

Time Series Methods in the R package **mlr**

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Abstract

The **mlr** package is a unified interface for machine learning tasks such as classification, regression, cluster analysis, and survival analysis. **mlr** handles the data pipeline of preprocessing, resampling, model selection, model tuning, ensembling, and prediction. This paper details new methods for developing time series models in **mlr**. It includes standard and novel tools such as autoregressive and Lambert $W \times F()$ transform data generating processes, fixed and growing window cross validation, and forecasting models in the context of univariate and multivariate time series. Examples from forecasting competitions will be given in order to demonstrate the benefits of a unified framework for machine learning and time series.

1 Introduction

There has been a rapid development in time series methods over the last 25 years [23] whereby time series models have not only become more common, but more complex. The R language [39] has several task views that list the extensive amount of packages available for forecasting, time series methods, and empirical finance. However, the open source nature of R has left users without a standard framework whereby packages will have their own sub-culture of style, syntax, and output. The **mlr** [7] package, short for Machine Learning in R, works to give a strong syntactic framework for the modeling pipeline. By automating many of the standard tools in machine learning such as preprocessing and cross validation, **mlr** reduces error from the user during the modeling process.

While there are some time series methods available in **caret** [18], development of forecasting models in **caret** is difficult due to computational constraints and design choices within the package. The highly modular structure of **mlr** makes it the best choice for implementing time series methods and models. This paper will show how using **mlr**'s strong syntactic structure allows for time series packages such as **forecast** [28], **rugarch** [19], and **BigVAR** [38] to use machine learning methodologies such as automated parameter tuning, data preprocessing, model blending, cross validation, performance evaluation, and parallel processing techniques for decreasing model build time.

2 Forecasting Example with the M4 Competition

Professional forecasters attempt to predict the future of a series based on its past values. Forecasting can be used in a wide range of tasks including forecasting stock prices [25], weather patterns [2], international conflicts [9], and earthquakes [47]. In order to evaluate **mlr**'s forecasting framework we need a large set of possible time series to make sure our methods generalize well.

The Makridakis competition [35] is a set of forecasting challenges organized by the International Institute of Forecasters and led by Spyros Makridakis to evaluate and compare the accuracy of forecasting methods. The most recent of the competitions, the M4 competition, contains 10,000 time series on a yearly, quarterly, monthly, and daily frequency in areas such as finance, macroeconomics, climate, microeconomics, and industry. To show examples of how **mlr**'s forecasting features works we will look at a particular climate series. The data is daily with the training subset starting on September 6th, 2007 and ending on September 5th, 2009 while the testing subset is from September 6th, 2009 to October 10th, 2009 for a total of 640 training periods and 35 test periods to forecast.

```
# Read in M4 Data
library(M4comp)
library(xts)
library(lubridate)
m4.climate <- M4[[8836]]
m4.train <- xts(m4.climate$past, as.POSIXct("2007-12-05") +
               days(0:I(length(m4.climate$past)-1)))
```

```

m4.test <- xts(m4.climate$future, as.POSIXct("2009-09-06") +
              days(0:I(length(m4.climate$future)-1)))
colnames(m4.train) <- "target_var"
colnames(m4.test) <- "target_var"

```

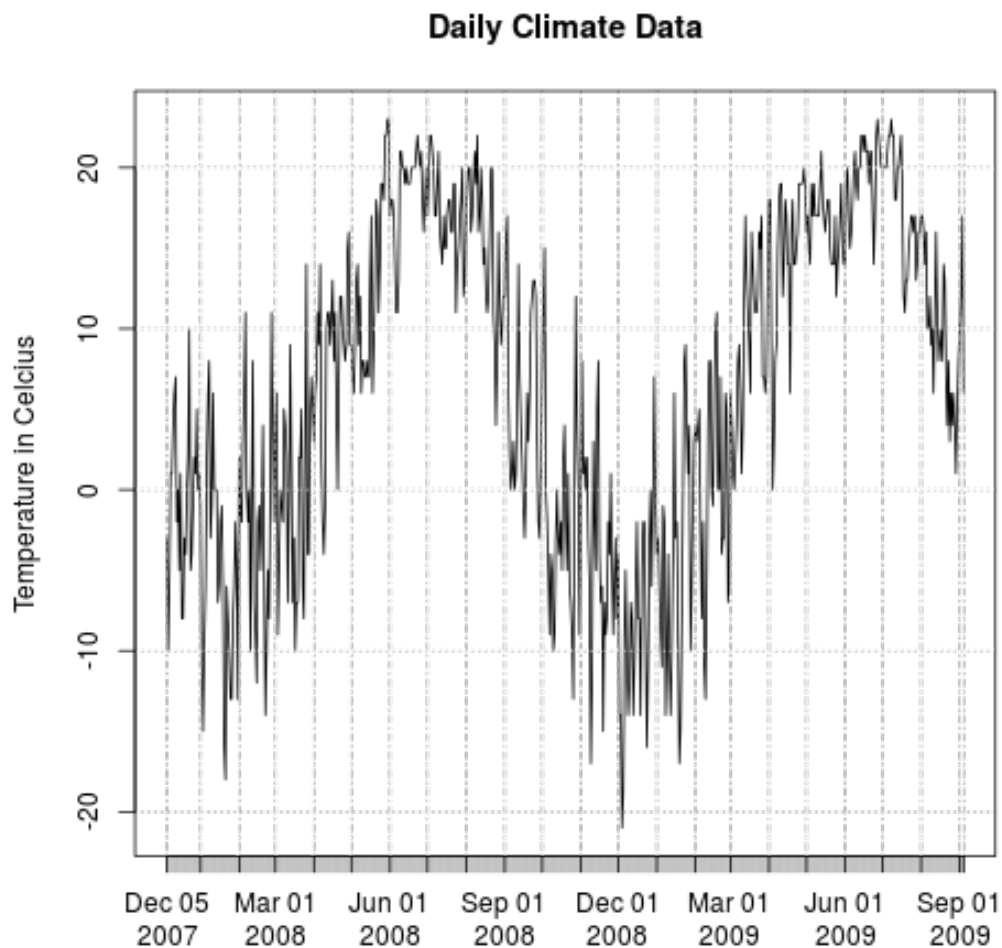


Figure 1: The training data from the M4 forecasting competition

This series was chosen for its obvious seasonality and time features, which will give the forecasting tools in **mlr** a challenge. Figure 1 is what most would

imagine when they think of a time series. There is a clear seasonal time trend with individual points moving about the seasonal periods. The data can be found in the package **M4comp** [3] under setsM4[8836].

For multivariate forecasting, we will use the EUStockMarkets data set from the **datasets** [40] package. It contains a subset of DAX, SMI, CAC, and FTSE European stock indices from July 1st, 1991 to August 24th, 1998 totaling 1828 training observations and 32 test observations.

```
# Read in Stock Market Data
data("EuStockMarkets")
EuStockMarkets.time = lubridate::date_decimal(
  as.numeric(time(EuStockMarkets)))
EuStockMarkets = xts::xts(as.data.frame(EuStockMarkets),
  order.by = EuStockMarkets.time)
eu.train = EuStockMarkets[1:1828,]
eu.test = EuStockMarkets[1829:1860,]
```

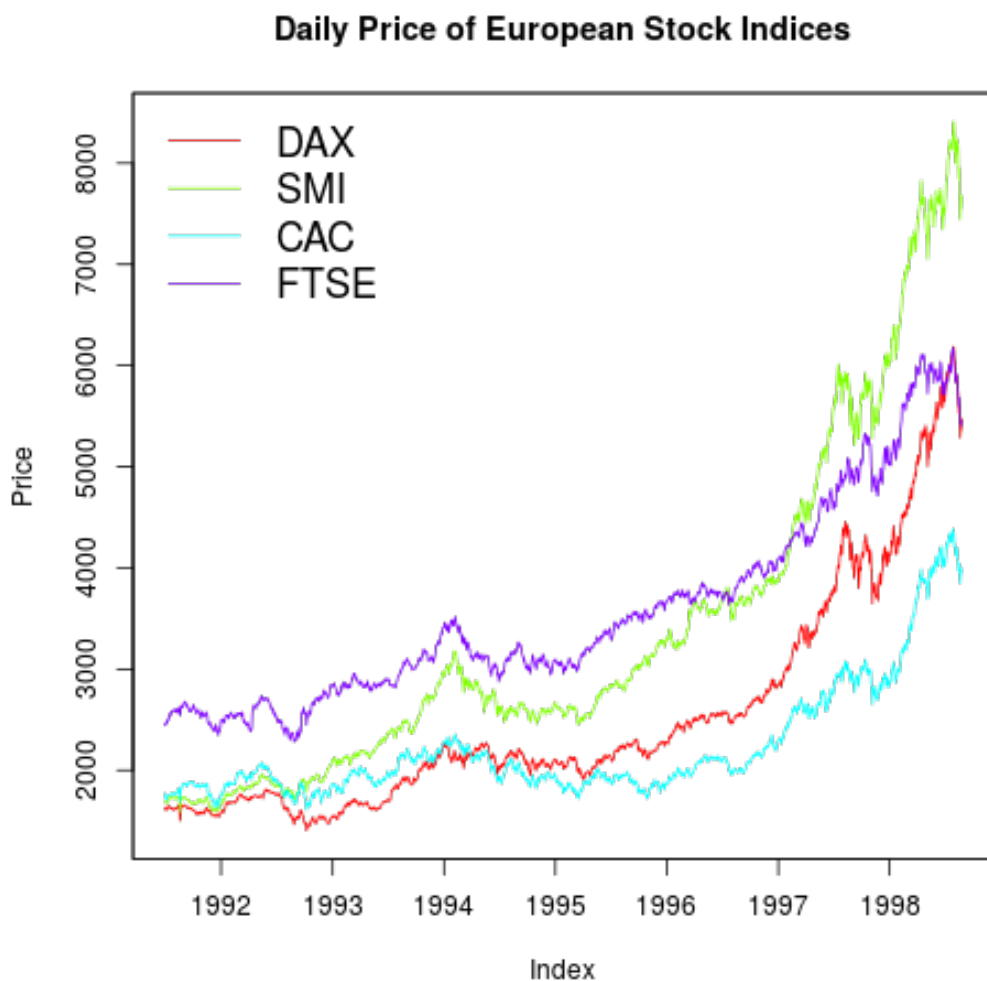


Figure 2: The training data from the EuStockMarkets dataset

Note that each stock index tends to follow a similar, but diverging, trend. This will be important to note when we perform windowing cross validation as it will let us see how well the models adapt to what appears to be nonstationary data.

