Active Learning Support Vector Machines to Classify Imbalanced Reservoir Simulation Data

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Abstract—Reservoir modeling is an on-going activity during the production life of a reservoir. One challenge to constructing accurate reservoir models is the time required to carry out a large number of computer simulations. To address this issue, we have constructed surrogate models (proxies) for the computer simulator to reduce the simulation time. The quality of the proxies, however, relies on the quality of the computer simulation data. Frequently, the majority of the simulation outputs match poorly to the production data collected from the field. In other words, most of the data describe the characteristics of what the reservoir is not (negative samples), rather than what the reservoir is (positive samples). Applying machine learning methods to train a simulator proxy based on these data faces the challenge of imbalanced training data. This work applies active learning support vector machines to incrementally select a subset of informative simulation data to train a classifier as the simulator proxy. We compare the results with the results produced by the standard support vector machines combined with other imbalanced training data handling techniques. Based on the support vectors in the trained classifiers, we analyze high impact parameters that separating good-matching reservoir models from bad-matching models.

I. INTRODUCTION

Petroleum reservoirs are normally large and geologically complex. In order to make management decisions that maximize oil recovery, reservoir models are constructed with as many details as possible. Two types of data that are commonly used in reservoir modeling are geophysical data and production data. Geophysical data, such as seismic and wire-line logs, describe earth properties, e.g. porosity, of the reservoir. In contrast, production data, such as water saturation and pressure information, relate to the fluid flow dynamics of the reservoir. Both data types are required to be honored so that the resulting models are as close to reality as possible. Based on these models, managers make business decisions that attempt to minimize risk and maximize profits.

The integration of production data into a reservoir model is usually accomplished through computer simulation. Normally, multiple simulations are conducted to identify reservoir models that generate fluid flows matching the historical production data. This process is called history matching.

History matching is a challenging task for the following reasons:

- Computer simulation is very time consuming. On average, each run takes 2 to 10 hours to complete.
- This is an inverse problem where more than one reservoir model can produce flow outputs that give acceptable match to the production data.

As a result, only a small number of computer simulation runs are conducted and the history matching results are associated with uncertainty.

To address this issue, we have constructed approximate models (proxies) for the computer simulator, based on the computer simulation data. Unlike the full reservoir simulator, which gives the flow of fluids of a reservoir, this proxy only labels a reservoir model as "good" or "bad", based on whether or not its flow outputs match well with the production data. In other words, this proxy acts as a classifier to separate "good" models from "bad" models in the reservoir descriptor parameter space. Using this "cheap" proxy as a surrogate of the full-simulator, we can examine a large number of reservoir models in the parameter space in a short period of time. Collectively, the identified good-matching reservoir models provide us with comprehensive information about the reservoir with a high degree of certainty.

The quality of a proxy, however, relies on the quality of computer simulation data. Frequently, the majority of the simulation outputs match poorly to the historical production data. In other words, most of the data describe the characteristics of what the reservoir is not (negative samples), rather than what the reservoir is (positive samples). Applying machine learning methods to train a simulator proxy based on these data faces the challenge of imbalanced training data. This work applies active learning support vector machines (SVMs) to incrementally select a subset of informative simulation data to train a classifier as the simulator proxy. We compare the results with the results produced by the standard support vector machines combined with other imbalanced training data handling techniques. Based on the support vectors in the trained classifiers, we analyze high impact parameters that separating good-matching reservoir models from bad-matching models.

The paper is organized as follows. Section II explains SVM, active learning SVM and other techniques used in this work to address the issue of learning using imbalanced data. In Section III, the reservoir field and the computer simulation data are described. We explain the performance measurements in Section IV. In Section V, the data preprocessing and the experimental setup are detailed. We present the results and give our analysis in Section VI. Finally, Section VII concludes the paper and outlines our future work.

II. METHODOLOGY

In this section, we first briefly describe the basic concept of two-class SVM classification [13], [3]. Extension of SVM
and other techniques we used to handle imbalanced training data will be explained in the subsections.

