Mathematical and Computational Initiatives from the University of Buenos Aires to Contribute to Decision-Making in the Context of COVID-19 in Argentina

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Abstract

With the arrival of the pandemic in Argentina in March 2020, a working group of scientists from two institutes belonging to the Faculty of Exact and Natural Sciences of the University of Buenos Aires and CONICET, together with colleagues from different academic institutions in the country, decided to put forth our experience and knowledge in data science and associated disciplines, towards helping with decision-making in the context of COVID-19. Data analysis within Argentina and other countries, scenario simulation, as well as rapid response projects- mainly in the province of Buenos Aires- were all within the scope of our aim. This review article outlines some of the activities carried out by our team throughout these pandemic months.

Keywords: data-driven COVID-19 response, agent-based simulations, effective reproductive number, prognosis models, Buenos Aires
Introduction

The COVID-19 pandemic, caused by the infection of the SARS-CoV-2 virus, has caused over one million deaths worldwide during 2020 (World Health Organization, n.d.). Thousands of academic groups around the world have been working on issues related to the pandemic since the beginning of 2020; the group of Neil Ferguson at Imperial College London being of the most cited (“Report 1 - Estimating the Potential Total Number of Novel Coronavirus (2019-nCoV) Cases in Wuhan City, China” n.d.) along with the Franco-Spanish engineer Tomás Pueyo, with his renowned anecdote of the hammer and dance as key tools to face COVID-19 (Pueyo 2020).

Due to the great difficulty that health organizations have faced in finding effective treatments and vaccines, and thus containing the progression of the pandemic, new approaches have emerged that seek to use information (previously existing or generated during the pandemic) to help the decision-making process of institutional stakeholders. These are multidisciplinary approaches that consider logistical, economic, political and social aspects.

From the applied math and computer science institutes -Instituto de Cálculo (IC) and the Instituto de Ciencias de la Computación (ICC)- of the Faculty of Exact and Natural Sciences of the University of Buenos Aires (FCEyN-UBA) and CONICET, a multidisciplinary group with extensive experience in the development and research of mathematical-computational tools, who, alongside scientists from other national academic institutions, established collaborations with different jurisdictions, predominantly in the Province of Buenos Aires, in order to facilitate decision-making during the pandemic.

In this article we summarize the main activities carried out by this scientific nucleus. It is a broad set of contributions, ranging from data analysis and recommendations to different districts, to rapid response projects in territory. These developments enable the acquisition of data and organization of available resources; quantify and predict the progress of the pandemic; and facilitate the quantitative analysis of scenarios to predict the effects of certain public policies. Several of the activities respond to specific needs or requests made by the main actors in this pandemic, including hospital directors, health ministry authorities, mayors, and governors, and were designed and co-developed with these actors. Other contributions were academic initiatives that were made public or available to decision makers.

The activities carried out (outlined below) are divided in two parts; in the first we highlight data analysis initiatives, and in the second the rapid response projects.

Data analysis:

- Evaluation of different models for the estimation of key indicators used by the authorities, such as the basic reproductive number \( R_0 \) and the effective number \( R_e \) (Section 1.1), the doubling time (DT) of the number of cases (Section 1.2).
- Analysis of patient data collected by the Ministry of Health of the Province of Buenos Aires for the development of models for Automatic Prognosis (Section 1.3.1), and to be able to estimate the probability that an infected individual needs hospitalization in Intensive Care Units (ICU) (Section 1.3.2).
- Development of a portal where various indicators of the evolution of the pandemic are presented, at different levels of resolution: national, provincial, municipal and other geographical or administrative groups (Section 1.4)
- Estimates of prevalence at the national, provincial and municipal levels based on Infection Fatality Rate (IFR) estimates and information on death counts (Section 1.5).
- Analysis of different types of immunity and their consequences, considering different indicators (Section 1.6).

Rapid response projects:

- Development of an application that permits allocation of hospital beds within a network of hospitals, and its use in the southeastern region of Greater Buenos Aires (Section 2.1)
- The Covid Prevention Center at the Faculty of Exact and Natural Sciences of the University of Buenos Aires (Section 2.2)
- Agent-based simulations in the Marcos Paz district of the Province of Buenos Aires (Section 2.3)

Fundamental variables such as data privacy, responsible use of data, as well as the urgency of solutions and the limited budget available were all taken into account throughout this challenging and necessary effort to mitigate the negative impact of the pandemic in Argentina.
1. Data analysis

1.1. Estimation of the reproductive number R of the pandemic over time

Background:

A widely accepted metric to characterize incidence is the reproductive number, or R, defined as the average number of encounters with successful transmissions generated by each infectious individual.

At the beginning of an epidemic, the entire population can be considered susceptible and the dynamics of social encounters is not yet affected by sanitary measures. In this situation R assumes its value known as the basic reproductive number, or R\(_0\). Afterwards, when there may be people immune to contagion or the population adopts special hygiene and distancing measures, R is called the effective reproductive number, or R\(_{\text{ef}}\), usually satisfying R\(_0\) \geq R\(_{\text{ef}}\) \geq 0

R\(_{\text{ef}}\) serves as a guideline to synthesize the state of the epidemic in terms of the dynamics of change in the number of active infected: R\(_{\text{ef}}\) values greater than one represent an exponential (or sub-exponential) growth phase whereas R\(_{\text{ef}}\) values less than one account for a phase of decline (or suppression) in which the epidemic tends towards its extinction.

Objective:

Our specific objective is to provide a methodology for the estimation of R\(_{\text{ef}}\) in distinct parts of the country.

Methodology:

There are several techniques to determine the R\(_{\text{ef}}\) value from the evidence that accumulates as an epidemic develops.

One family of methods is based on models that explicitly define the mechanics of contagion in a population, such as compartmental models (Vynnycky and White 2010). The population is divided into categories (compartments) such as Susceptible (S), Infected (I) and Recovered (R) (in its most compact version known as the SIR model), and a system of differential (continuous time) or difference (discrete time) equations define the dynamics of the evolution from one compartment to another. In this framework, R\(_{\text{ef}}\) (t) is a variable that depends on system parameters, such as in the following system of differential equations:

\[
\begin{align*}
\frac{dS}{dt} &= -\beta(t)I(t)S(t) \quad \text{(average infection rate of susceptibles)} \\
\frac{dI}{dt} &= \beta(t)I(t)S(t) - \gamma I(t) \quad \text{(average rate of change of infectious infections)} \\
\frac{dR}{dt} &= \gamma I(t) \quad \text{(average rate of recovery and/or death)} \\
N &= S(t) + I(t) + R(t) \quad \text{(constant total population)} \\
R_{\text{ef}}(t) &= \frac{\beta(t)}{\gamma}S(t) \quad \text{(effective reproductive number, dimensionless)}
\end{align*}
\]

Here \(\beta(t)\) expresses the average per-capita infection rate, which is dependent on social behavior, whereas \(\gamma\) is the average rate of recovery and/or death. Given that \(S(t)\) is the number of susceptible individuals and that it decreases monotonically as the epidemic progresses, this value is one of the determining factors so that, after a certain time, R\(_{\text{ef}}\) (t) crosses the threshold of value 1 indicating the beginning of the extinction of the process.

We opted for this type of model as it allows us to better understand the evolution of the system and make projections within certain time windows in the future, assuming hypothetical scenarios of change, typically through non-pharmacological interventions (i.e. public policies) reflected as changes in \(\beta(t)\).

We also apply an extension of this model, known as SEIR, which adds an Exposed (E) compartment to represent the infected population that is not yet contagious, to estimate R\(_{\text{ef}}\) (t) as the exposed subgroup is relevant in the evolution of the COVID-19 pandemic.

