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Deep phenotyping of 34,128 adult patients hospitalised with COVID-19 in an international network study

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Comorbid conditions appear to be common among individuals hospitalised with coronavirus disease 2019 (COVID-19) but estimates of prevalence vary and little is known about the prior medication use of patients. Here, we describe the characteristics of adults hospitalised with COVID-19 and compare them with influenza patients. We include 34,128 (US: 8362, South Korea: 7341, Spain: 18,425) COVID-19 patients, summarising between 4811 and 11,643 unique aggregate characteristics. COVID-19 patients have been majority male in the US and Spain, but predominantly female in South Korea. Age profiles vary across data sources. Compared to 84,585 individuals hospitalised with influenza in 2014-19, COVID-19 patients have more typically been male, younger, and with fewer comorbidities and lower medication use. While protecting groups vulnerable to influenza is likely a useful starting point in the response to COVID-19, strategies will likely need to be broadened to reflect the particular characteristics of individuals being hospitalised with COVID-19.

he ongoing coronavirus disease 2019 (COVID-19) pandemic is placing a huge strain on health systems worldwide. While a number of studies have provided information on the clinical characteristics of individuals being hospitalised with COVID-[1](#page-9-0)9¹⁻³, substantial uncertainty around the prevalence of comorbidities and prior medication use among this population remains. Moreover, it is not known whether those hospitalised with COVID-19 are systematically different from individuals hospitalised during previous influenza seasons.

COVID-19 shares similarities with influenza to the extent that both cause respiratory disease which can vary markedly in its severity and present with a similar constellation of symptoms, including fever, cough, myalgia, malaise, fatigue and dyspnoea. Early reports do, however, indicate that the proportion of severe infections and mortality rate is higher for COVID-19^{[4](#page-9-0)}. Older age and a range of underlying health conditions, such as immune deficiency, cardiovascular disease, chronic lung disease, neuromuscular disease, neurological disease, chronic renal disease and metabolic diseases, have been associated with an increased risk of severe influenza and associated mortality⁵. While age appears to be a clear risk factor for severe COVID-19[4](#page-9-0), other associations are not yet well understood. Comparisons with COVID-19 are further complicated by the heterogeneity in influenza itself, with different strains resulting in different clinical presentations and associated risks. Those hospitalised with the A(H1N1) pdm09 subtype of the influenza A virus during the associated influenza pandemic in 2009 were, for example, generally younger and with fewer comorbidities than those from preceding influenza seasons⁶.

Here we first aimed to describe the characteristics of patients hospitalised with COVID-19. In particular, we set out to summarise individuals' demographics, medical conditions, and medication use. Our second aim was to compare the characteristics of individuals hospitalised with COVID-19 to those of patients hospitalised with influenza in previous seasons.

Results

Patients hospitalised with COVID-19. A total of 34,128 individuals hospitalised with COVID-19 (CUIMC: 1759; HIRA: 7341; HM: 2078; PHD: 5257; SIDIAP: 16,347; UC HDC: 769; VA OMOP: 577) were included. In all, 68,829 summary characteristics, from 4811 (HM) to 11,643 (CUIMC) unique aggregate characteristics, were extracted and summarised. All are made available in an accompanying interactive website [\(http://evidence.](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/) [ohdsi.org/Covid19CharacterizationHospitalization/](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/)).

Cohorts from CUIMC, HM, PHD, SIDIAP, UC HDC and VA OMOP were predominantly male (52%, 60%, 52%, 54%, 54% and 94%, respectively, but mostly female for HIRA (60%). The age distributions of those hospitalised for COVID-19 are summarised in Fig. [1](#page-3-0) (alongside those hospitalised with influenza, see below). Different patterns are seen in the various contributing databases with, in particular, patients in South Korea (HIRA) seen to be younger than elsewhere.

The distribution of comorbidities in COVID-19 patients varied across sites and countries, see Table [1](#page-4-0). The mean Charlson comorbidity index of those hospitalised with COVID-19 in the US ranged from 3.1 for PHD to 5.4 for VA OMOP, from 0.8 for HM to 1.4 for SIDIAP in Spain, and was 2.7 in HIRA, covering South Korea. In the US, the proportion of those hospitalised with COVID-19 who had asthma ranged from 7 to 15%, from 24 to 43% for diabetes, from 28 to 49% for heart disease and from 8 to 18% for cancer. In Spain, between 4 and 7% for asthma, from 13 to 22% for diabetes, from 17 to 27% for heart disease and from 5 to 16% for cancer. In South Korea, 12% of those hospitalised had a history of asthma, 17% had diabetes, 13% heart disease and 4%

cancer. The prevalence of hypertension ranged from 37 to 70% in the US, from 30 to 46% in Spain and was 24% in South Korea. The prevalence of the full range of conditions summarised are shown in Fig. [2](#page-4-0), with all values reported at [http://evidence.ohdsi.](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/) [org/Covid19CharacterizationHospitalization/.](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/)

