: the correlation between any two models in the ensemble, and the strength of individual models. It is known that the ensemble method works best if base learners are independent of each other\(^ {6, 18}\). Increase of the correlation decreases the strength of the ensemble. Increase of the strength of individual models increases the strength of the ensemble. Unfortunately, increasing the percentage of descriptors used to construct each individual tree increases both correlation and strength of individual trees. We conducted a trial on a simulated data set (with 1000 observations, 100 variables and 7.5% activity rate). Nine percentages were considered for how
many descriptors to include in training: 10%, 20%, … 90%. The results are in favor of using 70% of the descriptors to construct individual trees for the ensemble.

Alternatively, other techniques could be employed for selecting base learners (indeed, we find the techniques used in RF to be desirable). The primary focus in this paper is combining the base learners by adjusting the threshold and using PA, and these procedures are applicable no matter how one chooses to create base learners.

For Factor (c), probability averaging is implemented in MBEnsemble because of its great appeal, as mentioned earlier. When using PA, the threshold $\delta$ becomes a tuning parameter that is used to control the prediction of being active. Different $\delta$s may be needed for different data, especially for imbalanced data. Furthermore, the ideal $\delta$ relies on the properties of the base learner, e.g. $\mu_1$, $\sigma_1$, $\mu_0$ and $\sigma_0$. It is difficult to decide a good $\delta$ before the analysis. Therefore, MBEnsemble is designed to automatically choose the optimal $\delta$ enroute to its analysis.

The procedure of MBEnsemble is listed as follows:

Loop A: for $i$ in (1:10): do 10-fold cross validation

Loop B: for $j$ in (1:100): use 100 decision trees for analysis

I. randomly select 70% of descriptors from the complete data matrix to get data $D_{ij}$

II. use the $i^{th}$ fold of $D_{ij}$ as a test set and the rest of $D_{ij}$ as the training set

III. run the decision tree on the training set to obtain the model $M_{ij}$

IV. use model $M_{ij}$ to estimate the probabilities of being active for the training set $P'_{ij}$

V. use model $M_{ij}$ to estimate the probabilities of being active for the test set $P_{ij}$

End Loop B

• use the PA scheme to aggregate $P'_{ij}$ and get $\bar{P}_{ij} = \frac{1}{100} \sum_{j=1}^{100} P'_{ij}$ for the prediction on all the folds except the $j^{th}$ fold

• find the optimal threshold $\delta_i$ that maximizes the value of $F_1$ (F-measure with $\alpha = 1$) based on $\bar{P}_{ij}$

• use PA on $P_{ij}$ to obtain $\bar{P}_{i} = \frac{1}{100} \sum_{j=1}^{100} P_{ij}$ and the optimal $\delta_i$ to make prediction on the $i^{th}$ fold data

End Loop A

The inner Loop B in MBEnsemble ensures multiplicity for the ensemble because it creates 100 different DT models. Combining those DT models supports the detection of multiple mechanisms. The outer Loop A of MBEnsemble controls cross validation and selection of control parameter $\delta$. As such, the outer loop works to decrease the variance of ensemble estimation and resists overfitting the data.

A natural question concerns possible overfitting due to our search for optimal thresholds in Loop A. As explained in the MBEnsemble pseudo-algorithm, 10-fold cross validation is used inside Loop A. Consequently, 10 optimal thresholds are determined. Each threshold is the optimal choice for the subset of 90% of the compounds used to construct trees in Loop B. With
this threshold and the trees constructed in Loop B, we make predictions on the remaining 10% of the compounds that are not used in the construction of trees. Prediction on this set is fair because the set is excluded from tree construction and search for the optimal threshold. Therefore, MBEnsemble is less prone to issues with overfitting.

With MBEnsemble, we do not need to specify $\delta$ beforehand and hence “optimal” thresholds can be identified through the analysis. Such properties will benefit the analysis on imbalanced data. Empirical results shown in the next section indicate how analyses on imbalanced data profit from MBEnsemble.

Results and Discussion

Data

In this section, we will discuss results of eight small datasets from the study of Bruce et al.\textsuperscript{12} as well as two larger assays obtained from PubChem\textsuperscript{8}. The earlier discussion on the choice of $\delta$ will continue and the importance of $\delta$ will be displayed through empirical results obtained from these datasets. Moreover, MBEnsemble results will be compared with results from RF (an ensemble method based on MV) and a single DT (using 0.5 as a threshold to distinguish between active and inactive compounds).

