The Use of Machine Learning to Predict Hospitalization of Covid-19: A Case Study in the State of Minas Gerais - Brazil

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Keywords: COVID-19, Hospitalization, Machine Learning.

Abstract: This work aims to verify the applicability of using Machine Learning techniques to predict hospitalization in confirmed cases of Covid-19. The study also intends to discover which attributes have the most significant impacts on hospitalization. The machine learning (ML) algorithms used in this experiment were Decision Tree, Random Forest, Neural Networks, and Naive Bayes. The data used for this experiment were made available by the government of Minas Gerais - Brazil, through open data. The model based on Random Forest obtained the best results, presenting the following metrics: Precision, Recall and F1-Score of 0.85, 0.84 and 0.84, respectively. In this experiment, essential characteristics for classifying the patient's hospitalization are Comorbidity, Age Group, and HDI. The results point to a good predictive ability, demonstrating the potential use of ML techniques to predict the hospitalization of people by COVID-19.

1 INTRODUCTION

The year 2020 was marked as a pandemic year. A Pandemic is a term that signals an epidemiological trend. Indicates that many outbreaks are happening simultaneously and spread far and wide. The Coronavirus Disease 2019 (COVID-19) has hit everyone indiscriminately. As a result of the pandemic, many people were infected, sick, hospitalized, and admitted to ICU, and many died.

Given the rapid transmission of the virus, there has been an abrupt and growing additional demand for hospitalizations worldwide, putting health systems under strain in many countries (Armstrong et al., 2020). Brazil has the Unified Health System (SUS)\textsuperscript{1}, which is the largest public and universal health system in the world, covering the entire country. About 75% of the Brazilian population does not have a private health plan and depends exclusively on SUS (Paim et al., 2011). In this context, the SUS was not prepared to meet the demand. As a result, the country went through a situation of the imminent collapse of its health system due to the lack of beds in Intensive Care Unit (ICU), equipment, essential supplies, and health professionals to ensure adequate care for patients in severe conditions affected by the virus (Santos et al., 2022b).

The great concern about contracting the disease revolves around the severity of the disease, which is variable, with some patients presenting few or no symptoms, some requiring hospitalization and hospitalization in the short, medium, or even long term, directly affecting public health. Given the above, the big question is how to predict whether an infected person will be hospitalized.

Machine Learning Models (ML) have been widely used in health. Simply put, ML is an area of Artificial Intelligence that attributes the ability to learn automatically to computational models. Thus, the main objective of its use in health is to use previous experiences to identify patterns and provide more assertive information for decision-making.

Given the scenario described and from the database of confirmed cases of COVID-19, made available by the government of Minas Gerais-Brazil, this work has as its primary objective to verify the
applicability of using ML techniques to predict the hospitalization of patients confirmed with Covid-19, as well as to present the most significant attributes in this prediction. The machine learning (ML) algorithms used in this experiment were Decision Tree, Random Forest, Neural Networks, and Naive Bayes. These algorithms were chosen because they are the most commonly used for data classification.

The remainder of this article is organized as follows: Section 2 brings the related works; the material and methods applied in the experiment are described in Section 3; the last sections get the analysis of the results, discuss and conclude the findings, indicating limitations of the work and proposing opportunities for further research.

2 RELATED WORKS

Jehi et al. (2020) proposed implementing a predictive algorithm to estimate, among COVID-19 patients, the risk of being admitted to a hospital with SARS-CoV-2 infection based on sex, age, COVID-19 symptoms, and comorbidities. Algorithm validation was evaluated internally using resampling bootstrap (1000 repetitions) to assess model discrimination and calibration. Each patient was stratified as a high, medium, or low-risk patient for the combined outcome of hospitalization and death, according to clinical and demographic characteristics highlighted by two predictive models developed by ATS Milan: the predictive model for hospitalization and the predictive model for overall mortality risk.

Gao et al. (2022) proposed MedML, a machine learning-based framework that aims to predict hospitalization and severity risk for pediatric patients with COVID-19 using electronic health records. It is a hybrid approach that combines medical knowledge graphs and data-driven feature extraction to predict better pediatric hospitalization and the severity of COVID-19 MedML extracts the most predictive attributes based on medical knowledge and propensity scores of more than 6 million medical concepts and incorporates the relationships between resources in graphs of medical knowledge through neural networks. As a result, MedML achieves an AUROC\(^2\) up to 7% higher and an AUPRC\(^3\) 14% higher compared to the best baseline machine learning models.

