

FUNDAMENTALS OF COMPUTATIONAL MATHEMATICS
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Project: Yield Curve Model Calibration

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1 The Nelson-Siegel model

The Nelson Siegel model is a model used for the prediction of yield curves, developed by Charles Nelson and Andrew Siegel in 1987 at the University of Washington.

This model aims to predict the movements of the yield curve basing itself on various terms to maturity, such as: flat, hump and S shapes.

This model is of widely interest in economics since the yield curve is a reliable indicator of market sentiment, since it can show expected inflation, movement of interest rates and the overall direction of the economy.

The formulated model is the following:

$$R(\tau) = \beta_0 + \beta_1 * \left[\frac{1 - \exp(-\frac{\tau}{\lambda})}{\frac{\tau}{\lambda}} \right] + \beta_2 * \left[\frac{1 - \exp(-\frac{\tau}{\lambda})}{\frac{\tau}{\lambda}} - \exp(-\tau/\lambda) \right]$$

With $R(\tau)$ equal to yield at time τ , while β_0 , β_1 and β_2 are parameters associated to the long-run yield level, slope of the yield and curvature of the yield respectively.

The last parameter, λ , is the one influencing the rate at which yields converge to the long-term level.

This model is widely used by economists because it has several characteristics that tend to make it a versatile and efficient model. First of all, it respects the restrictions imposed by the economic and financial theory, moreover its approximation avoids in-sample overfitting, such as that it leads to an increase in forecasting capacity.

In addition, versatility refers to the fact that it can take any yield curve form that has been empirically observed in the market.

For these reasons the model is extensively used by central banks and monetary policy makers, while fixed-income portfolio managers use it in order to immunize their portfolios.

Academic literature abounds with instances where researchers have derived significant conclusions by employing the Nelson-Siegel model: Diebold and Li (2006) benchmarked the Nelson-Siegel against other models in the term structure forecasts, the finding was that, especially for longer forecast horizons, it performs very well. Coroneo, Nyholm and Vidava-Koleva (2008) test to which degree the Nelson-Siegel model approximate an arbitrage-free model.

Although there exist also other models used to estimate the term structure of interest rates, most of them have been proven to have undesirable economic properties, moreover they have been seen to be black-box models, so they are not easy to be interpreted and understood by humans.

1.1 Estimation difficulties: The problems of optimization and multicollinearity

The estimation problems addressed to this model refer to his lack of linearity and to the presence of multicollinearity between variables.

Since the parameter that causes non – linearity in the model is λ , or the so-called ‘shape parameter’, many estimation approaches are based on the condition that λ is fixed, in order to linearize the model and perform the estimation by using Ordinary Least Squares.

We are going to investigate this approach more in the detail later, for now we just state that the main problem of this model is that it can become heavily collinear depending on the estimate or fixed shape parameter, λ .

The issue of multicollinearity is that, when dealing with it, the estimation of parameters performed by using least squares can lead to unstable and highly variable coefficient estimates.

Since the Nelson-Siegel model is highly non-linear, we need to estimate its parameters by using non-linear optimization techniques, like for example the Gradient Descent, always considering that this model tends to be very sensitive to the starting values used in the optimization. This issue refers to what is called the “Optimization problem”.

In order to investigate further these problems, we consider a more general representation of the model:

$$r(\tau) = \begin{bmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \end{bmatrix}' \begin{bmatrix} 1 \\ \lambda(1 - e^{-\tau/\lambda})/\tau \\ \lambda(1 - e^{-\tau/\lambda})/\tau - e^{-\tau/\lambda} \end{bmatrix} = \begin{bmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \end{bmatrix}' \begin{bmatrix} r_0 \\ r_1 \\ r_2 \end{bmatrix}.$$

Here we divided the model into three main components, r_0 , r_1 and r_2 that represent respectively the level, the slope, and the curvature of the spot curve.

Deeping further in the behavior of the model, we can see that when the time to maturity grows to infinity, the slope and the curvature components goes to zero and the long-term spot rate converges to a constant level of interest, that is β_0 .

β_1 is the spread, it measures the slope of the term structure, a negative value represents an upward.

The degree of curvature is determined by β_2 , which is the rate at which the slope and curvature components decay to zero.

λ here determines the location of the maximum/minimum value of the curvature component, it determines both the shape of the curvature component and the hump/through of the term structure.

When dealing with Ordinary Least Squares estimation, the multicollinearity problem refers to the high correlation between the slope (r_1) and the curvature (r_2) components, this correlation heavily depends on the choice of λ , that is usually set to be 1.37 or 3, other than that, it depends also on the choice of the time to maturity vector.

To summarize, Gilli et al. (2010) identifies three major problems with estimating this model by non-linear methods or OLS:

- The optimization problem is non-convex, showing multiple local optima.
- The model is badly conditioned for a certain range of parameters, the estimates are unstable before small price movements.
- The value of the decay factor (λ) directly affects the correlation between the loadings of the slope and curvature factors, the problem here could be in the forecasting use of the model.

1.2 Extended Nelson-Siegel function or Svensson model

In order to extend the Nelson – Siegel model and make it possible to capture economics change in case of market crisis, Svensson added an extra term to the original model, allowing for a second hump in the yield curve.

Thanks to this second hump, the extended model is able to capture an even broader and more complicated range of term structure shapes.

This fifth parameter is able to capture a greater variety of yield curve shapes, that can fit in a good way also yields in time of monetary crisis. Indeed, the second hump added by Svensson brought flexibility to the model and thereby improved the goodness of the fit.

The model is computed as follows:

$$R(\tau) = \beta_0 + \beta_1 \left[\frac{1 - \exp\left(-\frac{\tau}{\lambda_1}\right)}{\frac{\tau}{\lambda_1}} \right] + \\ \beta_2 \left[\frac{1 - \exp\left(-\frac{\tau}{\lambda_1}\right)}{\frac{\tau}{\lambda_1}} - \exp\left(-\frac{\tau}{\lambda_1}\right) \right] + \beta_3 \left[\frac{1 - \exp\left(-\frac{\tau}{\lambda_2}\right)}{\frac{\tau}{\lambda_2}} - \exp\left(-\frac{\tau}{\lambda_2}\right) \right]$$

The parameters present in the original model have the same interpretation that they had in the first one, while λ_2 and β_3 determine position and magnitude of a possible second hump in the yield curve.

2 The project

2.1 Data collection

We started by collecting the historical data of various European bonds, we selected four times to maturity: one, two, three, five and ten years, in order to cover the whole maturity range and distribute the data along the whole period.

We chose not to select bonds with very short maturities, because it is possible that they don't show sufficient liquidity.

The data selected are both from European and non-European countries, to be more precise we chose bonds from the following four countries: United States, Portugal, Germany and South Korea.

The selection of the countries was based on the need to incorporate bonds from diverse market economies. This approach allows for an examination of the Nelson-Siegel model's efficacy across different economic conditions and market dynamics.

United states bonds are believed to be one of the safest investments, moreover they are highly liquid and considered close to risk-free, since they are backed by the U.S. government's; given their stability and reliability they tend to be very popular in the financial market. For these reasons, the most commonly reported yield curve is that of US treasury bonds.

German bonds have kind of the same characteristics of the US ones in terms of stability and reliability in the Eurozone, they have a high credit rating, AAA, that states for a very low risk of default.

Given their reliability they are often used as a benchmark in the bond market, in the sense that their yields are often compared to those of other European bonds, moreover they serve as a reference point for assessing credit risk in the Eurozone, this characteristic make these bonds highly liquid.

Portugal bonds, on the other hands are less liquid than the German one, even if they are relatively liquid, and they are considered to be less reliable due to the economic conditions of the country.

For what concern the choice of South Korean bonds, we selected them because they tend to show very interesting yield curves which present an unusual behavior respect to the ones typical of developed economies, this may be due to the fact that the Korean currency is weak with respect to the US dollar.

The time frame we are referring to goes from the first of January 2022 to the first of December, 2023.

2.2 Creation of the dataframe

We started with the creation of a global Data frame in order to be able to work directly on it, at first, we imported on python the data downloaded from the website ‘investing.com’. Since the data we found where divided by each maturity, at the beginning, we had five data frames for each country.

The data we obtained included additional information beyond what is required for implementing the Nelson-Siegel model. This supplementary information encompasses explicit market values of the bonds, such as opening, high, and low prices, as well as percentage changes. In order to get rid of them we applied the function ‘*clear_df*’, which purpose is to drop useless columns and set the date of the month as index of the cleared dataframes. Please find this function in the script ‘*fun.py*’.

After filtering out unnecessary columns, we are left with a separate data frame for each maturity level and country. Each data frame contains a column representing bond prices indexed by month.

In order to perform the Nelson-Siegel model we need a data frame for each date, which reports the yield of the bonds with respect to the maturity. In order to obtain this, we developed the following function, ‘*join_df_date*’:

```
def join_df_date(df1, df2, df3, df4, df5, maturity1, maturity2, maturity3, maturity4, maturity5):
    all_df = []
    data = {'Maturity': [maturity1, maturity2, maturity3, maturity4, maturity5]}
    for i in range(len(df1)):
        price = {'Yield': [df1['Price'].iloc[i], df2['Price'].iloc[i], df3['Price'].iloc[i], df4['Price'].iloc[i], df5['Price'].iloc[i]]}
        df_new = pd.DataFrame({**data, **price})
        all_df.append(df_new)
    return all_df
```

Using this command we successfully merged our data frames with a column for price and the other one for maturity for each month, resulting in a unified list of data frames for each month. We created this type of data frame for each country with the same function.

For example, the data frame obtained for the German bonds referring to the date 1st of August 2023 is the following:

| Maturity | | Yield |
|----------|----|-------|
| 0 | 1 | 3.68% |
| 1 | 2 | 3.21% |
| 2 | 3 | 3.05% |
| 3 | 5 | 2.77% |
| 4 | 10 | 2.84% |

This is just a line of the data frame created for German bonds, which contains all the maturities together.

In the last step of data preparation, we created a dictionary in which the keys are countries' names and the values are the corresponding dataframes.

In order to do that we used the following command:

```
all_df_joint = {'Germany': all_df_joint_germany,
                 'Portugal': all_df_joint_portugal,
                 'South Korea': all_df_joint_sk,
                 'United States': all_df_joint_us}
```

Next, we aim to enhance these data frames by incorporating additional columns containing predicted yields derived from the Nelson-Siegel and the Nelson-Siegel Svensson models. To achieve this, utilized the function previously described theoretically, implemented in Python., whit that function we managed to compute the predicted R(t).

```
def compute_R(tau, params_NS=None, params_NSS=None, tau2=None):
    """
    Computes the R(t) value using the Nelson-Siegel model.

    Parameters:
    - time: column of dataframe
        The time parameter for which R(t) is to be computed.
    - params: array
        Parameters of the Nelson-Siegel model

    Returns:
    - float:
        The computed R(t) value.
    """
    if params_NS is not None:
        beta0, beta1, beta2, time = params_NS
        f1 = (1 - np.exp(-time / tau)) / (time / tau)
        f2 = (1 - np.exp(-time / tau)) / (time / tau) - np.exp(-time / tau)
        return beta0 + beta1 * f1 + beta2 * f2
    else:
        beta0, beta1, beta2, beta3, time = params_NSS
        f3 = (1 - np.exp(-time / tau2)) / (time / tau2) - np.exp(-time / tau2)
        return beta0 + beta1 * f1 + beta2 * f2 + beta3 * f3
```

The function `compute_R` is then looped across the data frame of each country contained in the global one, `'all_df_joint'`, to obtain the predictions of the yield curve for each date for every country according to the maturities at disposal.

In the function mentioned above we also added the possibility to predict the curve by means of the Nelson-Siegel-Svensson model as you can see in the ‘else’ statement, the advantages of this extended model are going to be exploited further in this paper.

The predictions that we are going to get with this curve won’t be any good but will change once we have optimized the function.

That is because this model relies on some parameters, that are:

- The time, for which the expected return has to be computed.
- The parameters insight the model, which are β_0 , β_1 , β_2 , λ_1 (β_3 and λ_2 for the Nelson-Siegel-Svensson).

The setting of the starting parameters is quietly important, especially for the optimization methods that we are going to use later to optimize our model. We underline how in particular the Gradient Descent method tends to be very sensitive to the initial parameters; inaccurate initial parameter values can lead the optimization function to diverge towards infinity.

We based our selection of appropriate values on insights from relevant academic papers, moreover our choice has been made after some trials of the behavior of the optimization methods with different starting values. We noticed that for example, by setting all the parameters to 0.01 the Gradient Descent method was diverging.

To kick off our exploration of optimal parameters, we adopted a randomized approach, selecting values from a range recommended by various academics, notably Gilli et al. (2010). This initial phase involved systematically testing different parameter combinations to identify the most promising configurations.

The ranges suggested by them are:

- $0 \leq \beta_0 \leq 15$
- $-15 \leq \beta_1 \leq 30$
- $-30 \leq \beta_2 \leq 30$
- $-30 \leq \beta_3 \leq 30$

We performed several trials with different combinations of values. At the end we found different starting values for each country, given the different economic conditions they lie in.

Focusing example of United States, the evidence that emerged from our empirical work, that we were able to find mostly by looking at the plots of the Nelson Siegel compared with the ones of the Historical data, was that values higher than 1 for the betas did not perform well. Consequently, we proceeded with subsequent trials using lower beta values based on this observation.

