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Associations of diabetes, hypertension and obesity with COVID-19 mortality: a systematic review and meta-analysis

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ABSTRACT

Introduction Despite a growing body of scholarly research on the risks of severe COVID-19 associated with diabetes, hypertension and obesity, there is a need for estimating pooled risk estimates with adjustment for confounding effects. We conducted a systematic review and meta-analysis to estimate the pooled adjusted risk ratios of diabetes, hypertension and obesity on COVID-19 mortality.

Methods We searched 16 literature databases for original studies published between 1 December 2019 and 31 December 2020. We used the adapted Newcastle-Ottawa Scale to assess the risk of bias. Pooled risk ratios were estimated based on the adjusted effect sizes. We applied random-effects meta-analysis to account for the uncertainty in residual heterogeneity. We used contour-funnel plots and Egger's test to assess possible publication bias.

Results We reviewed 34 830 records identified in literature search, of which 145 original studies were included in the meta-analysis. Pooled adjusted risk ratios were 1.43 (95% Cl 1.32 to 1.54), 1.19 (95% CI 1.09 to 1.30) and 1.39 (95% CI 1.27 to 1.52) for diabetes, hypertension and obesity (body mass index \geq 30 kg/m²) on COVID-19 mortality, respectively. The pooled adjusted risk ratios appeared to be stronger in studies conducted before April 2020, Western Pacific Region, low- and middle-income countries, and countries with low Global Health Security Index scores, when compared with their counterparts. **Conclusions** Diabetes, hypertension and obesity were associated with an increased risk of COVID-19 mortality independent of other known risk factors, particularly in lowresource settings. Addressing these chronic diseases could be important for global pandemic preparedness and mortality prevention.

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INTRODUCTION

The COVID-19 pandemic has caused over 753.4 million reported cases and over 6.8 million deaths globally as of 1 February 2023.¹ Early in the pandemic, older people, and people with pre-existing non-communicable diseases (NCDs) and related risk factors ('comorbid-ities'), including hypertension, diabetes and obesity, were found to be at higher risk of severe

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Early in the COVID-19 pandemic, older people and people with pre-existing non-communicable diseases and related risk factors were found to be at higher risk of severe COVID-19 illness and death. However, estimates of the strength of associations of diabetes, hypertension and obesity with COVID-19 mortality are highly variable, and additional findings, representative of the global context and adjusted for potential confounding effects, are needed.

WHAT THIS STUDY ADDS

 \Rightarrow In this comprehensive and rigorous systematic review and meta-analysis, we assessed the strength of adjusted associations of diabetes, hypertension and obesity with COVID-19 mortality using data of 145 observational studies conducted in 26 countries. We estimated that patients with diabetes, hypertension and obesity were at about 43%, 19% and 39% increased risk of COVID-19 mortality, respectively, independent of other known risk factors. Pooled adjusted risk ratios for the association of diabetes, hypertension and obesity with COVID-19 mortality were approximately 33%, 43% and 4%, smaller than the unadjusted risk ratios. The adjusted risk ratios appeared to be stronger in studies conducted before April 2020, in the Western Pacific region, in low- and middle-income countries, and in countries with lower Global Health Security Index scores, when compared with their counterparts.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ Our findings add to the body of evidence that shows the important relationship between underlying chronic diseases and mortality during the COVID-19 pandemic and support the need for further research on pathophysiological mechanisms. Efforts to reduce the prevalence and impact of chronic diseases and improve the function of core health systems are essential to population health in all countries at all times and would especially improve population resilience during times of pandemic threats. COVID-19 illness and death.²⁻⁴ This is not a new phenomenon, as viral respiratory infections (eg, influenza, Severe Acute Respiratory Syndrome or SARS and Middle Eastern Respiratory Syndrome or MERS) have previously been linked with a higher risk of severe outcomes among patients with comorbidities.⁵ The US Centers for Disease Control and Prevention defines "higher risk" for severe outcomes as an underlying medical condition or risk factor that has a published meta-analysis or systematic review demonstrating good or strong evidence for an increase in risk for at least one severe COVID-19 outcome.⁶ The risk of COVID-19 death increases as the number of comorbid conditions increases. The population-level consequences of COVID-19 illness are compounded by the increasing global burden of NCDs, which increases the potential benefit of reducing this burden through efforts targeted to prevention, early diagnosis, screening and treatment.^{7–9} To understand the magnitude of the dual epidemics of COVID-19 and NCDs, it is estimated that 349 million people, or 4% of the global population, are at high risk of severe COVID-19 due to age and pre-existing comorbidities.¹⁰ Moreover, the proportion varies across regions, ranging from 3.0% in Africa to 6.5% in Europe.¹⁰

Prior to the pandemic, the global prevalence of diabetes was estimated to be 9.3% among adults aged 20-79 years, with an increasing prevalence reaching 19.9% for those aged 65–79 years.¹¹ Global prevalence of hypertension was estimated to be 31.1% in the adult population.¹² Global prevalence of overweight and obesity combined is estimated to be 39.0% in the adult population, with 12.5% prevalence of obesity alone.¹³ Hypertension was identified early in the pandemic as a prevalent comorbidity among severely ill patients.¹⁴ After vaccines became available in 2021, hypertension continued to be an important comorbidity and was associated with a blunted serological response following vaccine administration in hypertensive versus normotensive patients.^{15 16} COVID-19 infected individuals with diabetes, a disease associated with chronic inflammation and hyperglycaemia, reportedly have a 2-3 fold increase in mortality from COVID-19 compared with people without diabetes.^{3 17 18} An exploratory study of UK medical records found the risk of dying from COVID-19 was almost three times higher for patients with type 1 diabetes and almost twice as high for type 2, versus those without diabetes.¹⁸ Obesity is both a disease and a major risk factor for many adverse health conditions, including diabetes and hypertension.¹⁹ With differences seen by age, race and sex, in populations with a high prevalence of obesity, as much as one-third of hypertension is reportedly due to obesity.²⁰ During the COVID-19 pandemic, obesity was found to be significantly associated with increased severity in terms of intensive care hospitalisation and mechanical ventilation and higher mortality among COVID-19 patients.²¹

Although, at the time of writing, the SARS-CoV-2 virus is still circulating globally, in many parts of the world, the pandemic is transitioning from response to recovery. Countries and public health decision makers must address common risk factors of NCDs and infectious diseases to decrease the economic burden of disease management and BMJ Glob Health: first published as 10.1136/bmjgh-2023-012581 on 14 December 2023. Downloaded from http://gh.bmj.com/ on April 28, 2024 by guest. Protected by copyright

to improve health outcomes as they evaluate the population level impact of COVID-19 on health systems and prepare for the next pandemic.²² ²³ Information on the consequences of pre-existing comorbidities has been reported throughout the pandemic, suggesting patterns of vulnerability within populations. Meta-analyses of high-quality studies with wide geographical representativeness are best suited to increase the accuracy of results used to inform health system recovery and strengthening. Therefore, in this study, we conducted a systematic review and meta-analysis to bring together the global evidence on the independent associations of diabetes, hypertension and obesity with mortality in COVID-19 patients and differences in these associations across regions, country-level characteristics and study-level characteristics.

METHODS

Search strategy and selection criteria

We conducted this systematic review and meta-analysis according to Conducting Systematic Reviews and Meta-Analyses of Observational Studies of Etiology (COSMOS-E) guidelines²⁴ and reported our results according to the Meta-analysis Of Observational Studies in Epidemiology checklist.²⁵ The details of eligibility criteria, study inclusion and exclusion criteria, data sources and search strategy, and study selection were developed with the assistance of an expert medical librarian at the CDC and delineated in our protocol, which was registered at PROSPERO and published previously.²⁶ In brief, we formulated our study eligibility criteria using the PECOS (Population/Participants, Exposures, Comparators, Outcomes and Study designs) description model.²⁴ Participants were male and female patients aged 18 years or older with laboratory-confirmed positive COVID-19 by molecular (PCR) or antigen test for COVID-19. Primary exposures were diabetes (defined as having a history of diagnosed diabetes by self-report or medical record or use of blood glucose lowering medications prior to the confirmation of COVID-19 or defined specifically in the study methods), hypertension (defined as having a history of diagnosed hypertension by self-report or medical record or use of blood pressure medications prior to the confirmation of COVID-19 or defined specifically in the study methods), and obesity (defined as having a history of established obesity with a body mass index (BMI)≥30 kg/m² prior to the confirmation of COVID-19 or as defined in individual studies). Comparators were patients with no history of preexisting diabetes, hypertension or obesity. The primary outcome was COVID-19 death, defined as people who have had a positive PCR or antigen test for COVID-19, died from a clinically compatible illness or syndrome attributable to COVID-19, and were not due to non-natural causes (eg, accidental, intentional self-harm, homicide).^{27 28} Meanwhile, the ICD-10 code U07.1 (COVID-19, virus identified) or U07.2 (COVID-19, virus not identified) was also used to define COVID-19 death. We considered cohort studies, case-control studies and cross-sectional studies

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to be eligible. Some randomised controlled trials for COVID-19 treatments and case series were carefully reviewed and considered to be eligible when sufficient data on specified 'exposures', 'comparators' and 'outcomes' were available. For studies labelled as case-series studies, we reassessed these studies and reclassified them to be either cohort studies (if they reported a follow-up time or attempt, or an HR), or cross-sectional studies if they did not.²⁹

We searched 16 databases (platforms) including MEDLINE (Ovid), Embase (Ovid), Global Health (Ovid), CAB Abstracts (Ovid), PsycINFO (Ovid), CINAHL (Ebsco), Academic Research Complete (Ebsco), Africa Wide Information (Ebsco), Scopus, PubMed Central, ProQuest Central (Proquest), WHO Virtual Health Library, Homeland Security COVID-19 collection, SciFinder (CAS), Clinical Trials and Cochrane Library for primary or original articles published between December 2019 and December 2020. Our rigorous and broad literature search strategy used key words or terms including, 'novel coronavirus, 2019 coronavirus, coronavirus disease, coronavirus 2019, betacoronavirus, COVID-19, COVID-19, nCoV, novel CoV, CoV 2, CoV2, sarscov2, sars-cov, sarscov, 2019nCoV, 2019-nCoV, severe acute respiratory syndrome or pneumonia outbreak or pandemic' and diabetes, obesity/overweight, hypertension, comorbidity, chronic disease, noncommunicable disease, cardiovascular disease, metabolic, predictor, risk factor or determinant' with no limitations on age, sex, publication type or language. Detailed search strategy and the number of records are presented in online supplemental text 1. After careful discussion, we decided not to search the grey literature and the reference lists of the included studies for additional records, because grey literature is not relevant to our research topic, and our literature search of 16 databases is likely to cover all potential original peer-reviewed articles since the start of COVID-19 pandemic in our defined time frame.

The initial search was carried out by the researchers, with technical assistance from an experienced medical librarian from CDC. All references were then collated in EndNote V.20. After the exclusion of duplicates using the function in EndNote V.20, the remaining articles were imported to Covidence Toolkit (a web-based collaboration software platform that streamlines the production of systematic and other literature reviews)³⁰ for further screening, review, data extraction and risk of bias assessment. For final inclusion, each study was assessed independently by two or more researchers, first by screening the title and abstract, and then through a full-text review. Disagreements on the selection of records between the two researchers were resolved by team discussion or by a third researcher.

Data analysis

Two researchers independently extracted data from each article. This included study-level characteristics such as first author and publication year, geographical location and setting, start and end dates, design, COVID-19 confirmation method, and data collection method. It also included detailed data on study participants, their exposures (diabetes, hypertension and obesity), and outcomes (mortality), and effect estimate measures reported as unadjusted, age-adjusted and age-adjusted and sex-adjusted, and multivariable-adjusted, as well as a list of covariates or potential confounders. Effect measures, including OR, HR,or relative risk (RR) and their 95% CI, were extracted directly from the studies when available. Disagreements in data extraction were resolved by a third researcher. For articles with missing data, we emailed the authors to request the data (eight requests sent and six responses received).

The Newcastle-Ottawa scale (NOS) was adapted to assess the risk of bias (quality) of included studies with a cohort, case–control or cross-sectional design (online supplemental text 2).^{31 32} Two researchers independently assessed the quality of studies. Disagreement between the two researchers in the quality assessment was resolved by a third researcher.

Overall pooled risk ratios (PRR) for the association between the exposure variables and the risk of COVID-19 death were conducted according to the type of risk ratio (OR, HR or RR) separately and according to adjustment for potential confounding effects (unadjusted vs multivariable-adjusted risk ratios) for each of the exposure variables (diabetes, hypertension and obesity), respectively. In the subgroup analyses, we combined studies with OR, HR and RR to ensure an adequate number of studies in each subgroup and estimated PRR as we considered HR and OR to be approximate measures of risk ratios given the low COVID-19 mortality rate globally.³³³⁴

We applied random-effects meta-analysis using a restricted maximum likelihood method35 36 and a Hartung-Knapp-Sidik-Jonkman (HKSJ) adjustment to the standard errors to account for the uncertainty in residual heterogeneity.³⁷⁻³⁹ We further applied an ad hoc Knapp-Hartung method to ensure that the HKSJadjusted SEs were appropriate given the unadjusted SEs.^{40 41} To assess the potential effects of geographical locations, socioeconomic factors and healthcare system on the associations between the exposure variables and the risk of COVID-19 death, subgroup analyses (stratified analyses, with ≥ 3 studies in each subgroup) were conducted by study design (cohort, case-control or crosssectional), study period (December 2019-April 2020 or May 2020-November 2020), WHO regions (Africa, South-East Asia, Americas, East Mediterranean, Europe, West Pacific inclusive of mainland China and West Pacific exclusive of mainland China), World Bank (WB) income level (high, upper-middle, lower-middle and low),⁴² NOS quality assessment score (high=8-9, medium=5-7, $low \le 5$) (Table S3),^{31 32} health index score (a measure of the extent to which people are healthy and have access to the necessary services to maintain good health, including health outcomes, health systems, illness and risk factors,

and mortality rates, with a higher score indicating a higher ranking),⁴³ and Global Health Security Index (GHSI) score (an index of a country's global health security capacity to prevent epidemics, with a higher score indicating a better health security and capability).⁴⁴ Meta-regression was conducted to assess the linear relationship between the continuous study-level and country-level indicators and the risk ratios using random-effects method.