3 Univariate and Multivariate Forecasting Tasks

4 Univariate Tasks

mlr uses the S3 object system to clearly define a predictive modeling task. Tasks contain the data and other relevant meta-information such as the task id and which variable you are targeting for supervised learning problems. Forecasting tasks are handled in **mlr** by the function `makeForecastRegrTask()`. The forecasting task inherits most of its arguments from `makeRegrTask`, but has two additional arguments.

data: Instead of a data frame, an `xts` object from `xts` [45] containing the time series.

frequency: An integer representing the periodicity of the time series. For example, daily data with a weekly periodicity has a frequency of 7, daily data with a yearly periodicity has a frequency of 365, and weekly data with a yearly frequency has a periodicity of 52.

```
library(mlr)

climate.task = makeForecastRegrTask(id = "M4 Climate Data",
                                   data = m4.train,
                                   target = "target_var",
                                   frequency = 183L)

climate.task

## Task: M4 Climate Data
## Type: fcregr
```

```

## Target: target_var
## Observations: 640
## Dates:
##   Start: 2007-12-05
##   End:   2009-09-04
## Frequency: 183
## Features:
## numerics  factors  ordered
##         0         0         0
## Missings: FALSE
## Has weights: FALSE
## Has blocking: FALSE

```

Like a regression task, this records the type of the learning problem and basic information about the data set such as the start and end dates, frequency, and whether we have missing values. Note that there are zero features in our task because we only have a target variable, which the model itself will use to build features.

4.1 Multivariate Tasks

One common problem with forecasting is that it is difficult to use additional explanatory variables or forecast multiple targets that are dependent on one another. If we are at time t and want to forecast 10 periods in the future, we need to know the values of the explanatory variables at time $t + 10$, which is often not possible. A new set of models [38] which treats explanatory variables endogenously instead of exogenously allows us to forecast not only our target, but additional explanatory variables. This is done by treating all the variables as targets, making them endogeneous to the model. To use these models, we create a multivariate forecasting task. The function `makeMultiForecastRegrTask()` has the same arguments as `makeForecastRegrTask()` with one exception. The `target` argument can contain either a single target variable, multiple target variables, or `All` which treats all variables endogeneously.


```

mfcreegr.univar.task = makeMultiForecastRegrTask(id = "bigvar",
                                                data = EuStockMarkets,
                                                target = "FTSE",
                                                frequency = 365L)

mfcreegr.univar.task

## Task: bigvar
## Type: mfcreegr
## Target: FTSE
## Observations: 1860
## Dates:
##   Start: 1991-07-01 02:18:27
##   End:   1998-08-24 20:18:27
## Frequency: 365
## Features:
## numerics  factors  ordered
##          3         0         0
## Missings: FALSE
## Has weights: FALSE
## Has blocking: FALSE

```

Like `makeForecastRegrTask()`, `mfcreegr.univar.task` has the standard output, but notice now that there are three features. Alternatively, `mfcreegr.all.task` contains multiple target values with no features. The difference between each of these multivariate tasks is that `mfcreegr.univar.task` will act similar to `makeForecastRegrTask()`, giving a univariate forecast output and evaluating the forecasts with univariate measures. When the target is `All` the model trained model will forecast and output all series and uses a multivariate forms of the univariate measures. Though the results appear different, both of these tasks will still forecast all of the underlying series, which allows us take exogenous models and treat them endogeneously for n-step ahead forecasts that use additional explanatory variables.

```

mfcregr.all.task = makeMultiForecastRegrTask(id = "bigvar",
                                             data = eu.train,
                                             target = "all",
                                             frequency = 365L)

mfcregr.all.task

## Task: bigvar
## Type: mfcregr
## Target: DAX SMI CAC FTSE
## Observations: 1828
## Dates:
##   Start: 1991-07-01 02:18:27
##   End:   1998-07-10 22:09:13
## Frequency: 365
## Features:
## numerics  factors  ordered
##          0         0         0
## Missings: FALSE
## Has weights: FALSE
## Has blocking: FALSE

```

5 Building and Tuning a forecast learner

5.1 Univariate Forecasting

The `makeLearner()` function provides a structured model building framework to the several forecasting models currently implemented in **mlr**. As an example, we will build the Trigonometric exponential smoothing state space model with Box-Cox transformation, ARMA errors, Trend and Seasonal Components (TBATS) [34].

TBATS is one of the most well known forecasting models and is available in **mlr** along with models such as BATS, ARIMA, ETS, several GARCH

variants, and autoregressive neural networks. In addition, Section 7.1 shows preprocessing features which allow arbitrary supervised machine learning models to be used in the context of forecasting. To implement the TBATS model we use `makeLearner()`, supplying the class of learner, order, the number of steps to forecast, and any additional arguments to be passed to `tbats` for **forecast**.

```
tbats.mod = makeLearner("fcregr.tbats", use.box.cox = TRUE,
                        use.trend = TRUE,
                        seasonal.periods = TRUE, max.p = 60, max.q = 60,
                        stationary = FALSE, use.arma.errors = TRUE,
                        h = 35, predict.type = "response")
```

We can also supply a predict type for forecasting models to either receive point estimates (**response**) or point estimates with quantiles of confidence intervals (**quantile**). To train the model we simply call `train`, supplying the forecasting model and task.

```
train.tbats = train(learner = tbats.mod, task = climate.task )
```

After training the model forecasts can be made by calling `predict()`. Optionally supplying the test data as an additional argument will allow **mlr** to return a nicely coalesced data frame with both the test and train data, returning an object containing meta information for the forecasts along with the prediction and test data in columns **truth** and **response**, respectively.

```
predict.tbats = predict(train.tbats, newdata = m4.test)
predict.tbats
## Prediction: 35 observations
```

```
## predict.type: response
## threshold:
## time: 0.00
##           truth response
## 2009-09-06      7 3.231406
## 2009-09-07     12 4.425507
## 2009-09-08     14 5.380601
## 2009-09-09     10 5.617823
## 2009-09-10     10 5.511217
## 2009-09-11     16 5.331298
## ... (35 rows, 2 cols)
```

To measure the performane of TBATS we call `performance()` with the Mean Absolute Scaled Error (MASE) [29] measure.

```
performance(predict.tbats, measure = mase,
             task = climate.task)

##           mase
## 0.0676601
```

MASE has favorable properties for calculating forecast errors relative to measures such as root mean squared error or median relative absolute error. Arguably one of the most important features, it's very interpretable. Let y_t and \tilde{y}_t be the target variable and prediction at time t until the final time T such that $t > T$ with $\epsilon_t = y_t - \tilde{y}_t$ being the forecast error.

$$\text{MASE} = \frac{\sum_{t=1}^T |\epsilon_t|}{\frac{T}{T-1} \sum_{t=2}^T |y_{t,\text{insample}} - y_{t-1,\text{insample}}|} \quad (1)$$

Where the denominator is the one step ahead naive forecast from the training data. When the numerator is equal to the denominator the model performed as good as a simple naive forecast method. A MASE score greater

than one indicates a model is performing worse than the naive forecasting method while scores less than one mean the model is performing better than the naive forecasting method.

The scale invariance of MASE means that it is independent of the scale of the data which allows models to be compared across data sets. The scale invariance of MASE has made it a favorite for comparing the accuracy of forecast methods [16] across datasets. While scaling in measures such as the Mean Absolute Percentage Error can cause poor behavior as the target variable goes to zero, MASE does not become skewed when the target variable approaches zero. This allows MASE to be use in situations in which zeros occur frequently or zero is not meaningful such as predicting temperature.

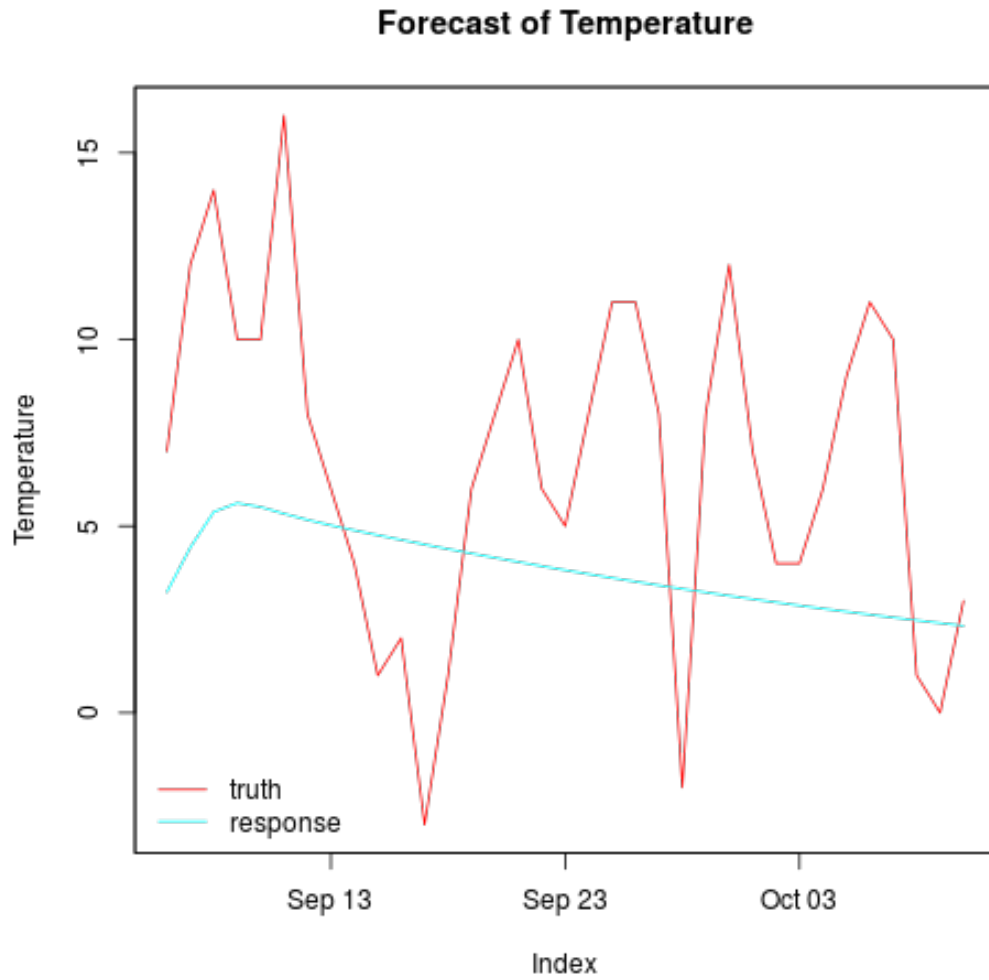


Figure 3: The forecasts response from TBATS against the true testing weather data

