

TSML (Time Series Machine Learning)

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ABSTRACT

Over the past years, the industrial sector has seen many innovations brought about by automation. Inherent in this automation is the installation of sensor networks for status monitoring and data collection. One of the major challenges in these data-rich environments is how to extract and exploit information from these large volume of data to detect anomalies, discover patterns to reduce downtimes and manufacturing errors, reduce energy usage, predict faults/failures, effective maintenance schedules, etc. To address these issues, we developed **TSML**. Its technology is based on using the pipeline of lightweight filters as building blocks to process huge amount of industrial time series data in parallel.

Keywords

Julia, Time Series, Machine Learning, Filters, Feature Extraction, Classification, Prediction, Aggregation, Imputation

1. Introduction

TSML[?] is a *Julia*[?] package for time series data processing, classification, and prediction. It provides common API for ML libraries from Python's *Scikit-Learn*, R's *Caret*, and native *Julia* MLs for seamless integration of heterogeneous libraries to create complex ensembles for robust time series prediction, clustering, and classification. **TSML** has the following features:

- (1) data type clustering/classification for automatic data discovery
- (2) aggregation based on date/time interval
- (3) imputation based on symmetric Nearest Neighbors
- (4) statistical metrics for data quality assessment and classification input features
- (5) ML wrapper with more than 100+ libraries from caret, scikit-learn, and julia
- (6) date/value matrix conversion of 1-D time series using sliding windows to generate features for ML prediction
- (7) pipeline API for high-level description of the processing workflow
- (8) specific cleaning/normalization workflow based on data type
- (9) automatic selection of optimised ML model
- (10) automatic segmentation of time-series data into matrix form for ML training and prediction
- (11) extensible architecture using just two main interfaces: fit and transform
- (12) meta-ensembles for automatic feature and model selection
- (13) support for distributed/threaded computation for scalability and speed

The **TSML** package assumes a two-column input for any time series data composed of *dates* and *values*. The first part of the workflow aggregates values based on the specified date/time interval which minimizes occurrence of missing values and noise. The aggregated data is then left-joined to the complete sequence of dates in a specified date/time interval. Remaining missing values are replaced by the median/mean or user-defined aggregation function of the *k*-nearest neighbors (*KNN*) where *k* is the symmetric distance from the location of missing value. This approach can be called several times until there are no more missing values.

For prediction tasks, **TSML** extracts the date features and convert the value column into matrix form parameterized by the size and stride of the sliding window. The final part joins the date features and the value matrix to serve as input to the ML with the output representing the values of the time periods to be predicted ahead of time.

TSML uses a pipeline which iteratively calls the *fit* and *transform* families of functions relying on *multiple dispatch* to dynamically select the correct algorithm from the steps outlined above. Machine learning functions in **TSML** are wrappers to the corresponding Scikit-Learn, Caret, and native Julia ML libraries. There are more than hundred classifiers and regression functions available using a common API.

2. TSML Workflow

TSML workflow borrows the idea of the *Unix* pipeline[?, ?]. The main elements in a pipeline are series of filters or transformers each performing one specific task and does it well.

To illustrate, below describes the main steps in using **TSML**. First, we create filters for csv reading, aggregation, imputation, and data quality assessment.

```
fname = joinpath(dirname(pathof(TSML)),
    "../data/testdata.csv")
csvfilter = CSVDateValReader(Dict(
    :filename=>fname,
    :dateformat=>"dd/mm/yyyy HH:MM"))
valgator = DateValgator(Dict(
    :dateinterval=>Dates.Hour(1)))
valputer = DateValNner(Dict(
    :dateinterval=>Dates.Hour(1)))
stfier = Statifier(Dict(:processmissing=>true))
```

We can then setup a pipeline containing these filters to process the csv data by aggregating the time series hourly and check the data quality using the *Statifier* filter.

```
apipeline = Pipeline(Dict(
    :transformers => [csvfilter, valgator, stfier]))
```

```
fit!(apipeline)
mystats = transform!(apipeline)
@show mystats
```

tstart	tend	sfreq	count	max	min	median	mean	q1	q2	q25	q75	q8	q9
DateTime	DateTime	Float64	Int64	Float64									
2014-01-01T00:00:00	2015-01-01T00:00:00	0.999886	3830	18.8	8.5	10.35	11.557	9.9	10.0	10.0	12.3	13.0	16.0
kurtosis	skewness	variation	entropy	autocor	pacf	bmedian	bmean	bq25	bq75	bmin	bmax		
Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64
0.730635	1.41283	0.200055	-1.09145e5	4.39315	1.04644	5.0	10.5589	3.0	6.0	1.0	2380.0		

Fig. 1. Data Quality Statistics. Column names starting with 'b' refer to statistics of contiguous blocks of missing data.

Calling the *fit* and *transform* in the pipeline iteratively calls the corresponding *fit* and *transform* within each filter. This common API relying on *Julia's* multi-dispatch mechanism greatly simplifies the implementations, operations, and understanding of the entire workflow. In addition, extending **TSML** functionality is just a matter of creating a new data type filter and define its own *fit* and *transform* functions.