Sensitivity analysis was carried out to assess the influence of individual studies on the PRR using influence plots, where one study was excluded at a time to see its effect on the overall estimate. Possible publication bias was assessed by contour-funnel plots and Egger's test.^{45–48} The tau-squared (τ^2) statistics were reported as a measure of between-study variance, while the I² statistic was reported as the proportion of total variability explained by between-study variance. All statistical analyses were carried out using the statistical software R V.4.2.2 and Stata V.16.1 (StataCorp).

Patient and public involvement

Patients or the public were not involved in the design, conduct, reporting or dissemination plans of our systematic review and meta-analysis. However, many contributing studies did involve patients and community stakeholders in the design and dissemination of their study results.

RESULTS

Characteristics of included studies

As we focused on synthesising adjusted estimates in this study, data from 145 studies conducted in 26 countries with adjusted risk ratios for the associations of diabetes, hypertension and obesity with COVID-19 mortality contributed to the quantitative analysis. We excluded 1329 studies with various reasons and additional 125 studies due to lack of data for the primary outcome (n=30), or for the primary exposures (n=12), or for adjusted risk ratios (n=83) (figure 1). Among 145 studies, 139 provided results from the fully adjusted models (age, sex, plus one or more comorbidities, complications, or other health risk factors) and 6 from age-adjusted and sex-adjusted models. The geographical distributions of the studies are presented in the map (figure 2). Countries with a large number of studies included the USA (N=40), China (N=23), Italy (N=15), Mexico (N=9), South Korea (N=9) and Spain (N=8). Most of the studies were started between December 2019 and April 2020 (97.2%), had a cohort design (79.3%), reported HR (40.0%) or OR (53.8%), used data from electronic health (medical) records (57.9%), had a high NOS score of 8 or 9 (73.8%), were from high (63.4%) or upper middleincome (32.4%) countries, had a health index score 70 or above (95.6%) and had a GHSI score 33.4 or above (97.2%) (table 1).

The median (ie, centre) and the IQR (defined as the difference between the 25th and 75th percentile) (ie, spread or dispersion) of the sample sizes are similar for diabetes and hypertension. Although the total number of studies for obesity (n=57) is smaller than those for diabetes (n=118) and hypertension (n=99), the median and the spread of the sample sizes in studies for obesity are larger than those for diabetes and hypertension (table 1).

Detailed characteristics of all 145 studies included in the meta-analysis are presented in online supplemental table S3. Because of a large number, details of the total excluded studies with reasons (n=1454) are not presented (available on request).

Meta-analysis

As expected, the overall pooled unadjusted risk ratios were larger than the adjusted risk ratios on COVID-19 mortality for diabetes (2.13, 95% CI 1.80 to 2.52; n=118), hypertension (2.07, 95% CI 1.74 to 2.47; n=99) and obesity (1.46, 95% CI 1.22 to 1.71; n=57) (figure 3). The overall pooled risk estimates using the OR slightly overestimated the risk estimates using HR and risk ratios (RR). The detailed numeric values of overall PRRs were presented in online supplemental table S1. In addition, details of the forest plots for the individual studies were shown in online supplemental figure S1.1 (diabetes), online supplemental file 1.2 (hypertension) and online supplemental figure S1.3 (obesity).

The pooled adjusted risk ratio for the association between diabetes and mortality was 1.43 (95% CI 1.32 to 1.54; n=118) with considerable heterogeneity $(\tau^2 = 0.12; I^2 = 0.94)$ (table 1). Sensitivity analysis indicated that the exclusion of any one of the studies did not significantly impact the overall PRR (online supplemental figure S2.1). Subgroup analysis showed a lower PRR in countries with a lower health index score, with a higher GHSI score, with a high-income level by WB, in studies with a cohort design, or with a high quality by NOS. In contrast, a higher PRR was observed in countries from the WHO WPR region (figure 4). The detailed numeric value of PRRs by subgroups was presented in online supplemental table S2.1. Meta-regression showed a negative association between the mean age of the participants (p=0.02) and GHSI score (p=0.02) with the risk ratios, and a positive association of health index score (p=0.003) with the risk ratios (table 2). There was no evidence of a funnel plot asymmetry in the association between diabetes and COVID-19 mortality (Egger's test p=0.29) (figure 5).

The pooled adjusted risk ratio for the association between hypertension and mortality was 1.19 (95% CI 1.09 to 1.30; n=99) with considerable heterogeneity ($\tau^2 = 0.12$; $I^2 = 0.91$) (table S1). Sensitivity analysis indicated that the exclusion of any one of the studies did not have any significant impact on the overall PRR (online supplemental figure S2.1). Subgroup analysis showed a lower PRR in studies with high quality, in the WB high



Figure 1 PRISMA flow diagram. PRISMA = Preferred Reporting Items for Systematic Reviews and Meta-Analyses.

income countries, and countries with a higher GHSI score, and a higher PRR in countries from the WHO WPR region (figure 4). Meta-regression showed a negative association of mean age of the participants (p=0.02) and GHSI score (p=0.04) with the risk ratios. There was no evidence of a funnel plot asymmetry in the association between hypertension and COVID-19 mortality (Egger's test p=0.25) (figure 5).

The pooled adjusted risk ratio for the association between obesity and mortality was 1.39 (95% CI 1.27 to 1.52; n=57) with considerable heterogeneity ($\tau^2 = 0.06$; $I^2 = 0.96$) (table S1). Sensitivity analysis indicated that the exclusion of any one of the studies did not significantly impact the overall PRR (online supplemental figure S2.3). Due to the small number of studies reporting adjusted obesity-COVID-19 mortality associations, some



Figure 2 Number and distribution of studies included in the meta-analysis by country. Colours in the map indicate the various number of studies included in the meta-analysis by country (n=143). Studies conducted in multiple countries (n=2) were not shown in this map.

subgroup analyses could not be conducted. Subgroup analysis showed a higher PRR in studies from the EUR region than from the AMR region, and in studies conducted in April or earlier than those conducted in May or later in 2020 (figure 4). Meta-regression showed a negative association of GHSI score (p=0.001) with the risk ratios. There was evidence of a funnel plot asymmetry in the association between obesity and COVID-19 mortality (Egger's test p=0.002) (figure 5). There was a suggestion of missing studies in the middle left-hand side of the contour-funnel plot (ie, small studies with high SE), broadly in the non-significance region (white area where p>0.1), making publication bias plausible. The detailed numeric value of PRRs for the subgroup analyses is presented in online supplemental table S2.

DISCUSSION

In this systematic review and meta-analysis, we estimated that persons with diabetes, hypertension and obesity were at about 43%, 19% and 39% increased risk of COVID-19 mortality, respectively, independent of other known risk factors. Our results showed that pooled adjusted risk ratios for the association of diabetes, hypertension and obesity with COVID-19 mortality were approximately 33%, 43% and 4% smaller than their unadjusted risk ratios. Moreover, the pooled adjusted risk ratios appeared to be stronger in studies conducted before April 2020, in the Western Pacific region, in low- and middle-income countries, and in countries with a lower GHSI score, when compared with the counterparts.

It is noteworthy to mention that the lower adjusted risk ratios for diabetes and hypertension on COVID-19 mortality than their unadjusted estimates as observed in this study confirm that unadjusted risk ratio could overestimate the real associations, as age, sex, health risk factors and other comorbidities and complications could be related to both the exposure measures and COVID-19 mortality. Across a number of published systematic reviews and meta-analyses, the majority reported the unadjusted estimates that failed to consider possible confounding effects and thus likely biased the strength or direction of the associations.^{49–53} As reported in a recent umbrella meta-analysis,49 the pooled unadjusted risk ratios for diabetes, hypertension and obesity with COVID-19 mortality were 2.09, 2.50 and 2.18, respectively, which were similar to the pooled unadjusted risk ratios in this study. In other umbrella meta-analyses, pooled unadjusted risk ratios for diabetes and hypertension on COVID-19 mortality were 1.87 and 1.79, respectively.^{50 51} The pooled unadjusted risk ratios for obesity on COVID-19 mortality ranged from 0.89 to 3.52.^{52 53} Umbrella reviews, which are reviews of previously published systematic reviews and meta-analyses, could be a cost-effective way to summarise information available on a specific topic.^{54 55} However, umbrella reviews might suffer from reliance on studies and reviews lacking in quality or data. Indeed, as shown in a recent umbrella meta-analysis, the majority of published systematic reviews and meta-analyses on the association between obesity and mortality in patients with COVID-19 presented critically low quality and very low certainty of the evidence.⁵³

Our results on the pooled adjusted risk ratios for diabetes and hypertension in relation to COVID-19 mortality are consistent with the summary RR estimates adjusted for multiple confounders reported in recently published meta-analyses with inclusion of studies published as of 2022.^{2 50 51 56} Therefore, our findings provide further evidence and support on the independent effects and highlighted importance of possible confounding effects for the association of diabetes and hypertension with COVID-19 mortality.

The association between BMI and COVID-19 mortality appeared to be inconsistent in published studies.²¹⁵³⁵⁷⁻⁵⁹ Persons with unclassified obesity (BMI \ge 30 kg/m²) or those with class III obesity (BMI \geq 40 kg/m²) were at risk of COVID-19 mortality, whereas those with obesity classes I $(30 \le BMI < 35 \text{ kg/m}^2)$ or II $(35 \le BMI < 40 \text{ kg/m}^2)$ were not at risk of COVID-19 mortality, as compared with those with normal BMI ($18.5 \le BMI < 25 \text{ kg/m}^2$) or without obesity.58 When BMI was modelled as a continuous measure, conflicting reports were found such that every 5 units (kg/m^2) increment in BMI increased the risk of COVID-19 mortality in one study,⁵⁸ whereas a continuous BMI measure was not associated with the risk of COVID-19 mortality in another study.⁵⁹ As observed in our analysis, most original studies on obesity and the risk of COVID-19 mortality were conducted in the countries with the highest level of obesity (ie, the USA and most of the western world).^{13 60 61} Our results on the pooled adjusted risk ratios for obesity (BMI \geq 30 kg/m²) and the risk of COVID-19 mortality are consistent with the summary RR in published meta-analyses.^{21 53 57 58} Nevertheless, caution is warranted when interpreting the associations between obesity as measured by BMI and COVID-19 mortality across different populations because

Table 1 Characteristics of the studies included in the meta-analysis							
Characteristic		Studies, n (%)*					
		Total	Diabetes	Hypertension	Obesity		
Overall		145 (100.0%)	118 (100.0%)	99 (100.0%)	57 (100.0%)		
Total N							
Study period							
Start date	December 2019–April 2020	141 (97.2%)	114 (96.6%)	96 (97.0%)	55 (96.5%)		
	May 2020–November 2020	4 (2.8%)	4 (3.4%)	3 (3.0%)	2 (3.5%)		
End date	February 2020–April 2020	84 (57.9%)	65 (55.1%)	47 (47.5%)	31 (54.4%)		
	May 2020–November 2020	61 (42.1%)	53 (44.9%)	52 (52.5%)	26 (45.6%)		
Sample size							
	Median	1000	1336	1157	2015		
	IQR	5053	6953	6964	10 117		
	95 to <1000	72 (49.7%)	52 (44.1%)	46 (46.5%)	24 (42.1%)		
	1000 to <10 000	47 (32.4%)	41 (34.7%)	33 (33.3%)	19 (33.3%)		
	≥10 000	26 (17.9%)	25 (21.2%)	20 (20.2%)	14 (24.6%)		
Mean or median age (years)							
	<60	62 (42.8%)	56 (47.5%)	47 (47.5%)	28 (49.1%)		
	≥60	83 (57.2%)	62 (52.5%)	52 (52.5%)	29 (50.9%)		
Male (%)							
	<50	41 (28.3%)	36 (30.5%)	24 (24.2%)	12 (21.1%)		
	≥50	104 (71.7%)	82 (69.5%)	75 (75.8%)	45 (78.9%)		
Study design							
	Cohort	115 (79.3%)	90 (76.3%)	78 (78.8%)	44 (77.2%)		
	Cross-sectional	28 (19.3%)	27 (22.9%)	21 (21.2%)	12 (21.1%)		
	Case-control	2 (1.4%)	1 (0.8%)	0 (0.0%)	1 (1.8%)		
Type of effect estimate							
	HR	58 (40.0%)	48 (40.7%)	35 (35.4%)	23 (40.4%)		
	OR	78 (53.8%)	63 (53.4%)	60 (60.6%)	30 (52.6%)		
	Relative risk (RR)	9 (6.2%)	7 (5.9%)	4 (4.0%)	4 (7.0%)		
Data source							
	Electronic health (medical) records	84 (57.9%)	63 (53.4%)	57 (57.6%)	26 (45.6%)		
	Administrative, registry, surveillance systems	49 (33.8%)	45 (38.1%)	34 (34.3%)	27 (47.4%)		
	Other†	12 (8.3%)	10 (8.5%)	8 (8.1%)	4 (7.0%)		
NOS score							
	8–9	107 (73.8%)	84 (71.2%)	71 (71.7%)	42 (73.7%)		
	5–7	34 (23.4%)	31 (26.3%)	26 (26.3%)	15 (26.3%)		
	<5	4 (2.8%)	3 (2.5%)	2 (2.0%)	0 (0.0%)		
Funding source							
	Industry funded	2 (1.4%)	1 (0.8%)	2 (2.0%)	1 (1.8%)		
	Independently funded	68 (46.9%)	57 (48.3%)	46 (46.5%)	24 (42.1%)		
	None or NA	45 (31.0%)	37 (31.4%)	28 (28.3%)	20 (35.1%)		
	Not reported	30 (20.7%)	23 (19.5%)	23 (23.2%)	12 (21.1%)		
WHO region							
	Africa	3 (2.1%)	2 (1.7%)	3 (3.0%)	1 (1.8%)		
					O		

Continued

Characteristic		Studies, n (%	⁄o)*		
	Americas-USA	40 (27.6%)	31 (26.3%)	29 (29.3%)	18 (31.6%
	Americas-outside USA	15 (10.3%)	15 (12.7%)	12 (12.1%)	14 (24.6%
	East Mediterranean	9 (6.2%)	9 (7.6%)	5 (5.1%)	2 (3.5%)
	Europe	42 (29.0%)	32 (27.1%)	27 (27.3%)	20 (35.1%
	South-East Asia	3 (2.1%)	2 (1.7%)	3 (3.0%)	0 (0.0%)
	Western Pacific-inclusive mainland China	23 (15.9%)	18 (15.3%)	15 (15.2%)	0 (0.0%)
	Western Pacific-exclusive mainland China	9 (6.2%)	8 (6.8%)	4 (4.0%)	2 (3.5%)
	Worldwide	1 (0.7%)	1 (0.8%)	1 (1.0%)	0 (0.0%)
WB income level					
	High	93 (64.1%)	73 (61.9%)	63 (63.6%)	42 (73.7%
	Upper middle	47 (32.4%)	41 (34.7%)	31 (31.3%)	14 (24.6%
	Lower middle	4 (2.8%)	3 (2.5%)	4 (4.0%)	1 (1.8%)
	Worldwide	1 (0.7%)	1 (0.8%)	1 (1.0%)	0 (0.0%)
Health Index Scor	e				
	≥80	66 (45.5%)	51 (43.2%)	41 (41.4%)	21 (36.8%
	70–79	72 (49.7%)	62 (52.5%)	51 (51.5%)	35 (61.4%
	<70	5 (3.4%)	3 (2.5%)	5 (5.1%)	1 (1.8%)
GHSI score					
	Most prepared (≥66.7)	46 (31.7%)	35 (29.7%)	34 (34.3%)	20 (35.1%
	More prepared (33.4–66.6)	95 (65.5%)	79 (66.9%)	61 (61.6%)	36 (63.2%
	Least prepared (0-33.3)	2 (1.4%)	2 (1.7%)	2 (2.0%)	1 (1.8%)

diabetes and obesi †Other types of data source include paper medical records, manual data collection and unspecified medical charts or records.

