Disentangling the Association of Hydroxychloroquine Treatment with Mortality in Covid-19 Hospitalized Patients through Hierarchical Clustering

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1. Introduction

Hydroxychloroquine (HCQ) is an antimalarial drug suggested to be effective in inhibiting Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) replication in vitro [1, 2]. Indeed, HCQ is characterized by antiviral, anti-inflammatory, and antithrombotic actions, contrasting the main disruptive effects of SARS-CoV-2 infection on the organism [3]. For this reason, it has been heavily used in treating patients affected by SARS-CoV-2 infection related disease (commonly known as Covid-19), especially in the first phases of the current pandemics, when Covid-19 was quite unknown [4].

Despite these elements and initial suggestive evidence of efficacy based on daily clinical practice, in the last months, the potential benefit of HCQ for Covid-19 patients has been harshly debated [3, 5]. In particular, evidence supporting protective effects from observational studies [6–13] was in contrast with that suggesting no effect at all by recent randomized clinical trials (RCTs) [14–18]. More recently, a meta-analysis combining both RCTs and observational studies over more than 44,000 patients supported a protective effect of HCQ, driven by the findings of observational studies [5]. A potential explanation for this discrepancy may be due to the usually high dosage administered in RCTs (800 mg/day), compared to lower dosages reported in observational studies supporting HCQ efficacy (≤400 mg/day), as hypothesized elsewhere [5]. As an alternative explanation, it is likely that the efficacy of HCQ treatment for Covid-19 may vary across patients and is influenced by subtypes of the disease, which in turn is largely dependent on patients’ characteristics and their nonlinear combinations [19]. In this “personalized medicine” view, response to HCQ may not be the same across all patients of the same age, or with similar circulating inflammation levels. In order to identify these combinations, the use of big data like Electronic Health Records (EHRs) and of machine learning (ML) algorithms to interpret hidden HCQ response patterns is of fundamental importance. ML is an umbrella term covering different algorithms designed for the identification of hidden patterns, information mining, and variable classification/estimation through modelling complex (including nonlinear) functions, usually adopted in a big data setting. These algorithms can be generally classified into supervised and unsupervised approaches. The former are designed to learn to predict specific outcomes after proper training of the algorithm in independent datasets, trying to model relationships and dependencies between the input variables (or features) and the target prediction output (or label). In unsupervised algorithms, the machine simply learns to identify hidden patterns across a high number of observations over many features, without the need for labels, and is more descriptive—rather than predictive—in nature. These algorithms have shown promising findings in public health, in the management of both chronic [20] and acute health conditions [21, 22], but also during the current pandemics. In particular, useful ML applications have been reported in the prediction of Covid-19 diagnosis and prognosis [23]. Notwithstanding this, to the best of our knowledge, only one study attempted so far to predict response to HCQ treatment within Covid-19 patients, through the application of a supervised ML technique (gradient boosting). Interestingly, the authors reported a reduction of in-hospital mortality within patients treated with HCQ, which was even more
pronounced within those patients predicted to benefit most from the drug, in line with expectations [19].

Here, we attempted a personalized Covid-19 patient characterization to better disentangle the beneficial effects of HCQ previously reported within the COVID-19 RISK and Treatments (CORIST) study, a large retrospective cohort of patients hospitalized for SARS-CoV-2 infection in Italy [13]. This approach consisted of i) identifying the existence of subtypes of Covid-19 patients through an unsupervised ML algorithm—hierarchical clustering—comparing their characteristics and their clinical risks, ii) testing the resulting patients’ clusters for association with mortality and modification of effect by treatment with HCQ, and iii) analyzing potential interactions between clusters and HCQ use. This approach represents a prominent example of how personalized medicine may support clinicians in Covid-19 treatment.

