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Abstract—Covid-19 is the most recent strain from the corona virus family that its rapid spread across the globe has caused a pandemic, resulting in over 200,000,000 infections and over 4,000,000 deaths so far. Many countries had to impose full lockdowns, with serious effects in all aspects of everyday life (economic, social etc.). In this paper, a computational framework is introduced, aptly named COVID-LIBERTY, in order to assist the study of the pandemic in Europe. Initially, the mathematics and details of the computational engine of the framework, a feed-forward, back-propagation Artificial Neural Network are presented. 5 European countries with similar population numbers were chosen and we examined the main factors that influence the spread of the virus, in order to be taken into consideration in the simulations. In this way lockdown, seasonal variability and virus effective reproduction were considered. The effectiveness of lockdown in the spread of the virus was examined and the Lockdown Index was introduced. Moreover, the relation of Covid-19 to seasonal variability was demonstrated and the parametrization of seasonality presented.

Keywords—COVID-LIBERTY, machine learning, Covid-19, Artificial Neural Network, Lockdown Index, Seasonal Variability, Effective Reproduction Number

I. INTRODUCTION

The Covid-19 respiratory disease, caused by the Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) was first discovered in Wuhan, China, near the end of December 2019. Following its global spread, it was declared as a public health emergency by the World Health Organization (WHO) on January 20\textsuperscript{th} 2020 [1]. In the course of one and a half year, it has affected 219 countries and there have been over 200,000,000 confirmed cases and over 4,000,000 deaths globally [2]. Since its discovery, scientists and medical experts around the world have been working towards treatment methods, control of the virus spread, and development of a vaccine. Most countries have experienced so far three distinct periods, where there was a
rise in the number of cases and deaths (pandemic waves): the first in spring 2020, the second in autumn 2020 and the third in winter-spring 2021 [3].

In recent years, Machine Learning (ML) methods are being utilized in various fields of medicine (for a review see [4]). Because of their nature, they are considered most suitable for application in Covid-19 research. A brief overview of ML applications related to Covid-19 follows.

ML methods, such as Convolutional Neural Networks, Support Vector Machines and Random Forests, have been utilized to propose improvements in fast diagnosis of cases, in order to augment the screening process of a patient with Computed Tomography images, X-Ray scans and blood samples [5].

In an attempt to improve Covid-19 diagnosis based on clinical data, the use of statistical analysis and ML was proposed to uncover relevant features in common clinical test results [6]. The SOM (self-organizing map) method was employed, along with Random Forest and LASSO regression models in order to classify whether a case is Covid-19 or common influenza and thus, find distinguishing features in the datasets.

SIR-based (Susceptible, Infectious, Recovered) models may help with predicting the spread of a disease, assessing the threat posed to a social group or predicting the mortality rate. Due to their high degree of complexity and the possibility of uncertainties in regards to the pandemic, SIR-based models may prove difficult to apply. To address these issues, ML methods have been employed in order improve the usability of these models [7].

A study on mortality rate in Korea [8] investigated possible factors associated with mortality and high risk through multivariable analysis of demographic and clinical data (age, income, underlying diseases, cardiac symptoms and others), identifying associations between older age, chronic lung disease, asthma with high risk of mortality and between young age, infection from personal contact with low mortality risk, among others. Five different ML classifiers were employed using the above data as predictors in order to assess a case’s mortality risk, each predictor showing different levels of importance between the classifiers. Despite the satisfying results, certain predictors may prove to be more important for other datasets (i.e., different countries).

In an article recently published in the Journal of Public Affairs [9], different ML algorithms were built with the aim to forecast cases in India. In particular, Nonlinear Regression, Decision Trees and Random Forests models were trained. Due to the insufficient data from India, data from China were used, because of similarities in both countries’ features (GDP, population, education level, etc.). Lockdown, severity of lockdown, number of deaths, recovered patients, confirmed cases and performed tests were the features which were utilized to train the model. The results analysis outlined the importance of a lockdown feature on training ML models and may be considered as evidence to the positive effects of lockdown.

