## TSSL Lab 4 - Recurrent Neural Networks

In this lab we will explore different RNN models and training procedures for a problem in time series prediction.

```
import numpy as np
import tensorflow as tf
from tensorflow import keras
from tensorflow.keras import layers
import pandas
import matplotlib.pyplot as plt

plt.rcParams["figure.figsize"] = (10,6) # Increase default size of plots
```

Set the random seed, for reproducibility

## 1. Load and prepare the data

We will build a model for predicting the number of sunspots. We work with a data set that has been published on Kaggle, with the description:

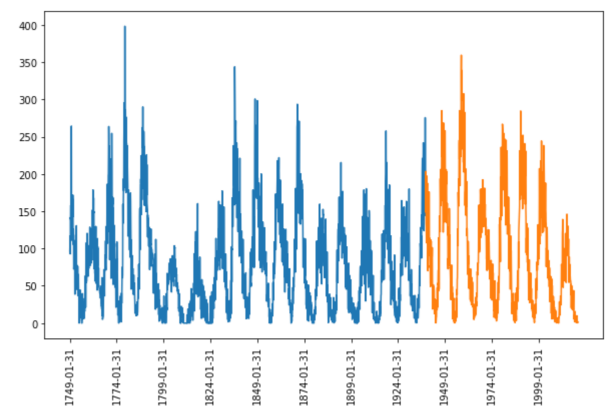
Sunspots are temporary phenomena on the Sun's photosphere that appear as spots darker than the surrounding areas. They are regions of reduced surface temperature caused by concentrations of magnetic field flux that inhibit convection. Sunspots usually appear in pairs of opposite magnetic polarity. Their number varies according to the approximately 11-year solar cycle.

The data consists of the monthly mean total sunspot number, from 1749-01-01 to 2017-08-31.

```
In [3]: # Read the data
data=pandas.read_csv('Sunspots.csv',header=0)
dates = data['Date'].values
y = data['Monthly Mean Total Sunspot Number'].values
ndata=len(y)
print(f'Total number of data points: {ndata}')

# We define a train/test split, here with 70 % training data
ntrain = int(ndata*0.7)
ntest = ndata-ntrain
print(f'Number of training data points: {ntrain}')
Total number of data points: 3252
Number of training data points: 2276
```

```
In [4]:
    plt.plot(dates[:ntrain], y[:ntrain])
    plt.plot(dates[ntrain:], y[ntrain:])
    plt.xticks(range(0, ndata, 300), dates[::300], rotation = 90); # Show only one tick
```



There is a clear seasonality to the data, but the amplitude of the peaks very quite a lot. Also, we note that the data is nonnegative, which is natural since it consists of counts of sunspots. However, for simplicity we will not take this constraint into account in this lab assignment and allow ourselves to model the data using a Gaussian likelihood (i.e. using MSE as a loss function).

From the plot we see that the range of the data is roughly [0,400] so as a simple normalization we divide by the constant MAX\_VAL=400.

```
In [5]:  MAX_VAL = 400 
 y = y/MAX_VAL
```

### 2. Baseline methods

Before constructing any sophosticated models using RNNs, let's consider two baseline methods,

- 1. The first baseline is a "naive" method which simply predicts  $y_{t} = y_{t-1}$ .
- 2. The second baseline is an AR(\$p\$) model (based on the implementation used for lab 1).

We evaluate the performance of these method in terms of mean-squared-error and meanabsolute-error, to compare the more advanced models with later on.

```
def evalutate_performance(y_pred, y, split_time, name=None):
    """This function evaluates and prints the MSE and MAE of the prediction.

Parameters
    ------
    y_pred : ndarrary
        Array of size (n,) with predictions.
    y : ndarray
        Array of size (n,) with target values.
    split_time : int
        The leading number of elements in y_pred and y that belong to the training d
```

```
# Compute error in prediction
resid = y - y_pred
# We evaluate the MSE and MAE in the original scale of the data, i.e. we add bac
train_mse = np.mean(resid[:split_time]**2)*MAX_VAL**2
test_mse = np.mean(resid[split_time:]**2)*MAX_VAL**2
train_mae = np.mean(np.abs(resid[:split_time]))*MAX_VAL
test_mae = np.mean(np.abs(resid[split_time:]))*MAX_VAL

# Print
print(f'Model {name}\n Training MSE: {train_mse:.4f}, MAE: {train_mae:.4f}\n
```