Murtas et al. (2021) aimed to develop a stratification tool to improve the management of patients with COVID-19 and the organization of health care.

To this end, a predictive algorithm was developed and applied to 36,834 patients with COVID-19 in Italy between March 8 and October 9, 2020, to predict their risk of hospitalization. The algorithm was developed following TRIPOD (Transparent reporting of a Multivariable Prediction Model for Individual Prognosis or diagnosis). Exposures considered were age, sex, comorbidities, and symptoms associated with COVID-19 (e.g., vomiting, cough, fever, diarrhea, myalgia, asthenia, headache, anosmia, ageusia, and dyspnea). As a result, the predictive model showed an excellent fit to predict hospitalization for COVID-19 and a good overall prediction accuracy score of Brier\(^4\) (0.14).

3 MATERIALS AND METHODS

In this section, we will present the method adopted to predict the hospitalization of patients confirmed by Covid-19 in a Brazilian state. The methodology involves selecting and extracting the database, preprocessing the dataset, and evaluating metrics. Figure 2 illustrates the steps of the methodology adopted in this work.

3.1 Obtaining the Database

The database used is available in the form of open data by the government of Minas Gerais - Brazil. According to the data dictionary, Table 1, the data

\[ BS = \frac{1}{n} \sum_{i=1}^{n} (pp_i - po_i)^2 \]

where \( n \) = the number of prediction-observation pairs; \( pp \) = predicted probability of occurrence of the event, and so is the observed probability of occurrence of the event (zero, the event did not occur, and one the event did happen).
extraction was performed from the official government systems SIVEP Grip (Severe Acute Respiratory Syndrome Notification System (SARS)), Esus-ve (tool for recording the notification of suspected cases of Covid-19) and REDCAP (RESEARCH ELECTRONIC DATA CAPTURE, a sophisticated platform for collecting, managing and disseminating research data). Data were updated until June 8, 2022, and extracted from the site on the same date.

The understanding of the base was carried out based on the available data dictionary, as well as a qualitative or quantitative analysis of the base. Two of the three available spreadsheets were used for this work. The first contains 500,001 records, and the second includes 88,101 records, thus totaling 588,102 records and 21 attributes.

3.2 Database Preprocessing

The pre-processing of the database can be defined as a set of activities that involve the preparation, organization, and structuring of the data. It is an essential step that precedes the accomplishment of the analyses and predictions since it will be decisive for the quality of the data analyzed. The following subsections describe each pre-processing step performed.

3.2.1 Selection of Attributes

Attribute selection performs a primary task and aims to select the more important attributes since non-relevant and/or redundant features can reduce the accuracy used by machine learning algorithms. For this work, the selection was made from understanding the data contained in Table 1. Of the 21 constant attributes, seven features and one class were selected. The selected attributes were: MICRO, DATE_NOTIFICATION, SEX, AGE, DATE_1st_SYMPTOM, RACE, COMORBIDITY, and the HOSPITALIZATION class.

3.2.2 Data Cleanup

The database may contain irrelevant or missing data. To deal with this situation, it is necessary to perform data cleaning. Both researchers and practitioners have recognized the need to detect errors, repair, and reason about querying and analyzing dirty and inconsistent data (Ilyas and Rekatsinas, 2022). In the case of the data presented, the instances containing missing data or “Not informed” conditions were removed, and the removal of inconsistent values were. A total of 112,917 instances were excluded in this process, so after cleaning the data, the base was left with 475,185 records.

3.2.3 Transformation, Normalization, and Discretization of Data

This step transforms the original data into more appropriate and suitable formats for machine learning algorithms. This is important, as some algorithms only work with numerical value entries, unable to predict others types of data entries. At this stage, the URS attribute was discarded, and the micro attribute was used because it had a more significant number of responses (89 different responses). This attribute MICRO (Microregion) has been transformed into two new attributes, namely: Population and HDI (Human Development Index) footnote The Human Development Index (HDI) is a general and synthetic measure used to classify the degree of economic development and the quality of life of countries. It was created in 1990 and published annually since 1993 by the United Nations Development Program – UNDP. The HDI varies on a scale from 0 to 1. The closer to 1, the greater the human development. This information was collected from data provided by the IBGE (Brazilian Institute of Geography and Statistics). For the HDI, there was no need to take any action since it is already normalized data.

