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Abstract

High-content screening is a powerful method to discover new drugs and carry out basic biological research. Increasingly, high-content screens have come to rely on supervised machine learning (SML) to perform automatic phenotypic classification as an essential step of the analysis. However, this comes at a cost, namely, the labeled examples required to train the predictive model. Classification performance increases with the number of labeled examples, and because labeling examples demands time from an expert, the training process represents a significant time investment. Active learning strategies attempt to overcome this bottleneck by presenting the most relevant examples to the annotator, thereby achieving high accuracy while minimizing the cost of obtaining labeled data. In this article, we investigate the impact of active learning on single-cell-based phenotype recognition, using data from three large-scale RNA interference high-content screens representing diverse phenotypic profiling problems. We consider several combinations of active learning strategies and popular SML methods. Our results show that active learning significantly reduces the time cost and can be used to reveal the same phenotypic targets identified using SML. We also identify combinations of active learning strategies and SML methods which perform better than others on the phenotypic profiling problems we studied.

Keywords

High-content screening, machine learning, active learning, phenotypic discovery, multiparametric analysis

Introduction

Developments in fluorescence labeling, automation, microscopy, data storage, and image analysis have paved the way for a new era of biological research using high-content screening (HCS). In a high-content screen, vast quantities of biological data are collected and analyzed to identify small molecules, peptides, or genes that alter the phenotype of a cell. HCS is used extensively by the pharmaceutical industry throughout all stages of the drug development process.¹ It has also emerged as a powerful approach for defining protein functions and understanding signaling pathways among academic investigators, thanks in large part to recent advances in genome-wide RNA interference (RNAi) technology.²

High-content screens capture a compound's molecular and phenotypic effects on a cell through a wealth of extracted image data. In traditional high-throughput screening methods, biological responses from thousands of cells are aggregated into a few carefully selected project-specific readout parameters. In contrast, high-content screens use image analysis to extract functional and morphometric data from individual cells on a massive scale. In the course of a typical genome-wide RNAi screen, researchers acquire millions of images and then extract and store abundant quantitative information to characterize the cells, such as intensity, size, morphology,

spatial distribution, and texture for computational analysis. With so much data, the search for positive results, or “hits,” can be daunting. Early approaches to hit identification ranked all compounds by a single parameter; those above a statistically determined threshold were considered hits.³ Filtering with additional parameters can improve the process but does not address the principal drawbacks of this approach: the difficulty of identifying which combinations of parameters are useful, human bias in choosing which parameters determine the hits, and the failure to exploit all of the information acquired from the image analysis.

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Supervised machine learning (SML) has helped to address this problem by simultaneously considering all the parameters collected during image analysis and automatically sorting compounds according to predicted hit quality.^{4,5} Recent studies have shown that SML methods identify target compounds with significantly more reliability than manual parameter selection.⁶ SML also eliminates human bias introduced during parameter selection by considering every parameter for its prediction. As SML methods have become more widely used in high-content screen analysis,^{7,8} they have demonstrated the power of multiparametric readouts to distinguish between true biological activity and nonbiological interference.

With machine learning, a new bottleneck has arisen in the analysis stage of the HCS process. Even without machine learning, experimenters face the arduous task of manually sifting through oceans of data in search of the right combination of parameters to generate a valid hit list. But if SML is used for hit identification, experimenters must provide the algorithms with labeled examples to build a predictive model in what is known as the training process. Typically, several hundred to several thousand examples are necessary for the classifier to achieve satisfactory performance, and providing more examples nearly always increases performance, although at a diminishing rate. Depending on the difficulty of the problem, labeling an individual cell can take on average between 3 and 15 s; therefore, the annotation process represents a significant time investment. Fatigue further complicates the issue: because annotation quality degrades over time, the biologist is often forced to perform the labeling in multiple sittings.

In a typical annotation setting, the expert, or “oracle,” is presented with an image from the high-content screen chosen at random. The expert then decides which of the cells that appear in the image to label, evaluates them, and assigns the appropriate labels. To speed up the process, some practitioners use a classifier trained on data from previously annotated images to make predictions for every cell in a new image, and the expert is asked to correct the predictions. In the first case, human selection bias means that the labeled population is often highly correlated and not representative of the true population. But it is more concerning that both schemes unintentionally waste the expert’s effort. A significant amount of time will be spent unknowingly labeling hundreds or even thousands of cells that have little impact on the decision boundary of the learning algorithm. Recently, active learning strategies have emerged as a way to improve classification performance using less training data, thereby making more efficient use of the expert’s valuable time. They do so by allowing the learning algorithm to select the data it uses to learn through relevant queries posed to the expert.

This article explores how active learning strategies can be leveraged to make more efficient use of the expert’s time, achieving high hit identification accuracy while minimizing the cost of obtaining labeled data. In particular, we compare four well-known active learning strategies and six

popular machine learning algorithms on data taken from three large-scale RNAi high-content screens representing a diverse set of phenotypic profiling problems. Our investigation measures performance over time using implementations from the popular software packages Advanced Cell Classifier⁸ and Weka.⁹ In addition, we provide a software-agnostic evaluation measuring classification performance for every query posed to the expert. Our results show that for the HCS assays we consider, there always exists an active learning strategy that will significantly increase the rate at which the classifier learns. In some cases, the time can be reduced by over a factor of 3. Although no one active learning strategy consistently ranked the best, some performed consistently better than others. And interestingly, a few strategies actually perform worse than passive learning if training time is taken into account.

Materials and Methods

Ribosome Biogenesis Assay

The biosynthesis of proteins in cells is performed by ribosomes, macromolecular complexes that consist of a large (60S) and a small (40S) subunit. In eukaryotes, the production of ribosomes is a complex, highly compartmentalized process that begins in the nucleolus. Both ribosomal subunits undergo separate maturation in the nucleolus and nucleoplasm before they are exported to the cytoplasm where final maturation occurs. Both subunits join to form the translational competent machinery.¹⁰ To investigate 40S biogenesis, a tet-inducible RPS2-YFP (40S) HeLa cell line was generated that allows us to observe the nuclear maturation of freshly synthesized precursors of the small ribosomal subunit. This image-based assay partially relies on RPS2-YFP localization as readout. Under normal biogenesis conditions, the reporter mainly localizes to the cytoplasm, reflecting mature ribosomes. If biogenesis defects occur either in the nucleolus or the nucleoplasm, the reporter localizes to the respective cellular compartment and the cytoplasmic signal is strongly reduced.¹¹ This assay was used in a genome-wide RNAi association study using the Qiagen Human genome-wide siRNA library (HsNmV1; Qiagen, Venlo, the Netherlands) and analyzed using HCS. Example images are provided in **Figure 1a**.