GHSI, Global Health Security Index; NOS, Newcastle-Ottawa Scale; WB, World Bank.

of the ethnic differences in BMI and its associations with disease risks.^{62–64}

As compared with the number of original studies included for diabetes and hypertension, we identified fewer studies for obesity, with several possible reasons. First, obesity was not recognised as a risk factor for COVID-19 mortality at the early stage of the pandemic,^{65 66} therefore, few studies reported results for obesity in the countries at the early pandemic.⁶⁵ ⁶⁶ Second, countries with a lower prevalence of obesity might be less likely to report data due to insufficient number of deaths by obesity status. It is evident in this study that few studies on obesity were identified in Asia and Africa. Third, various BMI scales used in the studies could make it difficult to compare results across studies or countries and synthesise data in meta-analyses. For example, whereas many studies used BMI \geq 30 kg/m² to define obesity (ie, overall obesity or unclassified obesity), a few studies used BMI as classified categories (ie, underweight: <18.5 kg/m², normal weight: 18.5 to $<25 \text{ kg/m}^2$, overweight: 25 to $<30 \text{ kg/m}^2$ m^2 , obesity class I: 30 to <35 kg/m², obesity class II: 35 to $<40 \text{ kg/m}^2$ and obesity class III: $\geq 40 \text{ kg/m}^2$) or a continuous scale.⁵⁹ Fourth, missing data on BMI in electronic

health (medical) record systems are common.⁶⁷ Fifth, it is possible that insignificant or negative results for obesity, particularly in small studies, might not be published or reported as suggested by the possible publication bias detected in our analysis.

Our pooled adjusted risk ratios suggest that patients with diabetes and obesity had about a 40% increased risk for COVID-19 mortality and those with hypertension about a 20% increased risk, independent of other known risk factors. While mechanisms for the increased risk of COVID-19 mortality in individuals with diabetes, hypertension and obesity remain elusive, our findings provide further motivation to support research on the underlying pathophysiology. Available laboratory and clinical studies suggest that overexpression of ACE2 in adipose tissue, impaired immune function, increased proinflammatory response and cytokine storm might play critical roles in the severity and mortality of COVID-19 in patients with diabetes, hypertension and obesity.^{68–70} Emerging evidence showed that SARS CoV-2 infection could increase the risk of developing new onset diabetes among survivors.^{71 72} The relationship between SARS CoV-2 infection and new onset diabetes is complex, however,



Pooled risk ratios (95% confidence intervals)

Figure 3 Overall pooled adjusted risk ratios for the associations of diabetes, hypertension and obesity with COVID-19 mortality. HR, hazard ratio. OR, odds ratio. RR, relative risk or risk ratio.

and not only is acquiring the virus associated with more severe outcomes,^{2 50} but a large and increasing body of epidemiological evidence shows an increase in diabetes incidence following infection.^{71 72} This is consistent with laboratory evidence showing that the virus infects and can kill pancreatic beta cells.⁷³

The elevated mortality risk among COVID-19 patients with comorbidities, particularly among those with uncontrolled diabetes or hypertension, suggests a correlation between prepandemic levels of control and the impact of these conditions on COVID-19 outcomes.^{74 75} Countries with better healthcare quality often have a higher proportion of individuals with controlled diabetes and hypertension. This could imply that variations in prepandemic control levels across countries play a role in COVID-19 mortality rates among those with comorbidities.

Although the differences in the strength of associations of diabetes, hypertension and obesity with COVID-19 mortality we observed across regions were lower than anticipated given the known differences in the control of these chronic conditions and quality of health services, they are still intriguing and appeared to be related to the timeline of COVID-19 spreading and virus strain mutations across countries or regions.⁷⁶ As the first country where the outbreak occurred, China had the strongest associations, followed by South Korea, European region, East Mediterranean region, South-East Asian region, followed by North America. One of the explanations for this could be improved knowledge of treating COVID-19 patients. Our study included articles published in the entire year of 2020, covering the initial months of the pandemic. Potential differences in the treatment of COVID-19 might be attributed to the evolving understanding of the condition and the identification of effective therapeutic options.⁷⁷ As the pandemic progressed, individuals affected later on received more informed care, especially regarding treating individuals with comorbidities.^{78–80} Another explanation could be the notion of a'quality penalty' imposed by overburdened healthcare services occurred early in the pandemic, where the benefits of treatment at high-quality facilities are diminished when the system is overwhelmed.⁸¹ Other factors, sometimes outside of pandemic preparedness efforts, such as adequacy and resiliency of healthcare systems could act as effect modifiers on the strength of observed association across countries or regions.

One of the interesting results in our study is the inverse association between the higher GHSI score and the lower

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Pooled risk ratios of the associations between underlying conditions and mortality: subgroup analysis

Pooled risk ratios (95% confidence intervals)

Figure 4 Pooled adjusted risk ratios for the associations of diabetes, hypertension and obesity with COVID-19 mortality by subgroups. GHSI, Global Health Security Index. AFR, Africa. AMR, Americas. EMR, Eastern Mediterranean. EUR, Europe. SEAR, South-East Asia. USA, United States of America. WB, World Bank. WHO, World Health Organization. WPR, Western Pacific.

strength in the associations of diabetes, hypertension and obesity with COVID-19 mortality. The GHSI is the first comprehensive assessment of countries' preparedness for infectious disease outbreaks such as COVID-19 based on the health security and related capabilities of 195 States Parties to the WHO 2005 International Health Regulations (IHR).⁴⁴ Our results were consistent with findings reported by others that higher country GHSI scores were associated with reduced deaths from communicable diseases (a composite of diarrhoeal disease, HIV, lower respiratory infection, meningitis and tuberculosis)⁸² and that greater levels of preparedness were associated with lower excess COVID-19 mortality after accounting for under-reporting and age structure.⁸³ Collectively, these findings suggest that GHSI could be a measure for the capacity of overall healthcare system readiness, emergency medical response and critical care for illness that can progress in severity such as COVID-19 when risk is amplified by comorbidities such as diabetes, hypertension and obesity. Indeed, based on the global experience of COVID-19, the Monitoring and Evaluation Framework of the IHR was updated in 2021 to integrate health

systems strengthening and health equity. Previously focused mainly on infection prevention and control, the updates recognise the importance of ensuring the provision of essential health services before, during and after an emergency to foster overall health system resilience.⁸⁴

The major strengths of this systematic review and meta-analysis were its comprehensiveness and rigour. It involved searching 16 literature bases and obtaining a large number of eligible studies. While the majority of articles found in our literature review are in English, eight articles in Chinese, French, Italian, Persian, Russian, Spanish and Turkish were also identified, translated into English, and reviewed by two or more researchers to minimise possible omission of published original studies. The large number of studies enabled us to assess variations in subgroups by study-level and country-level characteristics as well as across all seven WHO regions. There were also several limitations in this study. First, all original studies included in this study were observational studies; therefore, the presence of information bias is possible, particularly due to the inclusion of studies relying on self-reports and retrospective data. However, the recall bias would be

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 Table 2
 Meta-regression analysis* on the effect estimates for the associations of diabetes, hypertension and obesity with

 COVID-19 mortality by study-level and country-level indicators

Study-level and country- level indicators	Diabetes		Hypertension		Obesity	
		Р		Р		Ρ
	β (95% CI)†	value‡	β (95% CI)	value	β (95% CI)	value
Mean, age, years	-0.01 (-0.02 to -0.001)	0.02	-0.01 (-0.02 to -0.001)	0.03	-0.00 (-0.01 to 0.01)	0.34
Men, %	-0.00 (-0.01 to 0.001)	0.23	-0.00 (-0.01 to 0.01)	0.74	-0.00 (-0.01 to 0.01)	0.43
Study starting date, month	-0.03 (-0.09 to 0.02)	0.20	-0.03 (-0.08 to 0.03)	0.30	-0.03 (-0.08 to 0.02)	0.28
NOS score	-0.03 (-0.10 to 0.04)	0.37	-0.02 (-0.10 to 0.06)	0.64	0.01 (-0.09 to 0.11)	0.85
Health Index Score, 2019	0.02 (0.01 to 0.04)	0.003	0.01 (-0.01 to 0.02)	0.21	0.00 (-0.02 to 0.02)	0.71
GHSI score, 2019	-0.01 (-0.01 to -0.001)	0.02	-0.01 (-0.01 to -0.001)	0.04	-0.01 (-0.02 to -0.001)	0.001

*Meta-regression was conducted to assess the linear relationship between the explanatory variables (continuous study-level and countrylevel indicators) and the outcome variables (effect estimates) using a random-effects method.

†The regression coefficient (β) and 95% CI describe how the outcome variable (the effect estimate) changes with a unit increase in the explanatory variable (potential moderation effect).

 \ddagger The statistical significance (p value) of the regression coefficient is a test of whether there is a linear relationship between the explanatory variable and the outcome variable. Bold values indicate P<0.05.

NOS, Newcastle-Ottawa Scale; GHSI, Global Health Security Index.

expected to be minimal as data from electronic health (medical) records were used for most studies included in this meta-analysis. Second, although we focused on the use of adjusted risk ratios in our meta-analyses, residual confounding might be possible because some unobserved variables might not have been included in the original studies. Third, our meta-analyses relied on the adjusted risk ratios available in studies that used different sets of covariates, which might have contributed to the variations observed. Fourth, about half of the studies used OR as the risk measure, which could overestimate the associations. However, OR can be used approximately as an approximate measure of risk given the low mortality rate for COVID-19.^{33 34} Fifth, we adapted the NOS tool as a method to assess the quality or risk of bias of included studies. Due to the lack of a universally standardised scoring method, the NOS score for the individual study assessed in our study might differ from that in other similar analyses. The scores were produced by two researchers independently, and disagreement between two independent researchers was resolved by group discussion or by a third researcher, which would be expected to minimise the possibility of bias in quality assessment. Finally, our findings were limited to the studies published at the early phase of COVID-19 pandemic with highly publicised Alpha (B.1.1.7), Beta (B.1.351) and Gamma (P.1) variants of SARS-CoV-2 virus by the end of 2020. Future studies would be helpful to examine these associations in the later phases of COVID-19 pandemic with Delta (B.1.617.2) variant that hit hard in the spring of 2021 and Omicron (BA.1) variant that was identified in late November 2021 and overtook Delta as the dominant variant.85

Although diabetes, hypertension and obesity have been linked clinically with mechanistic and cellular plausibility,^{86 87} few studies have assessed the effects of the combination of these three comorbidities on the risk of COVID-19 mortality perhaps due to insufficient sample size. A large study from Mexico reporting all possible combinations of three comorbidities suggested that patients having two or three comorbidities could have increased risk for COVID-19 mortality compared with those with only one chronic condition.⁸⁸ As diabetes, hypertension and obesity are inter-related and increasingly prevalent conditions globally,^{11–13} integration of communicable and NCD prevention and treatment services could be a strategic measure to lessen the impact of future pandemics.^{7–9}

CONCLUSION

Our systematic review and meta-analysis suggests that patients with diabetes and those with obesity had about a 40% increased risk for COVID-19 mortality, while those with hypertension had a 20% increased risk, independent of other known risk factors for COVID-19 mortality. Our findings motivate further research into the underlying pathophysiology of the associations. The independent associations of diabetes, hypertension and obesity with COVID-19 mortality support the need for intervention and management of these chronic conditions to mitigate the risk of mortality from respiratory pathogens and other infectious agents. The significant differences in the strength of associations across countries or regions and by the GHSI scores highlight the importance of readiness and preparedness of healthcare systems, medical resources, clinical care provision and capacity. Healthcare systems need to be integrated and resilient enough that they can not only







Figure 5 Contour-funnel plots of meta-analyses for the associations of diabetes (DM, A), hypertension (HTN, B), and obesity (OB, C) with COVID-19 mortality. Yellow region = P < 0.01, light orange region = $0.01 \le P < 0.05$, dark orange region = $0.05 \le P < 0.1$, white region = $P \ge 0.1$. The vertical dashed line represents the overall pooled risk ratio. The diagonal dashed lines show the expected 95% confidence intervals around the overall pooled risk ratio. Each dot represents the effect size of a study.

react to emergencies but can proactively adapt so they are prepared to provide quality healthcare in every situation. Addressing the increasing burden of diabetes, obesity and hypertension is important both for the prevention of NCDs and for the resilience of populations in the face of pandemics, particularly those in low- and middle-income countries where healthcare access and resources can vary greatly.