A summary of the eight small datasets studied by Bruce et al.\textsuperscript{12} is shown in Table 2. The assay measurement for these original datasets is continuous and shows a uniform distribution. Bruce et al\textsuperscript{12} focused on balanced classification, so they created binary responses by thresholding the continuous assay response at the median. Our study focuses on classification in the presence of imbalanced class counts, so we applied thresholds other than the median. While we studied many thresholds, even those that resulted in activity rates as low as 10 percent, we only present results corresponding to a near 20\% activity rate. Due to ties in assay values at the threshold, the actual activity rates fluctuated around 20\%. Table 2 shows activity rates of the eight small datasets; only ACE and BZR do not have activity rate of 20\%.

Bruce et al.\textsuperscript{12} use two types of descriptors: 2.5D descriptors generated by Sutherland et al.\textsuperscript{20} and linear fragment descriptors. In this paper, we focus on the 2.5D descriptor set. Among the eight datasets, GPB, THER and THR are quite small. In these datasets, the number of descriptors nearly equals the number of compounds. Therefore, these three datasets have more challenges for QSAR modeling. Furthermore, GPB is believed to be the most difficult dataset among the eight datasets because it has only 66 compounds and the number of descriptor is greater than the number of compounds.

The two large assays are assay AID364 and assay AID371. Both assays are expected to experience modeling challenges. Assay AID364 is a cytotoxicity assay with 1.4\% activity rate; the data was downloaded from PubChem\textsuperscript{8} on June 4, 2006. Assay AID371 is an assay of A549 lung tumor cell growth inhibition with 8.4\% activity rate; the data was downloaded from PubChem\textsuperscript{8} on November 2, 2006. Because toxic reactions can occur in many different ways, multiple mechanisms are expected in both assays and we expect difficulty detecting all the mechanisms.

There are a large number of different sets of molecular descriptors for quantitatively representing chemical structure. Nevertheless, there is no consensus of opinion on types of input descriptors for QSAR models because a descriptor can achieve success for some targets but fail for other targets. Since the choice of descriptor is target-dependent, we report results of five types of descriptors for the two PubChem assays studied in this paper. With the descriptor generation engine of PowerMV (Liu et. al.\textsuperscript{21}), five sets of descriptors were generated for each assay: weighted Burden numbers (BN), pharmacophores fingerprints (PF), atom pairs
Results – Specification of \( \delta \)

In the previous section, we mentioned that the threshold \( \delta \) is an important tuning parameter when using PA to make predictions. To show the importance of \( \delta \), we run a pedagogic ensemble that consists of 100 decision trees with 10-fold cross validations. Predictions of the pedagogic ensemble are based on PA with a preset threshold. The major difference between the pedagogic ensemble and MBEnsemble is that MBEnsemble determines and uses the optimal threshold \( \delta_{opt} \) on the \( i^{th} \) fold data while the pedagogic ensemble uses a preset threshold on the complete dataset. The results of \( F_1 \) (F-measure with \( a = 1 \)) for the pedagogic ensemble with varying preset threshold \( \delta \) are reported in Tables 4 and 5. We actually report averages of 9 replications for the small datasets (in Table 4) and averages of 3 replications for the large assays (in Table 5). The value in bold denotes the highest \( F_1 \) that was achieved among the seven preset thresholds (\( \delta = 0, 0.1, \ldots, 0.6 \)) for the data set. Also shown are the optimal \( F_1 \) value \( F_1(\delta_{opt}) \) for this pedagogic study, as well as the threshold \( \delta_{opt} \) that achieves this optimum.