In the *Statifier* filter result, blocks of missing data is indicated by column names starting with *b*. Running the code above indicates that there are plenty of missing data blocks. We can add the *ValNner* filter to perform *k*-nearest neighbour imputation and check the statistics:

```
bpipeline = Pipeline(Dict(
    :transformers => [csvfilter, valgator,
                    valputer, stfier]))
fit!(bpipeline)
imputed = transform!(bpipeline)
@show imputed
```

tstart	tend	sfreq	count	max	min	median	mean	q1	q2	q25	q75	q8	q9
DateTime	DateTime	Float64	Int64	Float64									
2014-01-01T00:00:00	2015-01-01T00:00:00	0.999886	8761	18.8	8.5	10.0	11.1362	9.95	10.0	10.0	11.5	12.0	14.95
kurtosis	skewness	variation	entropy	autocor	pacf	bmedian	bmean	bq25	bq75	bmin	bmax		
Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64	Float64
2.37274	1.87452	0.187997	-2.36714e5	4.47886	1.06917	NaN							

Fig. 2. Statistics after Imputation. All statistics for blocks of missing data indicate NaN to indicate the stats of empty set. This implies that all missing blocks are imputed.

The result in Fig. ?? now indicates *NaN* for all missing data statistics column because the set of missing blocks count is now empty.

We can also visualise our time series data using the *Plotter* filter instead of the *Statifier* as shown in Fig ?. Looking closely, you will see discontinuities in the plot due to blocks of missing data.

```
pltr=Plotter(Dict(:interactive => true))
plpipeline = Pipeline(Dict(
    :transformers => [csvfilter, valgator, pltr]))
fit!(plpipeline)
transform!(plpipeline)
```

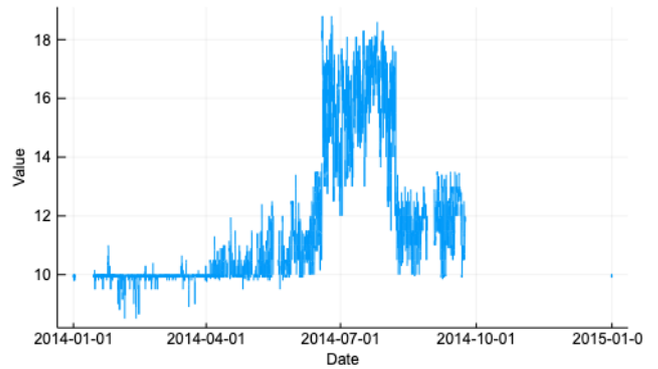


Fig. 3. Plot with Missing Data

Using the imputation pipeline described before, we can visualise the result by replacing the *Statifier* with the *Plotter* filter. Figure ?? shows the plot after imputation which gets rid of missing data.

```
bplpipeline = Pipeline(Dict(
    :transformers => [csvfilter, valgator,
                    valputer, pltr]))
fit!(bplpipeline)
transform!(bplpipeline)
```

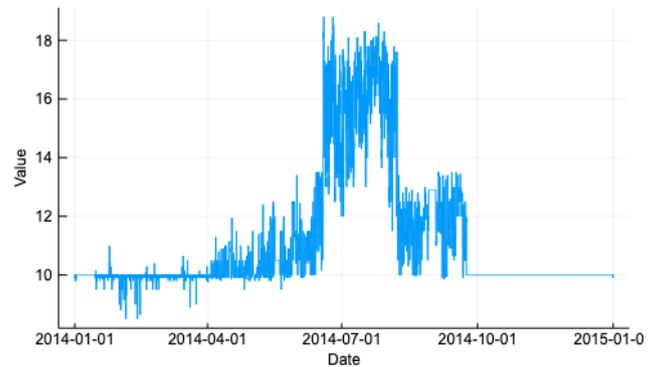


Fig. 4. Plot after Data Imputation

2.1 Processing Monotonic Time Series

In this subsection, we will show additional filters we use to handle monotonic data which are commonly employed in energy/water meter sensors as well as in footfall sensors. In the former case, the time series type is strictly monotonically increasing while in the latter case, the monotonicity happens daily by resetting the sensor to certain baseline at the end of the day. We use the filter called *Monotonicer* which automatically detects the two types of monotonic sensors and apply the normalisation accordingly.

```
mono = joinpath(dirname(pathof(TSML)),
    "../data/typedetection/monotonic.csv")
monocsv = CSVDateValReader(Dict(
    :filename=>mono,
    :dateformat=>"dd/mm/yyyy HH:MM"))
```

Let us plot the monotonic data with the usual workflow of aggregating and imputing the data first.

```
monopipeline = Pipeline(Dict(
    :transformers => [monofilecsv, valgator,
                     valputer, pltr]))
fit!(monopipeline)
transform!(monopipeline)
```

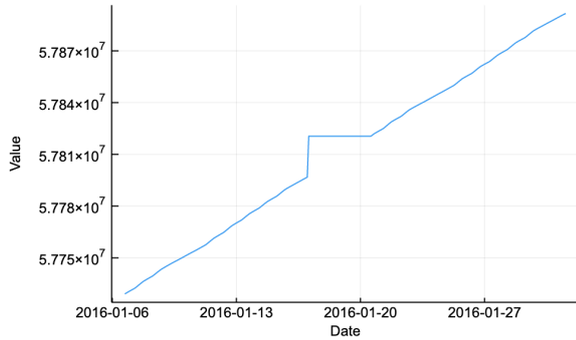


Fig. 5. Monotonic Time Series

Let us now normalise the data by adding the *Monotonicer* filter in the pipeline and plot it.

```
mononicer = Monotonicer(Dict())
monopipeline = Pipeline(Dict(
    :transformers => [monofilecsv, valgator, valputer,
                     mononicer, pltr]))
fit!(monopipeline)
transform!(monopipeline)
```