Results:

Assuming a known gamma (in its average value) and having the data for the reported number of infected individuals Ir (t), we perform a constant parameter fit to time periods, in such a way that the norm of the mean square error between I (t) (model) and Ir (t) (data) is minimized. This procedure implies identifying the minimum amount of time points tk, with k = 0, ..., M in which it is necessary to subdivide the time periods. Another option is to choose fixed divisions \(a \text{ priori}\) for the
time periods. The first strategy aims to identify the points in time when there were significant changes in social dynamics whereas the second is more useful to compare identical time periods between different jurisdictions.

Figure 1 shows an example of a fit for Argentina choosing 14-day stretches (a sufficient period to capture the data changes in the number of newly infected, recovered, and deaths). Using a semilogarithmic scale, and using official data from cases reported by symptom onset date, the fit of the model (filled line) with the data (points) is observed. The new daily confirmed cases and the accumulated confirmed cases are shown, together with the successive values fit for $R_{eff}(tk)$.

![Number of new confirmed cases (daily and accumulated) by symptom onset date](image)

**Figure 1.** Evolution of $R_{eff}(tk)$ for Argentina fitted to a SEIR-type model.

Applying the same criteria for several jurisdictions, we can compare their temporal evolutions as shown in Figure 2.
Conclusions:

The application of the variable-time-period and fixed-time-period methods showed consistent results. In turn, the daily updated online visual presentation proved to be informative for comprehensive analysis of the temporal progression of the stages of the epidemic in different jurisdictions of the country. The flexibility of easily defining new geographic regions of interest to be monitored through the effective reproduction number makes it easier to respond quickly to changing information demands.

1.2. Doubling times

Background:

A parameter widely used by governments around the world to know the rate at which infections are growing is the doubling time of cases (DT). In Argentina, the national government has defined phases of the pandemic also according to the DT (“Fases de Administración Del Aislamiento” 2020). According to this criterion, if in a district the doubling time exceeds 25 days, that district could change the phase in the isolation/lockdown process and, consequently, reopen activities that had been prohibited. However, uncertainty associated with inferred DT is rarely reported. This is important since, for example, a doubling time of $15.2 \pm 0.1$ days should be interpreted differently than one of $15.2 \pm 2$ days. Therefore, for a reliable diagnosis of the progress of the pandemic, it is also necessary to know the precision of the estimate.

Objective:

Our objective is to estimate the doubling time of confirmed cases and the confidence interval for this estimate. We propose to do this using two variants of the exponential growth model.

Methodology:

Let $y_n$ be the number of confirmed cases up to day $n$. To estimate the doubling time on day $n$, data from the last 7 days are taken, $y_{n-6}, \ldots, y_n$, and fit to the following models:

**Figure 2.** Temporal heat map showing evolutions of the effective reproductive number $R_{eff}(t)$ for the Argentine provinces (snapshot of 11/15/2020). This display is part of the automated analysis available on the website described in Section 1.4.
• model 1: \( \log(y_n) = \alpha + \beta n + \varepsilon_n \)

• model 2: \( y_n = \gamma \exp(\beta n) + \varepsilon_n \)

The expected doubling time is \( \log(2)/\beta \). Using the least squares method we calculate an estimator of \( \beta \), which we call \( \beta^* \), and we deduce that an estimator for the doubling time is \( \log(2)/\beta^* \).

In model 1, the distribution of \( \beta^* \) is known, assuming that the errors have a normal distribution. In model 2, however, it is only possible to find the approximate or asymptotic distribution of \( \beta^* \) and, since we want to base our estimate on only 7 observations, this approximation may not be very good. Nevertheless, we calculated the confidence intervals under both models and did a Monte Carlo study to study their level of empirical coverage. We performed experiments with normal errors and with heavy-tailed errors (\( t_3 \) distribution).

Results:

The results of the Monte Carlo study are summarized in Table 1. Based on these results, we decided to use only the intervals calculated with model 1.

A confidence interval of level \( 1-\alpha \) for the doubling time under this model is

\[
[\log(2)/(\beta^* + t_{n-2, \alpha/2} \text{SE}(\beta^*)), \log(2)/(\beta^* - t_{n-2, \alpha/2} \text{SE}(\beta^*))].
\]

We have observed that on some days the data entry is delayed and compensated for in the following days, causing a variability in the data that does not reflect the reality of the epidemic. For this reason we decided to replace the random variables \( y_n \) with variables \( z_n = (y_n + y_{n-1} + y_{n-2}) / 3 \).

Preliminary simulations showed that 7 days were sufficient and that confidence intervals could not be calculated with the non-linear method. For this reason, the subsequent analysis with real data was done with the linear estimation model.

Table 1. Estimation errors obtained by Monte Carlo simulations for each of the two models

<table>
<thead>
<tr>
<th></th>
<th>Model 1</th>
<th>Model 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal error</td>
<td>0.9499</td>
<td>0.8916</td>
</tr>
<tr>
<td>( t_3 ) Error</td>
<td>0.9497</td>
<td>0.8890</td>
</tr>
</tbody>
</table>

This method was used to estimate the doubling time of different districts of Argentina (see Figure 3). We created an application, available to the public, in which these results can be visualized ("Duplicación de Casos de COVID-19 En Argentina" n.d.):
Conclusions:

The estimation of the doubling time was similar for the two fitting methods. It is important to highlight that the doubling time is dependent on the model and should be interpreted as a measure of the “instantaneous velocity” of growth of infections, not as a prediction.

1.3. Analysis and prediction of the evolution of cases

Forecasting systems (or prognosis models) can assist health personnel in decision-making, by identifying cases for which immediate attention or frequent follow-up is required. In turn, the implementation of these models allows us to explore which are the most important attributes in the automatic decision, and in this way to discover potential new early markers of the disease.

The vast majority of people infected with the new coronavirus SARS-CoV-2 develop mild or moderate forms of COVID-19. However, some present severe symptoms that require admission to the Intensive Care Unit (ICU). In this context, it is necessary to identify who are those patients who might require an ICU bed and for this purpose we study which comorbidities have the greatest association with the fact of needing ICU admission.

In this section we describe two studies carried out using data provided by the Ministry of Health of the Province of Buenos Aires, within the framework of a cooperation agreement with the Faculty of Exact and Natural Sciences of the UBA. In section 1.3.1, the objective is to predict the probability that an individual will develop a severe case of the disease, using age, sex, and comorbidities as covariates. In section 1.3.2, on the other hand, the objective is to predict whether an individual should be admitted to the ICU or not in the course of the disease. Although the majority of severe cases are admitted to the ICU, not all are, which is evidenced by the fact that there is a percentage of deceased individuals who did not go through an ICU. This may explain the small differences in the conclusions in both sections regarding risk factors.
1.3.1. Automatic prognosis

Background:

Given the diversity of data and experiences, the existing prognosis models within the framework of COVID-19 differ both in the variable to be predicted and in the attributes used. Regarding the variables to predict, most of the models seek to predict mortality, however models have also been developed to predict severity, for example, in the case of hospital stays longer than 10 days or the need for ICU (Wynants et al. 2020; Gupta et al. 2020). This is particularly important when resources are scarce. A recent systematic review compares 22 of the existing models for predicting deterioration throughout the hospital stay and mortality. This review uses a sample for validation of 411 cases and concludes that no model exceeds the univariate analysis of the most explanatory attribute (Gupta et al. 2020). These are the oxygen saturation upon entering the hospital and the age for deterioration and mortality, respectively. In another article, Yan, Zhang, et al. (L. Yan et al. 2020) use a model based on decision trees in a sample of approximately 500 cases where it shows that using blood studies they obtain F1 values close to 1 in the prediction of mortality (L. Yan et al. 2020).

The reported studies of automatic prognostic models generally use small samples for training and validation, between 100 and 500 cases. A recent study explored the association of patient attributes with mortality in a much larger sample (N~10^7 people; (Williamson et al. 2020)). This study showed that age is the variable that best explains risk, increasing with age. Among the comorbidities, those with the greatest impact are diabetes, cancer, organ transplants, deficiencies in kidney or liver function, neurological diseases, previous strokes, or deficits in the immune system. Finally, other demographic or socioeconomic variables such as sex, ethnicity or socioeconomic context have a moderate incidence on risk. The findings in this study are consistent with automatic prognostic studies in more limited populations (Gupta et al. 2020; Wynants et al. 2020).