Moreover, the plots suggested to us that the value of β_3 should be negative.

After a considerable number of trials, we decided to select the following values, according to the higher performance they were showing for both the Gradient Descent and the Newton methods:

- $\beta_0 = 0.03$
- $\beta_1 = 0.015$
- $\beta_2 = 0.010$
- $\beta_3 = -0.0020$
- $\lambda_1 = 2.32$
- $\lambda_2 = 12.35$

Added the Nelson Siegel model values to our data frames we had all the data needed in order to compute the sum of squared difference between observed and model predicted yields and optimize it according to the parameters.

A similar approach was adopted to incorporate Nelson-Siegel Svensson values.

2.3 Parameter estimation

The sum of squared difference between observed and model predicted yields is computed as follows:

$$f(\beta_0, \beta_1, \beta_2, \lambda_1) = \sum_{i=1}^T (R(t_i) - R(t_i))^2$$

This is the function to compute the sum of squared residuals for the Nelson-Siegel model, once again in the python code we also presented the version for the Nelson-Siegel-Svensson model.

```
def compute_f(yields, tau, params_NS=None, params_NSS=None, tau2=None):
    """
    Computes the f(beta0, beta1, beta2, tau) or f(beta0, beta1, beta2, beta3, tau0, tau1) value using the Nelson-Siegel model.

    Returns:
    - float:
        The computed f(beta0, beta1, beta2, tau) or f(beta0, beta1, beta2, beta3, tau0, tau1) value.
    """
    if params_NS is not None:
        residuals = yields - compute_R(tau, params_NS=params_NS)
    else:
        residuals = yields - compute_R(tau, params_NSS=params_NSS, tau2=tau2)
    return np.sum(residuals**2)
```

Now we can proceed with the minimization of f , adjusting the parameters via de Gradient Descent and the Newton methods.

We imported these two methods at the beginning from a module we created to store all our functions, “fun.py”, in order for our code to be more compact.

3 Gradient-Descent method

We initiate the process with the Gradient Descent method, which operates on a continuous differentiable function, denoted as f , aiming to minimize it.

The procedure relies on a descent direction, d , and a step size, α .

The descent direction is the gradient of f calculated in the point before the one we are trying to update, while α is found by performing approximate line search, which ensures that we give to our descent direction a stable minimization following the ‘Armijo-Goldstein Condition’.

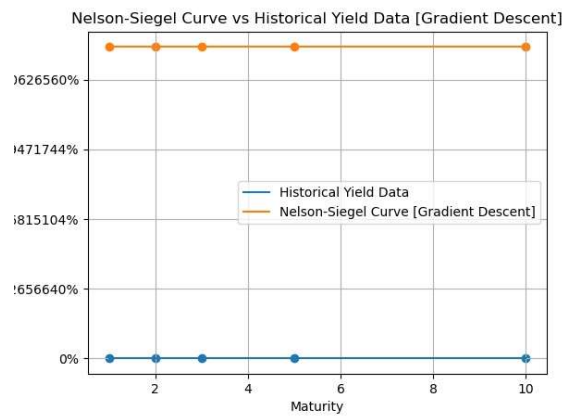
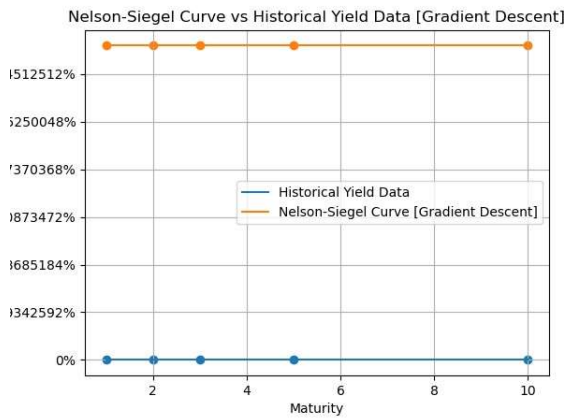
At the beginning, we did not use the approximate line search function, in python code: ‘`apx_LS = False`’. This approach is riskier but faster, this is because gradient descent function will decrease by taking giant steps, with the risk of missing the optimal solution, however the findings could also be really accurate.

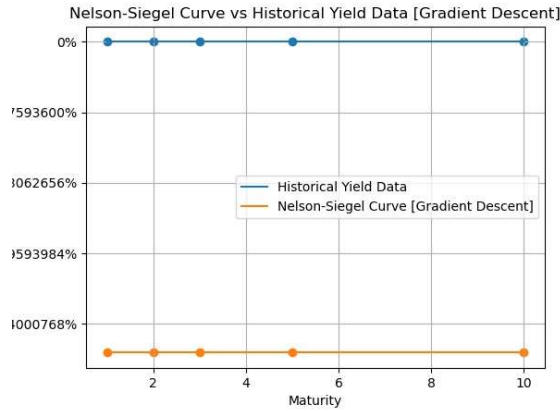
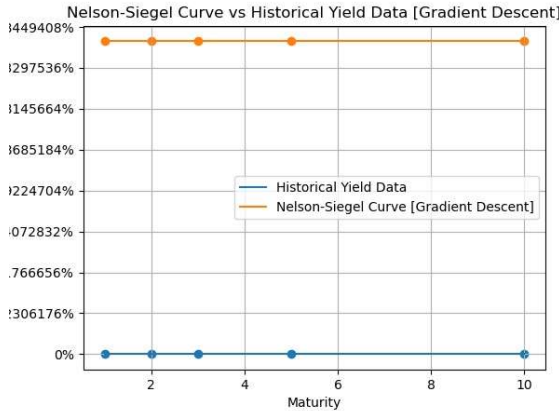
In order to ensure us to risk less with our optimization, we then proceeded by following the Armijo Goldstein condition.

The parameters we chose were the following ones:

- $\beta_0 = 0.1$
- $\beta_1 = 0.1$
- $\beta_2 = 0.1$
- $\beta_3 = 0.1$
- $\lambda_1, \lambda_2 = 1$

By using these conditions, we noticed that the Gradient Descent method was diverging. The following are some of the plots that we obtained, we present the example of the 1st of January 2022, for each country.





The same situation occurs every month.

After several tests we ended up with good starting values, different for each country given the different economic conditions.

For example, for US we went for the parameters presented at the beginning of this paper. We recall that this is just one of the optimal solutions.

In order to find these parameters, we started by looking at β_0 , at the beginning we performed two trials, one by increasing this parameter and the other one by decreasing it. We noticed that while with a smaller β_0 the Nelson Siegel was diverging, with a greater one it was converging to the actual yield curve.

We followed the same methodology also for the other β and λ . The predicted yield curve was more accurate in the case where all the β were smaller than the previous one. At the same time, we had a better convergence with a negative β_3 .

On the other hand, we needed greater λ for the two methods to converge to the actual yield curve.

We set $\alpha = 0.65$. We found that this value was a good meeting point between a good computational time and a good approximation. For example, $\alpha = 10$ was taking much less iterations but was overstepping a lot, while for an alpha smaller than 0.65 the time taken was too much.

As we already said, we set '*apx_LS = True*'.

Given that, the updated x is going to be the previous one plus alpha multiplied by the descent direction.

The procedure is going to stop either when the number of iterations reaches the maximum decided at the beginning or when the value of the last iteration is below a certain threshold decided at the beginning.

We established the maximum number of iterations to be 100.

The functions we are referring to are named in python as '*gradient_descent*' and '*apx_line_search*' and they are stored in the script '*fun.py*'.

Before performing the minimization, we initiated four lists, with the aim of storing the values found by the minimization. The lists are one for the parameters and one for the value of f , both of them were created for the Nelson Siegel and the Nelson Siegel Svensson model.

We ran a for loop to perform the gradient descent methods in each data frame for every country we are referring to. Once the optimization has been done, we collected the values of the parameters stored in the lists in a data frame that we saved in Excel. For the complete code please refer to the script ‘*Data.py*’.

We did the same also for the Nelson Siegel Svensson model, to compare which of the two models is more precise in fitting the historical data.

By running the code, we obtained that for most of the data provided the Gradient Descent method performed one-hundred iterations, which is the maximum number of iterations we decided at the beginning, clearly this states that the method is converging slowly.

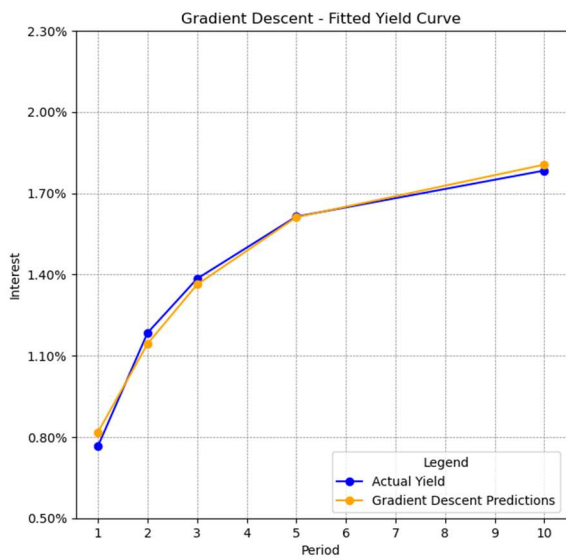
Anyway, even when the methods performed the maximum number of iterations, the graph results to be sufficiently precise.

Also moving to the plots one can clearly notice that, when the gradient descent performed less than one hundred iterations, the Nelson Siegel curve, and the Nelson Siegel Svensson curve as well, are fitting the historical yield data in a very precise way.

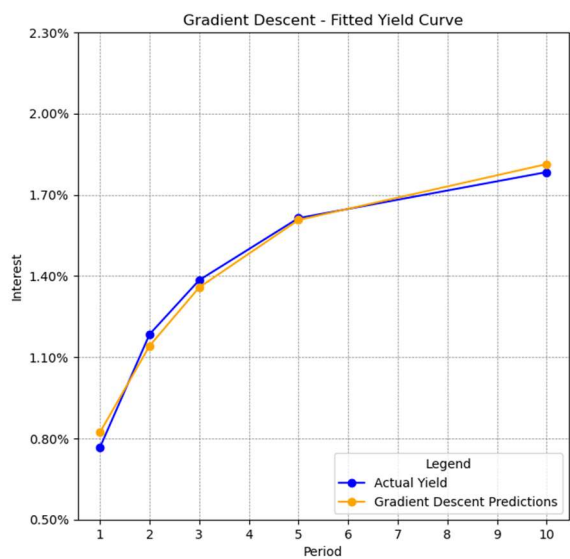
In order to plot the curves, we extracted the last parameters’ values from the data frames of parameters stored by the gradient descent method, this is because the last value should be the optimal one. Once we have obtained them, they can be used to compute the Nelson Siegel and the Nelson Siegel Svensson curve.

3.1 Gradient Descent method- Us

We are going to present some examples of different situations by looking at some of the plots that we obtained. All the plots can be found in the zip file. We start by showing the plots of the Nelson Siegel and Nelson Siegel Svensson curve for US bond on the 1st of January 2022, for which the Gradient Descent performed 100 iterations for both the Nelson Siegel and Nelson Siegel Svensson.

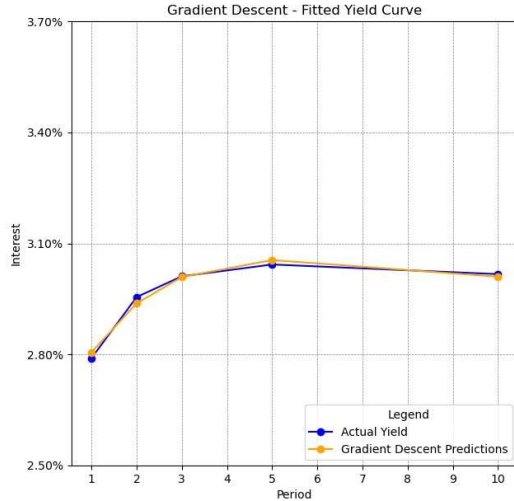


Nelson Siegel

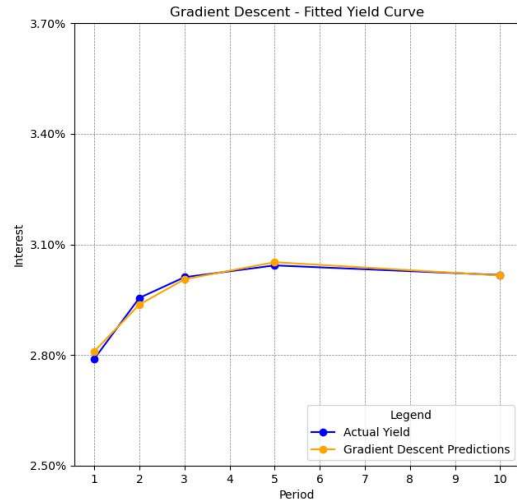


Nelson Siegel Svensson

It is evident that, if we take into consideration data that performed the maximum number of iterations, the plots are still reflecting the historical data. However, if we take into considerations plots of dates for which the gradient descent method performed less than 100 iterations, the curves are slightly more precise. This is the case of US bonds on the 1st of June 2022:



Nelson Siegel

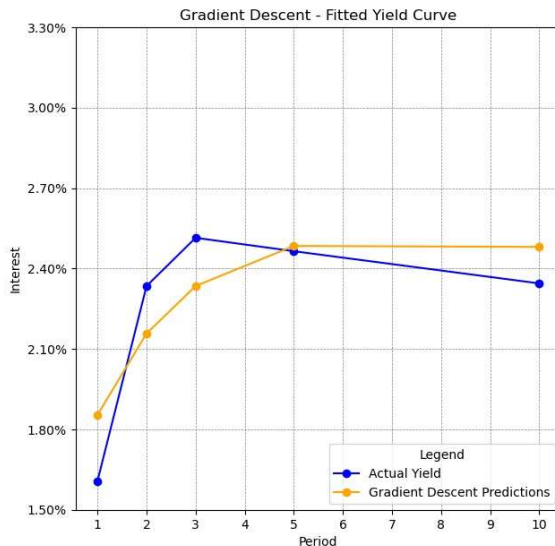


Nelson Siegel Svensson

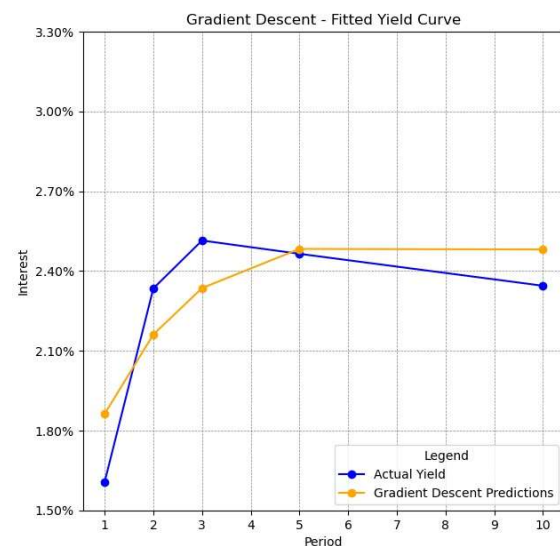
We underline that in this case the Gradient descent method performed 94 iterations for the Nelson Siegel and 100 for the Nelson Siegel Svensson.