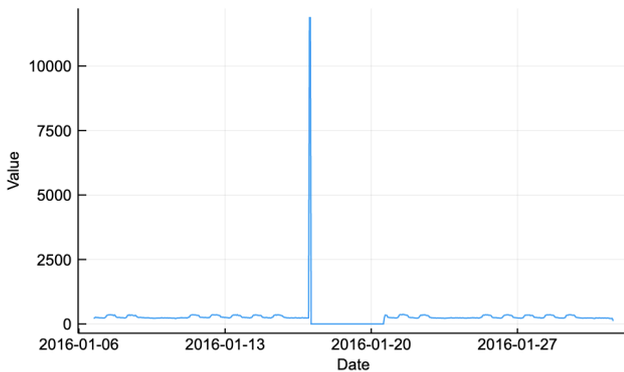


Fig. 6. Normalized Monotonic Time Series

The presence of outlier due to some random errors during meter reading becomes obvious after the normalisation. To remedy this issue, we can add the *Outliernicer* filter which detects outliers and replace them by using the *KNN* imputation technique used by the *DateValNNer* filter.

```
outliernicer = Outliernicer(
    Dict(:dateinterval=>Dates.Hour(1)))
monopipeline = Pipeline(Dict(
    :transformers => [monofilecsv, valgator, valputer,
                     mononicer, outliernicer, pltr]))
fit!(monopipeline)
transform!(monopipeline)
```

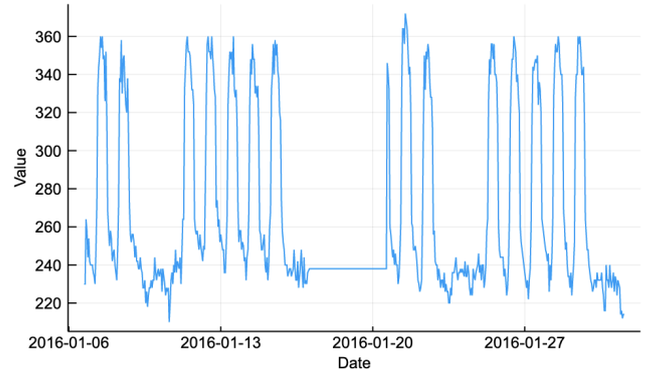


Fig. 7. Normalized Monotonic Time Series With Outlier Removal Filter

2.2 Processing Daily Monotonic Time Series

We follow similar workflow in the previous subsection to normalize daily monotonic time series. We visualise first the original data, apply the *Monotonicer* and *Outliernicer* filters, and plot the cleaned data to make sure monotonicity is gone.

```
dailymono = joinpath(dirname(pathof(TSML)),
    "../data/type-detection/dailymonotonic.csv")
dailymonocsv = CSVDateValReader(Dict(
    :filename=>dailymono,
    :dateformat=>"dd/mm/yyyy HH:MM"))
dailymonopipeline = Pipeline(Dict(
    :transformers => [dailymonocsv, valgator,
                     valputer, pltr]))
fit!(dailymonopipeline)
transform!(dailymonopipeline)
```

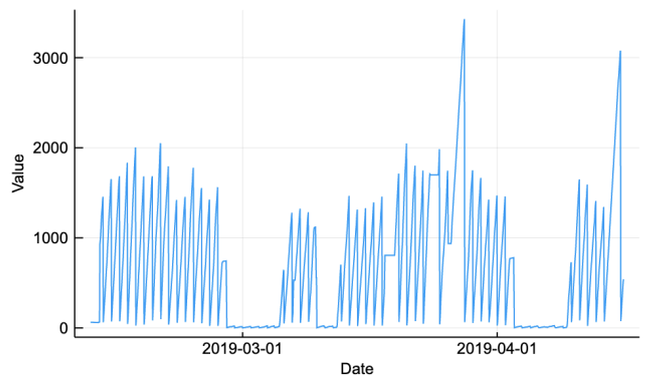


Fig. 8. Daily Monotonic Time Series

As seen in the plot, the values in daily monotonic time series is monotonically increasing and resets to certain baseline the following day and starts to rise again until the end of the daily cycle.

