Major earthquake event prediction using various machine learning algorithms

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Abstract—At least two basic categories of earthquake prediction exist: short-term predictions and forecast ones. Short term earthquake predictions are made hours or days in advance, while forecasts are predicted months to years in advance. The majority of studies are done on forecast, taking into consideration the history of earthquakes in specific countries and areas. In this context, the core idea of this work is to predict whereas an event is classified as negative or positive major earthquake by applying different machine learning algorithms. Eight different algorithms have been applied on a real earthquake dataset, namely: Random Forest, Naive Bayes, Logistic Regression, MultiLayer Perceptron, AdaBoost, K-nearest neighbors, Support Vector Machine, and Classification and Regression Trees. For each selected model, various hyperparameters have been selected, and obtained prediction results have been fairly compared using various metrics, leading to a reliable prediction of major events for 3 of them.

Index Terms—Machine learning, Major earthquake prediction

I. INTRODUCTION

Rapid global urbanization leads to an increase in earthquakes, a major catastrophe which has occupied 60% of deaths overall natural disasters [8]. Obviously, human cannot stop natural disasters, while the application of machine learning is a powerful and invaluable method and technique, used by researchers as new area of study in geology, in order to reduce, as much as they can, the loss of life and billions of dollars in infrastructure and housing costs. Traditional classification methods rely on statistical assumptions for earthquakes that turn out to be unsatisfactory in danger state prediction. Therefore, the use of machine learning algorithms is becoming more widespread, that adapt and learn a problem by simulating a biological or natural system.

Introducing the machine learning techniques in the field of geology and earthquake beats the traditional and standard methods used in earlier years and provides the scientists a new fresh method for assessing seismic risk and triggering future earthquakes. Data mining applications have achieved great success in geophysics and geology such as ecology, weather prediction, modeling, etc. and finally predicting the earthquakes. Two basic categories of earthquake prediction exist: short-term predictions and forecast ones. Short term earthquake predictions are made hours or days in advance, while forecasts are predicted months to years in advance. Due to the chaotic and complex phenomena of earthquake process, it is considered that short-term predictions are very difficult to address. Thus, the majority of studies are done on forecast, taking into consideration the history of earthquakes in specific countries and areas.

The core idea of this work is to predict whereas an event is classified as negative or positive major earthquake by applying different machine learning algorithms. It is well known that there is no best algorithm or one solution that fits all the problems and datasets for machine learning since the performance of algorithms depends on many factors. Some algorithms are best for small data, while others perform better for a tons of data sample. Some algorithms require categorical inputs, while others need quantitative. Another important criterion while choosing the algorithm is the complexity of the dataset and how many features the model needs to learn and predict. This is why, in this work, eight different algorithms have been applied on an earthquake dataset, namely: Random Forest, Naive Bayes, Logistic Regression, MultiLayer Perceptron, AdaBoost, K-nearest neighbors, Support Vector Machine, and Classification and Regression Trees. For each selected model, various hyperparameters have been tested, and obtained prediction results have been fairly compared using various metrics, leading to a reliable prediction of major events for 3 of them.

The remainder of this paper is organized as follows. The next section contains the basic recalls. Section III discusses related work done by researchers in the field of earthquake machine learning. In Section IV, we apply experiments on the testing set after training the data using machine learning algorithms and for some models, parameters were modified, then results were compared. Finally, Section V concludes the paper.
II. BASIC RECALLS

We begin by recalling the various classical machine learning methods that have been proved useful in predicting earthquakes, before detailing, in the next section, some state-of-the-art articles that have used them in this context.

A. Naive Bayes

The Naive Bayes algorithm, a method based on the work of Thomas Bayes, is a heavily simplified probabilistic model that calculates probabilities by counting combinations of values and frequency in a data set. Bayesian classification assumes that the data belongs to a particular class, then the probability for the hypothesis to be true is calculated. Naive Bayes operates on a strong independence assumption, in other terms, the conditional probability of one attributes will not affect the probability of the other. However, the hypothesis can be updated each time new evidence is added.

Bayes Theorem calculates the probability of an event A given another event B has occurred:

\[
P(A|B) = \frac{P(B|A) \cdot P(A)}{P(B)}
\]

where P(A) and P(B) are respectively probabilities of event A and event B, P(A|B) presents the probability of A given B, P(B|A) is the probability of B given A.