Furthermore, we noticed that in the case of bonds with high volatility the Nelson Siegel and Nelson Siegel Svensson model optimized by the gradient descent method are not able to perfectly predict the yield curve, even though the extended model tends to slightly outperform the original one.

Here is the example of US bonds on the 1st of March 2022:



Nelson Siegel



Nelson Siegel Svensson

In the location of the hump neither the original Nelson Siegel nor the extended version are able to fit the historical data. Here the model is converging very slow, if we set the possibility to run more iterations the Gradient Descent method would probably find the optimal parameters and the model would fit the historical data in a better way, but this procedure would require a huge computational time.

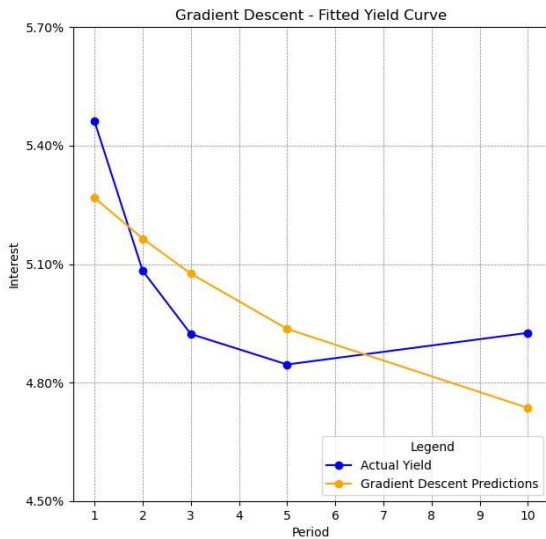
One issue that we noticed about the Newton method was in predicting Reversed Yield curves.

First of all, let's investigate what Reversed Yield curves are. When the yield curve inverts, it means that short-term interest rates are higher than long-term rates, the yield decreases the further away the maturity date is.

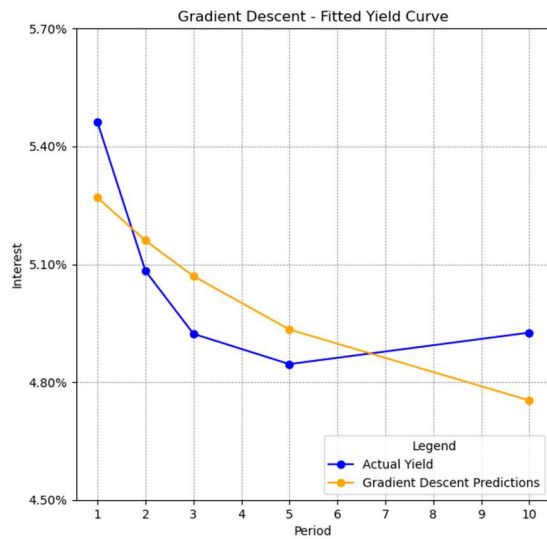
Inverted yields curve used to be an unusual occurrence, but in times of rising inflation this phenomenon tends to happen more frequently. Focusing on the US, in this time of raising interest rates, low maturity now tends to have higher yields.

That is why the inverted curve is a reliable indicator of a recession. It suggests that investors have little confidence in the near-term economic conditions and believe that interest rates will fall in the future.

Let's see the example of October 2023:



Nelson Siegel



Nelson Siegel Svensson

We will see that in the cases mentioned above the Newton method is more precise and we will explore the reasons behind this.

4 Newton method

The second method we implemented is the Newton method.

This procedure for functions optimization is similar to the Gradient Descent one, except that in this case the updating step is computed using second-order Taylor expansion. The ‘d’ we were referring to in the Gradient Descent method in this case is the opposite of the gradient at point x_i , while alpha is the inverse of the Hessian at point x_i .

In order for this method to be effective, it's crucial to verify that the Hessian matrix is Positive Definite. If it's not, the method cannot be applied as the Hessian would be non-invertible. To mitigate this issue, a common practice is to incorporate a regularization term, typically denoted as λI , into the matrix. This regularization term ensures the matrix becomes Positive Definite, allowing for the successful application of the method. The regularization term mentioned above has been called in the code ‘Damping factor’ and has been set to be 0.5.

Once again, to use this method, we implemented the code learned during laboratory hours, that we imported at the beginning of our script.

The function ‘*newton_method*’ is looped across all the data frames we created at the beginning for each country, in explicit we have twenty-three different data frames for every country we selected, one for every month of our time frame.

At each iteration the Nelson Siegel (and Nelson Siegel Svensson) parameters are optimized according to the Newton’s method, the number of iterations performed to achieve the optimal parameters are then returned by the ‘*newton_method*’ function itself.

We stored the values of every iteration in an Excel file, if it is of interest to you, please find it in the zip file.

Once we have done that, in order to plot the curves predicted by the Nelson Siegel and Nelson Siegel Svensson models with respect to the observed yield curve, we need to select the parameters found by the last iteration performed (that should be the optimal ones) and compute the expected returns by means of the original and extended version of the Nelson Siegel model with those parameters.

The following piece of code shows this procedure:

```
# Paste Optimal for Newton and plot it
for index, df in enumerate(df_joint_country):

    # Extract parameter values from df_params
    beta0 = df_params_newton.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][0]
    beta1 = df_params_newton.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][1]
    beta2 = df_params_newton.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][2]
    tau = df_params_newton.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][3]

    beta0_NSS = df_params_newton_NSS.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][0]
    beta1_NSS = df_params_newton_NSS.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][1]
    beta2_NSS = df_params_newton_NSS.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][2]
    beta3_NSS = df_params_newton_NSS.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][3]
    tau_NSS = df_params_newton_NSS.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][4]
    tau2_NSS = df_params_newton_NSS.iloc[:, 1:].ffill(axis=1).iloc[:, -1][index][5]

    # Create params_NS list
    params_NS_optimal = [beta0, beta1, beta2, tau]
    params_NSS_optimal = [beta0_NSS, beta1_NSS, beta2_NSS, beta3_NSS, tau_NSS, tau2_NSS]

    # Compute Nelson-Siegel curve values
    df['Nelson-Siegel'] = fun.compute_R(df['Maturity'], params_NS=params_NS_optimal)
    df['Nelson-Siegel-Svenson'] = fun.compute_R(df['Maturity'], params_NSS=params_NSS_optimal)

    # Plot the curve
    fun.plot_curve(maturities, df['Yield'], df['Nelson-Siegel'], name_country, 'Nelson-Siegel', 'Newton', dates[index])
    fun.plot_curve(maturities, df['Yield'], df['Nelson-Siegel-Svenson'], name_country, 'Nelson-Siegel-Svensson', 'Newton', dates[index])
```

From this code we are going to obtain the plots.

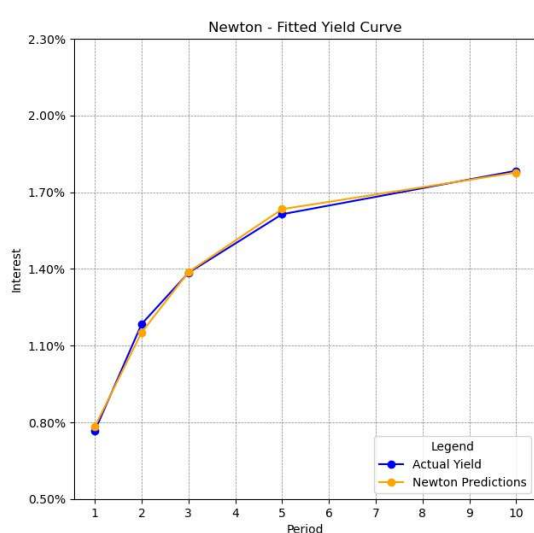
Let us present some difference and evidence between various yield curves

In Newton's methods, even when the algorithm reaches the maximum number of iterations, the resulting predicted plot consistently exhibits a remarkably precise fit to the historical yield data.

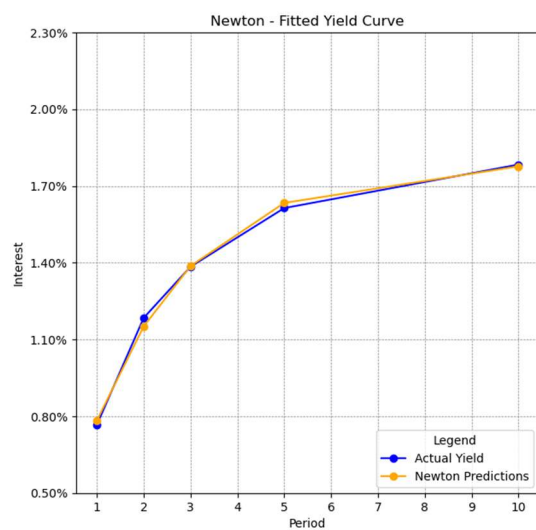
Moreover, the Nelson Siegel Svensson model tends to converge faster with respect to the original one.

4.1 Newton method US:

Here is the example for US bonds on the 1st of January 2022:



Nelson Siegel



Nelson Siegel Svensson

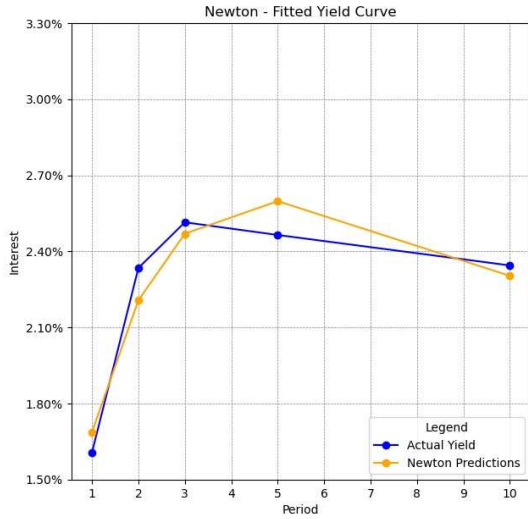
The Nelson Siegel and the Nelson Siegel Svensson curves closely mirror the historical yields data, exhibiting a remarkably strong correspondence.

The plots in this case are very similar to the ones of the Gradient Descent Methods.

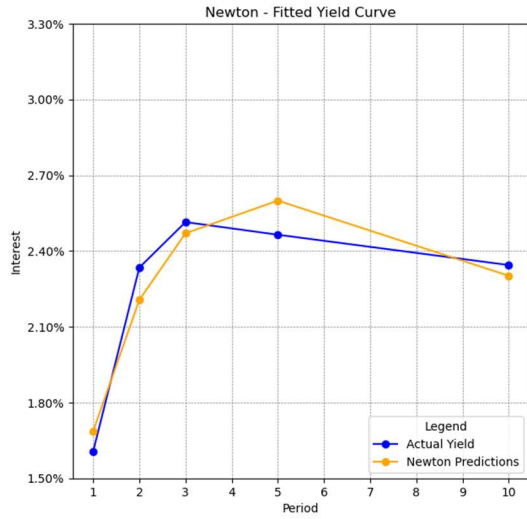
Both curves tend to be very precise, even if the extended model is slightly better.

While studying the Gradient Descent method we noticed that the model wasn't that good in fitting the curve of bonds with high volatility, if we apply the Newton method this issue emerges but is way less significant.