Artificial Neural Network (ANN) models have also been utilized in the scientific community’s attempt to analyze existing data and predict future trends of the disease. By utilizing epidemic data from the outbreak of the disease, researchers in [10] tested 14 ANN-based models to estimate the confirmed cases in China, Iran, South Africa, Italy, United States of America, Singapore and Japan. Each Network was trained solely on number of confirmed COVID-19 cases from previous days with the current day being the reference data, the first using only the previous day, and each subsequent Network going one day back, with the 14th being trained on 2 weeks of daily cases. The results showed that the Network trained on 2 weeks ranked first in terms of overall accuracy among all countries. The results themselves, however, were not satisfactory, possibly because using only reported cases as data is not enough to identify the difference in viral spread between different countries.

In this work, a modeling framework was designed around a feed-forward back-propagation ANN in order to study the evolution of Covid-19 in 5 European countries, namely Austria, Belgium, Greece, Portugal and Sweden. The framework was named COVID-LIBERTY: Cases OVer tests IDentifier-Lockdown Index Based Ensemble modeling with effective Reproduction and Temperature variabilitY. All countries have a population of the order of 10,000,000 with varying population densities (Sweden being the less and Belgium being the more densely populated of the 5, with Austria, Greece and Portugal being in between with similar orders of magnitude of population density) and they are located in different geographical parts of Europe, i.e. Greece and Portugal in the south, Austria in the centre and Belgium and Sweden in the north. All countries except for Sweden enforced periods of lockdown. These 5 countries were chosen in order to demonstrate the importance of lockdown in keeping down the numbers of infections and to put forward the proposition that seasonal variability is related to the ability of the virus to transmit,
as is the case with respiratory diseases (see e.g. [11]). In Part 1 of this work, it is established that lockdown and weather conditions are important factors that should be taken into consideration, along with the effective reproduction number. In the following sections, the architecture of the feed forward, back-propagation ANN, which constitutes the main computational engine of COVID-LIBERTY, is detailed. Then, the data acquisition procedure is given, followed by discussion on the virus effective reproduction. Then, separate analyses show the importance of lockdown and seasonal variability in the spread of the virus and the Lockdown Index and a parametrized effective country temperature are introduced. This paper acts as a precursor to Part 2, where the COVID-LIBERTY framework and the idea of ensemble modeling are introduced and tested.

It has to be noted that it is not the aim of the authors to compare their results with results from other similar models but rather to demonstrate the ability of the current framework to operate as a useful tool for predicting future trends of the pandemic. Its assessment and comparison with other similar tools will be performed in a future work. Moreover, different parameters to the ones used in other models have been utilized and new parametrizations have been derived exclusively for the scope of this work, thus making comparisons with other models difficult. It has to be noted that to the best of our knowledge, the idea of ensemble modeling within the same framework for the study of Covid-19, parametrizing a number of important factors and conditions, has not been utilized in any other current study.

II. THE COMPUTATIONAL ENGINE OF COVID-LIBERTY: BUILDING THE ANN MODEL

ANNs have been utilized for a number of years in a number of diverse applications, ranging from image processing [12] to tuberculosis studies [13]. For a recent comprehensive review, see [14]. In recent years, Convolutional Neural Networks (CNNs) have been used in areas where ANNs were dominant, especially when image classification and large datasets are involved [15]. However, ANNs are still widely utilized and perform better in cases where image recognition is not necessary or when datasets are limited [16].

In COVID-LIBERTY, a feed-forward, back-propagation ANN was utilized. The feed-forward, back-propagation algorithm is still the most popular algorithm employed in ANNs, due to its performance and simplicity to implement (see e.g., [17]). In a generic ANN (with input-hidden-output layers of neurons), each inter-neural connection is defined by a particular real number, a synaptic weight \( w_{ij} \). The total number of these weights is the degrees of freedom of the ANN and in these connections resides the ‘knowledge’ of a trained ANN. Each neuron internally performs two basic operations: (a) a combination which computes a linear summation over its input signals and (b) a scaling through an activation function which bounds the result of the summation to a required interval. The most commonly used activation function (employed in the present work, as well) is the non-linear sigmoid:

\[
y(x) = \frac{1}{1 + e^{-x}} \tag{1}
\]

whose range of values is the interval \([0, 1]\).