Semliki Forest Virus Assay

Semliki Forest virus (SFV) is an alphavirus (Togaviridae) transmitted by mosquitoes.¹² In humans and animals, it can cause serious diseases including long-lasting arthritis and lethal encephalitis. This virus replicates very efficiently in most cell culture systems, completing its life cycle in few hours. Here we used a recombinant SFV that was genetically engineered to express a green fluorescent protein.¹³ Soon after infection, the viral RNA genome is delivered in

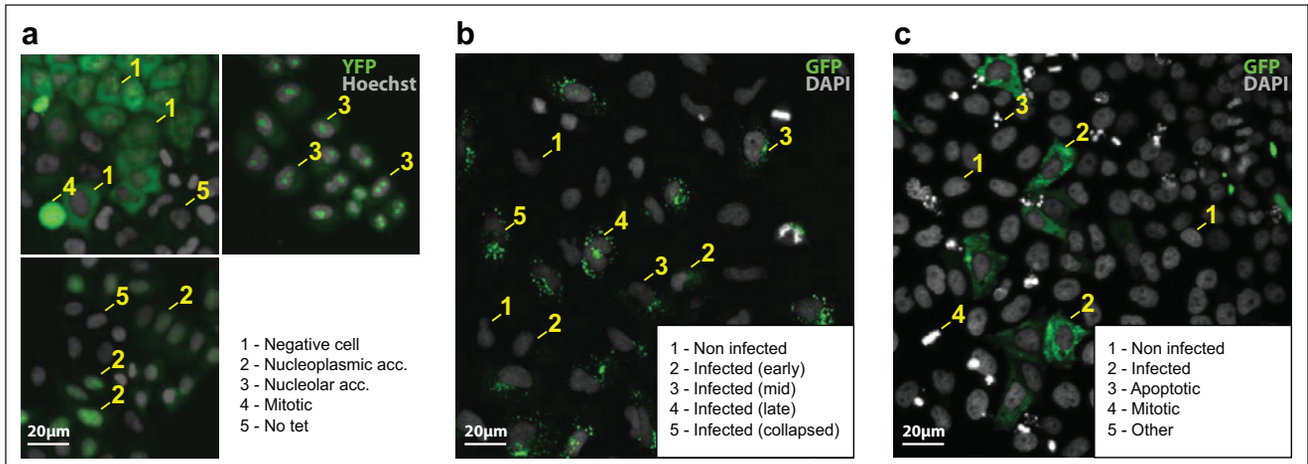


Figure 1. Examples images from the high-content assays considered in this study: (a) ribosome biogenesis, (b) Semliki Forest virus, (c) Uukuniemi virus. Annotations indicate the expert-labeled phenotype of a particular cell. A list of phenotypes is provided in the lower right of each pane. Phenotype 5—Other is not shown in (c) because it was used to collect errors in the image processing, which were extremely rare.

the cytoplasm of the cell. Here the ribosomes, cellular machineries responsible for the production of proteins, start to produce the viral proteins. One of these viral proteins was fused with green fluorescent protein. At the beginning of infection, a few small green spots appear in the cell. As the infection proceeds and more viral proteins are produced, these fluorescent spots become larger, brighter, and concentrate around the nucleus of the cell. Thus, by monitoring the intensity distribution and texture of the green fluorescence, it is possible to distinguish between the different stages of infection, as well as noninfected cells. This assay was used in a genome-wide RNAi association study using the Dharmacon Human genome-wide siRNA library and analyzed using HCS. Example images from the screen are provided in **Figure 1b** (corresponding segmentations appear in **Suppl. Fig. 1b**).

Uukuniemi Virus Assay

Uukuniemi virus (UUKV) belongs to the virus family of *Bunyaviridae* that contains more than 350 segmented, negative-stranded RNA viruses.¹⁴ As a nonpathogenic virus for humans, UUKV serves as a model for some severe human pathogenic bunyaviruses such as the Rift Valley fever virus.¹⁵ To study the host genes that are involved during the entry steps of UUKV, we conducted a genome-wide siRNA screen using the Qiagen Human genome-wide siRNA library. To monitor infected cells, we used indirect immunofluorescence against the viral nucleoprotein that is expressed in newly infected cells. After bunyaviruses enter into their host cells, their genome is released into the cytoplasm where the replication and the production of new viral proteins occur. Thus, we were able to classify infected and uninfected cells after depletion of a gene. Example images from the screen are provided in **Figure 1c**.

Microscopy

Images were acquired using two Molecular Devices (Sunnyvale, CA) ImageXpress Micro microscopes equipped with an automated plate loader and with 10x S Fluor 0.45NA objectives. The automated microscopes acquired nine sites per well arranged in a 3×3 slightly overlapping grid. Laser-based autofocusing was used to ensure that images were focused at every site. Two fluorescent channels were recorded for each high-content screen. For the ribosome biogenesis assay, a Hoechst channel was used for nuclear staining, and a YFP reporter is bound to the 40S subunit. For the SFV assay, a DAPI channel was used for nuclear staining, and a GFP channel reports the stage of viral infection. For the UUKV assay, a DAPI channel was used for nuclear staining, and a GFP channel was used to indicate if a cell was infected.

Image Analysis

The software analysis package CellProfiler¹⁶ was used to segment individual cells and extract features from images acquired for each screen. Several custom modules were used to increase the speed of the analysis. The image analysis for all three assays followed a common framework. First, the cellular nuclei were detected and segmented using Otsu adaptive thresholding and the watershed algorithm on the DAPI/Hoechst channel. Next, the nuclei were used as “seed” regions, and the cytoplasm is approximated as a ring surrounding the nucleus. From these regions denoting cellular compartments, various features were extracted for each cell describing the intensity, morphology, and texture. For the ribosome biogenesis assay, 26 features were collected, mostly intensity-based (eg, integrated intensity, mean intensity of the nucleus/cytoplasm, standard deviation of intensity of the nucleus/cytoplasm, intensity along the

Table 1. Active Learning Strategies and Supervised Learning Methods Considered in This Study.

Sampling Strategies	Machine Learning Methods
1. Passive learning	1. Naive Bayes
Uncertainty sampling	2. k -nearest neighbors
2. Least confident	3. Support vector machine
3. Margin sampling	4. LogitBoost
4. Entropy	5. Artificial neural network
Query by committee	6. Random forest
5-7. Vote entropy sampling with committee size $C = 3, 5,$ and 7	

edge of the nucleus, etc.). Texture features were also collected to describe the contrast, correlation, variance, entropy, gradient, and so forth. The UUKV assay used the same features as the ribosome biogenesis assay, with two additional features describing the nucleus intensity, for a total of 28 features. A total of 94 features were collected for the SFV assay. In addition to the features described above, several morphological features were collected such as area, perimeter, form factor, eccentricity, and so on. A full list of the extracted features is given in **Supplementary Table 2**. The texture features were extracted at multiple scales to ensure sensitivity to different sized objects. In addition, a custom implementation of an “a trous” transform¹⁷ was used to perform spot detection of viral clusters. The custom CellProfiler modules, the pipelines, and the data sets can be downloaded from www.highcontentanalysis.org.