Let's take for example the plots for US bonds on the 1st of March 2022:



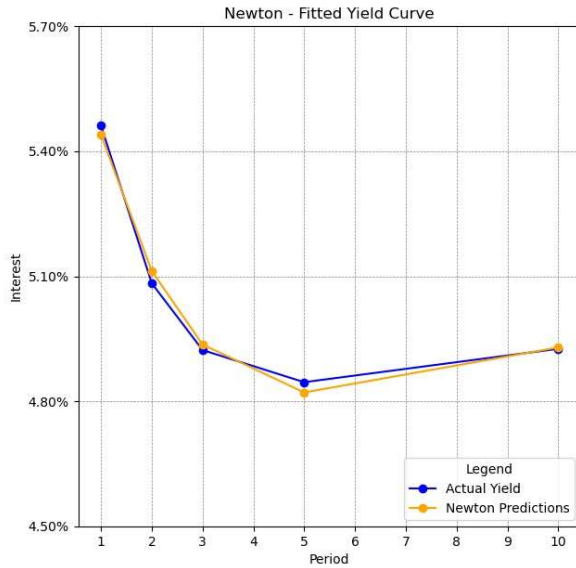
Nelson Siegel



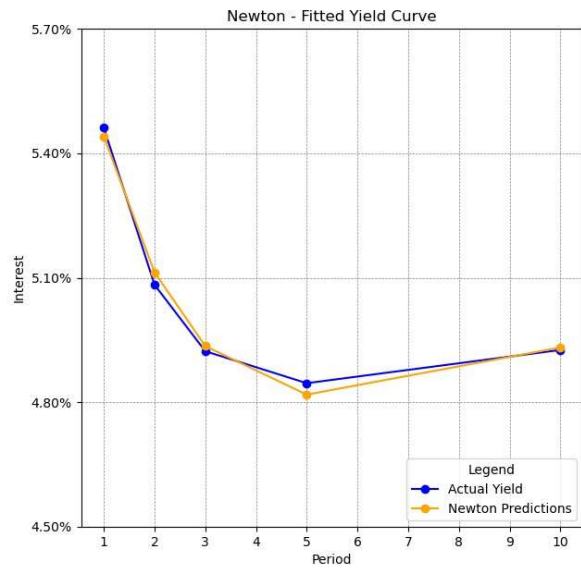
Nelson Siegel Svensson

In the case of the Gradient Descent, the Nelson Siegel model wasn't able to fit the historical yield in the location of the hump. However, with our current method, we observe an improved fit, more able to capture the characteristics of the hump.

The same occurs for inverted yield curves, let's see the case of October 2023:



Nelson Siegel



Nelson Siegel Svensson

This is given by the fact that the Newton method tends to converge faster to the optimal solution given the initial parameters, so the errors between the observed and predicted data are minimized in less iterations.

Furthermore, since the Newton method involves computing the Hessian, it effectively incorporates information about the curvature of the function. As a result, it can more accurately capture the presence of humps in the data, leading to improved fitting performance.

In the following paragraph we will present the plots for the other countries. Evidence that emerged for US bonds is going to be confirmed also for the other countries.

4.2 Portugal:

The yield curve for Portugal bonds is going to be different from the one of the US bonds; this is because the yields are lower with respect to the American ones and the countries are characterized by different economic backgrounds.

The parameters we selected for the Gradient Descent and Newton method for Portugal bonds are the following:

- $\beta_0 = 0.006$
- $\beta_1 = 0.4$
- $\beta_2 = 0.005$
- $\beta_3 = 0.01$
- $\lambda_1 = 1.5$
- $\lambda_2 = 1.6$

To find this optimal combination of parameters, we tried to change one of them at a time and see if they converged to the actual yield curve or not.

Our first guess was $(0.15, 1, 0.01, 0.01, 1.5, 1.6)$. We started by reducing and increasing the value of β_0 and we noticed that an increased value led to a divergent curve while a decreased one made the Nelson Siegel model converge to the actual yield. The same happened while decreasing the other betas.

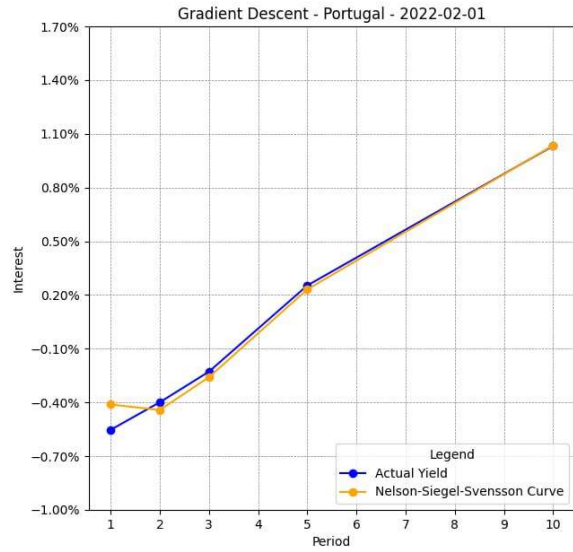
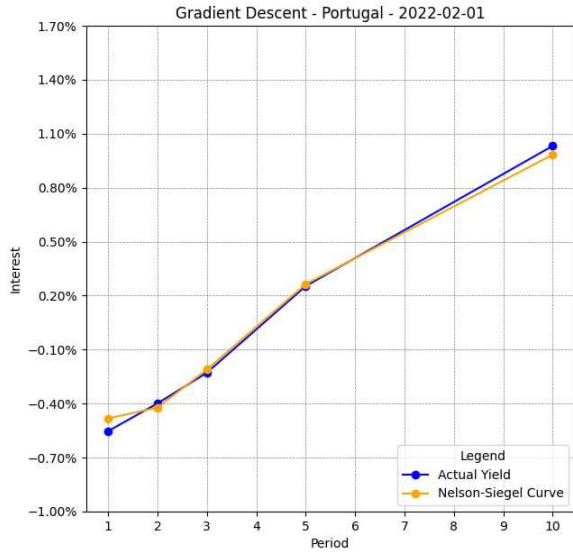
While for the lambda, a higher value led to a better fit of the predicted curve to the actual one, but with that the curve had an opposite trend with respect to the actual yield, so we decided to keep λ_1 equal to 1.5.

For what concern the approximate line search, we kept it ‘True’ and the alpha was set to 0.65.

4.2.1 Gradient descent Portugal:

The gradient descent performed around about 100 iterations for both models, so the method is converging slowly to the optimal values. Despite that the curves of predicted yields fit the actual ones in a pretty accurate way.

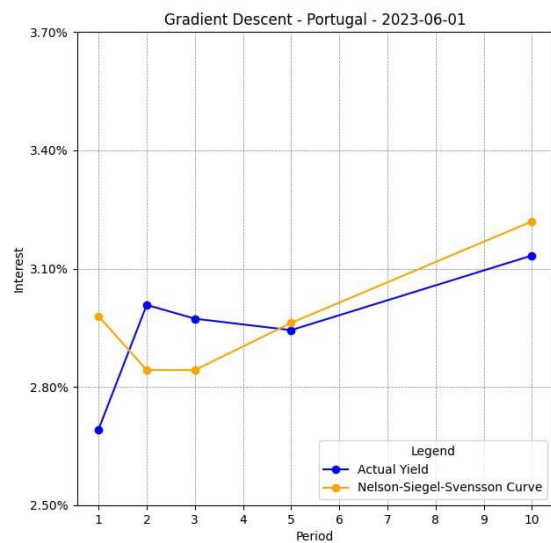
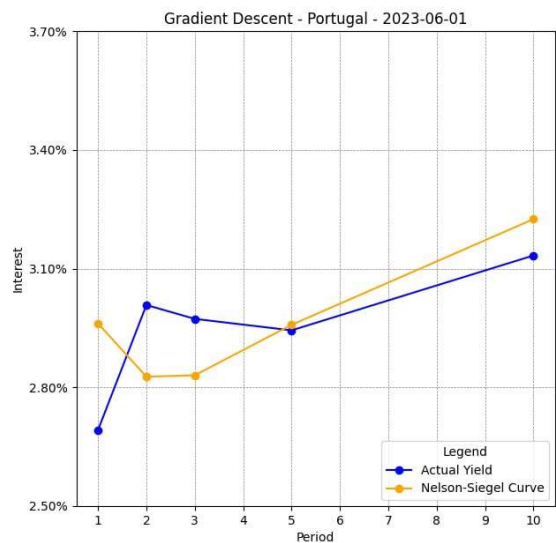
Let's look for example at the plots for the 1st of February 2022:



These graphs are pretty accurate, with the Nelson Siegel model fitting the actual yield curve in a less accurate way with respect to the extended one, especially for maturity over 5 years.

The issues encountered for US bonds in the presence of high volatility emerged also in the case of Portugal bonds.

Let's look at the example of the 1st of June 2023:

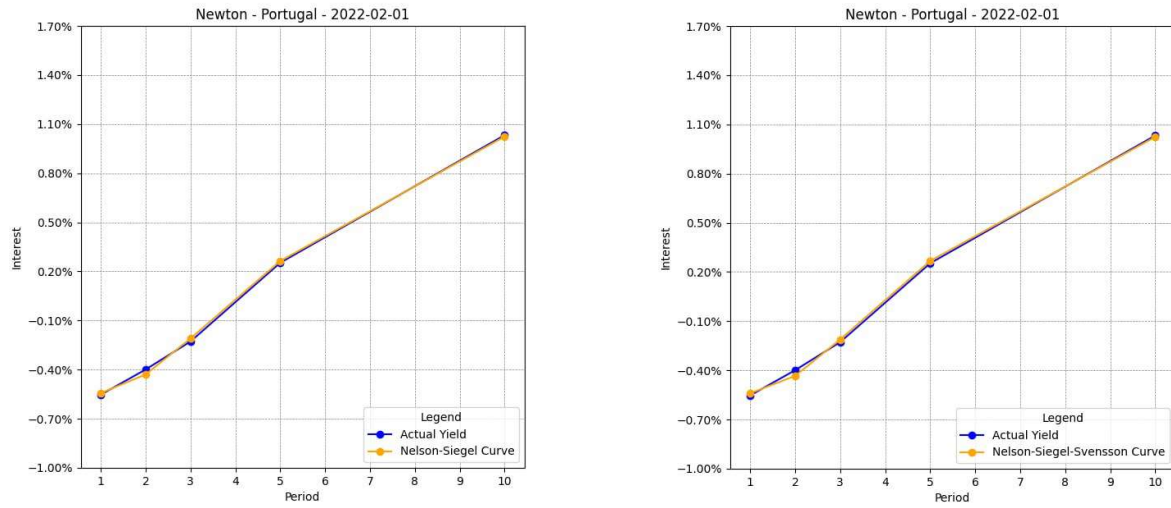


Once again, in the location of the hump, the models present difficulties in fitting the actual yield.

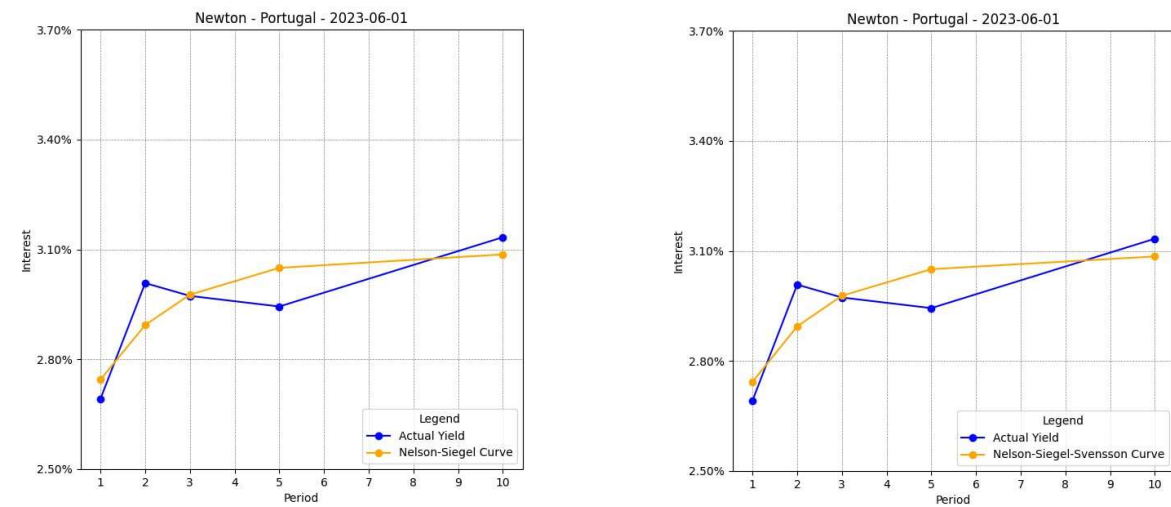
4.2.2 Newton Method Portugal:

By running the Newton method, we confirmed that it tends to be way more accurate with respect to the Gradient Descent one, and less sensitive to the initial parameters, even if it is more computationally complex.

These are the resulting plots for the 1st of February 2022:



And 1st of June 2023:



Even if we encountered some problems in fitting a yield curve also by means of the Newton method, the results are still more accurate with respect to the ones of the Gradient Descent.

4.3 Germany

Germany yield curves are more like the American ones, given the stability and reliability of the economic conditions of the country.

By looking at its plots we are not going to encounter a lot of curves that shows behavior very dissimilar to the classical ones, like on the other hand we encountered in the case of Portugal.

For what concern the parameters, we performed various tries in order to try one of the possible optimal combinations.

Our first guess was (4.99, 10.99, 4.99, 3.99, 5.99, 10.01) with alpha equal to 0.65.

We tried at first to decrease all the values of the betas separately, since we noticed that the predicted curve was converging to the actual yield, we kept decreasing them.

