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**Published paper**


[http://dx.doi.org/10.1080/02664763.2011.580335](http://dx.doi.org/10.1080/02664763.2011.580335)
A Classification Updating Procedure Motivated by High Content Screening Data

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Abstract

The current paradigm for the identification of candidate drugs within the pharmaceutical industry typically involves the use of high throughput screens (HTS). High content screening (HCS) is the term given to the process of using an imaging platform to screen large numbers of compounds for some desirable biological activity. Classification methods have important applications in high content screening experiments where they are used to predict which compounds have the potential to be developed into new drugs. In this paper a new classification method is proposed for batches of compounds where the rule is updated sequentially using information from the classification of previous batches. This methodology accounts for the possibility that the training data are not a representative sample of the test data and that the underlying group distributions may change as new compounds are analysed. This technique is illustrated on an example data set using linear discriminant analysis, k-nearest neighbour and random forest classifiers. Random Forests are shown to be superior to the other classifiers and are further improved by the additional updating algorithm in terms of an increase in the number of true positives as well as decreasing the number of false positives.

Keywords: Classification, Updating Algorithm, High Content Screening Experiments, Batch Learning, Random Forests, Linear Discriminant Analysis, K-Nearest Neighbour
1. Introduction

The work described in this paper is motivated by the need to use high content screens to identify candidate drugs. The current paradigm for the identification of candidate drugs within the pharmaceutical industry typically involves the use of high throughput screens. A high throughput screen with automated fluorescent imaging platform allows a large number of compounds to be tested in a biological assay in order to identify any activity inhibiting or activating a biological process.

High throughput fluorescent imaging platforms have several advantages over conventional screening techniques that rely on \textit{in vitro} techniques. The most important of these advantages is that the images contain a wealth of information that can be used to define fully the effects of a compound on cells. It is for this reason that fluorescent imaging assays have been termed high content screening \cite{4}. The study analysed here involves the use of a highly automated robotic system that administer compounds to the cellular assays (each consisting of approximately 300 cells) and then takes measurements of various aspects of cell activity by taking a high content image. These images are then analysed and quantified using advanced imaging algorithms to produce a set of variables.

Classification methods are used in the analysis of high content screening data to select those compounds that have the potential to be developed into future drugs. These compounds are denoted as hits and are checked visually. Compounds that have been misclassified as hits (false positives) are denoted false hits. It is important to minimise the number of false hits because the future development of these compounds generates unnecessary additional costs. Alternatively, missing a good hit (false
negative) may result in certain compounds with the potential to be developed further being ignored. Our procedure reduces the number of false hits and thus the amount of ‘wasted effort’ in manually checking false hit images and does not reduce the number of identified true hits (i.e. does not increase the number of false negatives).

In order to sample as diverse a chemical space as possible, each high content screen may extend to a million or more individual assays [11]. This, and the fact that only a small number of compounds in a screen (<1%) have the desired biological effect means that a number of interesting statistical challenges arise when analysing data.

Traditional multi-parametric methods of classification (e.g. linear discriminant analysis) make the assumption that the data used to train the classifier are randomly sampled from the same distribution as the points to be classified in the future [8]. However, in the case of high content screening experiments the training data are from compounds used to validate the experiment. These compounds are selected because of their known biological effects (i.e. both compounds that are known to activate and not to activate the biological process of interest). Hence these data points may not be a representative sample of the data to be classified in the future.

In this paper, we propose a new method for updating classification rules for data grouped into batches (compounds are processed through the experiment in batches and the data is analysed after each batch is complete). By sequentially updating the training data and classification rule we aim to increase the predictive capability of the model as the screen progresses. The resulting final model can then be applied to all the previous data to select any hits previously misclassified. This procedure will be
tested by applying it to a high content screening using linear discriminant analysis, k-nearest neighbours and random forests. The results of classifying this way will then be compared with that of the single parameter approach, classical versions of the multivariate classifiers.

2. Example Data Set

The data is derived from a high throughput screen to identify antagonists for a G-Protein Coupled Receptor (GPCR). The GPCR class of proteins represent a major class of drug targets. The assay used here is derived from a generic assay for GPCR activation. A cell line was generated that expressed the receptor of interest and fluorescently tagged protein β-arrestin. Upon activation of the receptor, β-arrestin will associate with the receptor at the cell membrane and then drive the internalisation of the receptor into intracellular vesicles. The appearance of the fluorescent label thus appears as a punctate distribution. In the presence of an antagonist of the receptor, the receptor does not associate with β-arrestin. Under these conditions, β-arrestin is uniformly distributed throughout the cell’s cytoplasm. The assay uses an automated imaging platform to visualise the fluorescence distribution within the cells in response to the test compounds. Image analysis algorithms are then used to quantify the distribution of fluorescence as to the degree to which the fluorescence appears punctate to identify active compounds within those screened.