For a binary SVM classification task, the training data are mapped into a higher dimensional feature space using a kernel function, such that a hyper-plane separating the two classes of data can be found. The hyper-plane consists of data \( x \) that satisfies \( w^T \phi(x) + b = 0 \), where \( w \) is the weight vector, \( x \) is the training data, \( \phi \) is the kernel function and \( b \) is the bias (see Figure 1).

While there are more than one hyper-planes that can separate the two classes of data, a SVM learner finds the one that has the maximal margin or the distance between the data that are closest to the hyper-plane (see Figure 1). Using the Lagrangian formation, \( w = \sum_{i=1}^{n} \alpha_i y_i \phi(x_i) \) gives the largest margin. The optimal separating hyper-plane is:

\[
w^T \phi(x) + b = \sum_{i=1}^{n} \alpha_i y_i \phi(x_i) + b
\]

\[
= \sum_{i=1}^{n} \alpha_i y_i \phi(x_i) + b = 0
\]

\[
= \sum_{i=1}^{n} \alpha_i y_i K(x, x_i) + b
\]

where \( n \) is the number of training data, \( y_i \) is the label value of data \( i \), \( K \) is the kernel matrix, and \( \alpha_i \) is the Lagrange multipliers, where \( \alpha_i \geq 0 \) and \( \sum \alpha_i y_i = 0 \).

Each training data \( x_i \) is associated with a Lagrange multiplier \( \alpha_i \). The larger the \( \alpha_i \) is, the more important that data is to the classification decision. Those data that lie close to the hyper-plane decision boundaries have nonzero \( \alpha_i \) and are called support vectors. These support vectors form the classifier that will be used to predict the class of new data.

When dealing with real-world data, which are normally noisy and uncertain, the hyper-plane margins can be relaxed by using a slack variable \( \xi \). Thus, a hyper-plane becomes:

\[
w^T \phi(x_i) + b \geq 1 - \xi_i, y_i = 1
\]

\[
w^T \phi(x_i) + b \leq -1 + \xi_i, y_i = -1
\]

\( x_i \geq 0 \forall i \)

An error occurs when the corresponding \( \xi \), for a data \( x_i \), exceeds 1 (see Figure 1). Under this setup, the optimal hyper-plane is not just with the maximal margin but also with the minimal total error. A SVM learner finds the optimal hyper-planes by maximizing the following objective function:

\[
|w|^2 + C \sum_{i=1}^{n} \xi_i
\]

where \( C \) is a regularization parameter that controls the trade-off between the two objectives. The larger the \( C \) is, the more penalty is assigned to the errors. Smaller \( C \) values relax this penalty and produce better results with noisy data [7]. Solving this multi-objective problem leads to the optimal hyper-plane that is the same as before (Equation 3). The only difference is that the Lagrange multiplier is constrained by \( C \), i.e. \( 0 \leq \alpha_i \leq C \). That is, the influence of an individual Lagrange multiplier \( \alpha_i \) has on the prediction decision is limited to \( C \).

**A. Asymmetric Soft Margins**

When the training data set is imbalanced, we can assign different cost \( C_+ \) and \( C_- \) to each of the two classes. The object function to optimize the hyper-plane thus becomes:

\[
|w|^2 + C_+ \sum_{i=1}^{n} \xi_i + C_- \sum_{i=1}^{n} \xi_i
\]

This approach was introduced by [14] to control the trade-off between false positives and false negatives. The idea was that by introducing different cost for positive and negative data, a bias of larger Lagrange multiplier \( \alpha_i \) is given to the class where the cost of misclassification is heavier. This, in turn, introduces a decision boundary which is much more distant from the ‘critical’ class than from the other. According to [11], the distance of a test data point from the boundary is related to its probability of mis-classification. Test points that are farther away from the hyper-plane are less likely to be misclassified.

In our study, we will assign \( C_+ \) a smaller value than the value of \( C_- \) to encourage the SVM learner to find a hyper-plane that classifies more positive samples correctly.

