Incidence, risk factors and prognostic effect of imaging right ventricular involvement in patients with COVID-19: a dose–response analysis protocol for systematic review

Chenghui Zhou 1, Baohui Lou,2 Hui Li,3 Xin Wang,4 Hushan Ao,1 Fujian Duan4

ABSTRACT

Introduction Emerging evidence has shown that COVID-19 infection may result in right ventricular (RV) disturbance and be associated with adverse clinical outcomes. The aim of this meta-analysis is to summarise the incidence, risk factors and the prognostic effect of imaging RV involvement in adult patients with COVID-19.

Methods A systematical search will be performed in PubMed, EMBase, ISI Knowledge via Web of Science and preprint databases (MedRxiv and BioRxiv) (until October 2021) to identify all cohort studies in adult patients with COVID-19. The primary outcome will be the incidence of RV involvement (dysfunction and/or dilation) assessed by echocardiography, CT or MRI. Secondary outcomes will include the risk factors for RV involvement and their association with all-cause mortality during hospitalisation. Additional outcomes will include the RV global or free wall longitudinal strain (RV-GLS or RV-FWLS), tricuspid annular plane systolic excursion (TAPSE), fractional area change (FAC) and RV diameter. Univariable or multivariable meta-regression and subgroup analyses will be performed for the study design and patient characteristics (especially acute or chronic pulmonary embolism and pulmonary hypertension). Sensitivity analyses will be used to assess the robustness of our results by removing each included study at one time to obtain and evaluate the remaining overall estimates of RV involvement incidence and related risk factors, association with all-cause mortality, and other RV parameters (RV-GLS or RV-FWLS, TAPSE, S’, FAC and RV diameter). Both linear and cubic spline regression models will be used to explore the dose–response relationship between different categories (>2) of RV involvement and the risk of mortality (OR or HR).

Ethics and dissemination There was no need for ethics approval for the systematic review protocol according to the Institutional Review Board/Independent Ethics Committee of Kuwaiti Hospital. This meta-analysis will be disseminated through a peer-reviewed journal for publication.

PROSPERO registration number CRD42021231689.

INTRODUCTION

By 31 January 2021, the COVID-19 pandemic had caused 103,495,393 infections and 2,236,883 deaths worldwide in 215 countries.1 Since the initial outbreak, the overall mortality risk of patients with COVID-19 has remained stable at approximately 2.0%. However, critically ill patients admitted to the intensive care unit (ICU) continue to exhibit a high mortality of up to 20%–50%.2–4 The major contributing complication is thought to be the cardiovascular involvement, manifesting as biomarker elevation of myocardial injury, ST-segment elevation, decrease in ventricular function and change in ventricular structure.5 In a retrospective analysis, Zhou et al showed that the incidence of heart failure after COVID-19 infection is up to 52% in deceased patients and approximately 12% in discharged patients.6

The right ventricles (RVs) have been recognised as passive conduits and volumetric organs, and are vulnerable to changes in preload and afterload. In clinical settings, therapy for patients with RV failure is less effective than for patients with left ventricular (LV) failure, resulting in poor medium-term
and long-term outcomes. Various pharmacological interventions have been studied with controversial results. Isolated reports suggest that administration of promising drugs may reduce ICU stays, but their effectiveness regarding mortality remains elusive. Hence, early identification of RV involvement could lead to risk stratification, early intervention and potentially improve the prognosis for critically ill patients with COVID-19.

The role of RV in cardiovascular medicine has become a heavily researched issue. In studies concerning COVID-19 related ventricular function, involvement of RV as opposed to LV may be more frequent and have more related ventricular function, involvement of RV as heavily researched issue. In studies concerning COVID-19 prognosis for critically ill patients with COVID-19. 

**Objectives**
The aim of this systematic review and meta-analysis is to summarise the evidence of incidence, risk factors and prognostic effect of imaging RV involvement in patients with COVID-19.

**METHODS AND ANALYSIS**

**Search strategy**
This systematic review and meta-analysis will be performed in accordance with the Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols. PubMed, EMBase, ISI Knowledge via Web of Science and preprint databases (MedRxiv and BioRxiv) (until October 2021) will be systematically searched to identify all cohort studies concerning imaging RV involvement in adult patients with COVID-19. The imaging methods will be echocardiography, cardiac CT and cardiac MRI. The reference lists of the retrieved articles will also be searched. The related search strategy with keywords is presented in **table 1**. The searching process is shown in **figure 1**.

**Type of participants**
Adult patients infected with COVID-19 will be selected as the study participants.

**Patient and public involvement**
Patients and/or the public will not be involved in the design, conduct or reporting or dissemination plans for this research.

**Type of studies**
We will include both retrospective and prospective cohort studies concerning imaging RV in adult patients with COVID-19. No language limitation will be set for the inclusion of eligible studies. Studies reporting imaging RV involvement in other coronary virus infections (SARS-CoV-2) will be included. The inclusion of eligible studies is shown in **table 1**.