Objective:

Develop a prototype of an automatic forecasting model of the severity of the condition that a certain person will have when contracting Covid.

Methodology:

A Random Forest model was developed for the prediction of disease severity, trained on a sample of ~ 10^5 cases, based on attributes such as age, sex, comorbidities, and symptoms.

Data provided by the Ministry of Health of the Province of Buenos Aires as of August 28 were used, in particular:

- Age and sex of the person
- Symptoms (19 values): New-onset anosmia, new-onset dysgeusia, dyspnea, odynophagia, fever, cough, arthralgia, headache, mental confusion, seizures, diarrhea, abdominal pain, chest pain, conjunctival injection, irritability, malaise, myalgia, food refusal, and vomiting.
- Comorbidities (22 values): Diabetes, pregnancy, chronic liver disease, previous neurological disease, previous cancer disease, asthma, chronic obstructive pulmonary disease (COPD), low birth weight, previous bronchiolitis, acute dialysis, chronic dialysis, ex-smoker, smoker, high blood pressure, congenital immunosuppression or acquired, heart failure, renal failure, previous community-acquired pneumonia (CAP), obesity, prematurity, tuberculosis, and a characteristic that indicates the lack of comorbidities.
- Hospitalization and death data: An indicator of whether the patient was hospitalized and, if so, whether it was in intensive care, and whether the patient died.
- All the attributes, except for age, which is measured in years, and fever, which takes three values (no fever, fever less than 38 ºC and fever greater than 38 ºC), take binary values where 1 codes if the symptom or comorbidity is present in the patient. Only confirmed COVID-positive cases were considered.

Random Forest, a standard machine learning method, chosen for its robustness, flexibility and interpretability, was used for modeling. The model was trained with the attributes listed above as input to predict the severity of each case. Severity was defined as mild if the patient recovers with or without hospitalization but without intensive care, and as severe if he/she requires intensive care and / or dies. We want to emphasize that this definition is an approximation to the severity of the case and it is not perfect since it is possible that there are cases that were admitted to the ICU only preventively without the condition being so serious (for example, in the case of pregnant women or small children). During model training, the cost corresponding to each sample was weighed by the inverse of the frequency corresponding to its class to compensate for the imbalance between the two classes.
For the development of the models, the cases with a symptom onset date (SOD) prior to July 31, 2020 were used, while for the final evaluation we used the data with SOD from August 1 to 15, 2020, discarding the more recent cases, since it is not yet known whether they will become severe cases. Only cases with at least one symptom were included in the study. The development data included 108,249 cases, of which 4.7% were severe cases. The selection of parameters was carried out with the cross-validation method on the development data. The evaluation was carried out by training a new model on the development data and calculating the performance on the evaluation data, which included 47,014 cases, of which 2.1% were severe. The percentage of severe cases is different in the developmental and evaluation data because the data were divided by date, and, in this data, the percentage of severe cases decreased over time. In order to understand which attributes were most relevant to the task, a sequential attribute selection algorithm was used (Mustafa 2017). Starting with the model that uses all the attributes of interest, attributes are discarded one at a time, discarding at each stage the one that least degrades or most improves system performance in development data. The algorithm thus generates an ordered list of attributes according to their importance for ranking in the development data.

Results:

Figure 4 shows the recall (percentage of serious cases that are detected as such) versus precision (percentage of cases detected as serious that were actually serious) curves obtained while varying the decision threshold applied to the system’s outputs for models trained with different lists of input attributes. The figure shows that age is an excellent predictor of the severity with which a case will develop and that sex adds relevant information to predict the severity of the case. These conclusions coincide with what was found in the literature (Williamson et al. 2020). On the other hand, adding comorbidities to age improves performance more than adding sex, and further including sex in addition to comorbidities makes only a marginal difference. This suggests that the information that the system extracts from the sex attribute is at least partially correlated with comorbidities. The resulting analysis of the attribute selection algorithm suggests that the most important comorbidities for classification are diabetes, obesity, kidney failure and previous oncological disease, which were also found to be relevant for prognosis in the literature. Finally, by adding the symptoms to the model, a significant gain is obtained. In this case, the selection algorithm indicates that dyspnea, odynophagia, and anosmia are the most relevant symptoms for classification. In the case of odynophagia and anosmia, the existence of the symptom occurs more frequently in mild cases than in severe cases, which had already been observed in the literature (C. H. Yan et al. 2020).

Figure 4. Recall and precision curves in evaluation data using different input attribute lists. The vertical dotted line corresponds to the percentage of severe cases. The references ‘com’ and ‘sym’ indicate comorbidities and symptoms, respectively.

Other symptoms and comorbidities in addition to those selected as important by the algorithm strongly correlate with the severity of the case, such as dialysis or heart failure when analyzed independently of the others. However, it is possible that these attributes are not chosen because they are highly correlated with others that the model does select, such as age. The next section (1.3.2) describes a work focused more specifically on the analysis of risk factors.
Conclusions:

The automatic prognosis system described in this section could be used to perform triage in cases of saturation of the health system that require postponing the care of some patients for the benefit of others with a worse prognosis. It is important to clarify that, since certain comorbidities that significantly affect prognosis occur very infrequently in the data, the model is probably not learning to correctly predict severity for those cases. For this reason, the output of the system should only be taken as a recommendation that must be confirmed by a health professional.

1.3.2. Comorbidities and risk of ICU admission due to COVID-19

Background:

In this section we aim to estimate the probability that an infected individual needs ICU admission using age, sex and the 22 comorbidities detailed in Section 1.3.1 as covariates. Our interest is focused on the analysis of risk factors.

This study is based on the data of confirmed cases collected by the Ministry of Health of the Province of Buenos Aires reported on July 7, 2020. In this section, the data analyzed corresponds to the confirmed cases of individuals under the age of 90 in the Province of Buenos Aires up until June 22, 2020, which comprises a total of 22,306 confirmed cases under study. In this analysis we discarded the confirmed cases between June 23 and July 7 since, throughout the development of the pandemic in Argentina, we have observed that, since a case is confirmed, it is necessary to allow a period of time to pass for the cases to evolve. The data indicate that for more than 99% of the individuals who were admitted to the ICU, this hospitalization occurred in 15 days or less from the confirmation of the diagnosis. An analysis of these times based on data from the entire country can be seen at http://covid.ic.fcen.uba.ar:3838/apps_ic/tiempoeventos/. The proportion of patients in this sample who required intensive care was 0.025 (2.5%).

Methodology:

In order to explore the association between the fact that a COVID-19 patient requires ICU or not and comorbidities, classification trees were used that constitute a descriptive tool based on graphs that use the branching method to represent the possible outcomes of a binary decision (James et al. 2013). A summary measure of the importance of the explanatory variables in a tree can be computed by decreasing the Gini index associated with each variable. A graphic representation of the importance of comorbidities, age and sex is presented in Figure 5. These measures were calculated from 1000 trees built using resampling of the original data and were computed as the average of the importance of each variable relative to the one of greatest importance. This descriptive study suggests age, obesity and hypertension as the variables of greatest importance.