The model does a successful job of catching the downward trend, but figure 3 shows the downward bias of the model which does not allow it to account for the sharp upward swings. Forecasting models are designed to predict the next n values that will appear in the series. This means that without a way to update the model with new data the model will have to be completely rebuilt for each

new set of forecasts. The `updateModel` function allows forecasters to avoid the expensive cost of retraining the model. The function updates the model with new points which are then allows for updated forecasts.

```
update.tbats = updateModel(train.tbats, climate.task, newdata = m4.test)
predict(update.tbats, task = climate.task)

## Prediction: 35 observations
## predict.type: response
## threshold:
## time: 0.00
##
##               response
## 2009-09-04 23:59:54 6.319959
## 2009-09-05 23:59:48 7.875401
## 2009-09-06 23:59:43 8.209328
## 2009-09-07 23:59:37 8.157638
## 2009-09-08 23:59:31 8.083591
## 2009-09-09 23:59:26 8.055280
## ... (35 rows, 1 cols)
```

5.2 Multivariate Forecasting

The package **BigVAR** [37] has been implemented for multivariate forecasting. **BigVAR** allows for estimation of high dimensional time series through a structured Lasso penalties method that finds the best autoregressive structure through cross-sections and time.

```
bigvar.mod = makeLearner("mfcreegr.BigVAR", p = 25, struct = "SparseLag",
                        gran = c(50, 60), h = 35, n.ahead = 35)
```

This section uses the multivariate forecast task which has all variables as targets, while section 9.2 performs the single target multivariate forecast with a stacked predictor.

```
train.bigvar = train(learner = bigvar.mod, task = mfcregr.all.task )
train.bigvar
```

```
## Model for learner.id=mfcregr.BigVAR; learner.class=mfcregr.BigVAR
## Trained on: task.id = bigvar; obs = 1828; features = 0
## Hyperparameters: p=25,struct=SparseLag,gran=50,60,h=35,n.ahead=35
```

Predictions for `multiForecast` methods have a similar output to `multiclass` methods, returning multiple truth and response variables. A multivariate version of MASE has been implemented which takes the mean of each MASE score for the individual variables.

```
predict.bigvar = predict(train.bigvar, newdata = eu.test)
predict.bigvar
```

```
## Prediction: 32 observations
## predict.type: response
## threshold:
## time: 0.02
##
```

		truth.DAX	truth.SMI	truth.CAC	truth.FTSE	response.DAX
##	1998-07-12 07:50:46	5905.15	8047.3	4252.1	5960.2	5852.841
##	1998-07-13 17:32:18	5961.45	8099.0	4304.4	5988.4	5820.194
##	1998-07-15 03:13:50	5942.06	8166.0	4311.1	5990.3	5801.712
##	1998-07-16 12:55:23	5975.88	8160.0	4333.1	6003.4	5790.565
##	1998-07-17 22:36:55	6018.89	8227.2	4339.9	6009.6	5781.540
##	1998-07-19 08:18:27	6000.84	8205.0	4319.2	5969.7	5772.087

```
##
```

		response.SMI	response.CAC	response.FTSE
##	1998-07-12 07:50:46	7982.503	4192.149	5997.825
##	1998-07-13 17:32:18	7960.779	4153.550	6037.958
##	1998-07-15 03:13:50	7955.264	4130.674	6063.687
##	1998-07-16 12:55:23	7958.119	4115.612	6083.094
##	1998-07-17 22:36:55	7965.061	4103.302	6100.075
##	1998-07-19 08:18:27	7973.586	4091.154	6116.419

```
## ... (32 rows, 8 cols)
```



```
performance(predict.bigvar, multivar.mase, task = mfcregr.all.task)

## multivar.mase
##      0.2209703
```

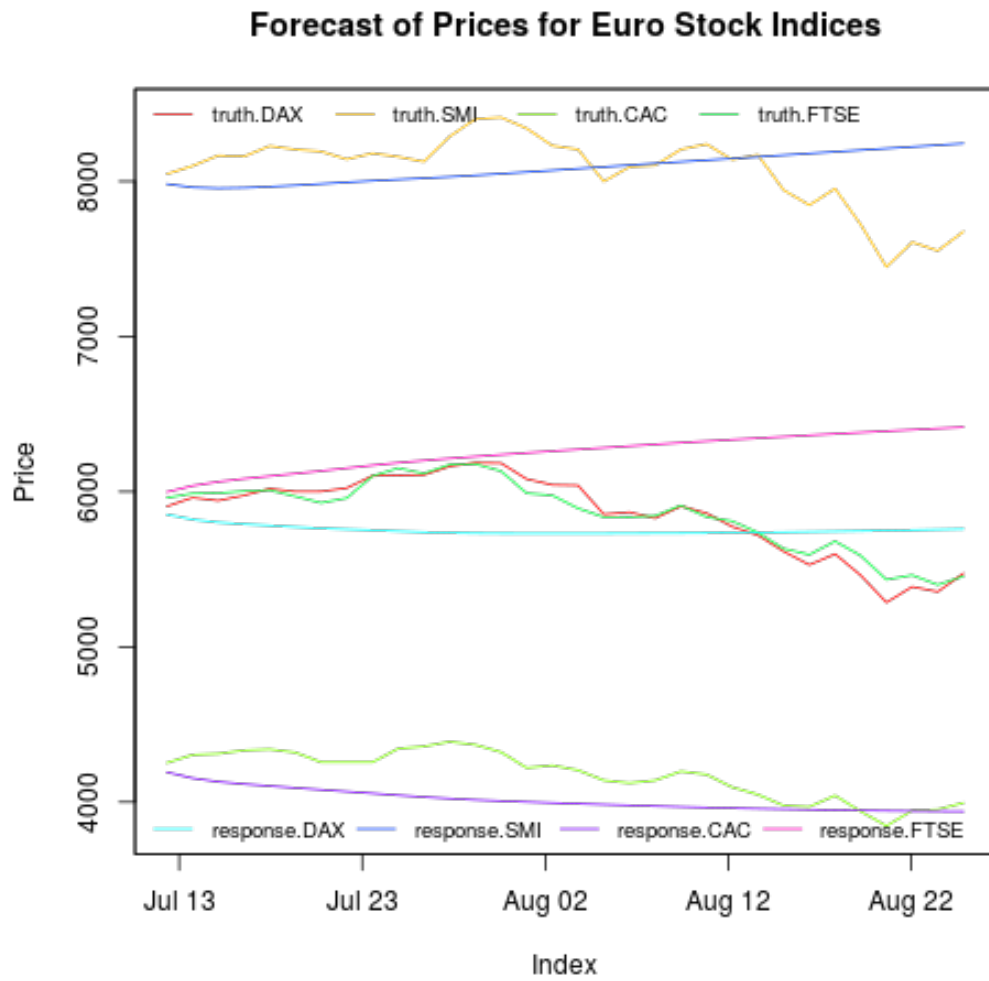


Figure 4: The forecasts response from BigVAR against the true test stock index data

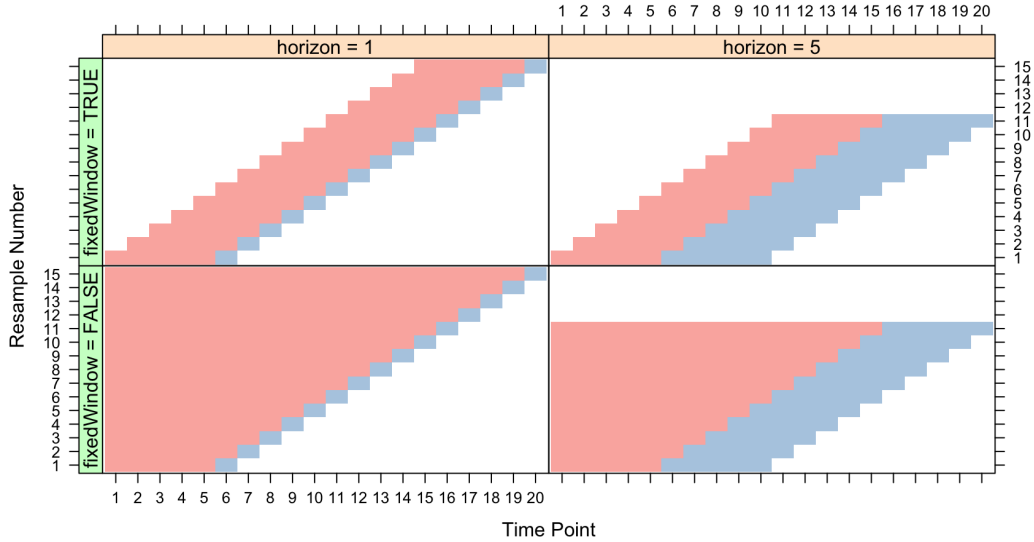
6 Resampling with Time

While TBATS is one of the most well known time series models, the order selection process or the ARIMA errors and whether to include trend, damped trend, or seasonal periods can be a subjective process that makes finding the best model difficult for users. One of the first proposals for automated forecasting methods comes from [13] for automatic order selection of ARIMA models. Innovations are obtained by fitting high order autoregressive models to the data and then computing the likelihood of potential models through a series of standard regressions. Proprietary algorithms from software such as **Forecast Pro** [43] and **Autobox** [42] are well known and have performed to high standards in competitions such as the M3 forecasting competition [35]. One of the most well known R packages for automated forecast is **forecast** [28] which contains several methods for automated forecasting including exponential smoothing based methods and step-wise algorithms for forecasting with ARIMA models.