To illustrate how for a pattern the values propagate through successive layers (input to hidden to output), imagine that \( I \) is the number of nodes of layer-\( i \) and \( J \) the number of nodes of layer-\( j \) (we are moving from layer- \( i \) to layer-\( j \)), where \( x_i \) are the values of layer-\( i \) that are fed in each node of layer-\( j \) to yield a value \( y_j \). First, each of the \( i \)-neurons will pass its value through the sigmoid function of Eq.(1) to yield the following output:

\[
x_{i}^{in} = \frac{1}{1 + e^{-x_i}}, \quad i = 1, 2, ..., I \tag{2}
\]

Therefore, each of the \( j \)-neurons receives a signal equal to:

\[
y_j = \sum_{i=1}^{I} w_{ij} x_{i}^{in} + w_{0j}, \quad j = 0, 2, ..., J \tag{3}
\]

In the previous equations, the subscript 0 corresponds to a bias, where \( x_{i}^{in} \) is taken to be equal to 1. The process described above is repeated until the values reach the output layer. The training process starts with a random distribution of the synaptic weights \( w_{ij} \) between layers. Via the process described above, a unique and wrong solution vector is produced at the output of the ANN, for a particular pattern. However, since the correct vector corresponding to the input vector for this pattern is known, a measure of the error can be calculated:

\[
E = \frac{1}{2} \sum_{k=1}^{K} (y_k - t_k)^2 \tag{4}
\]

where \( K \) is the number of output neurons and \( y_k \) and \( t_k \) are the components of their corresponding output and target values, respectively. The back-propagation algorithm with Gradient Descent is utilized in order to find the particular distribution of weights that minimize the value of \( E \). The
objective is to update $w_{ij}$ after the $n$th iteration by $\Delta^nw_{ij}$, such that the total error becomes smaller. This is achieved by determining $\Delta^nw_{ij}$ through the change in the signals $y_j$ corresponding to the neurons of layer-$j$ ($j=1,2,...,J$) (i.e., $\Delta y_j$) and $\Delta w_{ij}$ of the (n-1)th iteration (i.e., $\Delta^{n-1}w_{ij}$):

$$\Delta^nw_{ij} = \eta \cdot \Delta y_j + a \cdot \Delta^{n-1}w_{ij} \quad (5)$$

where $\eta$ and $a$ are the ‘learning rate’ and ‘learning momentum’, respectively. The values of these parameters, following the discussions in [18] and [19], are set to 0.2.

Next, the procedure is repeated for all patterns and thus an ‘epoch’ is completed. The procedure is then iterated over the necessary number of epochs to satisfy a convergence criterion. It is critical that during every epoch, the patterns are introduced at a random order. This ensures faster learning rates, avoids ‘memorizing’ and increases the capability of the ANN to handle situations it has not been trained for. After training has been completed, the obtained set of synaptic weights may be utilized in the ANN, in order to obtain predictions for patterns that the network has not been trained for.

For the current study, the ANN was also utilized during the initial stages of data acquisition, to fill in missing values for Sweden. Various configurations were tested, based on heuristics [20] and previous experience [19], in order to find the optimum combination of hidden layers (HL) and number of nodes in each HL, which gave the most accurate results in the shortest time. The configurations chosen for the ANN simulations were always 1 HL and either 20 nodes in the HL for 8 inputs and 1 output or 30 nodes in the HL for 4 inputs and 3 outputs. The convergence criterion at training was either for the cumulative error from all patterns to become 0.10% of its initial value or for 5,000,000 epochs in total to be completed. For all trainings performed, the error fell sharply to below 1% of its initial value within less than 1,000,000 epochs and then decreased slowly but it never reached the required 0.10% and all 5,000,000 epochs were completed. However, the maximum final error for all trainings was of the order of 0.69% of its initial value. Hence, the process of training was considered as successfully completed and we proceeded to the validation and prediction phase of the ANN.