2. Methods

2.1. Analyzed Cohort. The COVID-19 RISK and Treatments (CORIST) study includes 4,396 patients hospitalized for SARS-CoV-2 infection in 35 hospitals across Italy, between February 2020 and May 2020. Molecular diagnosis of SARS-CoV-2 infection was based on polymerase chain reaction (PCR) of viral DNA extracted and amplified from nasopharyngeal swabs. Within each participating hospital, clinical data were abstracted at one-time point from electronic medical records or charts and collected using either a centrally designed electronic worksheet or a centralized web-based database. Collected data included patients’ demographics, laboratory tests, medications in use, history of disease, and prescribed pharmacological therapy for Covid-19 treatment. For each participant, the study index date was defined as the date of hospital admission, while the study end point was death. Follow-up time was computed as the time between the index date and death, or alternatively between the index date and the date of discharge, applying right-censoring. Further details on the study are reported elsewhere [13, 24, 25].

2.2. Statistical Analyses

2.2.1. Cluster Analysis. All analyses were carried out in R 4.0.2 (https://www.r-project.org/) [26]. We applied a hierarchical clustering analysis on Covid-19 patients using their main clinical, lifestyles, and sociodemographic characteristics, which were suggested as the most influential on mortality risk by previous studies in the field [25, 27, 28]. These included age (years), sex, glomerular filtration rate (eGFR, mL/min/1.73 m²) and high-sensitivity plasma C-reactive protein levels (mg/L) at in-hospital admission, obesity (body mass index (BMI) ≥ 30 kg/m²), hypertension (Yes/No), and smoking status (never/previous/current smoker), as well as history of myocardial infarction, heart failure, chronic pulmonary disease, cancer, and diabetes (Yes/No). Missing data were imputed through a k-Nearest Neighbor approach, implemented in the kml() function of the VIM package (version 6.1.0; https://cran.r-project.org/web/packages/VIM/index.html), with k = 10 [29]. CRP was transformed on the natural logarithm scale to reduce skewness, and all the continuous variables were normalized through the normalize() function in Keras v2.3.0 (https://cran.r-project.org/web/packages/keras/index.html).

Cluster analyses were then performed on the 4,396 patients, using the variables specified above, through the Cluster package v2.1.0 (https://cran.r-project.org/web/packages/cluster/index.html). First, we computed a dissimilarity matrix based on Gower pairwise distance (Figure S1), through the daisy() function. Gower distance is a parameter in the [0; 1] range representing the average of pairwise dissimilarities across individuals (the higher the distance, the more the dissimilarities for a given pair of subjects) [30]. Second, we performed hierarchical clustering through the hclust() function applied to the Gower distance matrix computed above, which separated subjects based on their degree of pairwise dissimilarity, both from lowest to highest (agglomerative clustering) and from highest to lowest (divisive clustering). Third, we determined the appropriate number of clusters for patient classification, based on the Average Silhouette method (Figure S2). This computes the number of clusters, which maximizes the average silhouette width, a measure of the quality of clustering indicating how well each object lies within its cluster [31]. This method, applied through the fviz_nbclust() function of the Factoextra package (v1.0.7, https://cran.r-project.org/web/packages/factoextra/index.html), computed k = 2 as the optimal number of clusters. Finally, each patient was assigned to one of the two clusters determined above, through application of the cutree() function (Cluster package) to the results of the cluster analysis previously carried out.

2.2.2. Comparison of Clusters. First, we compared the classifications made through agglomerative and hierarchical clustering, which revealed high consistency (Odds Ratio = 34.0 [26.7-43.7], Fisher Exact Test p value < 10^-15). In light of this homogeneity of classification, and since divisive clustering has been reported to be more accurate and robust [32], all the subsequent analyses were performed on cluster classification identified through the latter approach.