The cells are also counterstained with a nuclear dye identify their location. Using additional image analysis algorithms, it is possible to quantify features of the cells not related to antagonism of the receptor. These include changes in nuclear morphology,
fluorescent label intensity and cell health. In combination, the features potentially report the ability of a test compound to specifically inhibit the receptor of interest, versus non-specific effects such as toxicity.

The data collected from this experiment is contained in three batches. The first 12,285 compounds were selected because of their known properties. This pre-screen batch makes up the training data. The remaining two batches of 33,941 and 33,408 compounds (labelled A and B respectively) yield the test data. 15 variables were measured for each compound; each represents a different aspect of cellular activity. The variables are derived by averaging over the measurements of individual cells in the image.

3. Analysis of High Content Screening Experiments

3.1 Single Parameter Approach

Early approaches for identifying hits from high content screening data involved the use of a single parameter. Hits are identified as those compounds whose measurements deviate from the majority of measurements on the same plate. The current practice is to select compounds that differ from the median by $c$ standard deviations, where $c$ is a preliminary chosen constant [6]. For the data set described in Section 2 hits are detected by filtering on the Fgrain (mean fractional fluorescence within granuli compared to total cell fluorescence) parameter. In this case, an observation is considered to be a statistical outlier if it is more than three standard
deviations away from the median of the corresponding plate. However, a low Fgrain value can also occur when there are false positives so all wells selected as hits have to be manually checked by eye so that these wells can be excluded [5]. Figure 1 shows the process of selecting hits using this approach.

3.2 Multi-Parameter Approach

Recent developments in the analysis of high content screening data have focused on investigating the implementation of multivariate classifiers. Huang and Murphy [9] and Zhou et al. [17] compare classification using K-nearest neighbours, neural networks, support vector machines, Gaussian mixture models and decision trees with HCS data from location proteomics and time-lapse fluorescence microscopy respectively. Svetnik et al. [15] made a comparison of the random forest classifier, proposed by Breiman [3], with other classifiers for predicting the activities of a compound based on a quantitative description of its molecular structure. The random forest was found to have the highest accuracy amongst all of the classifiers compared. For a general review of classifiers and statistical modelling of high content screening data, see [1, 18].

To show the potential of multi-parameter analysis of high content screening data a statistical pilot study [13] considered a refined selection of compounds from a data set previously analysed using the one-parameter approach. This refined analysis enabled the removal of ‘false positives’, arising from compounds that were, for example, intrinsically fluorescent or toxic. In this way, the number of selected compounds was
reduced and therefore enabled rapid progression of the most likely candidate drugs [5].

4. Updating Classification Rules

4.1 Changing Distributions

A fundamental assumption of classical classification techniques is that the various distributions do not change over time [7]. However, in many applications (including high content screens) this assumption is unrealistic and may lead to a decline in performance of a classifier over time. The evolution of class populations in high content screens is due to compounds being analysed in batches. Each of these new batches brings with it new compounds that may have different properties to those in the training data and those in previous batches. Therefore changes to the distributions of the classes and hence the posterior distributions of class membership may be required so that classifier performance does not deteriorate [10].

One way to combat this decline is to continually update classification rules when new information is available. The new method for updating high content screen classifiers in Section 4.2 takes the problem of changing distributions into account by updating after each batch of the experiment has been analysed.
4.2 New Updating Method

Here we outline the new method for updating the classification rules, which we use in the following section to classify data from a high content screening experiment.

The methodology for the new updating algorithm is as follows. The training data is initially classified using the single parameter plus visual checking approach described in Section 3.1 and a multivariate classifier is constructed using these data. This classifier is then used to classify those compounds which were screened as part of the first batch (in our case batch A) into groups of true hits, non-hits and false hits.

The compounds identified as true hits by the classifier are examined visually by the screening expert to verify the predictions. At this stage all true hits that have been misclassified have their classification labels corrected. A new training data set is now created by combining the data from the batch 1 (the original training data) with the visually checked compounds from batch A.