For medications, the proportion of those hospitalised with COVID-19 in the US who had been taking agents acting on the renin–angiotensin system over the 30 days up to their hospitalisation ranged from 18% to 39%, while the proportions taking immunosuppressants ranged from 4 to 6%, and from 21 to 51% for lipid-modifying agents over the same time period. In HIRA, 14% had been taking agents acting on the renin–angiotensin system, 1% immunosuppressants, and 16% lipid-modifying agents. In SIDIAP, 27% had been taking agents acting on the renin–angiotensin system, 2% immunosuppressants, and 24% lipid-modifying agents (Table [2\)](#page-5-0). Looking at one medication of particular interest (of which many can be explored in detail at [http://evidence.ohdsi.org/](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/) [Covid19CharacterizationHospitalization/\)](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/), use of hydroxychloroquine on the day of admission was 7% in HIRA and 41% in HM. The prevalence of the many medications summarised are shown in Fig. [2,](#page-4-0) with all values reported [http://evidence.ohdsi.org/](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/) [Covid19CharacterizationHospitalization/.](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/)

Removing the requirement of having a year prior history in datasets other than HM and Premier did not materially change the results (see [http://evidence.ohdsi.org/Covid19Characteriza](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/)[tionHospitalization/](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/) for full details).

A comparison of patients hospitalised with COVID-19 and patients hospitalised with influenza. A total of 84,585 patients hospitalised with influenza between 2014 and 2019 (2125 CUIMC, 49,977 PHD, 2947 SIDIAP, UC HDC 26,547 VA OMOP). In addition, 2443 patients hospitalised with influenza between 2009 to 2010 were included (170 CUIMC, 1689 SIDIAP, 584 VA OMOP) were also identified. Patient characteristics of those hospitalised with COVID-19 are compared to those of individuals hospitalised with influenza between 2014 and 2019 in Figs. [1,](#page-3-0) [3](#page-7-0) and [4](#page-7-0), and with those hospitalised with influenza between 2009 and 2010 in Supplementary Fig. 1 and Supplementary Fig. 2.

A greater proportion of those hospitalised with COVID-19 were male compared to those hospitalised with influenza between 2014 and 2019 for CUIMC, PHD, SIDIAP and UC HDC. Of those hospitalised between 2014 and 2019 with influenza 43, 44, 53 and 45% were male for each of these respective data sources. The ages of those hospitalised with COVID-19 generally appeared slightly younger compared to those hospitalised with influenza between 2014 and 2019, see Fig. [1.](#page-3-0)

Those hospitalised with COVID-19 had a comparable or lower prevalence of comorbidities compared to those hospitalised with influenza 2014–2019 (see Fig. [3](#page-7-0) top and Fig. [4](#page-7-0)). Chronic obstructive pulmonary disease (COPD), cardiovascular disease and dementia were all more common amongst those hospitalised with influenza compared to those hospitalised with COVID-19. Medication use was less common amongst COVID-19 patients (Fig. [3](#page-7-0) bottom and Fig. [4](#page-7-0)) with, for example, systemic corticosteroids and alpha-blockers amongst those more frequently used among those hospitalised with influenza.

Those hospitalised with influenza between 2009 and 2010 were typically younger compared to both COVID-19 and to influenza 2014–2019 admissions (see Supplementary Fig. 1 and Supplementary Table 2). COVID-19 patients were more likely to be male, with 40, 49 and 91% of those hospitalised between 2009 and 2010 for influenza being male for CUIMC, SIDIAP and VA OMOP. Comparisons of conditions and medications, however,

Fig. 1 Age of patients hospitalised with COVID-19 and of patients hospitalised with influenza. Individuals hospitalised with COVID-19 between December 2019 and April 2020 compared with those hospitalised with influenza between September 2014 to April 2019 (where available). Proportion of cohorts by 5-year age groups, with groups with counts of <10 omitted. CUIMC: Columbia University Irving Medical Center; HIRA: Health Insurance Review & Assessment; HM: HM Hospitales; PHD: Premier Healthcare Database; SIDIAP: The Information System for Research in Primary Care; UC HDC: University of Colorado Health Data Compass; VA OMOP: Department of Veterans Affairs. Influenza data for SIDIAP was only available from 2014 to 2017.

varied depending on the data source (see Supplementary Fig. 4 and Supplementary Table 2).

Discussion

Summary of key findings. The characteristics of 34,128 patients hospitalised with COVID-19 in US, South Korea, and Spain have been extracted from EHRs and health claims databases and summarised. Between 5000 and 12,000 unique aggregate characteristics have been produced across databases, with all made publicly available in an accompanying website.

Patients hospitalised with COVID-19 in the US and Spain were predominantly male with age distributions varying across data sources, while those in South Korea were mostly female and appreciably younger than patients in the US and Spain. Many comorbidities were common among individuals hospitalised with COVID-19 with, as an example, 37–70% of those hospitalised with COVID-19 in the US, 30–46% of those in Spain and 24% of those in South Korea having hypertension. Similarly, prior medication use was common with, for example, 18–39% in the US, 27% in Spain and 14% in South Korea, taking drugs acting on the renin–angiotensin system (ACE inhibitors and ARBs) in the 30 days up to their hospitalisation.