Forecasting in **mlr** takes a machine learning approach, creating a parameter set for a given model and using an optimization method to search over the parameter space. To do this, we will use a windowing resampling scheme to train over the possible models. Resampling schemes such as cross-validation, bootstrapping, etc. are common in machine learning for dealing with the bias-variance tradeoff [17] [44]. When there is a time component to the data, windowing schemes are useful in allowing a valid resampling scheme while still accounting for the time properties of the series. Figure 5 gives an example of fixed and growing window cross validation. Let h be a forecast horizon where the index of the starting and end points of the training data are start_i and

end_i where i is the index for each window. The first model will train on data between start_i and end_i while testing on data indexed between $\text{end}_i + h$. After each iteration the window slides forward h steps such that $\text{end}_{i+1} = \text{end}_i + h$. In the case of the fixed window, the starting index will also shift such that $\text{start}_{i+1} = \text{start}_i + h$ while start_i will remain constant in the growing case. This type of growing and fixed window resampling [27] are now available in the `resampling()` function of **mlr**.

Figure 5: Resampling with a window scheme as exemplified by **caret** [30]. The top graphs are fixed window cross validation while the bottom graphs are growing window cross validation.



A windowing resampling process is created in the function `makeResampleDesc()` by supplying the resampling type, horizon, initial window, the length of the series, and an optional argument to skip over some windows for the sake of time.

```
resampDesc = makeResampleDesc("GrowingCV", horizon = 35L,
                              initial.window = .7,
                              size = nrow(getTaskData(climate.task)),
                              skip = .004)

resampDesc

## Window description:
##  growing with 79 iterations:
##  448 observations in initial window and 35 horizon.
## Predict: test
## Stratification: FALSE
```

The methods in package **ParamHelpers** [6] allow **mlr** to have a simple and rigorous interface for creating parameter spaces. There are several types of tools to help us search our parameter space including grid search, random search [4], iterated F-racing, Simulated Annealing, and CMA-ES [1] to search the parameter space for the most optimal model.

```
parSet = makeParamSet(
  makeLogicalParam(id = "use.box.cox", default = FALSE,
                  tunable = TRUE),
  makeLogicalParam(id = "use.trend", default = FALSE,
                  tunable = TRUE),
  makeLogicalParam(id = "use.damped.trend", default = FALSE,
                  tunable = TRUE),
  makeLogicalParam(id = "seasonal.periods", default = FALSE,
                  tunable = TRUE),
  makeIntegerParam(id = "max.p", upper = 30, lower = 1,
```

```

        trafo = function(x) x*2),
makeIntegerParam(id = "start.p", upper = 30, lower = 1,
        trafo = function(x) x*2),
makeIntegerParam(id = "max.q", upper = 30, lower = 1,
        trafo = function(x) x*2),
makeIntegerParam(id = "start.q", upper = 5, lower = 0,
        trafo = function(x) x*2),
makeIntegerParam("max.P", lower = 0, upper = 5),
makeIntegerParam("max.Q", lower = 0, upper = 5),
makeDiscreteParam("ic", values = c("aicc", "aic", "bic")),
makeDiscreteParam("test", values = c("kpss", "adf", "pp")),
makeDiscreteParam("seasonal.test",
        values = c("ocsb", "ch")),
makeLogicalParam("biasadj", default = FALSE)
)

#Specify tune by grid estimation
ctrl = makeTuneControlIrace(maxExperiments = 500L)

```

Using `tuneParams()` the model is tuned for the task using the specified resampling scheme, parameter set, tune control, and measure. The `trafo` argument allows the tuning process to take values on a separate scale than the one described. In the tuning scheme above, `trafo` will search over 1, 4, 9, ..., 81, and 100 values of lag, selecting the model with the best lag structure. For this tuning task we use MASE [29] as a measure of performance¹. The model is also tuned in parallel using the package `parallelMap` [5] for its built in compatibility with `mlr`.

```

#Tune the TBATS Model
library("parallelMap")
parallelStartSocket(8)

```

¹Models with a seasonal difference > 0 may be favorably biased as we use the non-seasonal MASE score

```

configureMlr(on.learner.error = "warn")
set.seed(1234)
tbatsTune = tuneParams(makeLearner("fcregr.tbats", h = 35),
                        task = climate.task,
                        resampling = resampDesc, par.set = parSet,
                        control = ctrl, measures = mase)

parallelStop()
# Output the test mean of best model
tbatsTune$y

```

```

## mase.test.mean
##      0.06195313

```

Using **mlr**'s built in plotting routines the tuning parameters can be analyzed graphically using partial dependence plots [22]. Because of constraints which do not allow logical types in the partial dependence plots, the below code takes all of the logical parameters from **tbatsTune** and coerces them to numeric.

```

# Making graphics
plotHyperParsEffect(tbats.hyp, x= "max.q", y = "mase.test.mean",
                    plot.type = "line",
                    partial.dep.learn = "regr.randomForest")
plotHyperParsEffect(tbats.hyp, x= "max.P", y = "mase.test.mean",
                    plot.type = "line",
                    partial.dep.learn = "regr.randomForest")

```

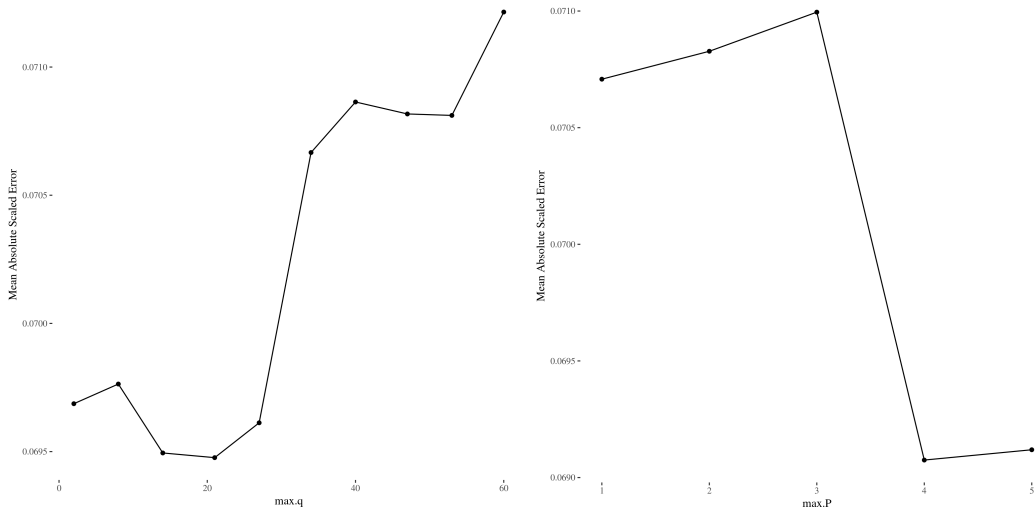


Figure 6: The figure on the left shows the dependence of the MASE score averaged over all windows for each value of the maximum number of moving average terms q . The figure on the right does the same, but for the autoregressive term p

For this model the best maximum number of moving average coefficients is two while the best number of seasonal autoregressive terms is five. The best model's parameters are extracted using `setHyperPars()` and passed to `train()` to go over the full data set.

```
# Train best model over full data
lrn = setHyperPars(makeLearner("fcregr.tbats", h = 35),
  par.vals = tbatsTune$x)
m = train(lrn, climate.task)
```

To make predictions the task test data are passed to `predict()`

```

climate.pred = predict(m, newdata = m4.test)
performance(climate.pred, measures = mase, task = climate.task)

##          mase
## 0.0562274

```

For comparison, TBATS is run in **forecast** with it's base parameters

```

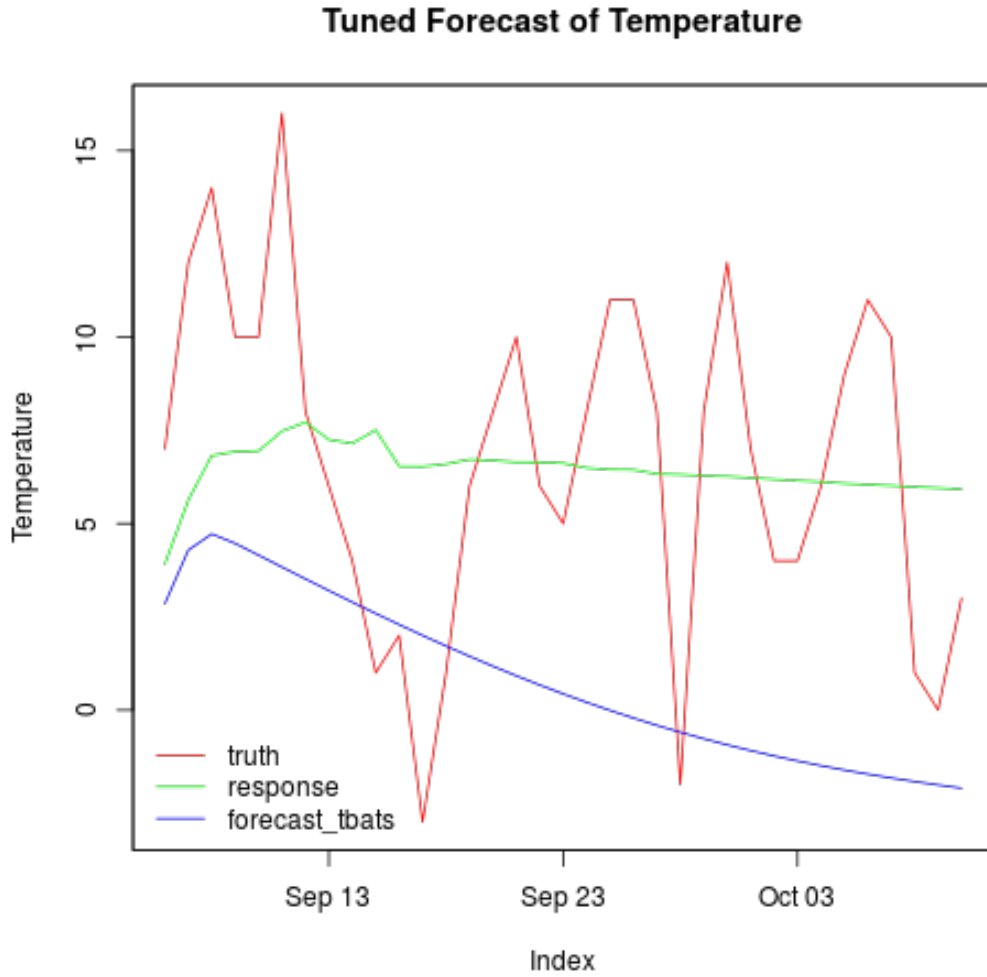
library(forecast)
forecast.train.tbats = tbats(ts(m4.train,frequency = 365))
forecast.tbats = forecast(forecast.train.tbats,h=35)
# Calculate MASE
sum(abs(coredata(m4.test)-forecast.tbats$mean)) /
  ((nrow(m4.test) / (nrow(m4.test) + 1)) *
    sum(abs(diff(coredata(m4.test))))))

## [1] 1.951852

```

Now it's possible to see the clear benefit of the forecast extension to **mlr**. While **forecast** does some automated tuning, this model would still require extensive manual testing in an attempt to find the best model. By automating this process we come to a faster solution with minimal work.