**Q1:** Implement the naive baseline method which predicts according to  $\hat{t}=y_{t-1}$ . Since the previous value is needed for the prediction we do not get a prediction at t=1. Hence, we evaluate the method by predicting values at t=2, t=2, t=2, t=3.

```
In [7]:

# Store the predictions in an array of length ndata-1. Note that there is a shift in # between the prediction and the observation sequence, since there is no prediction # Specifically, y_pred_naive[t] is a prediction of y[t+1], so the first element of y # second element of y, and so on. We will use the same "bookeeping convention" through you understand it!

y_pred_naive = y[0:ndata-1]#Starting from 0 index to ndata-1

evalutate_performance(y_pred_naive, # Predictions y[1:], # Correspondsing target values ntrain-1, # Number of leading elements in the input array name='Naive')

Model Naive
```

Training MSE: 776.5437, MAE: 19.3285 Testing MSE: 708.6360, MAE: 19.2256

Next, we consider a slightly more advanced baseline method, namely an AR(\$p\$) model.

```
# We import two functions that were written as part of lab 1
from tssltools_lab4 import fit_ar, predict_ar_1step

p=30 # Order of the AR model (set by a few manual trials)
ar_coef = fit_ar(y[:ntrain], p) # Fit the model to the training data

# Predict. Note that y contains both training and validation data,
# and the prediction is for the values y_{p+1}, ..., y_{n}.
y_pred_ar = predict_ar_1step(ar_coef, y)
```

Model AR
Training MSE: 603.8656, M

Training MSE: 603.8656, MAE: 17.3420 Testing MSE: 590.3732, MAE: 17.6221

## 3. Simple RNN

We will now construct a model based on a recurrent neural network. We will initially use the SimpleRNN class from *Keras*, which correspond to the basic Jordan-Elman network presented in the lectures.

Q2: Assume that we construct an "RNN cell" using the call layers.SimpleRNN(units = d, return\_sequences=True). Now, assume that an array X with the dimensions [Q,M,P] is fed as the input to the above object. We know that X contains a set of sequences (time series) with equal lengths. Specify which of the symbols \${Q,M,P}\$ that corresponds to each of the items below:

- The length of the sequences (number of time steps)
- The number of features (at each time step), i.e. the dimension of each time series
- The number of sequences

Furthermore, specify the values of \${Q,M,P}\$ for the data at hand (treated as a single time series).

*Hint:* Read the documentation for SimpleRNN to find the answer.

#### A2:

The length of the sequences (number of time steps): M, 3252(considering the entire data)

The number of features (at each time step), i.e. the dimension of each time series: P, for our problem 1.

The number of sequences: Q, for our problem it is 1.

**Q3:** Continuing the question above, answer the following:

- What is the meaning of setting units = d?
- Assume that we pass a single time series of length \$n\$ as input to the layer. Then what is the dimension of the *output*?
- If we would had set the parameter return\_sequences=False when constructing the layer, then what would be the answer to the previous question?

```
inputs = np.random.random([32, 10, 8]).astype(np.float32)
simple_rnn = tf.keras.layers.SimpleRNN(units=4, return_sequences=False)
output = simple_rnn(inputs)
output.shape
```

Out[10]: TensorShape([32, 4])

#### **A3**:

units = d is the dimentionality of the output space

In this case, X is [n, 1, feature] and if d is the number of units, the output will be of dimension [n, 1, d]

If return\_sequences=False, then the last output in the output sequence is returned, so the dimension will be [n, d]

In *Keras*, each layer is created separately and are then joined by a Sequential object. It is very easy to construct stacked models in this way. The code below corresponds to a simple Jordan-

Elman Network on the form,

 $\left( \frac{t}{t-1} + U \right) + U$  $C \mathbb{L} + c$ , \end{align\*}

Note: It is not necessary to explicitly specify the input shape, since this can be inferred from the input on the first call. However, for the summary function to work we need to tell the model what the dimension of the input is so that it can infer the correct sizes of the involved matrices. Also note that in Keras you can sometimes use None when some dimensions are not known in advance.

```
In [11]:
          d = 10 # hidden state dimension
          model0=keras.Sequential([
              # Simple RNN Layer
              layers.SimpleRNN(units = d, input_shape=(None,1), return_sequences=True, activat
              # A linear output layer
              layers.Dense(units = 1, activation='linear')
          1)
          # We store the initial weights in order to get an exact copy of the model when tryin
          model0.summary()
          init_weights = model0.get_weights().copy()
```

