



Heart–lung interactions in COVID-19: prognostic impact and usefulness of bedside echocardiography for monitoring of the right ventricle involvement

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Abstract

Due to the SARS-CoV-2 infection–related severe pulmonary tissue damages associated with a relative specific widespread thrombotic microangiopathy, the pathophysiologic role of heart–lung interactions becomes crucial for the development and progression of right ventricular (RV) dysfunction. The high resistance in the pulmonary circulation, as a result of small vessel thrombosis and hypoxemia, is the major cause of right heart failure associated with a particularly high mortality in severe COVID-19. Timely identification of patients at high risk for RV failure, optimization of mechanical ventilation to limit its adverse effects on RV preload and afterload, avoidance of medication-related increase in the pulmonary vascular resistance, and the use of extracorporeal membrane oxygenation in refractory respiratory failure with hemodynamic instability, before RV failure develops, can improve patient survival. Since it was confirmed that the right-sided heart is particularly involved in the clinical deterioration of patients with COVID-19 and pressure overload-induced RV dysfunction plays a key role for patient outcome, transthoracic echocardiography (TTE) received increasing attention. Limited TTE focused on the right heart appears highly useful in hospitalized COVID-19 patients and particularly beneficial for monitoring of critically ill patients. In addition to detection of right-sided heart dilation and RV dysfunction, it enables assessment of RV–pulmonary arterial coupling and evaluation of RV adaptability to pressure loading which facilitate useful prognostic statements to be made. The increased use of bedside TTE focused on the right heart could facilitate more personalized management and treatment of hospitalized patients and can contribute towards reducing the high mortality associated with SARS-CoV-2 infection.

Keywords COVID-19 · Echocardiography · Heart–lung interaction · Pulmonary vessel thrombosis · Right ventricle · Heart failure

Introduction

Coronavirus disease 2019 (COVID-19) is caused by the severe acute respiratory syndrome corona-virus 2 (SARS-CoV-2), a novel coronavirus related to the SARS-CoV and Middle East respiratory syndrome coronavirus (MERS-CoV) [1]. Compared with SARS and MERS, COVID-19 exerts more damages on the cardiovascular system, most notably life-threatening pulmonary vessel injury and cardiac dysfunction, with and without direct myocardial injury [1, 2].

The primary pulmonary injury and the subsequent cardiovascular complications constitute the key pathophysiology of COVID-19 [3]. Pulmonary vessel injuries with blood flow alterations, followed by right heart dilation and right ventricular (RV) failure, were found among the major causes of death related to COVID-19 [3–5]. In addition, critically ill COVID-19 patients are at particularly high risk of pulmonary embolism (PE) associated with massive RV pressure overload [6].

This article provides an overview on the clinical relevance of COVID-19-associated RV dysfunction and on the usefulness of right heart assessment by transthoracic echocardiography (TTE) in hospitalized patients. Special attention is focused on the benefits of bedside limited TTE in critically ill COVID-19 patients, where it is increasingly gaining importance for clinical decision-making.

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Major COVID-related pulmonary and cardiovascular injuries

Like other coronaviruses, SARS-CoV-2 mainly affects the lung tissue. In an early study, all hospitalized COVID-19 patients had pneumonia confirmed by chest CT. [7]. In large study, 48.8% of hospitalized patients had severe pneumonia requiring high-flow O₂ therapy [8]. Adults admitted to emergency departments with suspected COVID-19 that were subsequently confirmed were more than twice as likely to die or receive organ support as those who did not have COVID-19 confirmed, despite having similar age and comorbidities (except chronic lung disease) [9]. When compared to patients with pneumonia of other etiologies, those with COVID-19 appeared more likely to develop hypoxemic respiratory failure and acute respiratory distress syndrome (ARDS) which was also associated with higher mortality [10, 11]. In a large study, 41% of COVID-19 patients necessitating intensive care developed ARDS [12]. COVID-19-related ARDS (CARDS) revealed particularly high mortality rates (between 54 and 85%) [13–16]. However, for patients with ARDS of different other etiologies, a meta-analysis revealed for the years 2000–2015 an overall mortality rate of 45% [17]. The strikingly high mortality related to CARDS raises important questions regarding the reasons for this discrepancy. One of the most likely explanations for the high mortality could be the particularly high incidence and severity of cardiovascular complications associated with SARS-CoV-2 infections.