![Figure 5](image-url)
In order to make statistical inference, a fit was performed by logistic regression (Hosmer, Lemeshow, and Sturdivant 2013) categorizing the data into two classes, one representing those who required ICU and the other those who did not, and using comorbidities as explanatory variables. To that end, if \( y \in \{0, 1\} \) represents the dichotomous response and \( x \) represents the vector of covariates, the logistic regression model assumes that \( P(y = 1 \mid x) = \text{expit}(\beta^T x) \), where the function \( \text{expit}(t) = 1 / (1 + \exp(-t)) \). Its inverse is the function \( \text{logit}(p) = \log(p / (1 - p)) \) that involves odds. Indeed, if \( p \) is the probability of an event, the quantity \( p / (1-p) \) is called the odds of the event. For example, the probability of getting 6 on a die is 1/6, and the odds are 1/5. This means that for each time a 6 is rolled, it is expected that there will be approximately 5 times when a 6 is not returned. Figure 6 shows the observed proportions of confirmed cases requiring ICU admission by age and the logit of these same proportions. Observe that the proportion of infected who require intensive care decreases until approximately 30 years of age, then increases until around 70 years and finally appears to decrease for women, while for men it seems to continue to increase, although perhaps more slowly. These trends are most clearly seen on the logit scale. These observations lead us to include two splines in the logistic model to explain the effect of being over 30 years old and being over 70 years old and possible interactions between sex and age. The inclusion of a spline to obtain a log-linear relationship between age and the probability of developing a severe form of the disease was already considered by Williamson et al. (Williamson et al. 2020). Likewise, since there are very few infected over 90 years of age, we decided to restrict our analysis to confirmed cases under 90 years of age. Using several variable selection techniques (best subset selection, forward, backward, stepwise, and lasso) and cross-validation with various fit evaluation measures (AIC, ECM, AUC, deviance), we chose the procedure to fit the model and identify the variables to include. In Table 2 we report the significant coefficients and their confidence intervals with a global level of 0.05.

![Figure 6](image.png)

**Figure 6.** Proportion (left panel) and proportion in logit scale (right panel) of confirmed cases admitted to the ICU by age and sex (in light blue male and pink female).
Table 2. Odds ratios estimates and their confidence intervals of level 0.95.

<table>
<thead>
<tr>
<th>Risk factor</th>
<th>Estimated OR</th>
<th>2.5%</th>
<th>97.5%</th>
<th>Risk Factor</th>
<th>Estimated OR</th>
<th>2.5%</th>
<th>97.5%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diabetes</td>
<td>1.67</td>
<td>1.12</td>
<td>2.48</td>
<td>Heart Failure</td>
<td>1.75</td>
<td>1.06</td>
<td>2.91</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>7.31</td>
<td>1.70</td>
<td>31.38</td>
<td>Obesity</td>
<td>3.48</td>
<td>2.22</td>
<td>5.44</td>
</tr>
<tr>
<td>Previous Oncological Disease</td>
<td>2.65</td>
<td>1.47</td>
<td>4.78</td>
<td>Over 30</td>
<td>1.14</td>
<td>1.07</td>
<td>1.21</td>
</tr>
<tr>
<td>COPD</td>
<td>1.99</td>
<td>1.15</td>
<td>3.45</td>
<td>Over 70</td>
<td>0.95</td>
<td>0.91</td>
<td>0.99</td>
</tr>
<tr>
<td>Hypertension</td>
<td>1.72</td>
<td>1.23</td>
<td>2.41</td>
<td>Age</td>
<td>0.95</td>
<td>0.90</td>
<td>0.99</td>
</tr>
<tr>
<td>Imm. Congenital or Acquired</td>
<td>3.32</td>
<td>1.43</td>
<td>7.71</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Results:

The odds of needing ICU admission among individuals with a risk factor, say obesity, is the ratio between the probability that someone infected with the risk factor needs ICU hospitalization and the probability that they do not need it. In turn, the odds ratio (OR) is defined as the ratio of the odds of needing ICU for those infected who have the risk factor of interest and the odds for those who do not, keeping all other explanatory variables constant. For example, according to Table 2, the chance that an obese infected person needs ICU admission is estimated to be approximately three times the chance for a non-obese person of the same age, of the same sex and with all other risk factors equal. The confidence intervals tell us what is the precision of the chances that we have estimated. For example, we can say that the chance of needing intensive therapy for the obese is estimated to be between 2.22 and 5.44 times that of the non-obese, while the chance of needing intensive therapy for diabetics is estimated to be between 1.12 and 2.48 times that of non-diabetics. With the fitted model it is possible to estimate the probability that a person diagnosed with COVID-19 needs ICU admission, knowing their age, sex and comorbidities.

Conclusions:

The statistical analysis presented allows us to identify risk factors that are important for developing severe forms of COVID-19 that require ICU admission. From this study, the following emerge as relevant risk factors: diabetes, pregnancy, previous oncological disease, COPD, hypertension, congenital or acquired immunosuppression, heart failure, obesity, age and its two associated splines that indicate being over 30 years old and over 70 years old. This set includes most of the comorbidities detected as important for predicting severity in Section 1.3.1.

1.4. Data-based analysis and control panel

Background:

The management of the pandemic is nourished by multiple sources of quantitative information that are constantly updated. In an emergency context, the structures of the data sources are subject to changes, errors, delays, etc. At the same time, the amount of available information sources tends to grow over time, but the quality of the data is not usually homogeneous. For these reasons, there is a need for a centralized resource that applies consistent, repeatable, and comparable methods of ingesting, cleansing, curating, persisting, and visualizing data.

Objective:

The objective is to produce a reliable and unified repository that offers information on the temporal evolution of metrics on COVID-19 in Argentina. On this platform, additionally, various data analyses can be produced with different objectives, creating new more complex metrics.
Methodology:

Every day the data that is updated by the Ministry of Health of the Nation contains new information and also updates on information from the past, due to multiple reasons such as reporting delays or updating the health status of patients who were registered for the first time in the system a while back. Also, previous information that has no updates is repeated every day, generating a great deal of redundancy. In this sense, our system preserves a complete copy of each update, for statistical purposes on the evolution of the data entry process itself. This means that we currently have more than 170 million records. However, when consolidating a database for monitoring the epidemic, we only keep operational changes in the epidemiological condition of each citizen.

The analysis of relevant indicators of the epidemic requires different territorial groupings depending on the official, civil servant or data analyst who consumes the information. For example, from a health point of view, in the Province of Buenos Aires it is important to analyze cases according to the cordons of the Buenos Aires suburbs, and also by health regions (“Regiones Sanitarias” n.d.).

Results:

The information presented on the site is built from multiple data sources, including: Open data from the Ministry of Health of the Nation (“COVID-19. Casos Registrados En La República Argentina” n.d.), COVID-19 Situation Room (Ministry of Health of the Nation) (“Información Epidemiológica” 2020) and Open data on the epidemiological situation of COVID-19 from the Government of the City of Buenos Aires (“Casos COVID-19” n.d.), among others.

At the time of writing this article, the system has complete data sets updated daily since July 15, containing epidemiological information accounting for the symptom onset as early as from January 1, 2020.

The visible face of the system is the web page, accessible at (“COVID-19 Análisis Y Control Basado En Datos” n.d.). The system architecture is designed so that any data analysis group can create new content that can be viewed on this site, taking advantage of the data infrastructure developed.

There we provide the ability to define territorial units that we call “Conglomerates” that consist of relevant groupings of localities. This can be seen in the tab Province of Buenos Aires and AMBA (city of Buenos Aires plus its suburbs) -> Sanitary Regions and also -> suburban edges (Cordones conurbanos in spanish). At the moment the system presents information on 550 departments nationwide and has defined 26 conglomerates of interest.

Temporal evolutions are shown in the number of daily cases and the number of accumulated cases (both by symptom onset date) and daily and accumulated deaths (by death date). Proportions of geographic concentration of confirmed cases are also shown. The information can be filtered by geographical area and consulted on a linear or logarithmic scale, in total figures or per 100,000 inhabitants and in daily values or moving averages every 7 days. The graphics are interactive, in such a way that the user can modify relevant aspects of each visualization focusing on what interests him the most.

To date, the system has been consulted by more than 850 different users with more than 3,800 visits coming 95% from Argentina and also including visits from countries in South America, North America and Europe. Likewise, the visualizations updated daily serve as input to compile reports that are provided on request and also periodically to different public administrations, such as the Ministry of Health of the Province of Buenos Aires.