Figure 7: The TBATS model forecasts tuned in **mlr** Vs. the base TBATS model in forecast



To this author's knowledge, this is the first package in R that allows for automated tuning and training of GARCH models [15]. It is possible to pass and train multiple types of GARCH models while also tuning the models respective parameters. In this example `predict.type = "quantile"` is used

to estimate confidence intervals for the forecast.

```
# Make a tuning grid for GARCH
par_set = makeParamSet(
  makeDiscreteParam(id = "model", values = c("sGARCH", "csGARCH")),
  makeIntegerVectorParam(id = "garchOrder", len = 2L, lower = c(1,1),
    upper = c(4,4)),
  makeIntegerVectorParam(id = "armaOrder", len = 2L, lower = c(5,1),
    upper = c(8,3)),
  makeLogicalParam(id = "include.mean"),
  makeLogicalParam(id = "archm"),
  makeDiscreteParam(id = "distribution.model",
    values = c("norm","std","jsu")),
  makeDiscreteParam(id = "stationarity", c(0,1)),
  makeDiscreteParam(id = "fixed.se", c(0,1)),
  makeDiscreteParam(id = "solver", values = "nloptr")
)

#Specify tune by F-racing
ctrl = makeTuneControlRtrace(maxExperiments = 400L)

#Tuning in parallel
parallelStartSocket(6)
configureMlr(on.learner.error = "warn")
set.seed(1234)
garchTune = tuneParams(makeLearner("fcregr.garch", n.ahead= 35),
  task = climate.task, resampling = resampDesc,
  par.set = par_set, control = ctrl,
  measures = mase)

parallelStop()
garchTune$y

## mase.test.mean
##      0.07888857
```

Because the partial dependence plots are still in development they do not work well with `makeNumericVectorParams()` such as `garchOrder` and

armaOrder. Instead of looking at each p, q individually, the sum of the coefficient values for the p, q order of the GARCH component and the ARMA component are plotted.

```
# Make hyper parameter data
garch.hyperpar = generateHyperParsEffectData(garchTune,
      trafo = TRUE, include.diagnostics = FALSE,
      partial.dep = TRUE)
garch.hyperpar$data = garch.hyperpar$data[
      -which(garch.hyperpar$data$mase.test.mean ==
            max(garch.hyperpar$data$mase.test.mean)),]
garch.hyperpar$data$garchOrder1 = garch.hyperpar$data$garchOrder1 +
      garch.hyperpar$data$garchOrder2
garch.hyperpar$data$garchOrder2 = NULL
colnames(garch.hyperpar$data)[2] = "garchOrder"
garch.hyperpar$data$arOrder1 = garch.hyperpar$data$arOrder1 +
      garch.hyperpar$data$arOrder2
garch.hyperpar$data$arOrder2 = NULL
colnames(garch.hyperpar$data)[3] = "arOrder"
garch.hyperpar$data$include.mean = as.numeric(garch.hyperpar$data$include.mean)
garch.hyperpar$data$archm = as.numeric(garch.hyperpar$data$archm)

plotHyperParsEffect(garch.hyperpar, x= "garchOrder", y = "mase.test.mean",
      plot.type = "line",
      partial.dep.learn = "regr.randomForest")
plotHyperParsEffect(garch.hyperpar, x= "arOrder", y = "mase.test.mean",
      plot.type = "line",
      partial.dep.learn = "regr.randomForest")

## Saving 7 x 7 in image
## Saving 7 x 7 in image
```

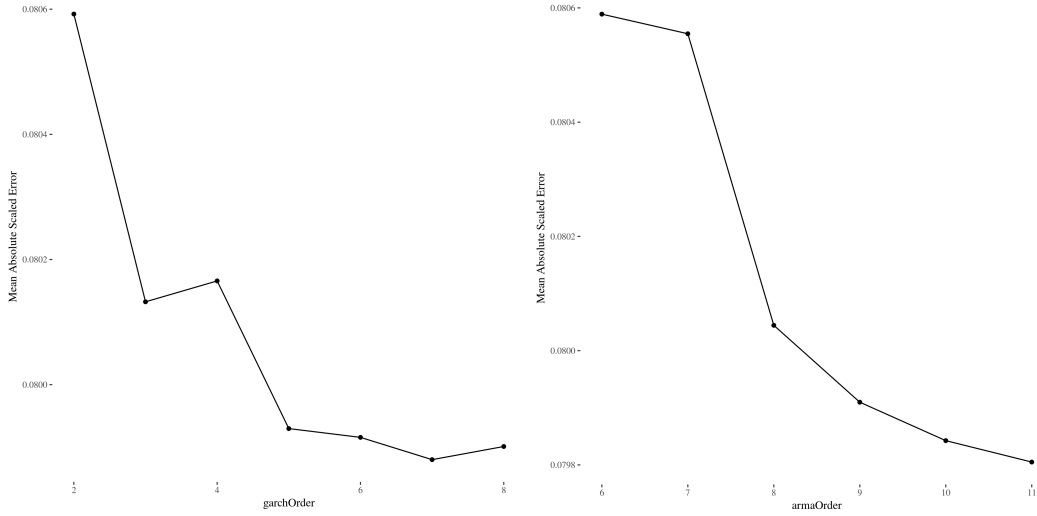


Figure 8: The figure on the left shows the dependence of the MASE score averaged over all windows for each value of the maximum number of p, q terms in the GARCH order. The figure on the right does the same, but for the ARMA order

The dependence plots show that for both the GARCH and ARMA orders, as the order is increased, the MASE score decreases. A better model would arise if order of both the GARCH and ARMA components increased up to twenty or more. Forecasting models with quantiles are treated the same as any other learner post-tuning. The best hyperparameters are taken and a final model is trained over the entire dataset.

```
# Get the hyperpars and train the best GARCH model
tuned.lrn = setHyperPars(makeLearner("fcregr.garch",
                                     predict.type = "quantile"),
                      par.vals = garchTune$x)
garch.train = train(tuned.lrn, climate.task)
```

```

climate.pred = predict(garch.train, newdata = m4.test)
climate.pred

## Prediction: 35 observations
## predict.type: quantile
## threshold:
## time: 0.01
##
##      truth response se.quantile0.05 se.quantile0.95
## 2009-09-06      7 3.870648    -2.8377519    10.57905
## 2009-09-07     12 5.940932    -0.7674667    12.64933
## 2009-09-08     14 6.962520     0.1060161    13.81902
## 2009-09-09     10 7.336334     0.3367573    14.33591
## 2009-09-10     10 7.473346     0.4736223    14.47307
## 2009-09-11     16 7.737601     0.7377337    14.73747
## ... (35 rows, 4 cols)

```

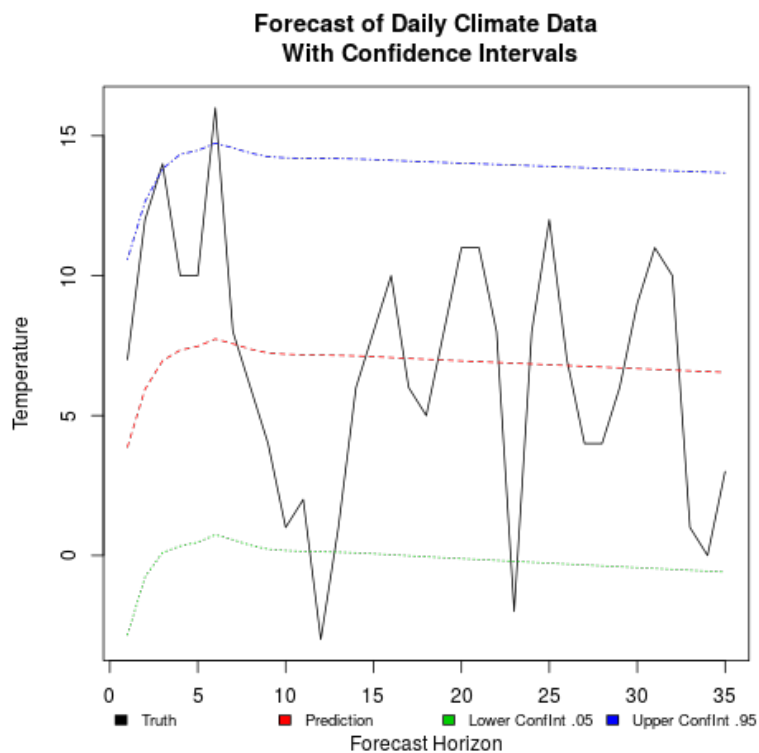


Figure 9: The GARCH model with confidence intervals

```
performance(climate.pred, measures = mase, task = climate.task)

##          mase
## 0.05589035
```

The simple model here performs relatively well. The extremes of the test data are almost all within the given bounds of the 95% confidence interval. With a bit more tuning and perhaps an alternative control for the search over the parameter space, the reader would easily find a better model. While GARCH is excellent at catching the conditional heteroskedasticity of the past series, the TBATS model performs well in terms of catching long or multiseasonal type periods. Section 9.1 goes over how to stack forecasting models to make use of the best characteristics of each.