B. K-nearest neighbors

K-nearest neighbors (KNN) is an algorithm that can be used for regression and classification, but which is more frequently used for classification predictive problems. KNN is based on the principle that instances with similar properties exist generally in close proximity. The class value of the unclassified instance can be predicted by observing the class of its nearest neighbors. The KNN finds the only adjustable parameter in this algorithm, the K nearest neighbors to the query instance and classify it by determining the most frequent class label. The model can be made less or more flexible by modifying K.

C. Logistic Regression

Logistic regression known as logit model estimates the relationship between one dependent variable and independent variables. It predicts the probability of an event utilizing a logit function.

The logistic regression is a linear regression of the explanatory variables \(x_i\):

\[
a_0 + a_1 \cdot x_1 + a_2 \cdot x_2 + a_3 \cdot x_3 + \ldots + a_k \cdot x_k \approx y,
\]

where the variable \(y\) could be (1) binary: only two possible outcomes such as good or bad. (2) Multinomial: three or more non-ordered categories such as single, married or divorced. Or (3) ordinal: three or more ordered categories such as food quality from 1 to 10.

D. MultiLayer Perceptron

Multilayer perceptron is a neural network [12] that consists at least of one hidden layer of nodes other than the input and output layers. This means, the layers of MLP should be a minimum of 3 layers including hidden, input and output. Each node of the layer, excluded the input, is called neuron. The neuron, a processing component, which is the main element of the neural network, collects information from a certain number of inputs, applies a weight, add a bias term and send the result to an active function which generates an output.

A multilayer perceptron model consists mainly of a linear activation function of all the neurons and backpropagation process for the training. The activation function maps the weighted inputs to the output of the neuron: it combines the input of the neuron with the weights, then adds a bias in order to generate the output. On the other hand, the backpropagation is supervised learning process that occurs with a continuous adjustment after each processing (based on the error in output) of the weights of the connections.

The backpropagation consists of two parts:

- Forward pass where the expected outputs that corresponds to given inputs are evaluated;
- Backward pass where partial derivative of the cost function is propagated through the network.

E. Support Vector Machine

SVM is a supervised machine learning algorithm, which is based on the concept of classification hyperplane, created to isolate between positive and other cases allowing for maximization. It plots each instance as a point in a \(n\)-dimensional space where \(n\) is the number of features (if the number of features is only 2, the hyperplane will be a simple line).

Moreover, the classification is done by finding the hyper-plane that best segregate the two classes with a maximum distance between points of both classes. Support Vectors are the closer points for hyperplane used to maximize the margin between hyperplane and data points. Also, they affect the orientation and position of the hyperplane.

F. Classification and Regression trees

Decision trees are used to explain a value from a series of discrete or continuous variables. These are fairly efficient, non-parametric and non-linear methods of partitioning individuals, producing groups of individuals as homogeneous as possible from the point of view of the variable to be predicted, taking into account a hierarchy of the predictive capacity of the variables considered. This hierarchy makes it possible to visualize the results in a tree, and to constitute explicit explanatory rules.

Several iterations are necessary. To each of them:

1) individuals are divided into \(k (=2)\) classes, to explain the output variable;
2) the first division is obtained by choosing the explanatory variable that will best separate individuals;
3) this division defines sub-populations, represented by nodes of the tree;
4) each node is associated with a proportion measure, which makes it possible to explain the belonging to a class or the meaning of an output variable;
5) the operation is repeated for each sub-population until no further separation is possible.

A decision tree can very quickly lead to overfitting, so it is necessary to prune the tree: stop at an adequate number of leaves when making the tree. In order to implement
all this, three main algorithms exist: CART, C4.5, and CHAID [7], [11], [16]. They proceed as follows:

- Choice of the decision variable:
  - measure of the $\chi^2$ difference to independence for CHAID and the $t$ of Chypruw;
  - Gini index (or split criterion) for CART
  - entropy for C4.5.

- Adjusting the size of the tree:
  - post-pruning for CART and C4.5: the purest tree is made with all the segmentation, then a criterion is used to compare trees of different sizes.
  - pre-pruning for CHAID: a stop rule is set to stop the construction.

