Evaluation of Machine Learning in Empirical Asset Pricing

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Abstract

Empirical Asset Pricing via Machine Learning

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Introduction

Motivations

Literature

Main Findings

Limitations of Machine Learning

Machine learning excels at prediction problems, namely estimating $E(r_{i,t+1}|\mathcal{F}_t)$, where $r_{i,t+1}$ is an asset's excess return over the risk free rate, and \mathcal{F}_t is the set of all information (including unobservable) available to market participants in this context.

This means that machine learning algorithms do not, nor do they aim to, explain how the market works in terms of underlying dynamics and equilibria. Though a machine learning algorithm may be able to identify patterns that otherwise cannot be easily found, an economist is still required to analyse these patterns to construct and hypothesize economic theory.

Methodology

Overall Design

Each model will be presented and explained so that a reader without any machine learning background can understand the basic idea behind the model. Computational methods such as algorithms however, will only have their general principles and background explained in the appendix. This is because there are many variations of algorithms available, and more importantly, specific understanding of how the algorithm works is not necessary.

All asset excess returns are modelled as an additive prediction error model:

$$r_{i,t+1} = E_t(r_{i,t+1}) + \epsilon_{i,t+1} \tag{1}$$

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where

$$E_t(r_{i,t+1}) = g^*(z_{i,t}) \tag{2}$$

Sample Splitting

Imperative to any machine learning technique is the establishment of how the dataset is to be split into training, validation and test sets. The training set is used to initially build the model and provide initial estimates of parameters, whereas the validation set is used to tune model parameters to optimise out of sample performance, thus preventing overfitting. Essentially, the validation set acts as a simulation of out of sample testing. The test set is used only for evaluation, and is thus truly out of sample.

There are three main approaches to splitting temporal data (such as financial data).

The first is to decide arbitrarily on a single training, validation and test set. This method is straightforward and the least computationally intensive, but is limited and inflexible in evaluating how models perform when more recent data is provided for training.

The second method is a "rolling window" method, where a fixed size or "window" for the training and validation set is first chosen. This window then incrementally move forwards in time to include more recent data, with a set of forecasts for the test sets made for all possible windows.

The third is a "recursive" method, which is the same as the rolling window method, but different in that the training set always contains previous data, with only the validation set staying fixed in size and "rolling" forwards. Hence, it is like a rolling window approach that has a growing training sample.

Both the rolling window and recursive schemes are very computationally intensive. Therefore, a hybrid of the methods where the training, validation and test samples are split, and hence models refit once each year. The training set is increased by one year each year, the validation set remains one year in length but moves forward by one year, and forecasts are only made using that model for the subsequent year. Cross validation was crucially not done to maintain the temporal ordering of the data.

Simple Linear Model

The least complex model considered is the simple linear regression model, otherwise known by its estimation method ordinary least squares (OLS). OLS struggles with high-dimensionality. Nevertheless, despite being expected to perform poorly it was implemented as a "control."

The simple linear model assumes that the underlying conditional expectation $g^*(z_{i,t})$ can be modelled as a linear function of the predictors and the parameter vector θ :

$$g(z_{i,t};\theta) = z'_{i,t}\theta \tag{3}$$

This model can capture non-linearities only if a new predictor set $z_{i,t}^*$ containing specified non-linear transformations or interaction terms. It is quite common to consider at least second order terms and two way interactions.

The baseline computational algorithm for this model is to minimize the standard least squares function:

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 (4)

Penalized Linear

Penalized linear models have the same underlying statistical model as simple linear models, only considering baseline untransformed predictors. They differ in their addition of a new penalty term in the loss function:

$$\mathcal{L}(\theta;.) = \underbrace{\mathcal{L}(\theta)}_{\text{Loss Function}} + \underbrace{\phi(\theta;.)}_{\text{Penalty Term}}$$
(5)

Several choices exist for the choice of the penalty function $\phi(\theta;.)$. This focus of this paper is the popular "elastic net" penalty, which takes the form:

$$\phi(\theta; \lambda, \rho) = \lambda(1 - \rho) \sum_{j=1}^{P} |\theta_j| + \frac{1}{2} \lambda \rho \sum_{j=1}^{P} \theta_j^2$$
(6)