Model: "sequential"

Layer (type)	Output Shape	Param #
simple_rnn_1 (SimpleRNN)	(None, None, 10)	120
dense (Dense)	(None, None, 1)	11
Total naname: 121	=======================================	========

Total params: 131 Trainable params: 131 Non-trainable params: 0

**Q4:** From the model summary we can see the number of paramters associated with each layer. Relate these numbers to the dimensions of the weight matrices and bias vectors \$\{W, U, b, C, c\}\$ in the mathematical model definition above.

A4:

# Question: What is the dimension of the hidden state? see notes

\$W\$ is \$(10,10)\$ as each hidden layer has 10 dimensions(10 units) - 100 parameters

\$U\$ is \$(1,10)\$ as each hidden layer has 10 dimensions(10 units) and observation is of 1 dimension - 10 parameters

\$b\$ is \$(1,10)\$ as each hidden layer has 10 dimensions(10 units) - 10 parameters

Hence, total number of parameters is 120 for Simple RNN layer.

\$C\$ is \$(10,1)\$ as each hidden layer has 10 dimensions(10 units) and observation is of 1 dimension - 10 parameters

\$c\$ is \$1\$ as each observation is of dimension 1 - 1 parameters

Hence, the total number of parameters is 11 for the Dense layer.

# 4. Training the RNN model

In this section we will consider a few different ways of handling the data when training the simple RNN model constructed above. As a first step, however, we construct explicit input and target (output) arrays for the training and test data, which will simplify the calls to the training procedures below.

The task that we consider in this lab is one-step prediction, i.e. at each time step we compute a prediction \$\hat y\_{t|t-1} \approx y\_t\$ which depend on the previous observations \$y\_{1:t-1}\$. However, when working with RNNs, the information contained in previous observations is aggregated in the *state* of the RNN, and we will only use \$y\_{t-1}\$ as the *explicit input* at time step \$t\$.

Furthermore, when addressing a problem of time series prediction it is often a good idea to introduce an explicit skip connection from the input  $y_{t-1}$  to the prediction  $\hat y_{t-1}$ . Equivalently, we can *define the target value* at time step t to be the residual  $\hat y_t := y_t - y_{t-1}$ . Indeed, if the model can predict the value of the residual, then we can simply add back  $y_{t-1}$  to get a prediction of  $y_t$ .

Taking this into consideration, we define explicit input and output arrays as shifted versions of the data series \$y\_{1:n}\$.

```
In [12]:
#Question: Why do we model the residual?
# Training data
x_train = y[:ntrain-1] # Input is denoted by x, training inputs are x[0]=y[0], ...,
yt_train = y[1:ntrain] - x_train # Output is denoted by yt, training outputs are yt

# Test data
x_test = y[ntrain-1:-1] # Test inputs are x_test[0] = y[ntrain-1], ..., x_test[ntes
yt_test = y[ntrain:] - x_test # Test outputs are yt_test[0] = y[ntrain]-y[ntrain-1]

# Reshape the data
x_train = x_train.reshape((1,ntrain-1,1))
yt_train = yt_train.reshape((1,ntrain-1,1))
x_test = x_test.reshape((1,ntest,1))
yt_test = yt_test.reshape((1,ntest,1))
```

# Option 1. Process all data in each gradient computation ("do nothing")

The first option is to process all data at each iteration of the gradient descent method.