This new updated training data is used to construct a new multivariate classifier for the classification of Batch B. This part of the algorithm accommodates the possibility that the training data is not representative of the test data by correcting the assumptions on underlying distributions made from the training data.

For each new batch of data the training data is updated using the previous batches until the final classifier that is constructed is the ‘best’ possible. At this stage it is
recommended that each of the batches are manually classified again to see if any true hits were misclassified during previous classification.

4.3 Algorithm

Let the pre-screen training data $L_0$ consist of data $\{(y_n, x_n), n = 1, \ldots, N\}$ where $N$ is total number of observations, the $x$’s are observations of $p$ variables and the $y$’s are the class labels (true hit, false hit, non-hit). Given $B$ batches of compounds to be classified, let $\{T_k\}, \ k = 1, \ldots, B$ be a sequence of test sets each consisting of $M_k$ observations of $p$ variables with unknown class labels. Let $\{L_k\}, \ k = 1, \ldots, B$ be a sequence of updated training sets that are created by the algorithm.

Step 1: Given the training set $L_0$, construct a classifier $\varphi_0(x, L_0)$, where given input $x$ the class membership $y$ is given by $\varphi(x, L)$.

Step 2: ($k = 1$) Classify the batch of test data $T_1$ using the classifier $\varphi_{k-1}$ to give class labels $\varphi_{k-1}(T_1, L_{k-1})$.

Step 3: Identify observations $x_i$ from the batch of test data $T_k$ such that $\varphi_{k-1}(x_i, L_{k-1}) = \text{True Hit}$.

Step 4: Check the classifications of the observations $x_i$ that were identified as true hits in step 3 and adjust any incorrect labels (this step is done by the screening expert).
**Step 5:** Construct a new training set $L_k$ consisting of the data from $L_{k-1}$ and those observations $x_i$ that were visually checked in step 4.

**Step 6:** Construct a classifier $\varphi_k$ using the new training set $L_k$.

Set $k = k+1$ and repeat steps 2-6 until $k = B$, the number of batches.

**Step 7:** Manually apply the classifier $\varphi_k$ to batches 1, ..., $B$ to identify any True Hits that have previously been misclassified.

5. Application

5.1 Classification

Linear discriminant analysis (LDA), random forests [3] and k-nearest neighbour (KNN) classifiers were applied both with and without our proposed algorithm to classify the batches of test data described in section 2 (for details of the linear discriminant analysis and k-nearest neighbours methodology see [7]). Due to the large number of compounds in the test batches it was only possible to check the classifications of those compounds that were classified as true hits. This is considered sufficient because we can compare the number of hits and false positives for each of the classifiers. The results of this analysis are compared with the single parameter approach in Table 1. Note that the results shown in Table 1 are produced by applying the final models to the two batches of data (i.e. the results of step 7 of the algorithm).
Also note that in Table 1 the true hits group has been broken down into hits and good hits. Good hits are defined as those compounds which show the greatest level of inhibition and are identified by the screening expert. The false hit group has been split into focus error, high background, over confluent, toxic, well dry, no visible image and low draq5. This allows more detailed comparisons to be made.

All of the analysis was conducted in the statistical package R [14]. The random forest classifier was implemented using the randomForest library [12] and the library class [16] was used for the k-nearest neighbour classifier. When implementing the k-nearest neighbour classifier a leave-one-out cross-validatory approach was used for selecting the value of the parameter k.

Table 1 shows the results of classifying the two batches of test data. When using the updating algorithm, the iterative stages of classification have been combined with classifications from the final models (this corresponds to step 7 of the algorithm). These results suggest that the new methodology performs better than both the single parameter approach and the traditional multiparameter approaches when using linear discriminant analysis and random forests. Neither the results of the k-nearest neighbour nor the updated k-nearest neighbour are an improvement on the single parameter approach. The single parameter, linear discriminant analysis, updated linear discriminant analysis, k-nearest neighbour, updated k-nearest neighbour, random forest and updated random forest identified 204, 196, 145, 185, 117, 279 and 148 compounds from the 2 batches respectively but 41.2%, 37.2%, 10.3%, 56.8%, 10.3%, 48.0% and 0.7% of these were found to have been misclassified. With the exception of the two k-nearest neighbour classifiers, all of the multivariate classifiers find more
true hits than the single parameter approach. The most noticeable improvement in the number of hits selected is found when using the updated random forest, using this methodology identifies 27 more hits than the single parameter approach, and with the number of false positives, the updated random forest only misclassifies one of the hit selected compounds compared to 12, 15, 73, 84, 105 and 134 for the updated k-nearest neighbour, updated linear discriminant analysis, linear discriminant analysis, single parameter, k-nearest neighbour and random forest respectively. However, the updated random forest failed to identify one good hit that was selected by the single parameter approach.