The elastic net has two hyperparameters: λ , which controls the overall magnitude of the loss, and ρ , which controls the shape of the penalization. The $\rho=0$ case corresponds to the popular LASSO and uses absolute (l_1) parameter penalization, which geometrically allows the coefficients to be shrunk to 0. This allows it to impose sparsity, and can be thought of as a variable selection tool. The $\rho=1$ case corresponds to ridge regression, which uses l_2 that shrinks all coefficients closer to 0, but not actually to 0. Ridge regression is therefore a shrinkage method which prevents coefficients from becoming too large and overpowering. For $0 < \rho < 1$, the elastic net aims to produce parsimonious models through both shrinkage and selection.

The hyperparameters λ and ρ are both tuned using the validation sample. See appendix for algorithm.

Classification and Regression Trees

Classification and regression trees are fully non-parametric models that can capture complex multi-way interactions. A tree "grows" in a series of iterations. With each iteration, a split ("branch") is made along one predictor such that it is the best split available at that stage (in terms of lowering the loss function). These steps are continued until either the stopping criterion is met (such as via regularization) or each observation is its own node. The eventual model slices the predictor space into rectagular partitions, and predicts the unknown function $g^*(.)$ with the average value of the outcome variable in each partition.

The prediction of a tree, \mathcal{T} , with K "leaves" (terminal nodes), and depth L is

$$g(z_{i,t};\theta,K,L) = \sum_{k=1}^{K} \theta_k \mathbf{1}_{z_{i,t} \in C_k(L)}$$
(7)

For this study, only recursive binary trees (the most common and easy to implement) are considered. The popular l_2 impurity was also chosen as the loss function (conceptually similar to mean squared error):

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$$H(\theta, C) = \frac{1}{|C|} \sum_{z_{i,t} \in C} (r_{i,t+1} - \theta)^2$$
(8)

where |C| denotes the number of observations in set C (partition). Given C, it is clear that the optimal choice for minimising the loss function is simply $\theta = \frac{1}{|C|} \sum_{z_{in}} \sum_{t \in C} r_{i,t+1} \text{ i.e. the average of the partition.}$

Trees, grown to a deep enough level, are highly unbiased and flexible. The tradeoff of course, is their high variance and instability. Thus, an ensemble method called "Random Forest" was used to regularize trees by combining many different trees into a single prediction.

Random Forests

Random Forests are an extension of trees that attempt to address some of their problems. A random forest algorithm creates B different bootstrap samples from the training dataset, fits an overfit regression tree to each using only a random subset m size from all available predictors (also known as dropout), and then averages their forecasts. The overfit trees means that the underlying trees has low bias, and the dropout procedure means that they have low correlation. Thus, averaging these low bias, uncorrelated trees results in a low bias, yet stable model. Specific details of the random forest algorithm used are detailed in the appendix.

Neural Networks

Neural networks are arguably the most complex type of model available, able to capture several non-linear interactions through many layers, hence its other name "deep learning." On the flipside, their high flexibility often means that they are among the most parameterized and least interpretable models, earning them the reputation as a black box model.

The scope of this paper is limited to traditional "feed-forward" networks. The feed forward network consists of an "input layer" of scaled data inputs, one or more "hidden layers" which interact and non-linearly transform the inputs, and finally an output layer

that aggregates the hidden layers and transform them a final time for the final output.

Neural networks with up to 5 hidden layers were considered, each named NNX where X represents the number of hidden layers. The number of neurons is each layer was chosen according to the geometric pyramid rule, i.e. NN1 has 32 neurons, NN2 has 32 and 16 neurons in the first and second hidden layers respectively, NN3 has 32, 16, and 8 neurons, NN4 has 32, 16, 8, and 4 neurons, and NN5 has 32, 16, 8, 4, 2 neurons respectively. All units are fully connected; that is, each neurons receives input from all neurons the layer before it.