The two clusters of patients identified were then compared for all anamnestic variables mentioned above, through Fisher’s Exact Test (for binary variables), Chi-squared test (for nonbinary categorical variables), and through Student’s t-test or Wilcoxon Rank Sum tests (for continuous variables meeting and not meeting parametric assumptions, respectively). Similarly, we compared Covid-19 disease severity, classified by recruiting centers in asymptomatic/mild, nonsevere pneumonia, severe pneumonia, and acute respiratory distress syndrome (ARDS). Moreover, we compared the use of six common drugs for Covid-19 treatment between the two clusters, including hydroxychloroquine, antihypertensive drugs, anti-interleukin-6 antibody, antivirals (Remdesivir, Lopinavir, Darunavir), and corticosteroids. These were reported as binary variables (Yes/No) and were, therefore, compared with clusters through Fisher Exact Tests.
2.2.3. Survival Analyses. Once the clusters were characterized, we modelled incident mortality risk as a function of patients’ clusters and use of HCQ through Cox Proportional Hazards (PH) models, using the \texttt{cox.zph()} function of the survival package (v3.2.7, see URLs: https://cran.r-project.org/web/packages/survival/index.html) [33]. Only patients with complete information needed in each model were included in the analysis (case-complete approach; see below). A preliminary check of the basic Cox PH assumptions revealed no influential observations based on dfbeta residuals (Figure S3a), while Schoenfeld residuals tests revealed a statistical violation of the proportionality of hazards assumption, although these did not show any evident trend at a visual inspection (Figure S3b). For this reason, we carried out Cox PH models both with and without including an interaction term with time-to-event, a strategy commonly used to overcome this violation [34]. Incrementally adjusted models were analyzed: i) a crude model including only patients’ clusters (Model 1; \(N = 4,319\)); ii) a model testing additive influence of clusters and HCQ use (Model 2: Model 1 + HCQ; \(N = 4,212\)); and iii) a model testing both additive and synergistic influence of clusters and HCQ use (Model 3: Model 2 + clusters*HCQ; \(N = 4,212\)). Additional sensitivity analyses were carried out to rule out potential confounding effects of additional drugs in use for Covid-19 treatment (Model 4: Model 3 + other drugs). Risk estimates were computed as hazard ratios (HR) with 95% confidence intervals (95% CI) of dying, and HR with \(p\) values below \(a = 0.05\) were considered significant. To quantify the potential for unmeasured confounding effects, the E-value was calculated for all the HRs observed in Model 3, as described in [35] (https://www.evalue-calculator.com/). This represents the minimum association required for a potential unmeasured confounder with both the exposure and the outcome to explain away the observed association. In other words, the higher the E-value, the harder it is to attribute an association to an unmeasured covariate [36].

2.2.4. Associations with Additional Endpoints. To better evaluate the associations of Covid-19 patients and HCQ use with negative outcomes other than death, we built a composite endpoint based on the occurrence of at least one of the following outcomes: in-hospital death, access to intensive care unit during hospitalization, or severe disease manifestation (either severe pneumonia or ARDS). In this case, the resulting binary variable was assigned a value of 1. Conversely, the variable got 0 “value if one of the following alternative conditions is applied: (i) none of the above mentioned outcomes was verified; (ii) a patient survived without recurring to intensive cares or (iii) without showing severe manifestations of the disease. Six patients with missing values on survival were removed. Then, we modelled the risk of manifesting a bad outcome through a logistic regression (\texttt{glm()} function in R), modelling both additive and interactive models of Covid-19 patients’ cluster and HCQ use, as above. This analysis was motivated by the fact that the curse of disease often differs across patients, e.g., with some subjects with less severe forms suddenly worsening their conditions until death and others having severe manifestations but still surviving, possibly thanks to intensive cares. Therefore, a composite outcome variable represented a robust way to measure potential risk/proactive effects of patients’ clusters and HCQ use.

3. Results

While both agglomerative and divisive clustering approaches were developed, all statistical analyses presented below are based on clusters identified through the same approach, since this showed a high homogeneity with the results of agglomerative clustering (see Methods section), and divisive clustering has been reported to be more accurate and robust [32, 34].