**B. Active Learning SVM**

SVM is a passive learning method if the training data are pre-selected for the learner to find the optimal separating hyper-plane. In an active learning framework, the learner actively seeks the training data, one after another, based on its needs. In other words, the learner can query/select informative new data to help train a better hyper-plane. Various research has explored this active learning framework in SVMs.

For example, Campbell, Cristianini and Smola[4] used the strategy that “starts by requesting the labels of a random subset of instances and subsequently iteratively requesting the label of that data point which is closest to the current hyper-plane” in their active learning SVM system. In terms of termination criterion, they “trained until either the margin band is empty or until a stopping point specified by the user has been met.” Their experimental results showed that their SVM learner processed a smaller number of training.
data to find the hyper-plane that performed as well or better (in terms of classification accuracy) than that found by the SVM using the random selection strategy. This indicates that the samples their method selected are more informative than the randomly selected samples. Schohn and Cohn also independently devised a similar selection method in their active learning SVM [10].

Tong and Koller [12] worked on the kind of training set that has some data labeled and some not. Active learning SVMs were used to select an unlabeled data sample, label it and then add it to the training set for the SVM learner to find a better separating hyper-plane. They devised 3 selection schemes:

- Simple Margin: select the data that is closest to the hyper-plane, which is the same as that used in [4] [10].
- MaxMin Margin: for each unlabeled data \( x \), compute the margin \( m^+ \) and \( m^- \) of the hyper-planes obtained when the data is labeled as +1 and -1 respectively; then select the unlabeled data whose \( \min(m^+, m^-) \) is the greatest.
- Ratio Margin: compute \( m^+ \) and \( m^- \) as that in the MaxMin Margin method. However, select the data for which \( \min(m^+, m^-) \) is the largest.

They reported that all 3 strategies performed better than the random selection strategy in that their strategies selected a smaller number of data for the SVM learner to find the optimal hyper-plane. Moreover, the MaxMin and Ratio margin strategies gave more stable performance on different data sets than the Simple margin strategy did. However, MaxMin and Ratio margins took longer to compute, hence lead to a longer SVM training time.

LASVM is an active learning SVM system developed by Bordes, Ertekin, Weston and Bottou [2] with the aim to speed up the SVM training time for applications which have a very large volume of data. The selection methods they used include:

- Gradient Selection: randomly select 50 data that have not been processed. Among them, the one that is most mis-classified, according to the current hyper-plane decision, is selected.
- Active Selection: randomly select 50 data that have not be processed. Among them, the one that is closest to the current hyper-plane decision boundaries is selected.
- Autoactive Selection: This replaces the sampling size of 50 in the Active Selection with a more flexible size. It continues to sample the unprocessed data until 5 of the samples are within the distance of \( 1 + \delta/2 \) (\( \delta \) is the gradient of the most \( \tau \)-violating pair in the current support vector set) to the hyper-plane decision boundaries or the maximum number of 100 samples is reached. Among them, the one that is closest to the current hyper-plane decision boundaries is selected.

Among the data sets they tested, Active and Autoactive selections performed well consistently. These two methods selected a smaller number of training data for LASVM to find an equivalent or better hyper-plane as that found by a standard SVM system (LIBSVM). The trained SVMs also contained a smaller number of support vectors than the SVM trained by LIBSVM. In contrast, Gradient selection only performed well when the data set was not noisy. This is understandable as a misclassified data, according to the current hyper-plane decision, might be actually classified correctly, since the data is noisy. Selecting those kind of misclassified data to add into the training set is less likely to help train an improved hyper-plane.

LASVM differs from the three previous mentioned active learning SVMs in the following two ways:

- The selection is based on a small number of random samples, instead of the entire unprocessed samples.
- Once a new data sample is selected and added into the training set, the current hyper-plane is updated, instead of being completely retrained.