The ReLU activation function was used for all hidden layers owing to its high computational speed, and hence popularity within recent literature. Other potential choices for activation functions such as sigmoid, softmax, tanh etc. were not used due to computational cost.

$$ReLU(x) = max(0, x)$$
 (9)

The neural networks detailed in this paper have the following general formula. Let $K^(l)$ denote the number of neurons in each layer $l=1,\ldots,L$. Define the output of neuron k in layer l as $x_k^(l)$. Next, define the vector of outputs for this layer as $x^(l)=(1,x_1^(l),\ldots,x_{K(l)}^(l))'$. The input layer is defined using predictors, $x^(0)=(1,z_1,\ldots,z_N)'$. The recursive output formula for the neural network at each neuron in layer l>0 is then:

$$x_k^{(l)} = ReLU(x^{(l-1)}\theta_k^{l-1}),$$
 (10)

with the final output

$$g(z;\theta) = x^{(L-1)'}\theta^{L-1}$$
 (11)

The neural network's weight and bias parameters are estimated by minimizing the penalized l_2 objective function of prediction errors.

Due to the highly complex nature of the loss function, and the high dimensionality caused by the large number of parameters, this optimisation problem is very computationally intensive to solve, generally done via algorithms such as Gauss-Newton steps and is called "gradient descent." A common solution is to use "stochastic gradient descent" (SGD) where instead optimising the loss function with respect to the entire training sample, only a small, random subset of the data (mini batches) is used at each optimisation step. This sacrifices some accuracy for a dramatic improvement in computational speed.

Due the noisiness (randomness) introduced by SGD, the learning rate (step size of each descent) needs to be shrunk towards zero as the gradient approaches zero to avoid the noisiness of the mini batch causing an "overshoot" of the optimum. A learning rate shrinkage algorithm was therefore employed.

"Batch normalization" is a technique for addressing a phenomenon known as internal covariate shift, a particularly prevalent problem in training deep, complex neural networks. Internal covariate shift is where the distributions of each layer's inputs change as the parameters of the previous layer change, resulting in the need for much slower learning rates and more careful initialization of parameters. By normalizing (de-meaning and variance standardizing) each training step (batch) input, the representation power of each unit is restored. Additionally, significant gains in computational speeds may also be achieved.

Finally, multiple random seeds (initializations) were used in training neural networks and the resulting predictions were averaged in an ensemble like fashion. This allows for regularization for the variance associated with the initial random seed.

Simulation Design

Simulate a latent factor model with stochastic volatility for excess return, r_{t+1} , for t = 1, ..., T:

$$r_{i,t+1} = g(z_{i,t}) + \beta_{i,t+1}v_{t+1} + e_{i,t+1}, \quad z_{i,t} = (1, x_t)' \otimes c_{i,t}, \quad \beta_{i,t} = (c_{i1,t}, c_{i2,t}, c_{i3,t})$$

$$e_{i,t+1} = \exp(\sigma_{i,t+1}/2)\varepsilon_{i,t+1},$$

$$\sigma_{i,t+1}^2 = \omega + \alpha_i e_{i,t+1}^2 + \gamma_i \sigma_{t,i}^2 + w_{i,t+1}.$$

Let v_{t+1} be a 3×1 vector of errors, and $w_{i,t+1}, \varepsilon_{i,t+1}$ scalar error terms. The matrix C_t is an $N \times P_c$ vector of latent factors, where the first three columns correspond to $\beta_{i,t}$, across the $1 \leq i \leq N$ dimensions, while the remaining $P_c - 3$ factors do not enter the return equation. The $P_x \times 1$ vector x_t is a multivariate time series, and ε_{t+1} is a $N \times 1$ vector of idiosyncratic errors.

One of my key concerns with the Gu et al. (2019) design is that the factors are uncorrelated across i, and, in particular, that the factors which do not matter in the return equation are uncorrelated with those that matter. This is not what is observed in practice.