We started to increase the tau, but in this way the Nelson Siegel model was diverging, so we worked the other way around and we decreased them.

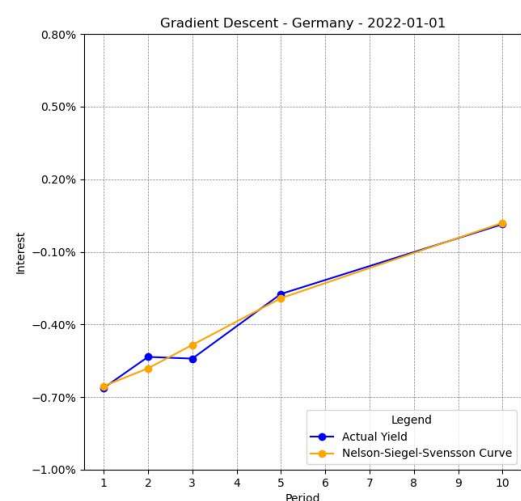
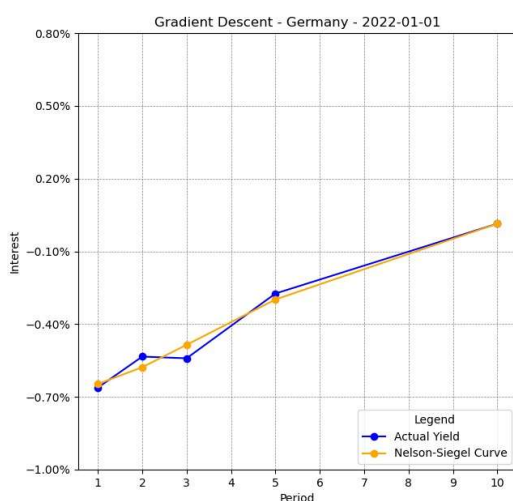
Once it seemed to us that we were moving closer to the optimal value, we worked on the step length, alpha, and we increased it a bit, setting it to 0.8.

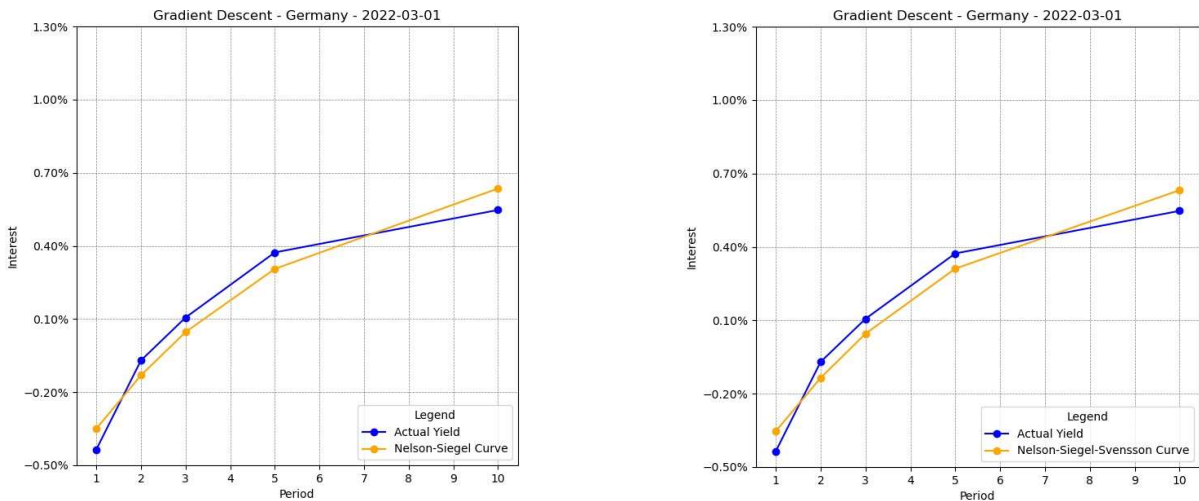
At the end, our chosen values are:

- $\beta_0 = 0.0049$
- $\beta_1 = -0.011$
- $\beta_2 = -0.013$
- $\beta_3 = -0.0039$
- $\lambda_1 = 1.96$
- $\lambda_2 = 5.05$

4.3.1 Gradient Descent Germany:

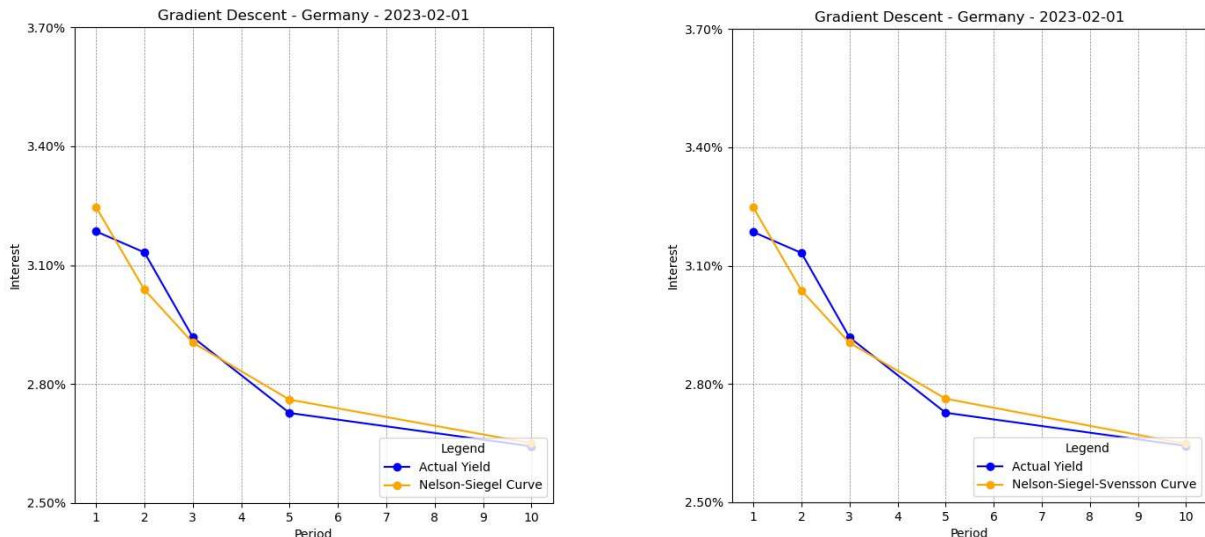
We present two cases, on the 1st of January 2022 the gradient descent performed 22 iterations, while in the 1st of March 2022 they were 100:





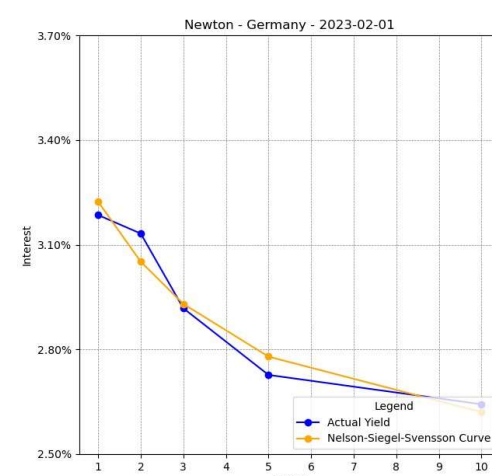
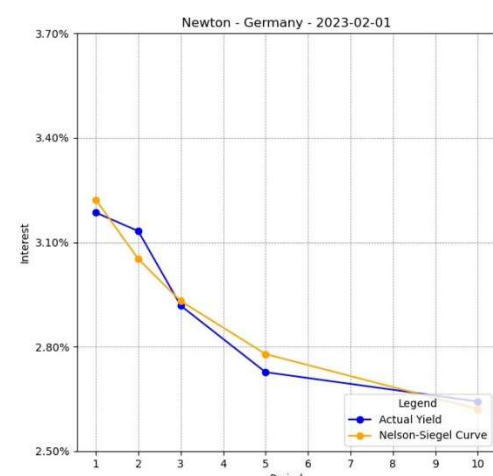
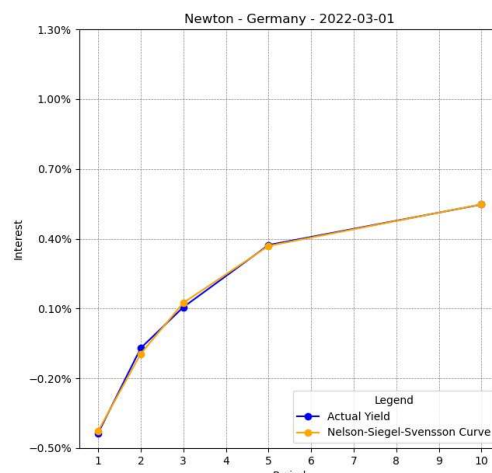
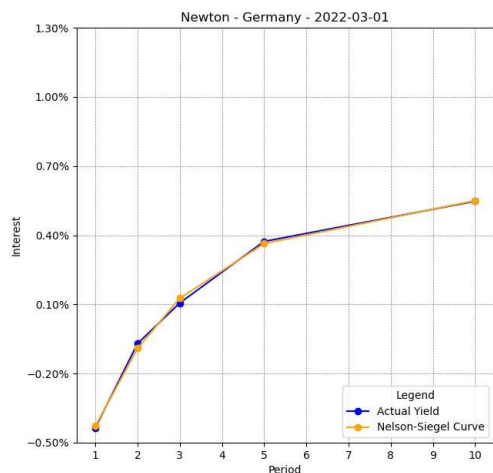
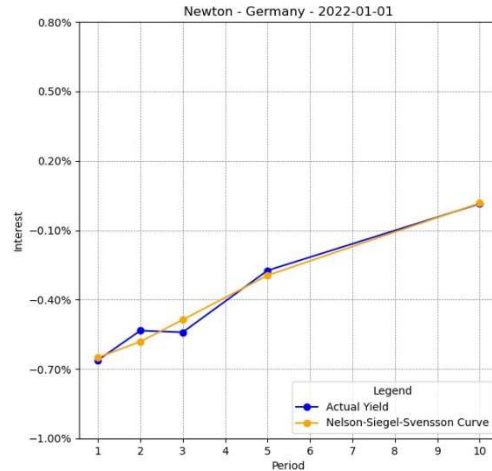
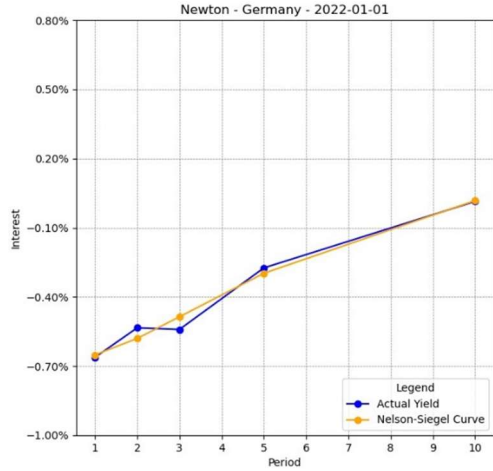
Even if in the first case the actual yield is fitted in a better way, also in the case of march the fit is pretty accurate.

Also, in the case of Germany, when the actual yield shows sudden changes, the predicted curves have some difficulties in fitting it. This is the case of the 1st of February 2023:



4.3.2 Newton Method Germany:

Once again, the Newton method shows to be more accurate:



4.4 South Korea

South Korea yield curves are less stable than those of other countries, especially in 2023 where almost all the curves are flat.

For what concern the parameters, we performed various tries in order to try one of the possible optimal combinations.

We first tried two set of parameters. The first is (0.04, -0.008, 0.007, -0.002, 2.32, 12.35) while the second was (0.02, 0.018, 0.017, -0.002, 2.32, 12.35) both with alpha equal to 0.65.

Both the set of parameters granted us good approximations of the yield curves. We then tried to use a lower value for both lambdas and we achieved even better results

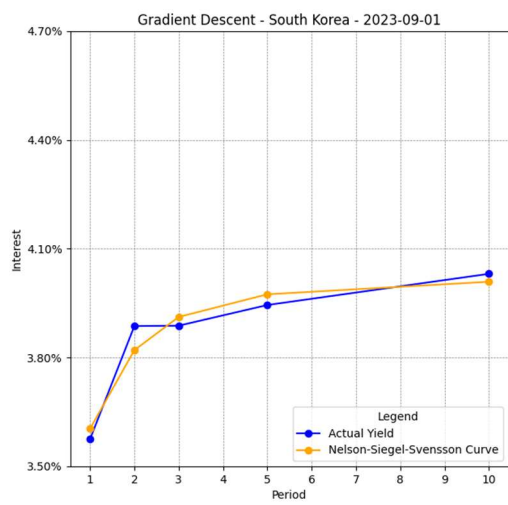
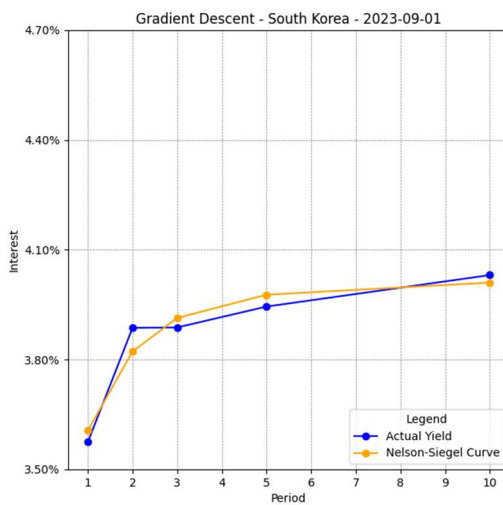
We decided to stop at that point because the fit in the graphs was enough good for our purpose. The final parameters are that from our first model but with lower values for lambdas.