We first consider the results in Table 4. The table shows that the values of \( F_1 \) heavily depend on the choice of \( \delta \): (1) when \( \delta = 0 \), the value of \( F_1 \) is small because the number of false positives reaches its maximum, which is equal to the number of inactive compounds, and far exceeds the number of true positives, which is equal to the number of active compounds; (2) the optimal threshold \( \delta_{opt} \) resulting in the highest value of \( F_1 \) is always less than 0.5 for all eight datasets - this confirms that using 0.5 as the threshold may not provide favorable performance for imbalanced data; (3) \( \delta \) and \( F_1 \) are positively correlated if \( \delta < \delta_{opt} \), while \( \delta \) and \( F_1 \) are negatively correlated if \( \delta > \delta_{opt} \), i.e. the relationship between \( \delta \) and \( F_1 \) appears to be unimodal, thus suggesting an algorithm aimed at determining optimum \( \delta \) (such as MBEnsemble) has likelihood for success; and (4) datasets with similar activity rate \( p \) can have quite different values of \( \delta_{opt} \). The inherent features of imbalanced data and the definition of F-measure account for observations (1) – (3), but not for observation (4). Therefore, we focus our discussion on observation (4).

As mentioned in the subsection of Probability Averaging, the appropriate choice of \( \delta \) relies on distributions of \( V_{1i} \) and \( V_{0i} \). Figure 2 displays estimated densities of \( V_{1i} \) and \( V_{0i} \) as dashed and dotted curves for datasets ACE and ACHE, and illustrates how the distributions of \( V_{1i} \) and \( V_{0i} \) affect the location of \( \delta_{opt} \). Both densities of \( V_{0i} \) for ACE and ACHE have exaggerated peaks around zero and hence most predictions on truly inactive compounds are correct when \( \delta \) is far enough from zero. On the other hand, both densities of \( V_{1i} \) have two peaks. For ACE, the peak around one is much higher than the peak around zero. This allows correct predictions on the majority of truly active compounds when \( \delta \) is far enough from one and zero, and in this case the optimal \( \delta \) is \( \delta_{opt} = 0.30 \). The distribution of \( V_{1i} \) for ACHE is contrary to that for ACE. Because of the high peak of \( V_{1i} \) around zero, it is difficult for ACHE to make correct predictions on most truly active compounds when \( \delta \) is far from zero. As a result, the value of \( \delta_{opt} \) for ACHE is greater than the value of \( \delta_{opt} \) for ACE (which equals 0.07).

The impact of \( \delta \) can be even more demonstrated for a single tree. Figure 3 shows a tree obtained from a subset of the ACE dataset. By default, \( \delta \) is set to 0.5 and this results in the predicted classes shown as the number listed for each leaf (terminal node) of the tree; one indicates that compounds falling the leaf predicted as active while zero indicates prediction as inactive. The numbers shown in parentheses for each leaf are the estimated probabilities of being active. Using \( \delta_{opt} = 0.30 \) as suggested by Table 4, we clearly see that one additional leaf (probability of 0.43) would predict compounds as active, thus possibly increasing the chance of identifying additional true actives.
The importance of specifying $\delta$ is more obvious in Table 5 since AID364 and AID371 are both extremely imbalanced. For AID364, the values of $F_1$ in bold are greater than at least 147% of the values of $F_1$ using $\delta = 0.5$. For AID371, the values of $F_1$ are almost zero when we use $\delta \geq 0.3$. Also, $F_1$ is very sensitive to the value of $\delta$ for AID371 with input of some descriptors; for example, for descriptor type AP, $F_1$ is 0.207 when $\delta = 0.1$ and falls to 0.007 when $\delta = 0.2$. All these observations again indicate that using $\delta = 0.5$ may be dubious for extremely imbalanced data and the choice of $\delta$ determines prediction quality.

Results – Comparisons

By specifying a preset threshold before the analysis, the above pedagogic ensemble exhibits the role of $\delta$ for prediction on imbalanced data but is not preferred over the datadriven MBEnsemble. Table 4 shows that the optimal threshold $\delta_{opt}$ varies with different datasets despite these sets having approximately equal activity rates and Table 5 implies that the value of $\delta_{opt}$ can change when using different molecular descriptors for the same assay. Therefore, it is difficult for the pedagogic ensemble to determine a reasonable $\delta$ only based on simple data properties such as the observed activity rate, the number of compounds, and the number of descriptors.