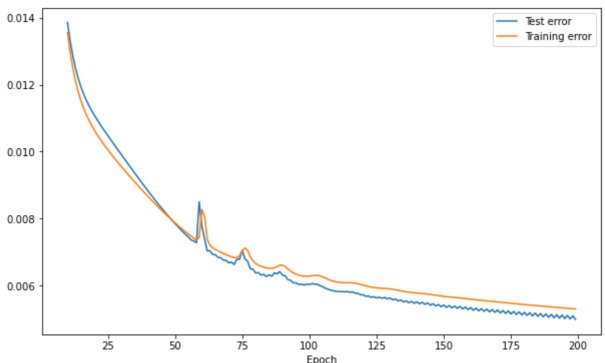
```
In [13]: model1 = keras.models.clone_model(model0) # This creates a new instance of the same model1.set_weights(init_weights) # We set the initial weights to be the same for al
```

**Q5:** What should we set the *batch size* to, in order to compute the gradient based on the complete training data sequnce at each iteration? Complete the code below!

*Note*: You can set verbose=1 if you want to monitor the training progress, but if you do, please **clear the output of the cell** before generating a pdf with your solutions, so that we don't get multiple pages with training errors in the submitted reports.

We plot the training and test error vs the iteration (epoch) number, using a helper function from the tssltools\_lab4 module.

```
from tssltools_lab4 import plot_history
start_at = 10 # Skip the first few epochs for clarity
plot_history(history1, start_at)
```



**Q6:** Finally we compute the predictions of \$\{y\_t\}\$ for both the training and test data uning the model's predict function. Complete the code below to compute the predictions.

*Hint*: You need to reshape the data when passing it to the predict to comply with the input shape used in *Keras* (cf. above).

Hint: Since the model is trained on the residuals \$\tilde y\_t\$, don't forget to add back \$y\_{t-1}\$ when predicting \$y\_t\$. However, make sure that you dont "cheat" by using a non-causal predictor (i.e. using \$y\_t\$ when predicting \$y\_t\$)!

```
In [16]: # Predict on all data using the final model.
    x_data1 = y[:ndata-1]
    x_data1 = x_data1.reshape((1,ndata-1,1))
```

```
# We predict using y_1, \ldots, y_{n-1} as inputs, resulting in predictions of the values # That is, y_pred1 should be an (n-1,) array where element y_pred[t] is based only of y_pred1 = model1.predict(x_data1).flatten() + y[1:ndata]
```

Using the prediction computed above we can plot them and evaluate the performance of the model in terms of MSE and MAE.

```
In [17]:
           def plot_prediction(y_pred):
               # Plot prediction on test data
               plt.plot(dates[ntrain:], y[ntrain:])
               plt.plot(dates[ntrain:], y_pred[ntrain-1:])
               plt.xticks(range(0, ntest, 300), dates[ntrain::300], rotation = 90); # Show onl
               plt.legend(['Data','Prediction'])
               plt.title('Predictions on test data')
In [18]:
           # Plot prediction
           plot_prediction(y_pred1)
           # Evaluate MSE and MAE (both training and test data)
           evalutate_performance(y_pred1, y[1:], ntrain-1, name='Simple RNN, "do nothing"')
          Model Simple RNN, "do nothing"
            Training MSE: 56.5074,
                                       MAE: 6.2866
            Testing MSE: 45.5036,
                                       MAE: 5.6976
                                              Predictions on test data
                                                                                          Data
                                                                                          Prediction
          0.8
          0.6
          0.4
          0.2
          0.0
                 1938-09-30
                                         1963-09-30
                                                                 1988-09-30
```

### Option 2. Random windowing

Instead of using all the training data when computing the gradient for the numerical optimizer, we can speed it up by restricting the gradient computation to a smaller window of consecutive time steps. Here, we sample a random window within the traing data and "pretend" that this window is independent from the observations outside the window. Specifically, when processing the observations within each window the hidden state of the RNN is initialized to zero at the first time point in the window.

In [19]:

To implement this method in Python, we will make use of a *generator function*. A generator is a function that can be paused, return an intermediate value, and then resumed to continue its execution. An intermediate return value is produces using the <code>yield</code> keyword.