A detailed comparison of the three non-updating methods used shows that linear discriminant analysis and random forests identify more hits than the single parameter approach but the k-nearest neighbour classifier does not. Linear discriminant analysis also misclassifies fewer compounds as hits but the random forest misclassifies a greater number of compounds with those wells that are toxic and over confluent being the main problem. Additional analysis compared the compounds that were identified as hits and good hits by the four different methods. The aim of this was to see if there were any hits that had been falsely classified as non-hits by the updated random forest, in other words to find if there were any false negatives. The results of this analysis show that there is a minimum of 8 good hits and 29 hits classified as non-hits.

Steps 4 and 7 of the algorithm require the input of a screening expert to visually check some of the images. This slows down what would otherwise be an automated process. The results in Table 1 do not take this process into account as they are the
final classifications at Step 7 of the algorithm. Table 2 compares the number of true
hits found by each classifier to the number of images that were required to be checked
in order to achieve the classification for the two batches of test data. The aim of this
analysis was to look at the effort required by the expert to check images with respect
to the number of true hits found. The updated random forest identifies the most true
hits but also requires the largest number of images to be checked (59% of images
checked were found to be true hits). Conversely, linear discriminant analysis has the
largest percentage of images checked that turn out to be hits (63%) but it identifies 24
less hits that the updated random forest.

5.2 ROC Analysis

ROC analysis of the classification results was carried out to compare the sensitivity
and specificity of the different classifiers. Figure 2(a) is a ROC plot for the
classification results shown in Table 1. As the true classifications are not known,
sensitivity and specificity have been calculated using only compounds checked
visually by the screening expert.

Comparing the seven classifiers in Figure 2(a) shows that in each case the updated
classifier has higher sensitivity and specificity than the corresponding non-updated
classifier. The updated random forest has the highest sensitivity and specificity of all
the classifiers which suggest this is the best classifier of those compared.
Figure 2(b) is a ROC plot for the results shown in Table 2. Sensitivity has been calculated using the number of images that were required to be check to get the final classifications; i.e.

\[
\text{Sensitivity} = \frac{\text{Number of true positives}}{\text{Number of images checked}}
\]

Specificity was calculated using the usual method (and is the same as for Table 1). Comparing the seven classifiers in this plot shows that updated random forests and updated k-nearest neighbours have higher sensitivity and specificity than the corresponding non-updated classifier but linear discriminant analysis has higher sensitivity than updated linear discriminant analysis. Overall, linear discriminant analysis has the highest sensitivity (63%) with the updated random forest and the single parameter both having the same value (59%). However, the updated random forest identifies 24 more true hits than linear discriminant analysis and 27 more true hits than the single parameter approach.

### 5.3 Sensitivity of Batch Orderings

An important consideration with any model is that of sensitivity. In our case we are interested in assessing how sensitive the classification algorithm is to the ordering of the batches. In other words, does the algorithm produce the same classification results regardless of the order of the batches? To empirically investigate the sensitivity of the batch orderings we compared the results of classifying the compounds using the
updated random forest as the batch order changed. We divided each of the two batches of test data into two sub-batches (A1, A2, B1 and B2); we then randomly assigned an order to the sub-batches and used the updating methodology to predict the class of the compounds. This process was repeated 8 times so that the predictions made by the model for each sub-batch order could be compared.

Table 3 shows the results of investigating the sensitivity of the classification results when permuting batch orders. It can be seen that there is some variation in the predicted classifications for the different batch orders. The most noticeable difference appears to be between those orderings which start with batch A and those which start with batch B. A detailed comparison of this difference shows that when a sub-batch from batch A is the first to be classified there are more hits from batch A identified than when a sub-batch from batch B is the first to be classified. Although the results of classification vary as the batch order is permuted we suggest that the updating algorithm approximately converges to the same classifications.