Let us now reuse the *Monotonicer* filter in the previous subsection to normalise the data.

```
dailymonopipeline = Pipeline(Dict(
  :transformers => [dailymonocsv, valgator,
                   valputer, mononicer, pltr]))
fit!(dailymonopipeline)
transform!(dailymonopipeline)
```

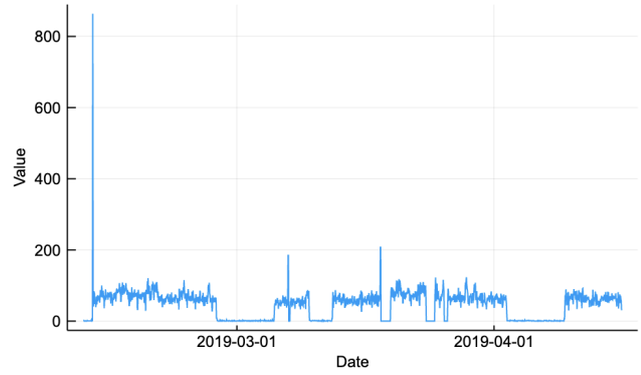


Fig. 9. Normalized Daily Monotonic Time Series

To remove outliers, we can reuse the *Outliernicer* filter in the previous subsection and plot the cleaned data.

```
dailymonopipeline = Pipeline(Dict(
  :transformers=>[dailymonocsv, valgator, valputer,
                 mononicer, outliernicer, pltr]))
fit!(dailymonopipeline)
transform!(dailymonopipeline)
```

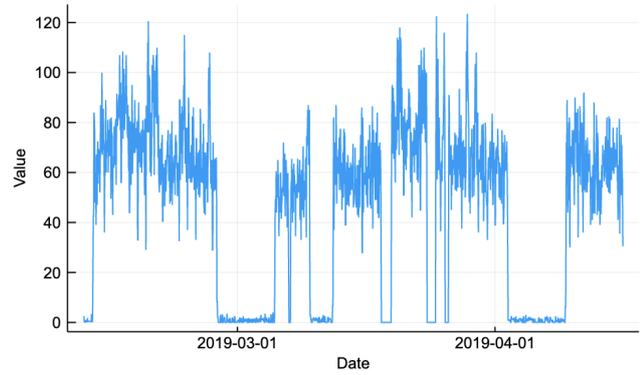


Fig. 10. Normalized Daily Monotonic Time Series with Outlier Detector

3. Time Series Classification

We can use the knowledge we learned in setting up the **TSML pipeline** and *filters* or *transformers* to build higher level operations to solve a specific industrial problem. One major problem which we consider relevant because it is a common issue in IOT (Internet of Things) is the time series classification. This problem is prevalent nowadays due to the increasing need to use many sensors to monitor status in different aspects of industrial operations and maintenance of cars, buildings, hospitals, supermarkets, homes, and cities.

Rapid deployment of these sensors result to many of them not properly labeled or classified. Time series classification is a significant first step for optimal prediction and anomaly detection. Identifying the correct sensor data types can help in the choice of what the most optimal prediction model to use for actuation or pre-emption to minimise wastage in resource utilisation. To successfully perform the latter operations, it is necessary to identify first the time series type so that appropriate model and cleaning routines can be selected for optimal model performance. The *TSClassifier* filter aims to address this problem and its usage is described below.

First, we setup the locations of files for training, testing, and saving the model. Next, we start the training phase by calling *fit* which loads file in the training directory and learn the mapping between their statistic features extracted by *Statifier* with their types indicated by a substring in their filenames. Once the training is done, the final model is saved in the *model* directory which will be used for testing accuracy and classifying new time series datasets.

The code below initialises the *TSClassifier* with the locations of the training, testing, and modelling repository. Training is carried out by the *fit* function which extracts the stat features of the training data and save them as a DataFrame input to the *RandomForest* classifier. The trained model is saved and used during testing.

```
trdirname = joinpath(dirname(pathof(TSML)),
  "../data/realdatatsclassification/training")
tstdirname = joinpath(dirname(pathof(TSML)),
  "../data/realdatatsclassification/testing")
modeldirname = joinpath(dirname(pathof(TSML)),
  "../data/realdatatsclassification/model")
tscl = TSClassifier(Dict(
  :trdirectory=>trdirname,
  :tstdirectory=>tstdirname,
  :modeldirectory=>modeldirname,
  :num_trees=>75)
)
fit!(tscl)
predictions = transform!(tscl)
@show testingAccuracy(predictions)
```

Figure ?? shows: a) a snapshot of the output during training and testing which extracts the statistical features of the time series; and b) the testing performance of the classifier. The training and testing data are labeled based on their sensor type for easier validation. The labels are not used as input during training. The classification workflow is purely driven by the statistical features. The prediction indicates 80% accuracy.

4. Extending TSML with Scikit-Learn and Caret

In the latest **TSML** version (2.3.4 and above), we refactored the base **TSML** to only include pure *Julia* code implementations and moved the external libs and binary dependencies into the **TSMLextra** package. One major reason is to have a smaller code

```
getting stats of RetTemp51.csv
getting stats of AirOffTemp4.csv
getting stats of AirOffTemp5.csv
getting stats of Energy5.csv
getting stats of Pressure5.csv
getting stats of RetTemp31.csv
.....
.....
fname  predtype
String SubStrin...
1 AirOffTemp4.csv AirOffTemp
2 AirOffTemp5.csv AirOffTemp
3 Energy5.csv Energy
4 Pressure5.csv Pressure
5 RetTemp31.csv Energy
```

Fig. 11. Extraction of Statistical Features for Training and Prediction. By convention, the *TSClassifier* validates the ground truth based on the filename which contains the label of the sensor type disregarding the numerical component.

base so that it can be easily maintained and rapidly deployed in a dockerized solution for *Kubernetes* or IBM's *OpenShift* cluster. Moreover, smaller codes make static compilation fast for smaller docker image in cloud deployment.