Generators are used in *Keras* to implement inifinite loops that feed the training procedure with training data. Specifically, the <code>yield</code> statement of the generator should return a pair <code>x, y</code> with inputs and corresponding targets from the training data. Each epoch of the training procedure will then call the generator for a total of <code>steps\_per\_epoch</code> such <code>yield</code> statements.

```
def generator_train(window_size):
    while True:
        """The upper value is excluded in randint, so the maximum value that we can
        Hence, the maximum end point of a window is ntrain-1, in agreement with the
        when working with one-step-ahead prediction."""
        start_of_window = np.random.randint(0, ntrain - window_size) # First time i
        end_of_window = start_of_window + window_size # Last time index of window (
        yield x_train[:,start_of_window:end_of_window,:], yt_train[:,start_of_window
In [20]:
model2 = keras.models.clone_model(model0) # This creates a new instance of the same
```

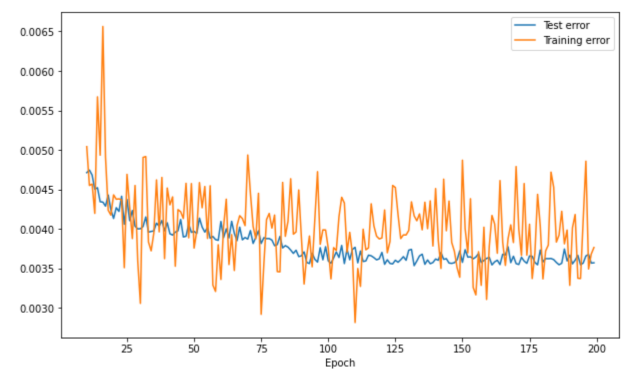
model2.set\_weights(init\_weights) # We set the initial weights to be the same for al

Q7: Assume that we process a window of observations of length window\_size at each

iteration. Then, how many gradient steps per epoch can we afford, for computational cost per epoch to be comparable to the method considered in Option 1? Set the steps\_per\_epoch parameter of the fitting function based on your answer.

Similarly to above we plot the error curves vs the iteration (epoch) number.

```
In [22]: plot_history(history2, start_at)
```

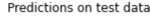


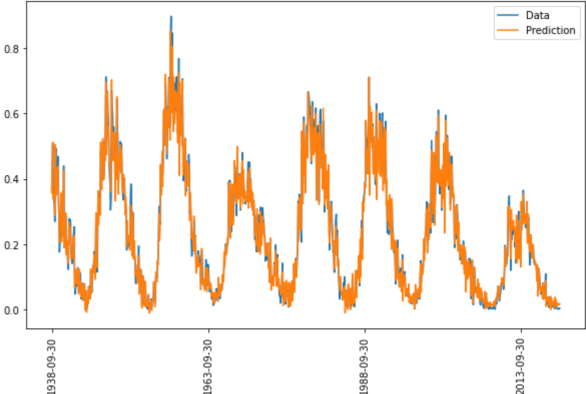
**Q8:** Comparing this error plot to the one you got for training Option 1, can you see any *qualitative* differences? Explain the reason for the difference.

**A8:** For the "Do Nothing" option, we observe a general decrease in error across epochs for both test an training data, while for Random windowing, there is no such trend in training or test data. This is because of the training approach where we compute the gradients for the entire data in Option 1 as opposed to computing the gradients for sequences of data with random starting points in Option 2. Furthermore, when processing the observations within each window the hidden state of the RNN is initialized to zero at the first time point in the window, hence it is fresh restart of the optimization process and there is no memory of previous hidden states. This is the reason why the errors across epochs keep fluctuating in Option 2.

**Q9:** Compute a prediction for all values of  $\{y_2, \dots, y_n\}$  analogously to **Q6**.

```
In [23]:
          # Predict on all data using the final model.
          x_{data1} = y[:ndata-1]
          x_data1 = x_data1.reshape((1,ndata-1,1))
          # We predict using y_1, \ldots, y_{n-1} as inputs, resulting in predictions of the values
          y_pred2 = model2.predict(x_data1).flatten() + y[1:ndata]
In [24]:
          # Plot prediction on test data
          plot_prediction(y_pred2)
          # Evaluate MSE and MAE (both training and test data)
          evalutate_performance(y_pred2, y[1:], ntrain-1, name='Simple RNN, windowing')
         Model Simple RNN, windowing
           Training MSE: 132.9366,
                                      MAE: 8.4102
           Testing MSE: 130.4852,
                                      MAE: 8.3267
```