3.1. Characteristics of the Clusters. We identified two clusters of Covid-19 patients (with \(N = 3,913\) and 483, Figure 1). A comparison of the continuous variables used for their determination is reported in Figures 2(a)–2(d). The larger cluster was younger (mean (SD) age: 65.2 (15.6) vs 77.9 (9.2) years; \(t\)-test = -25.9, \(p < 10^{-15}\)), with better kidney function (eGFR: 77.9 (26.9) vs 52.6 (25.6) mL/min/1.73 m\(^2\); \(t\)-test = 20.3, \(p < 10^{-15}\)) and lower circulating inflammation levels (CRP: 34.6 (62.4) vs 36.5 (58.6) mg/L; Wilcoxon test = 847,660, \(p = 2.5 \times 10^{-14}\)). Conversely, BMI did not show strong differences between the two clusters (28.0 (4.2) vs 27.5 (4.2) Kg/m\(^2\); \(t\)-test = 2.3, \(p = 0.02\)). Moreover, patients belonging to the larger cluster were less frequently men (60% vs 75%) and smokers (11.5% vs 19.5%) and showed a lower prevalence of chronic health conditions like myocardial infarction, heart failure, diabetes, hypertension, cancer, and lung disease (all \(p < 0.0001\)), while no significant difference was observed in the prevalence of obesity (Table 1).

Clusters were also associated with severe Covid-19 disease manifestations, with 65.8% of patients in the smaller cluster presenting with either severe pneumonia or ARDS, compared to 45.9% in the larger cluster (Chi-squared = 76.4, \(p < 10^{-15}\); Table S1). For the characteristics mentioned above, the large and small clusters will be hereafter named as “low risk” and “high risk” cluster. When we compared the use of specific drugs, in the high risk cluster, we observed a less frequent use of HCQ (\(p < 0.001\)) and of Lopinavir/ Darunavir (\(p < 0.05\)) and a more frequent use of corticosteroids and antihypertensive medications (\(p < 0.001\), compared to the low risk cluster (Table S2).

3.2. Combined Influence of Clusters and HCQ Use on Mortality Risk. In Cox PH regressions modelling mortality risk as a function of clusters (Model 1), we analyzed 4,319 patients with a case-complete approach, with a total of 799 deaths and a total of 73,924 person-days follow-up (median 13 days). In this model, patients belonging to the high risk cluster showed a significant increase of incident mortality risk, compared to those of the low risk cluster (HR [CI] = 3.81 [3.12–4.65]; Table 2). This association remained stable in a Cox regression modelling additive effects of clusters and HCQ use (Model 2:
$N = 4,212,743$ events, a total of 72,239 person-days follow-up, median 14 days). Indeed, the high risk cluster was associated with a significant increase of mortality (3.80 [3.08-4.67]), while HCQ use was associated with a significant independent reduction (0.51 [0.43-0.61]). When we modelled additive and interactive associations of clusters and HCQ in a single model.
users, both in the additive and in the interactive model with increased risk for cluster 2 and decreased risk for HCQ use, we observed results inline with survival analyses, as a function of clusters and disease—i.e., severe Covid-19 manifestations, access to intensive care unit or death—as a function of clusters and potential unmeasured confounding effects, with E-values of 3.1, 2.8, and 2.5 for the associations of mortality with the risk cluster, HCQ use, and cluster*HCQ interaction in Model 3. Moreover, these associations remained substantially stable in Cox PH models including additional drugs in use (Table 2, Model 4), as well as in those including an interaction term with time (Table S3).

### 3.3. Associations with a Combined Covid-19 Outcome

When we modelled the risk of bad clinical outcomes of the disease—i.e., severe Covid-19 manifestations, access to intensive care unit or death—as a function of clusters and HCQ use, we observed results in line with survival analyses, with increased risk for cluster 2 and decreased risk for HCQ users, both in the additive and in the interactive model (Table 3). While the cluster*HCQ interaction showed only a trend of significance (p = 0.08), HCQ still presented a significant protective association in the low risk cluster (OR [CI] = 0.67 [0.56-0.79], p = 4.0 × 10^{-6}) and with a milder nonsignificant reduction in the high risk cluster (OR [CI] = 0.98 [0.66-1.46], p = 0.92).