Comparisons with previous cohorts of patients admitted to hospital for seasonal influenza in recent years suggest that COVID-19-related admissions are seen more often in younger patients and with a higher proportion of men. In the US and Spain, those hospitalised with COVID-19 were generally either of comparable health or healthier than patients hospitalised with influenza. Consistent differences were noted in the prevalence of respiratory disease, cardiovascular disease and dementia, each more common amongst those hospitalised with influenza in all of the contributing databases. Similarly, the use of corticosteroids and alpha-blockers was consistently higher amongst influenza patients. Those hospitalised with influenza in 2009–2010, during the pandemic associated with H1N1, were seen to be younger than both those hospitalised with influenza in more recent years and patients hospitalised with COVID-19.

Findings in context. A number of studies have previously provided information on individuals hospitalised with COVID-19. While cohorts have generally been majority male, the prevalence of comorbidities has varied. In a study of 1099 individuals who tested positive for COVID-19 in China, of whom 94% were hospitalised, 58% were male, with 7% having diabetes, 15% hypertension and 1% cancer^{[7](#page-9-0)}. In another study of 191 patients with COVID-19 in two hospitals in Wuhan, China, 62% were male, 19% had diabetes, 30% had hypertension and 1% had cancer⁸. In a study which identified 1999 individuals who tested positive for COVID-19 and were hospitalised in New York, 63% were male, 25% had diabetes, 10% COPD and 45% a cardiovascular condition⁹. In another US study of 1482 patients admitted to hospital with COVID-19 in March 2020, 55% were male, with 28% having diabetes, 11% having COPD and 28% having cardi-ovascular disease^{[10](#page-9-0)}. Our findings add to this emerging body of evidence. The results from our study also provide an illustration of the variation in patient characteristics across contexts, with heterogeneity seen both across the cohorts from the US and between the US, Spain and South Korea.

Table 1 Conditions of individuals hospitalised with COVID-19.

Exact proportions have not been reported where counts were <10. Conditions were identified over the year prior and up to, and including, day of hospitalisation (note, HM and PHD had no prior time requirement with conditions primarily recorded on the day of hospital admission).

CUIMC Columbia University Irving Medical Center, HIRA Health Insurance Review & Assessment, HM HM Hospitales, PHD Premier Healthcare Database, SIDIAP The Information System for Research in Primary Care, UC HDC University of Colorado Health Data Compass, VA OMOP Department of Veterans Affairs, COPD chronic obstructive pulmonary disease, GERD gastroesophageal reflux disease.

Fig. 2 Prevalence of conditions and medication use among COVID-19 patients. Individuals hospitalised with COVID-19 between December 2019 and April 2020. Conditions from up to a year prior, medication use from day of hospitalisation. Each dot represents one of these covariates with the colour indicating the type of condition/medication. CUIMC: Columbia University Irving Medical Center; HIRA: Health Insurance Review & Assessment; HM: HM Hospitales; PHD: Premier Healthcare Database; SIDIAP: The Information System for Research in Primary Care; UC HDC: University of Colorado Health Data Compass; VA OMOP: Department of Veterans Affairs.

The comparison with influenza made in our study adds important context when considering the characteristics of those hospitalised with COVID-19. Individuals hospitalised with COVID-19 appear to be more likely younger and male, and in the United States and Spain, to have fewer comorbidities than those hospitalised with influenza in previous years. Indeed, those Table 2 Prior medications of individuals hospitalised with COVID-19.