#### Option 3. Sequential windowing with stateful training

As a final option we consider a model aimed at better respecting the temporal dependencies between consequtive windows. This is based on "statefulness" which simply means that the RNN remembers its hidden state between calls. That is, if model is in stateful mode and is used to process two sequences of inputs after each other, then the final state from the first sequence is used as the initial state for the second sequence.

```
In [25]: # To enable stateful training, we need to create model where we set stateful=True in
model3=keras.Sequential([
    # Simple RNN Layer with stateful=True
    layers.SimpleRNN(units = d, batch_input_shape=(1,None,1), return_sequences=True,
    # A Linear output Layer
    layers.Dense(1, activation='linear')
])
model3.set_weights(init_weights)
```

**Q10:** When working with stateful training we need to make some adjustments to the training data generator.

- 1. First, the RNN model doesn't keep track of the actual time indices of the different windows that it is fed. Hence, if we feed the model randomly selected windows, it will still treat them as if they were consecutive, and retain the state from one window to the next. To avoid this, we therefore need to make sure that the generator outputs windows of training data that are indeed consecutive (and not ranomdly selected as above).
- 2. When training the model we will process the whole training data multiple times (i.e. we train for multiple epochs). However, if we have statefulness between epochs this would effectively result in a "circular dependence", where the final state at time step  $t = n_{\star}$

would be used as the initial state at time \$t=1\$. To avoid this, we can manually reset the state of the model by calling model.reset states().

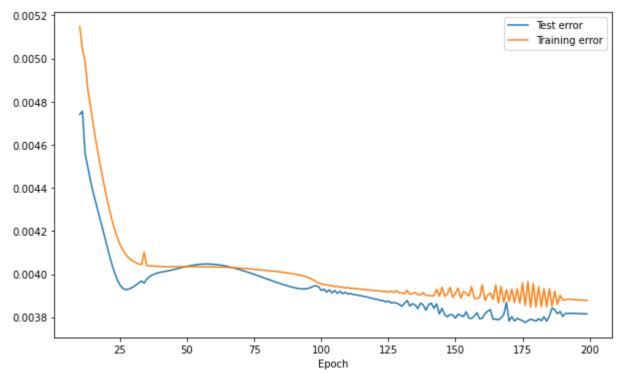
Taking this two points into consideration, complete the code for the stateful data generator below.

```
In [26]:
          def generator_train_stateful(window_size, model):
              """In addition to the window size, the generator also takes the model as input s
              that we can reset the RNN states at appropriate intervals."""
              # Compute the total number of windows of length window_size that we need to cove
              # Note 1. The Length of x_train (and yt_train) is ntrain-1 since we work with 1
              # Note 2. The final window could be smaller than window_size, if (ntrain-1) is
              number_of_windows = int((ntrain-1)//window_size)
              while True:
                  for i in range(number_of_windows):
                      # First time index of window (inclusive)
                      start_of_window = i * window_size
                      # Last time index of window (exclusive, i.e. this is the index to the fi
                      # Note 3. Python allows using end_of_window > ntrain-1, it will simply t
                      end_of_window = (i * window_size) + window_size
                      yield x_train[:,start_of_window:end_of_window,:], yt_train[:,start_of_wi
                  model.reset_states()
```

With the generator defined we can train the model.

Similarly to above we plot the error curves vs the iteration (epoch) number.

```
In [28]: plot_history(history3, start_at)
```



**Q11:** Comparing this error plot to the one you got for training Options 1 and 2, can you see any *qualitative* differences?