Instead, we will choose a simulation mechanism for C_t that gives some correlation across the factors and across time. To that end, first consider drawing normal random numbers for each $1 \le i \le N$ and $1 \le j \le P_c$, according to

$$\overline{c}_{ij,t} = \rho_i \overline{c}_{ij,t-1} + \epsilon_{ij,t}, \ \rho_i \mathcal{U}[1/2,1].$$

Then, define the matrix

$$B := \Lambda \Lambda' + \frac{1}{10} \mathbb{I}_n, \ \Lambda_i = (\lambda_{i1}, \dots, \lambda_{i4})', \ \lambda_{ik} \sim N(0, 1), \ k = 1, \dots, 4,$$

which we transform into a correlation matrix W via

$$W = \text{diag}^{-1/2}(W)W\text{diag}^{-1/2}(W).$$

To build in cross-sectional correlation, from the $N \times P_c$ matrix \bar{C}_t , we simulate characteristics according to

$$\widehat{C}_t = W\overline{C}_t.$$

Finally, we can construct the "observed" characteristics for each $1 \leq i \leq N$ and for $j = 1, \ldots, P_c$ according to

$$c_{ij,t} = \frac{2}{n+1} \operatorname{rank}(\bar{c}_{ij,t}) - 1.$$

For simulation of x_t we consider a VAR model

$$x_t = Ax_{t-1} + u_t,$$

where we have three separate specifications for the matrix A:

$$\begin{pmatrix}
(1) \ A = \begin{pmatrix}
.95 & 0 & 0 \\
0 & .95 & 0 \\
0 & 0 & .95
\end{pmatrix}$$

$$\begin{pmatrix}
(2) \ A = \begin{pmatrix}
1 & 0 & .25 \\
0 & .95 & 0 \\
.25 & 0 & .95
\end{pmatrix}$$

$$(3) \ A = \begin{pmatrix}
.99 & .2 & .1 \\
.2 & .90 & -.3 \\
.1 & -.3 & -.99
\end{pmatrix}$$

We will consider four different functions $g(\cdot)$

(1)
$$g(z_{i,t}) = (c_{i1,t}, c_{i2,t}, c_{i3,t} \times x'_t) \theta_0$$
, where $\theta_0 = (0.02, 0.02, 0.02)'$

(2)
$$g(z_{i,t}) = \left(c_{i1,t}^2, c_{i1,t} \times c_{i2,t}, \operatorname{sgn}\left(c_{i3,t} \times x_t'\right)\right) \theta_0$$
, where $\theta_0 = (0.04, 0.035, 0.01)'$

(3)
$$g(z_{i,t}) = \left(1[c_{i3,t} > 0], c_{i2,t}^3, c_{i1,t} \times c_{i2,t} \times 1[c_{i3,t} > 0], \operatorname{logit}(c_{i3,t})\right) \theta_0$$
, where $\theta_0 = (0.04, 0.035, 0.01)'$

(4)
$$g(z_{i,t}) = (\hat{c}_{i1,t}, \hat{c}_{i2,t}, \hat{c}_{i3,t} \times x'_t) \theta_0$$
, where $\theta_0 = (0.02, 0.02, 0.02)'$

Need to work out the corresponding cr0ss-sectional R^2 in this case. We can then tune θ^0 to be this close to Gu et al. (2019), as well as the predictive R^2 . This will require some work.

Follow Gu et al. (2019) in regards to the choice of N, T, P_c

Performance Evaluation

Predictive performance for individual excess stock returns were assessed using the out of sample \mathbb{R}^2 :

$$R_{OOS}^2 = 1 - \frac{\sum_{(i,t)\in\mathcal{T}_3} (r_{i,t+1} - \hat{r}_{i,t+1})}{\sum_{(i,t)\in\mathcal{T}_3} r_{i,t+1}^2}$$
(12)

where \mathcal{T}_3 indicates that the fits are only assessed on the test subsample, which is never used for training or tuning.

Variable Importance

Pending

Study

Data

Pending

Model

All machine learning methods are designed to approximate the empirical model $E_t(r_{i,t+1}) = g * (z_{i,t})$ defined in equation (2). The baseline set of stock-level covariates $z_{i,t}$ as:

$$z_{i,t} = x_t \otimes c_{i,t} \tag{13}$$

where $c_{i,t}$ is a $P_c \times 1$ matrix of characteristics for each stock i, and x_t is a $P_x \times 1$ vector of macroeconomic predictors (and are this common to all stocks, including a constant). $z_{i,t}$ is a $P \times 1$ vector of features for predicting individual stock returns $(P = P_c P_x)$ and includes interactions between individual characteristics and macroeconomic characteristics.

Trevor Hastie, 2013

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Trevor Hastie, J. F., Robert Tibshirani. (2013). The elements of statistical learning.