They also compared the performance of the hyper-planes learned under the selection based on a small sample size (50) or a large sample size (all), and with the hyper-plane retrained or updated at each iteration. They reported that if the data set size is large, all setups produced SVMs that had a similar prediction accuracy. But when the data set size is small, the SVMs trained by selection within a small sample size of 50 or with hyper-plane updated (instead of retrained) after a new data point was added to the training set had a small accuracy loss.

In [8], Ertekin, Huang, Bottou and Giles applied LASVM to imbalanced data to improve the SVM prediction on minority class data. They used the Active selection method to select new data from a pool of 50 random samples. They also implemented the termination criterion of [4] by comparing the newly selected data with the support vectors in its distance to the hyper-plane. If the new data (which is the closest to the hyper-plane among the 50 random samples) is not closer to the hyper-plane than any of the current support vectors, this indicates that there is no data points within the margin band and the learning process stopped.

They tested their version of LASVM on a number of imbalanced data sets. They also applied a standard SVM system (LIBSVM) combined with other techniques that handle imbalanced data, such as asymmetric soft margins and SMOTE (explained in the next subsection). They reported that all techniques produced SVMs that gave similar prediction performance. However, their LASVM processed a smaller number of data and the produced hyper-plane contained a smaller number of support vectors.

In our work, we will design experimental setups (detailed in Section V) similar to theirs and run the same version of LASVM on our data set.

C. SMOTE

SMOTE stands for Synthetic Minority Oversampling Technique, which was developed to improve the prediction accuracy of classifiers trained under imbalanced data [6]. For each minority sample, the algorithm first finds its \( k \)-nearest minority neighbors. Among these neighbors, \( j \) of them are
randomly selected. Each of the \( j \) neighbors is paired with the original sample to create one synthetic data point using the following three steps:

- Take the difference between the feature vectors of the original sample and sample \( j \);
- Multiply this difference vector by a random number between 0 and 1;
- Add this vector to the feature vector of the original sample;

The value \( k \) and \( j \) are parameters which can be varied depending on the amount of over-sampling needed. For example, if 200% of the minority class data are needed, the \( j \) is 2. The value of \( k \) is set as 5 in their examples. For the imbalanced data sets they tested, SMOTE worked better (measured by the AUC of the trained classifiers, which will be explained in Section IV) than the under-sampling of the majority class method in most of the tested data sets. They explained that the reason why SMOTE technique helped a learning method to produce a better classifier was because it caused the classifier to build larger decision regions that contain nearby minority class points. In contrast, the duplication of minority data method led to smaller and more specific decision regions. We will include this oversampling technique to create a more balanced data set for SVM training in our study.

III. RESERVOIR SIMULATION DATA

The simulation data were based on a large reservoir that has been in production for a long time. Due to the long production history, the data collected from the field were not consistent and the quality was not reliable. Although we do not know for sure what causes the production measurements to be inaccurate, we could speculate on newer technology but more likely it is just poor measurement taken from time to time.

This reservoir is overlain by a significant gas cap. Figure 2 shows the gas oil contact (GOC), which is the surface separating the gas cap from the underneath oil, and the water oil contact (WOC), which is the surface that separates oil from the water below, of the reservoir. The space between GOC and WOC surfaces is the oil volume to be recovered. The field also has 4 geological faults, illustrated in Figure 3, which affect the oil flow patterns. Those faults have to be considered in the computer flow simulation.

As a mature field with most of its oil recovered, the reservoir had residual pore space which can be used for storage. One proposed plan was using the pore space to store the gas produced (as a side product) from the neighboring oil fields. In this particular case, the gas produced had no economical value and re-injecting it back into the field was one environmental-friendly method to dispose the gas.

In order to evaluate the feasibility of the plan, the cumulative volume of gas that can be injected (stored) in year 2031 needed to be evaluated. This evaluation would assist managers in making decisions such as how much gas to transport from the neighboring oil fields and the frequency of the transportation.

The cumulative volume of the gas that can be injected in the field is essentially the cumulative volume of the oil that will be produced from the field, since this is the amount of space that will become available for gas storage. To answer that question, a production forecast study of this field in the year of 2031 was conducted.