The visualizations (updated daily) serve as input for compiling reports that are provided upon request as well as periodically to different public administrations, such as the Ministry of Health of the Province of Buenos Aires (see Section 1.1, Figure 1) along with an analysis of delay in reporting of deaths (see below).
Figure 7. Screenshot of the site showing daily confirmed cases and daily deaths for the Province of Buenos Aires, AMBA, AMBA not including CABA, Cordones 1 and 2 of the Greater Buenos Aires and CABA. It includes an analysis of the proportion of deaths produced in CABA, in the Province of Buenos Aires and in the rest of the country.

It can be noted that towards the end of the graphs there is a vertical dotted line that separates the last 12 days of data from the rest of the information, and that the moving averages end there. This is due to the fact that the site uses the Date of Symptom Onset (acronym in Spanish: FIS) as the date base, instead of the Report Date for cases. This is the case since the FIS provides a more stable and reliable indicator of the real dynamics of the epidemic, as it is less subject to the variability of the bureaucratic data entry process. As a counterpart, using FIS has the negative aspect that there is a delay, estimated at 12 days, until all cases that share the same FIS are reported.

Figure 8. Detail of the evolution of daily confirmed cases for Municipalities of the Province of Buenos Aires.

Finally, we show an example of data analysis that is possible exclusively if a history of all daily data updates is kept and an analysis of changes of states is applied for each individual over time (that is, it is not enough to access to “the last snapshot” of the public access databases).

Figure 9 shows the evolution of the death reporting process in CABA and PBA, focusing on the delay produced in the reporting of deaths that occur every day. The colors denote the difference between the date of death and the date of upload, for all cases that were entered into the system, day by day. This information can be found in the site’s analysis section.
The comparison is relevant for highlighting the different strategies that jurisdictions use to deal with a known problem that is present in the entire data entry system, as is the case in the delay in the reporting of deaths when health administrative systems are overwhelmed.

It arises from the analysis that the Province of Buenos Aires opted for an immediate, high intensity transparency and remediation, while the Autonomous City of Buenos Aires opted for a slow and dosed remediation strategy.

**Figure 9.** Reporting of deaths (Screenshot of site). Left: Autonomous City of Buenos Aires, Right: Province of Buenos Aires, Above: Number of deaths reported per day, Center: Percentages (normalized to 100%) of delays suffered by deaths reported each day, Below: Cumulative evolution of deaths reported.

**Conclusions:**

This project has shown very satisfactory results at the moment, since it has proven to be useful to add relevant information and instantaneously to numerous debates and discussions about the evolution of COVID-19 in our country, both in academic environments and with political officials and with public opinion. The site time series are used as input to adjust simulation models used to project possible future scenarios and suggest strategies for managing the epidemic. In this sense, having a unified and reliable panel of data such as the one presented in this section has been successfully covering a key need for advising decision-makers in public policies through mathematical-computational methods.

### 1.5. Estimation of prevalence at the national and district levels

**Background:**

We want to estimate the real percentage of infected population in each district using as input data the deaths discriminated by age range and by sex, and the lethality for each of these ranges. This also allows inferring the relationship between infected and confirmed cases.

**Objective:**

Knowing the percentage of infected population in each district is of interest when making public policy decisions, such as testing policies, tracking, restrictions on mobility, etc.

We will estimate these values for the country as a whole, for the 24 provinces of Argentina and for the 135 municipalities of the Province of Buenos Aires (PBA). The calculation for each municipality of PBA is particularly important given our close collaboration with the provincial government and the different municipalities of the Province.
Methodology:

This analysis was based on a seroprevalence study in Spain where the IFR (infection fatality rate) was obtained by age decile and by sex (Pastor-Barriuso et al. 2020). The choice of using the Spanish prevalence comes from the fact that it is the study with the highest level of detail that has been reported in the literature; however, other studies in different parts of the world show similar values. The summary of the infection fatality rate for each age decile and sex can be seen in Table 3.

Table 3. IFR by age decile and sex in Spain, estimated in (Pastor-Barriuso et al. 2020).

<table>
<thead>
<tr>
<th>Age Decile</th>
<th>0-9</th>
<th>10-19</th>
<th>20-29</th>
<th>30-39</th>
<th>40-49</th>
<th>50-59</th>
<th>60-69</th>
<th>70-79</th>
<th>80-89</th>
<th>90+</th>
</tr>
</thead>
<tbody>
<tr>
<td>M</td>
<td>0.002%</td>
<td>0.005%</td>
<td>0.004%</td>
<td>0.013%</td>
<td>0.012%</td>
<td>0.03%</td>
<td>0.02%</td>
<td>0.09%</td>
<td>0.05%</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>0.002%</td>
<td>0.005%</td>
<td>0.004%</td>
<td>0.013%</td>
<td>0.012%</td>
<td>0.03%</td>
<td>0.02%</td>
<td>0.09%</td>
<td>0.05%</td>
<td></td>
</tr>
<tr>
<td>M</td>
<td>0.38%</td>
<td>1.62%</td>
<td>0.62%</td>
<td>6.11%</td>
<td>2.68%</td>
<td>14.21%</td>
<td>5.47%</td>
<td>27.59%</td>
<td>10.51%</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>0.20%</td>
<td>1.62%</td>
<td>0.62%</td>
<td>6.11%</td>
<td>2.68%</td>
<td>14.21%</td>
<td>5.47%</td>
<td>27.59%</td>
<td>10.51%</td>
<td></td>
</tr>
</tbody>
</table>

We first estimate what we call “theoretical IFR” that comes from considering the Spanish IFR according to Table 3 and the age and sex distribution of the population of Argentina and its different jurisdictions. With these data we can obtain this theoretical IFR, which assumes that infections in all age ranges and in both sexes are homogeneously distributed. Once this theoretical IFR is obtained, plus the population of the district under consideration and the number of deaths to date, the percentage of the population actually infected can be estimated 2 weeks ago (we assume an average of 2 weeks between the reporting of a case and the date of possible death). As we know the reported number of confirmed cases, then we can estimate the coefficient of relationship between real and confirmed cases, which we will call the Quiros Factor (QF) (Figar et al. 2020) (Note that the QF is equal to 1 / IDR, where the IDR is the Infection to Detection Ratio, that is, the proportion of the unit of detected cases, defined in the literature). Given this QF and the confirmed cases in the last 2 weeks, we can estimate the percentage of infected population to date.

Next, we perform a fit on these calculations, using as input data the actual deaths, with their age ranges and sex, for each jurisdiction, in order to estimate what we call the current IFR.

Using these data on deaths as a starting point, plus the Spanish prevalence by age range and sex, we can estimate the number of real infected in each age range and sex, in each district, two weeks behind the moment in which we are making the estimate, which gives us the percentage of the infected population. With this percentage we can deduce the current IFR, and if we also use the number of confirmed cases, we can also estimate the adjusted QF (by age and sex of the deceased). Assuming that the QF is maintained for the last 2 weeks, we can now estimate the real percentage of infected to date, using again the data of confirmed cases in the last 2 weeks.

An additional observation: given that the number of deaths in the 0-19 age group is very low and that the prevalence taken in Spain for these groups was also calculated with very few cases, we have not used the prevalences of the 0-9 and 10-19 deciles in the estimation of the current IFR, and to estimate the number of real cases in both groups we have taken the percentages of infected from the 60-69 group for the 10-19 group and from 70-79 for the 0-9 group (for each district), assuming that a parallel can be established between the level of care of those respective groups and therefore that the percentage of infections has been relatively even.

Results:

Table 4 shows the situation in Argentina as of 10/30/20. The data are displayed for the country as a whole, the city of Buenos Aires, the suburbs of Buenos Aires, the rest of the Province of Buenos Aires (Interior, in spanish) and the rest of the country. There it can be seen that by that date, taking the estimate derived from the calculation that uses ages and sexes of actual deaths, Argentina already had approximately 24% of its population infected at the national level, a value that far exceeds the percentages of infected in the countries of Europe in the first wave of March / May. In the city of Buenos Aires...
and the suburbs of Buenos Aires, this value rose to approximately 37% (also strongly exceeding the level of contagion in other large urban conglomerates in the world), while in the rest of the Province of Buenos Aires and in the rest of the country that value fell to values that are around 15%.