Multiparametric Analysis Using SML

With so many features available in a high-content screen, it is too difficult to take every feature into account while manually performing hit identification. Although early works restricted themselves to just a few interpretable features, researchers have turned to machine learning to consider all the features collected during the image analysis. A few studies have made use of unsupervised learning techniques such as clustering¹⁸ and self-organizing maps,¹⁹ but the majority have opted for supervised techniques.^{4,5,7,8} SML infers a mapping function from labeled training data, which consists of a set of training example pairs (a feature vector describing the cell and the desired class label). The supervised learning algorithm uses the training data to produce a function that maps unseen instances to the desired output. This requires the supervised learning algorithm to generalize to unseen situations in a reasonable way.

In this study, we consider six popular SML methods: Naive Bayes, KNN, SVM, LogitBoost, ANN, and random forest. The first, Naive Bayes, builds a prior probability of the class distribution and uses Bayes rule to update this using kernel density estimates from features of other examples. It is well

suited for high-dimensional feature vectors and, despite its simplicity, can outperform more sophisticated methods. The second, k -nearest neighbors, or KNN, is a nonparametric instance-based learning method that predicts class membership based on the k closest training examples. KNN can be a powerful predictive model, but the training process does not scale well for large data sets. A support vector machine, or SVM, is a method for finding the best separating hyperplane for linearly separable problems. It can be extended to nonlinearly separable problems through the use of kernel functions and is one of the most popular and dependable machine learning methods. LogitBoost is an adaptation of AdaBoost set in a statistical framework that minimizes logistic loss on decision stumps. Adaboost is a meta-algorithm that trains a series of simple classifiers, each tuned to correct for mistakes made by the previous classifiers. It can be a powerful predictor for difficult problems. Artificial neural networks, or ANNs, are networks of artificial neurons that map several weighted inputs to a desired output, inspired by neurons in the brain. Random forests train thousands of decision trees on randomly chosen subsets of the feature space and make predictions by running the new instance to the terminal node of all the trees and performing some statistical measurement. For this study, we used the WEKA⁹ implementations of the above machine learning methods with parameters selected for the best performance using an extensive cross-validated random search. A summary of the supervised learning methods considered in this study is provided in **Table 1**.

The standard procedure for collecting the labeled data necessary for the SML method to learn the mapping function is sometimes referred to as *passive learning*. The expert is presented with a set of instances randomly sampled from the data and is asked to assign labels to each. Alternatively, the expert is allowed to choose which samples to label. In this study, an instance corresponds to an individual cell, the labels are nominal, and the expert is provided with an image selected at random and allowed to choose which cells to annotate. The resulting set of labeled instances, called the *training set*, is supplied to the algorithm for learning, or model training. SML algorithms require a large training set to perform well, and prediction accuracy increases as the size of the training set increases. The labeling process is strenuous and must usually be broken into multiple sittings to ensure fatigue does not compromise the quality of the data. These factors constitute a significant time investment from the experts, making the passive learning process a major bottleneck in HCS.

Active Learning Strategies

Passive learning does not make efficient use of the expert's time. There is no guarantee on the usefulness of the labels provided by the expert, regardless of whether the expert chose the cells to label or if they were selected at random. To help eliminate this bottleneck, we consider several active learning strategies to make more efficient use of the expert's

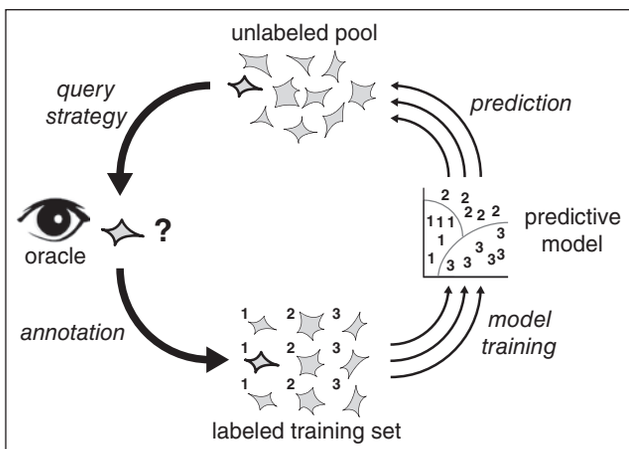


Figure 2. The active learning cycle. Starting from the top, an instance or set of instances is sampled from the unlabeled pool using a query strategy and presented as queries to the oracle (expert). Using expert knowledge, the oracle provides labels to the queries, and then the labeled instance(s) are added to the labeled training set. The labeled training set is a collection of all previously annotated labeled instances. Next, the predictive model is retrained with the updated labeled training data. After retraining, the SML algorithm makes predictions for each instance in the unlabeled pool (or a subset of it), which are then used by the query strategy to measure the informativeness of each instance when choosing the next query. In this way, the cycle iteratively improves classification performance by querying the expert with the most useful unlabeled example(s) and retraining the model.

time. Active learning is an iterative process that aims to minimize the cost of obtaining labeled data by carefully selecting the most informative examples for labeling. Consequently, active learning strategies promise to produce a more accurate predictor in significantly less time than passive learning.

Active learning can be broadly categorized into three frameworks: pool-based sampling, stream-based sampling, and membership query synthesis. Pool-based sampling is appropriate when large collections of unlabeled data can be gathered all at once. In pool-based sampling, there exists a large pool of unlabeled data U and a small set of labeled data L . Unlabeled samples are selectively drawn from the unlabeled pool according to an informativeness measure and presented to the oracle (expert) for labeling.²⁰ This process is referred to as a *query*. It is common to repeat this in an iterative cycle, as depicted in **Figure 2**. An instance or set of instances is sampled from the unlabeled pool using some selection strategy and presented as queries to the oracle. The oracle uses expert knowledge to provide labels, and then the instance(s) are added to the labeled data. Next, the predictive model is retrained with the updated labeled data. After retraining, the SML algorithm makes predictions on the unlabeled pool, which are used by the selection strategy to measure the informativeness of each instance when choosing the next query. In this way, the cycle iteratively

improves classification performance by querying the expert with the most useful unlabeled example(s) and retraining the model.

For high-content screens, pool-based sampling is typically the most appropriate framework because the assay and image analysis provide an extremely large pool of unlabeled data. But it is worthwhile to explain briefly the other frameworks, as some situations may arise in which they are also useful. Stream-based sampling is used when instances are drawn one at a time from the data source; real-time stock market data are a good example. In this case, the task of the active learning is to decide whether to query the oracle or discard the data, based on the expected usefulness.²¹ In the membership query synthesis framework, the oracle is presented with either real examples or synthetically generated examples from the input space.²² Although this approach is not immediately useful for high-content screens, progress in the creation of valid cellular generative models means that it might one day be practical.

