

FYS-STK3155 Project 3

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Abstract

In the attempt to find out whether we are alone in the universe or not, we take a look at other stars and try to fit a model to predict exoplanets. We use stars' flux observations with a binary classification of 0 and 1, labelling whether the stars has an exoplanet or not. In this project we have developed four different machine learning models; logistic regression, neural network, decisions trees and gradient boosted trees. We compare these models in their ability to detect exoplanets, under optimized circumstances. We were able to predict exoplanets with an accuracy of 98.1% using logistic regression, 99.8% using a neural network, 98.4% using a decision tree, and 100% using a gradient boosted tree with the XGBoost framework. We have showed the gradient boosting XGBoost to be the best performing model for such binary classification problems.

I. INTRODUCTION

Two possibilities exist: either we are alone in the Universe or we are not. Both are equally terrifying.

Arthur C. Clarke

In this project we will use machine learning methods to look for exoplanets. The best place to look for extraterrestrial life is on other planets. There are thousands of planets orbiting other stars in the universe, called exoplanets. But, given the large distance in our universe it makes it difficult to simply "look" for planets. There are many features and methods in astronomy that could be used to detect these exoplanets. One of which is using the flux dip of the star.

The data set we will be using contains the flux of 5657 stars in which only 42 stars have exoplanets orbiting them (Winterdelta, 2017). The flux data can be used to detect whether a exoplanet is orbiting the star, this is called the exoplanet transit method (Wilson, 2017). We will use this data in different machine learning methods and see what methods is best suited for detecting exoplanets. The methods we will use with the dataset is Logistic regression, Feed Forward Neural Network (FFNN), decision tree, and extreme gradient boosted decision tree.

II. THEORY

A. Dataset

The data is from NASA's Kepler space telescope. The cameras of the satellite stare at one area of space for 80 days (Winterdelta, 2019). Thus for 80 days light of one area of space is collected. We can use this continuous data sequence and look for dips in the light flux

from stars. This is called the exoplanet transit method (Wilson, 2017). Essentially, the dip in flux from a star could mean that there is a planet orbiting it. However, it could also mean its another object passing, maybe a very large asteroid, or maybe even disturbances in the camera. Nevertheless, we will use this method to train our different machine learning models and eventually see how well the methods are able to learn whether a flux dip implies an exoplanet in orbit.

The data set is formatted as a table/matrix where each row is a data sample (a star) and each column/feature is a flux measurement for that row's star. The data contains 3197 such features for each star. The total data set has a total of 5657 stars where 42 of them have a confirmed exoplanet in orbit. The data set comes pre-split into a training set with 5087 stars where 37 of them have a confirmed exoplanet, and a testing set with 570 stars where 5 of them have a confirmed exoplanet.

B. Scaling

We apply a standardization scaling to our data set. This is a scaling method that, for each feature, subtracts the whole feature by its mean, then divides it by the standard deviation:

$$x_{ij} = \frac{x_{ij} - \bar{x}_{:j}}{\sigma(x_{:j})}$$

where $\bar{x}_{:j}$ and $\sigma(x_{:j})$ are respectively the mean and the standard deviation of the feature column j defined as

$$\bar{x}_{:j} = \frac{1}{n_{rows}} \sum_i^{n_{rows}} x_{ij}$$
$$\sigma(x_{:j}) = \frac{1}{n_{rows}} \sqrt{\sum_i^{n_{rows}} (x_{ij} - \bar{x}_{:j})^2}$$

This method ensures that each feature has a mean value of zero and a standard deviation of one.

C. Measuring prediction performance

C1. Confusion matrix

The confusion matrix is a matrix which shows the four major prediction scores: the false negative, true negative, false positive, and true positive. In this sense a negative prediction is one that predicts the class 0 (no exoplanet), and positive is the prediction of 1 (exoplanet). We choose to normalize our matrix resulting in the confusion matrix being defined as

$$\text{Confusion matrix} = \frac{1}{n} \begin{bmatrix} TN & FP \\ FN & TP \end{bmatrix} \quad (1)$$

where n is the total number of predictions, and each of the metrics are defined as below:

$$\begin{aligned} TN &= \text{true negative} = \frac{\# \text{ of correct negative predictions}}{\text{Total } \# \text{ of negative predictions}} \\ FN &= \text{false negative} = \frac{\# \text{ of false negative predictions}}{\text{Total } \# \text{ of negative predictions}} \\ TP &= \text{true positive} = \frac{\# \text{ of correct positive predictions}}{\text{Total } \# \text{ of positive predictions}} \\ FP &= \text{false positive} = \frac{\# \text{ of false positive predictions}}{\text{Total } \# \text{ of positive predictions}} \end{aligned}$$