The ‘POPULATION’ attribute was presented in absolute value according to each microregion. The same was discretized in 3 bands: small, medium, and large. For the small population, a value of fewer than 50 thousand inhabitants was considered; for the medium-sized population, values equal to or greater than 50,000 inhabitants and less than 100,000 inhabitants were considered; for the large population, a population similar to or greater than 100,000 inhabitants was considered.

The “GENDER” attribute was coded as Female =1 and Male =0.

The RACE attribute was binarized and transformed into five columns, white, brown, black, yellow, and indigenous, thus representing the presence or absence of the characteristic.

The AGE attribute was discretized as follows: Children = Under 12 years old, Adolescents = 12 to 18 years old6, Young = 19 to 29 years old7, Adults =

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6These transformations carried out for children and adolescents were created according to the ECA (Child and Adolescent Statute). The ECA was made through Federal Law no. 8,069, July 13, 1990, which regulates article 227 of the Federal Constitution of 1988.

7The transformation to “youth” is following the Youth Statute. This statute was created through Federal Law No. 12,852, of August 5, 2013.
Table 1: Data dictionary of the database provided by the government of a Minas Gerais - Brazil.

<table>
<thead>
<tr>
<th>ATTRIBUTE</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>ID</td>
<td>Unique exam code</td>
</tr>
<tr>
<td>URS</td>
<td>Name of the URS (Regional Health Unit) where the test was performed. There is the possibility of 28 different answers regarding the URS of the Brazilian State considered.</td>
</tr>
<tr>
<td>MICRO</td>
<td>Name of the Microregion in which the exam was performed. There is the possibility of 89 different answers regarding the Microregions of the Brazilian State considered.</td>
</tr>
<tr>
<td>MACRO</td>
<td>Name of the Macro-region where the exam was performed. There is the possibility of 14 different answers regarding the Macroregions of the Brazilian State considered.</td>
</tr>
<tr>
<td>NOTIFICATION_DATE</td>
<td>Date of notification of the patient’s Exam result</td>
</tr>
<tr>
<td>CLASSIFICATION_CASE</td>
<td>Result of the exam performed by the patient (if is a confirmed case for COVID-19)</td>
</tr>
<tr>
<td>GENDER</td>
<td>Gender of the patient who underwent the examination (male, female or not informed)</td>
</tr>
<tr>
<td>AGE</td>
<td>Age of the patient who took the exam</td>
</tr>
<tr>
<td>AGE GROUP</td>
<td>Patient’s age group</td>
</tr>
<tr>
<td>CITIES_RESIDENCE</td>
<td>Name of the patient’s municipality of residence. In this field, there is the possibility of 853 different answers.</td>
</tr>
<tr>
<td>CODE</td>
<td>IBGE code related to the patient’s municipality of residence. There is the possibility of 853 different answers.</td>
</tr>
<tr>
<td>EVOLUTION</td>
<td>Evolution of the patient’s condition</td>
</tr>
<tr>
<td>EVOLUTION_DATE</td>
<td>Date of evolution of the patient’s condition</td>
</tr>
<tr>
<td>DATE_1_SINTOM</td>
<td>Date of 1st symptoms of the case.</td>
</tr>
<tr>
<td>HOSPITALIZATION</td>
<td>Situation of hospitalization in the patient’s clinical bed</td>
</tr>
<tr>
<td>UTI</td>
<td>Situation of the patient’s ICU bed admission</td>
</tr>
<tr>
<td>RACE</td>
<td>Patient’s race</td>
</tr>
<tr>
<td>ETHNICITIES</td>
<td>Names of indigenous ethnicities. Name and ethnicity code of the patient, when indigenous</td>
</tr>
<tr>
<td>COMORBIDITY</td>
<td>Presence or absence of patient comorbidity</td>
</tr>
<tr>
<td>DATE_UPDATE</td>
<td>Data regarding update data Status</td>
</tr>
<tr>
<td>SOURCE_INFORMATION</td>
<td>Laboratory that performed the examination of the patient</td>
</tr>
</tbody>
</table>

For the attributes “NOTIFICATION_DATE” and “DATA_1o_SINTOM” two new columns were created, being the month and year.

The COMORBIDITY attribute has the answers “Yes”, “No”, and “Not informed” (considered missing data). Thus, this was binarized as follows: Yes = 1 and No = 0.