### 4. Discussion

In the present work, we report differential influence of HCQ treatment on Covid-19 mortality through a hierarchical clustering analysis applied to patients hospitalized for SARS-CoV-2 infection. This revealed the existence of two separate clusters of Covid-19 patients, based on their clinical and sociodemographic characteristics: one of younger patients with less comorbidities, lower circulating inflammation, and better renal function (“low risk” cluster), and one of older and more comorbid patients, more prevalently men and smokers (“high risk” cluster). The former cluster showed a higher prevalence of severe manifestations of Covid-19, ranging from severe pneumonia to ARDS. Moreover, survival analyses showed an almost four-fold increase of in-hospital mortality for the high risk compared to the low risk cluster. Although a previous study attempted to identify subtypes of Covid-19 patients, associating them with disease severity [37], this represents the first attempt to use clustering in disentangling the effect of HCQ on different

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**Table 1: Comparison of main categorical variables between the two clusters identified.**

| Category (%) | Cluster 1 – low risk N = 3,913 | Cluster 2 – high risk N = 483 | p for difference |
|--------------|-------------------------------|-------------------------------|-----------------|
| Men | 2,346 (60.0%) | 362 (74.9%) | 6 × 10^{-11} |
| Smoke | | | |
| Current smokers | 450 (11.5%) | 94 (19.5%) | <10^{-15} |
| Previous smokers | 268 (6.8%) | 94 (25.3%) | 0.73 |
| Obesity (BMI ≥ 30 Kg/m²) | 546 (13.9%) | 64 (13.3%) | <10^{-15} |
| Myocardial infarction | 127 (3.2%) | 335 (69.4%) | <10^{-15} |
| Heart failure | 171 (4.4%) | 315 (65.2%) | <10^{-15} |
| Diabetes | 621 (15.9%) | 276 (57.1%) | <10^{-15} |
| Hypertension | 1,828 (46.7%) | 453 (93.8%) | <10^{-15} |
| Cancer | 392 (10.0%) | 89 (18.4%) | 2 × 10^{-07} |
| Lung disease | 415 (10.6%) | 207 (42.8%) | <10^{-15} |

P for difference resulting from comparison of the clusters—through Fisher’s Exact Test (for binary variables) or Chi-squared test (for nonbinary categorical variables, i.e., smoke)—are reported, along with absolute and % frequency of each condition within each cluster.

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**Table 2: Results of Cox PH regressions modelling incident mortality risk.**

| Model | N (deaths) | Cluster 2 vs 1 | HCQ Yes vs no | Cluster*HCQ |
|-------|------------|---------------|---------------|-------------|
| Model 1: Death ~ cluster | 4,319 (799) | 3.81 [3.12-4.65] (<10^{-15}) | | |
| Model 2: Death ~ cluster + HCQ | 4,212 (743) | 3.80 [3.08-4.67] (<10^{-15}) | 0.51 [0.43-0.61] (8.8 × 10^{-15}) | |
| Model 3: Death ~ cluster + HCQ + Cluster*HCQ | 4,212 (743) | 2.45 [1.69-3.54] (4.9 × 10^{-4}) | 0.46 [0.38-0.55] (-10^{-15}) | 1.90 [1.21-2.96] (2.1 × 10^{-4}) |
| Model 4: Death ~ cluster + HCQ + Cluster*HCQ + other drugs | 3,736 (664) | 1.65 [1.20-2.26] (2.2 × 10^{-3}) | 0.52 [0.43-0.63] (2.0 × 10^{-11}) | 1.98 [1.36-2.89] (4.0 × 10^{-4}) |