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Supplementary Materials

Abbreviations used in the supplementary materials:

AFR = African Region AMR = American Region C-C = case-controlCOSMOS-E = Conducting Systematic Reviews and Meta-Analyses of Observational Studies of Etiology C-S = cross-sectionalDM = diabetes mellitus EHR = electronic health (medical) record EMR = East Mediterranean Region ES = effect sizeEUR = European Region GHSI = Global Health Security Index HI = high incomeHR = hazard ratio HTN = hypertension LMI = lower middle income MOOSE = meta-analyses Of Observational Studies in Epidemiology NOS = Newcastle-Ottawa Scale Ob = obesityOR = odds ratioPRR = pooled risk ratio PRISMA = Preferred Reporting Items for Systematic Reviews and Meta-Analyses RR = relative risk or risk ratio ARS = administrative/registry/surveillance or (case) reporting system SEAR = Southeast Asian Region UK = United Kingdom UMI = upper middle income USA = United States of America WB = World BankWHO = World Health Organization WPR = West Pacific Region

Supplementary Text 1 Search Strategy

Time period: December 1st, 2019, through December 31st, 2020.

Key words or terms:

- 1. (COVID-19 and all possible variations) AND
- 2. (Diabetes, obesity, hypertension, and all relevant terms) OR
- 3. (Comorbidity, comorbid disease or illness or condition, underlying disease or illness or condition, chronic disease or illness or condition, noncommunicable disease or NCD, predictor, risk or risk factor, determinant, cardiovascular, and metabolic).

No restrictions in language, gender, age, publication types.

Databases: all 16 databases.

Database	Strategy	Records	Update 09/16/2020	Update 01/15/2021
Medline (OVID) 1946-	novel coronavir* OR novel corona virus* OR 2019 coronavirus OR coronavirus disease OR coronavirus 2019 OR betacoronavir* OR covid19 OR covid 19 OR nCoV OR novel CoV OR CoV 2 OR CoV2 OR sarscov2 OR sarscov OR sarscov OR 2019nCoV OR 2019-nCoV OR wuhan virus* OR ((wuhan OR hubei OR huanan) AND (severe acute respiratory OR pneumonia*) AND outbreak* AND (201912*.dt OR 2020*.dt)) OR ((coronavirus OR pandemic).mp AND (201912*.dt OR 2020*.dt))	5856	1586	7932
	Diabetes OR diabetic OR blood glucose OR glyc?emic control OR glucose control OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR waist circumference OR BMI OR body mass index OR hypertension OR hypertensive OR high blood pressure OR comorbid* OR co-morbid* OR pre- existing OR preexisting OR underlying OR chronic disease* OR chronic illness* OR chronic condition* OR noncommunicable disease* OR cardiovascular disease* OR predictor* OR determinant* OR risk factor* OR metabolic			
Embase (OVID) 1988-	(novel coronavir* OR novel corona virus* OR 2019 coronavirus OR coronavirus disease OR coronavirus 2019 OR betacoronavir* OR covid19 OR covid 19 OR nCoV OR novel CoV OR CoV 2 OR CoV2 OR sarscov2 OR sars-cov OR sarscov OR 2019nCoV OR 2019-nCoV OR wuhan virus*) OR ((wuhan OR hubei OR huanan) AND (severe acute respiratory OR pneumonia*) AND outbreak*) OR ((coronavirus OR pandemic).mp AND 2020*.dc) AND Diabetes OR diabetic OR blood glucose OR glyc?emic control OR glucose control OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR waist circumference OR BMI OR body mass index OR hypertension OR hypertensive OR high blood pressure OR comorbid* OR co-morbid* OR pre- existing OR preexisting OR underlying OR chronic disease* OR cardiovascular disease* OR predictor* OR determinant* OR risk factor* OR metabolic Not pubmed/medline	6461 -4050 duplicates =2411 unique items	2816 -1677 duplicates =1139 unique items	11477 -5134 duplicates =6343 unique items
Global Health (OVID)	(novel coronavir* OR novel corona virus* OR 2019 coronavirus OR coronavirus disease OR coronavirus 2019 OR betacoronavir* OR covid19 OR covid 19 OR nCoV OR novel CoV OR CoV 2 OR CoV2 OR sarscov2 OR sars-cov OR sarscov OR 2019nCoV OR 2019-nCoV OR wuhan virus*) OR (((wuhan OR hubei OR huanan) AND (severe acute respiratory OR pneumonia*) AND outbreak*) AND 2020*.up) OR ((coronavirus OR pandemic).mp AND 2020*.up) AND Diabetes OR diabetic OR blood glucose OR glyc?emic control OR glucose control OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR waist circumference OR BMI OR body mass index OR hypertension OR hypertensive OR high blood pressure OR comorbid* OR co-morbid* OR pre- existing OR preexisting OR underlying OR chronic disease* OR chronic illness*	1102 -744 duplicates =358 unique items	273 -107 duplicates =166 unique items	3225 -1597 duplicates =1628 unique items

		1		
	OR chronic condition* OR noncommunicable disease* OR cardiovascular disease* OR predictor* OR determinant* OR risk factor* OR metabolic			
CAB Abstracts	(novel coronavir* OR novel corona virus* OR 2019 coronavirus OR coronavirus	501	125	685
(OVID)	disease OR coronavirus 2019 OR betacoronavir* OR covid19 OR covid 19 OR nCoV OR novel CoV OR CoV 2 OR CoV2 OR sarscov2 OR sarscov OR sarscov	-463	-121	-669
	OR 2019nCoV OR 2019-nCoV OR wuhan virus*) OR ((wuhan OR hubei OR	duplicates	duplicates	duplicates
	huanan) AND (severe acute respiratory OR pneumonia*) AND outbreak*) OR ((coronavirus OR pandemic) mp AND 2020* up)	=38	=4	=16
	AND	unique items	unique items	unique items
	Diabetes OR diabetic OR blood glucose OR glyc?emic control OR glucose control			
	OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR			
	adipos* OR waist circumference OR BMI OR body mass index OR hypertension			
	existing OR preexisting OR underlying OR chronic disease* OR chronic illness*			
	OR chronic condition* OR noncommunicable disease* OR cardiovascular disease*			
PsycInfo	(novel coronavir* OR novel corona virus* OR 2019 coronavirus OR coronavirus	159	74	609
(OVID) 1087	disease OR coronavirus 2019 OR betacoronavir* OR covid 19 OR covid 19 OR	106	47	254
1907-	OR 2019nCoV OR 2019-nCoV OR wuhan virus*) OR (((wuhan OR hubei OR	duplicates	duplicates	duplicates
	huanan) AND (severe acute respiratory OR pneumonia*) AND outbreak*) AND 2020* up) OR ((coronavirus OR pandemic) mp AND 2020* up)	-53	-27	-355
	AND	unique items	unique items	unique items
	Diabetes OR diabetic OR blood glucose OR gluce?emic control OR glucose control			
	OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR			
	adipos* OR waist circumference OR BMI OR body mass index OR hypertension OR hypertensive OR high blood pressure OR comorbid* OR co-morbid* OR pre-			
	existing OR preexisting OR underlying OR chronic disease* OR chronic illness*			
	OR chronic condition* OR noncommunicable disease* OR cardiovascular disease*			
CINAHL	("novel coronavir*" OR "novel corona virus*" OR "2019 coronavirus" OR	1668	264	1225
(EbscoHost)	betacoronavir* OR covid19 OR "covid 19" OR nCoV OR "novel CoV" OR "CoV 2" OR CoV2 OR sarscov2 OR sarscov OR sarscov OR 2019nCoV OR 2019-nCoV	-602	-105	-569
	OR "wuhan virus*") OR (((wuhan OR hubei OR huanan) AND ("severe acute	duplicates	duplicates	duplicates
	respiratory" OR pneumonia*) AND outbreak*) AND PY 2020) OR ((coronavirus OR pandemic) AND PY 2020)	=766	=259	=656
	AND	unique items	unique items	unique items
	Diabetes OR diabetic OR "blood glucose" OR "glyc?emic control" OR "glucose			
	control" OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR			
	OR hypertension OR hypertensive OR "high blood pressure" OR comorbid* OR			
	co-morbid* OR pre-existing OR preexisting OR underlying OR "chronic disease*"			
	OR "cardiovascular disease*" OR predictor* OR determinant* OR "risk factor*"			
	OR metabolic			
Andomia	Exclude Medline records	1644	617	2595
Research	betacoronavir* OR covid19 OR "covid 19" OR nCoV OR "novel CoV" OR "CoV	1044	047	2383
Complete	2" OR CoV2 OR sarscov2 OR sarscov OR sarscov OR 2019nCoV OR 2019-nCoV	-1096 duplicatos	-448 duplicates	-1979 duplicates
	respiratory" OR pneumonia*) AND outbreak*) AND PY 2020) OR ((coronavirus	uupiteates	uplicates	uupiteates
	OR pandemic) AND PY 2020)	=548	=199	=606
	AND	unique nems	unique nemis	unique nems
	Diabetes OR diabetic OR "blood glucose" OR "glyc?emic control" OR "glucose control" OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR			
	overweight OR adipos* OR "waist circumference" OR BMI OR "body mass index"			
	OR hypertension OR hypertensive OR "high blood pressure" OR comorbid* OR co-morbid* OR pre-existing OR preexisting OR underlying OR "chronic disease*"			
	OR "chronic illness*" OR "chronic condition*" OR "noncommunicable disease*"			
	OR "cardiovascular disease*" OR predictor* OR determinant* OR "risk factor*" OR metabolic			
Africa Wide	("novel coronavir*" OR "novel corona virus*" OR "2019 coronavirus" OR	6	0	15
Information	betacoronavir* OR covid19 OR "covid 19" OR nCoV OR "novel CoV" OR "CoV 2" OR CoV2 OR sarscov2 OR sars-cov OR sarscov OR 2019nCoV OR 2019-nCoV	-1		-3
	OR "wuhan virus*") OR (((wuhan OR hubei OR huanan) AND ("severe acute	duplicates		duplicates
	respiratory" OR pneumonia*) AND outbreak*) AND PY 2020) OR ((coronavirus OR pandemic) AND PY 2020)	=5		=11
	AND	unique items		unique items

	Diabetes OR diabetic OR "blood glucose" OR "glyc?emic control" OR "glucose control" OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR "waist circumference" OR BMI OR "body mass index" OR hypertension OR hypertensive OR "high blood pressure" OR comorbid* OR co-morbid* OR pre-existing OR preexisting OR underlying OR "chronic disease*" OR "chronic illness*" OR "chronic condition*" OR "noncommunicable disease*" OR "catiovascular disease*" OR predictor* OR determinant* OR "risk factor*"			
Scopus	TITLE-ABS("novel coronavir*" OR "novel corona virus*" OR "2019 coronavirus" OR betacoronavir* OR covid19 OR "covid 19" OR ncov OR "CoV 2" OR cov2 OR sarscov2 OR sars-cov OR sarscov OR 2019ncov OR 2019-nCoV OR "novel CoV" OR "wuhan virus") AND TITLE-ABS(Diabetes OR diabetic OR "blood glucose" OR "glyc?emic control" OR "glucose control" OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR "waist circumference" OR BMI OR "body mass index" OR hypertension OR hypertensive OR "high blood pressure" OR comorbid* OR co-morbid* OR pre-existing OR preexisting OR underlying OR "chronic disease*" OR "chronic illness*" OR "chronic condition*" OR "noncommunicable disease*" OR "cardiovascular disease*" OR predictor* OR determinant* OR "risk factor*" OR metabolic)	4021 -3551 duplicates =470 unique items	94 -73 duplicates =21 unique items	1038 -702 duplicates =336 unique items
РМС	 ("novel coronavir*"[Title/Abstract] OR "novel corona virus*"[Title/Abstract] OR "2019 coronavirus"[Title/Abstract] OR "betacoronavir*"[Title/Abstract] OR "covid19"[Title/Abstract] OR "covid19"[Title/Abstract] OR "cov2"[Title/Abstract] OR "cov2"[Title/Abstract] OR "cov2"[Title/Abstract] OR "cov2"[Title/Abstract] OR "cov2"[Title/Abstract] OR "sars-cov"[Title/Abstract] OR "sars-cov"[Title/Abstract] OR "2019-nCoV"[Title/Abstract] OR "novel CoV"[Title/Abstract] OR "sars-cov"[Title/Abstract] OR "sars-cov"[Title/Abstract] OR "2019-nCoV"[Title/Abstract] OR "novel CoV"[Title/Abstract] OR "wuhan virus"[All Fields]) AND Diabetes[Title/Abstract] OR diabetic[Title/Abstract] OR "blood glucose" [Title/Abstract] OR "glyc?emic control" [Title/Abstract] OR "glucose control" [Title/Abstract] OR myperglyc?emia[Title/Abstract] OR blood glucose" [Title/Abstract] OR hyperglyc?emia[Title/Abstract] OR adipos*[Title/Abstract] OR "waist circumference" [Title/Abstract] OR BMI[Title/Abstract] OR "body mass index"[Title/Abstract] OR hypertension[Title/Abstract] OR "hypertensive[Title/Abstract] OR "high blood pressure"[Title/Abstract] OR comorbid*[Title/Abstract] OR "chronic disease*"[Title/Abstract] OR "chronic disease*"[Title/Abst	918 -676 duplicates =242 unique items	243 -172 duplicates =71 unique items	791 -727 duplicates =64 unique items
ProQuest Central	TI,AB("novel coronavir*" OR "novel corona virus*" OR "2019 coronavirus" OR betacoronavir* OR covid19 OR "covid 19" OR nCoV OR "novel CoV" OR "CoV 2" OR CoV2 OR sarscov2 OR sarscov OR sarscov OR 2019nCoV OR 2019- nCoV) AND TI,AB(Diabetes OR diabetic OR "blood glucose" OR "glyc?emic control" OR "glucose control" OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR "waist circumference" OR BMI OR "body mass index" OR hypertension OR hypertensive OR "high blood pressure" OR comorbid* OR co-morbid* OR pre-existing OR preexisting OR underlying OR "chronic disease*" OR "chronic illness*" OR "chronic condition*" OR "noncommunicable disease*" OR "cardiovascular disease*" OR predictor* OR determinant* OR "risk factor*" OR metabolic)	1238 -844 duplicates =394 unique items	483 -339 duplicates =144 unique items	1256 -760 duplicates =496 unique items
SBT COVID-19 Library This library covers (PrePrints - Medrxiv, BIOrxiv, Chemrxiv, SSRN, Scielo -, WHO COVID-19 database, Homeland	Diabetes OR diabetic OR "blood glucose" OR "glyc?emic control" OR "glucose control" OR hyperglyc?emia OR hypoglyc?emia OR obesity OR obese OR overweight OR adipos* OR "waist circumference" OR BMI OR "body mass index" OR hypertension OR hypertensive OR "high blood pressure" OR comorbid* OR co-morbid* OR pre-existing OR preexisting OR underlying OR "chronic disease*" OR "chronic illness*" OR "chronic condition*" OR "noncommunicable disease*" OR "cardiovascular disease*" OR predictor* OR determinant* OR "risk factor*" OR metabolic	Preprints = 1602 WHO = 913 HLSC = 25 SciFinder = 82	Preprints = 26 WHO = 0 HLSC = 0 SciFinder =0 Clinicaltrials = 384	No longer being updated

Security COVID-	Clinicaltrials		
19 collection,	= 326		
SciFinder,			
Clinicaltrials)			
Total	13646	3467	18443

Notes: Duplicates were identified using the Endnote automated "find duplicates" function with preference set to match on title, author, and year, and removed from your Endnote library. There will likely be additional duplicates found that Endnote was unable to detect.