CUIMC (n: 1759) HIRA (n: 7341) HM (n: 2078) PHD (n: 5257) SIDIAP (n: 16,347) UC HDC (n: 769) VA OMOP (n: 577) Antineoplastic and immunomodulating agents Year up to, and including day of, hospitalisation 9.6% 4.5% – – 3.4% 12.9% 13.2% 30 days up to, and including day of, hospitalisation 6.1% 2.6% – – 2.7% 9.6% 7.1% Day of hospitalisation 4.9% 2.0% 3.8% 2.1% 2.5% 8.2% 4.9% Agents acting on the renin-angiotensin system
Year up to, and including day 27.3% Year up to, and including day of, hospitalisation 27.3% 15.9% – – 30.2% 39.8% 49.6% 30 days up to, and including day of, hospitalisation 18.0% 14.2% – – 26.6% 33.0% 38.5% Day of hospitalisation 15.9% 13.4% 9.9% 7.4% 25.7% 31.5% 33.1% Antiepileptics Year up to, and including day of, hospitalisation 19.0% 9.1% – – 10.6% 27.2% 33.4% 30 days up to, and including day of, hospitalisation 9.9% 5.2% – – 8.1% 21.6% 22.7% Day of hospitalisation 9.1% 4.2% 3.3% 11.3% 7.7% 20.2% 16.6% Anti-inflammatory and antirheumatic products Year up to, and including day of, hospitalisation 23.6% 63.9% – – 30.9% 52.9% 46.1% 30 days up to, and including day of, hospitalisation 10.5% 30.0% – – 9.9% 38.6% 16.8% Day of hospitalisation 7.8% 17.7% 3.8% 7.9% 6.3% 35.1% 9.9% Antithrombotic agents Year up to, and including day of, hospitalisation 44.9% 34.6% – – 25.7% 72.0% 46.1% 30 days up to, and including day of, hospitalisation 29.9% 17.1% – – 21.9% 67.5% 16.8% Day of hospitalisation 28.3% 11.4% 38.2% 38.4% 21.3% 66.8% 9.9% Beta-blocking agents Year up to, and including day of, hospitalisation 26.9% 10.1% – – 13.8% 32.8% 44.4% 30 days up to, and including day of, hospitalisation 16.8% 6.5% – – 12.4% 24.7% 36.0% Day of hospitalisation 15.2% 5.6% 5.7% 17.8% 11.9% 23.8% 31.0% Calcium channel blockers Year up to, and including day of, hospitalisation 23.5% 16.7% – – 12.4% 24.3% 35.0% 30 days up to, and including day of, hospitalisation 15.0% 14.6% – – 10.5% 20.5% 27.0% Day of hospitalisation 13.4% 14.0% 6.8% 9.7% 10.1% 19.9% 24.3% **Diuretics** Year up to, and including day of, hospitalisation 27.6% 10.2% – – 24.6% 33.3% 42.6% 30 days up to, and including day of, hospitalisation 19.2% 7.5% – – 21.5% 27.4% 33.1% Day of hospitalisation 17.7% 6.7% 7.6% 14.1% 20.7% 26.4% 28.4% Drugs for acid related disorders Year up to, and including day of, hospitalisation 35.4% 68.8% – – 30.4% 53.6% 52.0% 30 days up to, and including day of, hospitalisation 20.6% 36.5% – – 22.7% 44.5% 36.2% Day of hospitalisation 17.9% 29.3% 39.7% 21.8% 21.5% 42.4% 28.9% Immunosuppressants Year up to, and including day of, hospitalisation 5.6% 2.0% – – 2.3% 7.5% 5.7% 30 days up to, and including day of, hospitalisation 3.9% 0.7% – – 1.9% 6.1% 3.5% Day of hospitalisation 3.7% 0.4% 2.5% 1.5% 1.7% 5.9% 2.6% Insulins and analogues Year up to, and including day of, hospitalisation 17.8% 3.7% – – 5.5% 31.9% 22.9% 30 days up to, and including day of, hospitalisation 10.9% 2.7% – – 4.9% 26.9% 18.4%

Study participants from HM and PHD were not required to have a year of prior history and so only medications at the time of hospitalisation are summarised.

CUIMC Columbia University Irving Medical Center, HIRA Health Insurance Review & Assessment, HM HM Hospitales, PHD Premier Healthcare Database, SIDIAP The Information System for Research in Primary Care, UC HDC University of Colorado Health Data Compass, VA OMOP Department of Veterans Affairs.

hospitalised with COVID-19 were consistently seen to be less likely to have COPD, cardiovascular disease and dementia than those hospitalised with influenza in recent years.

This study has also added important information on medication use by individuals hospitalised with COVID-19, based on electronic health records and claims data. There is tremendous interest in the risks and benefits of medications such as ACE inhibitors and ARBs for COVID-19, and whether other medications, such as ibuprofen, should be avoided. However, to date, there has been little evidence as to what proportion of those hospitalised with COVID-19 have previously been taking such medications. Our findings shed light on this area, and highlight the importance of further research on the benefits and harms associated with continued use of such treatments, especially those that are commonly taken amongst individuals with COVID-19. It has been seen here, for example, that between 1 in 10 and 2 in 5 of those hospitalised with COVID-19 were taking medicines acting on the renin–angiotensin system in the month before their hospital admission. The consequences of temporarily discontinuing such treatments on cardiovascular risks and mortality remain unknown $¹¹$. Interestingly, corticosteroid use, recently shown to be</sup> effective to treat COVID-19¹², was consistently seen to be less prevalent in patients hospitalised with COVID-19 compared to those hospitalised with influenza across databases, as were alphablockers which some have hypothesised to have a beneficial effect in COVID- $19¹³$.

It should be noted that the characteristics of individuals with COVID-19 have been described in this study at the particular point in time of admission to hospital. While this is of particular interest given its intrinsic link with healthcare utilisation, this only provides a snapshot of the whole picture. Those testing positive for COVID-19 in the community without progressing to hospitalisation can be expected to be younger and with fewer comorbidities than those hospitalised $9,14$ $9,14$ $9,14$, while those progressing to intensive care can be expected to be older and in worse general health $3,15$ $3,15$ $3,15$. In addition, those being referred to or admitted to intensive care also seem more likely to be male $3,15$. Admission to hospital (and intensive care) is also influenced by a range of supply-side factors, such as availability of beds and criteria for admission, and so the characteristics of those hospitalised do not necessarily only reflect the characteristics accompanying severe illness. These factors likely explain some of the heterogeneity seen in those hospitalised with COVID-19 in this study. Geographic variation in populations and transmission dynamics can also be expected to explain the variation across sites. This particularly relevant for patients from South Korea, where given the management of COVID-19 in the country, the patient population is more closely reflective of those infected during early outbreaks.