C2. Accuracy

For classification problems the accuracy is defined as:

$$\begin{aligned} \text{Accuracy} &= \frac{\text{Number of correct predictions}}{\text{Total number of predictions}} \\ &= \frac{\sum_{i=1}^n I(t_i = y_i)}{n} \end{aligned} \quad (2)$$

where I is the indicator function that simply checks if they are equal:

$$I(t = y) = \begin{cases} 1, & t = y \\ 0, & t \neq y \end{cases} \quad (3)$$

this can also be written out as

$$\text{Accuracy} = \frac{TN + TP}{FN + TN + FP + TP} \quad (4)$$

D. Logistic regression

Logistic regression is a suitable method to use for this particular problem of finding whether there is a planet in orbit or not. For each data point get squeezed between 0 and 1 through the Sigmoid function which is defined as (Hjorth-Jensen, 2023)

$$f(x) = \frac{1}{1 + e^x}, \quad (5)$$

The resulting values ranges from 0 to 1, the function is plotted in figure 3.

$$f(x) \rightarrow [0, 1] \quad (6)$$

E. Feed Forward Neural Network

For the neural network method we will test Feed Forward Neural Network (FFNN). A FFNN has an input layer which the data is fed into as a matrix \mathbf{X} , at least one hidden layer, and at last a output layer (Pladsen, Qanbari, & Vattøy, 2023). The basic schematic is illustrated in figure (1).

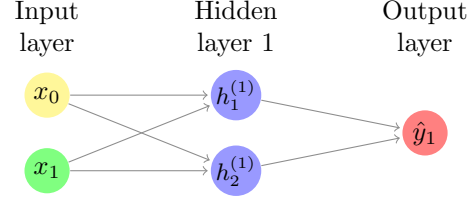


Figure 1: Neural network schematic.

In the hidden hidden layer the input data \mathbf{X} is multiplied with weights \mathbf{W}_l and a bias \mathbf{b}_l is added resulting in the equation (equation 7) (Pladsen et al., 2023).

$$\mathbf{z}_l = \mathbf{X}\mathbf{W}_l + \mathbf{b}_l \quad (7)$$

The output of the function \mathbf{z}_l in the hidden layer is passed through an activation function resulting in a function (equation 8)

$$\mathbf{a}_l = f(\mathbf{z}_l) \quad (8)$$

The activation function \mathbf{a}_l is then multiplied by weights in the output layer \mathbf{W}_L and a bias is added \mathbf{b}_L resulting in the function \mathbf{z}_L which could also be passed through an activation function depending on the problem at hand (Pladsen et al., 2023).

E1. Activation functions

Activation functions are an essential component of a neural network. Inspired from neurons in biological systems where neural signalling is dependent on the activation level: a neuron does not fire upon any activation, it requires a certain threshold before firing (Hjorth-Jensen, 2023). Activation functions in neural networks are set perform in a similar way. They add important features to the model, such as improving the training convergence of the network (Dubey, Singh, & Chaudhuri, 2021). Without activation functions, a neural network would be simply be a regression model (GeeksforGeeks, 2023). In other words, the activation functions do non-linear transformations on the input data that allows the network to learn the abstract features of the data (Dubey et al., 2021; GeeksforGeeks, 2023; Pladsen et al., 2023).

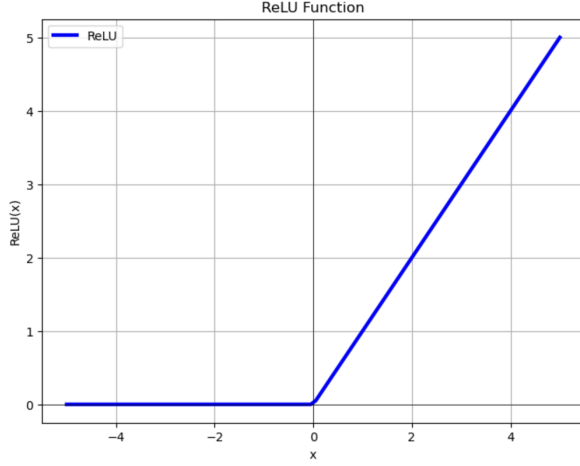


Figure 2: The ReLU function.

The ReLU (Rectified linear unit) function (figure 2) is one often used for activation and is defined as

$$f(x) = \begin{cases} x, & x > 0 \\ 0, & x \leq 0 \end{cases} \quad (9)$$

This activation provides an output that ranges from 0 to infinity

$$f(x) \in [0, \infty] \quad (10)$$

The ReLU function has a limitation which is the dying of neurons caused by negative values. If the amount of negative values are significant in the data set then, most of the data is set to zero in the activation function (Dorsaf, 2021; Dubey et al., 2021). However, as explored in (Pladsen et al., 2023) it proved to be computationally efficient and lead to high prediction accuracy.