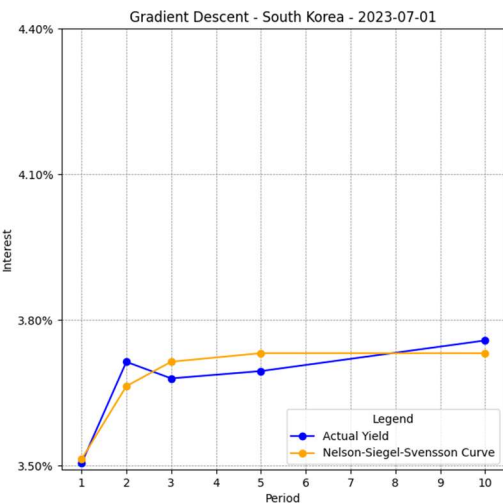
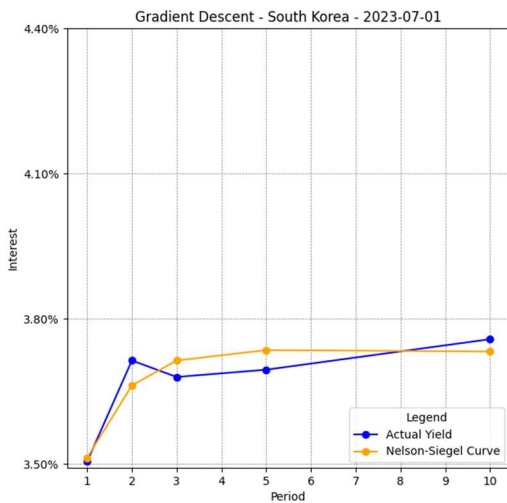
At the end, our chosen values are:

- $\beta_0 = 0.04$
- $\beta_1 = -0.008$
- $\beta_2 = 0.007$
- $\beta_3 = -0.002$
- $\lambda_1 = 1$
- $\lambda_2 = 5$

4.4.1 Gradient Descent South Korea:

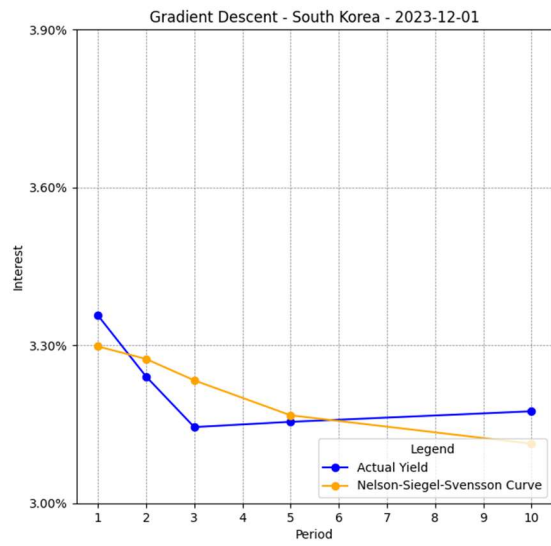
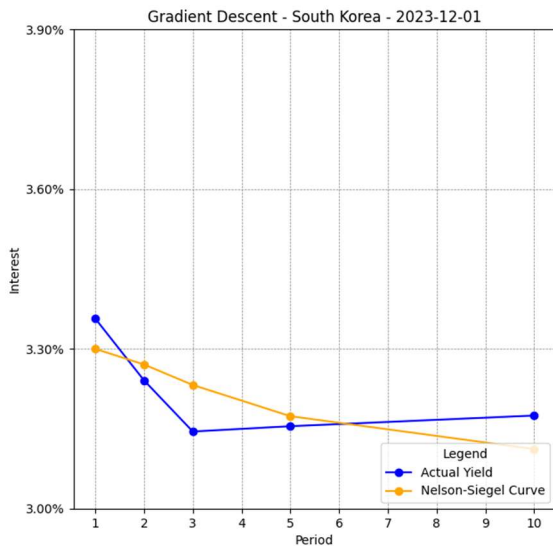
We present two cases, on the 1st of September 2023 the gradient descent performed 44 iterations, while in the 1st of July 2023 they were 60:





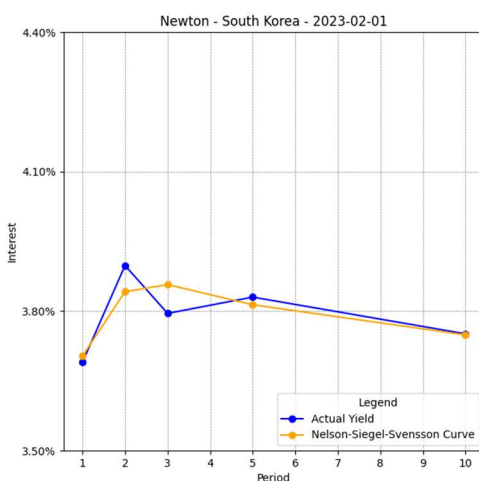
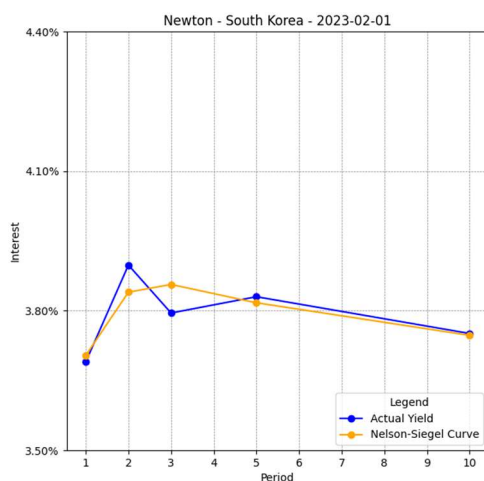
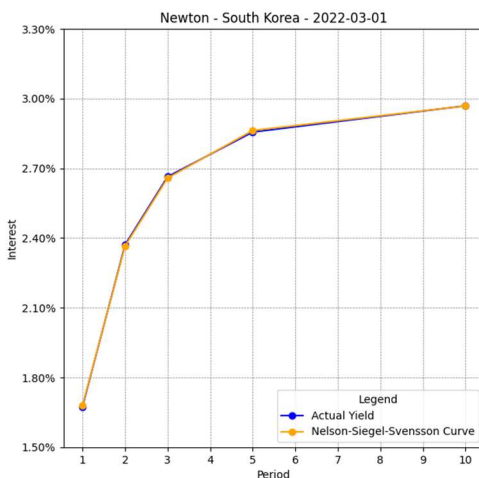
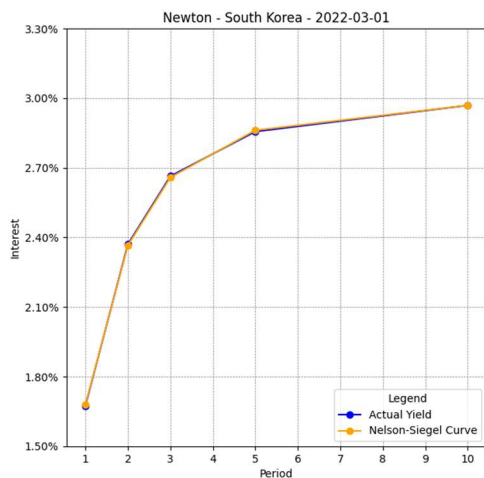
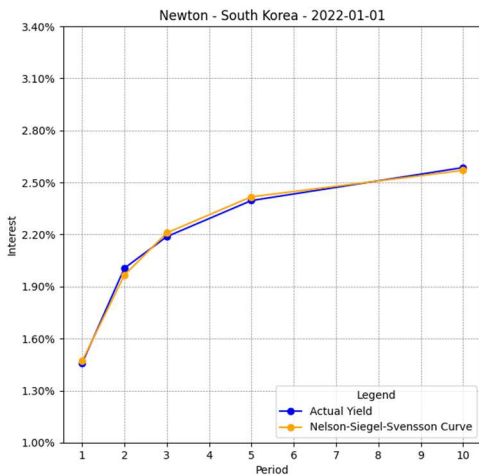
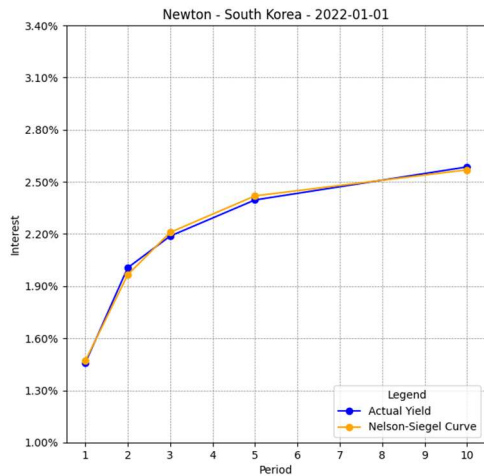
Even if in the first case the actual yield is fitted in a better way, also in the case of July the fit is pretty accurate.

As usual, Gradient descent fails in fit perfectly the yield curve when we face sudden changes. This is the case of the 1st of December 2023:



4.4.2 Newton Method South Korea:

Once again, the Newton method shows to be more accurate:



5 Comparison between Gradient Descent and Newton methods

Based on the results we have presented so far, it's evident that disparities exist between the outcomes obtained using the Gradient Descent method and those using Newton's method. These discrepancies extend beyond mere implementation nuances and likely stem from inherent divergences in the optimization strategies employed by each method.

First, we noticed that in the case of the Gradient Descent method convergence to the optimal values is way slower, while Newton always converges in less iterations. This difference is given by the fact that the former method doesn't consider information about the curvature of the function, while the latter is able to capture them by employing the Hessian.

Newton's method has stronger constraints in terms of the differentiability of the function than Gradient descent. If the second derivative of the function is undefined in the function's root, then we can apply gradient descent on it but not Newton's method.

Moreover, the fact that the second derivative is required to apply the Newton method makes it more computationally expensive with respect to the Gradient Descent one, especially in the case of high-dimensional problems.

Newton and gradient descent method are gradient based algorithms, given that, a starting point is chosen and then the algorithm is forced to explore a small area around it. For this reason, other methods tend to outperform with respect to them, since we are trying to optimize a function with multiple local optima.

The gradient based method depends heavily on the starting points, so if the ones chosen are not the correct ones, this can lead to heavy errors in the optimization. However, from a financial perspective, employing the model in a financial frame allows a good initial guess of the parameters, leading to the possibility to make use of gradient-based methods in an efficient way.

Moreover, for what concern the Svensson model, proof have been found that gradient based methods are unable to exploit the extra term of the extended model, so they seem not to be that efficient while studying this algorithm due to some limitations. In the plots we presented before about the Gradient Descent it appears this empirical evidence, we can see that most of the time the Nelson Siegel Model tend to be more precise than the extend one.

After considering gradient descent and Newton's method, we decided to implement two additional unconstrained optimization methods in our analysis. These alternative methods combine the benefits of the two aforementioned methods without having to incur their typical disadvantages.

6 Other approaches implemented in the optimization of the model:

We will see the most common approaches used to get an estimation of the model parameters.

We underline that, the approaches that are most used by analyst are, ordered by number of times used:

1. Ordinary Least Squares
2. Quasi-Newton method: BFGS algorithm
3. Levenberg-Marquardt method

6.1 Ordinary least squares– Ridge regression approach:

Due to the convenience and the simplicity of linearizing the model, grid search or OLS approach are the most used estimation procedure to optimize the Nelson-Siegel model.

This approach is performed by fixing the parameter of the shape λ .

The main issues recurring in these approaches are that they can occur in the complication that the Nelson-Siegel model becomes heavily collinear, and this depends on the fixed shape parameter.

This problem is solved by performing the so-called ridge regression: in explicit a regularization term is added to the traditional least squares objective function of linear regression.

The regularization term is composed by a regularization parameter, which controls the strength of the regularization, multiplied by the sum of the squared values of the coefficients.

The aim of this regularization term is to help to prevent overfitting and to stabilize the model by providing a balance between a well fit to the data and the necessity of keeping coefficients far from being too large.

In this approach the non-linear problem is transformed into a linear one by fixing λ , that is the parameter responsible of the lack of linearity in the model; once the model has been linearized, in order to obtain parameters that ensures the best fit, they are estimated by OLS, conditional upon a grid of the fixed shape parameter.

The parameters are estimated by minimizing the sum of squared error, by using a grid search in order to determine the optimal λ , which is chosen according to the value of the R^2 , and linear regression.

Once the optimal λ has been detected, there is the need to test the degree of multicollinearity of the two factors, if this degree is too high the results need to be re-estimated using ridge regression.

The main advantage of this method is that it does not depend on any starting values, while non-linear optimization techniques give estimates that are very sensitive to the starting value of the optimization.

On the other hand, as we stated before, in this case it must be kept in mind that the Nelson-Siegel model is very sensitive to the choice of the λ (which is usually set to be 1.37, Diebold and Li (2003) or 3, Fabozzi et al. (2006)). To understand the magnitude of this issue de Pooter (2007) studied that with different λ fixed, the remaining parameters can take extreme values.

Moreover, the correlation between the two parameters varies depending on the remaining maturity of the financial instruments chosen in the bootstrap.