All active learning methods must evaluate the informativeness of unlabeled instances and choose which instances to present to the oracle. How this problem is formulated is known as the *query strategy*. Many query strategies exist in the literature. Expected model change attempts to estimate which instance would impart the greatest change to the current model if its label were known.²³ Expected error reduction attempts to measure not how much the model is likely to change but how much its generalization error is likely to be reduced.²⁴ Similarly, expected variance reduction attempts to indirectly reduce generalization error by minimizing the variance of the output. In this work, we focus on the two most popular query strategies: uncertainty sampling and query by committee.

Uncertainty Sampling. The simplest query strategy, uncertainty sampling, attempts to select instances about which it is the least certain how to label. For probabilistic SML methods, it is straightforward to use the predicted class probabilities to measure uncertainty directly. One method is to query the instance whose prediction is the least confident,²⁵

$$x^* = \arg \max_x 1 - P_{\theta}(\hat{y} | x), \quad (1)$$

where x refers to an unlabeled instance, y refers to its associated label, x^* refers to the most informative instance chosen for the query, and $\hat{y} = \arg \max_y P_{\theta}(y | x)$ is the class label with the highest posterior probability predicted by the model θ . An interpretation of this query strategy is that it selects as the most informative instance the one that has most likely been mislabeled. However, a drawback to this approach is that it considers only the most probable class label, discarding information about the confidence of the remaining class labels.

To correct for this, some researchers use a query strategy that considers the first and second most probable class label, known as margin sampling,²⁶ in which the most informative instance is given by

$$x^* = \arg \min_x P_\theta(\hat{y}_1 | x) - P_\theta(\hat{y}_2 | x), \quad (2)$$

where \hat{y}_1 and \hat{y}_2 are the first and second most probable class labels predicted by the model, respectively. This strategy attempts to estimate the margin of each instance, or the distance to the decision boundary. Instances in which the model has little trouble differentiating between the first two class labels are assumed to have a large margin and are considered less informative. Instances with small margins are more ambiguous and are more likely to be useful when posed as queries to the oracle.

Margin sampling still considers only the two most probable class labels. A more general query strategy that considers all predicted class labels uses the entropy as a measure of uncertainty,²⁷

$$x^* = \arg \max_x - \sum_i P_\theta(\hat{y}_i | x) \log P_\theta(\hat{y}_i | x), \quad (3)$$

where k indexes all possible labels. In information theory, entropy is a fundamental measure of uncertainty and allows this approach to consider all the class labels predicted by the SML model θ .

Query by Committee. A more theoretically motivated query strategy first proposed by Seung et al.,²⁸ query by committee, constructs a “committee” of models $C = \{\theta^1, \dots, \theta^C\}$ that represent competing hypotheses, each trained on the same labeled set L . This query strategy selects the most informative instance by determining the instance with the strongest disagreement among the various committee members. Many variations of query by committee appear in the literature, but they all must (1) be able to generate committee models whose hypotheses are consistent with the current labeling in L and (2) be able to measure disagreement among committee members. A popular method for discriminative learning models to generate approximately valid committee members is query by bagging.²⁹ It uses bagging, a learning ensemble meta-algorithm, to construct committee members by sampling labeled instances with replacement from L and then trains models on each set of sampled data. To measure disagreement among the committee members, we can adapt the entropy-based uncertainty sampling method to votes from the committee members. Accordingly, the most informative instance determined by entropy of the committee voting is given by

$$x^* = \arg \max_x - \sum_i \frac{V(y_i)}{C} \log \frac{V(y_i)}{C}, \quad (4)$$

where $V(y_i)$ is the number of votes that class label y_i receives from the various committee members and C is the size of the committee.

Table 1 summarizes the active learning strategies considered in this study. In total, we compare seven different strategies including passive learning, three variants of uncertainty sampling, and three variants of query by committee. Query by committee using vote entropy is tested for three, five, and seven committee members. In all cases, committee members are trained using instances sampled with replacement from the labeled training set L , where $m = \frac{3}{4} |L|$ and $|\cdot|$ is the cardinality operator. Each of the six active learning strategies considered here (and passive learning) were tested in combination with the six SML methods described above.

Data Sets

We collected an expert-labeled data set for each of the assays described above. A summary is provided in **Supplementary Table 1**. The ribosome biogenesis data set contains a pool of 4492 labeled cells, each with 26 descriptive parameters (or features). The SFV data set consists of 5119 labeled cells, each with 94 features. The UUKV data set contains 500 labeled cells, each with 28 features. Each HCS assay identified five phenotypic classes, described below. In all our experiments, 66% of the data set was used for training the model (using active or passive strategies) and 33% was reserved to test the model. Annotations were performed using Advanced Cell Classifier.⁸ The data sets and their descriptions can be downloaded from www.highcontentanalysis.org.

For the ribosome biogenesis assay, cells belong to one of the following classes. Examples of each class appear in **Figure 1a**.

- *Negative*: No biogenesis defect in the cell; the reporter is localized in the cytoplasm
- *Nucleoplasmic accumulation*: A biogenesis defect occurred in the cell; the reporter is localized in the nucleoplasm
- *Nucleolar accumulation*: A biogenesis defect occurred in the cell; the reporter is localized in the nucleoli
- *Mitotic*: The cell is undergoing mitosis
- *No tet*: The reporter failed to activate in this cell

For the SFV assay, cells belong to one of the following classes. Examples of each class appear in **Figure 1b**.

- *Noninfected*: No reporter presence indicates that the cell is not infected
- *Infected (early)*: A few faint localized reporters indicate early stage of infection
- *Infected (mid)*: More numerous, larger reporters indicate mid stage of infection
- *Infected (late)*: Larger, brighter reporters (sometimes clustered) indicate the late stage of infection

- *Infected (collapsed)*: Clustered, bright reporters accompany a dramatic change in cell morphology after cell collapse.

For the UUKV assay, cells belong to one of the following classes. Examples of each class appear in **Figure 1c**.

- *Noninfected*: No reporter presence indicates that the cell is not infected
- *Infected*: The presence of the viral nucleoprotein causes the marker to be expressed in the cytoplasm
- *Apoptotic*: Fragmented morphology of the nucleus indicates programmed cell death
- *Mitotic*: The cell is undergoing mitosis
- *Other*: Used to collect cells that occasionally encountered image segmentation problems.