The Sigmoid function (equation 5) is another widely used activation function, see its graph in figure 3. However, as explored in (Pladsen et al., 2023) the Sigmoid function has some limitations. One of which is the saturation of neurons when the activation becomes 0 and 1. The derivative of the Sigmoid function is:

$$f'(x) = f(x)(1 - f(x))$$

thus for activations that are 0 or 1 the derivative becomes zero and training of the neuron essentially stops (Dorsaf, 2021). Thus the Sigmoid function could hinder training and the calculation of optimal parameters.

The hyperbolic tangent function (tanh) is another one used for activation, it is plotted in figure 4 and is defined as

$$f(x) = \tanh(x) = \frac{2}{1 + e^{-2x}} - 1 \quad (11)$$

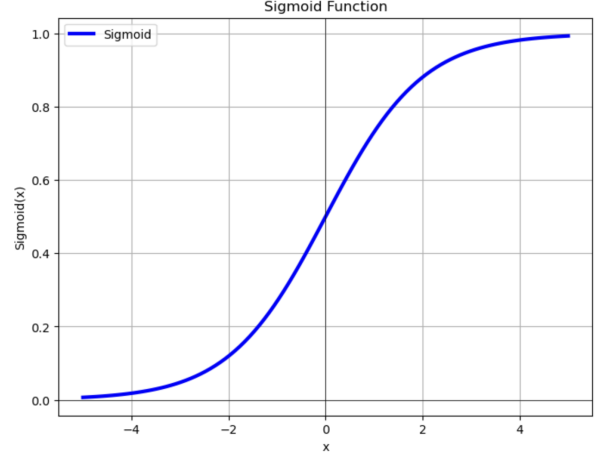


Figure 3: The Sigmoid logistic function

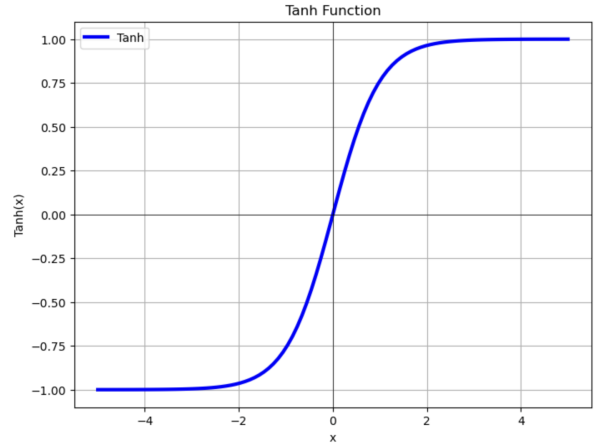


Figure 4: The hyperbolic tangent function

Its output has the range

$$f(x) \in [-1, 1] \quad (12)$$

The final activation function we wish to compare is no activation at all, that is to say

$$f(x) = x$$

this is often called the *Identity* activation function in relation to neural network code.

E2. Backpropagation

We use the algorithm known as backpropagation to train our feed-forward neural network. For a network we want to optimize the weights \mathbf{W} and the biases \mathbf{b} in the different layers by using the error for the output layer L generally defined as 13 and for a general layer l defined as 14. More about this can be found in (Pladsen et al., 2023), but Scikit-learn can also be used to easier analyse datasets.

$$\begin{aligned}\delta^L &= \frac{\partial C}{\partial \mathbf{z}} = \frac{\partial \mathbf{a}}{\partial \mathbf{z}} \frac{\partial C}{\partial \mathbf{a}} \\ &= f'(\mathbf{z}^L) \frac{\partial C}{\partial \mathbf{a}}\end{aligned}\quad (13)$$

$$\begin{aligned}\delta^l &= \delta^{l+1}(\mathbf{W}^{l+1})^T \frac{\partial \mathbf{a}^l}{\partial \mathbf{z}^l} \\ &= \delta^{l+1}(\mathbf{W}^{l+1})^T f'(\mathbf{z}^l)\end{aligned}\quad (14)$$

F. Decision tree

A classification decision tree, or just a classification tree, is a machine learning method that creates a binary tree structure of conditions to classify the input as one of the output classes.