With MBEnsemble, we do not need to determine the threshold beforehand and optimal thresholds can be automatically determined through the performance of base learners on training datasets. As a result, MBEnsemble is a better candidate as a family ensemble on imbalanced data. Moreover, its incorporation of careful cross validation, to avoid model building and assessment using the same set makes MBEnsemble resistant to overfitting. Next, we will show the comparison results of $F_1$ from MBEnsemble, Random Forest (RF: randomForest using ntree = 100, nodesize = 5 and default settings in R), a single decision tree (DT: tree using default settings in R) and random guessing as described in the subsection Assessment Using the F-Measure. Our ChemModLab website, http://eccr.stat.ncsu.edu/ChemModLab/Default.aspx, provides a computing platform for QSAR modeling based on different methods, including RF and DT discussed here. Due to heavy computing, the MBEnsemble is not yet available on the ChemModLab website.

Table 6 gives the average $F_1$ of nine replications for the eight small datasets. Using modeling approach as a four-level factor (with levels MBEnsemble, RF, DT, Random) and folds (from the 10-fold cross-validation exercise) as a second factor, an analysis of variance (ANOVA) was run with subsequent application of Tukey’s HSD to obtain multiple comparisons between modeling approaches. For each dataset, the best statistically equivalent methods are in bold. Recall that a higher value of $F_1$ implies better performance of the model. The values of $F_1$ are dependent on the dataset — ACE has the most successful performance and BZR gets the least successful performance. With “optimal thresholds”, MBEnsemble is one of the most effective methods for all eight datasets. For all of these small datasets, the difference between the $F_1$ of MBEnsemble and the minimum $F_1$ is between 10% and 65%. Even after accounting for uncertainty and variability. MBEnsemble is statistically better than RF for five of the eight datasets (BZR, COX2, DHFR, THERM, THR) and statistically equivalent to RF for the other three.

Surprisingly, RF does not gain improvement over DT for all datasets. DT has higher value of $F_1$ on four out of eight datasets, but its $F_1$ is not statistically higher than the $F_1$ of RF except for dataset THR. The last column in Table 6 gives the performance for random guessing, which is used as the baseline method for model assessment. After accounting for variation, RF is equivalent to random guessing for BZR and THR, and DT is equivalent to random guessing for GPB. These results are contrary to the results obtained by Bruce et al. They showed that ensembles(including RF) were superior methods and preferred to DT with respect to performance on balanced datasets. The datasets for Table 6 are all imbalanced. The
disagreement between the balanced datasets and imbalanced datasets implies that it is possible for both RF and DT to fail in modeling imbalanced datasets. Next, we continue assessment for the three methods and inspect the performance of RF on extremely imbalanced datasets.

Table 7 compares $F_1$ of MBEnsemble, RF and DT on the two PubChem assays, which both have relatively low activity rate. Because of the relatively large number of compounds in these assays and the low activity rates, observed $F_1$ values tend to be rather small. For example, a method that results in perfect recall ($TP=49$, $FN=0$) and a hit rate ten times better than random guessing ($FP=301$) for AID364 still results in the very low $F_1=0.25$. So while the numbers in Table 7 are much smaller than those in Table 6, it should not be assumed that all results in Table 7 are unsuccessful attempts for prediction of these assays. In fact, several models provide very effective prediction for these PubChem assays.

Table 7 shows that MBEnsemble has stable performance on those assays and achieves the best performance. While both MBEnsemble and RF aggregate the results of 100 trees, the $F_1$ of MBEnsemble is between 0.201 and 0.316 and the $F_1$ of RF can vary a lot when using different molecular descriptors on the same assay. For example, $F_1$ for RF can increase 379% when changing molecular descriptors from PF (or AP) to BN on AID364. From Table 7, MBEnsemble provides dramatic improvement over RF, except when using BN descriptors. Statistical tests show that the difference in $F_1$ between MBEnsemble and RF is not significant for BN descriptors. Moreover, similar to Table 6, RF is not necessarily better than the DT in Table 7 and is statistically equivalent to or worse than random guessing for many cases. So Table 7 confirms possibly poor results when using RF on imbalanced data as studied in the paper.