**Table 1** Search strategy in various databases (PubMed, EMBase, ISI Knowledge via Web of Science, MedRxiv and BioRxiv)

<table>
<thead>
<tr>
<th>Database</th>
<th>Search items</th>
</tr>
</thead>
<tbody>
<tr>
<td>PubMed</td>
<td>No.</td>
</tr>
<tr>
<td></td>
<td># 1 (((right ventricle) OR (right ventricular)) OR (right heart)) OR (cardiac function) OR (ventricular function) OR (((echocardiographic) OR (computed tomography)) OR (magnetic resonance imaging))</td>
</tr>
<tr>
<td></td>
<td># 2 (COVID-19) OR (SARS-CoV-2)</td>
</tr>
<tr>
<td></td>
<td># 3 # 1 and # 2</td>
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<tr>
<td>EMBase</td>
<td>right ventricle OR right ventricular OR right heart OR cardiac function OR ventricular function OR echocardiographic OR computed tomography OR magnetic resonance imaging</td>
</tr>
<tr>
<td></td>
<td># 2 ‘COVID-19 19’ OR ‘sars cov 2’</td>
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<tr>
<td></td>
<td># 3 # 1 and # 2</td>
</tr>
<tr>
<td>ISI Knowledge via Web of Science</td>
<td>TOPIC: (right ventricle) OR TOPIC: (right ventricular) OR TOPIC: (right heart) OR TOPIC: (cardiac function) OR TOPIC: (ventricular function) OR TOPIC: (echocardiographic) OR TOPIC: (computed tomography) OR TOPIC: (magnetic resonance imaging)</td>
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<tr>
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<td># 2 TOPIC: (COVID-19) OR TOPIC: (SARS-CoV-2)</td>
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<td># 3 # 1 and # 2</td>
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<tr>
<td>MedRxiv</td>
<td>(((right ventricle) OR (right ventricular)) OR (right heart)) OR (cardiac function) OR (ventricular function) OR (((echocardiographic) OR (computed tomography)) OR (magnetic resonance imaging))</td>
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<td>BioRxiv</td>
<td>(((right ventricle) OR (right ventricular)) OR (right heart)) OR (cardiac function) OR (ventricular function) OR (((echocardiographic) OR (computed tomography)) OR (magnetic resonance imaging))</td>
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</tbody>
</table>
or Middle East respiratory syndrome coronavirus) will be excluded. Studies without reporting the outcomes of interest will also be excluded.

**Type of outcomes**
The primary outcome will be the incidence of RV involvement (dysfunction and/or dilation) assessed by echocardiography, CT or MRI. The definition criteria of RV dysfunction will be as follows: (1) fractional area change (FAC) <35%, tricuspid annular plane systolic excursion (TAPSE) <17 mm and/or S’<10 mm/sec; (2) 3D-(RV ejection fraction) RVEF <45%; (3) RV free wall longitudinal strain (RV-FWLS) >−20%. The definition criteria of RV dilation will be as follows: (1) RV to LV basal diameter classified as mild (0.67–0.9), moderate (1.0) and severe (>1.0) enlargement; (2) RV basal diameter >41 mm.

Secondary outcomes will include the risk factors for RV involvement the association with all-cause mortality during hospitalisation. Additional outcomes will be RV global longitudinal strain (RV-GLS) or RV-FWLS, TAPSE, S’, FAC, RVEF, and RV diameter.

**Data extraction**
Data extraction will be completed by two independent authors (BL and HL). A third author (XW) will make the final decision in case of disagreements. Data extraction will consist of study design (author, publication year, country, retrospective or prospective type and sample size), patient characteristics (mean age, male proportion, diabetes proportion, hypertension proportion, hyperlipidaemia proportion, smoking proportion, coronary artery disease proportion, previous myocardial infarction, chronic heart failure, atrial fibrillation, previous peripheral vascular disease, previous stroke or transient ischaemic accident, acute or chronic kidney dysfunction, previous lung disease, pulmonary hypertension proportion, LV ejection fraction, elevated cTn proportion, elevated BNP proportion, acute LV failure proportion, beta-blocker usage, statin usage, ACE inhibitor/angiotensin receptor blocker usage, calcium channel blocker usage, aspirin usage, sacubitril/valsartan and diuretics), follow-up period, type of RV involvement, different categories of RV involvement and imaging methods (echocardiography, CT or MRI).

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**Figure 1** Searching process flowchart for eligible trials Selection. MERS, Middle East respiratory syndrome coronavirus; RV involvement, right ventricular involvement.
**Risk of bias assessment**

The methodological quality evaluation of the studies will be evaluated by the Newcastle-Ottawa quality assessment scale.29