As illustrated in figure (5) such a tree starts with a root node. Each node branches into two new nodes each, and the output nodes are called leaf nodes or just leaves. The middle nodes that are not a root node or a leaf are called internal nodes (Hjorth-Jensen, 2023; IBM, n.d.). The root node and internal nodes each contain a logical condition to classify which branch direction the data should continue in. The leaf represents an output class, which in our case it would be either "no exoplanet" or "exoplanet".

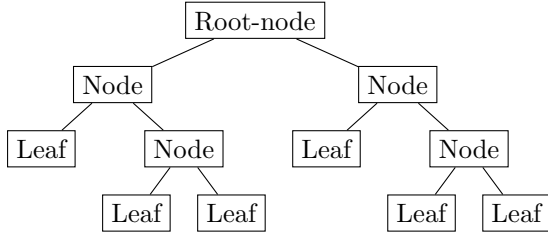


Figure 5: Schematic of a decision tree structure

Decision trees are prone to overfitting, therefore it is common to set a threshold for how deep the tree goes (IBM, n.d.). In general the node conditions can be yes/no, however for a data set such as ours with numerical flux observation, the conditions would be numeric.

F1. Node split criterion

The two most common methods of evaluating each node branching/splitting quality are the Gini impurity/index (equation 15) and the information entropy (equation 16) (Hjorth-Jensen, 2023). They both are based on the following probability distribution function

$$p_k = \frac{1}{n} \sum_{i=1}^n I(y_i = k)$$

where p_k represents the number of observations of a given class $k = 0, 1, 2, \dots, K$ from n total observations. For our binary classification problem we would have $k = 0, 1$ ($K = 1$).

The Gini index, also called the Gini impurity, measures the probability for a random instance being misclassified when chosen randomly (Dash, 2023). It is defined as (equation 15):

$$g = \sum_{k=1}^K p_k(1 - p_k) \quad (15)$$

The information entropy is defined as

$$s = - \sum_{k=1}^K p_k \log p_k \quad (16)$$

The information entropy bases a split on the information gain, which is a measure of the purity of the possible classification (Dash, 2023). This is based on thermodynamic entropy from physics and is suited well for decision trees, because it bases the splits on whether or not it causes an information gain (a more pure classification) (Dash, 2023). The entropy values are between 0 and 1 (IBM, n.d.), where 0 is low entropy and 1 is high entropy. Thus the goal is make a classification based on low entropies.

F2. Finding optimal depth

Starting from the the root node and splitting the three downward can make a large tree with many branches. How large could the tree become? In theory the splitting could go on until all branches represent a classification (Hjorth-Jensen, 2023). However, for increasing tree complexity the classification becomes less pure, meaning fewer data points end up branches and have a lower purity (IBM, n.d.).

In order to avoid overfitting we could find the optimal depth with respect to the accuracy of the model. It is also possible to develop a complex tree and then prune the tree back a few depths. In our model we test various depths and use a grid search for the optimal depth and regularization term.

G. Gradient boosting

Gradient boosting is a method combining iterative gradient descent with simple decision trees. It uses an ensemble of so-called weak predictors or weak learners, which in this case would be a simple decision tree, usually with a low depth. For each iteration it creates a new tree based on the previous tree's result, with a learning rate η (gradient descent).

Learning from the ensemble of weak predictors, the final gradient boosted decision tree usually outperforms a simple non-boosted tree. We choose a number of boosting rounds and a maximum tree depth for the final result.

III. METHODS

As mentioned our star dataset (section A) is already split into a training and testing set, however we need to preprocess it. The stars with a confirmed exoplanet are labeled with the label 2, while the stars without confirmed exoplanets have the label 1. We first change these labels to 0 and 1 so we can easily use the Sigmoid function to get the prediction outputs between $[0, 1]$. The label 1 means the star has a confirmed exoplanet.

The two classes are also extremely unbalanced; the training set contains 5050 non-exoplanet stars, but only 37 confirmed exoplanet stars. This will heavily impact our training and predictions. To solve this we use a technique called Synthetic Minority Over-sampling Technique (SMOTE) (Chawla, Bowyer, Hall, & Kegelmeyer, 2002). For this we use the Python library `Imbalanced-learn`: `imblearn.over_sampling.SMOTE`. This balances out the number of occurrences of each class. We chose to do this to just the training set and then re-split that into a new training and testing set. One could also include the original testing data by combining the data before doing this preprocessing. We use Scikit-learn’s `sklearn.model_selection.train_test_split` with a testing size of 20% for our split.

For each of the different classification methods we try out scaling the data vs not scaling before training, to see if it improves performance or not. We use Scikit-learn’s standardization implementation `sklearn.preprocessing.StandardScaler`.