Clinical and prognostic relevance of cardiovascular complications

In addition to pulmonary damages, SARS-CoV-2 can cause both direct and indirect cardiovascular injuries, including acute cor pulmonale and cardiogenic shock, thrombotic complications (including PE), acute coronary syndromes, myocardial injury, and severe arrhythmias [2, 18, 19]. Microthrombosis in various organs, which is the most prominent clinical feature of COVID-19, was demonstrable in 305 (91.3%) of 334 autopsies [20]. Autopsies revealed a major contribution of pulmonary vessel damages to COVID-19-related deaths [21–23]. Pulmonary small vessel thrombi and platelet agglutination are distinctive features of COVID-19 being also related to its poor prognosis [19, 22]. Between 80 and 100% of the lungs examined at autopsy revealed platelet–fibrin thrombi in the small pulmonary arterial and venous vasculature [22, 24, 25]. Histological analyses of the pulmonary vessels of deceased SARS-CoV-2-infected patients found that microangiopathy with widespread thrombosis (including alveolar capillary microthrombi) were 9 times more frequent than in the lungs from patients who died from influenza [23]. One study revealed

microthrombosis within small pulmonary arteries of all persons who died from COVID-19, and in all cases the cause of death was found within the lungs or the pulmonary vascular system [26]. The incidence of PE in COVID-19 patients is also particularly high (around 7% in hospitalized patients and up to one-third of those requiring intensive care unit) and is associated with a significantly higher mortality risk compared to COVID-19 patients without PE [27, 28].

Simultaneous occurrence of ARDS and high D-dimer plasma levels in COVID-19 appeared associated with high mortality. In two studies, all deceased patients with COVID-19 had high D-dimer values [25, 29]. D-dimer values were found up to 14 times higher in deceased patients with COVID-19 than in survivors with moderate COVID-19 [29]. In patients with high D-dimers and low compliance of the respiratory system, the 28-day mortality was found 4.3 times higher compared to those with low D-dimers and high compliance (56% and 13%, respectively) [30]. CT also revealed filling defects or occlusions of the pulmonary vasculature that were more prominent in patients with high D-dimer levels (94% of them had bilateral, diffuse areas of hypoperfusion, consistent with the presence of thrombi or emboli) indicating that pulmonary vascular thrombosis could be the main cause of COVID-19-related death [30]. Pulmonary vascular thrombosis and the relatively high incidence of PE can also explain the high prevalence of pulmonary hypertension (PH) with dilated RV and relatively small left ventricular (LV) in symptomatic COVID-19, as well as the autopsy findings were RV dilation with concurrently reduced LV diameters was the most significant macroscopic cardiac abnormality [19, 22, 31]. In one study, RV dilation with RV:LV diameter ratios > 1.1 was found at autopsy in 41% of patients who died of COVID-19 [28]. In hospitalized patients, RV dilation and systolic pulmonary arterial pressure (PAPs) appeared significantly correlated with D-dimer levels [32].

Myocardial injuries, revealed by elevated cardiac biomarkers, were detected in up to 30% of hospitalized patients with COVID-19, and in those with preexisting cardiovascular diseases, the prevalence of elevated biomarkers reached 55% [2]. The contribution of cardiac dysfunction to the mortality of COVID-19 patients, either alone or in conjunction with respiratory failure, can reach 43% [33]. In-hospital COVID-related death was found twice as high in patients with HF history than in those without [34]. In one study, 20% of the hospitalized COVID-19 patients showed increased high-sensitive troponin I (hs-TnI) blood levels with or without additional abnormalities in electrocardiography (ECG) or echocardiography (ECHO) and in these patients the mortality reached 51%, whereas in those with normal hs-TnT, the mortality rate was only 4.5% [33]. Increased hs-TnI was identified as an independent risk factor for COVID-19-related mortality [35, 36]. In a multicenter study, 62% of the hospitalized COVID-19 patients had