Study limitations. The study was based on routinely collected data and so, as always, data quality issues must be considered. For instance, individuals were considered as having COVID-19 at the time of hospitalisation only if they had a test result or diagnosis indicating the disease, which will have led to the omission of individuals who can be suspected to have had the disease but lacking confirmation of it. Medical conditions may have been underestimated as they were based on the presence of condition codes, with the absence of such a record taken to indicate the absence of disease. Meanwhile, medication records indicate that an individual was prescribed or dispensed a particular drug, but this does not necessarily mean that an individual took the drug as originally prescribed or dispensed. Our study could be subject to exposure misclassification with false positives if a patient had a medication dispensing event but did not ingest the drug, but may also be subject to false negatives for non-adherent patients who continued their medication beyond the days of supply due to stockpiling. Medication use estimates based on the data collected at the time of hospitalization is particularly sensitive to misclassification, and may conflate baseline concomitant drug history with immediate treatment upon admission.

Time periods were inclusive of the day of hospitalisation, so that information on conditions captured on the day of hospitalisation and medication use up to and including the day of hospitalisation were captured. Other time frames could though have been considered for information prior to hospitalisation. Moreover, outcomes after the index date have not been summarised as these were outside the scope of this particular study. Associations between particular patient characteristics and the risks of hospitalisation and outcomes following hospitalisation were also beyond the scope of this study, and remain an area for further research.

Comparisons of individuals hospitalised with COVID-19 with individuals previously hospitalised with influenza have limitations. In particular, observed differences may be explained by changes in clinical practice or data capture procedures over time, rather than by differences in the individuals themselves. This is likely particularly relevant for comparisons of medication use which, in particular, can be expected to vary over time. Changes in practice in response to the COVID-19 pandemic, such as referrals which may not be observed in our data and thresholds for hospitalisation, also may in part explain some of the observed differences in patient profiles.

Interpretation. Rates of comorbidities and medication use are high among individuals hospitalised with COVID-19. Those individuals hospitalised with COVID-19 do not, however, appear to be in worse general health than those typically hospitalised with influenza. Indeed, in many cases, individuals hospitalised

Fig. 3 Standardised mean difference in conditions (top) and medication use (bottom) among COVID-19 patients compared to 2014–2019 influenza patients. Individuals hospitalised with COVID-19 between December 2019 and April 2020 compared with those hospitalised with influenza between September 2014 and April 2019. Conditions from up to a year prior, medication use from day of hospitalisation. Each dot represents one of these covariates with the colour indicating the type of condition/medication and the size of the dot reflecting the prevalence of the variable in the COVID-19 study populations. CUIMC: Columbia University Irving Medical Center; PHD: Premier Healthcare Database; SIDIAP: The Information System for Research in Primary Care; VA OMOP: Department of Veterans Affairs.

with COVID-19 were seen to be younger and healthier than patients hospitalised with seasonal influenza. Patients hospitalised for COVID-19 are also more likely to be male in comparison to those hospitalised with influenza. Protecting those groups known to be vulnerable to influenza is likely to be a useful starting point to minimize the number of hospital admissions needed for COVID-19, but such strategies may need to be broadened so as to reflect the particular characteristics of individuals seen here to have been hospitalised with COVID-19.

Methods

Study design. This is a cohort study based on routinely collected primary care and hospital electronic health records (EHRs), hospital billing data, and insurance claims data from the US, South Korea and Spain. The data sources used were mapped to the Observational Medical Outcomes Partnership (OMOP) Common Data Model (CDM)¹⁶. The open-science Observational Health Data Sciences and Informatics (OHDSI) network maintains the OMOP CDM, and its members have developed a wide range of tools to facilitate analyses of such mapped data^{[17](#page-9-0)}. Two particular benefits of this approach were that contributing centres did not need to share patient-level data and common analytical code could be applied across databases.

Data sources. Data from the US, South Korea, and Spain underpinned the study. EHR data from the US came from the Columbia University Irving Medical Center (CUIMC), covering NewYork-Presbyterian Hospital/Columbia University Irving Medical Center, University of Colorado Health Data Compass (UC HDC), which includes the UCHealth System with data from 12 hospitals, and United States Department of Veterans Affairs (VA OMOP), which includes 170 medical centres. In addition, data from a US hospital billing system database came from the Premier Hospital database (PHD). EHR data from Spain came from The Information System for Research in Primary Care (SIDIAP), a primary care records database that covers $~80\%$ of the population of Catalonia, Spain¹⁸, and the inpatient care database of HM Hospitales (HM), a hospital group which includes 15 general hospitals from all over Spain, with detailed hospital admission information for COVID-19 patients from March 1 to April 20, 2020. Data from South Korea came from Health Insurance Review & Assessment (HIRA), a repository of national claims data which is collected in the process of reimbursing healthcare providers^{[19](#page-9-0)}. In addition, the analysis was also performed on US EHR data from Tufts-Clinical Academic Research Enterprise Trust (CLARET), which covers Tufts Medical Center (TMC), and the STAnford medicine Research data Repository (STARR-OMOP), which includes data from Stanford Health Care^{[20](#page-9-0)}, and on data from the Daegu Catholic University Medical Center, a teaching hospital in Daegu, South Korea, covered by Federated E-health Big Data for Evidence Renovation Network (FEEDER-NET). These latter sites are not reported here due to smaller study populations, but the results are reported in the accompanying interactive website [\(http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/\)](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/).