Associations between incident mortality risk, Covid-19 clusters identified, and use of Hydroxychloroquine (HCQ) were tested in the incremental models and in a sensitivity analysis including all the drugs used for Covid-19 treatment. No other covariates were included in the analysis. Hazard Ratios with 95% confidence intervals (HR [CI]) and relevant p-values (in brackets) are reported. Significant HRs (p < 0.05) are highlighted in bold.
types of patients, by testing associations with incident inhospital mortality risk. Specifically, we tested and observed both additive and interactive associations of HCQ and Covid-19 subtypes. Indeed, the high risk cluster was consistently associated with increased mortality across all models, while treatment with HCQ was generally associated with a halving of death risk, in line with previous evidence from both observational [6–13] and experimental studies [19]. While we already reported evidence suggesting a protective influence of HCQ against mortality in a largely overlapping sample [13], here, we have further deepened this relationship by testing and reporting a significant association between cluster-by-HCQ interaction and mortality, which was driven by a differential association within the two clusters. Indeed, the low risk cluster showed a significant “protective” influence of HCQ on in-hospital deaths, while the high risk cluster showed a concordant but nonsignificant association. This represents an element of novelty of the present study, since, in our previous work, we observed a “protective” association between HCQ and mortality within the totality of patients (about 75% of the current sample size), and when stratifying by age, sex, and other characteristics [13], but not within different subtypes of patients combining all these characteristics together in a nonlinear setting, as can be built through unsupervised ML algorithms. Moreover, here, our evidence is supported also by concordant associations with a composite and possibly more robust outcome of the disease, based on the occurrence of death, access to ICU, and severity of manifestations.

Recently, an approach based on the definition of subtypes of Covid-19 patients has been already proven to be successful in identifying which patients benefit most from HCQ treatment, in a multicenter trial involving six US hospitals and 290 patients hospitalized for Covid-19, the IDENTIFY study [19]. HCQ treatment was associated with higher survival in the treated harm, and especially within those patients that were predicted to benefit most based on a supervised ML algorithm applied to their characteristics, which included blood pressure, heart rate, temperature, respiratory rate, oxygen saturation, white blood cell and platelet count, lactate, blood urea nitrogen, creatinine, and bilirubin levels [19]. Interestingly, lactate and creatinine levels were the most important features in this algorithm [19], the latter representing an index of renal function, which was also a characteristic feature of the low risk cluster in the present study, where HCQ was more effective. Moreover, patients eligible for HCQ treatment as derived by the algorithm of [19] were shown to be younger and less comorbid than the whole population studied, in line with the evidence reported in the present work, suggesting that HCQ treatment may be more effective for younger patients with better general health conditions.

Table 3: Results of logistic regressions modelling Covid-19 composite bad outcome risk.

| Model                                  | N   | Cluster 2 vs 1          | HCQ             | Cluster+HCQ          |
|----------------------------------------|-----|-------------------------|-----------------|----------------------|
| Bad outcome ~ cluster                   | 4,373 | 2.53 [2.09-3.08] (p<10^{-15}) | 0.71 [0.61-0.83] (p=2.1 × 10^{-5}) | 1.46 [0.95-2.26] (p=0.08) |
| Bad outcome ~ cluster + HCQ             | 4,265 | 1.94 [1.35-2.79] (3.8 × 10^{-4}) | 0.67 [0.56-0.79] (4.0 × 10^{-4}) | 1.47 [0.92-2.36] (0.10) |
| Bad outcome ~ cluster + HCQ + Cluster+HCQ | 4,265 | 1.80 [1.21-2.68] (3.8 × 10^{-3}) | 0.55 [0.45-0.66] (9.0 × 10^{-10}) | 1.47 [0.92-2.36] (0.10) |

The composite bad outcome was defined as one of the following: death, access to intensive care unit, or severe Covid-19 manifestation (either severe pneumonia or ARDS). Associations were tested in three incremental models and in a sensitivity analysis including all the drugs used for Covid-19 treatment, as for Cox PH regressions. Odds Ratios with 95% confidence intervals (OR [CI]) and relevant p-values (in brackets) are reported. Significant ORs (p < 0.05) are highlighted in bold.