Performance Evaluation

The efficiency of an active learning strategy is typically measured by computing the accuracy of the classifier after every query. By plotting the accuracy against the query index, we can generate a learning curve (depicted in **Fig. 3**), and integrating the area under the learning curve (AUC) provides a useful measure of how quickly the active learning strategy improves classifier performance.²⁶ However, measuring performance this way neglects an important practical component of the evaluation, namely, the time cost to the expert. Labeling a cell and retraining the classifier requires time and effort from the expert, and this fact should be accounted for in the performance evaluation. It may well be that an active learning strategy that appears to be more efficient when measured per query is actually inefficient when we consider how much time it took to label and train.

Therefore, to provide a more practical measure of learning efficiency, we measure the performance of an active learning strategy by computing the AUC over time instead of per query. This provides a better measurement of whether we have achieved our ultimate goal of reduced training and annotation time, but it comes with a caveat. The annotation time and model training time are linked to the data, the software, and the machines used for testing. Thus, the time measurements we report are specific to the problems considered in this study, although we made an effort to choose representative assays and popular implementations of SML methods.

Because our evaluation considers the time needed to retrain the SML model, a new question arises: what is the most efficient number of queries to present to the expert before retraining the SML model and proceeding with the next iteration of the active learning cycle? To answer this, we also run tests to investigate what is the optimal number of queries per cycle for each combination of active learning strategies and SML methods on all the HCS assays.

Results and Discussion

We tested 42 combinations of SML methods and active learning strategies (six supervised learners and six active learning strategies, plus passive learning). Each combination was evaluated for $q_c = \{1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60\}$ queries per active learning cycle, for a total of 672 tests. The sheer number of individual tests required too much time from the experts; therefore, we had the experts create a shared pool of labeled data beforehand and recorded the annotation time. For each test, we then sampled two-thirds of the instances from the shared pool and used these data to simulate the active learning process, and we held out the remaining one-third for validation. Each test was initialized with a small labeled training set of 10 instances, then proceeded to apply 1 of the 42 active learning cycles with q_c queries per cycle, and was iterated until 300 queries were added to the labeled set (300 queries is approximately the size of the training set for the smallest data set, the UUKV), and then tested on the validation set. We repeated each test five times and report the average results. Testing took 10,320 CPU hours (9 days) on 48 cores. Finally, we repeated every test using the per-query measurements to provide a generic analysis that is not tied to the specific software and machines we used.

A summary of the results is provided in **Figure 4** (results reported in **Fig. 4** use three queries per cycle). Each column represents a combination of an active learning strategy and SML method, grouped by SML methods. The first three rows show results for the three high-content screens, and the last row shows the average of the three. The number above each bar graph is the average normalized AUC measured over time. Passive learning performance is indicated by a white bar, and the best active learning strategy for each SML method is indicated by a dark bar. The overall best performing strategy for each high-content screen is indicated by an asterisk. A similar table is provided in **Figure 5** (results reported in **Fig. 5** use one query per cycle), but the scores are measured per instance instead of over time. The average labeling time for each data set is provided in **Supplementary Table 1**. In **Supplementary Figure 4**, we provide the average cycle times for least confident sampling and various SML models.

Learning curves for each high-content screen are shown in **Figure 3a–c**. In each case, the most efficient active learning strategy and SML method combination are provided, along with other active learning strategies for the same SML method. Similar curves are provided in **Figure 3d–f** measured per instance instead of over time. A complete set of learning curves is provided in **Supplementary Figure 2**. We also show the time and number of instances required to achieve fixed accuracy thresholds in **Supplementary Tables 3 and 4**.

In **Supplementary Figure 5**, we compare the efficiency of various values of q_c , the number of queries per active learning cycle, where efficiency is measured by the average

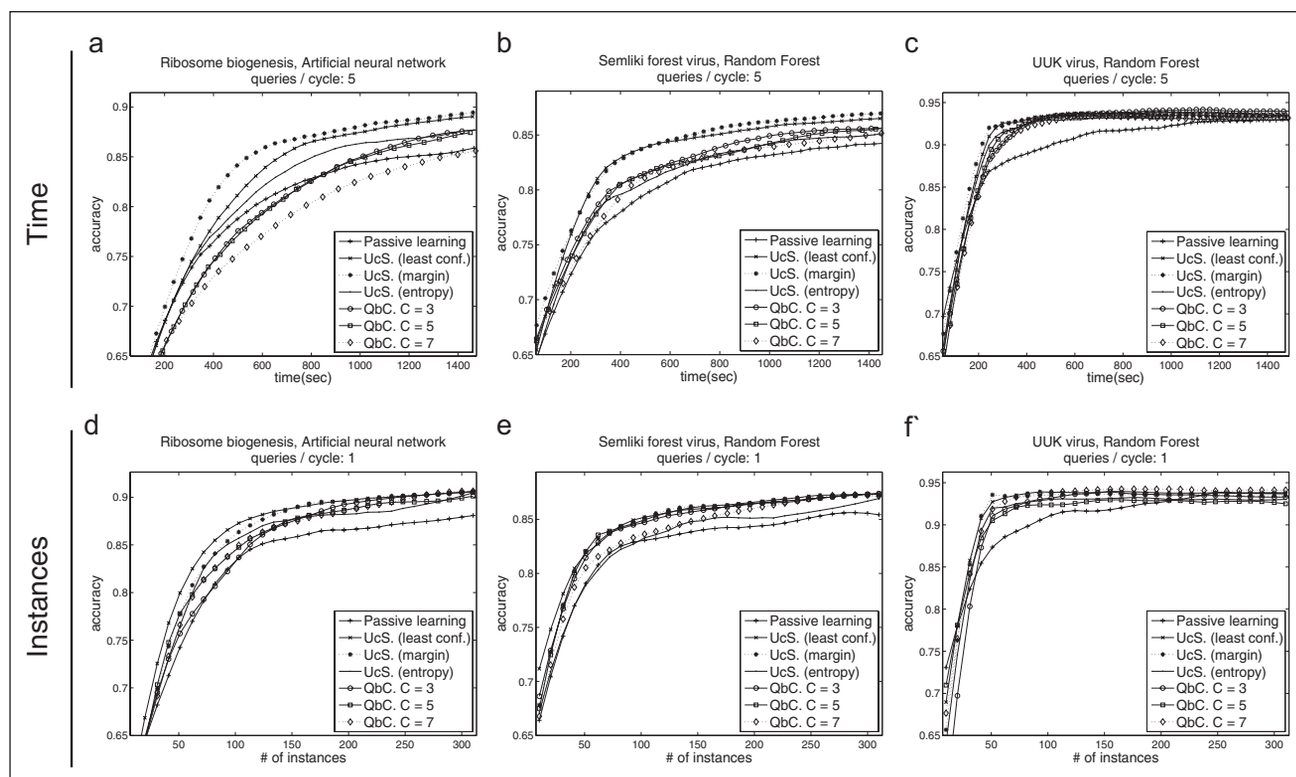


Figure 3. Learning curves for various active learning strategies. The top row shows learning curves as accuracy versus time, and the bottom row shows the accuracy versus number of instances. The curves shown here were selected based on the highest mean area under the learning curve. A complete set is provided in **Supplementary Figure 2**. (a) For the ribosome biogenesis assay, the most efficient strategy was margin sampling on an artificial neural network; (b) for the Semliki Forest virus, the most efficient strategy was margin sampling on random forests; and (c) for the Uukuniemi virus, the most efficient strategy was also margin sampling on random forests.

normalized AUC measured over time. A complete set of figures for all active learning/SML method combinations is provided in **Supplementary Figure 3**.