Total records before removing duplicates = 35,556; total records after removing duplicates via Endnote 20 = 34,830; total records after further removing duplicates via Covidence = 30,586.

Supplementary Text 2

Adapted Newcastle-Ottawa Scale (NOS) for assessing the quality of nonrandomised studies in metaanalyses

A. CASE-CONTROL STUDIES

<u>Note</u>: A study can be awarded a maximum of one star for each numbered item within the Selection and Exposure components. A maximum of two stars can be given for the Comparability component.

Selection (Maximum 4 stars)

- 1) <u>Is the case definition adequate</u>?
 - a) yes, with independent validation*
 - b) yes, e.g., record linkage or based on self-reports
 - c) no description
- 2) <u>Representativeness of the cases</u>
 - a) consecutive or obviously representative series of cases*
 - b) potential for selection biases or not stated
- 3) Selection of Controls
 - a) community controls*
 - b) hospital controls
 - c) no description
- 4) Definition of Controls
 - a) no history of disease (endpoint)*
 - b) no description of source

Comparability (Maximum 2 stars)

- 1) Comparability of cases and controls on the basis of the design or analysis
 - a) study controls for age (the most important factor)*
 - b) study controls for age plus any additional factor** (This criterion could be modified to indicate specific control for a second important factor)
 - c) study does not control for any confounders or no information provided

Exposure (Maximum 3 stars)

- 1) Ascertainment of exposure
 - a) secure record (e.g., surgical records)*
 - b) structured interview where blind to case/control status*
 - c) interview not blinded to case/control status
 - d) written self-report or medical record only
 - e) no description
- 2) <u>Same method of ascertainment for cases and controls</u>
 - a) Yes*
 - b) no
- 3) <u>Non-Response rate</u>
 - a) same rate for both groups*

- b) non respondents described
- c) rate different and no designation

B. COHORT STUDIES

<u>Note</u>: A study can be awarded a maximum of one star for each numbered item within the Selection and Outcome components. A maximum of two stars can be given for the Comparability component.

Selection (Maximum 4 stars)

- 1) <u>Representativeness of the exposed cohort</u>
 - a) truly representative of the average (describe) in the community*
 - b) somewhat representative of the average _____ in the community*
 - c) selected group of users (e.g., nurses, volunteers)
 - d) no description of the derivation of the cohort
- 2) Selection of the non-exposed cohort
 - a) drawn from the same community as the exposed cohort*
 - b) drawn from a different source
 - c) no description of the derivation of the non-exposed cohort
- 3) Ascertainment of exposure
 - a) secure record (e.g., surgical records)*
 - b) structured interview*
 - c) written self-report
 - d) no description
- 4) Demonstration that outcome of interest was not present at start of study
 - a) Yes*
 - b) no

Comparability (Maximum 2 stars)

- 1) <u>Comparability of cohorts on the basis of the design or analysis</u>
 - a) study controls for _____ (select the most important factor)*
 - b) study controls for the most important factor plus any additional factor** (This criterion could be modified to indicate specific control for a second important factor)
 - c) study does not adjust for any relevant confounders/risk factors or no information provided

Outcome (Maximum 3 stars)

- 1) Assessment of outcome
 - a) independent blind assessment*
 - b) record linkage*
 - c) self-report
 - d) no description
- 2) <u>Was follow-up long enough for outcomes to occur</u>
 - a) yes (select an adequate follow up period for outcome of interest)*
 - b) no
- 3) Adequacy of follow up of cohorts
 - a) complete follow up all subjects accounted for*

- 8
- b) subjects lost to follow up unlikely to introduce bias small number lost > _____ % (select an adequate %) follow up, or description provided of those lost)*
- c) follow up rate < $\$ % (select an adequate %) and no description of those lost
- d) no statement

C. CROSS-SECTIONAL STUDIES

<u>Note</u>: This scale has been adapted from the Newcastle-Ottawa Quality Assessment Scale for case-control studies and cohort studies to provide quality assessment of cross-sectional studies. A study can be awarded a maximum of one star for each numbered item within the Selection component. A maximum of two stars can be given for the Comparability and Outcome components.

Selection (Maximum 4 stars)

- 1) Representativeness of the sample
 - a) truly representative of the average in the target population* (all subjects or random sampling)
 - b) somewhat representative of the average in the target group* (non-random sampling)
 - c) selected group of users/convenience sample.
 - d) no description of the derivation of the included subjects

2) Sample size

- a) justified and satisfactory (including sample size calculation)*
- b) not justified
- c) no information provided
- 3) Ascertainment of the exposure (risk factor)
 - a) Secure record (e.g., surgical record)*
 - b) structured interview*
 - c) written self-report
 - d) no description
- 4) Non-respondents
 - a) proportion of target sample recruited attains pre-specified target or basic summary of non-respondent characteristics in sampling frame recorded*
 - b) unsatisfactory recruitment rate, no summary data on non-respondents
 - c) no description of the response rate or the characteristics of the responders and the non-responders

Comparability (Maximum 2 stars)

- 1) Comparability of cohorts on the basis of the design or analysis
 - a) study controls for _____ (select the most important factor)*
 - b) study controls for the most important factor plus any additional factor** (This criterion could be modified to indicate specific control for a second important factor)
 - c) study does not adjust for any relevant confounders/risk factors or no information provided

Outcome (Maximum 3 stars)

1) Assessment of outcome

- a) independent blind assessment**
- b) record linkage*
- c) self-report
- d) no description
- 2) Statistical test
 - a) statistical test used to analyse the data clearly described, appropriate and measures of association presented including confidence intervals and probability level (p-value)*
 - b) statistical test is not appropriate, not described, or incomplete

Total NOS scores:8-9 stars: high quality or low risk of bias5-7 stars: moderate quality or moderate risk of bias

<5 stars: low quality or high risk of bias.

Supplementary Figures

Fig. S1.1 – Forest Plots for the Association of Diabetes with COVID-19 Mortality



Fig. S1.2 – Forest Plots for the Association of Hypertension with COVID-19 Mortality



Fig. S1.3 – Forest Plots for the Association of Obesity with COVID-19 Mortality



Fig. S2.1 – Influence Plot for the Association of Diabetes with COVID-19 Mortality



Fig. S2.2 – Influence Plot for the Association of Hypertension with COVID-19 Mortality

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Jackson_BR_USA_2020			Shi S CHN 2020		
Jain AC IND 2020			Sun H CHN 2020		
Jiang Y CHN 2020		1	1 Sun V CHN 2020	1 P	T I
Kaeuffer C FRA 2020		- <u>- </u>	Sul_1_CHN_2020_		
Kim DW KDB 2020			Tantoi_ST_USA_2020	•	
Kim SW KOR 2020			Thempson L ODD 2020		
Kim T 11Så 2020	L.		Inompson_3_GBR_2020_	-i -i	i
King E USA 2020			van_Gerwen_m_USA_2020_	•	t
Kung S IND 2020			wang_Z_USA_2020_		+
Kunai_5_IND_2020_			workinggroup_ESP_2020		
Lee_SG_KOR_2020	(*		•i	
Mancilla-Galindo J MEX 2020	1	•	Xie_J_USA-2020 _	••	
Martos-Benitez_FD_MEX_2021		•	Yang_Q_CHN_2020	•	
Mejia_F_PER_2020		•	Yu_C_CHN_2020		
	1.1	12	13	11 40	
	17.1	1.2	1.0	1.1 52	1.6

Fig. S2.3 – Influence Plot for the Association of Obesity with COVID-19 Mortality

Obesity and mortality (ini	nuence plot)	19
Ahlstrom_B_SWE_2020	deSouza_C_BRA_2020_	
Al-Salameh_A_FRA_2020b	Miller_J_USA_2020 _	.
Alguwalhes_AM_SAU_2020	Mirani_M_ITA_2020	
Almazeedi_S_KWT_2020	Mukherjee_V_USA_2020	
Baqui_P_BRA_2020	Nachega_JB_COG_2020	• <u>+</u>
Bellan M ITA 2020	Nakeshbandi_M_USA_2020 _	
Berenquer J ESP 2020	Pacheco-Pantoja_E_MEX_2020	•
Biscarini S ITA 2020	Palaiodimos_L_USA_2020	
Breland JY USA 2021	Parikh_R_USA_2020 _	
Carrillo-Vena ME MEX 2020	Parra-Bracamonte_GM_MEX_2020	
	Pena_JE_MEX_2020	
	Petiti NN USA 2020	
Dicasteinuovo A TIA 2020	Polverino_F_JTA_2020	
Filardo_TD_USA_2020	Posso_M_ESP_2020_	
Fried_MW_USA_2020	Prado-Galbarro_FJ_MEX_2020	
Giacomelli A ITA 2020	Reilev_M_DNK_2020 _	
Gil-Rodrigo_A_ESP_2020 _	Rodriguez-Gonzalez_CG_ESP_2020	
Gutierrez_J_MEX_2020	Rossi_AP_ITA_2020 _	
Hajifathalian_K_USA_2020 _	Rossi_PG_ITA_2020 _	· · · · · · · · · · · · · · · · · · ·
ez-Galdamez_DR_MEX_2020	Rottoli_M_ITA_2020 _	
Izurieta_HS_USA_2020	Saand AR USA 2020	
Kaeuffer_C_FRA_2020 _	Seiglie_J_USA_2020 _	
Kim_SW_KOR_2020	Singh_S_USA_2020	
Kim_SY_KOR_2020	Sousa_GJB_BRA_2020 _	- <u> </u>
Lanini_S_ITA_2020 _	Tavares_C_BRA_2020 _	
ancilla-Galindo_J_MEX_2020	van_Gerwen_M_USA_2020 _	
artos-Benítez_FD_MEX_2021	Wang_Z_USA_2020 _	
McNeill JN USA 2020	Xie J USA-2020 _	
Mejia_F_PER_2020	Yazdanpanah_Y_FRA_2020	•
		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

Supplementary Tables

Table S1 – Overall Pooled Effect Estimates for the Association of Diabetes, Hypertension, and Obesity with

 COVID-19 Mortality

Exposure	Effect Estimate	N	PRR* (95% CI)	τ ² (95% CI)	<i>I</i> ² (95% CI)
Diabetes	Unadjusted	54	2.13 (1.80, 2.52)	0.31 (0.21, 0.53)	0.99 (0.99, 0.99)
Diabetes	Adjusted (overall)	118	1.43 (1.32, 1.54)	0.12 (0.10, 0.21)	0.94 (0.93, 0.95)
Diabetes	Adjusted (OR)	63	1.59 (1.40, 1.81)	0.17 (0.12, 0.34)	0.96 (0.95, 0.96)
Diabetes	Adjusted (HR/RR)	55	1.24 (1.15, 1.32)	0.02 (0.01, 0.11)	0.79 (0.73, 0.84)
Hypertension	Unadjusted	43	2.07 (1.74, 2.47)	0.28 (0.18, 0.47)	0.99 (0.99, 0.99)
Hypertension	Adjusted (overall)	99	1.19 (1.09, 1.30)	0.12 (0.09, 0.21)	0.91 (0.89, 0.92)
Hypertension	Adjusted (OR)	60	1.19 (1.04, 1.35)	0.17 (0.11, 0.33)	0.91 (0.89, 0.92)
Hypertension	Adjusted (HR/RR)	39	1.18 (1.06, 1.30)	0.06 (0.03, 0.16)	0.91 (0.89, 0.93)
Obesity	Unadjusted	28	1.45 (1.22, 1.71)	0.12 (0.07, 0.32)	0.85 (0.80, 0.89)
Obesity	Adjusted (overall)	57	1.39 (1.27, 1.52)	0.06 (0.04, 0.18)	0.96 (0.96, 0.97)
Obesity	Adjusted (OR)	30	1.46 (1.29, 1.65)	0.06 (0.03, 0.24)	0.98 (0.97, 0.98)
Obesity	Adjusted (HR/RR)	27	1.31 (1.14, 1.51)	0.06 0.03, 0.24)	0.77 (0.66, 0.84)

Note: * PRR = pooled risk ratio. τ^2 = a measure of between-study variance; I^2 = a statistic of the proportion of total variability explained by between-study variance. Estimates are from random-effects meta-analysis using restricted maximum likelihood method and Hartung-Knapp-Sidik-Jonkman adjusting for the standard errors.

HR = hazard ratio; OR = odds ratio; RR = relative risk.