7 Forecasting with Machine Learning Models

7.1 Forecasting with Regression Tasks

The forecasting extension of **mlr** includes a preprocessing function that allows forecasting with supervised machine learning models. the function `createLagDiffFeatures()` allows for $AR(p, d)$ structures to be imbedded in machine learning models.

```
climate.regr.task = makeRegrTask(id = "lagged gbm",
                                data = as.data.frame(m4.train),
                                target = "target_var")
climate.task.lag = createLagDiffFeatures(climate.regr.task,
                                         lag = 1L:24L,
                                         difference = 1L,
                                         na.pad=FALSE)

climate.task.lag

## Supervised task: lagged gbm
```

```
## Type: regr
## Target: target_var
## Observations: 615
## Features:
## numerics  factors  ordered
##          24        0        0
## Missings: FALSE
## Has weights: FALSE
## Has blocking: FALSE
```

Notice that `createLagDiffFeatures()` returns a new task with the lagged variables as the new features. Once the lagged task is created the model is trained or tuned like any other.

```
# make Gradient Boosting Machine
lag.gbm = makeLearner("regr.gbm", par.vals = list(n.trees = 20000,
                                                  shrinkage = .000001,
                                                  interaction.depth = 15,
                                                  bag.fraction = .7))
gbm.train = train(lag.gbm, climate.task.lag)
```

The `forecast()` function allows machine learning models to do arbitrary n -step ahead forecasts. Let the one step ahead forecast be defined by

$$\hat{y}_{t+1} = \sum_{i=1}^p (\rho_i \Delta_d y_i + \epsilon_i) \quad (2)$$

where ρ_i is the autoregressive parameter of order p and d is the lag of the difference operator Δ . Then the n -step ahead forecast is defined as

$$\hat{y}_{t+n} = \sum_{i=t+1}^n (\rho_i \Delta_d \hat{y}_i + \epsilon_i) \quad (3)$$

```
# Forecast with GBM
gbm.forecast = forecast(gbm.train, h = 35L,
                        newdata = as.data.frame(m4.test))
```

```
gbm.forecast

## Prediction: 35 observations
## predict.type: response
## threshold:
## time: 7.30
##   truth response
## 1      7 7.655469
## 2     12 8.234003
## 3     14 8.234003
## 4     10 8.234003
## 5     10 8.234003
## 6     16 8.234003
## ... (35 rows, 2 cols)

performance(pred = gbm.forecast,
            measures = mase,
            task = climate.regr.task)

##      mase
## 0.0558791
```

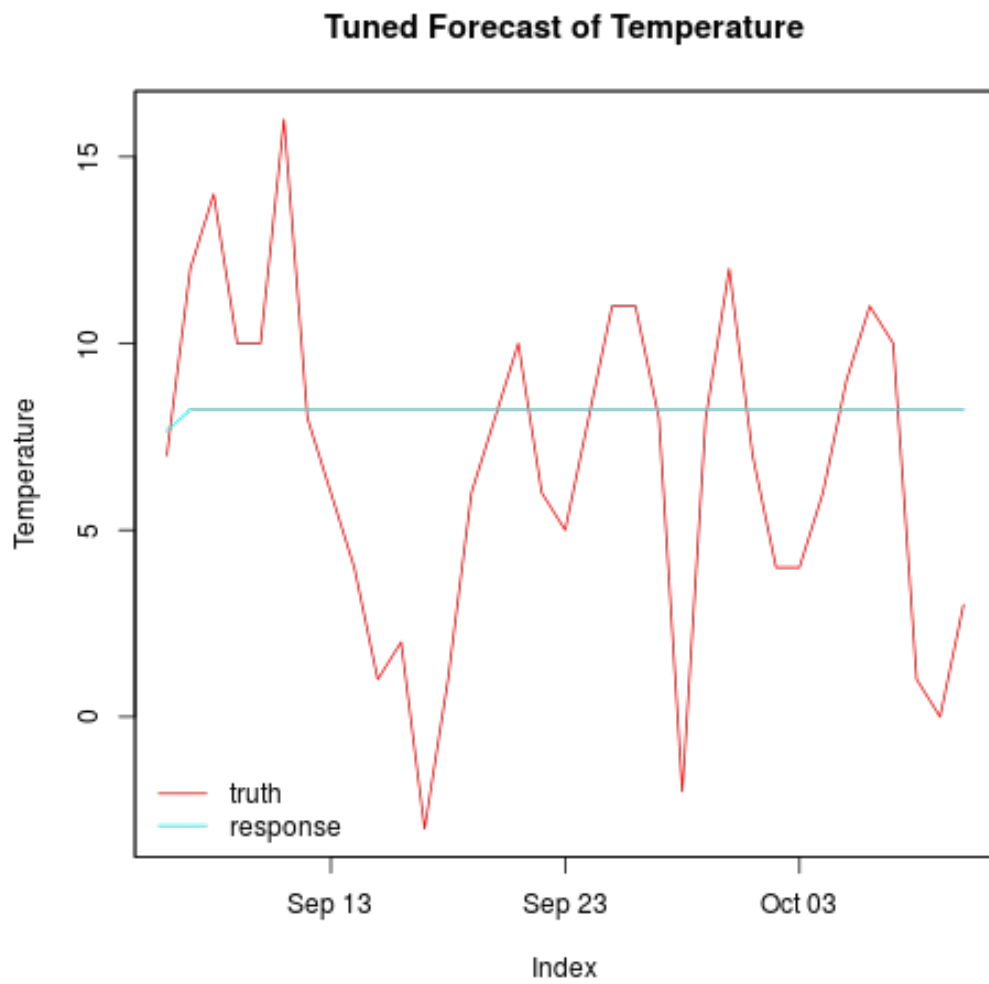



Figure 10: The forecast of the M4 data using a Gradient Boosting Machine

The forecasts drift quickly to an average value, which may be due to the modeling placing the highest value on the most recent data.

7.2 Forecasting with Classification Tasks

Forecasting for binary or multiclass outcomes [31] is a common problem in the real world. However, research in this area of forecasting only started picking up speed in the last decade [14]. The forecasting extension of **mlr** gives econometricians the ability to take all the classification models in **mlr** and apply them to the forecasting context. For developing trading strategies, we normally have a discrete set of choices such as to buy, sell, or hold onto a stock. Using forecasting in **mlr** we can now train classification models that forecast these choices [32]. To example this, a simple buy, sell, or hold trading strategy will be built Using the `EuStockMarkets`'s DAX index. If the stock goes up by 5% in a day a buy is executed, down 5% a sell is executed, and otherwise no action will be taken.

```
# Make Trading Strategy
DAX = EuStockMarkets$DAX/lag(EuStockMarkets$DAX,
                             7,na.pad = FALSE) - 1
trade.strat = ifelse(DAX > .05, "Buy",
                    ifelse(DAX < -.05, "Sell", "Hold"))
trade.strat = trade.strat[8:1860]
euro.classif.data = data.frame(trade.strat = trade.strat ,
                              row.names = index(trade.strat))
euro.classif.train = euro.classif.data[1:1838,,drop = FALSE]
euro.classif.test  = euro.classif.data[1839:1853,,drop = FALSE]
classif.task = makeClassifTask(data = euro.classif.train,
                              target = "DAX")

# Make Lagged Task and Learner
classif.task.lag = createLagDiffFeatures(classif.task,
                                         lag = 1L:565L,
                                         na.pad = FALSE)
classif.learn = makeLearner("classif.boosting", xval = 1,
```

```
mfinal = 200, minsplit = 10)
```

```
classif.train = train(classif.learn, classif.task.lag)
classif.fc = forecast(classif.train, h=15, newdata = euro.classif.test)
classif.fc
performance(classif.fc)
```

```
## Prediction: 15 observations
## predict.type: response
## threshold:
## time: 422.87
##   truth response
## 1  Hold      Hold
## 2  Hold      Hold
## 3  Sell      Hold
## 4  Hold      Hold
## 5  Hold      Hold
## 6  Hold      Hold
## ... (15 rows, 2 cols)
##           mmce
## 0.4666667
```

With the models and methodologies available in **mlr**, forecasting binary outcomes is now as simple as any other model. These tools can be used for further research in areas such as directions of stock movement [33] and forecasting extreme values [10].

8 Lambert W Transforms

Many machine learning and time series models rely on the assumption that our data or errors fit a normal distribution. This assumption becomes precarious when modeling the asymmetric and fat-tailed data of the real world. Lambert

$W \times F()$ transforms are a family of generalized skewed distributions [20] that have bijective and parametric functions that allow heavy tailed and asymmetric data to appear more Gaussian [21].

Let U be a continuous random variable with cdf $F_U(u|\beta)$ and pdf $f_U(u|\beta)$ given β is a parameter vector. Define a continuous location-scale random variable $X \sim F_X(x|\beta)$. A location-scale skewed Lambert $W \times F_X$ random variable is defined as

$$Z = U \exp \left(\frac{\delta}{2} (U^2) \right), \delta \geq 0 \quad (4)$$

And the heavy-tailed Lambert $W \times F_X$ random variable can be defined as

$$Z = U \exp \left(\frac{\delta}{2} (U^2)^\alpha \right), \delta \geq 0 \alpha > 0 \quad (5)$$

Given that $U = (X - \mu_X)/\sigma_X$ where μ_X , σ_X , δ , and α are the mean and standard deviation of X and the parameters to control skewness and asymmetry, respectively. When $\delta = 0$, equation 4 reduces to a standard normal distribution. Equation 4 is the general form of Tukey's h distribution [26] and the basis for Morgenthaler and Tukey's [36] skewed, heavy tailed family of hh random variables.

$$Z = \begin{cases} U \exp \left(\frac{\delta_l}{2} (U^2)_l^\alpha \right), \delta_l \geq 0 \alpha_l > 0 \\ U \exp \left(\frac{\delta_r}{2} (U^2)_r^\alpha \right), \delta_r \geq 0 \alpha_r > 0 \end{cases} \quad (6)$$