Table 4. Situation of Argentina in terms of prevalence and IFR in the main districts of the country by the end of October 2020.

<table>
<thead>
<tr>
<th>Districts</th>
<th>Population (2020)</th>
<th>Confirmed Cases</th>
<th>Deaths</th>
<th>Cases incidence</th>
<th>Theoretical IFR (homogeneous)</th>
<th>IFR (current)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Estimated infected population (using theoretical IFR)</td>
<td>Estimated infected population (with age and sex of deaths)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>IFR (current)</td>
<td>Quiles Factor (homogeneous)</td>
</tr>
<tr>
<td>Argentina</td>
<td>45,376,763</td>
<td>1,157,174</td>
<td>36,792</td>
<td>2.19%</td>
<td>11.15%</td>
<td>0.61%</td>
</tr>
<tr>
<td>Buenos Aires City</td>
<td>3,075,664</td>
<td>146,758</td>
<td>4,907</td>
<td>4.57%</td>
<td>17.68%</td>
<td>0.90%</td>
</tr>
<tr>
<td>Buenos Aires suburb</td>
<td>13,830,369</td>
<td>458,346</td>
<td>14,093</td>
<td>3.16%</td>
<td>19.55%</td>
<td>0.56%</td>
</tr>
<tr>
<td>Province of Bs. As. (Interior)</td>
<td>3,910,772</td>
<td>166,033</td>
<td>2,324</td>
<td>1.75%</td>
<td>7.28%</td>
<td>0.82%</td>
</tr>
<tr>
<td>Real of the country</td>
<td>24,759,976</td>
<td>463,498</td>
<td>8,403</td>
<td>1.41%</td>
<td>8.14%</td>
<td>0.56%</td>
</tr>
</tbody>
</table>

As can be seen in Table 4, the current IFR values are higher than the theoretical IFRs, due to the fact that infections have not been homogeneous by age range and sex. What is perceived is something to be expected; older people have taken better care of themselves and consequently in proportional terms they have been less infected, lowering the IFR values in practically all districts (for example at the national level the theoretical IFR is 0.61, whereas the current is 0.33).

As for the real QF at the national level, this number is around 9, in line with what has been seen in other parts of the world. This means that one case would be confirmed for every 9 infections.

Conclusions:

By the end of October, the number of real infected (deduced from the lethality by age range and sex, plus the reported death data) both in the city of Buenos Aires and in the Buenos Aires suburbs was already very high. This group accounts for 40% of the total population (which has implied a high number of deaths per 100,000 inhabitants, placing the country as a whole in the top-10 in the world by the end of October). This level of contagion in Buenos Aires and its surroundings resulted in a sharp drop in the weekly number of infected, mainly due to the sharp drop in the number of susceptible people, as well as the maintenance of personal care (and despite a sharp rise in the mobility). In the rest of the country, the situation still presents comparably lower values of infected (around 15% if we look at both the interior of the Province of Buenos Aires and the rest of the country), which forces us to take extreme measures of personal care, restriction of mobility and permanent tracing of close contacts, to keep the contagion curve (and consequently of deaths) at low levels.

1.6. Immunity: from biological to social causes

Background:

Some speculation has been raised that a certain degree of herd immunity is the main cause of the decline in infections in Europe after the initial outbreak. There were claims that between 10% and 20% of infections are enough to stop growth dynamics, instead of the 60-70% indicated by simple calculations (this value is estimated at 100 * (R0 - 1) / R0). Among the speculations about herd immunity, several possible causes are mentioned: some previous immunity in the entire population due to cross-reaction with other coronaviruses, highly inhomogeneous contagion patterns, among others. However, most of these speculations fell through when the real numbers of acquired immunity were obtained: less than 10% in most cities. The parameter known as the IFR (infection-fatality ratio), which can be estimated with a sufficient degree of precision from the death count as seen in the previous subsection, even in the absence of large seroprevalence studies, also shows low prevalence values in all European countries. Taking these facts into account, the question arises as to the main cause of the rapid decline in the infection rate in European countries.
Objective:

We have investigated an alternative explanation, in which the focus of the decline in cases shifts from biological to social causes, from herd immunity to “fright” immunity.

Results:

The term fright immunity reflects the awareness of individuals after learning about the consequences of COVID-19, including deaths and serious sequelae. When the “fright” occurs, the majority isolate themselves and begin to behave in a much more cautious manner. The effective reproduction number $R_{ef}(t)$ for the entire subpopulation of “frightful” is much less than one. However, a small fraction of individuals persist in dangerous behavior (whom we call “reckless” -mostly young- people who are not afraid and are less likely to self-isolate), which although somewhat less than before the pandemic still implies a reproduction number $R_{ef}(t)$ greater than unity. Many of these individuals shared social and physical proximity, and the virus found a fairly simple way to spread. It is important to note that these fractions are not defined institutionally, nor are they necessarily determined by their employment status. Many exempt essential workers behaved according to very strict rules and protocols of care, and vice versa. Daredevils being a smaller and much more mobile fraction of the group, achieve (as a subgroup) true herd immunity in the months after the high pandemic peak. However, the frightened majority never reach high infection numbers, although a small fraction of them were infected by the mobile population. This small fraction of a large total implies that the total number of quarantined infected is similar to that of reckless, showing one of the remarkable characteristics of the COVID-19 pandemic: some transmit, others suffer. This is accentuated due to the higher fatality among older adults, with generally less risky behavior.

It is possible to establish a simple model based on a two-population SEIR, where the beta transmission parameter is broken down into four values $\beta_{11}, \beta_{12}, \beta_{21}, \beta_{22}$, corresponding to transmission between populations 1 (frightened and quarantined) and 2 (reckless and mobile). Figure 10 shows a schematic model for Spain in which both populations are described as homogeneous. From the IFR, deaths and seroprevalence, actual infection figures can be derived. The fraction of reckless individuals ($f_2$) is a parameter adjusted to the actual data as well as the reproduction numbers $R_1=\beta_{11}/\gamma$, $R_2=\beta_{22}/\gamma$ and the cross transmission parameter $\beta_{21}$.

Figure 10. Fright immunity pattern for Spain. Red line: Daily infections of the mobile population. Green line: Daily infections of the quarantined population. Black line: sum of infections. Squares: real data, corrected for seroprevalence factors and IFR. Population= 47x10^6, $R_1=0.67$, $R_2=3.57$, $f_2=0.0253$, $\beta_{21}=8.3x10^{-3}\beta_{12}=0$
This type of behavior should be considered as a cartoon, to the extent that there is undoubtedly a whole gradient of behaviors associated with more or less the extreme positions of the model. When the fitting procedure is applied to various countries or cities in Europe (United Kingdom, Germany, Belgium, France, Madrid, London) it can be seen that in all cases the fitting yields two well differentiated values of $R$ corresponding to a behavioral situation. On the contrary, when the data of the Autonomous City of Buenos Aires is treated with this same model, the adjusted $Rs$ are 1.6 and 1.13 respectively, both greater than 1 and close to each other, which shows that there aren’t two clearly differentiated populations as in European countries. This low level of “fright” is very compatible with the success in the objective sought to “flatten the curve” of infections. The gradual arrival of cases and deaths, avoiding collapse of the health system significantly reduced the perception of danger, and encouraged a behavior closer to the norm, only partially inhibited by restrictions on circulation and certain activities. The analysis must be differentiated with respect to popular/urban slum neighborhoods, where barriers to transmission are minimal and $R$ values are much higher, than the expected level for the natural epidemic curve of COVID. The strategies used to mitigate this situation were ineffective, reaching a cumulative prevalence of more than 50% in a few weeks (eg Barrio Carlos Mugica) (Figar et al. 2020). A very different situation occurs in the traditional neighborhoods of the City. As seen in the previous chapter, at the end of October the accumulated prevalence of CABA can already be estimated to be above 35%. However, the fraction of infected is very different for the various age ranges.