An attribute called “ILLNESS TIME” was created, which is the calculation of the difference between the attributes “NOTIFICATION_DATE” and “DATE_1o_SINTOM”. Negative values or values greater than 150 days were considered inconsistent and were removed from the database. This time was normalized using the Min-Max method.

According to Ciaburro et al. (2018), min-max normalization performs a linear transformation on the original data. This technique gets all data scaled in the range (0, 1), Equation 1.

\[
X_{\text{Scaled}} = \frac{X - X_{\text{min}}}{X_{\text{max}} - X_{\text{min}}} \tag{1}
\]

Also, according to Ciaburro et al. (2018), the min-max normalization preserves the relationships between the original data values. The cost of limiting this range is that we will end up with more minor standard deviations, which can suppress the effect of outliers.

Thus, after these pre-processing steps, the base presented 475,185 instances, 15 input attributes, and 1 Class attribute. The attributes are POPULATION, HDI, DATE_NOT_MONTH, DATE_NOT_YEAR, GENDER, AGE GROUP, DATE_1_SINT_MONTH, DATE_1_SINT_YEAR, ILLNESS TIME, COMORBIDITY, BROWN RACE, RACE WHITE, YELLOW RACE, BLACK RACE, INDIGENOUS RACE, and HOSPITALIZATION class.

3.2.4 Redundancy Elimination

Redundancy in machine learning preprocessing is the presence of attributes that are highly correlated with each other. Removing redundancy is helpful because it can reduce training time and increase model accuracy. Furthermore, eliminating redundancy can simplify the model, making it easier to interpret.

The `drop_duplicates()` function was used to exclude redundant data from the database, which eliminates equal instances from the database. In this way, the unique values in the base were kept; that is, there was no data loss. In this process, we went from 475,185 instances to 185,742; approximately 61% of the database corresponds to repeated cases.

3.2.5 Missing Data Imputation

After removing the redundancy, missing data (null) was evaluated. There are different methods of imputing missing data. In this case, the `KNNImputer` was used.

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8 According to the Elderly Person Statute, it was created by Federal Law No. 10,741 of October 1, 2003.
used, which assigns the missing values using the Euclidean distance. **KNNimputer** is a scikit-learn class used to fill in or predict missing values in a dataset. The $n_{	ext{neighbors}}=2$ (number of data points to be included closer to the missing value) was used as a parameter for this process.

### 3.3 Description of Methods

The study was conducted according to the workflow shown in Figure 2. To obtain the best predictive capacity, models based on four ML algorithms were used: Decision Tree, Random Forest, Neural Networks, and Naive Bayes. The models were implemented in Python using the scikit-learn learning library, developed explicitly for the practical application of MI. After some tests, we present the hyperparameters used in the experiment. The system's default hyperparameters were utilized for the Naive Bayes method.

The following hyperparameters were used for the Decision Tree: $\text{Criterion} = \text{entropy}$ (the criterion for measuring the quality of a node's split); $\text{Max\_depth} = 5$ (the maximum depth of the tree); $\text{Max\_features} = \text{auto}$ (the maximum number of features to be considered for the best split); $\text{Min\_samples\_leaf} = 7$ (the minimum number of samples needed to be in a leaf node); $\text{Min\_samples\_split} = 6$ (the minimum number of samples required to split a node); $\text{Splitter} = \text{random}$ (the criterion used to choose the best division).

For Neural Networks, the following hyperparameters were used: $\text{Activation} = \text{relu}$ (Activation function for hidden layer); $\text{Alpha} = 0.001$ (Regularization Term); $\text{Hidden\_layer\_sizes} = 20$ (Number of neurons in the hidden layer); $\text{Learning\_rate\_init} = 0.001$ (Initial learning rate); $\text{Max\_iter} = 700$ (Maximum number of iterations); $\text{Solver} = \text{adam}$ (algorithm for optimization).

For Random Forest, the following hyperparameters were used: $n_{\text{estimators}}=100$ (the number of trees); $\text{Criterion} = \text{entropy}$, (the criterion used to measure the quality of a node's split); $\text{Max\_depth} = 10$ (the maximum depth of the tree); $\text{Max\_features} = 10$ (the number of features to consider when looking for the best split); $n_{\text{jobs}}=5$ (the number of cores to be used to build the trees).

### 3.4 Model Quality Assessment Metrics

The following metrics will be used to measure the quality of the models: $\text{Recall}^9$, $\text{Precision}^{10}$, and $F$-measure$^{11}$.