**Data synthesis**

The incidence of imaging RV involvement including RV dysfunction and/or dilation in each study will be calculated. The OR or HR and 95% CI in each study for the risk factors of RV involvement and association with all-cause mortality will be extracted. For continuous outcomes of RV-GLS, RV-FWLS, TAPSE, FAC and RV diameter (reported as the mean±SD, median and IQR, or median and range), we will calculate mean differences for each study and used weights (the inverse variance of the estimate) to pool the estimate (weighted mean difference) with 95% CI. The DerSimonian and Laird random-effects model will be used in the pooled analysis for potential clinical inconsistency. Univariable or multivariable meta-regression and subgroup analyses will be conducted for related risk factors (especially acute or chronic pulmonary embolism and pulmonary hypertension) and the association of RV involvement with all-cause mortality. Sensitivity analyses will be used to assess the robustness of our results by removing each included study at one time to obtain and evaluate the remaining overall estimates of RV involvement incidence and related risk factors, association with all-cause mortality, and other RV parameters (RV-GLS, RV-FWLS, TAPSE, FAC and RV diameter). If different severity categories are reported, we will choose the category without RV involvement as a reference. We will pool the ORs by combining all the categories of RV involvement for comparison with and without RV involvement categories using the DerSimonian and Laird random-effects model. If a study reports multiple categories (>2 categories), we will calculate the OR using data on the number of cases and non-cases in all of the categories. Both linear and cubic spline regression models will be used to explore the dose–response relationship between different categories of RV involvement and the risk of mortality (OR or HR).21 22 Publication bias assessment will be performed with the Begg’s and Egger’s tests. P<0.05 (two-sided) will be considered statistically significant. All statistical analyses will be performed in Stata software (V.10.0, StataCorp) and RevMan software (V.5.0, Cochrane Collaboration, Oxford, UK).

**DISCUSSION**

Although the primary target of COVID-19 is the respiratory system, the cardiovascular system could also be affected by the following major mechanisms: (1) COVID-19 directly binds to the renin-ACE2 receptor in the cardiovascular system,23 interferes with the related downstream signaling pathway, significantly affects the neurohumoral regulation of the cardiovascular system and directly causes myocardial injury. (2) COVID-19 binds to ACE2 receptors in the respiratory system,25 which directly causes lung injury and affects respiratory function. Hypoxia could directly affect the balance of myocardial oxygen supply and demand, or indirectly cause right heart damage by pulmonary hypertension. (3) In many critical cases, COVID-19 can cause acute systemic inflammatory reactions and cytokine storms,24 resulting in systemic organ dysfunction or failure to cause myocardial injury directly or indirectly. All of these factors could potentially affect the structure and function of the RV.

The imaging tools for RV structure and function assessment includes echocardiography, CT and cardiac Magnetic Resonance Imaging (CMR) in a direct way with more information than other indirect methods (such as electrocardiography, X-ray or central venous pressure). Although CT and CMR possess some unique advantages, 2D with 3D echocardiography could also provide non-invasive, convenience, fast and comprehensive RV assessment with low cost for subsequent therapeutic treatment guidance.25 In addition, point-of-care evaluation using echocardiography is quite meaningful for critical patients with COVID-19 infection in intensive care unit. Specifically, echocardiographic evaluation for patients with COVID-19 could reduce the time of transportation and thereby the cross-over contamination in hospital. In this study, TAPSE, FAC, RV-GLS, RV-FWLS and RV diameter will be chosen as the second outcomes. These echocardiographic parameters are regular and useful for RV structural and function assessment, and have been recommended in the recent guideline for patients with COVID-19.26–28

The major strength of this systematic review and meta-analysis will be the first comprehensive summary of the risk factors and the prognostic effect of imaging RV involvement in patients with COVID-19. Moreover, this meta-analysis will focus on the incidence of imaging RV involvement with different outcomes in patients with COVID-19. There are several limitations in our analysis. First, the majority of included study will be retrospective in design. Thus, inherit bias cannot be ruled out. Second, the sample size and the number of studies may be small due to the limited usage of imaging tools for cardiac assessment in patients with COVID-19. Third, we could not rule out the potential influence of different definitions of RV dysfunction in COVID-19. Fourth, the impact of different severities of COVID-19 associated with RV involvement on adverse clinical outcomes needs further studies.

**ETHICS AND DISSEMINATION**

There is no need for ethical approval for the systematic review protocol according to the Institutional Review Board/Independent Ethics Committee of Fuwai Hospital. This meta-analysis will be disseminated through a peer-reviewed journal for publication.

**Author affiliations**

1Department of Anesthesiology, State Key Laboratory of Cardiovascular Disease, Fuwai Hospital, National Center for Cardiovascular Diseases, Chinese Academy of Medical Sciences & Peking Union Medical College, Beijing, China

2Department of Radiology, Beijing Hospital, National Center of Gerontology; Institute of Geriatric Medicine, Chinese Academy of Medical Sciences, Beijing, China

[92x234]RV involvement on adverse clinical outcomes needs further

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2Department of Radiology, Beijing Hospital, National Center of Gerontology; Institute of Geriatric Medicine, Chinese Academy of Medical Sciences, Beijing, China
Contributors CZ and FD contributed to the conception and design of the study, and revision of the protocol. The manuscript of the protocol was drafted by BL. HL and XW will independently search and select the eligible studies and extract the data from the included studies. HL and HA will assess methodological quality and the risk of bias. All the authors approved the protocol publication.

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Competing interests None declared.

Patient consent for publication

Patient and public involvement Patients and/or the public were not involved in the design, conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

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ORCID iD

Chenghui Zhou http://orcid.org/0000-0001-6428-6069

REFERENCES

2020