G. Random Forest

Decision trees have the following main flaws: performance too heavily dependent on the initial sample, and a tree topology that can change completely with the input of some additional observations.

To overcome these problems, several trees are used. And to avoid having equal trees, randomness is added: each tree has a fragmented vision of the problem, randomly drawn from the input observations, and from the explanatory variables. More precisely, the assembly of decision trees built on the basis of a random draw among the observations is the tree bagging algorithm. The random forests (RFs), proposed by Leo Breiman [3], [9], add a feature sampling to the tree bagging [10].

H. AdaBoost

While RFs build several trees in parallel, boosting also builds $k$ trees (or other basic algorithms [6]), but it does so in series. The $k+1$ tree will have access to its predecessor, or more precisely to the latter's error: it will concentrate its effort on correcting these errors. For a classification problem, prediction is no longer a majority vote, but a weighted sum of each of the weak algorithms.

The first implementation of boosting, proposed by Yoav Freund and Robert Shapire, is called AdaBoost (adaptive boosting [19]), which starts from the following idea: a meta-algorithm works successively weak algorithms, each having access to a different distribution of the problem, focusing on observations that are difficult to process, and thus forcing its successor to treat them correctly. The term boosting refers, in a broad sense, to methods operating on this principle of serial assembly of weak learners.

In the case of RF, weak learners are unit decision trees, built in a totally independent way. Each algorithm has the same importance for the final vote. Boosting is a little less democratic, and achieves a sum weighted by the final vote. The weighting coefficients $\alpha_i$, in AdaBoost, depend only on the errors $\epsilon_i$ of each weak learner, as follows: $\alpha_i = \frac{1}{2} \ln \left( \frac{1-\epsilon_i}{\epsilon_i} \right)$. And at the AdaBoost level, the weak learner $i+1$ receives a different distribution of the data than the $i$-th, the latter's errors having been overweighted.

III. RELATED WORK

Despite all the efforts made by researchers, it may never be possible to know the exact time of earthquakes because there is some random component in such a catastrophe. In this field of research, errors are obviously predominant, even if exceptions exist. T. Mastuwaza et al. [14], researchers in the center for prediction of earthquakes and volcanic eruptions in Japan, compared for instance the recurrent seismograms every 5.3 $\pm$ 0.53 years and summed up their study by predicting that an earthquake will occur on November 2001 in Sanriku (Japan) with a probability larger than 0.99. And actually, as expected, on 13 November 2001 an earthquake of magnitude 4.8 happened.

Many machine learning based earthquakes research studies have been done with various motivations and goals such as determining the occurrence of earthquake, predicting the magnitude, the time, the location, detection of damage, and many other classification problems and methods. A technique done by A. Negarestani, 2002 [4] using layer neural network estimated the radon concentration in soil related to the environmental parameters. The change of soil radon is not only an earthquake precursor (anomaly phenomena in the earth) but is also controlled by the environmental parameters such as temperature, humidity, rainfall, etc. The data was obtained from a location in Thailand, processed using neural network of two hidden layers and analyzed. The result of this study indicates the ability to distinguish between time variation in radon concentration raised by earth anomaly phenomena such as earthquake and those caused by environmental parameters. This technique, in comparison with linear computational methods, can estimate better the radon variations related to environmental parameters.

Another investigation done by Rout-Leduc et al. in 2017 [18] emphasized that predicting the magnitude and timing before an earthquake fails is a main goal for geoscientist. This was done by listening to acoustic hidden signals that precede lab quakes in a laboratory that resemble to the earth: they apply Random Forest model to a continuous time-series data in a laboratory where each tree will predict the time remaining before the next failure of lab quake based on statistical features derived from time windows. In comparison with Naive Bayes model that gives a performance of 0.3, the RF achieves an accuracy of 0.89 which is largely better.