There are cases where the main task of time series classification requires more complex ensemble model using hierarchy or tree structure where members are composed of heterogeneous ML learners derived from binaries in different languages. For illustration purposes, we will show how to ensemble ML libraries from *Scikit-Learn* and *Caret* using *TSML* meta-ensembles that support the *fit* and *transform* APIs.

4.1 Parallel TSML Using Distributed Workflow

We will use *Julia*'s built-in support for parallelism by using the *Distributed* standard library. We also let *Julia* detect the number of processors available and activate them using the following statements:

```
using Distributed
nprocs() == 1 && addprocs()
```

With several workers active, we use the *@everywhere* macro to load the necessary filters and transformers to all workers.

```
@everywhere using TSML
@everywhere using TSMLextra
@everywhere using DataFrames
@everywhere using Random
@everywhere using Statistics
@everywhere using StatsBase: iqr
@everywhere using RDatasets
```

With all the necessary *TSML* functions loaded, we can now setup the different MLs starting with some learners from *Caret* and *Scikit-Learn*. The list is not exhaustive for demonstration purposes.

```
# Caret ML
@everywhere caret_svmlinear =
    CaretLearner(Dict(:learner=>"svmlinear"))
```

```
@everywhere caret_treebag =
    CaretLearner(Dict(:learner=>"treebag"))

# Scikit-Learn ML
@everywhere sk_knn =
    SKLearner(Dict(:learner=>"KNeighborsClassifier"))
@everywhere sk_gb =
    SKLearner(Dict(:learner=>
        "GradientBoostingClassifier",
        :impl_args=>Dict(:n_estimators=>10)))
@everywhere sk_extratree =
    SKLearner(Dict(:learner=>"ExtraTreesClassifier",
        :impl_args=>Dict(:n_estimators=>10)))
@everywhere sk_rf =
    SKLearner(Dict(:learner=>
        "RandomForestClassifier",
        :impl_args=>Dict(:n_estimators=>10)))
```

Let us setup ML instances from a pure *Julia* implementation of learners and ensembles wrapped from the *DecisionTree.jl* package [?, ?, ?, ?].

```
# Julia ML
@everywhere jrf = RandomForest()
@everywhere jpt = PrunedTree()
@everywhere jada = Adaboost()

# Julia Ensembles
@everywhere jvote_ens=VoteEnsemble(Dict(
    :learners=>[jrf, jpt, sk_gb, sk_extratree, sk_rf]))
@everywhere jstack_ens=StackEnsemble(Dict(
    :learners=>[jrf, jpt, sk_gb, sk_extratree, sk_rf]))
@everywhere jbest_ens=BestLearner(Dict(
    :learners=>[jrf, sk_gb, sk_rf]))
@everywhere jsuper_ens=VoteEnsemble(Dict(
    :learners=>[jvote_ens, jstack_ens,
        jbest_ens, sk_rf, sk_gb]))
```

Next, we setup the pipeline for training and prediction.

```
@everywhere function predict(learner,
    data, train_ind, test_ind)
    features = convert(Matrix, data[:, 1:(end-1)])
    labels = convert(Array, data[:, end])
    # Create pipeline
    pipeline = Pipeline(
        Dict(
            :transformers => [
                OneHotEncoder(), # nominal to bits
                Imputer(), # Imputes NA values
                StandardScaler(), # normalize
                learner # Predicts labels on instances
            ]
        )
    )
    # Train
    fit!(pipeline, features[train_ind, :],
        labels[train_ind])
    # Predict
    predictions = transform!(pipeline,
        features[test_ind, :])
    # Assess predictions
    result = score(:accuracy,
        labels[test_ind], predictions)
    return result
end
```

Finally, we setup the *parallelmodel* function to run different learners distributed to different workers running in parallel relying on *Julia*'s native support of parallelism. Take note that there are two parallelisms in the code. The first one is the distribution of

task in different trials and the second one is the distribution of tasks among different models for each trial. It is interesting to note that with this relative compact function definition, the *Julia* language makes it easy to define a parallel task within another parallel task in a straightforward manner without any problem.

```
function parallelmodel(learners::Dict,
                      data::DataFrame; trials=5)
  models=collect(keys(learners))
  ctable=@distributed(vcat) for i=1:trials
    # Split into training and test sets
    Random.seed!(3i)
    (trndx, tstndx) = holdout(size(data, 1), 0.20)
    acc=@distributed(vcat) for model in models
      res=predict(learners[model],
                 data, trndx, tstndx)
      println("trial ", i, ", ", "model", " => ",
             round(res))
      [model res i]
    end
    acc
  end
  df = ctable |> DataFrame
  rename!(df, :x1=>:model, :x2=>:acc, :x3=>:trial)
  gp=by(df, :model) do x
    DataFrame(mean=mean(x.acc), std=std(x.acc),
              n=length(x.acc))
  end
  sort!(gp, :mean, rev=true)
  return gp
end
```