Prior to carrying out the production forecast, the reservoir model has to be updated through the history matching process. We consulted a field engineer to select 10 reservoir parameters and their value ranges to conduct computer simulation. These parameters, as shown in Table I, are unitless except WOC and GOC, which have feet as their unit. Critical Gas Saturation (SGC) is the gas saturation values for each phase (water and oil) and has a value between 0 and 1. Skin (SKIN), which is the rock formation damage caused by the drilling of new gas injector wells, has a value between 0 and 30. The other 6 parameters are multipliers whose values are in \( \log_{10} \) scale.

During computer simulation, the 10 parameter values are used to define reservoir properties in each grid of the 3-D model in two different ways. The values of the 4 regular parameters are used to replace the default values in the original 3-D reservoir model while the 6 multipliers are applied to the default values in the original model. Computer simulations are then performed on the updated 3-D reservoir model to generate flow outputs.

Based on the uniform design method, values of the 10 parameters were decided to conduct 600 computer simulation
runs. Among them, 593 were successful in that the runs continued until year 2031. The other 7 runs failed for various reasons, such as power failures, and terminated before the runs reached year 2031.

During the computer simulation, various flow data were generated. Among them, only field water production rate (FWPR) and field gas production rate (FGPR) of 30 production years were used for history matching. The other flow data were ignored because we do not trust the quality of their corresponding production data collected from the field.

FWPR and FGPR collected from the field were compared to the computer simulation outputs at each run. The error $E$, defined as the mismatch between the two, is the sum of squared errors calculated as follows:

$$E = \sum_{i=1}^{30} (FWPR_{obs_i} - FWPR_{sim_i})^2 + (FGPR_{obs_i} - FGPR_{sim_i})^2$$

Here, $obs$ indicates production data while $sim$ indicates computer simulation outputs. The largest $E$ that can be accepted as good match is 1.2. Additionally, if a model has $E$ smaller than 1.2 but has any of its $FWPR$ or $FGPR$ simulation outputs match badly to the corresponding production data (difference is greater than 1), the production data were deemed to be unreliable and the entire simulation record is disregarded. Based on this criterion, 12 data were removed.

We then conducted an outlier study on the remaining simulation data by examining the consistency of inputs and outputs patterns [15]. There were 7 data points falling outside the patterns and were removed from the data set. The final set contained 564 simulation data. Among them, 63 were labeled as “good” models while 501 were labeled as “bad” models. In the rest of the paper, we will call the data labeled as “good” models positive samples and the data labeled as “bad” models negative samples.

IV. PERFORMANCE MEASURES

In binary classification problems, to which this work belongs, performance measures based on classification accuracy are meaningless, since a classifier can simply predict all data as the majority class to obtain a reasonably good accuracy. Yet, this classifier is practically useless in identifying the minority class data, which is more important in most cases. In our case, if the simulator proxy can not identify good reservoir models, this proxy can not be used to substitute the computer simulator.

There are several performance measures developed to evaluate classifiers trained under imbalanced data. We adopted several of them to provide a more subjective view of the quality of the trained classifiers. Most of these measures are calculated from the values in a confusion matrix (Table II).

Precision gives the percentage of data that is predicted as positive is indeed positive. Recall gives the percentage of the positive data that is correctly predicted as positive. Precision and recall often have an inverse relationship, where the increase of one may reduce the other. Usually, these two measures are used in combination. For example, $F$-score is the weighted harmonic mean of precision and recall.

- $Precision = TP / (TP + FP)$.
- $Recall = TP / (TP + FN)$.
- $Fscore = 2 \times Precision \times Recall / (Precision + Recall)$.
- $AUC = area under ROC curve$.