A. Cooner et al. [5] made a study after the earthquake happened on January 2010 near Port-au-Prince, Haiti, 7.0 moment magnitude with 316,000 deads [15], in order to detect the damage caused by this devastating event by applying Random Forests (400 trees), Neural Networks (two hidden layers of 20 neurons each) and Radial Basis Function Neural Network (150 Gaussian functions) algorithms. The remote sensing data was sourced from DigitalGlobe Foundation and the goal was to compare between a pre-disaster image captured in 2009 and a post-disaster image after the earthquake. Damaged and undamaged pixels were used for training datasets, then preprocessed. The accuracy found was best in RBFNN with 77.26%. ANN and RF, for their part, had accuracy of 74.14% and 76.14% respectively.

Another study achieved by K. M. Asim et al. [1] predicted the magnitude of earthquake in Hindukush region by using historic seismic activated with machine learning.
classifiers. They applied four machine learning algorithms on a dataset extracted from temporal distribution of past earthquake: Pattern recognition neural network, Recurrent Neural Network, Random Forest (50 trees) and linear programming boost ensemble. The accuracy to predict the earthquake occurrence in this study was best using Pattern recognition neural network or Linear Programming Boost Ensemble with 65% of accuracy. However, for Recurrent Neural Network and Random Forest, the scores were of 58% and 62% respectively.

J. Reyes et al. [17] presented in their study some neural networks to predict earthquakes in Chile in four different areas. The database was obtained from the Chiles National Seismological Service. They applied one neural network with 7 input neurons and 15 neurons in hidden layer, and then compared the prediction results with K-Nearest Neighbors (KNN), Support Vector Machine (SVM), and k-means. They finally found that none of the algorithms obtained a better result than their approach in more than one area.

A. Li and L. Kang [13] proposed a method called PR-KNN that combined between Polynomial Regression and K Nearest Neighbors models to predict the aftershock with magnitude greater or equal to 4.0. They selected the values in training sample based on the KNN algorithm, then these attributes have been modeled by applying polynomial regression method. The experimental data was collected from Wenchuan website. Finally, PR-KNN was compared with Distance-Weighted KNN regression and traditional KNN regression algorithms, which shown that the maximum relative error was reduced by 7.751% and 6.012% respectively.

IV. EXPERIMENTS

Further introduction for the world of machine learning, and explanations of the mathematical concepts for algorithms used in this paper are in [2].

A. Experimental protocol

In this study, a single time series set of data taken from an Earthquake data center in Northern California has been considered, where the first reading was on 1967 and the last in 2003. Each data point is an average reading for 1 hour. Let us notice that there is no use of future or past information while making the prediction. Each prediction employs only the data onto one single time series of this earthquake center. Thus, by analyzing the huge number of data point, the events were classified between negative and positive major earthquake. It is known that major events or major shocked are followed by small earthquakes in the same geographical area called aftershocks, and only reading above 5 on the Rictor scale are considered major event. The objective of this dataset is not to detect the aftershocks, but to classify between negative or positive earthquake major event. Negative output are instances having readings less than 4 and preceded, in the past 512 hours, by at least 20 non-zero readings.

On the other hand, a positive case is defined by a major event which was preceded, for at least 512 hours, by another major event.

The experiment is based on machine learning algorithms or in other terms, statistical learning theory that involves a training test (size 322) with associated inputs and outputs. Then, the trained model is utilized to evaluate a testing set (size 139) which is totally independent of the training set.

As stated previously, we used the following algorithms, for the sake of comparison:

- Random Forest (RF);
- Support Vector Machine (SVM);
- Logistic Regression (LR);
- Naive Bayes (NB);
- KNN;
- Multilayer Perceptron;
- AdaBoost;
- CART.

All the methods were applied on the training and testing sets using Matlab release 2018b and Weka, on a 2.7 GHz core i7 processor with 8 GB RAM. In obtained results of classification, several evaluation criteria have been considered:

- TP or true positive: number of times the algorithm classifies the event as 1 and it is actually 1.
- FP or false positive: number of times the algorithm classifies the event as 1 and it is actually 0.
- TN or true negative: number of times the algorithm classifies the event as 0 and it is actually 0.
- FN or false negative: number of times the algorithm classifies the event as 0 and it is actually 1.
- Mean absolute error:

  \[
  MAE = \frac{1}{n} \sum_{j=1}^{n} |y_j - x_j|
  \]

  where \( n \) is the number of all the values, \( y \) is the actual value, \( x \) is the predicted value.