Overall Performance

The results appearing in **Figure 4** indicate that for the three phenotypic profiling problems considered in this study, there always exists an active learning strategy that will speed up the annotation process. However, there is no universally most efficient combination of active learning strategy and SML method. Random forests with margin sampling was the most efficient method for the SFV and UUKV data sets, but ANN with margin sampling was the most efficient for ribosome biogenesis. This suggests that margin sampling is the best overall active learning strategy, but if we average the performance over all three data sets, random forests with least confident samplings reigns supreme. Interestingly, if we remove time consideration from our evaluation, as shown in **Figure 5**, the overall trends remain similar, but least confident is the best performing active learning strategy for all three data sets. Regardless of the data, random forests coupled with least confident or margin sampling appear to be consistently good choices and would probably be the safest recommendation for a novel data set.

Our investigation into the optimal number of queries to present to the oracle every active learning cycle shows an overall trend, evident in **Supplementary Figure 5**. On average, for low q_c , active learning strategies increase in efficiency as more queries are added to each cycle (**Suppl. Fig. 6**). But this trend quickly plateaus, and then efficiency decreases as more queries are added. There is no universally most efficient value for q_c , but it does appear that each learning method has an optimal choice that depends on the data and the speed of the learner. However, if we take the mean score of all the methods on all the data sets as a measure, $q_c = 3$ is the most efficient number of queries per cycle. This number best balances the expert labeling time and the retraining time of the SML model. An interesting note is the relative poor performance of entropy sampling relative to its uncertainty sampling family members, least confident and margin sampling (**Fig. 6a,b**). We speculate that the reason for this discrepancy is that entropy sampling allows the active learning strategy to suggest examples further from the decision boundary than the other two methods.³⁰ Although for some problems this is a desirable property (e.g., discovery of novel phenotypes), it appears to degrade efficiency on the HCS data sets we considered.

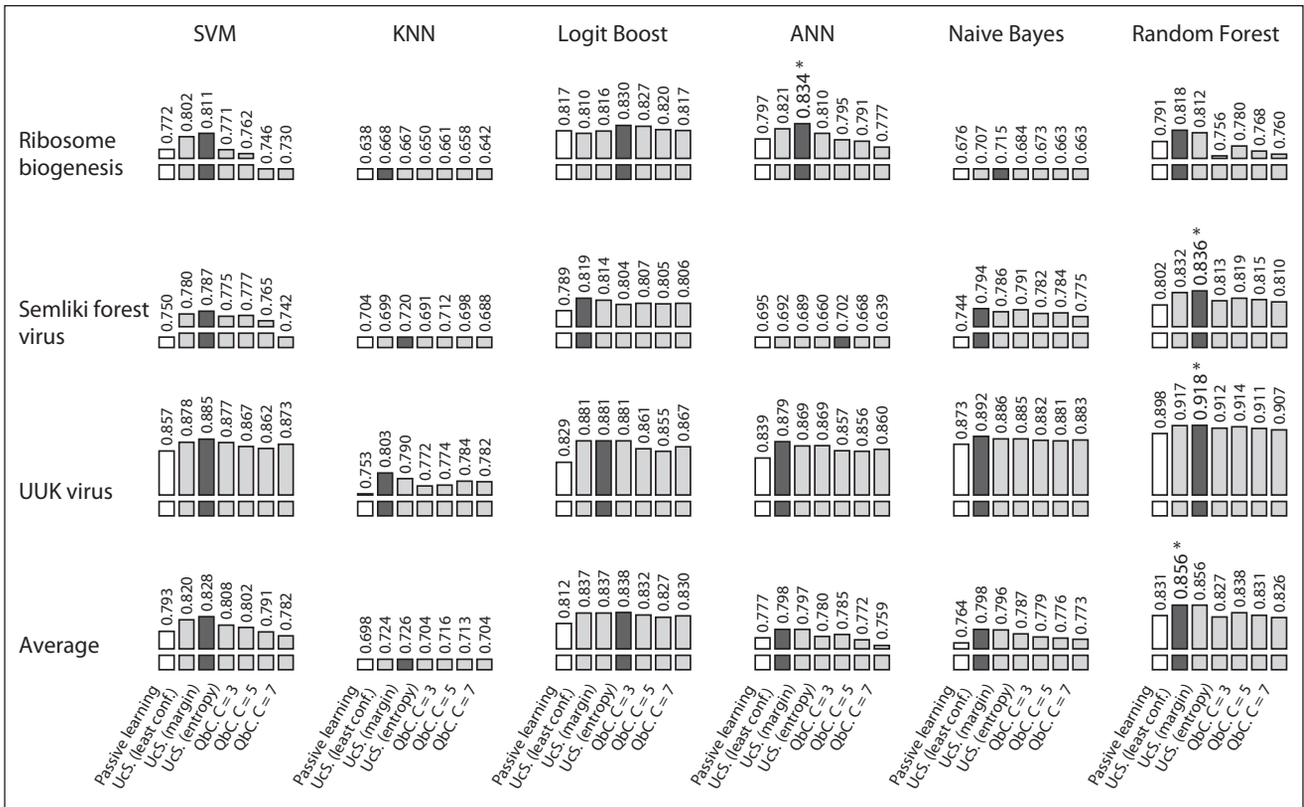


Figure 4. Summary of active learning strategy efficiency (based on time). Performance is measured using the normalized area under the learning curve (AUC) over time. Each column represents a combination of an active learning strategy and supervised machine learning (SML) method, grouped by SML method. The first three rows show results for the three high-content screens, and the last row shows the average of the three. The number above each graph is the average AUC over five tests. Passive learning performance is indicated by a white bar, and the best active learning strategy for each SML method is indicated by a dark bar. The overall best performing strategy for each high-content screen is indicated by an asterisk.

Ribosome Biogenesis

The ribosome biogenesis assay was one of the more difficult phenotypic classification problems. The nucleoplasmic and nucleolar phenotypes are difficult to distinguish, and the overall heterogeneity of cells is strong. **Figure 4** clearly identifies margin sampling with an artificial neural network as dominant over other active learning strategies. In **Figure 3a**, we can see that this approach is able to achieve the same accuracy as passive learning in one-third the time, a significant time savings. Interestingly, in the same plot, all three query-by-committee strategies were less efficient than passive learning for $q_c = 5$, meaning that adopting these strategies would cost the expert time. If we turn to **Supplemental Figure 3**, we can see that query by committee is universally less efficient than passive learning for ANN, a surprising result. We can attribute this to the significant training time of an ANN and the fact that query by committee must train several committee members, whereas only one learner is needed for uncertainty sampling.