For the evaluation of the quality of the learning models, 10% of the data was reserved for testing. In the remaining 90% of the data, 10-fold cross-validation was applied. Figure 2 presents a summary of the entire process performed.

### 4 RESULTS AND DISCUSSIONS

The coronavirus is one of the pathogens that act on the human respiratory system, causing severe acute respiratory syndrome (SARS) and Middle East respiratory syndrome (MERS), and it is considered a pathogen of great importance to public health (Rothan and Byrareddy, 2020). Since the first case of SARS-CoV-2 infection, modeling the epidemic growth pattern has been crucial to understanding the pandemic’s evolution and guiding the implementation of prevention and control measures (Sciannameo et al., 2022).

The outbreak caused by the coronavirus was declared in Brazil a pandemic, on March 11, 2020, by the World Health Organization (WHO) and, thus, brought significant challenges to public health. In Brazil, the first case of COVID-19 was confirmed on February 26, and the first death on March 17. Furthermore, the Brazilian Ministry of Health recognized national community transmission on March 20, 2020 (Santos et al., 2022a).

Regarding clinical manifestations, the Epidemiological Surveillance Guide of the Brazilian Ministry of Health (da Sáude do Brasil, 2021) clarifies that SARS-CoV-2 infection can range from asymptomatic cases and mild clinical manifestations to moderate, severe, and critical conditions. It also highlights that although most people with covid-19 develop mild (40%) or medium (40%) symptoms, approximately 15% may develop severe symptoms that require oxygen support, and about 5% may have the necessary form of the disease. In this situation, the individual may experience complications such as respiratory failure, sepsis and septic shock, thromboembolism, and multiple organ failure, including acute liver or heart damage, requiring intensive care. These same rates are presented by the World Health Organization (Organization et al., 2020).

According to the Hospital Policy and Management Observatory (OPGH) (Albuquerque, 2021), in 2020, there was an increase of almost 55% compared to 2019 in hospitalization for treatment of infectious and parasitic diseases (IPD) in the SUS and was the first cause of hospitalization in 2020.
At different times, the COVID-19 pandemic in Brazil led the health system to exhaust its installed capacity. Thus, it became necessary to capture the extent of the damage and build knowledge to subsidize care reorganization choices to deal with unmet routine needs and the new demands imposed by Covid. So, it is essential to trace the pattern of use in the Unified Health System (SUS) hospital network during this pandemic (Portela et al., 2021). Analyzing this pattern will enable more assertive decision-making regarding the patient’s hospitalization.

For the analyzed dataset, we executed the models of the five proposed ML methods (Decision Tree, Random Forest, Neural Networks, and Naive Bayes) to understand the profile of people hospitalized for COVID-19. Table 2 shows the demographic data of the study.

### Table 2: Demographic questions.

<table>
<thead>
<tr>
<th>Question</th>
<th>N</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>93,042</td>
<td>50.09</td>
</tr>
<tr>
<td>Female</td>
<td>92,700</td>
<td>49.91</td>
</tr>
<tr>
<td>Age group</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12 less</td>
<td>10,467</td>
<td>5.64</td>
</tr>
<tr>
<td>12 to 18</td>
<td>9,961</td>
<td>5.36</td>
</tr>
<tr>
<td>19 to 29</td>
<td>27,656</td>
<td>14.89</td>
</tr>
<tr>
<td>30 to 59</td>
<td>74,121</td>
<td>39.91</td>
</tr>
<tr>
<td>60 or more</td>
<td>63,537</td>
<td>34.21</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>80,335</td>
<td>43.25</td>
</tr>
<tr>
<td>Brown</td>
<td>76,385</td>
<td>41.12</td>
</tr>
<tr>
<td>Black</td>
<td>20,985</td>
<td>11.30</td>
</tr>
<tr>
<td>Yellow</td>
<td>7,725</td>
<td>4.16</td>
</tr>
<tr>
<td>Indigenous</td>
<td>312</td>
<td>0.17</td>
</tr>
<tr>
<td>Population</td>
<td></td>
<td></td>
</tr>
<tr>
<td>50k less</td>
<td>31,879</td>
<td>17.16</td>
</tr>
<tr>
<td>50k to 99,999</td>
<td>52,052</td>
<td>28.02</td>
</tr>
<tr>
<td>100k or more</td>
<td>101,811</td>
<td>54.81</td>
</tr>
</tbody>
</table>

Observing Figure 3, which presents the results of the testing phase of the prediction models, the model based on Random Forest obtained the best results, considering that the model has an excellent predictive capacity for this dataset. Thus, we used the Random Forest method to discuss the results better.