- Root mean squared error:

  \[
  RMSE = \sqrt{\frac{1}{n} \sum_{j=1}^{n} (y_j - x_j)^2}
  \]

  Prediction value: Percentage of correctly classified instances.

B. Obtained results

For all algorithms, we tried to change the batch size, but the score was not affected by this parameter.

1) Naive Bayes: By applying Naive Bayes algorithm, 66.9% of instances were classified correctly, however 33.1% were not accurate.

The prediction of the major event was the worst between all the algorithms tested with 62.58%. 52 instances were classified wrong out of 139.

2) K-nearest neighbors: We applied for the KNN algorithm different number of K (neighbors) on the earthquake dataset as shown in Table I (MAE stands for Mean Absolute Error, while RMSE is for Root Mean Squared Error). The convergence of percentage of prediction occurs after using 5 neighbors. The highest accuracy reached goes for 3 neighbors with 75.53% (Figure 1) of right instances classified.
3) Logistic Regression: The prediction of the major event was the worst between all the algorithms tested with 62.58% where 52 instances were classified wrong out of 139. However, after changing the ridge to 1, the accuracy increases to 68.34% and the number of wrongly classified instances becomes 44.

4) MultiLayer Perceptron: We use in this experiment two hidden layers with three different couple of nodes: (10,20), (20,30) and (40,50). The comparison between the prediction value, MAE and RMSE using these different couples of nodes is illustrated in Table II. After choosing (20,30) as the number of nodes for two hidden layers, we apply many tests on other parameters and realize that the best score for prediction and MAE was found by using 20 epochs (Figure 2), by setting learning rate to a value of 0.6 (Figure 3), the momentum to a value of 1 (Figure 4) and by using sigmoid activation function. After changing the values of the batch size, we found that this parameter does not have any impact on the results of the experiment.

5) Classification and Regression Trees: CART algorithm leads to 70.5% of accuracy while 28 instances were not classified correctly.

6) Support Vector Machine: In our experiment, we used two different functions in SVM: PolyKernel and Normalized Poly Kernel with 64.02% and 74.82% accuracy respectively as shown in Table III.

7) Random Forest: Different number of trees are applied on the dataset. Table IV and Figure 5 show the statistical features for each number of trees. The best percentage of prediction is 76.97 using 3 trees and after reaching 100 trees, the algorithm converges and reaches a stable prediction of 74.82%.

8) AdaBoost: By applying AdaBoost on the earthquake dataset, 35 instances were classified wrongly and 72.66% of instances are predicted correctly.

<table>
<thead>
<tr>
<th>Nb of nodes</th>
<th>Prediction</th>
<th>MAE</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>10,20</td>
<td>73.38</td>
<td>0.2937</td>
<td>0.4713</td>
</tr>
<tr>
<td>20,30</td>
<td>71.22</td>
<td>0.3226</td>
<td>0.4853</td>
</tr>
<tr>
<td>40,50</td>
<td>70.50</td>
<td>0.3168</td>
<td>0.4860</td>
</tr>
</tbody>
</table>

TABLE II
STATISTICAL FEATURES USING MLP

<table>
<thead>
<tr>
<th>Method used</th>
<th>MAE</th>
<th>RMSE</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>PolyKernel</td>
<td>0.3597</td>
<td>0.5998</td>
<td>64.02%</td>
</tr>
<tr>
<td>Normalized Poly Kernel</td>
<td>0.2518</td>
<td>0.5018</td>
<td>74.82%</td>
</tr>
</tbody>
</table>

TABLE III
COMPARISON OF STATISTICAL FEATURES BEFORE AND AFTER NORMALIZATION OF KERNEL FUNCTION IN SUPPORT VECTOR MACHINE
Table IV