7 Quasi-Newton Methods and BFGS algorithm

We solved our optimization problem first using gradient descent and then using Newton method. We highlighted the major differences between these 2 models. Summarizing, the Newton method:

- Does not need a learning rate parameter
- Converges much faster than gradient descent
- Is sensitive to initial conditions, more than gradient descent, especially if our objective function is non-convex.
- Is very computationally expensive, with a computational time of $O(n^3)$, due to the fact that we have to compute the Hessian and its inverse

We can obtain a method which is a sort of hybrid between gradient descent and Newton's method, where we can have faster convergence than gradient descent, but lower operational cost per iteration than Newton's method. This class of optimization methods is called *quasi-Newton methods*.

Recall that in Newton's method, we have to make the following update at each iteration:

$$x_{k+1} = x_k - (D^2 f(x_k))^{-1} \nabla f(x_k)$$

where $D^2 f(x_k)$ is a positive definite Hessian. If instead of the Hessian $D^2 f(x_k)$ we use an approximation $B_k \approx D^2 f(x_k)$, we can have a much faster algorithm comparing to Newton's method. This happens because the positive definite matrix B is updated iteration to iteration using information computed from previous steps, so we compute fewer new quantities at each iteration.

A common feature to all the quasi-Newton methods is that the Hessian approximation B must satisfy the *quasi-Newton condition* (or *secant equation*):

$$B_{k+1}[x_{k+1} - x_k] = \nabla f(x_{k+1}) - \nabla f(x_k)$$

which is obtained from the first order Taylor expansion of $\nabla f(x_{k+1})$ about $\nabla f(x_k)$. This condition essentially states that the product of the approximation of the inverse Hessian (B_k^{-1}) and the change in the variable space ($x_{k+1} - x_k$) should approximate the difference in gradients ($\nabla f(x_{k+1}) - \nabla f(x_k)$). In other words, it expresses a relationship between changes in the variable space and changes in the gradient.

We can understand quite easily this condition in one dimension case, where we replace the second derivative with its finite difference approximation, but things could complicate when dealing with n-dimensional secant condition.

The problem now is that our equation only has n components, while B is in general a symmetric $n \times n$ matrix with $n(n+1)/2$ components. We are dealing with an undetermined system (while in the one-dimension case we had a square system).

The quasi-Newton method solves this problem by imposing further constraints on B to solve for it. These additional constraints depend on the specific quasi-Newton method used. In this case, we focus on the BFGS method, which is one of the most popular quasi-Newton methods.

7.1 BFGS optimization

The name of the method comes from the names of its creators: Broyden, Fletcher, Goldfarb, and Shanno, who each came up with the algorithm independently in 1970.

To determine a scheme for B in $n > 1$ dimensions, we will need additional constraints. Two of them are positive-definiteness and symmetry of B (these properties should be valid in each update).

A third property we want is for B_{k+1} to be sufficiently close to B_k at each update $k + 1$. We make use of the matrix norm to characterize this property. But we recall from the Newton method that we need the Hessian's inverse (and not the Hessian itself), so we can compute it directly:

$$\min_{B_{k+1}^{-1}} \|B_{k+1}^{-1} - B_k^{-1}\|$$

$$\text{subject to } B_{k+1}^{-1 T} = B_{k+1}^{-1}$$

and $\Delta x_k = B_{k+1}^{-1} y_k$,

where $\Delta x_k = x_{k+1} - x_k$ and $y_k = \nabla f(x_{k+1}) - \nabla f(x_k)$.

Instead of requiring the full Hessian matrix at the point x_{k+1} to be computed as B_{k+1} , the approximate Hessian at stage k is updated by the addition of two symmetric rank-one matrices:

$$B_{k+1} = B_k + U_k + V_k$$

In order to maintain the symmetry and positive definiteness of B_{k+1} , the update form can be chosen as:

$$B_{k+1} = B_k + \alpha u u^\top + \beta v v^\top$$

We impose the secant condition $B_{k+1} \Delta x_k = y_k$ and we choose $u = y_k$ and $v = B_k \Delta x_k$. We then have:

$$\alpha = \frac{1}{y_k^T \Delta x_k}$$

$$\beta = -\frac{1}{\Delta x_k^T B_k \Delta x_k}$$

Substituting α and β into the previous equation we have the BFGS update:

$$B_{k+1} = B_k + \frac{y_k y_k^T}{y_k^T \Delta x_k} - \frac{B_k \Delta x_k \Delta x_k^T B_k^T}{\Delta x_k^T B_k \Delta x_k}$$

7.2 Code implementation of BFGS algorithm

We started the code by defining 4 empty vectors to store values for parameters and f values for NS and NSS model.

We then implemented a for cycle in which, for each country at each point in time:

1. We minimize the function “*fun.compute_f()*”, using “*scipy.optimize*” module and the method “*BFGS*”
2. The results are saved in two lists
3. The bonds’ yields with NS and NSS are computed on the last results and saved into two dataframes
4. The graph of NS and NSS model with BFGS method are plotted against our sample data
5. The parameters and f values for NS and NSS are appended into the corresponding lists
6. The lists are saved into excel files

```

for name_country, df_joint_country in all_df_joint.items():
    # BFGS method
    params_values_bfgs = []
    f_values_bfgs = []
    params_values_bfgs_NSS = []
    f_values_bfgs_NSS = []

    for index, df in enumerate(df_joint_country):
        results_NS = minimize(lambda params: fun.compute_f(df['Yield'], df['Maturity'], params_NS=params), params_NS, method='BFGS')
        results_NSS = minimize(lambda params: fun.compute_f(df['Yield'], df['Maturity'], params_NSS=params), params_NSS, method='BFGS')
        params_NS_BFGS = results_NS.x
        params_NSS_BFGS = results_NSS.x
        df['Nelson-Siegel'] = fun.compute_R(df['Maturity'], params_NS=params_NS_BFGS)
        df['Nelson-Siegel-Svensson'] = fun.compute_R(df['Maturity'], params_NSS=params_NSS_BFGS)
        fun.plot_curve(maturities, df['Yield'], df['Nelson-Siegel'], name_country, 'Nelson-Siegel', 'BFGS', dates[index])
        fun.plot_curve(maturities, df['Yield'], df['Nelson-Siegel-Svensson'], name_country, 'Nelson-Siegel-Svensson', 'BFGS', dates[index])

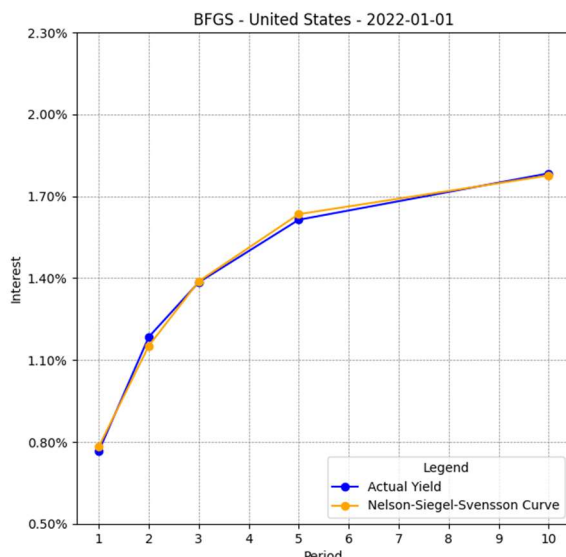
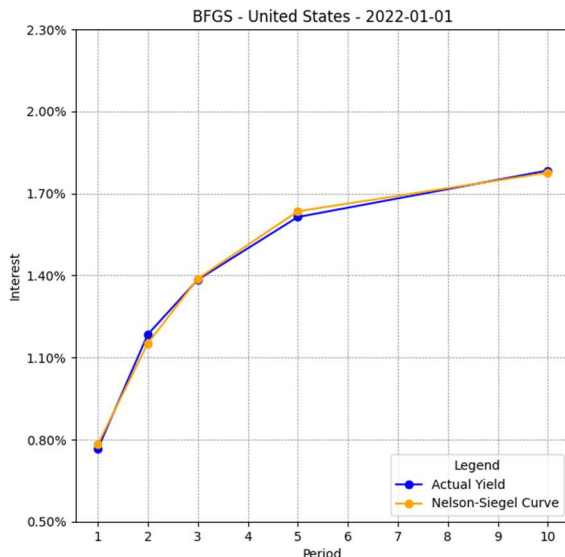
        params_values_bfgs.append(results_NS.x)
        f_values_bfgs.append(results_NS.optimality)
        params_values_bfgs_NSS.append(results_NSS.x)
        f_values_bfgs_NSS.append(results_NSS.optimality)

    # Save everything in Excel
    fun.excel(params_values_bfgs, name_country, 'Nelson-Siegel', 'BFGS', 'Parameters')
    fun.excel(f_values_bfgs, name_country, 'Nelson-Siegel', 'BFGS', 'Function')
    fun.excel(params_values_bfgs_NSS, name_country, 'Nelson-Siegel-Svensson', 'BFGS', 'Parameters')
    fun.excel(f_values_bfgs_NSS, name_country, 'Nelson-Siegel-Svensson', 'BFGS', 'Function')

```

In minimizing the function, we set the vector of starting points to be equal to that we used for gradient descent, as we don’t need to search for an optimal starting point to optimize with BFGS.

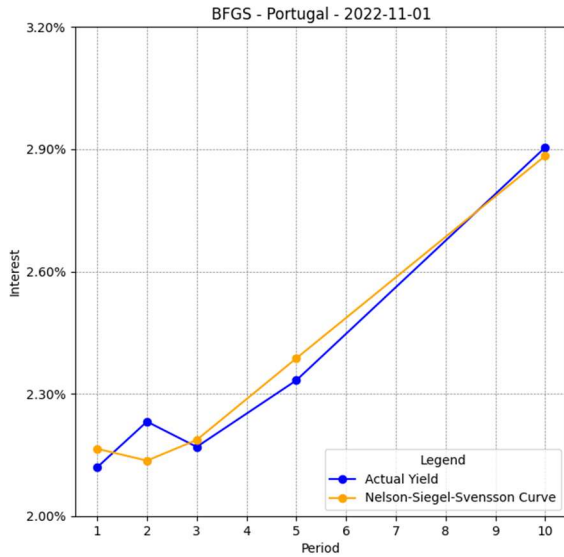
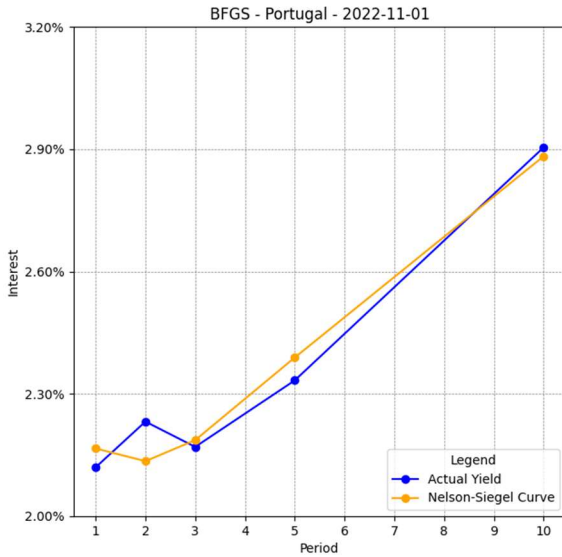
Let’s now observe the resulting graphs. Here is the example for US bonds on the 1st of January 2022:



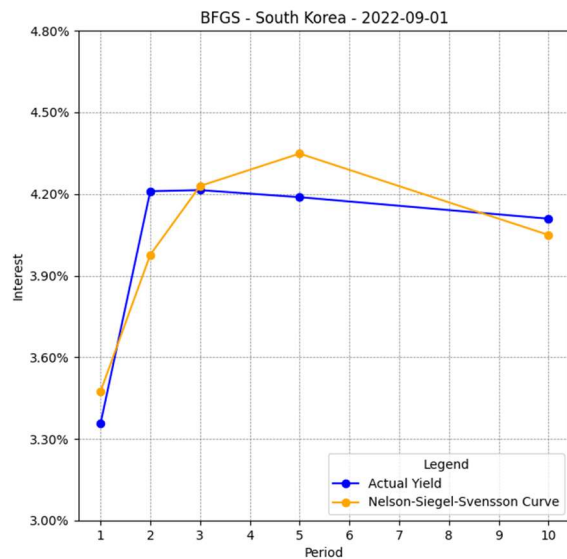
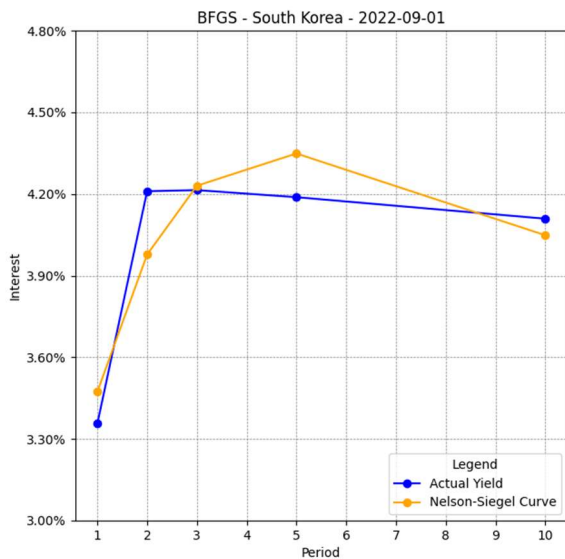
As in the Newton case, here both NS and NSS curves fit almost perfectly the historical yields data,. Unfortunately, we cannot appreciate the difference between the 2 models just by looking at these graphs. The main difference between BFGS and the previously mentioned methods is the faster convergence: NS model converges in just 3 iterations with BFGS method, while both Gradient descent and Newton took 100 iterations (which we recall is the maximum value we set for iterations). NSS using BFGS is slower than NS, taking 5 iterations to optimize the function.