4.1. Strengths and Limitations. Although, to the best of our knowledge, this study represents the largest and broadest cluster analysis on Covid-19 patients and a novel approach in analyzing the influence of a pharmacological treatment on Covid-19 mortality and outcomes, it also presents few limitations.

First, the observational retrospective design does not allow us to completely control for confounders and randomization of treatments across individuals. The former aspect is quite unlikely, since a potential residual confounder should be strongly associated with in-hospital mortality to take away observed associations in the interactive model, as suggested by the computed E-values [35, 36]. As for drug therapy, we cannot rule out that assignment to specific treatments was driven also by clinical conditions of the participants, as usually found in common clinical practice. For the same reason, the protective association observed for HCQ may be hypothesized to be driven by other coadministered medications. However, here, HCQ and patients’ cluster showed significant independent associations, which remained substantially stable across models and survived correction for other drugs in use for Covid-19 treatment. Lastly, our evidence is in contrast with RCTs published so far [14–18], which are commonly conceived as the gold standard for establishing drug efficacy and safety. While we generally agree with this view, we would like to underline that these studies did not randomize patients to treatment arms based on combinations of their features, but rather based on single characteristics such as age and sex. This may be the reason for this discrepancy, along with the hypothesis that the high dosage of HCQ administered in RCTs may be harmful for
patients, compared to lower dosages reported in observational studies supporting HCQ efficacy [5]. Of interest, a recent critical review underlined the aspect of suboptimal randomization methods of RCTs, which often do not take into account the whole patient profile and disease severity and may lead to misleading conclusions [38].

5. Conclusions

Overall, the evidence supported here and elsewhere [19] suggests that HCQ treatment may be more effective in specific subtypes of Covid-19 patients and indicates machine learning as a useful approach to identify the most “promising” patients in terms of success rate of this treatment. In the future, further studies on independent datasets are warranted, possibly using supervised ML techniques as in other clinical settings (e.g., [39, 40]), to validate this hypothesis and test the feasibility of predicting responsiveness to HCQ before intervention. Ideally, a trial administering low dosages of HCQ (≤400 mg/day) and randomizing subjects based on their Covid-19 subtype profile rather than on single characteristics may be warranted to clarify the effects of HCQ on mortality risk in SARS-CoV-2 infection, especially within those patients with a “low risk” profile.

This may help solving current controversies on the use of HCQ as a medication for Covid-19 and maximize the efficacy of treatment strategies for this yet largely unknown disease, especially in low-income and developing countries with poorer national health systems.

Data Availability

The data used to support the findings of this study may be released upon application to the CORIST collaboration board, who can be contacted through corresponding author.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

Authors’ Contributions

The raw data analyzed in the present work may be made available under approval by each local center involved in the study, in a way which does not affect patients’ privacy. ADiC, RDC, and LI conceived the CORIST study. AG, ADiC, and LI conceptualized the present work. AG performed statistical analyses and wrote the first draft of the manuscript, under the supervision of ADiC and LI. All the co-authors contributed to collection, curation, and elaboration of the analyzed data, and/or to a critical review and editing of the manuscript. Di Castelnuovo and Gialluisi contributed equally to this manuscript.

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Supplementary Materials

Figure S1. Cross-patient dissimilarity matrix based on Gower distance. Figure S2. Optimal number of clusters (k) to apply in the hierarchical clustering analysis. Figure S3. Check for basic assumptions of Cox PH models. Table S1. Contingency table of Covid-19 cluster by disease severity. Table S2. Contingency tables of Covid-19 clusters by drugs used for treatment. Table S3. Results of Cox PH regressions modelling incident mortality risk, including interaction terms with time. (Supplementary Materials)

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