Each database was mapped to the OMOP CDM, which was developed as a means of standardising the structure, content and semantics of observational databases²¹. The OHDSI network has maintained and further developed this common data model since 2014, developing tools to both facilitate the mapping of source data to the OMOP CDM and analysing it once mapped. While the journey of implementing the extract, transform, and load process varies across sites (given factors such as infrastructure, size of the database, the complexity of the mapping, and the technical expertise available), the destination is the same for each site; a database which conforms to the requirements of the OMOP CDM. The mapping of source codes to standard concepts is done using OMOP standardised vocabularies, which are regularly updated 22 . In OHDSI studies, such as this one, data partners run the same analysis package using a distributed network approach, where analyses are run locally and only aggregated summary statistics are then shared. OHDSI held a COVID-19 studyathon in March 2020, during which this study was initiated. To note, the OMOP CDM has also been used to facilitate network COVID-19 studies by the 4CE consortium, in which trajectories of laboratory test measurements among COVID-19 patients were described²³, and is being used by the N3C consortium to help harmonise EHR data on COVID-19 patients^{[24](#page-9-0)}.

Study participants. Patients hospitalised between December 2019 and April 2020 with COVID-19 were identified on the basis of having a hospitalisation along with a confirmatory diagnosis or test result of COVID-19 within a time window from 21 days prior to admission up to the end of their hospitalisation. This time window was chosen so as to include those who had the diagnosis made prior to their hospitalisation and allow for a delay in test results or diagnoses to be made or recorded. Patients were also required to be aged 18 years or older at the time of hospitalisation. The same algorithm was used to identify COVID-19 cases across sites, except for CUIMC where the algorithm was adjusted to account for local coding practice. The codes used to identify COVID-19 are described in more detail in Supplementary Methods.

Analogous criteria were used for identifying individuals hospitalised with influenza between September 2014 and April 2019, with individuals identified on the basis of having a hospitalisation along with a confirmatory diagnosis or test result of influenza within a time window from 21 days prior to admission up to the end of their hospitalisation. For SIDIAP, influenza cases were only available up to the end of 2017. An additional cohort of those hospitalised with influenza between September 2009 and April 2010 was also identified. The motivation for this latter group was that the 2009–2010 flu epidemic included many cases of A(H1N1) pdm09 infection, which had different clinical characteristics and associated severity compared to the seasonal flu. Each individual's first hospitalisation with COVID-19 or a particular flu season was considered.

Except for the HM and Premier databases, individuals were required to have a minimum of 365 days of prior observation time available for the primary analysis, to allow for a comprehensive capture of baseline diagnoses and medications up to their hospitalisation. Individuals' observation period in the OMOP CDM reflects a period during which any clinical event that happens to the patient is expected to be recorded. The way this is operationalised does though depend on the type of source data being used. For claims data, for example, observation periods are generally inferred from the enrolment periods to a particular plan, while for EHR observation periods generally begin at the time of interaction with the health system. Consequently, as the requirement for prior observation time could exclude persons with little prior healthcare utilisation or without sustained health insurance, we also characterised cohorts without this restriction in a sensitivity analysis. Given the nature of data collection, individuals in HM and Premier had no prior observation time available and so no requirement for prior observation time was imposed.

Patient features and data analysis. Age at hospitalisation and sex distributions were summarised. Medication use was calculated over three time periods: (1) from a year prior up to, and including, the day of hospitalisation, (2) from 30 days prior up to, and including, the day of hospitalisation and (3) the day of hospitalisation. Only the latter time period was reported for HM and Premier. Drug eras were calculated to give the span of time when an individual is assumed to be exposed to a particular active ingredient. These begin on the start date of the first drug exposure and end on the observed end date if available, or were inferred (for example, based on the number of days of supply). A persistence window of up to 30 days was permitted between two medication records for them to be considered as part of the same drug era. Individual medications were categorised using Anatomical Therapeutic Chemical (ATC) groupings. All drugs are reported in full in a dedicated interactive website (see 'Results' section), but specific classes are reported here based on recent interest due to their potential effects (positive or negative) on COVID-19 susceptibility or severity: agents acting on the renin–angiotensin system (including angiotensin-converting enzyme (ACE) inhibitors and angiotensin II receptor blockers (ARBs)), antiepileptics, beta-blocking agents, calcium channel blockers, diuretics, drugs for acid related disorders, immunosuppressants, insulins, analogues and lipid-modifying agents (such as statins). Prevalence of medication use for each time window was determined by the proportion of persons who had at least one day during the time window overlapping with a drug era for each medication or drug class of interest. Conditions were identified on the basis of SNOMED codes, with all descendent codes included. Similarly, all recorded diagnoses are available for consultation in the accompanying website, but a list of key conditions is reported here based on recent reports of associations with COVID-19 infection or outcomes.