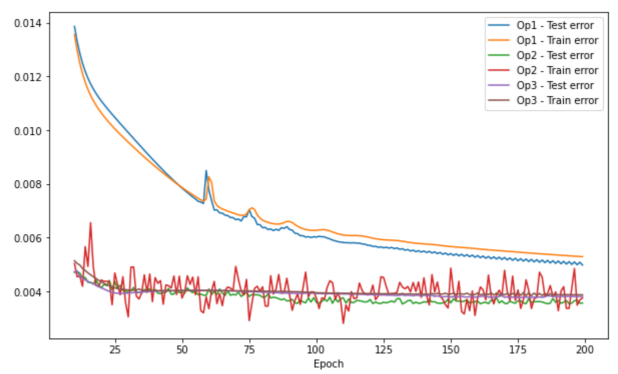
*Optional:* If you have a theory regarding the reason for the observed differences, feel free to explain!

```
plt.plot(history1.epoch[start_at:], history1.history['val_loss'][start_at:], label =
    plt.plot(history1.epoch[start_at:], history1.history['loss'][start_at:], label = 'Op

plt.plot(history2.epoch[start_at:], history2.history['val_loss'][start_at:], label =
    plt.plot(history2.epoch[start_at:], history2.history['loss'][start_at:], label = 'Op

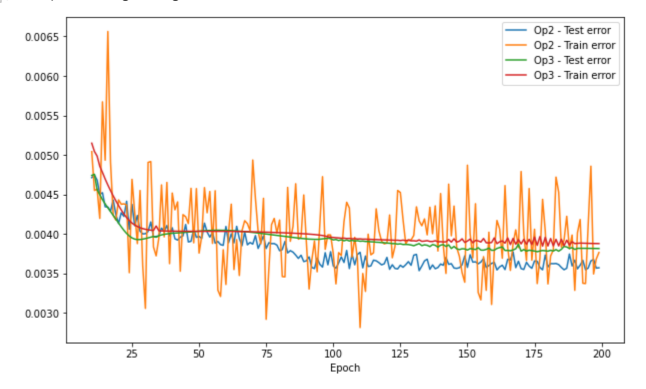
plt.plot(history3.epoch[start_at:], history3.history['val_loss'][start_at:], label =
    plt.plot(history3.epoch[start_at:], history3.history['loss'][start_at:], label = 'Op
    #plt.legend(['Test error', 'Training error'])
    plt.xlabel('Epoch')
    plt.legend()
```

Out[29]: <matplotlib.legend.Legend at 0x21c0ce49df0>



```
plt.plot(history2.epoch[start_at:], history2.history['val_loss'][start_at:], label =
   plt.plot(history2.epoch[start_at:], history2.history['loss'][start_at:], label = 'Op
   plt.plot(history3.epoch[start_at:], history3.history['val_loss'][start_at:], label =
   plt.plot(history3.epoch[start_at:], history3.history['loss'][start_at:], label = 'Op
   #plt.legend(['Test error','Training error'])
   plt.xlabel('Epoch')
   plt.legend()
```

Out[30]: <matplotlib.legend.Legend at 0x21c0d003f70>



**A11:** We achieve better results right at the start of epochs with Option 2 and 3 in comparison to Option 1. This is because in Option 2 and 3, we are dissecting the data into smaller windows, and hence the gradient calculations are more efficient, while in Option 1, the gradient calculations is

computationally heavy and the errors are high at the start of the epochs and it takes quite a lot of epochs for the errors to reduce.

W.r.t the difference in Option 2 and 3, in option 3, we make sure that the generator outputs windows of training data that are consecutive, and furthermore, we also pass the final hidden state to the next window during training. This makes the error less erratic in comparison to the Option 2.

**Q12:** Compute a prediction for all values of  $\{y_2, \dots, y_n\}$  analogously to **Q6**.

```
In [31]:
           # Predict on all data using the final model.
           x_{data1} = y[:ndata-1]
           x_{data1} = x_{data1.reshape((1,ndata-1,1))}
           # We predict using y_1, ..., y_{n-1} as inputs, resulting in predictions of the values
           y_pred3 = model3.predict(x_data1).flatten() + y[1:ndata]
In [32]:
           # Plot prediction on test data
           plot_prediction(y_pred3)
           # Evaluate MSE and MAE (both training and test data)
           evalutate_performance(y_pred3, y[1:], ntrain-1, name='Simple RNN, windowing/stateful
          Model Simple RNN, windowing/stateful
            Training MSE: 163.4925,
                                        MAE: 9.2348
            Testing MSE: 172.0340,
                                        MAE: 9.6996
                                             Predictions on test data
                                                                                          Data
                                                                                          Prediction
          0.8
          0.6
          0.4
          0.2
          0.0
                                                                                        2013-09-30
                                                                1988-09-30
```

### 5. Reflection

**Q13:** Which model performed best? Did you manage to improve the prediction compared to the two baseline methods? Did the RNN models live up to your expectations? Why/why not? Please reflect on the lab using a few sentences.

#### A13:

Which model performed best? The RNN model with Option 1, i.e. do nothing approach performed the best which gave the least test MSE in comaprison to other approaches.

Did you manage to improve the prediction compared to the two baseline methods? Yes, which is evident from the test MSEs.