We benchmark the performance of the different machine learners by creating a dictionary of workers containing instances of learners from *Caret*, *Scikit-Learn*, and *Julia* libraries. We pass the dictionary of learners to the `parallelmodel` function for evaluation.

```
learners=Dict(
  :jvote_ens=>jvote_ens, :jstack_ens=>jstack_ens,
  :jbest_ens=>jbest_ens, :jrf=>jrf, :jada=>jada,
  :jsuper_ens=>jsuper_ens,
  :crt_svmlinear=>caret_svmlinear,
  :crt_treebag=>caret_treebag,
  :skl_knn=>sk_knn, :skl_gb=>sk_gb,
  :skl_extratree=>sk_extratree, :sk_rf=>sk_rf
)

datadir = joinpath("tsdata/")
tsdata = extract_features_from_timeseries(datadir)
first(tsdata, 5)

respar = parallelmodel(learners, tsdata; trials=3)
```

```
From worker 2: trial 2, crt_treebag => 100.0
From worker 2: trial 3, skl_extratree => 100.0
From worker 3: trial 1, crt_treebag => 100.0
From worker 3: trial 2, jrf => 100.0
From worker 4: trial 3, jrf => 25.0
From worker 6: trial 1, crt_svmlinear => 100.0
From worker 6: trial 2, crt_svmlinear => 75.0
From worker 6: trial 3, crt_svmlinear => 75.0
```

Fig. 12. Output During Training with Several Workers

The data used in the experiment are sample snapshots of the data in our building operations. For reproducibility, the data can be found in the `juliacon2019-paper` branch of **TSML** in *GitHub*: `/data/benchmark/tsclassifier`. There are four time series types,

	model	mean	std	n
	Any	Float64	Float64	Int64
1	skl_extratree	91.6667	14.4338	3
2	crt_treebag	83.3333	28.8675	3
3	crt_svmlinear	83.3333	14.4338	3
4	jvote_ens	75.0	0.0	3
5	jrf	66.6667	38.1881	3
6	jbest_ens	66.6667	38.1881	3
7	skl_gb	66.6667	14.4338	3
8	jsuper_ens	66.6667	38.1881	3
9	skl_rf	66.6667	14.4338	3
10	skl_knn	66.6667	14.4338	3
11	jada	25.0	25.0	3
12	jstack_ens	16.6667	28.8675	3

Fig. 13. @distributed: Classification Performance in 3 Trials

namely: *AirOffTemp*, *Energy*, *Pressure*, and *RetTemp*. We took a minimal number of samples and classes for the sake of discussion and demonstration purposes in this paper.

Figures ?? and ?? show a snapshot of running workers exploiting the distributed library of *Julia* and the classification performance of each model, respectively. There are 8 workers running in parallel over 12 different machine learning classifiers.

From the results, *ExtraTree* from *Scikit-Learn* has the best performance with 91.67% accuracy followed by *TreeBag* and *SVMLinear* from *Caret* library with 83.33 % accuracy for both. With this workflow, it becomes trivial to search for optimal model by running them in parallel relying on *Julia* to do the low-level tasks of scheduling and queueing as well as making sure that the dynamically available compute resources such as cpu cores and memory resources are fairly optimised.

4.2 Parallel TSML Using Threads Workflow

With *Julia* 1.3, lightweight multi-threading support in *Julia* becomes possible. We will be using the pure *Julia*-written ML models because installing external dependencies such as *Caret* MLs through *RCall* package has some issues with the alpha version of *Julia* 1.3 at this point in time. We will update this documentation and add more MLs once the issues are resolved.

The main difference in the workflow between *Julia*'s distributed computation model compared to the threaded model is the presence of `@everywhere` macro in the former for each function defined to indicate that these function definitions shall be exported to all running workers. Since threaded processes share the same memory model with the *Julia* main process, there is no need for this macro. Instead, threading workflow requires the use of *mutex* locking in the update of the global dataframe that accumulates the prediction performance of models running in their respective threads. In similar observation with the distributed framework, the `threadedmodel` function contains two parallelism: threads in different trials and threads among models in each trial. The function is surprisingly compact to implement threads within threads without issues and the main bottleneck happens only during the update operation of the global `ctable` dataframe.

```
function threadedmodel(learners::Dict,
    data::DataFrame; trials=5)
    Random.seed!(3)
    models=collect(keys(learners))
    global ctable = DataFrame()
    @threads for i=1:trials
        # Split into training and test sets
        (train_ind, test_ind) =
            holdout(size(data, 1), 0.20)
        mtx = SpinLock()
        @threads for themodel in models
            res=predict(learners[themodel],
                data, train_ind, test_ind)
            println(themodel, " => ", round(res), ",
                thread=", threadid())
            lock(mtx)
            global ctable=vcat(ctable,
                DataFrame(model=themodel, acc=res))
            unlock(mtx)
        end
    end
    df = ctable |> DataFrame
    gp=by(df, :model) do x
        DataFrame(mean=mean(x.acc),
            std=std(x.acc), n=nrow(x))
    end
    sort!(gp, :mean, rev=true)
    return gp
end
```

Let us define a set of learners that are written in pure *Julia* for this thread experiment.

```
# Julia ML
jrf = RandomForest(Dict(:impl_args=>
    Dict(:num_trees=>500)))
jpt = PrunedTree()
jada = Adaboost(Dict(:impl_args=>
    Dict(:num_iterations=>20)))

## Julia Ensembles
jvote_ens=VoteEnsemble(Dict(:learners=>
    [jrf, jpt, jada]))
jstack_ens=StackEnsemble(Dict(:learners=>
    [jrf, jpt, jada]))
jbest_ens=BestLearner(Dict(:learners=>
    [jrf, jpt, jada]))
jsuper_ens=VoteEnsemble(Dict(:learners=>
    [jvote_ens, jstack_ens, jbest_ens]));
```