After choosing the model, it is necessary to understand the quality of its results and make the model results more explainable. To this end, the importance of each attribute selected by the model for the prediction was verified, where the most critical feature was comorbidity, followed by age group.

The importance of each attribute was calculated using Random Forest’s feature_importances method. This method is applied to Random Forest models and returns a list containing the importance of each feature for the model. The importance of attributes is calculated according to the average impurity reduction when a decision vertex is divided on an element.

The importance of the attributes was displayed through a bar graph (Figure 4), where the length of the rectangle represents the weight of the feature.

This graph represents the weight of each attribute for the classification of a patient’s hospitalization. Thus, in this experiment, essential characteristics for classifying the patient’s hospitalization are: Comor-
Feitoza et al. (2020) state that about 20% to 51% of patients with COVID-19 were detected with a chronic disease already installed. Ranzani et al. (2021) shows in its study that a large proportion, 74%, of hospitalized patients with COVID-19 in Brazil had one or two comorbidities. Regarding the Age Group, it can be seen in Figure 5, taken from the database, that the elderly and adults have a greater tendency to be hospitalized by Covid-19 than children, adolescents, and young people.

Lana et al. (2021) presented in their study the over-risk calculation and the incidence rate of hospitalization and deaths per thousand inhabitants for Covid-19 by gender, age group, and comorbidity. The study shows that from the age group of 45-49 years ($\text{SRfe}^{12} = 1.133$), there is a gradual increase in the risk of hospitalization, reaching $\text{SRfe} = 8.466$ among those aged 90 years or older, which demonstrates that age is a strong and well-established risk factor for hospitalization and death from COVID-19.

Regarding the HDI Maciel et al. (2020) analyzed the spatial distribution of the incidence of Covid-19 and its correlation with the municipal human development index (MHDI) of municipalities in a Brazilian state. They concluded that the incidence of COVID-19 was heterogeneous and unequally distributed for the Brazilian state studied and that the cities with the highest coefficients of incidence of COVID-19 had higher MHDI values. Thus, HDI can be presented as a relevant factor for the patient with Covid-19.

The graph represented by Figure 6, drawn from the data of this study, collaborates with the findings above and shows the increase in hospitalization and the rise to the increase in HDI.

The analysis showed a positive correlation between the incidence of hospitalization for Covid-19 and the HDI; that is, increases in the HDI also increase the incidence of hospitalization for Covid-19.

5 FINAL CONSIDERATIONS

This work demonstrates the potential use of ML techniques to predict hospitalizations for COVID-19. The results presented in this study are considered promising since the ML models performed well in classifying patients. In addition, ML models showed results on time, which is extremely important regarding response time. Therefore, ML models can be used as additional tools in decision-making regarding patients hospitalized for COVID-19.

Implementing an ML model can provide a more accurate assessment of the hospitalization of a patient with COVID-19 and assist the health agent in decision-making. If systematically implemented, and with the breadth of clinical data collected, it is believed that better identification and care of patients infected by COVID-19 will be possible.

As a suggestion for future work, the ML models presented in this study can be improved with other ML hyperparameters and/or methods, such as the Deep Neural Network, which gives a better representation of data and, therefore, can help to improve the performance of the model. In addition, other open datasets can also be analyzed to understand better the profile of patients hospitalized for COVID-19.

5.1 Study Limitation

Knowing that the clinical spectrum of COVID-19 is vast, the a priori clinical knowledge about the associations between symptoms and hospitalization for COVID-19 was limited for this study since the field “Comorbidity” presented only the yes or no values. Thus, we can point out that the major limitation of this research is the lack of data for an evaluation of other important variables in this predictive process, such as the patient’s clinical condition, diabetes, high

\[ \text{SRfe}^{12} \]

The stratified over-risk is defined for group i with the denominator of the incidence rate for the age group (fe) in question, that is, $\text{SR}_{i,fe} = \frac{\Theta_{i,fe}}{\Theta_{0,fe}}$, where $\Theta_{i,fe}$ and $\Theta_{0,fe}$ represent the incidence rate of group i and the general population for the age group of interest. The $\text{SRE}_{i,fe}$ distribution for group i has derived analogously to the over-risk distribution for group i.
blood pressure, chronic kidney disease, fever, cough, shortness of breath, fatigue, lung disease, among others.

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