III. DATA ACQUISITION AND THE IMPORTANCE OF EFFECTIVE REPRODUCTION

All data in this study were collected for the period between March 1, 2020 and April 30, 2021. For four countries, namely Austria, Belgium, Sweden and Portugal data went up to January 10, 2021, whereas for Greece data were collected up until April 30, 2021, in order to study the third wave of the pandemic. Data up until December 20, 2021, were utilized both in Parts 1 and 2 of this work. In Part 1 these data were used to establish the important factors which should be considered in the computational framework, i.e., effective reproduction, lockdown and seasonal variability. Then, in Part 2 the COVID-LIBERTY framework was trained on data up until December 20, 2020, to obtain predictions for the Christmas period (December 21, 2020 until January 10, 2021) and introduce the algorithm of ensemble modeling. Moreover, a further study was conducted for Greece (utilizing a training dataset with data until March 1, 2021), in order to demonstrate the ability of the framework to accurately predict the disease evolution during the prolonged third pandemic wave (March-April 2021). Data in all graphs are presented from March 15, 2020 onwards, when consecutive non-zero values of all variables were present in the dataset.

Covid-19 data concerning the 5 countries of this study were retrieved from [21] for daily numbers of performed Covid-19 tests, daily numbers of reported positive cases and daily numbers of reported deaths. In general, full datasets were available for all countries, with occasional, not consecutive, values missing. For such missing values, the average of their border values (i.e., of the previous and the following day) was chosen in order to fill the gaps. For Austria, the numbers of daily performed Covid-19 tests were missing before April 2, 2020 hence, it was decided for Austria to use data after that date. For Sweden, data on daily numbers of performed tests were absent (only weekly numbers were given) hence, in order not to introduce any further error, we did not perform any analysis for Sweden concerning the number of daily positive cases with respect to the number of daily performed tests. There were also missing data for the daily cases in Sweden for 73 days between August 28 and December 20, 2020. These days were not all consecutive but were split into one period of 3 days, sixteen periods of 4 days and one period of 6 days, scattered between periods where data were available. The final day for all periods contained the sum of positive cases of all consecutive days. For the case of the single periods (i.e. those of 3 and 6 days), the cumulative values were equally split between the 3 and 6 days. For the sixteen periods, where only the cumulative number of positive cases was available on the $4^{th}$ day, it was decided to train the COVID-
LIBERTY ANN with data from March 16, 2020 (when Sweden started reporting consecutive non-zero positive cases) to August 28, 2020 in order to be able to predict the missing values. For this reason, that interval was split into periods of 7 days and patterns for the training of the network were formulated, where as inputs the values of the first three days were taken, plus the sum of the values of the following 4 days. The outputs requested from the ANN were the values of the 4th, 5th and 6th days. Obviously, the value of the 7th day may be calculated through the subtraction of the 4th, 5th and 6th predicted values from the cumulative value given for the final day. Hence, 173 such patterns were formulated and the ANN was trained with 4 inputs, 3 requested outputs, 1 HL and 30 nodes in the HL with final training error of 0.65% after 5,000,000 epochs. Then, with the obtained weights, the 73 missing values were predicted and the dataset of daily reported positive cases for Sweden was completed.

In Figures 1, 2 and 3 the data until December 20, 2020 for all countries are presented (except for daily tests for Sweden), along with their 7-days moving averages, which follow closely the trend of the data curves. It was decided, based on the discussion in [18], to utilize moving averages of the data for future predictions, rather than their daily values. The 7-days moving averages were preferred to the 5- and 10-days moving averages, because it was a good compromise between the two, following closely the data curve, without smoothing it out in a way that the trend of the data was lost.

It was deemed necessary to include in our study a measure of the spread rate of the disease, which would give an indication of its future evolution. One such number, widely used, is the effective reproduction number $R_t$. $R_t$ is defined as the average number of secondary infectious cases that may potentially result from a primary infectious case. For instance, a value of 2 for $R_t$ means that an infected person may infect 2 more persons, who in turn may infect 2 persons more each and so on. For the method followed in order to derive the values of $R_t$ see [22]. It should be noted that for the derivation of $R_t$, the population density of a country (i.e., the number of people per unit area) is not taken into consideration. There is also a 7-day lag in the $R_t$ values, because there is an incubation period of up to 14 days [23]. This means that every case is considered to have contracted the virus on average 7 days prior to the date that the case is reported. Values of $R_t$ were retrieved from [24] from March 19, 2020 onwards.