Additionally, the performance of DT on AID371 is not comparable to any of the two ensemble methods, or even to random guessing. DT in this comparison uses 0.5 as the threshold. As mentioned in the earlier sections, $\delta = 0.5$ may not be an appropriate threshold for imbalanced data. Figure 4 illustrates why the $F_1$ of DT is almost zero on AID371 as revealed through estimated probabilities of being active when the compound is truly active ($V_{ij}$). For AP and PF, all DT-estimated $V_{ij}$ are lower than 0.5 and this results in $TP = 0$ and hence $F_1 = 0$. For BN, FP and CAP, the mean of DT-estimated $V_{ij}$ is far below 0.5 and only a few “outliers” are greater than 0.5. So the value of $TP$ is much smaller than the value of $FN$ and this results in low value of $F_1$ for BN, FP and CAP.

We conclude that MBEnsemble outperforms the other studied methods for both the small datasets having approximately 20% activity rate as well as the larger PubChem assays having less than 10% activity rate.

Summary

This paper introduces an ensemble method, MBEnsemble, for building QSAR models. MBEnsemble selects a threshold based on the behaviors of its base learners on training sets, rather than using a preset threshold in the decision rule for declaring that a compound is active. Imbalanced data benefits from this ensemble approach that allows flexibility with regard to thresholds. According to eight small datasets and two larger PubChem assays, MBEnsemble is the best of the studied methods on imbalanced data, even in the presence of multiple mechanisms within the PubChem assays.

The F-measure is used as the primary measure of assessment due to its relevance for awarding methods that correctly identify actives and avoid faulty decisions. Comparisons to other assessment measures show the benefits of the F-measure for QSAR studies. This F-measure comparison shows that MBEnsemble is at least as good as, and often better than RF and DT.
MBEnsemble is not perfect. It is computationally intensive yet does not always provide statistically significant improvements over the computationally attractive DT and RF for some datasets. Nevertheless, despite the current limitations, MBEnsemble provides stable performance on imbalanced data in the presence of multiple mechanisms. Empirical results also confirmed it is not suitable to use majority voting or a preset threshold for classification and prediction in the presence of the imbalanced data studied in this paper. Clearly, therefore, MBEnsemble is a powerful tool for developing QSAR models.

More importantly, some effective and essential components implemented in MBEnsemble (e.g., determining optimal thresholds and use of probability averaging) are directly transportable to other base learners, thus allowing much broader application and potential impact. For example, the KNN algorithm can be made equally scalable as a decision tree for binary classification, but KNN has the additional benefit of flexible decision surfaces instead of the hyper-rectilinear decision surfaces implemented by decision trees. KNN could be used as the basis of an ensemble approach that incorporates probability averaging and selection of the number of neighbors according to F-measure optimization. This and other ensemble approaches will be the subject of future investigations.

Acknowledgments

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References

19. Tan, PN.; Steinbach, M.; Kumar, V. Introduction to Data Mining. Addison Wesley; Boston, MA: 2005.
Figure 1. Improvement in accuracy rate due to PA relative to MV’s accuracy rate, i.e. $\frac{\theta_{E,PA} - \theta_{E,MV}}{\theta_{E,MV}}$. The improvement depends on $m$, the number of independent learners in the ensemble, and $\theta$, the accuracy rate for base learners.
Figure 2.
The left plot shows the results obtained from dataset ACE and the right plot shows the results obtained from dataset ACHE. The dashed curve is the density plot for $V_{1i}$, the estimated probability of being active when the compound is truly active as reflected in the observed activity measurement; the dotted curve is the density plot for $V_{0i}$, the estimated probability of being active when the compound is truly inactive as reflected in the observed activity measurement; the solid curve shows the F-measure as a function of $\delta$; and the grey vertical line is a base line displaying the location of $\delta_{\text{opt}}$. 
Figure 3.
A tree constructed for dataset ACE from randomly selecting 70% of the molecular descriptors to build the tree. The first number listed for each leaf is the prediction for that node, based on a default threshold of 0.5; 1 means active, 0 means inactive. The second number (shown in parentheses) is the estimated probability of being active.
Figure 4.
Box plots of DT estimated probability of being active when the compound is truly active as reflected in the observed activity measurement ($V_{1i}$) for AID371.
Table 1

Confusion Matrix

<table>
<thead>
<tr>
<th></th>
<th>Predicted Active Class</th>
<th>Predicted Inactive Class</th>
</tr>
</thead>
<tbody>
<tr>
<td>Truly Active Class</td>
<td>TP</td>
<td>FN</td>
</tr>
<tr>
<td>Truly Inactive Class</td>
<td>FP</td>
<td>TN</td>
</tr>
</tbody>
</table>
Table 2