We use the accuracy (equation 2) to evaluate the prediction performances using Scikit-learn’s `sklearn.metrics.accuracy_score`. In addition, we plot the confusion matrices (equation 1 for the results using `sklearn.metrics.confusion_matrix`). These are applied to the prediction on the testing set after being trained on the training set.

A. Logistic regression

From Scikit-learn we use `sklearn.linear_model.LogisticRegression`, and apply a L2 regularization penalty term λ with 1000 epoch iterations. To find out if scaling the data improves the prediction we start off with $\lambda = 10^{-4}$.

We then plot the testing set accuracy with respect to λ to find the optimal value. Here we choose to use 100 epochs to reduce the computation time. Using the found optimal value we then do our final prediction with 1000 epochs.

B. Classification neural network

From Scikit-learn we now use `sklearn.neural_network.MLPClassifier` for our classification network. We use the ADAM method (Pladsen et al., 2023) for scaling the learning rate.

The parameters we need to optimize are the L2 regularization λ , the initial learning rate η_0 , the activation

function, the number of batches $n_{batches}$, number of hidden layers n_{layers} and the number of nodes in each hidden layer n_{nodes} . We can write the last two as a merged vector where each element is the number of nodes for that hidden layer:

$$\mathbf{n}_{nodes} = (n_{nodes, \text{layer 1}}, n_{nodes, \text{layer 2}}, \dots)$$

The number of nodes for the input layer is the number of features in the training set, which is 3197, and for the output layer we only have one classification label (either 0 or 1) so there is only one node.

Initially to test if scaling improves performance we use the following parameters and architecture

$$\begin{aligned} \lambda &= 0.001 \\ \eta_0 &= 0.1 \\ n_{batches} &= \text{auto (Scikit-learn)} \\ \text{activation} &= \text{ReLU (equation 9)} \\ n_{epochs} &= 100 \\ \mathbf{n}_{nodes} &= (10) \end{aligned}$$

Afterwards we try out different numbers of hidden layers in our network. Then, we test the activation functions other than ReLU, which is: Sigmoid (equation 5), Identity (equation 1), and finally tanh (equation 11). Furthermore, we do a grid search plotting accuracy to η_0 and λ . Lastly, we do a similar grid search for $n_{batches}$ and \mathbf{n}_{nodes} for the optimal number of layers.

Using the optimal parameters we do our final prediction with 1000 epochs.

C. Classification tree

From Scikit-learn we use `sklearn.tree.DecisionTreeClassifier` for our classification tree.

Initially we use a max tree depth of 5 using the Gini index (equation 15). Afterwards we test the information entropy (equation 16) method as the criterion.

Using the optimal splitting method we plot the accuracy as function of the maximum tree depth, then use that found value to make our final prediction.

D. Gradient boosted classification tree

We use the extreme gradient boosting Python library `XGBoost` and use the `xgboost.XGBClassifier` for the boosted tree.

Initially we use a maximum tree depth of 5 with 100 boosting rounds to test out scaling, with no regularization. Afterwards we create a grid plot to find the optimal values for these two parameters. We also want to find the optimal λ for L2 regularization if it improves the performance, we use these optimal parameters to make our final prediction.

IV. RESULTS

A. Logistic regression

For the chosen initial parameters, we found that scaling does improves the accuracy:

No scaling : 0.899

Scaling : 0.979

We then plot the accuracy with respect to the L2 regularization λ shown in figure 6, we found the optimal value to be

$$\lambda = 10^{-5}$$

Making our final prediction we conclude the accuracy of the logistic regression method to be

Accuracy = 0.981

with the confusion matrix plotted in figure 7.

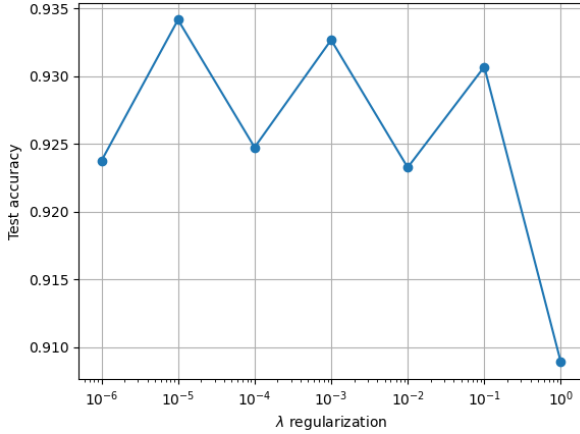


Figure 6: Logistic regression accuracy plotted as a function of the L2 regularization λ .