bio-marker evidence of myocardial injury (high troponin), but myocardial injury alone was not a predictor of increased mortality [13]. In those with high troponin accompanied by ECHO abnormalities, the inhospital mortality reached 31.2%, and high troponin in combination with ECHO evidence of RV dysfunction was associated with a threefold increased risk of death [13]. In another study, the mortality rate for patients without any underlying cardiovascular disease (CVD) and normal TnT levels was 13%, whereas in patients with an underlying CVD plus elevated TnT, the mortality reached 69.4% [37]. A prospective study also revealed hs-TnI as the most relevant predictive factor for ICU referral in COVID-19 patients and hs-TnI appeared significantly correlated with sPAP values and TAPSE [32]. Given that in PE not related to COVID-19, the combined effect of RV afterload mismatch and myocardial ischemia can lead to progressive RV dysfunction associated with elevated TnT [38]; the high troponin levels and their prognostic value in severe COVID-19 could also be explained by the deleterious impact of the pulmonary microthrombosis-induced RV myocyte damage related to the high wall stress caused by acute overload of the RV [32]. The detection of RV dysfunction in 50% of the patients with COVID-19 associated with elevated TnI, but normal LV function, supports this assumption [39]. In a pediatric study, the RV free wall longitudinal strain was the strongest predictor of myocardial injury defined by elevated brain natriuretic peptide (BNP) and/or TnI levels [40].

Acute heart failure (HF) can be the primary presenting manifestation of COVID-19 [14, 18]. It was present in 23% of patients with COVID-19 at their initial presentation and in 52% of those who finally died in the ICU [14]. There is evidence that pulmonary microvascular damages and occlusion by neutrophil aggregates and platelet–fibrin thrombi responsible for the particularly severe acute respiratory failure may have also an important contribution to the development of COVID-19-related acute HF [21].

Given the high risk of life-threatening acute cardiorespiratory failure triggered by the extensive pulmonary vessel thrombosis, clinical deterioration of the respiratory function or the lack of improvement despite of supportive therapy, together with laboratory evidence of severe inflammation and hypercoagulable state, as well as the appearance of the first ECHO and/or ECG signs of RV pressure overload should prompt rapid diagnostic procedures to identify patients with pulmonary in situ microthrombosis or PE [41]. In this regard, particularly noteworthy is the detection of ECG signs suggesting acute RV pressure overload in 30% of hospitalized patients with COVID-19 [42]. This indicates that ECG could be particularly useful for rapid identification of patients at higher risk for RV failure already at the time of admission to the hospital and close monitoring of hospitalized patients by ECG could also be useful for the timing of ECHO examinations.

LV dysfunction appeared not to be a major cause for COVID-19-associated HF [16, 19, 22]. One study found no differences in the incidence of congestive HF between COVID-19 and non-COVID-19 patients with ARDS [11]. An ECHO study revealed no differences in the LV ejection fraction (LVEF) between patients treated in the ICU and those who did not need intensive care [19]. In another study, most patients with elevated troponin had preserved LV function [13]. In an autopsy study where thrombotic microangiopathy was identified in all lungs, the coronary arteries showed no significant stenoses or acute thrombus formation [22].

Although acute myocarditis is a recognized severe complication of the disease, the initially presumed relevant contribution of to the morbidity and mortality of COVID-19 is still debated. In a large recent multi-center study, acute myocarditis was confirmed only in < 1% of patients with COVID-19 [43]. An autopsy study showed no evidence of myocarditis and the lack of relevant myocardial necrosis which could impair myocardial pump function explains the relatively few cases of COVID-19-associated severe LV dysfunction [22].

Pathophysiology of cardiovascular involvement in COVID-19

Similar to SARS-CoV, SARS-CoV-2 also uses the angiotensin-converting enzyme 2 (ACE2) receptor to invade pulmonary alveolar cells and other intra- or extra-pulmonary cells with ACE2 receptors [2, 44]. The higher affinity of binding to the ACE2 receptor revealed by SARS-CoV-2 may contribute to the increased transmissibility of COVID-19 and the usually higher degree infection severity [44]. ACE2 receptors are highly expressed in cardiovascular tissue, and SARS-CoV-2 may spread via the bloodstream and produce direct viral injury [2]. However, the pathophysiology underlying cardiac involvement during SARS-CoV-2 infection is likely to be multifactorial (direct viral toxicity, cytokine-mediated endothelial damage and thrombotic events, overloading-induced RV dysfunction) [2].