Summary of Datasets

<table>
<thead>
<tr>
<th>data set</th>
<th>Compound type</th>
<th>#comp</th>
<th>#actives</th>
<th>active rate</th>
<th># descriptors</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACE</td>
<td>angiotensin converting enzyme</td>
<td>114</td>
<td>23</td>
<td>16%</td>
<td>56</td>
</tr>
<tr>
<td>ACHE</td>
<td>acetyl-cholinesterase inhibitors</td>
<td>111</td>
<td>22</td>
<td>20%</td>
<td>63</td>
</tr>
<tr>
<td>BZR</td>
<td>benzodiazepine receptor</td>
<td>163</td>
<td>29</td>
<td>18%</td>
<td>75</td>
</tr>
<tr>
<td>COX2</td>
<td>cyclooxygenase-2 inhibitors</td>
<td>322</td>
<td>66</td>
<td>20%</td>
<td>74</td>
</tr>
<tr>
<td>DHFR</td>
<td>dihydrogolate reductase inhibitors</td>
<td>397</td>
<td>79</td>
<td>20%</td>
<td>70</td>
</tr>
<tr>
<td>GPB</td>
<td>glycogen phosphorylase b</td>
<td>66</td>
<td>13</td>
<td>20%</td>
<td>70</td>
</tr>
<tr>
<td>THERM</td>
<td>thermolysin inhibitors</td>
<td>76</td>
<td>15</td>
<td>20%</td>
<td>64</td>
</tr>
<tr>
<td>THR</td>
<td>thrombin inhibitors</td>
<td>88</td>
<td>18</td>
<td>20%</td>
<td>66</td>
</tr>
</tbody>
</table>

* Binary activity was obtained by thresholding the continuous assay measurement at the 84th or 80th or 82nd percentile.
**Table 3**

Summary of Assays

<table>
<thead>
<tr>
<th>Assay</th>
<th>Descriptor Type</th>
<th># descriptors</th>
</tr>
</thead>
<tbody>
<tr>
<td>BN</td>
<td>AID364</td>
<td>24</td>
</tr>
<tr>
<td>PF</td>
<td></td>
<td>121</td>
</tr>
<tr>
<td>AP</td>
<td># compounds = 3381</td>
<td></td>
</tr>
<tr>
<td>FP</td>
<td># actives = 49</td>
<td>395</td>
</tr>
<tr>
<td>CAP</td>
<td>activity rate = 1.4%</td>
<td></td>
</tr>
<tr>
<td>BN</td>
<td>AID371</td>
<td>24</td>
</tr>
<tr>
<td>PF</td>
<td># compounds = 3312</td>
<td></td>
</tr>
<tr>
<td>AP</td>
<td># actives = 278</td>
<td>382</td>
</tr>
<tr>
<td>FP</td>
<td>activity rate = 8.4%</td>
<td></td>
</tr>
<tr>
<td>CAP</td>
<td># descriptors</td>
<td>1578</td>
</tr>
<tr>
<td>BN</td>
<td></td>
<td>24</td>
</tr>
<tr>
<td>PF</td>
<td></td>
<td>119</td>
</tr>
<tr>
<td>AP</td>
<td></td>
<td>382</td>
</tr>
<tr>
<td>FP</td>
<td></td>
<td>580</td>
</tr>
<tr>
<td>CAP</td>
<td></td>
<td>1487</td>
</tr>
</tbody>
</table>
### Table 4
Average $F_1$ from Nine Replicate Runs of 10-Fold Cross Validations of Ensembling 100 DTs with PA and Varying Preset $\delta$ for the Small DataSets