6. Discussion

Before closing with an overall review and conclusions of the work presented we discuss various aspects of the work which may benefit from some further study. Although the work that has been presented has focused on application with one particular dataset from high content screening experiments we can see that there are other situations where our suggested methodology could prove to be effective.
As discussed in section 4.1, population drift may lead to a decline in performance of a classifier over time. We have attempted to combat this problem by continually updating the classifier each time new information is available; however, we suggest that a substantial drift in the populations may cause the model to change sufficiently so that misclassification occurs for the training data and the previously classified test batches. In other words, the model may change so that compounds that were previously classified correctly are now misclassified. It may be possible it identify when this occurs by checking each new model to see how well it predicts the classifications of the original data. In such cases, an alternative approach based on the method of Biernacki et al. [2] may be appropriate.

The results of investigating the sensitivity of batch ordering in Section 5.2 suggest that the model approximately converges to the same classification. However, we suggest that it may be of interest to investigate this further by considering larger numbers of batches. It is possible to optimise the ordering of the batches before they are classified by the algorithm, however, any increase in accuracy would have to be balanced against the time taken to implement it.

The single parameter approach (Section 3.1) used to identify hits in the pre-screen data uses a filter based on $c$ standard deviations from the median. In this case the value of $c$ was chosen to be three by the screening expert so that the number of images required to be checked was not too high. By increasing this number more hits may be identified and this may have an effect on the overall classification results. The sensitivity of this parameter is something that may be investigated further.
When we applied the multivariate classifiers in Section 5 we did not directly consider the incorporation of prior knowledge nor the unequal cost of misclassification. In some cases these could be specified and incorporated into the classifiers at various points in the algorithm. There would be the opportunity to vary the costs at each step in the algorithm but this has not been explored further in the current study.

In conclusion, random forests were shown to be superior to the single parameter approach and the other classical multivariate classifiers. They were further improved by the additional updating algorithm in terms of an increase in the number of true positives and a decrease in the number of false negatives.

7. Acknowledgements

The authors would like to thank members of GSI-ASTL AstraZeneca, Charnwood for access to the data set used and Chris Harbron of the Discovery Statistical Sciences group at AstraZeneca, Alderley Park for his help and advice with this work.
8. References


Table 1: Observed classifications of hit selected compounds

<table>
<thead>
<tr>
<th></th>
<th>Single Parameter Approach</th>
<th>LDA</th>
<th>Updated LDA</th>
<th>KNN</th>
<th>Updated KNN</th>
<th>Random Forest</th>
<th>Updated Random Forest</th>
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<td><strong>True Hits</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
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<td>73</td>
<td>77</td>
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<td></td>
<td>51</td>
<td>50</td>
<td>53</td>
<td>35</td>
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<td>80</td>
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<td>31</td>
<td>3</td>
<td>2</td>
<td>23</td>
<td>3</td>
<td>7</td>
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<td>21</td>
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<td>1</td>
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<td>1</td>
<td>45</td>
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<tr>
<td>Toxic</td>
<td></td>
<td>10</td>
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<td>2</td>
<td>25</td>
<td>1</td>
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<td>6</td>
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Table 2: Comparing the number of hits found to the number of images checked for different classifiers

<table>
<thead>
<tr>
<th>Classifier</th>
<th>True Hits Found</th>
<th>Images Checked</th>
<th>% Images Checked That Were Found to be True Hits</th>
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<td>196</td>
<td>63</td>
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<tr>
<td>LDA (Updated)</td>
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<td>KNN</td>
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<td>185</td>
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</tr>
<tr>
<td>KNN (Updated)</td>
<td>95</td>
<td>196</td>
<td>48</td>
</tr>
<tr>
<td>Random Forest</td>
<td>145</td>
<td>272</td>
<td>53</td>
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<tr>
<td>Random Forest (Updated)</td>
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<td>250</td>
<td>59</td>
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Table 3: Observed classifications of hit selected compounds using updated random forests with different batch orders

<table>
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<tr>
<th>Batch Order</th>
<th>Hit</th>
<th>Good Hit</th>
<th>Non-Hit</th>
<th>Focus Error</th>
<th>High Background</th>
<th>Over Confluent</th>
<th>Toxic</th>
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<tbody>
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<td>A1, B1, A2, B2</td>
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<td>1</td>
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<td>1</td>
</tr>
<tr>
<td>A1, B2, A2, B1</td>
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<td>50</td>
<td>1</td>
<td>1</td>
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<td>0</td>
<td>0</td>
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<td>49</td>
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<tr>
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Figure 1: Hit selection using a single parameter
Figure 2(a): ROC Plot showing the classification results from Table 1
Figure 2(b): ROC Plot showing the classification results from Table 2