Semliki Forest Virus

The SFV data present a different kind of difficulty because they attempt to model a continuous process with discrete

classes; therefore, there can be significant confusion both by the learning method and by the expert. Looking at **Figure 4**, it is clear that random forests and LogitBoost are the most efficient learning algorithms for this problem. Although active learning gives a significant boost in efficiency for this data set, the particular choice of active learning method is less important than the SML method. This is interesting because the SFV data set contained 96 features, the most of the three data sets. The best performers, random forests and LogitBoost, are both meta-learning algorithms built using decision trees, which are known to give good performance for high-dimensional data.

Uukuniemi Virus

The UUKV was the easiest of the three data sets. What is noteworthy here is not so much the performance of the learning strategies but the performance of the annotator. The pregenerated label pool for this assay was annotated quickly and without much discretion in choosing which cell to label. As a result, the gap between passive learning and all active learning methods is significant (**Fig. 3c** and **Fig. 4**). This suggests that there is some advantage to allowing the expert to choose which instances to label instead of sampling at random. Because the

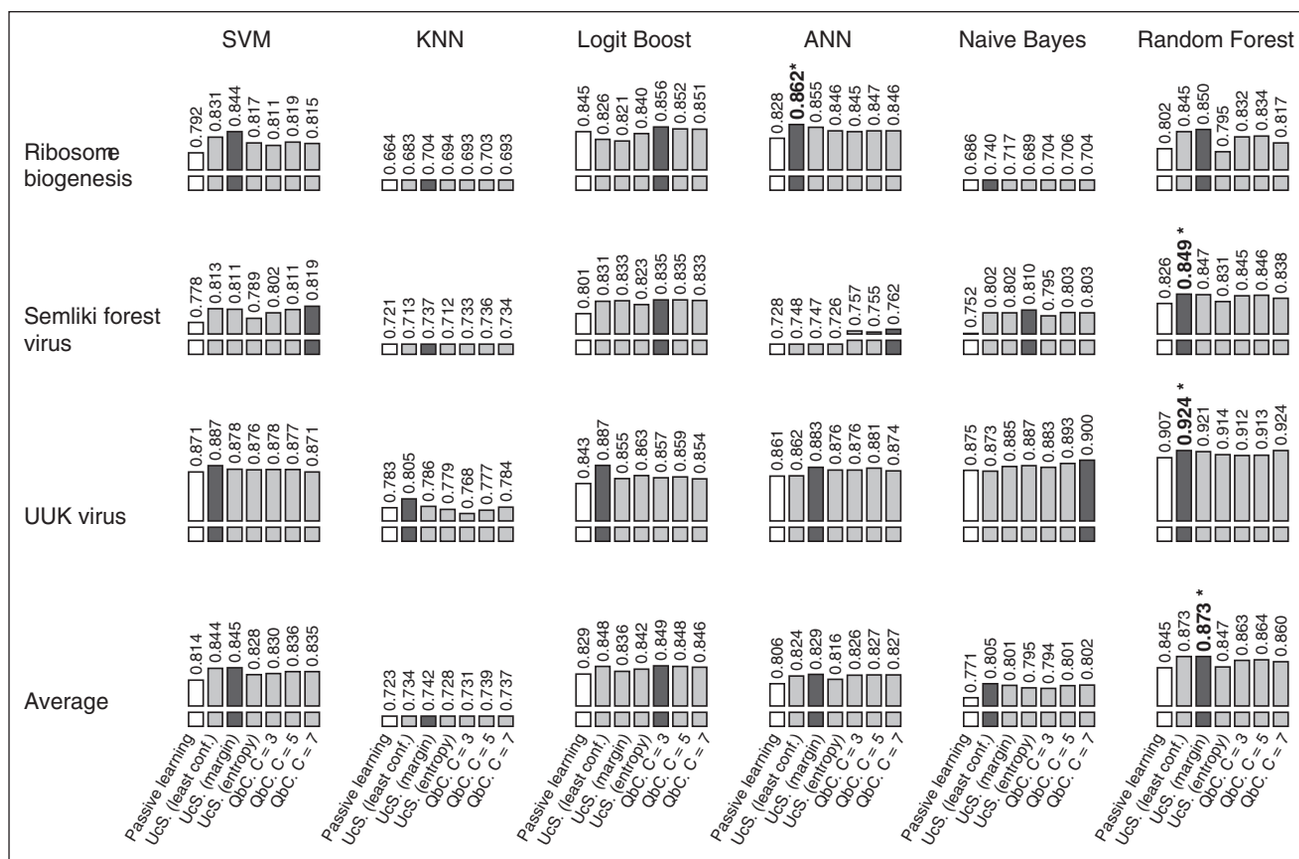


Figure 5. Summary of active learning strategy efficiency (based on queries). Performance is measured using the normalized area under the learning curve (AUC) per query. Each column represents a combination of an active learning strategy and supervised machine learning (SML) method, grouped by SML method. The first three rows show results for the three high-content screens, and the last row shows the average of the three. The number above each graph is the average AUC over five tests. Passive learning performance is indicated by a white bar, and the best active learning strategy for each SML method is indicated by a dark bar. The overall best performing strategy for each high-content screen is indicated by an asterisk.

experts were more picky (less random) in the ribosome biogenesis and Semliki Forest data sets, passive learning was more competitive for those problems.

Discussion

Our tests have shown that for three representative phenotypic profiling problems, active learning can increase the speed and accuracy of the analysis process. Although there is no universally most efficient method, margin sampling with random forests with $q_c = 3$ performed best on average for the problems we considered in this work (Suppl. Fig. 6). We also note that active learning methods that seem to be efficient when measured per query might turn out to be inefficient when real-time costs are considered, and this should be taken into account before using active learning. Another important point to recognize is that allowing the expert to choose which cells to label biases the training set, and the labeled population is no longer representative of the real population. Although active learning does not eliminate bias from the labeled data, one can argue that it is a more

justifiable bias because samples are chosen according to informativeness for the SML model, not human taste. In Supplementary Figure 7, we show that the annotator selection bias does not always strongly affect performance.