Unlike the other methods, here we cannot observe any notable difference between the graphs of NS and NSS. This happens for all countries at all points in time.

Here an example for Portugal bonds on the 1st of November 2022:



And another example for South Korean bonds on the 1st of September 2022:



What instead seems to be clear (especially from these 4 graphs) is that the BFGS finds itself in difficulty in cases where bond yields do not constitute a real curve. In this case the BFGS method approximates the returns with a curvilinear shape and, consequently, turns out to be less precise.

8 The Levenberg-Marquardt (LM) algorithm

The last method we used to fit our data is the Levenberg-Marquardt algorithm (from now on we'll refer to it as "LM"), which is used to solve non-linear least squares problems. This algorithm is a blend of two minimization methods: the gradient descent method and the Gauss-Newton method. It was first published in 1944 by Kenneth Levenberg¹ and then rediscovered in 1963 by Donald Marquardt².

The main advantages of LM are:

- Fast convergence: The LM algorithm can converge faster than standard gradient descent method.
- Robustness: The LM algorithm is more robust than Newton algorithm, which means that in many cases it finds a solution even if it starts very far off the final minimum.
- Efficiency in small datasets: Particularly effective in scenarios with limited data.

While the main limitations are:

- High computational time: The LM algorithm tend to be slower than Newton algorithm and, for large datasets, it is therefore better to use another algorithm with a lower computational time.
- Calibration of lambda: Choosing the right damping factor can be challenging and may require experimentation.

Let's now observe the algorithm's formulation. Given a set of m empirical pairs (x_i, y_i) of independent and dependent variables, we want to find the parameters β of the model curve $f(x, \beta)$ so that we minimize the sum of the squares of the residuals $S(\beta)$:

$$\operatorname{argmin}_{\beta} \sum_{j=1}^m [y_i - f(x_i, \beta)]^2$$

Like with the other methods, we have to provide an initial guess for the vector β , which will be:

- A uniformed standard guess like $B^T = (1, 1, \dots, 1)$ in case with only one minimum
- A guess as close as possible to the final solution in case with multiple minima

In each iteration step, the parameter vector β is replaced by a new estimate $\beta + \delta$. The function $f(x_i, \beta + \delta)$ is approximated as:

$$f(x_i, \beta + \delta) \approx f(x_i, \beta) + J_i \delta$$

¹ Levenberg, Kenneth (1944). "A method for the Solution of Certain Non-Linear Problems in Least Squares". *Quarterly of Applied Mathematics*. 2 (2): 164–168.

² Marquardt, Donald (1963). "An Algorithm for Least-Squares Estimation of Nonlinear Parameters". *SIAM Journal on Applied Mathematics*. 11 (2): 431–441

where $J_i = \frac{\partial f(x_i, \beta)}{\partial \beta}$ is the gradient of f with respect to β .

The sum of square residuals is minimized at a zero gradient with respect to β . The first-order approximation of $f(x_i, \beta + \delta)$ is:

$$S(\beta + \delta) \approx \sum_{j=1}^m [y_j - f(x_j, \beta) - J_j \delta]^2$$

If we take derivative of this approximation with respect to δ and set the result equal to zero, we get:

$$(J^T J) \delta = J^T [y - f(\beta)]$$

where J is the Jacobian matrix of size $m \times n$.

The multiplication $(J^T J)$ yields a $n \times n$ square matrix which, in combination with vector of size n resulting from the product on the right-hand side of the equation, form a set of n linear equations, which can be solved for δ .

The most important point of this model is the addition of the damping factor λ to the last equation, which becomes:

$$(J^T J + \lambda I) \delta = J^T [y - f(\beta)]$$

where I is the identity matrix.

Small values of λ results in a Newton update, while large values of λ results in a gradient descent update. λ is initialized to be large so that first updates are small steps in the steepest-descent direction. If any iteration happens to result in a worse approximation, then λ is increased.

Otherwise, as the solution improves, λ is decreased, the LM method approaches the Gauss-Newton method, and the solution typically accelerates to the local minimum.

Unfortunately, for large values of λ , the step will be taken approximately in the direction opposite to the gradient. If either the length of the calculated step δ or the reduction of sum of squares from the latest parameter vector $\beta + \delta$ fall below predefined limits, iteration stops, and the last parameter vector β is considered to be the solution.

To make the solution scale invariant Marquardt's algorithm solved a modified problem with each component of the gradient scaled according to the curvature. This provides larger movement along the directions where the gradient is smaller, which avoids slow convergence in the direction of small gradient. Fletcher (1971) simplified the form, replacing the identity matrix with the diagonal matrix consisting of the diagonal elements of $J^T J$

$$(J^T J + \lambda \text{diag}(J^T J) \delta = J^T [y - f(\beta)]$$

8.1 Code implementation of LM algorithm and comparison with BFGS

Here we show the full code implementation of LM, which is very similar to that of BFGS method.

```
for name_country, df_joint_country in all_df_joint.items():
    # Levenberg-Marquardt method
    params_values_list = []
    f_values_list = []
    params_values_list_NSS = []
    f_values_list_NSS = []

    #Levenberg-Marquardt method
    for index, df in enumerate(df_joint_country):
        results1 = least_squares(lambda params: fun.compute_f_lm(df['Yield'], df['Maturity'], params_NS=params), params_NS, n=df['Nelson-Siegel'])
        results2 = least_squares(lambda params: fun.compute_f_lm(df['Yield'], df['Maturity'], params_NSS=params), params_NSS, n=df['Nelson-Siegel-Svensson']) = fun.compute_R(df['Maturity'], params_NS=results1.x)
        df['Nelson-Siegel-Svensson'] = fun.compute_R(df['Maturity'], params_NSS=results2.x)
        time = range(1, len(df['Yield'])) + 1
        fun.plot_curve(time, df['Yield'], df['Nelson-Siegel'], 'Nelson-Siegel', 'LM', dates[index])
        fun.plot_curve(time, df['Yield'], df['Nelson-Siegel-Svensson'], 'Nelson-Siegel-Svensson', 'US', dates[index])

        params_values_list.append(results1.x)
        f_values_list.append(results1.optimality)
        params_values_list_NSS.append(results2.x)
        f_values_list_NSS.append(results1.optimality)

    # Convert lists to dataframes
    df_params = pd.DataFrame(params_values_list)
    df_f = pd.DataFrame(f_values_list)
    df_params_NSS = pd.DataFrame(params_values_list_NSS)
    df_f_NSS = pd.DataFrame(f_values_list_NSS)

    # Save everything in Excel
    fun.excel(params_values_list, name_country, 'Nelson-Siegel', 'LM', 'Parameters')
    fun.excel(f_values_list, name_country, 'Nelson-Siegel', 'LM', 'Function')
    fun.excel(params_values_list_NSS, name_country, 'Nelson-Siegel-Svensson', 'LM', 'Parameters')
    fun.excel(f_values_list_NSS, name_country, 'Nelson-Siegel-Svensson', 'LM', 'Function')
```

As for BFGS, we set the vector of starting points to be equal to that we used for gradient descent, as we don't need to search for an optimal starting point to optimize with LM.

The main difference between our code for LM method and that for BFGS is that here we don't use the Nelson-Siegel-Svensson model.

The NSS requires residuals to be at least equal to the number of parameters we need to minimize. For NSS model we need to minimize 6 parameters ($\beta_0, \beta_1, \beta_2, \beta_3, \lambda_1$ and λ_2) but in our dataframes we have just 5 maturities (1,2,3,5 and 10 years).

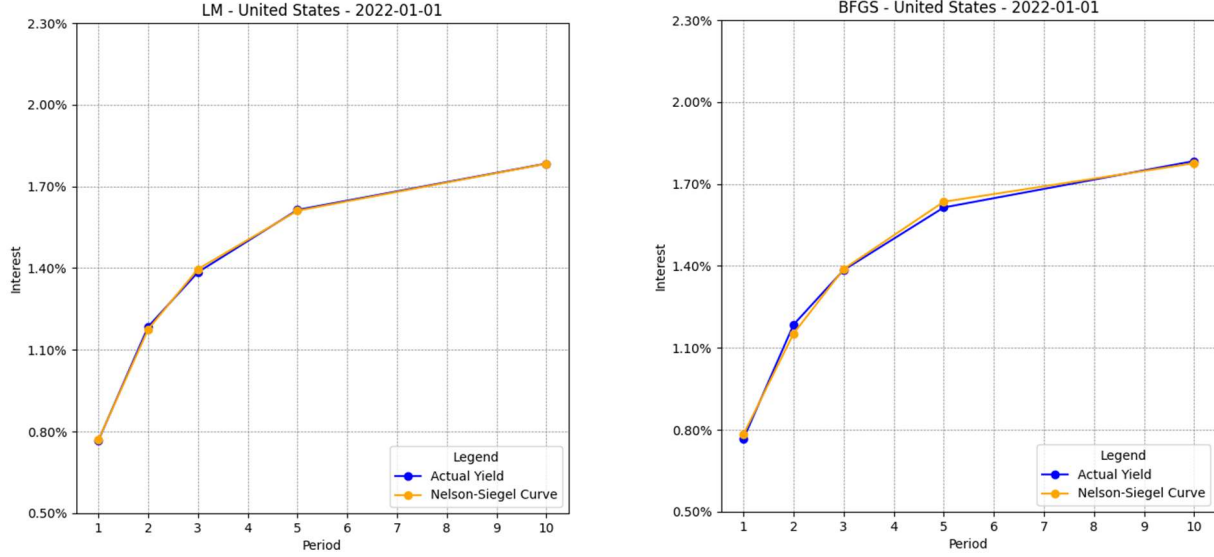
In the initial phases of our work, we had 6 maturities (1,2,3,5,7, and 10 years), but we removed the 7 years maturity when we noticed that South Korea does not have a 7 years bond from which we could take data.

Therefore, we have just 5 residuals and this is why we cannot compute NSS model using LM method.

In real world we would not have this problem, as we could take data from the shortest (1 month) to the longest (i.e. 30 years or 50 years) maturities.

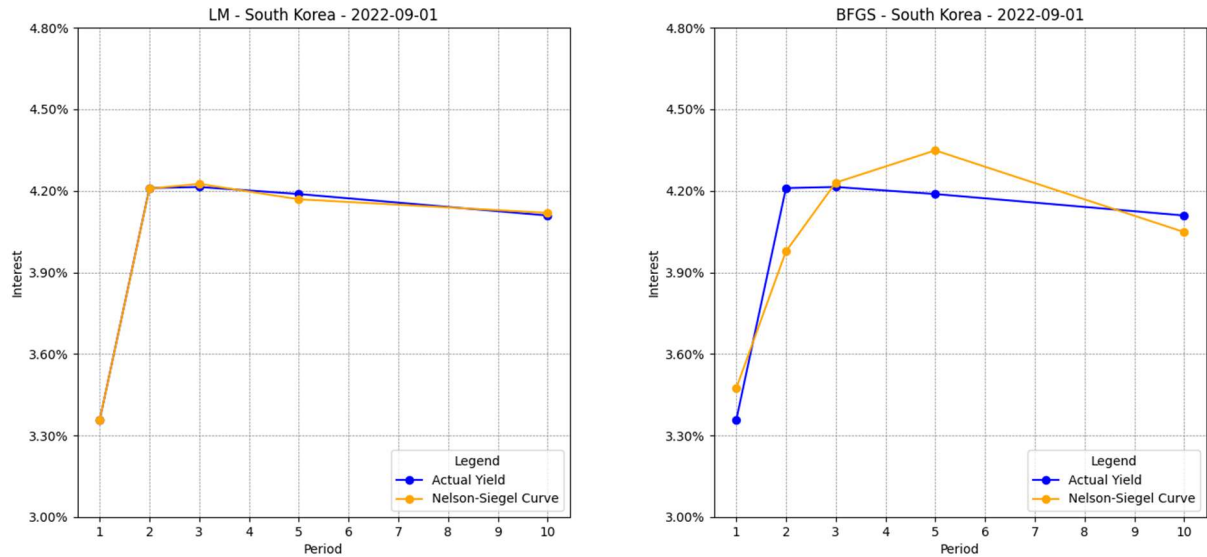
So, for the LM method, we will just show graphs for the NS model, and we will make a comparison with the other methods to see the differences in the fit to sample data.

We can now observe how this last model performs on our sample data. Here we show the graph for US bonds on the 1st of January 2022 (comparison between LS method and BFGS method for NS model):



The fit in the LM case is almost perfect, improving the improving the already excellent result obtained with BFGS. Furthermore, this is achieved with the same number of iterations as BFGS, i.e. 3.

In the LM case what convinced us the most is that this method has a very good fit to sample also when there is some volatility. We report again a comparison between LM and BFGS but now on South Korean bonds on the 1st of September 2022:



We clearly see the difference in fit between the 2 methods. LM does not tend to create a curve on bond yields like BFGS does. The result is an almost perfect fit also when volatility shows up.

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