Let us run in parallel the different models using the same dataset with that of the distributed workflow.

```
using Base.Threads

learners=Dict(
    :jvote_ens=>jvote_ens,
    :jstack_ens=>jstack_ens,
    :jbest_ens=>jbest_ens,
    :jrf => jrf, :jada=>jada,
    :jsuper_ens=>jsuper_ens);

datadir = joinpath("tsdata/")
tsdata = extract_features_from_timeseries(datadir)

resthr = threadedmodel(learners, tsdata; trials=10)
```

Figures ?? and ?? show example snapshot of the running threads and the final result of classification, respectively. In this experi-

```
jvote_ens => 75.0, thread=8
jvote_ens => 100.0, thread=7
jbest_ens => 75.0, thread=3
jbest_ens => 50.0, thread=5
jbest_ens => 75.0, thread=1
jbest_ens => 50.0, thread=7
jrf => 50.0, thread=3
jbest_ens => 100.0, thread=2
jada => 75.0, thread=3
jbest_ens => 100.0, thread=6
jbest_ens => 75.0, thread=8
jbest_ens => 75.0, thread=4
jrf => 100.0, thread=5
jada => 100.0, thread=5
jrf => 100.0, thread=1
jrf => 100.0, thread=7
jada => 100.0, thread=1
jada => 50.0, thread=7
```

Fig. 14. Output During Training Using Several Threads

	model	mean	std	n
	Symbol	Float64	Float64	Int64
1	jada	85.0	17.4801	10
2	jrf	82.5	20.5818	10
3	jvote_ens	77.5	24.8607	10
4	jbest_ens	75.0	20.4124	10
5	jsuper_ens	75.0	28.8675	10
6	jstack_ens	47.5	24.8607	10

Fig. 15. @threads: Classification Performance in 10 Trials

ment, Adaboost has 85.0% accuracy followed by Random Forest with 82.5% accuracy.

5. Applications Using Public IoT Datasets

This section summarizes the results of applying the TSML workflow for classification tasks using data in this site: <http://www.timeseriesclassification.com/>. Among the hundreds of datasets available, we handpicked only 4 datasets, namely: ElectricDevices, RefrigerationDevices, FordB, and Earthquakes.

ElectricDevices dataset covers 251 households, sampled in 2-minute intervals over a month. The target is to collect how consumers use electricity within the home based on the devices they use at a particular time of the day. The dataset is highly imbalanced with the following total count in each class, respectively: 1394, 4187, 1606, 2639, 4275, 1252, 1284. This dataset is a good test on the robustness of the classifier to handle bias during training. The class distribution in training is consist of: 727, 2231, 851, 1471, 2406, 509, 728. For testing, the class distribution is: 667, 1956, 755, 1165, 1869, 743, 556. Using this highly imbalanced dataset is a good way to determine which among the classifiers has the best method to deal with this sampling bias. There are 96 input features and 6 output classes in a total of 16637 samples.

RefrigerationDevices dataset is based on similar study with that of ElectricalDevices dataset. The dataset uses the same set of households and sampling protocol for gathering electric consumption data from three types of refrigeration devices: Fridge/Freezer, Refrigerator, Upright Freezer. Unlike in ElectricDevices, the class distribution in RefrigerationDevices are balanced, i.e., 250 for each class in training and 125 for each class during testing. There are 720 input features and 3 output classes in a total of 750 samples.

Table 1. Classification Problems

Problem	Classes	Imbalance	Dimension
Electric Devices	6	Y	16637 x 96
Refrigeration Devices	3	N	750 x 720
Earthquakes	2	Y	461 x 512
FordB	2	N	4446 x 500

Table 2. Electric Devices

Model	MeanFscore	Std	Trials
c-rf	0.56	0.00	10
c-treebag	0.54	0.01	10
s-extratree	0.52	0.01	10
j-vote	0.51	0.01	10
j-rf	0.51	0.01	10
...

The Earthquakes classification problem requires predicting whether a major event is about to occur based on the most recent readings in the surrounding area. The data is aggregated hourly with the Rictor scale reading over 5 indicating a major event. There are 368 negative cases versus 93 positive cases of major events. This dataset is highly imbalanced in class distribution which makes predicting whether a major event occurs very problematic. The training set is composed of 264 negative examples and only 58 positive examples. The testing data is composed of 104 negative cases and only 35 positive cases. There are 512 input features and 2 output classes in a total of 461 samples. The highly imbalanced distribution of this dataset will provide a good way to determine which among the classifiers are robust to deal with this bias.

FordB dataset is a classification problem to diagnose whether a certain symptom exists in an automotive system. There are 500 measurements of engine noise. The training data were collected in typical operating conditions while the test data were collected under noisy conditions. The dataset is slightly imbalanced consisting of 1860 class1 vs 1776 class2 in training while 401 class1 and 409 class2 in testing. There are 500 input features and 2 output classes in a total of 4446 samples.