<table>
<thead>
<tr>
<th>dataset</th>
<th>0</th>
<th>0.1</th>
<th>0.2</th>
<th>0.3</th>
<th>0.4</th>
<th>0.5</th>
<th>0.6</th>
<th>$\delta_{opt}$</th>
<th>$F_1(\delta_{opt})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACE</td>
<td>0.336</td>
<td>0.658</td>
<td>0.680</td>
<td><strong>0.696</strong></td>
<td>0.692</td>
<td>0.659</td>
<td>0.615</td>
<td>0.30</td>
<td>0.696</td>
</tr>
<tr>
<td>ACHE</td>
<td>0.331</td>
<td><strong>0.516</strong></td>
<td>0.505</td>
<td>0.494</td>
<td>0.468</td>
<td>0.453</td>
<td>0.384</td>
<td>0.07</td>
<td>0.522</td>
</tr>
<tr>
<td>BZR</td>
<td>0.302</td>
<td><strong>0.451</strong></td>
<td>0.445</td>
<td>0.413</td>
<td>0.362</td>
<td>0.352</td>
<td>0.298</td>
<td>0.17</td>
<td>0.465</td>
</tr>
<tr>
<td>COX2</td>
<td>0.340</td>
<td>0.517</td>
<td>0.524</td>
<td><strong>0.531</strong></td>
<td>0.503</td>
<td>0.484</td>
<td>0.427</td>
<td>0.28</td>
<td>0.540</td>
</tr>
<tr>
<td>DHFR</td>
<td>0.332</td>
<td>0.540</td>
<td>0.589</td>
<td><strong>0.607</strong></td>
<td>0.596</td>
<td>0.567</td>
<td>0.488</td>
<td>0.34</td>
<td>0.608</td>
</tr>
<tr>
<td>GPB</td>
<td>0.329</td>
<td>0.542</td>
<td><strong>0.566</strong></td>
<td>0.560</td>
<td>0.499</td>
<td>0.471</td>
<td>0.363</td>
<td>0.27</td>
<td>0.577</td>
</tr>
<tr>
<td>THERM</td>
<td>0.330</td>
<td>0.523</td>
<td><strong>0.548</strong></td>
<td>0.536</td>
<td>0.540</td>
<td>0.517</td>
<td>0.491</td>
<td>0.25</td>
<td>0.561</td>
</tr>
<tr>
<td>THR</td>
<td>0.340</td>
<td>0.444</td>
<td>0.482</td>
<td>0.504</td>
<td><strong>0.526</strong></td>
<td>0.522</td>
<td>0.451</td>
<td>0.45</td>
<td>0.535</td>
</tr>
</tbody>
</table>

**Bold**: the highest $F_1$ that was achieved among the seven preset thresholds ($\delta = 0.0, 0.1, \ldots, 0.6$)
Table 5
Average $F_1$ from Three Replicate Runs of 10-Fold Cross Validations of Ensembling 100 DTs with PA and Varying Preset $\delta$ for the Large Assays

<table>
<thead>
<tr>
<th>Assay</th>
<th>Descriptor</th>
<th>$\delta$ 0</th>
<th>$\delta$ 0.1</th>
<th>$\delta$ 0.2</th>
<th>$\delta$ 0.3</th>
<th>$\delta$ 0.4</th>
<th>$\delta$ 0.5</th>
<th>$\delta$ 0.6</th>
<th>$\delta_{opt}$</th>
<th>$F_1(\delta_{opt})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>AID364</td>
<td>BN</td>
<td>0.029</td>
<td>0.189</td>
<td>0.238</td>
<td>0.284</td>
<td>0.236</td>
<td>0.182</td>
<td>0.103</td>
<td>0.31</td>
<td>0.289</td>
</tr>
<tr>
<td></td>
<td>PF</td>
<td>0.029</td>
<td>0.199</td>
<td>0.213</td>
<td>0.262</td>
<td>0.192</td>
<td>0.114</td>
<td>0.078</td>
<td>0</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>AP</td>
<td>0.029</td>
<td>0.254</td>
<td>0.378</td>
<td>0.281</td>
<td>0.361</td>
<td>0.273</td>
<td>0.179</td>
<td>0.101</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>FP</td>
<td>0.029</td>
<td>0.270</td>
<td>0.281</td>
<td>0.261</td>
<td>0.239</td>
<td>0.191</td>
<td>0.160</td>
<td>0.17</td>
<td>0.302</td>
</tr>
<tr>
<td></td>
<td>CAP</td>
<td>0.029</td>
<td>0.241</td>
<td>0.281</td>
<td>0.239</td>
<td>0.191</td>
<td>0.160</td>
<td>0.17</td>
<td>0.302</td>
<td></td>
</tr>
</tbody>
</table>