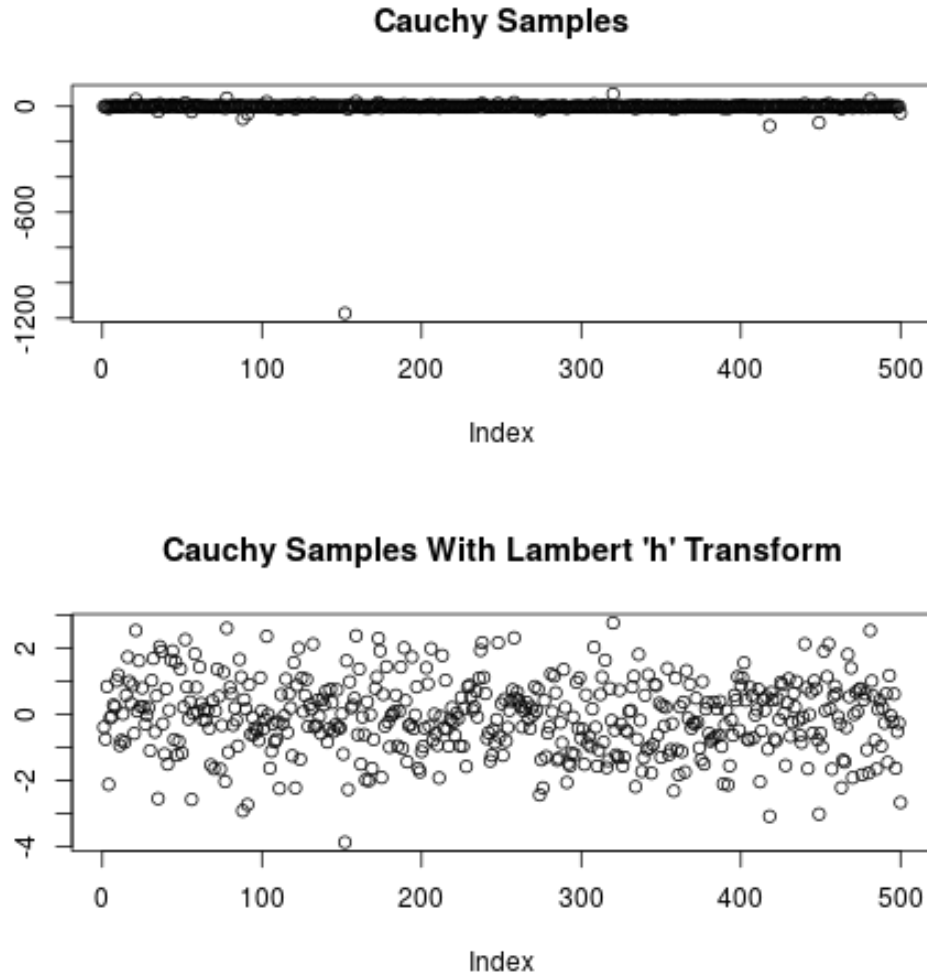


Figure 11: The top plot are samples from the cauchy distribution where the bottom plot represents the cauchy transformed variables with the hh distribution to make the samples approximately normal.

The function `Gaussianize()` is available in the package **LambertW** and has been made into a preprocessing function in **mlr**. Instead of calling `makeLearner()` to create a model, the function `makePreprocWrapperLambert()` can be used to create the model. Wrapping the model in this way allows the

pre-processing function to be embedded with the model itself. This stops users from accidentally biasing their models due to improperly applying pre-processing processes. For instance, if a user demeaned their entire data set and then split the data into train and test subsets, the training data will be biased because demeaning the model over both the train and test data gives the training data information about the mean of the test data. What should happen instead is, the user first splits the data into training and test data, and then demeans each separately. Similarly, cross-validation follows the above schema whereby each cross validation will be biased if the user demeaned all of the training data at one time. It is very easy for a user, who assumes they have made a good faith attempt to not bias their model, to receive overconfident results. Embedding the preprocessing within the model itself allows **mlr** to overcome this.

The preprocessign scheme is broken down into two main components. One function to gather and estimate necessary parameters in the training data, then a function to apply the learned preprocess over the training and test data. In the context of Lambert $W \times F()$ transforms, the h , hh , or s distribution that gaussianizes the data is estimated from the training data. Then the estimated parameter values from the training set are used during prediction to gaussianize the test observations. The code below follows this methodology, creating the model with Lambert preprocessing, training the model, and then performing prediction. The end result to the user appears the same, but a significant amount of bias is reduced in the background.

```

# Make a Learner with Lambert WxF() learner
lamb.lrn = makePreprocWrapperLambert("classif.lda", type = "h")
lamb.lrn

## Learner classif.lda.preproc from package MASS
## Type: classif
## Name: ; Short name:
## Class: PreprocWrapperLambert
## Properties: numerics,factors,prob,twoclass,multiclass
## Predict-Type: response
## Hyperparameters: type=h,methods=IGMM,verbose=FALSE

lamb.trn = train(lamb.lrn,iris.task, subset = 1:120)
lamb.pred = predict(lamb.trn, iris.task, subset = 121:150)

# Do the non-LW version
trn = train(makeLearner("classif.lda"),iris.task, subset = 1:120)
pred = predict(trn, iris.task, subset = 121:150)
performance(lamb.pred)

## mmce
## 0.1

performance(pred)

## mmce
## 0.1

```

9 Stacking Forecasting Learners

Stacking is a form of ensemble learning [12] in which a learning algorithm is trained on the predictions of several other learning algorithms. Let $y_{i,m}$ be the prediction at time i of model m . Given an aggregation function ϕ , a stacked forecast learner [11] is represented as

$$\tilde{y}_{i+1} = \phi(\tilde{y}_{i+1,1}, \tilde{y}_{i+1,2}, \dots, \tilde{y}_{i+1,m}, \sum_{j=1}^m \epsilon_{i+1,j}) \quad (7)$$

For a simple ϕ such as the ensemble average then equation 7 becomes

$$\tilde{y}_{i+1} = \frac{\tilde{y}_{i+1,1} + \tilde{y}_{i+1,2} + \dots + \tilde{y}_{i+1,m}}{m} \quad (8)$$

In section 9.1 the simple model average is used to show how stacked forecast models are built in **mlr**. Section 9.2 does a more advanced method of ensemble averaging involving the forecast of endogeneous variables.

9.1 Stacking Univariate Learners

For this example the models TBATS, GARCH, and ARFIMA [24] are stacked together and averaged on the climate task data. A resample description is made, and the function `makeLearners()` is used to start multiple learners at the same time.

```
resamp.sub = makeResampleDesc("GrowingCV",
                             horizon = 35L,
                             initial.window = .90,
                             size = nrow(getTaskData(climate.task)),
                             skip = .01
                             )
lrns = makeLearners(c("fcregr.tbats", "fcregr.garch",
                     "fcregr.arfima"))
```

The function `makeStackedLearner()` takes the initialized learners and sets the meta information for stacking. This method uses simple model averaging

such as 8, however a super learner [46] can be used here, where $\phi()$ becomes another machine learning model.

```
stack.forecast = makeStackedLearner(base.learners = lrns,  
                                   predict.type = "response",  
                                   method = "average")
```

Each of the stacked learners are tuned over the cross product of all model parameters. This leads to a change in design where, given that some models may have the same argument names, the full name of the model is placed before the argument. This leads to longer code, but tuning over the cross product of the models allows for a more honest perspective of how each model interacts in the stack.

```
# Simple param set for tuning sub learners  
ps = makeParamSet(  
  makeDiscreteParam("fcregr.tbats.h", values = 35),  
  makeDiscreteParam("fcregr.garch.n.ahead", values = 35),  
  makeDiscreteParam("fcregr.arfima.h", values = 35),  
  makeDiscreteParam("fcregr.arfima.estim", values = "ls"),  
  makeDiscreteParam(id = "fcregr.garch.model",  
                    values = c("csGARCH")),  
  makeIntegerVectorParam(id = "fcregr.garch.garchOrder",  
                        len = 2L, lower = c(1),  
                        upper = c(6)),  
  makeIntegerVectorParam(id = "fcregr.garch.armaOrder",  
                        len = 2L, lower = c(1),  
                        upper = c(4)),  
  makeDiscreteParam(id = "fcregr.garch.distribution.model",  
                    values = c("norm", "std", "jsu")),  
  makeDiscreteParam("fcregr.tbats.test",  
                    values = c("kpss", "adf", "pp")),  
  makeIntegerParam("fcregr.tbats.max.P", lower = 0, upper = 3),
```

```

    makeIntegerParam("fcregr.tbats.max.Q", lower = 0, upper = 2)
  )
  ctrl = makeTuneControlRtrace(maxExperiments = 400L)
  ## tuning
  library(parallelMap)
  parallelStartSocket(7)
  configureMlr(on.learner.error = "warn")
  set.seed(1234)
  fore.tune = tuneParams(stack.forecast, climate.task,
                        resampling = resamp.sub,
                        par.set = ps, control = ctrl,
                        measures = mase, show.info = FALSE)

  parallelStop()
  fore.tune

```

The rest of the modeling process flows in a way similar to the standard training and predicting schema. the function `setHyperPars2()` takes the best parameter models from the tuning process and assigns it to the final model to train over all of the data. Training and prediction are handled in the same manner as univariate forecasters.

```

# get hyper params
stack.forecast.tune = setHyperPars2(stack.forecast,fore.tune$x)
# Train the final best models and predict
stack.forecast.mod = train(stack.forecast.tune,climate.task)
stack.forecast.pred = predict(stack.forecast.mod,
                             newdata = m4.test)

stack.forecast.pred

## Prediction: 35 observations
## predict.type: response
## threshold:
## time: 0.00
##           truth response
## 2009-09-06      7 4.818892

```

```
## 2009-09-07    12 6.996835
## 2009-09-08    14 7.721747
## 2009-09-09    10 7.918065
## 2009-09-10    10 7.880770
## 2009-09-11    16 8.030163
## ... (35 rows, 2 cols)

performance(stack.forecast.pred,mase,climate.task)

##          mase
## 0.06057101
```