**STATISTICAL FEATURES FOR DIFFERENT NUMBER OF TREES FOR RANDOM FOREST ALGORITHM**

<table>
<thead>
<tr>
<th>Nb of trees</th>
<th>Prediction</th>
<th>MAE</th>
<th>RMSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>74.82</td>
<td>0.3108</td>
<td>0.4313</td>
</tr>
<tr>
<td>2</td>
<td>74.82</td>
<td>0.3237</td>
<td>0.4379</td>
</tr>
<tr>
<td>3</td>
<td>76.97</td>
<td>0.312</td>
<td>0.4161</td>
</tr>
<tr>
<td>4</td>
<td>74.1</td>
<td>0.3123</td>
<td>0.4233</td>
</tr>
<tr>
<td>5</td>
<td>72.66</td>
<td>0.3373</td>
<td>0.4417</td>
</tr>
<tr>
<td>10</td>
<td>69.06</td>
<td>0.3313</td>
<td>0.4375</td>
</tr>
<tr>
<td>100</td>
<td>74.82</td>
<td>0.3297</td>
<td>0.4193</td>
</tr>
<tr>
<td>200</td>
<td>74.82</td>
<td>0.3186</td>
<td>0.4122</td>
</tr>
</tbody>
</table>

Fig. 5. Percentage of prediction for different number of trees using Random Forest Algorithm

C. Comparing results of all algorithms

For each algorithm, we collected the True Positive, False Positive, True Negative, and False Negative rates, together with Mean Absolute Error, Root Mean Squared Error and Percentage of accuracy in Table V.

Table V

**STATISTICAL FEATURES COMPARISON BETWEEN 8 ALGORITHMS**

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>TP</th>
<th>FP</th>
<th>TN</th>
<th>FN</th>
<th>MAE</th>
<th>RMSE</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF</td>
<td>5</td>
<td>2</td>
<td>102</td>
<td>30</td>
<td>0.312</td>
<td>0.4161</td>
<td>76.97</td>
</tr>
<tr>
<td>NB</td>
<td>9</td>
<td>20</td>
<td>84</td>
<td>26</td>
<td>0.3257</td>
<td>0.5548</td>
<td>66.90</td>
</tr>
<tr>
<td>LR</td>
<td>6</td>
<td>29</td>
<td>89</td>
<td>15</td>
<td>0.3851</td>
<td>0.3161</td>
<td>68.34</td>
</tr>
<tr>
<td>MLP</td>
<td>0</td>
<td>35</td>
<td>104</td>
<td>0</td>
<td>0.2518</td>
<td>0.5018</td>
<td>74.82</td>
</tr>
<tr>
<td>AdaBoost</td>
<td>14</td>
<td>14</td>
<td>90</td>
<td>24</td>
<td>0.2958</td>
<td>0.4177</td>
<td>72.66</td>
</tr>
<tr>
<td>KNN</td>
<td>2</td>
<td>1</td>
<td>103</td>
<td>33</td>
<td>0.3002</td>
<td>0.4565</td>
<td>75.53</td>
</tr>
<tr>
<td>SVM</td>
<td>0</td>
<td>104</td>
<td>35</td>
<td>0.2518</td>
<td>0.5018</td>
<td>74.82</td>
<td></td>
</tr>
<tr>
<td>CART</td>
<td>7</td>
<td>13</td>
<td>91</td>
<td>28</td>
<td>0.3078</td>
<td>0.4607</td>
<td>70.50</td>
</tr>
</tbody>
</table>

The algorithm that performs better in term of accuracy is Random Forest with 76.97% (see Figure 6), which is very close to KNN with 75.53% and both MLP and SVM with 74.82%.

SVM, MLP, and AdaBoost perform between 72.66% and 74.82%, while 70.5% was the prediction for CART. On the other hand, MultiLayer Perceptron and Support Vector Machine are the algorithms having the minimum average of the errors in the prediction set with 0.25 as value of MAE (Figure 8). However, the algorithms giving the worst prediction percentage were Naive Bayes and Logistic Regression with 66.9% and 66.9% respectively. The comparison between TP, FP, TN, and FN is proposed in Figure 7.

V. Conclusion

Eight machines learning algorithms have been tested for our work to classify the major earthquake events between negative and positive. The study has been applied to a dataset collected from a center in California, which was recording inputs for 36 years. Every machine learning technique shows different results from each other. KNN, Random Forest and MLP are the best by producing the least false output (FP) while SVM, KNN, MLP and Random Forest classify the higher number of output correctly. Future work involves working on case studies based on
real intervention data and add consideration to features selection methods.

REFERENCES