![Figure 11](attachment:image.png)

**Figure 11.** Estimated prevalence (as of October 16, 2020) for the different age ranges in the City of Buenos Aires.

Figure 11 shows that younger individuals have behaved more recklessly and reached important prevalence levels, although much more slowly than the reckless fraction in Spain and other European countries, whereas older adults have had much less contact with the virus, having understood the associated risk, particularly high at these ages. This low prevalence of the elderly further indicates that for the high number of deaths in the city of Buenos Aires and its suburbs, the number of real cases among young individuals has been several times higher than those that occurred in Spain and other European countries.

**Conclusions:**

A mathematical model is exhibited that could explain the reason for the decline in the contagion curve in European countries as of May, despite the low prevalence number. The causes of this behavior move from biological to social reasons. While in Europe there is a sharp peak with saturation of the health system, followed by an abrupt drop in cases as a result of the measures taken and the awareness of the population, the epidemic curve in the city of Buenos Aires and its suburbs is completely different; there is no generalized fear despite the high number of cases and deaths, since these appear gradually. While young people present a prevalence close to “herd” conditions, a sign of a relatively more mobile behavior during the isolation imposed by the government, which helps to lower the number of daily cases (surely added to the seasonal factor that may have accentuated the descent), the elderly have remained with little contact. The low prevalence among the age ranges most prone to severe COVID-19 represents an additional risk if transmission is not stopped at very low levels before weather conditions become more favorable for the spread of the virus.
2. Rapid response projects

2.1. Hospitals in networks: the case of Del Cruce Hospital in Florencio Varela, Province of Buenos Aires

Background:

Because of the advance of the pandemic, hospitals in the southeastern region of Greater Buenos Aires, which includes the municipalities of Almirante Brown, Quilmes, Florencio Varela and Berazategui, began to be restructured to work as a single functional unit (Figure 12). One of the objectives of forming this network was to organize and thus optimize the occupation of hospital beds in the context of the looming pandemic. The network has 7 hospitals, the central hospital of which is the Hospital del Cruce de Florencio Varela. During the pandemic, 3 Modular hospitals and 4 Early Care Units (ECU) were also built, some of which operate within the same hospitals.

Figure 12. Location of the hospitals in the southeastern region of Greater Buenos Aires, whose central hospital is the Del Cruce Hospital in Florencio Varela. The modular hospitals and ECUs that were built during the development of this project are included.

Objective:

Our aim was to build a software to organize patients and hospital beds for this hospital’s network which could be used by any other hospital network in the country. We also aim that the software offer the capacity to collect relevant information for future analysis.

Methodology:

Due to the urgency to start using the tool, we performed a first version (minimum viable product) in a few days, and then the development was expanded as more requirements were defined. Figure 13 shows the main screen of the system. The image corresponds to a panel with fictitious data.

Although the system responded to the requirements of this hospital network, it was developed so that any hospital network can use it (“Software as a service”). The system is located in a place accessible from any computer. Everything is customizable and self-manageable.
Figure 13. Main Panel of Hospital Beds. The hospitals in the network (7 hospitals plus 7 Modular / ECU). The hospital beds organized in Types are observed: General, Intermediate or Critical. The color of each bed corresponds to its Status: COVID-19, Suspect, Others, Free, Disabled. Each circle on the map corresponds to a hospital / Modular / UPA in the network, the area and the color of the circle correspond to the number of beds in that state. The panel allows users to download data. The image corresponds to a panel with fictitious data.

The system also allows entering patient data, which can be seen in the Patient Panel (Figure 14). This panel is used to organize the patients and beds of each unit. It should be clarified that the system does not collect sensitive information from patients such as name, ID, address, telephone, etc.
Conclusions:

The tool is widely used within the hospital network (Yacobitti et al., n.d.). It is making it possible to organize beds and patients within hospitals, as well as transfers between them. The project continues being improved to include new functionalities and requirements. The link to the system is available at (“Hospitales” n.d.).

2.2. COVID-19 Prevention Center at the Faculty of Exact and Natural Sciences (UBA)

Background:

One of the prevention programs developed by the Province of Buenos Aires is the identification and monitoring of close contacts (CC) of confirmed or suspected cases of COVID-19 to ensure that they do not continue to spread, and at the same time that they can receive the assistance they need.

On July 22 of this year, a University Telemedicine Center was put into operation in the Faculty of Exact and Natural Sciences of the University of Buenos Aires, in order to carry out the identification and monitoring of CCs. This prevention center provides support to the Municipality of Vicente López and the Municipality of Tigre. This Center is part of a program of the Province that involves another 34 university centers with the same dynamics (all of them created within national public universities).

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1 A CC (as defined by the PBA) is considered to be any person who has remained within a distance of less than 2 meters with a confirmed or suspected case of COVID-19, while exhibiting symptoms or within 48 hours prior to the onset of symptoms, for a period of at least 15 minutes.
Objective:

End the chain of infections by ensuring that individuals who have tested positive for COVID-19 as well as their close contacts remain isolated in their homes. Early detection of potential infected cases. Assist people who are confined during the days of their confinement.

Methodology:

The Center of Exactas is made up of a group of more than 40 students and alumni of our Faculty. It operates from Monday to Saturday in 4 fixed shifts of 4 hours, each of which is coordinated by a person responsible for daily tasks. All work is registered on a digital platform developed by the PBA. Both the physical facilities and the individual behaviors comply with the sanitary measures established by a strict Hygiene and Safety protocol provided by the academic unit of the institution and approved by the Province.

The daily work methodology at the center consists of:

Identification:

i. Telephone communication with the suspected or confirmed COVID-19 person.
ii. Determination of the date of symptom onset.
iii. Survey of close contacts
iv. Entry of CC information into the PBA application.

Accompaniment:

i. Telephone communication with the CC.
ii. Verification of the date of last contact with the epidemiological link.
iii. Ask about the appearance of one or more symptoms compatible with COVID-19.
iv. Survey on the needs for food assistance and/or hygiene supplies.
v. Classification of the evolution of the follow-up according to: continuation in isolation, discharged from follow-up or becoming a suspicious case due to the presence of symptoms.

In addition to the routine questions, the calls provide a space for advice and containment to people during the strict isolation they have to face, as well as an opportunity to relieve needs in the provision of food and hygiene supplies, among others. This information is duly compiled and transmitted daily to those responsible for the municipalities; who in turn use it as a source of information in their internal management.

Since the start of operation of the CeTeU and for the period of 16 weeks, 31070 effective calls have been made in total, of which 6560 correspond to suspected or confirmed people to identify their CCs (an average of 71 people per day), obtaining 3931 people with the CC classification (an average of 42 people per day). This shows us that approximately every 10 suspects contacted by the center identifies 6 CCs. Although the data are not conclusive, a possible explanation for this relationship is that a good part of the suspected cases are discarded because their swabs turned out to be negative. On the other hand, a total of 20,579 follow-up calls were accumulated, with an average of 224 calls per day for this task. Figure 15 shows the total per week of the suspects contacted along with the total per week of identified CCs, whereas Figure 16 shows the total per week of follow-up calls made.
Figure 15. Number of suspected/confirmed (red) cases contacted and close contacts (orange) identified per week.

Figure 16. Number of follow-ups with close contacts per week.

Although it is still premature to measure the real impact of the work carried out by the Center, there are certain positive indicators. A first indication can be seen in Figure 17, the confirmed cases in Vicente López show a stabilization towards the end of July (when the center began its activities), while the accumulated cases of the other 12 municipalities in same Sanitary Region (administrative organization that uses PBA to group municipalities) continued to grow steadily.
Figure 17. Evolution of daily confirmed cases over time for the municipality of Vicente López (in red) and for the group of other municipalities of Sanitary Region V (in green). Vertical line indicates start of contact tracing (Source: Antonio Milanese of Municipality of Vicente López).