B. Classification neural network

For the chosen starting configuration (see methods B) we found that scaling the data is better: the accuracies with ReLU activation and $\mathbf{n}_{nodes} = (10)$ were

No scaling : 0.930

Scaling : 0.990

We then found increasing the number hidden layer from one decreased this accuracy (with scaling) to

$\mathbf{n}_{nodes} = (10, 10, 10)$: 0.523

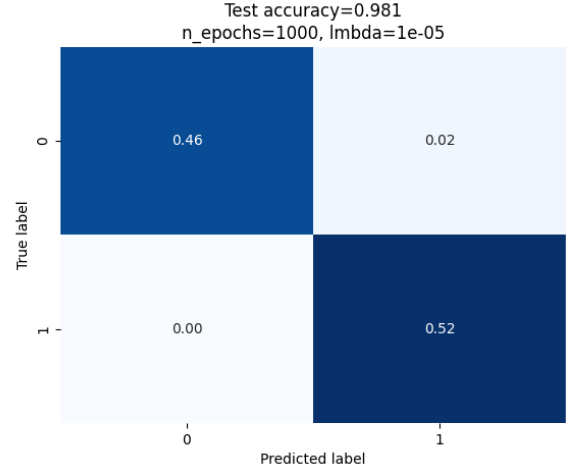


Figure 7: Confusion matrix of the logistic regression's final prediction.

We then tried out the different activation functions other than ReLU and found with $\mathbf{n}_{nodes} = (10)$ the following accuracies

Identity : 0.801

Sigmoid : 0.533

Tanh : 0.528

meaning ReLU showed the best performance by far.

The grid search for optimal η_0 and λ values is plotted in figure 8 where we found the optimal

$$\eta_0 = 0.01$$

$$\lambda = 10^{-5}$$

The final grid search to find the optimal number of batches and nodes is plotted in figure 9 where we found the optimal values as

$$n_{batches} = 80$$

$$\mathbf{n}_{nodes} = (40) \text{ (one layer with 40 nodes)}$$

Using these found optimal parameters our final prediction for this neural network method proved to be

Accuracy = 0.998

The confusion matrix is plotted in figure 10.

C. Classification tree

For the classification (decision) tree we start using the Gini index with a max tree depth of five which results in the accuracies

No scaling : 0.940

Scaling : 0.939

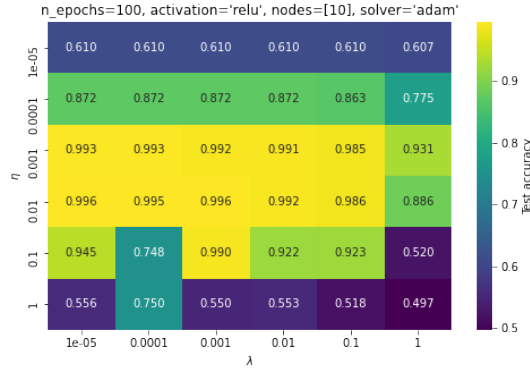


Figure 8: Neural network grid search to find the optimal values for the parameters initial learning rate η_0 and L2 regularization term λ .

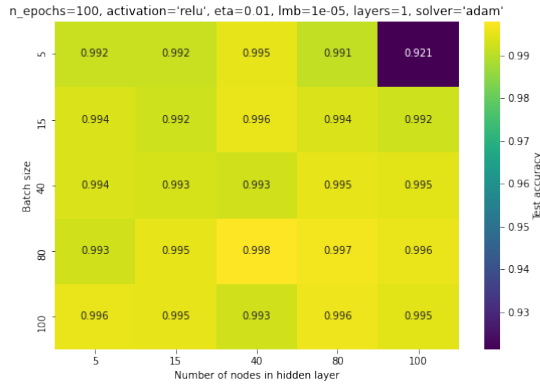


Figure 9: Neural network grid search to find the optimal values for the parameters batch size $n_{batches}$ and number of nodes n_{nodes} in the one hidden layer.

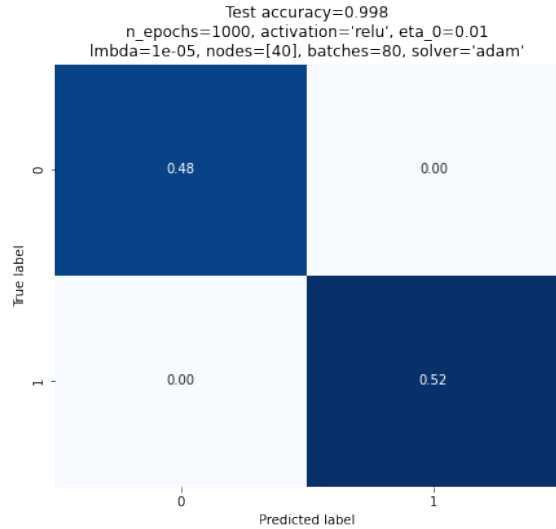


Figure 10: Confusion matrix of the neural network's final prediction.

showing that in this case there is barely any difference when scaling. We move on without scaling replacing the criterion with entropy (equation 16) which showed slightly worse performance:

Entropy : 0.934

Using the Gini index with no scaling we find the optimal maximum depth considering both computational time and performance to be 30, the analysis is plotted in figure 11.