Study- or Country- Level Variable	Subgroups	N	PRR* (95% CI)	τ ² (95% CI)	<i>I</i> ² (95% CI)
Type of risk ratio	OR	63	1.59 (1.40, 1.81)	0.17 (0.12, 0.34)	0.96 (0.95, 0.96)
	HR	48	1.26 (1.17, 1.36)	0.02 (0.01, 0.13)	0.73 (0.64, 0.79)
	RR	7	1.08 (0.94, 1.24)	0.01 (0.00, 0.15)	0.36 (0.00, 0.73)
Study period	May 2020 - November 2020	48	1.38 (1.26, 1.51)	0.07 (0.05, 0.15)	0.97 (0.96, 0.97)
	December 2019 - April 2020	70	1.48 (1.31, 1.68)	0.17 (0.12, 0.35)	0.82 (0.78, 0.86)
Study design	Cohort	90	1.37 (1.28, 1.46)	0.05 (0.04, 0.13)	0.79 (0.75, 0.83)
	Other	28	1.53 (1.20, 1.96)	0.31 (0.20, 0.76)	0.98 (0.97, 0.98)
Study quality	Low	8	1.56 (0.70, 3.49)	0.71 (0.26, 4.23)	0.87 (0.76, 0.93)
	Medium	54	1.50 (1.32, 1.70)	0.16 (0.11, 0.31)	0.96 (0.96, 0.97)
	High	56	1.30 (1.21, 1.40)	0.02 (0.01, 0.11)	0.79 (0.73, 0.84)
WHO region	EMR	9	1.31 (0.99, 1.72)	0.05 (0.00, 0.35)	0.38 (0.00, 0.72)
	EUR	32	1.35 (1.19, 1.53)	0.07 (0.04, 0.21)	0.93 (0.91, 0.94)
	AMR	46	1.27 (1.17, 1.37)	0.05 (0.03, 0.08)	0.95 (0.95, 0.96)
	AFR/SEAR	4	1.32 (0.81, 2.15)	0.00 (0.00, 2.11)	0.06 (0.00, 0.86)
	WPR	26	2.26 (1.76, 2.89)	0.24 (0.10, 0.57)	0.85 (0.80, 0.89)
WPR	-China	18	2.42 (1.64, 3.56)	0.41 (0.15, 1.05)	0.90 (0.85, 0.93)
	-South Korea	8	2.15 (1.79, 2.60)	0.00 (0.00, 0.15)	0.00 (0.00, 0.68)
WB income level	High	72	1.30 (1.21, 1.40)	0.05 (0.03, 0.11)	0.93 (0.92, 0.94)
	Upper middle	41	1.65 (1.38, 1.97)	0.22 (0.14, 0.49)	0.95 (0.94, 0.96)
	Lower middle	3	1.44 (0.47, 4.38)	0.07 (0.00, 8.80)	0.36 (0.00, 0.79)
Health index score tertile	1 st	39	1.29 (1.18, 1.41)	0.06 (0.03, 0.11)	0.96 (0.95, 0.97)
	2 nd	39	1.28 (1.17, 1.42)	0.04 (0.02, 0.12)	0.93 (0.91, 0.94)
	3 rd	38	1.87 (1.55, 2.27)	0.24 (0.13, 0.47)	0.86 (0.82, 0.89)
GHSI score tertile	1 st	39	1.66 (1.35, 2.04)	0.27 (0.15, 0.54)	0.89 (0.86, 0.92)
	2^{nd}	39	1.44 (1.29, 1.60)	0.07 (0.04, 0.18)	0.93 (0.91, 0.94)
	3 rd	38	1.24 (1.15, 1.35)	0.03 (0.02, 0.08)	0.96 (0.95, 0.97)

Table S2.1 – Pooled Effect Estimates for the Association between Diabetes and COVID-19 Mortality by

 Subgroups

Note: * PRR = pooled risk ratio. τ^2 = a measure of between-study variance; I^2 = a statistic of the proportion of total variability explained by between-study variance. Estimates are from random-effects meta-analysis using restricted maximum likelihood method and Hartung-Knapp-Sidik-Jonkman adjusting for the standard errors.

HR = hazard ratio; OR = odds ratio; RR = relative risk; GHSI = Global Health Security Index.

Study- or Country- Level Variable	Subgroups	N	PRR* (95% CI)	τ ² (95% CI)	I ² (95% CI)
Type of risk ratio	OR	60	1.19 (1.04, 1.35)	0.17 (0.11, 0.33)	0.91 (0.89, 0.92)
	HR	35	1.19 (1.06, 1.33)	0.06 (0.03, 0.18)	0.92 (0.89, 0.93)
	RR	4	1.06 (0.65, 1.73)	0.06 (0.00, 1.00)	0.67 (0.03, 0.89)
Study period	May 2020 - November 2020	46	1.12 (1.04, 1.21)	0.03 (0.02, 0.13)	0.86 (0.82, 0.89)
	December 2019 - April 2020	53	1.24 (1.07, 1.44)	0.21 (0.13, 0.36)	0.93 (0.91, 0.94)
Study design	Cohort	78	1.17 (1.07, 1.27)	0.07 (0.05, 0.18)	0.87 (0.84, 0.89)
	Other	21	1.22 (0.95, 1.56)	0.25 (0.13, 0.58)	0.96 (0.95, 0.97)
Study quality	Low	8	1.07 (0.60, 1.91)	0.29 (0.07, 2.25)	0.71 (0.40, 0.86)
	Medium	40	1.30 (1.10, 1.53)	0.20 (0.13, 0.40)	0.95 (0.94, 0.96)
	High	51	1.09 (1.01, 1.18)	0.02 (0.01, 0.10)	0.62 (0.49, 0.72)
WHO region	AMR	41	1.11 (1.02, 1.21)	0.04 (0.02, 0.13)	0.89 (0.85, 0.91)
	EUR	27	1.13 (0.99, 1.30)	0.07 (0.04, 0.21)	0.94 (0.92, 0.95)
	EMR	5	0.92 (0.58, 1.47)	0.00 (0.00, 0.54)	0.00 (0.00, 0.79)
	AFR/SEAR	6	1.20 (0.74, 1.92)	0.07 (0.00, 2.28)	0.59 (0.00, 0.83)
	WPR	19	1.52 (1.08, 2.15)	0.39 (0.17, 0.90)	0.92 (0.88, 0.94)
WPR	-China	15	1.57 (1.01, 2.45)	0.49 (0.20, 1.27)	0.93 (0.89, 0.95)
	-South Korea	4	1.30 (0.71, 2.37)	0.09 (0.00, 1.99)	0.64 (0.00, 0.88)
WB income level	High	62	1.07 (0.99, 1.16)	0.05 (0.03, 0.14)	0.90 (0.88, 0.92)
	Upper middle	31	1.42 (1.17, 1.71)	0.20 (0.11, 0.42)	0.91 (0.88, 0.93)
	Lower middle	4	1.06 (0.43, 2.60)	0.05 (0.00, 7.90)	0.47 (0.00, 0.82)
Health index score tertile	1 st	33	1.16 (1.05, 1.28)	0.04 (0.03, 0.15)	0.89 (0.86, 0.92)
	2 nd	32	1.02 (0.92, 1.13)	0.02 (0.01, 0.22)	0.60 (0.40, 0.73)
	3 rd	32	1.38 (1.12, 1.69)	0.24 (0.13, 0.46)	0.92 (0.89, 0.94)
GHSI score tertile	1 st	33	1.29 (1.03, 1.61)	0.28 (0.14, 0.52)	0.90 (0.87, 0.92)
	2 nd	32	1.22 (1.10, 1.36)	0.05 (0.03, 0.16)	0.92 (0.90, 0.94)
	3 rd	32	1.01 (0.93, 1.10)	0.02 (0.01, 0.19)	0.62 (0.44, 0.74)

Table S2.2 – Pooled Effect Estimates for the Association between Hypertension and COVID-19 Mortality by Subgroups

Note: * PRR = pooled risk ratio. τ^2 = a measure of between-study variance; I^2 = a statistic of the proportion of total variability explained by between-study variance. Estimates are from random-effects meta-analysis using restricted maximum likelihood method and Hartung-Knapp-Sidik-Jonkman adjusting for the standard errors.

HR = hazard ratio; OR = odds ratio; RR = relative risk; GHSI = Global Health Security Index.

Study- or Country- Level Variable	Subgroups	N	PRR* (95% CI)	τ ² (95% CI)	<i>I</i> ² (95% CI)
Type of risk ratio	HR	23	1.34 (1.13, 1.59)	0.09 (0.04, 0.33)	0.80 (0.71, 0.86)
	OR	30	1.46 (1.29, 1.65)	0.06 (0.03, 0.24)	0.98 (0.97, 0.98)
	RR	4	1.23 (1.03, 1.48)	0.00 (0.00, 0.07)	0.00 (0.00, 0.85)
Study period	May 2020 - November 2020	26	1.25 (1.14, 1.37)	0.03 (0.01, 0.11)	0.98 (0.98, 0.98)
	December 2019 - April 2020	31	1.61 (1.38, 1.87)	0.08 (0.04, 0.31)	0.72 (0.60, 0.80)
Study design	Cohort	44	1.43 (1.27, 1.62)	0.08 (0.05, 0.26)	0.91 (0.89, 0.93)
	Other	13	1.34 (1.18, 1.51)	0.03 (0.01, 0.12)	0.95 (0.93, 0.97)
Study quality	High	28	1.46 (1.21, 1.77)	0.12 (0.07, 0.51)	0.78 (0.69, 0.85)
	Medium	27	1.36 (1.24, 1.50)	0.04 (0.02, 0.08)	0.98 (0.98, 0.98)
	Low	2	-	-	-
WHO region	EUR	20	1.63 (1.32, 2.01)	0.10 (0.04, 0.44)	0.75 (0.61, 0.84)
	EMR	2	-	-	-
	AMR	32	1.31 (1.19, 1.45)	0.05 (0.03, 0.12)	0.98 (0.97, 0.98)
	WPR	2	-	-	-
	AFR/SEAR	1	-	-	-
WPR	China	0	-	-	-
	South Korea	2	-	-	-
WB income group	High	42	1.34 (1.18, 1.52)	0.07 (0.05, 0.28)	0.75 (0.66, 0.81)
	Upper middle	14	1.49 (1.33, 1.67)	0.03 (0.01, 0.09)	0.94 (0.91, 0.96)
	Lower middle	1	-	-	-
Health index score tertile	1 st	19	1.42 (1.27, 1.58)	0.04 (0.02, 0.09)	0.99 (0.98, 0.99)
	2 nd	19	1.17 (0.99, 1.38)	0.04 (0.02, 0.41)	0.64 (0.41, 0.78)
	3 rd	19	1.67 (1.36, 2.06)	0.07 (0.02, 0.41)	0.53 (0.20, 0.72)
GHSI score tertile	1 st	19	1.55 (1.26, 1.92)	0.06 (0.02, 0.60)	0.58 (0.31, 0.75)
	2 nd	19	1.54 (1.38, 1.72)	0.03 (0.01, 0.14)	0.91 (0.88, 0.94)
	3 rd	19	1.09 (1.01, 1.17)	0.00 (0.00, 0.19)	0.63 (0.40, 0.78)

Table S2.3 – Pooled Effect Estimates for the Association between Obesity and COVID-19 Mortality by

 Subgroups

Note: * PRR = pooled risk ratio. τ^2 = a measure of between-study variance; I^2 = a statistic of the proportion of total variability explained by between-study variance. Estimates are from random-effects meta-analysis using restricted maximum likelihood method and Hartung-Knapp-Sidik-Jonkman adjusting for the standard errors.

HR = hazard ratio; OR = odds ratio; RR = relative risk; GHSI = Global Health Security Index.

Table S3 – Characteristics of Studies Included in the Meta-Analysis on the Associations of Diabetes, Hypertension, and Obesity with COVID-19 Mortality, December 2019 – December, 2020 (n=145)

Study ID	Country	WHO region	WB income level	Health index score	GHSI score	Start date	End date	Data source	Study design		Exposure	e	Sample size	Mean age, y	Men, %	Effect estimate type	Funding source	NOS score
										DM	HTN	OB						
Agarwal_S_USA_20201	USA	AMR	HI	73.9	76.2	3/11/2020	5/7/2020	EHR	Cohort	No	Yes	No	1,279	67.9	49.3	OR	Not reported	9
Ahlstrom_B_SWE_2020 ²	Sweden	EUR	HI	82.1	66.4	1/31/2020	5/27/2020	ARS	Cohort	No	Yes	Yes	9,905	61.0	74.0	HR	Independent	9
Al_Wahaibi_A_OMN_20203	Oman	EMR	HI	75.2	40.9	2/24/2020	7/19/2020	ARS	C-S	Yes	Yes	No	68,967	40.0	74.9	OR	None or NA	4
Alguwaihes_AM_SAU_20204	Saudi Arabia	EMR	HI	74.5	45.0	5/15/2020	7/15/2020	EHR	C-S	Yes	Yes	Yes	439	55.0	68.3	HR	Independent	7
Allameh_SF_IRN_20205	Iran	EMR	UMI	74.8	39.5	2/20/2020	3/19/2020	EHR	Cohort	Yes	Yes	No	396	56.9	61.8	RR	Not reported	8
Almazeedi_S_KWT_20206	Kuwait	EMR	HI	76.9	40.1	2/24/2020	4/20/2020	EHR	Cohort	Yes	Yes	Yes	1,096	41.0	81.0	OR	Independent	9
Al-Salameh_A_FRA_2020a7	France	EUR	HI	80.5	62.6	10/1/2020	4/21/2020	EHR	Cohort	Yes	No	No	432	73.0	55.1	HR	None or NA	9
Al-Salameh_A_FRA_2020b8	France	EUR	HI	80.5	62.6	1/24/2020	5/1/2020	EHR	Cohort	No	No	Yes	329	81.0	59.6	OR	Not reported	9
Amit_M_ISR_20209	Israel	EUR	HI	82.8	50.7	3/5/2020	4/27/2020	EHR	Cohort	Yes	Yes	No	156	72.0	69.0	OR	None or NA	9
Baqui_P_BRA_202010	Brazil	AMR	UMI	72.0	51.0	2/27/2020	5/4/2020	ARS	C-S	Yes	No	Yes	7,371	55.2	45.5	HR	None or NA	8
Barron_E_GBR_202011	UK	EUR	HI	78.8	68.3	3/1/2020	5/11/2020	ARS	C-S	Yes	No	No	61,414,470†	40.9	49.9	OR	None or NA	8
Bellan_M_ITA_202012	Italy	EUR	HI	81.1	51.9	3/1/2020	4/28/2020	EHR	Cohort	No	No	Yes	407	71.0	59.0	OR	None or NA	9
Bepouka_BI_COD_202013	D.R. Congo	AFR	LMI	48.6	26.0	3/23/2020	6/15/2020	EHR	Cohort	Yes	Yes	No	141	49.6	67.4	HR	Not reported	9
Berenguer_J_ESP_202014	Spain	EUR	HI	80.5	60.4	1/31/2020	3/17/2020	EHR	Cohort	No	Yes	Yes	4,035	70.0	61.0	HR	Independent	9
Bhargava_A_USA_202015	USA	AMR	HI	73.9	76.2	3/8/2020	6/14/2020	EHR	Cohort	No	Yes	No	265	50.4	52.8	OR	None or NA	4
Biscarini_S_ITA_202016	Italy	EUR	HI	81.1	51.9	2/21/2020	3/31/2020	ARS	Cohort	Yes	Yes	Yes	427	67.0	68.1	HR	Independent	9
Boulle_A_ZAF_202017	South Africa	AFR	UMI	56.6	47.5	3/1/2020	6/9/2020	ARS	Cohort	No	Yes	No	2,978	53.4	37.9	HR	Independent	9
Breland_JY_USA_202118	USA	AMR	HI	73.9	76.2	3/2/2020	5/20/2020	EHR	Cohort	No	No	Yes	9,347	65.0	91.0	OR	Independent	8
Cai_Y_CHN_202019	China	WPR	UMI	82.8	49.0	1/20/2020	3/3/2020	EHR	Cohort	Yes	Yes	No	941	57.0	48.0	HR	Independent	9
Cao_Y_CHN_202020	China	WPR	UMI	82.8	49.0	1/5/2020	2/22/2020	EHR	C-S	Yes	Yes	No	101	56.6	66.3	OR	Independent	8
Carrillo- Vega_MF_MEX_2020 ²¹	Mexico	AMR	UMI	72.1	55.1	2/28/2020	4/23/2020	ARS	C-S	Yes	Yes	Yes	9,946	48.2	57.8	OR	Independent	7
Cernigliaro_A_ITA_202022	Italy	EUR	HI	81.1	51.9	1/3/2020	6/26/2020	ARS	C-S	Yes	No	No	2,847	50.0	49.5	OR	Not reported	5
Chang_MC_KOR_2020 ²³	South Korea	WPR	HI	84.1	65.9	2/1/2020	4/10/2020	EHR	Cohort	Yes	No	No	106	67.6	50.1	HR	Independent	9
Cheng_X_CHN_2020 ²⁴	China	WPR	UMI	82.8	49.0	1/11/2020	2/20/2020	ARS	Cohort	Yes	Yes	No	220	59.5	48.2	HR	Independent	8
Chilimuri_S_USA_202025	USA	AMR	HI	73.9	76.2	3/9/2020	4/9/2020	EHR	Cohort	Yes	Yes	No	375	63.0	63.0	OR	None or NA	9
Ciardullo_S_ITA_2020 ²⁶	Italy France,	EUR	HI	81.1	51.9	2/22/2020	5/15/2020	EHR	Cohort	Yes	Yes	No	373	72.0	65.4	RR	None or NA	9
Group_FRA_2021 ²⁷	switzerland, and Belgium	EUR				2/25/2020	5/4/2020	EHR	Cohort	Yes	Yes	No	4,244	63.0	74.0	HR	Independent	9