Receiver Operating Characteristic (ROC) displays the relationship between sensitivity ($TP / (TP + FN)$) and specificity ($TN / (TN + FP)$) of all possible thresholds for a binary classifier, applied to previous unseen testing data. In other words, ROC is a plot of the TP rate against the FP rate as the decision threshold is changed. The area under a ROC ($AUC$) is the numerical measure of a classifier’s prediction ability to separate the positive and negative data. Since $AUC$ evaluates a classifier across the entire range of decision thresholds, it gives a good indication of the classifier’s performance under the situation where the data distribution is imbalanced [9]. The larger the $AUC$ of a classifier is, the better its performance is.

V. EXPERIMENTAL SETUP

We first normalized the data to have values between 0 and 1. The resulting 564 data were then split into training (2/3) and testing (1/3) sets. We used the 451 training data (50 were positive and 401 were negative samples) to conduct 5-fold cross-validation to identify SVM parameter values that gave the best $AUC$. The selected parameter values were then used to train the final SVM model using the 451 data. The quality of the final SVM model was evaluated based on its prediction on the 113 testing data (13 were positive and 100 were negative samples).
We used two public domain SVM tools to conduct our experiments: LIBSVM [5] and LASVM [2]. LIBSVM is a state-of-the-art SVM solver while LASVM is an active learning SVM system. We conducted six sets of experiments: (a) LIBSVM baseline (b) LIBSVM with asymmetric soft margins (c) LIBSVM on SMOTE data (d) LIBSVM with asymmetric soft margins on SMOTE data (e) LASVM (f) LASVM with asymmetric soft margins. All these experiments used the radial basis function (RBF) kernel $K(u, v) = e^{-\gamma(u-v)\cdot(u-v)}$ to conduct SVM training. The values of other SVM parameters (see Table III) were decided by 5-fold cross-validation.

We applied the SMOTE process to the 50 positive training data to generate 400 synthetic data, 8 for each positive sample. This led to a total of 450 positive and 401 negative samples in the final training set. For setups 5 and 6, LASVM started with an initial training set of 10 data, 5 were positive and 5 were negative samples, which were randomly selected from the entire training data set.

We carried out the 6 sets of experiments and applied the learned SVMs on the 113 testing data. The results are given in Table IV. Note that in addition to the performance measurements discussed in Section IV, we also provide classification accuracy (acc) for reference.

## VI. Results and Analysis

As shown in Table IV, the LIBSVM baseline, setup (a), gave the worst performance, in terms of F-score and AUC. The other 5 setups, which included one or two of the imbalanced data handling techniques discussed previously, trained SVMs that gave similar performance. Among them, LIBSVM combined with asymmetric soft margins and the SMOTE technique, setup (d), gave the best F-score and AUC. This is similar to that reported in [1] where the SVM trained under asymmetric soft margins and SMOTE data gave better performance than the SVM trained under either of the two techniques alone. However, the AUC and F-score differences between setups (b), (c) and (d) are marginal.

LASVM, setup (e) and (f), did not work better than LIBSVM combined with the SMOTE technique, setup (c) and (d). This result is similar to that reported in [2] where in small data sets, which our data set belongs to, LASVM does not provide performance advantage over other techniques. However, it selected a smaller number of data samples (see Table V) to train a SVM with a smaller number of support vectors (see Table VI) than that based on SMOTE data. A similar result was also reported in [8].

It has been suggested that the performance gain of the active learning SVM on imbalanced data, compared to the standard SVM (setup (a)), is due to its ability to supply the learner a more balanced training data set [8]. This is shown to be untrue in our case. The ratio of the positive and negative samples selected by LASVM (setup (e) and (f)) is the same as that of the entire training set: 1:8. This indicates that the active learning method is choosing more informative samples to overcome the learning bias created by the majority class data. Tong and Koller also observed a similar data selection ratio as ours in their active learning SVM system [12].

LIBSVM combined with the SMOTE technique (setup (c) and (d)) produced classifiers with the best AUC. This indicates that there is a certain smoothness in the inputs space. The SVM learner was able to use the gradient information to find the optimal hyper-plane.