| AID371 | BN         | 0.155      | 0.227        | 0.098        | 0.045        | 0.023        | 0.007        | 0.005        | 0.11          | 0.235           |
|        | PF         | 0.155      | 0.194        | 0.043        | 0            | 0            | 0            | 0            | 0             | 0.08            |
|        | AP         | 0.155      | 0.207        | 0.007        | 0            | 0            | 0            | 0            | 0             | 0.11            |
|        | FP         | 0.155      | 0.226        | 0.157        | 0.021        | 0.005        | 0            | 0            | 0             | 0.13            |
|        | CAP        | 0.155      | 0.241        | 0.211        | 0.045        | 0.012        | 0            | 0            | 0             | 0.13            |

**Bold**: the highest $F_1$ that was achieved among the seven preset thresholds ($\delta = 0, 0.1, \ldots, 0.6$)
Table 6
Average $F_1$ from Nine Replicate Runs of MBEnsemble, Random Forest and Decision Tree.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>MBEnsemble</th>
<th>RF</th>
<th>DT</th>
<th>Random</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACE</td>
<td>0.703</td>
<td>0.684</td>
<td>0.599</td>
<td>0.288</td>
</tr>
<tr>
<td>ACHE</td>
<td>0.486</td>
<td>0.474</td>
<td>0.442</td>
<td>0.284</td>
</tr>
<tr>
<td>BZR</td>
<td>0.392</td>
<td>(0.294)</td>
<td>0.350</td>
<td>0.262</td>
</tr>
<tr>
<td>COX2</td>
<td>0.513</td>
<td>0.446</td>
<td>0.450</td>
<td>0.291</td>
</tr>
<tr>
<td>DHFR</td>
<td>0.591</td>
<td>0.530</td>
<td>0.497</td>
<td>0.285</td>
</tr>
<tr>
<td>GPB</td>
<td>0.513</td>
<td>0.493</td>
<td>(0.310)</td>
<td>0.283</td>
</tr>
<tr>
<td>THERM</td>
<td>0.580</td>
<td>0.484</td>
<td>0.521</td>
<td>0.283</td>
</tr>
<tr>
<td>THR</td>
<td>0.499</td>
<td>(0.302)</td>
<td>0.461</td>
<td>0.290</td>
</tr>
</tbody>
</table>

**Bold:** best statistically equivalent methods after multiplicity adjustments (); statistically equivalent to random guessing.
### Table 7
Average $F_1$ from Three Replicate Runs of MBEnsemble, Random Forest and Decision Tree

<table>
<thead>
<tr>
<th>Assay</th>
<th>Descriptor</th>
<th>MBEnsemble</th>
<th>RF</th>
<th>DT</th>
</tr>
</thead>
<tbody>
<tr>
<td>AID364 Random: $F= 0.029$</td>
<td>BN</td>
<td>0.252</td>
<td>0.249</td>
<td>0.182</td>
</tr>
<tr>
<td></td>
<td>PF</td>
<td>0.223</td>
<td>(0.052)</td>
<td>0.137</td>
</tr>
<tr>
<td></td>
<td>AP</td>
<td>0.239</td>
<td>(0.052)</td>
<td>0.122</td>
</tr>
<tr>
<td></td>
<td>FP</td>
<td>0.316</td>
<td>0.173</td>
<td>0.188</td>
</tr>
<tr>
<td></td>
<td>CAP</td>
<td>0.277</td>
<td>0.115</td>
<td>0.169</td>
</tr>
<tr>
<td>AID371 Random $F= 0.144$</td>
<td>BN</td>
<td>0.211</td>
<td>0.243</td>
<td>0.030</td>
</tr>
<tr>
<td></td>
<td>PF</td>
<td>0.201</td>
<td>0.094</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>AP</td>
<td>0.218</td>
<td>(0.124)</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>FP</td>
<td>0.229</td>
<td>(0.158)</td>
<td>0.031</td>
</tr>
<tr>
<td></td>
<td>CAP</td>
<td>0.255</td>
<td>(0.122)</td>
<td>0.023</td>
</tr>
</tbody>
</table>

**Bold:** best statistically equivalent methods after multiplicity adjustments ( ): methods statistically equivalent to random guessing.