Cummings_MJ_USA_2020 ²⁸	USA	AMR	HI	73.9	76.2	3/2/2020	4/1/2020	EHR	Cohort	Yes	Yes	No	257	62.0	67.0	HR	Independent	9
Dennis_JM_GBR_2021 ²⁹	UK	EUR	HI	78.8	68.3	3/1/2020	7/27/2020	ARS	Cohort	Yes	Yes	Yes	19,256	67.0	60.1	HR	Independent	9
deSouza_C_BRA_202030	Brazil	AMR	UMI	72.0	51.0	7/26/2020	8/1/2020	ARS	Cohort	Yes	Yes	Yes	9,807	70.2	47.5	OR	Not reported	8
DiCastelnuovo_A_ITA_202031	Italy	EUR	HI	81.1	51.9	2/19/2020	5/23/2020	EHR	Cohort	Yes	Yes	Yes	3,894	67.0	61.7	HR	None or NA	9
Eshrati_B_IRN_202032	Iran	EMR	UM	74.8	39.5	2/22/2020	3/25/2020	ARS	Cohort	Yes	No	No	3,188	55.1	60.4	HR	Independent	9
Filardo_TD_USA_202033	USA	AMR	HI	73.9	76.2	3/9/2020	4/8/2020	EHR	Cohort	Yes	No	Yes	270	58.0	67.4	RR	None or NA	8
Fox_T_USA_202034	USA	AMR	HI	73.9	76.2	3/1/2020	4/24/2020	EHR	C-S	No	Yes	No	355	66.2	51.0	OR	None or NA	8
Fried_MW_USA_202035	USA	AMR	HI	73.9	76.2	2/15/2020	4/20/2020	ARS	C-S	Yes	Yes	Yes	11,721	65.0	53.4	OR	Industry	7
Fumagalli_C_ITA_202036	Italy	EUR	HI	81.1	51.9	2/22/2020	4/10/2020	EHR	Cohort	Yes	Yes	No	516	67.0	66.9	HR	None or NA	8
Galloway_JB_GBR_202037	UK	EUR	HI	78.8	68.3	3/1/2020	4/17/2020	EHR	Cohort	Yes	Yes	No	1,157	71.0	57.6	HR	None or NA	9
Giacomelli_A_ITA_202038	Italy	EUR	HI	81.1	51.9	2/21/2020	4/20/2020	EHR	Cohort	No	No	Yes	233	61.0	69.1	HR	None or NA	9
Gil-Rodrigo_A_ESP_202039	Spain	EUR	HI	80.5	60.4	3/1/2020	4/30/2020	EHR	Cohort	Yes	Yes	Yes	1,000	62.3	56.2	OR	None or NA	6
Goodman_KE_USA_202040	USA	AMR	HI	73.9	76.2	4/15/2020	6/15/2020	ARS	Cohort	Yes	No	No	66,646	62.8	52.9	RR	Independent	9
Grasselli_G_ITA_202041	Italy	EUR	HI	81.1	51.9	2/20/2020	5/30/2020	EHR	Cohort	Yes	Yes	No	3,988	63.0	80.0	HR	Independent	9
Gupta_S_USA_202042	USA	AMR	HI	73.9	76.2	3/4/2020	4/4/2020	EHR	Cohort	Yes	Yes	No	2,215	60.5	64.8	OR	Independent	8
Gutierrez_J_MEX_202043	Mexico	AMR	UMI	72.1	55.1	2/28/2020	9/16/2020	ARS	C-S	Yes	Yes	Yes	654,858	46.1	52.2	OR	None or NA	8
Haase_N_DNK_202044	Denmark	EUR	HI	80.6	67.3	3/10/2020	6/16/2020	EHR	Cohort	No	Yes	No	323	68.0	74.0	HR	Industry	9
Hajifathalian_K_USA_202045	USA	AMR	HI	73.9	76.2	3/4/2020	4/9/2020	EHR	Cohort	No	No	Yes	770	63.5	60.8	RR	Not reported	9
Harrison_SL_USA_202046	USA	AMR	HI	73.9	76.2	1/20/2020	5/26/2020	EHR	Cohort	Yes	No	No	31,461	50.0	45.5	OR	Independent	8
Galdamez_DR_MEX_202047	Mexico	AMR	UMI	72.1	55.1	2/15/2020	6/27/2020	ARS	C-S	Yes	Yes	Yes	211,003	45.7	54.7	OR	Not reported	8
Huang_S_CHN_202048	China	WPR	UMI	82.8	49.0	12/30/2019	4/19/2020	EHR	Cohort	No	Yes	No	310	62.0	56.0	OR	Not reported	9
Hui_Y_CHN_202049	China	WPR	UMI	82.8	49.0	1/28/2020	3/10/2020	EHR	Cohort	Yes	No	No	167	65.0	65.3	HR	Independent	9
Iaccarino_G_ITA_202050	Italy	EUR	HI	81.1	51.9	3/9/2020	4/9/2020	ARS	C-S	Yes	Yes	No	1,591	66.5	64.0	OR	Independent	7
Ioannou_GN_USA_202051	USA	AMR	HI	73.9	76.2	2/28/2020	5/14/2020	EHR	Cohort	Yes	Yes	No	10,131	63.6	91.0	HR	Independent	9
Izurieta_HS_USA_202052	USA	AMR	HI	73.9	76.2	4/1/2020	5/8/2020	ARS	C-S	Yes	Yes	Yes	25,333,329†	73.0	44.0	OR	Independent	8
Jackson_BR_USA_202053	USA	AMR	HI	73.9	76.2	3/1/2020	3/31/2020	EHR	Cohort	Yes	Yes	No	297	60.0	50.0	OR	Independent	9
Jain_AC_IND_202054	INDIA	SEAR	LMI	67.1	43.6	4/15/2020	6/15/2020	EHR	Cohort	Yes	Yes	No	425	49.0	73.4	OR	None or NA	7
Jiang_Y_CHN_202055	China	WPR	UMI	82.8	49.0	1/30/2020	4/10/2020	EHR	Cohort	Yes	Yes	No	281	70.0	50.9	OR	Independent	9
Kaeuffer_C_FRA_2020 ⁵⁶	France	EUR	HI	80.5	62.6	3/20/2020	3/20/2020	EHR	Cohort	Yes	Yes	Yes	1,045	66.0	59.0	OR	Independent	7
Kim_DW_KOR_202057	South Korea	WPR	HI	84.1	65.9	1/20/2020	3/26/2020	ARS	C-S	Yes	Yes	No	9,148	46.0	39.0	OR	Independent	7
Kim_SW_KOR_202058	South Korea	WPR	HI	84.1	65.9	2/18/2020	7/10/2020	EHR	Cohort	Yes	Yes	Yes	2,254	57.0	35.8	HR	Independent	8
Kim_SY_KOR_202059	South Korea	WPR	HI	84.1	65.9	1/20/2020	4/30/2020	ARS	Cohort	No	No	Yes	4,057	50.0	42.5	HR	Independent	8
Kim_T_USA_202060	USA	AMR	HI	73.9	76.2	3/1/2020	5/12/2020	EHR	Cohort	Yes	Yes	No	10,861	65.0	59.6	OR	Independent	9
Klang_E_USA_202061	USA	AMR	HI	73.9	76.2	3/1/2020	5/17/2020	EHR	Cohort	Yes	Yes	No	572	60.0	69.4	OR	Not reported	9

Kocayigit_I_TUR_202062	Turkey	EUR	UMI	75.1	49.8	3/20/2020	4/10/2020	EHR	Cohort	Yes	No	No	169	65.8	46.7	OR	Not reported	7
Kunal_S_IND_202063	India	SEAR	LMI	67.1	43.6	1/30/2020	5/7/2020	EHR	Cohort	No	Yes	No	108	51.2	64.8	OR	None or NA	5
Lanini_S_ITA_202064	Italy	EUR	HI	81.1	51.9	1/29/2020	3/28/2020	EHR	Cohort	No	No	Yes	379	61.7	72.0	OR	Independent	9
Lee_SG_KOR_202065	South Korea	WPR	HI	84.1	65.9	3/26/2020	5/15/2020	ARS	C-S	Yes	Yes	No	7,339	47.1	40.1	OR	None or NA	7
Li_H_CHN_2020 ⁶⁶	China	WPR	UMI	82.8	49.0	1/22/2020	3/17/2020	EHR	Cohort	Yes	No	No	453	61.0	52.0	HR	Independent	8
Liu_J_CHN_202067	China	WPR	UMI	82.8	49.0	12/29/2019	2/28/2020	EHR	Cohort	Yes	No	No	1,190	57.0	53.4	OR	None or NA	7
Liu_M_CHN_202068	China	WPR	UMI	82.8	49.0	1/1/2020	3/4/2020	EHR	Cohort	Yes	No	No	665	58.0	47.8	OR	Independent	9
Liu_Z_CHN_2020 ⁶⁹ Mancilla-	China	WPR	UMI	82.8	49.0	2/8/2020	4/15/2020	EHR	Cohort	Yes	No	No	934	62.0	48.6	HR	None or NA	9
Galindo_J_MEX_2020 ⁷⁰	Mexico	AMR	UMI	72.1	55.1	2/28/2020	5/30/2020	ARS	Cohort	Yes	Yes	Yes	83,779	46.3	56.6	HR	None or NA	9
Mansour_A_IRN_2020 ⁷¹ Martos-	Iran	EMR	UMI	74.8	39.5	2/25/2020	4/21/2020	EHR	C-S	Yes	No	No	353	61.7	57.5	OR	None or NA	7
Benitez_FD_MEX_202172	Mexico	AMR	UMI	72.1	55.1	1/1/2020	5/13/2020	ARS	C-S	Yes	Yes	Yes	38,324	46.9	58.3	OR	None or NA	7
McNeill_JN_USA_202073	USA	AMR	HI	73.9	76.2	2/28/2020	4/27/2020	EHR	Cohort	No	No	Yes	781	61.0	58.0	OR	Independent	7
Mejia_F_PER_202074	Peru	AMR	UMI	76.4	53.8	3/29/2020	6/11/2020	OTH	Cohort	Yes	Yes	Yes	369	59.0	65.3	HR	None or NA	6
Mikami_T_USA_202075	USA	AMR	HI	73.9	76.2	3/12/2020	4/17/2020	EHR	Cohort	Yes	Yes	No	3,708	66.0	57.0	HR	Not reported	6
Miller_J_USA_202076	USA	AMR	HI	73.9	76.2	3/7/2020	4/30/2020	EHR	Cohort	Yes	Yes	Yes	2,316	64.5	51.8	OR	Independent	7
Mirani_M_ITA_202077	Italy	EUR	HI	81.1	51.9	2/20/2020	4/9/2020	EHR	Cohort	Yes	Yes	Yes	387	66.0	66.7	HR	Not reported	8
Moon_SJ_KOR_202078	South Korea	WPR	HI	84.1	65.9	1/20/2020	5/15/2020	ARS	C-S	Yes	No	No	5,307	56.0	39.0	OR	Independent	7
Munoz_P_ESP_202079	Spain	EUR	HI	80.5	60.4	3/1/2020	5/10/2020	OTH	Cohort	No	Yes	No	100	61.5	52.0	OR	Independent	9
Mukherjee_V_USA_202080	USA	AMR	HI	73.9	76.2	3/10/2020	5/18/2020	EHR	Cohort	Yes	Yes	Yes	137	59.0	72.3	HR	Not reported	9
Munblit_D_RUS_202081	Russia	EUR	UMI	71.6	47.1	4/8/2020	5/28/2020	EHR	Cohort	Yes	Yes	No	3,480	56.0	50.5	OR	Independent	7
Nachega_JB_COG_2020 ⁸²	D.R. Congo	AFR	LMI	48.6	26.0	3/10/2020	7/31/2020	ARS	Cohort	Yes	Yes	Yes	766	46.0	65.3	HR	Independent	9
Nachtigall_I_DEU_202083	Germany	EUR	HI	81.1	65.7	2/12/2020	6/12/2020	OTH	Cohort	Yes	No	No	1,904	73.0	51.5	HR	Independent	9
Nakeshbandi_M_USA_2020 ⁸⁴	USA	AMR	HI	73.9	76.2	3/10/2020	4/13/2020	EHR	Cohort	Yes	Yes	Yes	504	68.0	53.0	RR	Not reported	9
Nogueira_PJ_PRT_2020 ⁸⁵	Portugal	EUR	HI	77.6	58.7	1/1/2020	4/21/2020	ARS	C-S	Yes	No	No	20,293	52.1	41.3	OR	None or NA	7
Orioli_L_BEL_2020 ⁸⁶	Belgium	EUR	HI	80.6	61.9	3/1/2020	5/6/2020	ARS	C-S	Yes	No	No	192	67.0	50.0	HR	None or NA	4
Orwa_A_MYS_2020 ⁸⁷ Pacheco-	Worldwide	World				12/30/2019	4/21/2020	ARS	C-S	Yes	Yes	No	828	49.4	59.1	OR	None or NA	6
Pantoja_E_MEX_2020 ⁸⁸	Mexico	AMR	UMI	72.1	55.1	2/28/2020	4/30/2020	ARS	Cohort	Yes	Yes	Yes	19,224	46.6	58.2	OR	Not reported	7
Palaiodimos_L_USA_202089	USA	AMR	HI	73.9	76.2	3/9/2020	4/12/2020	EHR	Cohort	Yes	No	Yes	200	64.0	49.0	OR	None or NA	9
Panagiotou_OA_USA_202190	USA	AMR	HI	73.9	76.2	3/16/2020	9/15/2020	EHR	Cohort	Yes	Yes	No	5,256	79.0	39.0	OR	Independent	9
Parikh_R_USA_202091	USA	AMR	HI	73.9	76.2	3/1/2020	5/1/2020	EHR	Cohort	No	No	Yes	160	60.4	65.6	OR	None or NA	9
Park_BE_KOR_202192	South Korea	WPR	HI	84.1	65.9	2/15/2020	4/24/2020	ARS	Cohort	Yes	Yes	No	2,269	55.5	36.0	OR	Independent	9
Park_JG_KOR_2020 ⁹³ Parra-	South Korea	WPR	HI	84.1	65.9	2/20/2020	4/14/2020	EHR	Cohort	Yes	No	No	289	72.0	46.0	HR	Independent	8
Bracamonte_GM_MEX_202094	Mexico	AMR	UMI	72.1	55.1	1/13/2020	7/17/2020	ARS	Cohort	Yes	Yes	Yes	331,298	44.0	53.8	OR	Not reported	8
Pena_JE_MEX_202095	Mexico	AMR	UMI	72.1	55.1	2/28/2020	11/13/2020	ARS	C-S	Yes	Yes	Yes	121,225	50.0	59.8	OR	Not reported	7