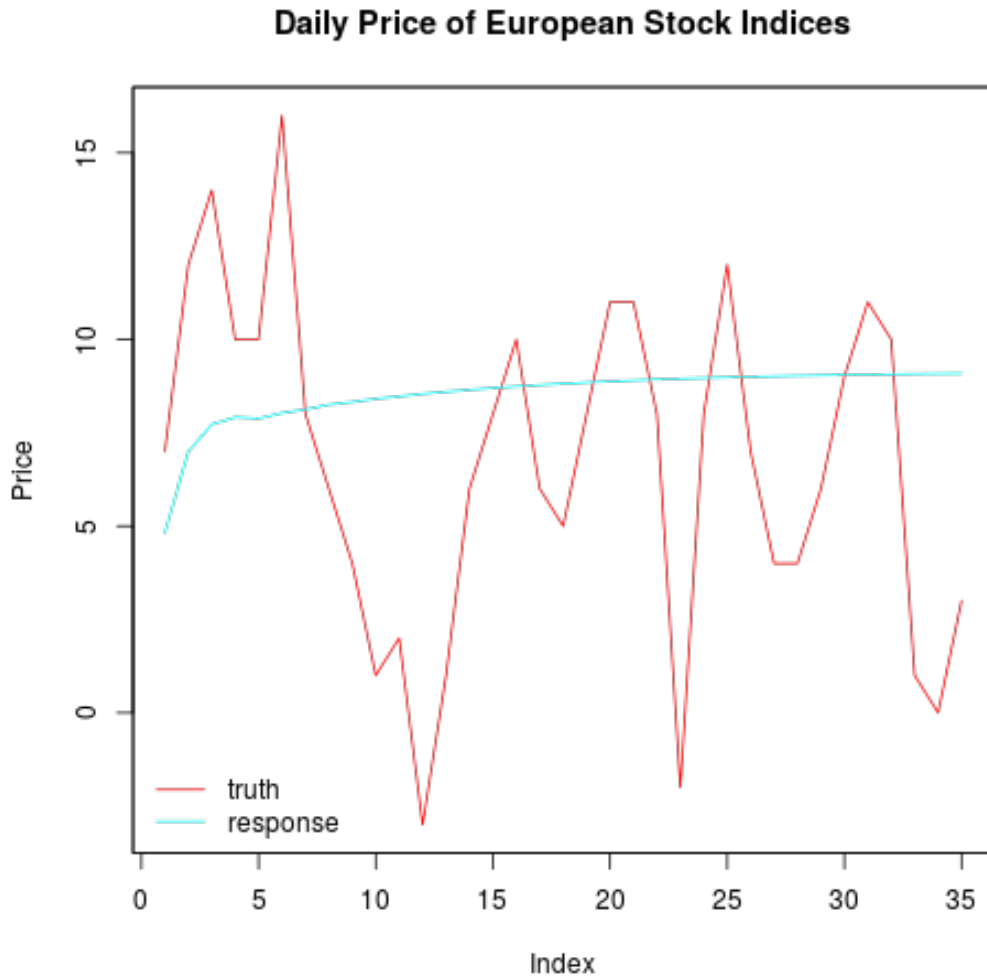


Figure 12: The forecast of the M4 data using the Stacked ARFIMA, TBATS, and GARCH model

9.2 Multivariate Stacked Learner

When there is a single target variable with multiple predictors stacked learning can be used with multivariate forecasters to forecast the predictors and have a machine learning model train over the forecasts of all variables. For this,

equation 7 can be modified to include forecasts of other predictors x_i, k where k is the index for each predictor variable

$$\tilde{y}_{i+1} = \phi(\tilde{y}_{i+1,1}, \dots, \tilde{y}_{i+1,m}, \tilde{x}_{i+1,1}, \dots, \tilde{x}_{i+1,k}, \sum_{j=1}^m \epsilon_{i+1,j}, \sum_{j=1}^k \epsilon_{i+1,j}) \quad (9)$$

In the example below, a boosted glm [8] is used as a super learner over a sparse lag multivariate VAR model to forecast FTSE prices. A resampling strategy is creating for both the underlying stacked learner and the super learner.

```
multfore.task = makeMultiForecastRegrTask(id = "bigvar",
                                          data = eu.train,
                                          target = "FTSE")

resamp.sub = makeResampleDesc("GrowingCV",
                              horizon = 32L,
                              initial.window = .90,
                              size = nrow(getTaskData(multfore.task)),
                              skip = .01
)

resamp.super = makeResampleDesc("CV", iters = 3)
```

In `makeStackedLearner()`, the super learner argument contains the boosted glm model.

```
base = c("mfcregr.BigVAR")
lrns = lapply(base, makeLearner)
lrns = lapply(lrns, setPredictType, "response")
lrns[[1]]$par.vals$verbose = FALSE
```

```
stack.forecast = makeStackedLearner(base.learners = lrns,
                                   predict.type = "response",
                                   super.learner = makeLearner("regr.glmboost",
                                                             family = "Laplace"),
                                   method = "growing.cv",
                                   resampling = resamp.sub)
```

Just as with univariate stacked forecasting models, a parameter set is created for the multivariate VAR model and tuning is done with `tuneParams()`.

```
ps = makeParamSet(
  makeDiscreteParam("mfcregr.BigVAR.p", values = 9),
  makeDiscreteParam("mfcregr.BigVAR.struct",
                    values = "SparseLag"),
  makeNumericVectorParam("mfcregr.BigVAR.gran", len = 2L,
                         lower = 35, upper = 50),
  makeDiscreteParam("mfcregr.BigVAR.h", values = 32),
  makeDiscreteParam("mfcregr.BigVAR.n.ahead", values = 32)
)

## tuning
library(parallelMap)
parallelStartSocket(4)
configureMlr(on.learner.error = "warn")
set.seed(1234)
multfore.tune = tuneParams(stack.forecast, multfore.task,
                           resampling = resamp.sub, par.set = ps,
                           control = makeTuneControlGrid(),
                           measures = mase, show.info = FALSE)
parallelStop()
```

Once the tuning is complete, the best model can be extracted with `setHyperPar2()` and the final model will be trained on all of the data. Since the multivariate model is used to produce forecasts for an single variable, univariate MASE is

used instead of the multivariate form of MASE.

```
stack.forecast.f = setHyperPars2(stack.forecast,  
                                multfore.tune$x)  
multfore.train = train(stack.forecast.f,multfore.task)
```

```
multfore.pred = predict(multfore.train,  
                        newdata = as.data.frame(eu.test))  
multfore.pred  
  
## Prediction: 32 observations  
## predict.type: response  
## threshold:  
## time: 0.02  
##  
##          truth response  
## 1998-07-12 07:50:46 5960.2 5629.466  
## 1998-07-13 17:32:18 5988.4 5629.378  
## 1998-07-15 03:13:50 5990.3 5629.423  
## 1998-07-16 12:55:23 6003.4 5629.519  
## 1998-07-17 22:36:55 6009.6 5629.655  
## 1998-07-19 08:18:27 5969.7 5629.759  
## ... (32 rows, 2 cols)  
  
performance(multfore.pred, mase, task = multfore.task)  
  
##          mase  
## 0.2363337
```

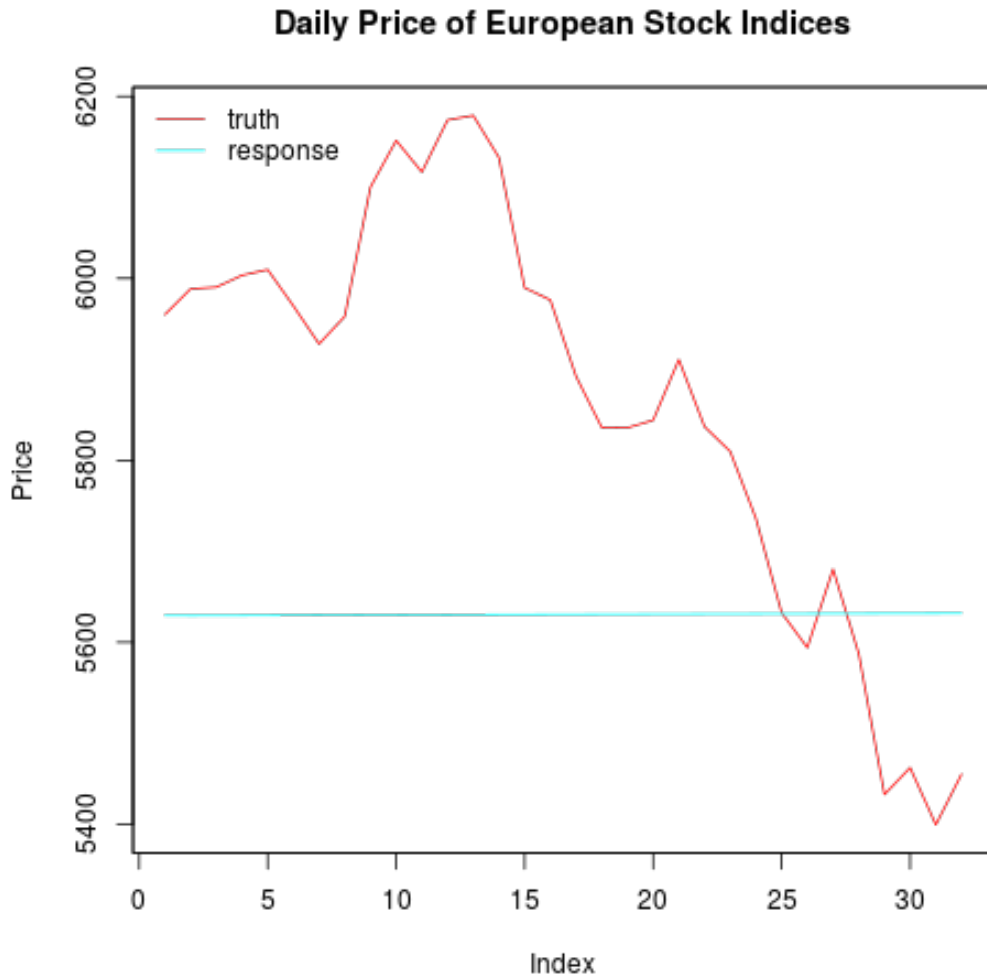


Figure 13: The forecast of the M4 data using the Stacked ARFIMA, TBATS, and GARCH model

10 Conclusion

The results of this paper show that creating a unified interface for forecasting models in R allows for better models through an automated methodology of resampling, preprocessing, model selection, stacking tuning, and training.

Building on the wide range of forecasting packages available in R, automating tasks such as windowing cross validation and model selection allow applied forecasters to spend less time dealing with the beauracracy of modeling and more time testing new models. New methods such as multivariate stacked learners, Lambert W transforms, and the ability to create arbitrary $AR(p, d)$ machine learning models allows researchers to easily experiment with new ideas. While the example models here are not perfect, this was mostly due to time. It will be easy for researchers to beat the models created in this paper.

Future research based on this package would involve tuning these models to see how useful they are in the real world. The classification forecast learners made available in the forecasting extension of **mlr** is a rising field, and with the ease of making models in **mlr** new research in this area can progress be easily replicatable. Updates to this package will include more multivariate and univariate forecast learners as well as new methods to stack models such as Bayesian averaging [41].

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