Age distributions in each cohort were plotted. The proportion of a cohort having a particular characteristic was described, with the prevalence of all medications and conditions captured in the databases depicted using Manhattanstyle plots. Standardised mean differences (SMD) were calculated when comparing characteristics of study cohorts, with a SMD of above 0.1 generally considered to indicate a meaningful difference in the prevalence of a covariate^{[25](#page-9-0)}. The SMDs for all variables were plotted in Manhattan-style plots. In addition, the prevalence of particular conditions or medications of interest among those hospitalised with COVID-19 (Y axis) were compared to those hospitalised with influenza (X axis) in scatter plots, with dots on the top-left indicating a higher prevalence among those hospitalised with COVID-19 and dots on the bottom-right indicating a higher prevalence among those hospitalised with influenza.

Ethical approval. All the data partners received Institutional Review Board (IRB) approval or exemption. STARR-OMOP had approval from IRB Panel #8 (RB-53248) registered to Leland Stanford Junior University under the Stanford Human Research Protection Program (HRPP). The use of VA data was reviewed by the Department of Veterans Affairs Central Institutional Review Board (IRB) and was determined to meet the criteria for exemption under Exemption Category 4(3) and approved the request for Waiver of HIPAA Authorization. The research was approved by the Columbia University Institutional Review Board as an OHDSI network study. The IRB number for use of HIRA data was AJIB-MED-EXP-20- 065. HM Hospitales and SIDIAP analyses were approved by the Clinical Research Ethics Committee of the IDIAPJGol (project code: 20/070-PCV). The UC-HDC data use was reviewed by Colorado Multi-Institutional Review Board (COMIRB) and was determined to meet the criteria for exemption under Exemption Category

4(3) and approved the request for Waiver of HIPAA Authorization (protocol # 20- 0730).

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The aggregated results set, that does not include patient-level health information, is available via [http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/.](http://evidence.ohdsi.org/Covid19CharacterizationHospitalization/) Data partners (Columbia University Irving Medical Center [CUIMC], Health Insurance Review & Assessment [HIRA], HM Hospitales [HM], Premier Hospital database [PHD], The Information System for Research in Primary Care [SIDIAP], United States Department of Veterans Affairs [VA OMOP] and University of Colorado Health Data Compass [UC HDC]) contributing to this study remain custodians of individual patientlevel health information.

Code availability

The analytic code used in the study has been made available at [https://github.com/ohdsi](https://github.com/ohdsi-studies/Covid19HospitalizationCharacterization.)[studies/Covid19HospitalizationCharacterization](https://github.com/ohdsi-studies/Covid19HospitalizationCharacterization.)²⁶. This study package is run from R and is a OHDSI CohortDiagnostics-type package²⁷. This makes use of other OHDSI packages, such as FeatureExtraction²⁸, to extract patient characteristics for user-specified cohorts in the common data model.