Figure 4 presents the data set of $R_t$ for all countries and its 7-days moving averages, until December 20, 2021. As observed, data were missing for Greece for the period between May 22 and June 06, 2020. In order to fill the missing data, the two border values were considered (i.e., the values of May 21 and June 07), their average was calculated and all missing values were considered to take that value. This approach was preferred to regression, because the behaviour of $R_t$ is observed to be quite erratic hence, regression will not offer better error minimization and better approximation of the missing values. The importance of the variations of $R_t$ may be understood, when they are compared to the variations of the number of reported cases, where it can be seen that for all countries there was a several days’ lag between the peak of $R_t$ and the peak of the number of cases. The quantification of this variation will be made more explicit in Part 2 of this work, where $R_t$ is explicitly compared to the number of reported cases.

Population statistics data (i.e. population of each country and population density) were retrieved from [2]. Population density represents the number of people per unit area.

Meteorological data were retrieved from [25]. Only temperature data were retrieved, since as a first approximation, only temperature was considered in this study as a measure of seasonality. All countries chosen for this study fall well within the synoptic meteorological scale (i.e., similar weather conditions to all parts of the country for time scales of a few days at a time, see e.g. [26]). Hence, a daily effective temperature was calculated for each country by averaging the mean daily temperatures of 4 different geographical locations. These locations were chosen to be either in the North-East-South-West or only in the North-South directions, depending on the morphology of each country. In this way, the locations considered were: for Austria Graz, Innsburg, Salzburg and Vienna; for Belgium Anwerp, Kotrijk, Malmedy and Vieux Viron; for Greece Athens, Henaklion, Patras and Salonica; for Portugal Elvas, Faro, Lisbon and Porto; for Sweden Gothenburg, Malmö, Stockholm and Umeå. Taking into consideration the Covid-19 incubation period as for $R_t$ it was decided to attribute to each moving average of daily reported positive cases a 3-day moving average country temperature, 7 days prior to the date the cases were reported. The 3-day moving average for temperature was chosen to represent the average duration of synoptic scale meteorological phenomena. Through this parametrization,
we believe that the meteorological/seasonal conditions at the
time of infection were correctly represented.

IV. THE IMPORTANCE OF LOCKDOWN

As already mentioned, of the countries considered in
this study, only Sweden did not officially enforce any
lockdown. By lockdown, a restrictive policy is meant that
forces people to stay where they are and various aspects
of everyday life are restricted (people movement
restrictions in place, commercial shops/restaurants
closed, schools/universities operating tele-education, etc.).
Lockdown was proposed early on, as a preventive measure
for the spread of the disease [27]. Lockdown may be
classified as soft (some commercial shops, apart from
supermarkets, are in operation, limited restrictions in
people movement, etc.) or hard (no shops apart from
supermarkets are in operation, strict restrictions in people
movement and travel, restrictions to some industrial
operations). The periods of lockdown for each country
were obtained through [28] and [3]. All lockdown
information was valid until January 10, 2021, for Austria,
Belgium, Sweden and Portugal and until April 30, 2021 for
Greece. Austria imposed a hard lockdown from March 16
until April 20, 2020. Then, a second hard lockdown was
introduced from November 17, 2020, which turned into soft
between December 6 and December 26, 2020 and since that
date a hard lockdown was again in place. Belgium
introduced a hard lockdown on March 18 until April 17,
2020, which then turned into soft and eventually ended on
June 07, 2020. Then a soft lockdown was reintroduced on
October 22, 2020, it turned into hard on November 1, 2020
and continued as such. Greece introduced a hard lockdown
on March 19 until May 4, 2020. Then, it reintroduced a hard
lockdown on November 6, 2020, this turned into soft from
December 14, 2020 until January 1, 2021 and then it turned
into hard and continued as such. In Portugal, only one
period of hard lockdown was considered between April 2
and May 18, 2020. Then, in October, some minor
restrictions were introduced (earlier closing times for shops
and restaurants) that could not be classified as lockdown.
The periods of lockdown (from March 15 to December 20,
2020) for these countries may be graphically seen in Figure
5, where the 7-day moving average of reported cases is
presented, without differentiating on each panel between
hard and soft lockdowns. This however, was taken into
consideration in the COVID-LIBERTY framework with the
formulation of a Lockdown Index. This will be discussed
later in this section.