Petrilli_CM_USA_202096	USA	AMR	HI	73.9	76.2	3/1/2020	4/8/2020	EHR	Cohort	Yes	Yes	No	5,279	54.0	49.5	HR	Independent	9
Pettit_NN_USA_202097	USA	AMR	HI	73.9	76.2	3/1/2020	4/18/2020	EHR	Cohort	Yes	Yes	Yes	238	58.5	47.5	OR	Not reported	9
Polverino_F_ITA_202098	Italy	EUR	HI	81.1	51.9	3/25/2020	4/22/2020	OTH	C-S	Yes	Yes	Yes	3,179	69.0	68.3	OR	Independent	8
Posso_M_ESP_202099	Spain	EUR	HI	80.5	60.4	2/23/2020	5/12/2020	EHR	Cohort	Yes	Yes	Yes	834	78.2	46.5	OR	None or NA	8
Galbarro_FJ_MEX_2020 ¹⁰⁰	Mexico	AMR	UMI	72.1	55.1	2/27/2020	4/27/2020	ARS	Cohort	Yes	Yes	Yes	15,529	55.0	57.8	HR	None or NA	9
Qin_W_CHN_2020101	China	WPR	UMI	82.8	49.0	12/19/2019	2/20/2020	EHR	Cohort	No	Yes	No	582	64.0	50.3	OR	Independent	6
Qin_W_CHN_2021102	China	WPR	UMI	82.8	49.0	1/31/2020	3/6/2020	EHR	Cohort	No	Yes	No	262	63.5	46.9	HR	Independent	9
Rastad_H_IRN_2020103	Iran	EMR	UMI	74.8	39.5	2/20/2020	3/25/2020	EHR	Cohort	Yes	No	No	2,957	54.8	53.7	OR	Independent	8
Reilev_M_DNK_2020104	Denmark	EUR	HI	80.6	67.3	2/27/2020	5/19/2020	ARS	Cohort	Yes	Yes	Yes	11,122	48.0	42.0	OR	Independent	7
Rethemiotaki_I_CHN_2020 ¹⁰⁵	China	WPR	UMI	82.8	49.0	12/19/2019	2/20/2020	ARS	C-S	Yes	Yes	No	44,672	55.0	64.0	OR	Not reported	7
Izquierdo_M_ESP_2020 ¹⁰⁶	Spain	EUR	HI	80.5	60.4	3/16/2020	4/10/2020	OTH	Cohort	Yes	No	No	238	64.7	55.0	HR	Independent	9
Gonzalez_CG_ESP_2020 ¹⁰⁷	Spain	EUR	HI	80.5	60.4	3/1/2020	3/24/2020	OTH	Cohort	Yes	Yes	Yes	1,208	65.0	58.0	OR	Independent	9
Rodriguez- Molinero_A_ESP_2020 ¹⁰⁸	Spain	EUR	HI	80.5	60.4	3/12/2020	5/2/2020	EHR	Cohort	Yes	Yes	No	418	65.4	57.0	OR	None or NA	9
Rodriguez- Nava_G_USA_2020 ¹⁰⁹	USA	AMR	HI	73.9	76.2	3/1/2020	5/25/2020	OTH	Cohort	No	Yes	No	313	68.0	58.0	HR	Not reported	9
Rosenthal_N_USA_2020110	USA	AMR	HI	73.9	76.2	4/1/2020	5/31/2020	ARS	C-S	Yes	Yes	No	64,781	57.0	49.0	OR	Independent	7
Rossi_AP_ITA_2020111	Italy	EUR	HI	81.1	51.9	3/8/2020	3/30/2020	ARS	Cohort	No	No	Yes	95	62.5	82.1	HR	Not reported	9
Rossi_PG_ITA_2020112	Italy	EUR	HI	81.1	51.9	2/27/2020	4/2/2020	ARS	Cohort	Yes	Yes	Yes	2,653	63.2	50.1	HR	Independent	9
Rottoli_M_ITA_2020113	Italy	EUR	HI	81.1	51.9	3/1/2020	4/27/2020	ARS	Cohort	Yes	Yes	Yes	482	66.2	63.0	HR	None or NA	9
Rozaliyani_A_IDN_2020114	Indonesia	SEAR	UMI	72.7	49.2	3/2/2020	4/29/2020	ARS	Cohort	Yes	Yes	No	4,052	45.8	54.0	OR	Not reported	9
Saand_AR_USA_2020115	USA	AMR	HI	73.9	76.2	3/15/2020	5/30/2020	ARS	Cohort	Yes	No	Yes	495	68.0	58.4	HR	None or NA	9
Salacup_G_USA_2020116	USA	AMR	HI	73.9	76.2	3/1/2020	4/24/2020	EHR	Cohort	Yes	Yes	No	242	66.0	51.0	OR	Not reported	9
Santos_MM_BRA_2020117	Brazil	AMR	UMI	72.0	51.0	2/20/2020	6/2/2020	ARS	Cohort	Yes	No	No	80,123	51.0	57.0	HR	Not reported	9
Seiglie_J_USA_2020118	USA	AMR	HI	73.9	76.2	3/11/2020	4/30/2020	ARS	Cohort	Yes	Yes	Yes	450	63.3	57.6	OR	Independent	9
Shah_C_USA_2020119	USA	AMR	HI	73.9	76.2	1/1/2020	5/31/2020	EHR	Cohort	Yes	Yes	No	487	68.4	56.1	OR	None or NA	9
Shah_P_USA_2020120	USA	AMR	HI	73.9	76.2	3/2/2020	5/6/2020	EHR	Cohort	Yes	Yes	No	522	63.0	41.8	OR	Not reported	8
Shang_J_CHN_2020121	China	WPR	UMI	82.8	49.0	12/25/2019	3/20/2020	EHR	Cohort	Yes	Yes	No	584	59.0	47.4	HR	Independent	9
Sheshah_E_SAU_2020122	Saudi Arabia	EMR	HI	74.5	45.0	5/1/2020	7/31/2020	OTH	Cohort	Yes	Yes	No	300	49.7	86.3	OR	Independent	8
Shi_S_CHN_2020123	China	WPR	UMI	82.8	49.0	1/1/2020	2/23/2020	EHR	Cohort	Yes	Yes	No	671	63.0	48.0	HR	Independent	8
Singh_S_USA_2020124	USA	AMR	HI	73.9	76.2	1/20/2020	5/31/2020	EHR	C-C	No	No	Yes	16,224	50.0	39.0	RR	Independent	8
Smith_AA_USA_2020125	USA	AMR	HI	73.9	76.2	3/1/2020	4/22/2020	EHR	Cohort	Yes	No	No	346	66.9	56.0	RR	None or NA	8
Sousa_GJB_BRA_2020126	Brazil	AMR	UMI	72.0	51.0	2/20/2020	4/14/2020	ARS	Cohort	Yes	No	Yes	2,070	44.0	49.0	HR	None or NA	7
Sun_H_CHN_2020127	China	WPR	UMI	82.8	49.0	1/29/2020	3/5/2020	EHR	Cohort	No	Yes	No	244	69.0	54.5	OR	Not reported	9
Sun_Y_CHN_2020128	China	WPR	UMI	82.8	49.0	1/15/2020	4/15/2020	EHR	Cohort	Yes	Yes	No	3,400	61.0	49.0	OR	Not reported	8

Sutter_W_FRA_2020129	France	EUR	HI	80.5	62.6	2/26/2020	4/20/2020	EHR	C-C	Yes	No	No	1,206	71.2	61.8	HR	None or NA	8
Tartof_SY_USA_2020130	USA	AMR	HI	73.9	76.2	2/13/2020	5/2/2020	EHR	Cohort	Yes	Yes	No	6,916	49.1	45.0	RR	Independent	9
Tavares_C_BRA_2020131	Brazil	AMR	UMI	72.0	51.0	2/26/2020	6/30/2020	ARS	C-S	Yes	Yes	Yes	89,405	58.9	56.5	OR	None or NA	7
Thompson_J_GBR_2020132	UK	EUR	HI	78.8	68.3	3/12/2020	5/19/2020	EHR	Cohort	No	Yes	No	470	68.7	54.0	OR	None or NA	9
van_Gerwen_M_USA_2020133	USA	AMR	HI	73.9	76.2	3/20/2020	5/13/2020	EHR	Cohort	Yes	Yes	Yes	2,015	56.8	55.3	OR	None or NA	9
Wang_Z_USA_2020134	USA	AMR	HI	73.9	76.2	3/1/2020	4/15/2020	EHR	C-S	Yes	Yes	Yes	3,273	65.0	57.0	HR	None or NA	7
Workinggroup_ESP_2020135	Spain	EUR	HI	80.5	60.4	1/31/2020	4/27/2020	ARS	C-S	Yes	Yes	No	218,652	61.0	43.8	OR	Not reported	7
Wu_R_CHN_2020136	China	WPR	UMI	82.8	49.0	12/10/2019	3/18/2020	ARS	Cohort	Yes	Yes	No	21,392	50.0	52.0	HR	Independent	7
Xie_J_USA-2020137	USA	AMR	HI	73.9	76.2	3/30/2020	4/5/2020	OTH	Cohort	Yes	Yes	Yes	287	61.5	43.0	OR	Independent	9
Yang_Q_CHN_2020138	China	WPR	UMI	82.8	49.0	1/1/2020	2/29/2020	EHR	Cohort	No	Yes	No	226	53.9	51.8	HR	None or NA	8
Yazdanpanah_Y_FRA_2020139	France	EUR	HI	80.5	62.6	1/24/2020	3/15/2020	ARS	Cohort	Yes	No	Yes	246	65.0	57.0	HR	Independent	9
You_H_KOR_2020140	South Korea	WPR	HI	84.1	65.9	1/20/2020	3/31/2020	ARS	Cohort	Yes	No	No	5,473	45.0	44.6	OR	Not reported	9
Yu_C_CHN_2020141	China	WPR	UMI	82.8	49.0	1/14/2020	3/26/2020	OTH	Cohort	Yes	Yes	No	1,464	64.0	50.3	OR	Independent	9
Zandkarimi_E_IRN_2020142	Iran	EMR	UMI	74.8	39.5	2/22/2020	5/18/2020	EHR	Cohort	Yes	No	No	1,831	57.7	55.7	HR	Independent	9
Zhang_J_CHN_2020143	China	WPR	UMI	82.8	49.0	1/1/2020	3/17/2020	EHR	Cohort	Yes	No	No	312	57.0	44.9	HR	Independent	9
Zhang_Y_CHN_2020144	China	WPR	UMI	82.8	49.0	1/29/2020	3/12/2020	OTH	Cohort	Yes	No	No	258	64.2	54.0	HR	Independent	8
Zhu_L_CHN_2020145	China	WPR	UMI	82.8	49.0	12/30/2019	3/20/2020	OTH	Cohort	Yes	No	No	7,337	54.0	47.4	HR	Independent	4

Note: UK = United Kingdom, USA = United States of America, HI = high income, UMI = upper middle income, LMI = lower middle income, GHSI = global health security index, WHO = World Health Organization, WB = World Bank, AFR = African Region, SEAR = Southeast Asian Region, AMR = American Region, EMR = East Mediterranean Region, EUR = European Region, WPR = West Pacific Region, EHR = electronic health (medical) record, ARS = administrative/registry/surveillance or (case) reporting system, C-C = case-control, C-S = cross-sectional, DM = diabetes mellitus, HTN = hypertension, OB = obesity, ES = effect size, OR = Odds ratio, HR = hazard ratio, RR = relative risk, NOS = Newcastle-Ottawa Scale.

+ Population size.

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