On the other hand, the evolution of $R$ also shows a greater decrease in Vicente López than in the group of parties in the same Sanitary Region, as can be seen in Figure 18.
Figure 18. Evolution of the $R$ in Vicente López (bottom), and in the set of municipalities of Sanitary Region V to which Vicente López belongs (top). These indicators come from the parameter adjustment method based on simulation models described in Section 1.1 which is automatically calculated daily by the analysis, control and visualization system described in Section 1.4.
Other indications are of a qualitative nature, such as the strong awareness that is made through each call, either explaining the need for strict isolation of the CCs and the appropriate prevention measures that they must take. The social response to these calls is in the vast majority of cases positive and grateful. On the other hand, the Municipality of Vicente López has also expressed great appreciation for the work carried out in the center and the contribution of the information provided. In the case of Tigre, the beginning is very recent, but without a doubt, the participation of the Center allows the Municipality to identify and monitor a greater number of cases and, above all, earlier contact.

Conclusions:

In conclusion, it can be said that the experience of the University Telemedicine Center at the Faculty of Exact and Natural Sciences of the University of Buenos Aires is satisfactory and is generating an impact that may soon be even more quantifiable. Meanwhile, it is a useful tool in prevention tasks, being the first line of data acquisition used for management in the containment and mitigation of the disease.

2.3. Towards an agent-based simulation platform for the municipality of Marcos Paz, in the Province of Buenos Aires

Background:

The compartmental models described in Section 1.1 incur a level of abstraction that does not allow the study of detailed dynamics at the territorial level. The compartments, the reproductive number R, and the structure of the model itself are a macroscopic abstraction, representing the emergent behavior of networks of complex microscopic interactions between people of different ages who participate in very different ways in the epidemic process. These interaction networks are manifested in the territory according to the socioeconomic structure of each jurisdiction. At this level of detail, aspects such as the location of public buildings, shops and manufacturing plants, communication routes with other municipalities, location and capacity of hospitals, population density in residential areas and capacity dynamics towards central areas of greater activity, to name a few.

Analyzing social dynamics at this level is of particular interest to municipalities that must make decisions about regulations to manage the epidemic, which must contemplate the specific reality of the jurisdiction (a municipality supported by summer tourism activity, as in the case of Mar del Plata, will develop strategies different from a Mediterranean municipality focused on the food industry, as in the case of Marcos Paz).

Objective:

We therefore propose the development of a comprehensive platform based on simulation models that allows assisting decision-making with an explicit territorial level of detail, with a particular focus on the Municipality of Marcos Paz in the second cordon of the suburbs of the Province of Buenos Aires.

Methodology:

Spatially explicit simulation based on individual agents (Taillandier et al. 2019; Vynnycky and White 2010) allows to exercise hypothetical scenarios where interventions on reality take the form, for example, of restrictions on the number of vehicles that are allowed to cross the municipality from other jurisdictions, number of resident excepted personnel who are allowed to carry out working hours outside the municipality, quantity and size of public buildings with authorization for the capacity of citizens and their limitations on the number of people simultaneously sharing certain spaces, etc. The agents, in addition to forming family groups located according to the density of residence indicated by the censuses, run the epidemic through the susceptible, exposed, asymptomatic infected, symptomatic infected (possibly mild or severe), recovered, and deceased states.

Results:

We have developed a simulation model, together with its visualization infrastructure, for the Municipality of Marcos Paz (“Inicio - Municipio de Marcos Paz” n.d.), with the capacity to explicitly simulate the 65,000 inhabitants of the municipality and flows of travelers from/to other municipalities. Through close interaction with Marcos Paz’s municipal administration and its Ministry of Health, we carried out incremental phases of fitting of the model to capture the main socioeconomic dynamics that describe the mobility of people and the evolution of the metrics of the epidemic.

An example of a “snapshot” of activity simulated using our spatially explicit agent model is shown in Figure 19.
Figure 19. Agent-based simulation visualization for the Municipality of Marcos Paz in the province of Buenos Aires.

Conclusions:

This tool shows a great wealth for communication, interaction and discussion of possible results and scenarios with decision makers of public policies. Although it is in the initial stages of development, its structure opens the doors to provide a framework for future analysis of socio-economic problems that require explicit representation at the urban scale in the Municipality of Marcos Paz.

3. General conclusions and discussion

The pandemic has put the entire society on edge, mainly due to the rapid spread of the virus, influenced by the high mobility that exists due to commercial and tourist flows. That is why it took most nations by surprise, wreaking havoc on the global health and economic system.

The international scientific community reacted quickly by looking for palliative methods and vaccines, and developing models to be able to anticipate or at least keep up with the development of the pandemic. At the local level, the scientific community responded very quickly and proactively, generating solutions of all kinds, from detection systems, different types and mechanisms of tests, treatments and different proposals to contain contagions and to understand the behavior of the pandemic, to the development of the official application for prevention and public health care (Cuidar app (“Sistema Y Aplicación Cuidar” 2020)), in which scientists from our groups actively participated.

In this work we have presented the main contributions that a group of scientists from the Applied Math and Computer Science Institutes of the Faculty of Exact and Natural Sciences of the University of Buenos Aires and CONICET, together with colleagues from other institutions, in the fight against the pandemic in Argentina. Quantitative tools were developed that are helping to understand the state of the pandemic, as well as testing scenarios to evaluate different containment policies. Our work also included policies in the territory, such as the developments for hospitals and the tracking and contact identification centers (the latter an essential task but one that was not implemented on the scale that would have been necessary at the national level). We have also provided consulting and scientific-technological support to municipalities and provincial states, as well as organized a weekly seminar (29 to date) to highlight and debate these initiatives with experts of diverse disciplines (“Instituto de Cálculo” n.d.).

If we were to analyze Argentina’s behavior in response to the pandemic, we could say that the early quarantine dictated by the national government on March 20 allowed a slow growth of the contagion curve (see Figure 1), which in turn gave
rise to a strengthening of the health system. As a favorable result and unlike what happened in most European countries, the healthcare system was not saturated and potential deaths due to sanitary collapse were avoided. In contrast, this “flattening” of the curve, where the slow but permanent growth of infections is seen, led to the accumulated number of infected at the national level throughout the months being very high (see Section 1.5), similar to what has been observed in the number of deaths, turning Argentina into a country with a very high death toll per 100,000 inhabitants. These figures could have been reduced with a national strategy aimed at the suppression of the virus, based on the tracing of close contacts of confirmed and suspected cases, and the search for asymptomatic infected persons using intelligent testing techniques. Some of these actions had good local results (as can be seen, for example, in Section 2.2), but the strategy was not expanded nationally and in time to achieve a drastic decrease in infections.

We hope to be near the end of the pandemic, either because it is suppressed through efficient tracing and testing, because immunity is obtained through a vaccination campaign, because effective treatments are developed, or through a combination of these efforts; nevertheless, accurate information is essential to make the best decisions to control the spread of the virus. It is possible that these modeling techniques could also be useful for the organization of vaccination campaigns and drug delivery, as well as for dealing with other types of diseases with similar characteristics.

In summary, we consider that the contributions made by our working group have been very valuable in the fight against COVID-19 in Argentina. This is evidenced in the receptivity that the proposals had in different local and provincial governments, and in the strong impact that a good part of these developments had in the regional and national media.

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The main participation of each of the co-authors of the publication in the different sections of the work is detailed below.

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Section 2.2: Guillermo Durán, Natalia Fernández, Diego Garbervetsky, Rodrigo Goldsmit, Carolina Grillo Vidal, Mario Lozano, Mauricio Mendiluce y Sol Minoldo.

Section 2.3: Rodrigo Castro y Ezequiel Puerta.

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