Appendix

 ${\bf Algorithms}$

Trees.

Algorithm 1: Classification and Regression Tree

Initialize;

for d from 1 to L do

for
$$i$$
 in $C_l(d-1), l = 1, \dots, 2^{d-1}$ do

For each feature j = 1, 2, ..., P, and each threshold level α , define a split as $s = (j, \alpha)$ which divides $C_l(d-1)$ into C_{left} and C_{right} :

$$C_{left}s = \{z_j \le \alpha\} \cap C_l(d-1); C_{right}s = \{z_j > \alpha\} \cap C_l(d-1)$$

Define the impurity function:

$$\mathcal{L}(C, C_{left}, C_{right}) = \frac{|C_{left}|}{|C|} H(C_{left}) + \frac{|C_{right}|}{|C|} H(C_{right})$$

where

$$H(C) = \frac{1}{|C|} \sum_{z_{i,t} \in C} (r_{i,t+1} - \theta)^2, \theta = \frac{1}{|C|} \sum_{z_{i,t} \in C} r_{i,t+1}$$

and |C| denotes the number of observations in set C

Find the optimal split

$$s^* \leftarrow \underset{s}{argmin} \mathcal{L}(C(s), C_{left}(s), C_{right}(s))$$

Update nodes (partition the data):

$$C_{2l-1}(d) \leftarrow C_{left}(s^*), C_{2l}(d) \leftarrow C_{right}(s^*)$$

end

 \mathbf{end}

Result: The prediction of a regression tree is:

$$g(z_{i,t}; \theta, L) = \sum_{k=1}^{2^L} \theta_k \mathbf{1}_{z_{i,t} \in C_k(L)}; \theta_k = \frac{1}{|C_k(L)|} \sum_{z_{i,t} \in C_k(L)} r_{i,t+1}$$

Random Forest.

Algorithm 2: Random Forest

for b from 1 to B do

Draw bootstrap samples $(z_{i,t}, r_{i,t+1}), (i,t) \in Bootstrap(b)$ from the dataset Grow a tree T_b using Algorithm, using only a random subsample, say \sqrt{P} of all features. Denote the resulting bth tree as

$$\hat{g}_b(z_{i,t}, \hat{\theta}_b, L) = \sum_{k=1}^{2^L} \theta_b^k \mathbf{1}_{z_{i,t} \in C_k(L)}$$

end

Result: The final random forest prediction is given by the output of all trees:

$$\hat{g}_b(z_{i,t}; L, B) = \frac{1}{B} \sum_{b=1}^{B} \hat{g}_b(z_{i,t}, \hat{\theta}_b, L)$$

Neural Networks. There are numerous stochastic gradient descent algorithms available.

Algorithm 3: Early stopping via validation

Initialize j = 0, $\epsilon = \infty$ and select the patience parameter p (max iterations)

while j < p do

Update θ using the training algorithm Calculate the prediction error from the validation sample, denoted as ϵ'

$$\begin{array}{c|c} \textbf{if} \ \epsilon' < \epsilon \ \textbf{then} \\ j \leftarrow 0 \\ \epsilon \leftarrow \epsilon' \\ \theta' \leftarrow \theta \\ \textbf{else} \\ j \leftarrow j+1 \\ \textbf{end} \end{array}$$

end

Result: θ' is the final parameter estimate

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Algorithm 4: Batch Normalization for one activation over one batch

Input: Values of x for each activation over a batch $\mathcal{B} = x_1, x_2, \dots, x_N$

$$\mu_{\mathcal{B}} \leftarrow \frac{1}{N} \sum_{i=1}^{N} x_i$$

$$\sigma_{\mathcal{B}}^2 \leftarrow \frac{1}{N} \sum_{i=1}^{N} (x_i - \mu_{\mathcal{B}})^2$$

$$\hat{x}_i \leftarrow \frac{x_i - \mu_{\mathcal{B}}}{\sqrt{\sigma_{\mathcal{B}}^2 + \epsilon}}$$

$$y_i \leftarrow \gamma \hat{x_i} + \beta := BN_{\gamma,\beta}(x_i)$$

Result: $y_i = BN_{\gamma,\beta}(x_i) : i = 1, 2, ..., N$