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References

- 1. Chen, N. et al. Epidemiological and clinical characteristics of 99 cases of 2019 novel coronavirus pneumonia in Wuhan, China: a descriptive study. Lancet 395, 507–513 (2020).
- 2. Wang, D. et al. Clinical characteristics of 138 hospitalized patients with 2019 novel coronavirus–infected pneumonia in Wuhan, China. JAMA 323, 1061–1069 (2020).
- 3. Bhatraju, P. K. et al. Covid-19 in critically ill patients in the seattle region case series. N. Engl. J. Med. <https://doi.org/10.1056/NEJMoa2004500> (2020).
- 4. Verity, R. et al. Estimates of the severity of coronavirus disease 2019: a modelbased analysis. Lancet 3099, 1–9 (2020).
- 5. Mertz, D. et al. Populations at risk for severe or complicated influenza illness: systematic review and meta-analysis. BMJ Br. Med. J. 347, f5061 (2013).
- 6. Reed, C. et al. Complications among adults hospitalized with influenza: a comparison of seasonal influenza and the 2009 H1N1 pandemic. Clin. Infect. Dis. 59, 166-174 (2014).
- 7. Guan, W. et al. Clinical characteristics of coronavirus disease 2019 in China. N. Engl. J. Med. <https://doi.org/10.1056/NEJMoa2002032> (2020).
- 8. Zhou, F. et al. Clinical course and risk factors for mortality of adult inpatients with COVID-19 in Wuhan, China: a retrospective cohort study. Lancet 395, 1054–1062 (2020).
- 9. Petrilli, C. M. et al. Factors associated with hospitalization and critical illness among 4103 patients with COVID-19 disease in New York City. medRxiv <https://doi.org/10.1101/2020.04.08.20057794> (2020).
- 10. Garg, S., Kim, L., Whitaker, M., O'Halloran, A. & Cummings, C. Hospitalization rates and characteristics of patients hospitalized with laboratory-confirmed coronavirus disease 2019—COVID-NET, 14 States, March 1–30, 2020. MMWR Morb. Mortal Wkly Rep. [https://doi.org/10.15585/](https://doi.org/10.15585/mmwr.mm6915e3) [mmwr.mm6915e3](https://doi.org/10.15585/mmwr.mm6915e3) (2020).
- 11. Whiting, P. et al. What are the risks and benefits of temporarily discontinuing medications to prevent acute kidney injury? A systematic review and metaanalysis. BMJ Open 7, e012674 (2017).
- 12. Horby, P. et al. Effect of dexamethasone in hospitalized patients with COVID-19: preliminary report. medRxiv <https://doi.org/10.1101/2020.06.22.20137273> (2020).
- 13. Vogelstein, J. T. et al. Alpha-1 adrenergic receptor antagonists for preventing acute respiratory distress syndrome and death from cytokine storm syndrome. Preprint at <https://arxiv.org/abs/2004.10117> (2020).
- 14. CDC COVID-19 Response Team. Preliminary estimates of the prevalence of selected underlying health conditions among patients with coronavirus disease 2019—United States, February 12–March 28, 2020. MMWR Morb. Mortal Wkly Rep. 69, 382 (2020).
- 15. Grasselli, G. et al. Baseline characteristics and outcomes of 1591 patients infected with SARS-CoV-2 admitted to ICUs of the Lombardy Region, Italy. JAMA <https://doi.org/10.1001/jama.2020.5394> (2020).
- 16. Voss, E. A. et al. Feasibility and utility of applications of the common data model to multiple, disparate observational health databases. J. Am. Med. Inf. Assoc. 22, 553–564 (2015).
- 17. Hripcsak, G. et al. Observational health data sciences and informatics (OHDSI): opportunities for observational researchers. Stud. Health Technol. Inform. 216, 574–578 (2015).
- 18. García-Gil, M. D. M. et al. Construction and validation of a scoring system for the selection of high-quality data in a Spanish population primary care database (SIDIAP). Inform. Prim. Care 19, 135–145 (2011).
- 19. Kim, J.-A., Yoon, S., Kim, L.-Y. & Kim, D.-S. Towards actualizing the value potential of Korea Health Insurance Review and Assessment (HIRA) data as a resource for health research: strengths, limitations, applications, and strategies for optimal use of HIRA data. J. Korean Med. Sci. 32, 718–728 (2017).
- 20. Datta, S. et al. A new paradigm for accelerating clinical data science at Stanford Medicine. Preprint at <https://arxiv.org/abs/2003.10534> (2020).
- 21. Overhage, J. M., Ryan, P. B., Reich, C. G., Hartzema, A. G. & Stang, P. E. Validation of a common data model for active safety surveillance research. J. Am. Med. Inform. Assoc. 19, 54–60 (2011).
- 22. Observational Health Data Sciences and Informatics. The Book of OHDSI (Independently published, 2019).
- 23. Brat, G. A. et al. International electronic health record-derived COVID-19 clinical course profiles: the 4CE consortium. npj Digit. Med. 3, 109 (2020).
- 24. Melissa, H., Christopher, C. & Kenneth, G. The National COVID Cohort Collaborative (N3C): rationale, design, infrastructure, and deployment. J. Am. Med. Informatics Assoc. <https://doi.org/10.1093/jamia/ocaa196> (2020).
- 25. Austin, P. C. An introduction to propensity score methods for reducing the effects of confounding in observational studies. Multivar. Behav. Res. 46, 399–424 (2011).
- 26. Burn, E. et al. Deep phenotyping of 34,128 patients hospitalised with COVID-19 and a comparison with 81,596 influenza patients in America, Europe and Asia: an international network study. medRxiv [https://doi.org/10.1101/](https://doi.org/10.1101/2020.04.22.20074336) [2020.04.22.20074336](https://doi.org/10.1101/2020.04.22.20074336) (2020).
- 27. OHDSI. CohortDiagnostics. <https://github.com/OHDSI/CohortDiagnostics>.
- 28. OHDSI. FeatureExtraction. [https://github.com/OHDSI/FeatureExtraction.](https://github.com/OHDSI/FeatureExtraction)

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Study conception: E.B., S.C.Y., L.S., D.V., A.G., F.N., D.P.V. and P.R.; data analysis: E.B., A.S., S.C.Y., T.F., S.D. and J.P.; initial drafting of the paper: E.B., D.P.V. and P.R. All authors (E.B., S.C.Y., A.G.S., K.K., H.A., M.T.F.A., A.A., H.A., O.A., T.M.A., M.A., C.A, J. M.B., J.C., A.C.C., A.D., F.J.D., T.D.S., S.D., T.F., S.F.B., W.G., A.G., J.H., G.H., V.H., H.J., Y.J., C.Y.J., B.S.K.H., D.K., S.K., Y.K., S.K., J.C.E.L., H.L., K.E.L., R.M., M.E.M., P.P.M., D. R.M., K.N., F.N., A.O., R.W.P., J.P., J.D.P., A.P.U., G.R., C.R., Y.R., P.R., L.M.S., M.S., N. H.S., A.S., S.S., M.S., M.A.S., J.N.S., D.V., S.V., H.W., A.E.W., B.B.Y., L.Z., O.Z., D.P.A. and P.R.) were involved in the design of the work, interpretation of the data and revising the draft paper.

Competing interests

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Additional information

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