Using this configuration we obtain the final prediction accuracy

Accuracy = 0.984

where the confusion matrix is plotted in figure 12.

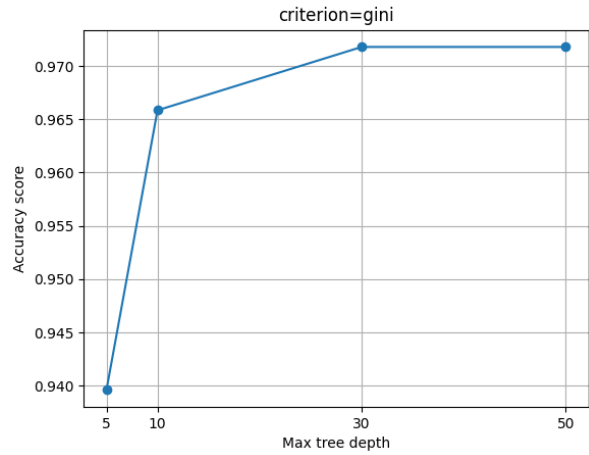


Figure 11: The classification tree accuracy plotted as a function of the maximum tree depth. Anything more than 30 depth proved to not increase the accuracy. It is possible that the phenomena of overfitting can be shown by extending the x-axis, however the computational time is already huge for these values.

D. Gradient boosted classification tree

Using the XGBoost classification tree with depth = 5 and 100 boosting rounds we immediately obtained the accuracies

No scaling: 1.0000

Scaling: 0.9995

showing that no scaling is also slightly better in this case.

This accuracy is perfect, we still plot a grid search to see how the accuracy varies (figure 13). We find that we keep our perfect accuracy with reduced computation time by using 30 boosting rounds with a tree depth of 5. We decided there was no reason to further tweak parameters as we already achieved perfect accuracy.

The final prediction confusion matrix is plotted in figure 14.

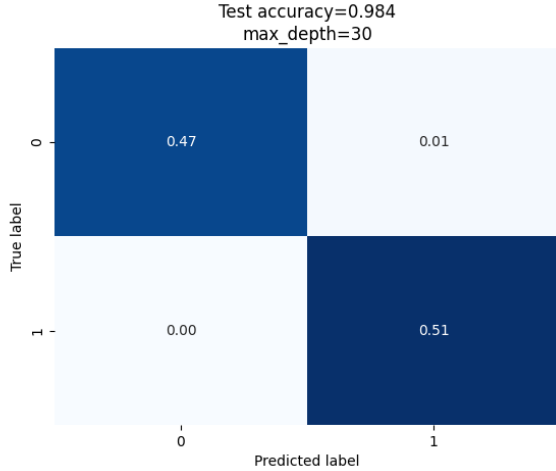


Figure 12: Confusion matrix of the classification tree’s final prediction.

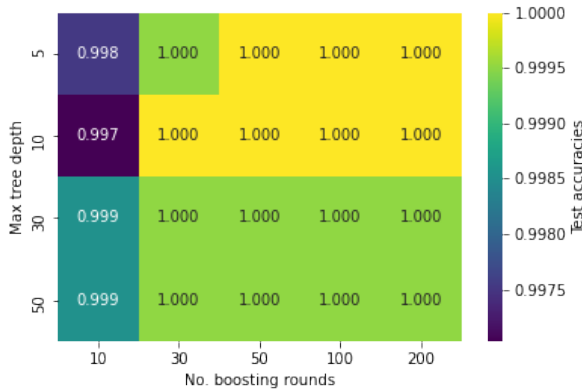


Figure 13: XGBoost tree grid search to find the optimal number of max tree depth and number of boosting rounds.

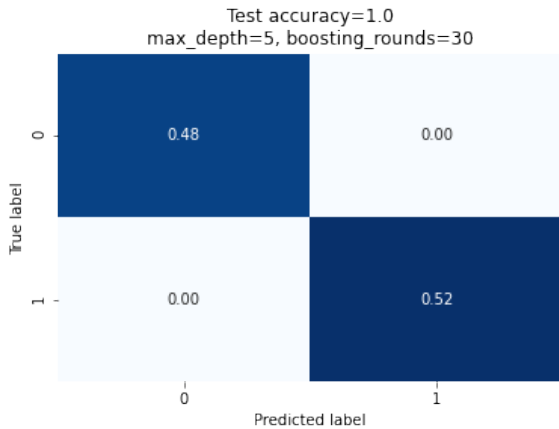


Figure 14: Confusion matrix of the XGBoost tree’s final prediction.