There are some limitations to our study that should be mentioned. The classes were known a priori because of the structure of our experiments. In practice, classes are usually discovered on the fly, although this should not significantly affect the results. We also note that the approaches considered in this work do not provide a way to discover automatically novel classes or recognize mistakes in previous annotations based on the discovery of a new class. The size of the data pool available to the active learning methods is smaller in our study than would be used in practice, because of the limited availability of data. Active learning strategies may be more efficient than indicated as more plentiful data may contain more useful instances. Finally, one important topic we did not address in this study is stopping criteria. All active learning methods suffer from diminishing returns as more instances are added; in theory, a good stopping criterion can predict when the cost of providing another label

outweighs its expected performance improvement. Unfortunately, formulating a good stopping criterion is difficult in practice. However, it is possible to provide the expert with an estimate of the accuracy of the current model after each cycle, allowing the expert to make a more informed decision on when to stop training.

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References

1. Swinney, D. C.; Anthony, J. How Were New Medicines Discovered? *Nat. Rev. Drug Discov.* **2011**, *10*, 507–519.
2. Stertz, S.; Shaw, M. L. Uncovering the Global Host Cell Requirements for Influenza Virus Replication via RNAi Screening. *Microbes Infect.* **2011**, *13*, 516–525.
3. Krishnan, M. N.; Ng, A.; Sukumaran, B.; et al. RNA Interference Screen for Human Genes Associated with West Nile Virus Infection. *Nature* **2008**, *455*, 242–245.
4. Banerjee, I.; Yamauchi, Y.; Helenius, A.; et al. High-Content Analysis of Sequential Events during the Early Phase of Influenza A Virus Infection. *PLoS One* **2013**, *8*, e68450.
5. Neumann, B.; Walter, T.; Heriche, J. K.; et al. Phenotypic Profiling of the Human Genome by Time-Lapse Microscopy Reveals Cell Division Genes. *Nature* **2010**, *464*, 721–727.
6. Durr, O.; Duval, F.; Nichols, A.; et al. Robust Hit Identification by Quality Assurance and Multivariate Data Analysis of a High-Content, Cell-Based Assay. *J. Biomol. Screen.* **2007**, *12*, 1042–1049.
7. Misselwitz, B.; Strittmatter, G.; Periaswamy, B.; et al. Enhanced CellClassifier: A Multi-Class Classification Tool for Microscopy Images. *BMC Bioinform.* **2010**, *11*, 30.
8. Horvath, P.; Wild, T.; Kutay, U.; et al. Machine Learning Improves the Precision and Robustness of High-Content Screens: Using Nonlinear Multiparametric Methods to Analyze Screening Results. *J. Biomol. Screen.* **2011**, *16*, 1059–1067.
9. Witten, I.; Frank, E. *Data Mining: Practical Machine Learning Tools and Techniques*; Morgan Kaufmann: Burlington, MA, 2005.
10. Tschochner, H.; Hurt, E. Pre-Ribosomes on the Road from the Nucleolus to the Cytoplasm. *Trends Cell. Biol.* **2003**, *13*, 255–263.
11. Wild, T.; Horvath, P.; Wyler, E.; et al. A Protein Inventory of Human Ribosome Biogenesis Reveals an Essential Function of Exportin 5 in 60S Subunit Export. *PLoS Biol.* **2010**, *8*, e1000522.
12. Fields, B. N.; Knipe, D. M.; Howley, P. M. *Fields' Virology*; Wolters Kluwer Health/Lippincott Williams & Wilkins: Baltimore, MD, 2007.
13. Spuul, P.; Balistreri, G.; Kaariainen, L.; et al., Phosphatidylinositol 3-Kinase-, Actin-, and Microtubule-Dependent Transport of Semliki Forest Virus Replication Complexes from the Plasma Membrane to Modified Lysosomes. *J. Virol.* **2010**, *84*, 7543–7557.
14. Walter, C. T.; Barr, J. N. Recent Advances in the Molecular and Cellular Biology of Bunyaviruses. *J. Gen. Virol.* **2011**, *92*(pt 11), 2467–2484.
15. Lozach, P. Y.; Kuhbacher, A.; Meier, R.; et al. DC-SIGN as a Receptor for Phleboviruses. *Cell Host Microbe* **2011**, *10*, 75–88.
16. Carpenter, A. E.; Jones, T. R.; Lamprecht, M. R.; et al. CellProfiler: Image Analysis Software for Identifying and Quantifying Cell Phenotypes. *Genome Biol.* **2006**, *7*, R100.
17. Olivo-Marin, J.-C. Extraction of Spots in Biological Images Using Multiscale Products. *Pattern Recognition* **2002**, *35*, 1989–1996.
18. Collinet, C.; Stoter, M.; Bradshaw, C. R.; et al. Systems Survey of Endocytosis by Multiparametric Image Analysis. *Nature* **2010**, *464*, 243–249.
19. Kummel, A.; Selzer, P.; Siebert, D.; et al. Differentiation and Visualization of Diverse Cellular Phenotypic Responses in Primary High-Content Screening. *J. Biomol. Screen.* **2012**, *17*, 843–849.
20. David, D. L.; William, A. G. A Sequential Algorithm for Training Text Classifiers. In *Proceedings of the 17th Annual International ACM SIGIR Conference on Research and Development in Information Retrieval*; Springer-Verlag: New York, 1994.
21. David, C.; Les, A.; Richard, L. Improving Generalization with Active Learning. *Mach. Learn.* **1994**, *15*, 201–221.
22. Dana, A. Queries and Concept Learning. *Mach. Learn.* **1988**, *2*, 319–342.
23. Settles, B.; Craven, M.; Friedland, L. Active Learning with Real Annotation Costs. In *NIPS Workshop on Cost-Sensitive Learning*, **2008**.
24. Nicholas, R.; Andrew, M. Toward Optimal Active Learning through Sampling Estimation of Error Reduction. In *Proceedings of the Eighteenth International Conference on Machine Learning*; Morgan Kaufmann: Burlington, MA, 2001.
25. Lewis, D.; Catlett, J. Heterogeneous Uncertainty Sampling for Supervised Learning. In *International Conference on Machine Learning (ICML)*; The International Machine Learning Society, New Brunswick, NJ, 1994; pp 148–156.
26. Burr, S.; Mark, C. An Analysis of Active Learning Strategies for Sequence Labeling Tasks. In *Proceedings of the Conference on Empirical Methods in Natural Language Processing*, Association for Computational Linguistics: Honolulu, HI, 2008.
27. Shannon, C. E. A Mathematical Theory of Communication. *SIGMOBILE Mob. Comput. Commun. Rev.* **2001**, *5*, 3–55.
28. Seung, H. S.; Opper, M.; Sompolinsky, H. Query by Committee. In *Proceedings of the Fifth Annual Workshop on Computational Learning Theory*; ACM: Pittsburgh, PA, 1992.
29. Naoki, A.; Hiroshi, M. Query Learning Strategies Using Boosting and Bagging. In *Proceedings of the Fifteenth International Conference on Machine Learning*; Morgan Kaufmann: Burlington, MA, 1998.
30. Settles, B. *Active Learning*. Morgan & Claypool: San Rafael, CA, 2011.