### A. Model Interpretation

In data mining, one of the objectives is to understand the knowledge extracted in the learned model and apply that knowledge to the domain system. Unfortunately, SVM is a black-box model, which is not easy to interpret. However, since the model is essentially the critical samples (support vectors) for classification, we can examine the support vectors, particularly their relationship to the entire training set in the input space, to gain some insights of the reservoir characteristics. To be conservative in our interpretation, we only use the support vectors selected by all 6 SVM learning systems to conduct our analysis. The number of such kind of support vectors is 32.

Figure 4 gives the mean and standard deviation of the 10 reservoir parameter values calculated from the 32 support

### Table III

**Parameters decided by cross-validation.**

<table>
<thead>
<tr>
<th>SETUP</th>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
<th>(d)</th>
<th>(e)</th>
<th>(f)</th>
</tr>
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<tbody>
<tr>
<td>$\gamma$</td>
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<td>$\times$</td>
<td>$\times$</td>
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### Table IV

**Performance evaluation on 113 testing data.**

<table>
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<th>SETUP</th>
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<th>(b)</th>
<th>(c)</th>
<th>(d)</th>
<th>(e)</th>
<th>(f)</th>
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<td>TP</td>
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<td>10</td>
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</tr>
<tr>
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<td>3</td>
<td>3</td>
<td>6</td>
<td>4</td>
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</tr>
<tr>
<td>Acc</td>
<td>87.61%</td>
<td>86.73%</td>
<td>83.19%</td>
<td>84.07%</td>
<td>87.61%</td>
<td>87.61%</td>
</tr>
<tr>
<td>Pre</td>
<td>44.44%</td>
<td>43.43%</td>
<td>38.46%</td>
<td>48.00%</td>
<td>46.61%</td>
<td>47.37%</td>
</tr>
<tr>
<td>Rec</td>
<td>30.77%</td>
<td>76.92%</td>
<td>78.92%</td>
<td>76.92%</td>
<td>53.85%</td>
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<tr>
<td>F-Score</td>
<td>36.36%</td>
<td>57.14%</td>
<td>51.28%</td>
<td>56.63%</td>
<td>50%</td>
<td>56.25%</td>
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<tr>
<td>AUC</td>
<td>80.07%</td>
<td>89.93%</td>
<td>92.08%</td>
<td>92.23%</td>
<td>86.46%</td>
<td>90.15%</td>
</tr>
</tbody>
</table>

### Table V

**Number of data processed.**

<table>
<thead>
<tr>
<th>SETUP</th>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
<th>(d)</th>
<th>(e)</th>
<th>(f)</th>
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</thead>
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<td>851</td>
<td>851</td>
<td>441</td>
<td>441</td>
</tr>
</tbody>
</table>

### Table VI

**Number of support vectors.**

<table>
<thead>
<tr>
<th>SETUP</th>
<th>(a)</th>
<th>(b)</th>
<th>(c)</th>
<th>(d)</th>
<th>(e)</th>
<th>(f)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive SVs</td>
<td>50</td>
<td>48</td>
<td>120</td>
<td>48</td>
<td>42</td>
<td>44</td>
</tr>
<tr>
<td>Negative SVs</td>
<td>163</td>
<td>248</td>
<td>121</td>
<td>114</td>
<td>95</td>
<td>106</td>
</tr>
<tr>
<td>Total SVs</td>
<td>215</td>
<td>296</td>
<td>241</td>
<td>162</td>
<td>137</td>
<td>150</td>
</tr>
</tbody>
</table>
Vectors and from the entire training set. For WOC and GOC, we report the mean and standard deviation of the oil column (WOC-GOC) instead, as our previous study [15] has identified that oil column is an important feature to distinguish a good-matching reservoir model from a bad-matching reservoir model. Note that all parameter values have been normalized to have values between 0 and 1.