Figure 6 presents the 7-day moving averages of
reported cases and deaths for all 5 countries from March 07
to December 20, 2020 and the 7-days moving averages of \( \mathcal{R}_t \)
from March 23 to December 20, 2020. As observed,
Belgium exceeded significantly in absolute numbers the
number of reported cases and deaths of all other countries,
whereas \( \mathcal{R}_t \) displayed an erratic behavior, similar for all
countries, after the initial stages of the pandemic,
oscillating between 0.35 and 1.75. However, in these
results, the population density of each country was not
taken into consideration. Since all countries have similar
populations (ranging from just over nine million for Austria
to about eleven and a half million for Belgium), it was
decided to convert these results with respect to population
density. If all numbers (daily cases, deaths and \( \mathcal{R}_t \) rates) are
divided by the population density of the respective
countries, we believe that these relative numbers give a
good indication of the effect of Covid-19, depending on
how sparsely or densely populated a country is. The
population densities of all 5 countries are given in Table 1.

Table 1. Population densities of countries
considered in this study (retrieved from [2]).

<table>
<thead>
<tr>
<th>Country</th>
<th>Population density (people/km(^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Austria</td>
<td>109</td>
</tr>
<tr>
<td>Belgium</td>
<td>383</td>
</tr>
<tr>
<td>Greece</td>
<td>81</td>
</tr>
<tr>
<td>Portugal</td>
<td>111</td>
</tr>
<tr>
<td>Sweden</td>
<td>25</td>
</tr>
</tbody>
</table>

Figure 7 shows the converted with respect to population
density 7-days moving averages of daily cases, deaths and
\( \mathcal{R}_t \) rates. It is very clear from the results, that Sweden was
well above all other countries in relative numbers. The
relative number of cases for Sweden always remained
above that of the other countries, had a small peak towards
the end of June and from September onwards until
December 20, 2020 (when the last data for Sweden were
collected) followed a sharply increasing trend, in direct
contrast with the behaviour of the other countries. In the
relative number of deaths, two large peaks were observed,
towards the beginning and the end of the examined period
(first and second pandemic waves), which were 2 to 4 times
higher than the highest peak of the rest of the countries.
The relative \( \mathcal{R}_t \) rate of Sweden was always much higher
than those of the other countries, which exhibited a similar
behaviour with most of their relative \( \mathcal{R}_t \) rates intersecting
V. THE IMPORTANCE OF SEASONALITY

By observing the evolution of the Covid-19 pandemic in Europe from its start until recently, it is believed that weather conditions/seasonal variations play a crucial role in the ability of the virus to infect more people. The idea that climate and weather conditions are important in the evolution of diseases is not new (see e.g., [29]). It has been recently argued that seasonality might induce two outbreaks of respiratory-linked diseases per year [30]. Of the atmospheric variables that characterize weather (i.e., surface temperature, atmospheric pressure, precipitation, humidity, wind speed and direction, see [31]), temperature is the most readily available and is the one exhibiting the required variability in order to describe seasonal changes (see e.g., [26]). Hence, in this study seasonal changes were correlated to the increase/decrease in the number of reported positive cases and deaths through temperature. An effective daily temperature for each country was calculated and attributed to reported cases and deaths, as described in Section 3.

Figure 8 presents the 7-day moving average of daily cases and the corresponding temperatures during periods when no lockdown was in place for all 5 countries, during summer-autumn 2020. The results are given outside lockdown periods, in order to isolate the effect of seasonal variability. It is clear from the results that there existed a trend of inverse proportionality between number of cases and temperature (i.e., when temperatures decreased, the number of cases seemed to be increasing). This trend became clearer when cases were presented with respect to temperatures in Figure 9. Both cases and temperatures were normalized between 0 and 1, utilizing their respective maxima and minima for each country. The inverse proportionality between cases and temperatures was evident, with the lower temperatures corresponding to higher number of cases, whereas for higher temperatures, the number of cases was significantly reduced. Regression analysis with least squares was applied and a $y=a/x+b$ curve was fitted to the data. The coefficients $a$ and $b$ for all five countries are given in Table 2.

Table 2. Coefficients of equation $y=a/x+b$, where $x$ is normalized temperature and $y$ is the normalized number of reported cases for all countries in this study.