Table ?? summarizes the differences and similarities among the different classification problems under consideration. Two of the problems have highly imbalanced dataset, namely: Electric Devices and Earthquakes. Both FordB and Refrigeration Devices have no or slightly less imbalanced data.

One thing to note is that the Refrigeration dataset has almost the same number of features with its total number of samples. We expect that this will result into a much harder classification problem even though it does not suffer from imbalanced class distribution. Ideally, there must be more samples than features in order for the classifier to properly extract the correct subset of features for robust classification.

5.1 Results

Due to the data imbalance, the typical accuracy measurement will not be able to capture the performance of the algorithms because its value may be overshadowed by the dominant class. In this regard, we use F-score to measure the performance of a given classifier to each of its classes and get the mean of these F-scores.

Table ?? shows the top 5 performing classifiers together with the 2 worst performing classifiers. The first letter of each classifier's name indicates whether the classifier comes from (c)aret,

Table 3. Refrigeration Devices

Model	MeanFscore	Std	Trials
s-gb	0.51	0.00	10
c-rf	0.51	0.01	10
c-treebag	0.50	0.01	10
j-super	0.50	0.02	10
s-rf	0.48	0.03	10
...

Table 4. Earthquakes

Model	MeanFscore	Std	Trials
c-treebag	0.86	0.01	10
c-rf	0.86	0.00	10
c-rpart	0.86	0.00	10
s-extratree	0.85	0.01	10
s-gb	0.85	0.00	10
...

Table 5. FordB

Model	MeanFscore	Std	Trials
c-rf	0.66	0.01	10
s-knn	0.62	0.00	10
j-vote	0.61	0.01	10
j-rf	0.61	0.01	10
j-super	0.60	0.03	10
...

(s)cikitLearn, or (j)ulia. The table indicates that *Random Forest* and *TreeBag* from *Caret* performed the best followed by *ExtraTree* of *ScikitLearn*. Julia's *Vote* ensemble and *Random Forest* complete the top 5. On the other hand, Julia's *Adaboost* and *Caret*'s *RPart* are the worst classifiers for this problem.

Using similar naming convention, Table ?? indicates that *Gradient-Boost* from *ScikitLearn* and *Random Forest* from *Caret* are the best classifiers for the Refrigeration Devices problem. The two worst algorithms are the same as in Table ??.

For Earthquakes classification problem (Table ??), the top 3 best classifiers are dominated by those from *Caret* library, namely: *Tree-Bag*, *Random Forest*, and *RPart*. The worst performers are from *Caret*'s *SVML* and *Julia*'s *PartitionTree*.

For the FordB dataset (Table ??), the best performers are: *Random Forest* from *Caret*, *KNN* from *ScikitLearn*, and *Vote Ensemble* from *Julia*. The worst performers are similar to Tables ?? and ??.

5.2 Discussion

Comparing the performances of different classifiers among the four problems indicate that the most difficult problem to classify is the Refrigeration Devices while the easiest one is the Earthquakes problem. As we expected, there is not enough sample for the classifiers to learn the mapping in Refrigeration dataset relative to its feature dimension. One way to alleviate this problem is to remove features that are highly correlated or perform PCA to reduce the feature dimension. This is beyond the scope of the current paper which focuses mainly on the applicability of TSML MLs to different set of problems.

Among the classifiers, the *Random Forest* from *Caret* is the top performer in all problems. On the other hand, *TreeBag* from *Caret* and *Gradient Boosting* from *ScikitLearn* are the top performers in

Refrigeration Devices and Earthquakes, respectively. It is interesting to note that among the *Random Forest* implementations, the top performer is the *Caret's* version written by Breiman[?] who was the original author of the said algorithm in *fortran*. The superior performance of Breiman's *Random Forest* can be highlighted in the FordB classification performance. *kNN*, the next best classifier is 4% lower than Breiman's *Random Forest*. In all other cases, the next best performer has almost same performance with Breiman's *Random Forest*.

As we expected based on past studies, the different ensemble models dominated the top 5 performers. While not shown in the table, the most consistent worst performer is the *Caret's RPart*. This is consistent to Breiman's observation that *Random Forest* performs well by using unstable or inferior ML models in its leaves. In *Caret's Random Forest*, the leaves are composed of *RPart* models which has the poorest performance in all problems. The combination of boosting and bagging weak learners such as *RPart* make the *Random Forest* robust from datasets imbalances.

6. Summary and Conclusion

Packages for time series analysis are becoming important tools with the rapid proliferation of sensor data brought about by **IoT**. We created **TSML** as a time series machine learning framework which can easily be extended to handle large volume of time series data. **TSML** exploits the following *Julia* features: multiple dispatch, type inference, custom data types and abstraction, and parallel computations.

TSML main strength is the adoption of *UNIX* pipeline architecture containing filters or transformers to perform both preprocessing and modelling tasks. In addition, **TSML** uses a common machine learning API for both internal and external ML libraries, distributed and threaded support for modeling, and a growing collection of filters for preprocessing, classification, clustering, and prediction.

Extending **TSML** can easily be done by creating a custom data type filter and defining its corresponding *fit* and *transform* operations which the **TSML** pipeline iteratively calls for each filter in the workflow.