Selected by the uniform design method, the parameter values of the training data are consistent with mean around 0.5 and standard deviation around 0.3. The parameter values of the support vectors, however, are more diverse. Among them, WOC-GOC and YERM have the mean values that are higher than the means of the training data. Similarly, TRANS and SGC have their mean values that are lower than the means of the training data. This indicates that these 5 reservoir parameters are important features separating a good-matching reservoir model from a bad-matching reservoir model. This result is consistent with that found in our previous study on the same reservoir [15]. Note that we ignore SKIN parameter as it is related to the future drilling of new injector wells.

We analyze the 32 support vectors farther by dividing them into two groups: good-matching reservoir models and bad-matching reservoir models. Figure 5 gives the parameter mean values of the two groups. As shown, the same 5 parameters have their good-matching model mean and bad-matching model mean close to each other, indicating they are similar in behavior. In other words, these 5 reservoir parameters have values in the support vector group that are similar to each other but are different from that in the non-support-vector group. This supports our analysis that they are important reservoir features to classify a good-matching reservoir model from a bad-matching model. Note that we ignore XYPERM as its mean in the support vector group is similar to the mean in the training data, hence can not be considered as an important feature for classification.

Figure 6 gives the cross-plot between WOC-GOC and YPERM for the training data and the support vectors. It shows that good-matching models have oil column height (WOC-GOC) between 750 and 825. Also, their YPERM is higher than 1.07. We will validate these reservoir characteristics by studying more good-matching reservoir models interpolated by the trained SVM classifier proxy in our future work.

Figure 7 gives the cross-plot between TRANS and SGC for the training data and the support vectors. As shown, good-matching reservoir models have their TRANS and SGC cross the entire value ranges. We therefore can not conclude their value ranges in this reservoir. We will conduct more analysis on these two reservoir characteristics by studying more good-matching reservoir models interpolated by the trained SVM classifier proxy in our future work.
Reservoir modeling and simulation play an important role in the management of petroleum energy exploration and production. Although the surrogate model approach has great potentials to provide more accurate reservoir models, it can help management making more reliable decisions, its effectiveness is dependent upon the construction of high quality proxies. One challenge that makes quality proxies unattainable is the inadequacy of the computer simulation data, which frequently only provide partial views of the reservoir characteristics. One typical case is the majority of the simulation data describe the characteristics of what the reservoir is not (negative samples), rather than what the reservoir is (positive samples). Such kind of data set is difficult for most machine learning methods to produce a proxy model that can separate reservoir models that match historical production data well from those that don’t. This work addresses this problem by applying an active learning SVM system to construct a classifier as the proxy model. The result of our investigation is very encouraging.

During the training process, the active learning method intelligently selected informative data, one after another to help the SVM learner improve the hyper-plane. The resulting model is a classifier that performs well on the testing data, based on the AUC measure. This indicates that the technique is suitable in handling imbalanced reservoir simulation data produced by computers. Other investigated techniques also worked well. This gives us multiple tools to use in the future depending on the size and quality of the data set. Additionally, the SMOTE technique, which relies on the smoothness in the input space, helped the SVM learner to build a quality classifier. This suggests that this technique can be effective on other type of data, as long as the smoothness holds in the input space. We will test this method on other type of data in conjunction with other machine learning methods to produce models represented in different forms.

We also examined the support vectors in the trained classifiers to analyze parameters that have high impact on reservoir classification. Our analysis shows that oil column (WOC-GOC), YPERM, TRANS and SGC have values in the support vector group that are similar to each other but are different from that in the non-support-vector group. This indicates that they are important characteristics separating good-matching reservoir models from bad-matching reservoir models. This result is consistent with what we have found in our previous study on this reservoir.

In terms of these 5 critical parameters’ value ranges in this reservoir, the oil column is approximately between 750 and 825 feet and the Y-perm multiplier is greater than 1.07, based on the provided good-matching reservoir model data. The fault transmissibility multiplier (TRANS) and the critical gas saturation (SGC) values, however, can not be derived from these good-matching reservoir model data. In our future work, we will examine more good-matching reservoir models interpolated by the trained SVM classifier proxy. With a larger number of good-matching models, the reservoir characteristics related to these 5 critical parameters will be estimated more accurately.

REFERENCES