<table>
<thead>
<tr>
<th>Countries</th>
<th>$a$</th>
<th>$b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Austria</td>
<td>0.086</td>
<td>-0.089</td>
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We cannot perform the exact same analysis for the summer of 2021, since we are halfway through it and there exists the new Delta variant, which, with its very high degree of transmissivity, has made the number of infections rise significantly (see e.g. [32], [33], [34]). However, the number of deaths is particularly low, compared to the rise in infections, and this is believed to be due to weather conditions, since the herd immunity threshold of 80% due to vaccination has yet to be reached in Europe (see [3] and [35]). A rigorous analysis will be performed at a later stage (beginning of autumn), when all data from summer 2021 will be available.

The performed analyses indicate that weather conditions do play a significant role and in the COVID-LIBERTY framework seasonality will be considered through the parametrization discussed in Section 3.

VI. CONCLUSIONS

In Part 1 of this work, full analyses were performed in order to establish the link between lockdown / seasonal variability and the spread of Covid-19. Their effects were appropriately parametrized through LI (Section 4) and an effective daily temperature (Section 3), in order to be utilized in the COVID-LIBERTY framework. The importance of the effective reproduction number was also discussed. Moreover, the foundations for COVID-LIBERTY were laid by fully describing the feed-forward, back propagation ANN, which constitutes the engine of the framework. To the best of our knowledge, the proposed approach is the only one found in literature thus far, which considers a number of (readily available from public datasets) important parameters for the spread of the disease and incorporates them as parametrizations in a computational framework to predict the evolution of the pandemic. In Part 2, the setup of the COVID-LIBERTY framework will be presented, along with the algorithm of ensemble modeling. Two different periods of the pandemic will be studied, in order to assess the ability of the framework to predict the evolution of the disease.

REFERENCES


<table>
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<th>Country</th>
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<th>Script Value</th>
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<tr>
<td>Sweden</td>
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<td>0.0075</td>
</tr>
</tbody>
</table>


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Contribution of individual authors to the creation of a scientific article (ghostwriting policy)

Nicholas Christakis conceived the idea, wrote the code in Fortran77 for the ANN, organized all simulations and wrote the paper.
Michael Politis, Evangelos Pilios and Panagiotis Paraschis retrieved all weather and Covid-19 data for Sweden, performed the ANN simulations for Sweden and produced all graphs.
Panagiotis Tirchas, Minas Achladianakis and Eleftherios Avgenikou retrieved and analysed all Covid-19 data for Belgium and Portugal.
Argyri Kyriakaki, Christina Kalafati Matthaiou and Maria Kalykaki retrieved and analysed all Covid-19 data for Greece and Austria.
George Kossioris helped in organizing the layout of the paper and simulations and wrote the conclusions.

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Figure 1. Daily reported Covid-19 positive cases and 7-days moving averages for (a) Austria, (b) Belgium, (c) Greece, (d) Portugal and (e) Sweden.

Figure 2. Daily reported deaths due to Covid-19 and 7-days moving averages for (a) Austria, (b) Belgium, (c) Greece, (d) Portugal and (e) Sweden.
Figure 3. Daily performed Covid-19 tests and 7-days moving averages for (a) Austria, (b) Belgium, (c) Greece and (d) Portugal.

Figure 4. Daily $R_t$ and 7-days moving averages for (a) Austria, (b) Belgium, (c) Greece, (d) Portugal and (e) Sweden.
Figure 5. 7-days moving averages of reported positive cases between March 7 and December 20, 2020 with lockdown periods being marked.

Figure 6. Comparisons between countries of 7-days moving averages of (a) reported cases, (b) reported deaths and (c) $R_t$. 
Figure 7. Comparisons between countries of 7-days moving averages of converted with respect to population density: (a) reported cases, (b) reported deaths and (c) $R_t$.

Figure 8. 7-days moving averages of reported cases and corresponding temperatures for periods when no lockdown was in place for (a) Austria, (b) Belgium, (c) Greece, (d) Portugal and (e) Sweden.
Figure 9. Normalized number of reported cases vs. normalized temperatures of Figure 8 for (a) Austria, (b) Belgium, (c) Greece, (d) Portugal and (e) Sweden.