V. DISCUSSION

A. Logistic regression

We performed an analysis using logistic regression on the dataset with and without scaling the training data before training. We found that scaling improved the prediction performance by 9% (absolute difference). We did not predict this as we had been warned that usually one does not use scaling for binary classification purposes like these, however for the simple regression it seemed to help. In addition, the final prediction had a higher accuracy than we originally expected from the simplest method in this comparison, at 98% correct.

From the λ -plot in figure 6 we see that the general trend shows that the lower values were better, however it fluctuated quite a bit so that the $\lambda = 10^{-5}$ was 1% better than the lowest tested value $\lambda = 10^{-6}$.

B. Classification neural network

We found that with ReLU activation, scaling improved the prediction accuracy by 6%. This accuracy (0.99) is already outperforming the simple logistic regression by 9%, but we can improve this even more by finding the optimal parameters.

Continuing with scaling, we saw that the Sigmoid and tanh activations performed surprisingly bad at around 53% accuracy. This is just slightly better than pure guessing, which we did not expect. We originally thought the Sigmoid function would perform similarly to ReLU, maybe slightly worse ((Pladsen et al., 2023)). Also unexpected, the identity activation (which was no activation function at all) performed second best after ReLU by a 19% reduction. Our expectation in order of thought performance was ReLU, Sigmoid, tanh, then identity but the results showed ReLU, identity, then Sigmoid and tanh basically useless activations.

The accuracy also decreased when increasing the number of hidden layers from one, which is why we focused on one layer throughout the rest of the method’s execution. This was somewhat expected from our previous project Pladsen et al. (2023), where increasing the number of layers did not automatically mean better performance.

The final prediction accuracy proved to be 1.7% better than logistic regression. This is what we expected because the regression method is pretty rudimentary/dumb. The accuracy difference was not that significant, which is reflected in the confusion matrices shown in figure 7 and 10.

C. Classification tree

We used `sklearn.tree.DecisionTreeClassifier` to classify the data. In this case scaling vs no scaling has almost no impact, scaling reduced the accuracy by only 0.1%. We found that the node splitting criterion performed al-

most the same, the optimal being the gini index which performed 0.7% better than the entropy criterion.

Plotting the accuracy as a function of the tree depth we saw that increasing the depth increased the accuracy up to a certain point, then it became stagnant. We expect the accuracy to eventually begin dropping as the depth increases more as a sign of overfitting, however the computational time was already so high that we stopped at the depth of 50 layers.

This method surprisingly performed worse than the neural network, we initially expected this to be better for a binary classification problem like this. It performed 1.4% worse than the neural network, and only 0.3% better than the logistic regression.

D. Gradient boosted classification tree

The XGBoost tree was by far the best predictor for our data set with a perfect accuracy of 1 or 100% correct. In this case scaling the data actually slightly reduced this accuracy to 0.9995, which is only a 10^{-4} % difference. As seen in the grid search plot (figure 13), there were many different combinations giving a perfect prediction accuracy (this plot was rounded to three decimals so some of them rounded up to one). Most relevant for us it showed we could reduce our number of boosting rounds from 100 to 30 while still having perfect accuracy (rounded), saving computational time for the final prediction and eventual other predictions. Already having a perfect accuracy for multiple configurations, we found no further reason to keep tweaking the other XGBoost parameters like the learning rate, the regularization, the growing policy and many others.

This result is exactly as we expected, because we have read that the XGBoost algorithms show incredible results for classification purposes, and recently XGBoost is winning many machine learning competitions.

VI. CONCLUSION

Our main goal in this report was to compare different machine learning methods, and ultimately find the accuracy of predicting whether a star has an exoplanet or not. For the four different methods we found the following final accuracies:

Logistic regression: 98.1%

Neural network: 99.8%

Decision tree: 98.4%

XGBoost: 100%

This showed that XGBoost was the best predictor method, outperforming the second best neural network method with 0.2%. However, in general all methods performed pretty similarly; the biggest deviation of 1.9% from XGBoost was using the logistic regression method,

which is what you would expect from such a relatively simple method.

Appendix A. Github repository

<https://github.com/LassePladsen/FYS-STK3155-projects/tree/main/project3>

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