Did the RNN models live up to your expectations? Why/why not? Yes, as it is able to capture both the local trend and the long range temporal dependencies as it is able to use the hidden states(containing the long range temporal dependencies) and the observations to compute the next hidden state & observation.

# 6. A more complex network (OPTIONAL)

If you are interested, feel free to play around with more complex models and see if you can improve the predictive performance! It is very easy to build stacked models in *Keras*, see the example below.

```
# A stacked model with 3 layers of LSTM cells, two Dense layers with Relu activation
model4 = tf.keras.models.Sequential([
    tf.keras.layers.LSTM(64, batch_input_shape=(1,None,1), return_sequences=True, stat
    tf.keras.layers.LSTM(64, batch_input_shape=(1,None,1), return_sequences=True, stat
    tf.keras.layers.Dense(32, activation="relu"),
    tf.keras.layers.Dense(16, activation="relu"),
    tf.keras.layers.Dense(1),
])
model4.summary()
```

Model: "sequential\_2"

Layer (type)	Output Shape	Param #
lstm (LSTM)	(1, None, 64)	16896
lstm_1 (LSTM)	(1, None, 64)	33024
lstm_2 (LSTM)	(1, None, 64)	33024
dense_2 (Dense)	(1, None, 32)	2080
dense_3 (Dense)	(1, None, 16)	528
dense_4 (Dense)	(1, None, 1)	17

Total params: 85,569
Trainable params: 85,569
Non-trainable params: 0

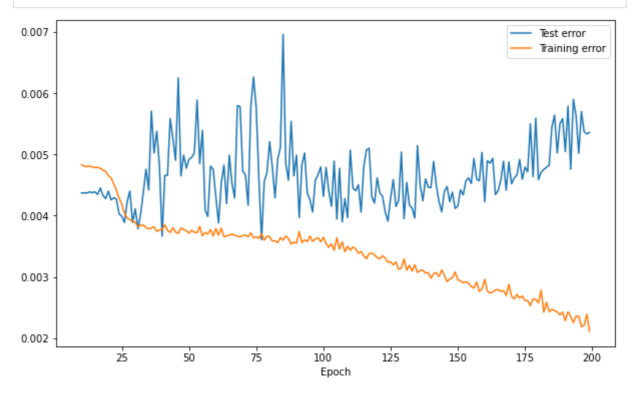
We can store the best model in a file, so that we can load it after analyisng the training procedure.

```
checkpoint_filepath = './'
model_checkpoint_callback = tf.keras.callbacks.ModelCheckpoint(
    filepath=checkpoint_filepath,
```

```
save_weights_only=True,
monitor='val_loss',
save_best_only=True) # Save only the best model, determined by the validation l
```

Train the model

In [36]: plot\_history(history, start\_at)



**Q14 (optional):** Based on the training and test error plots, are there signs of over- or underfitting?

**A14:** As we have less data, we don't need such a complex model, and as a result the model is overfitting.

We load the best model from checkpoint.

```
In [37]: model4.load_weights(checkpoint_filepath)
```

Out[37]: <tensorflow.python.training.tracking.util.CheckpointLoadStatus at 0x21c148f3c10>

```
In [38]: # Predict on all data using the final model.
    x_data1 = y[:ndata-1]
    x_data1 = y[:ndata-1].reshape((1,ndata-1,1))

# We predict using y_1,...,y_{n-1} as inputs, resulting in predictions of the values
    y_pred4 = model4.predict(x_data1).flatten() + y[1:ndata]
```

In [39]:
# Predict on all data using the final model.
# We predict using y\_1,...,y\_{n-1} as inputs, resulting in predictions of the values
y\_pred4 = model4.predict(y[:-1].reshape(1, ndata-1, 1)).flatten() + y[:-1]

In [40]: # Plot prediction on test data
plot\_prediction(y\_pred4)

# Evaluate MSE and MAE (both training and test data)
evalutate\_performance(y\_pred4, y[1:], ntrain-1, name='Stacked RNN, windowing/statefu
#Question: Why is the MSE so high?

Model Stacked RNN, windowing/stateful Training MSE: 572.8246, MAE: 17.1681 Testing MSE: 554.7079, MAE: 17.0709

Predictions on test data