The pathophysiologic role of the heart–lung interactions appeared mainly involved in the development and progression of severe RV dysfunction (Table 1; Fig. 1), which in turn has a major contribution to the poor prognosis of CARDS. Due to the particularly distinctive hyper-inflammatory and pro-thrombotic state, extensive pulmonary in situ microthrombosis and PE rather than pneumonia appeared to be more often not only the major cause of respiratory failure but also the major cause of acute decompensated HF, where the excessively increased RV afterload has a crucial pathogenic role. Acute RV failure is a serious clinical challenge regardless of its etiology because it complicates the use of recommended treatment strategies in critically ill patients

Table 1 Heart–lung interactions in patients with COVID-19

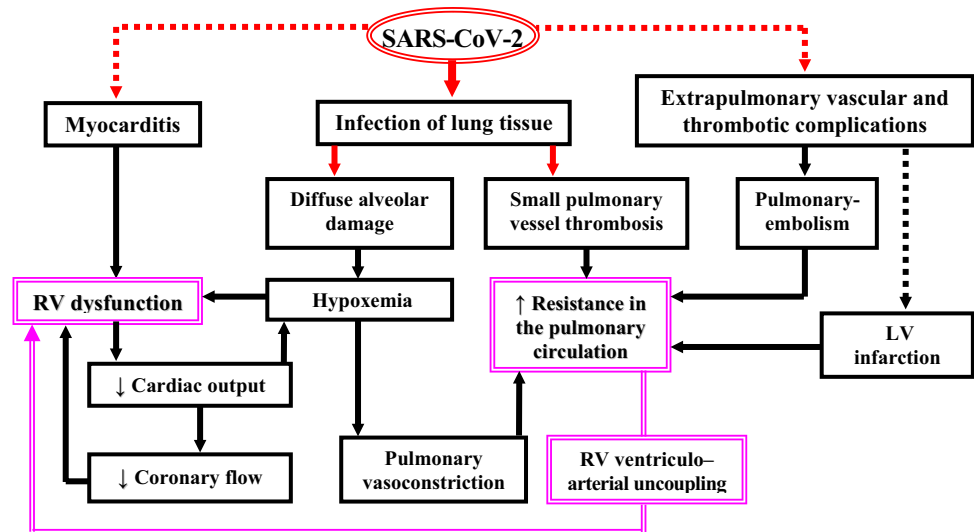
Studies	Relevance of heart–lung interactions in COVID-19
Fox et al. [22, 31]	<ul style="list-style-type: none"> • Diffuse alveolar damage accompanied by thrombosed small vessels in all lungs, as well as RV dilation present in 41% of the deceased patients were the most significant autopsy findings
Dolhnikoff et al. [24]	<ul style="list-style-type: none"> • Autopsy revealed a variable number of small fibrinous thrombi in small pulmonary arterioles in areas of both damaged and more preserved lung parenchyma in 8 out of 10 cases
García-Cruz et al. [59]	<ul style="list-style-type: none"> • In the ICU, the most frequent ECHO findings were elevated PAPs, RV dilatation and RV dysfunction (detected in 69.5%, 28% and 27% of patients, respectively) • PAPs > 35 mmHg increased the probability of in-hospital mortality
Zeng et al. [19]	<ul style="list-style-type: none"> • 29% of patients treated in the ICU had PAPs values > 40 mmHg
Mahmoud-Elsayed et al. [60]	<ul style="list-style-type: none"> • The chief ECHO abnormalities in 78 investigated patients were RV dilatation and RV dysfunction in 41% and 27%, respectively. RV impairment was associated with increased D-dimer levels • LV function was hyperdynamic or normal in most (89%) patients
Goudot et al. [32]	<ul style="list-style-type: none"> • In hospitalized patients (56% of them in the ICU) hs-TnI and D-dimer levels were significantly correlated with both PAPs and TAPSE. Additionally, D-dimer levels correlated also with RV dilation ($p=0.01$) • The high predictive value of elevated hs-TnI for ICU referral in patients without LV alterations, together with the correlation of D-dimer with RV dilatation, confirms the relevance of COVID-19-associated pulmonary microvasculature thrombosis-induced RV afterload increase with the resulting high wall tension-induced RV myocardial damage and dysfunction
Giustino et al. [13]	<ul style="list-style-type: none"> • 26% of patients with biomarker evidence of myocardial injury had ECHO signs of RV dysfunction which was associated with a three-fold increased risk of death
Argulian et al. [61]	<ul style="list-style-type: none"> • ECHO revealed RV dilation in 31% of the hospitalized patients with similar prevalence of major co-morbidities, laboratory markers of myocardial injury or inflammation, and also similar LV size and EF compared with those without RV dilation • Nearly 4 times higher mortality rate in patients with RV dilation than in those without dilation (41% and 11%, respectively). On multivariate analysis, RV enlargement was the only variable significantly associated with mortality
Li et al. [62]	<ul style="list-style-type: none"> • ECHO monitoring revealed that compared with survivors, non-survivors displayed enlarged right heart chambers, reduced RV function and elevated PAPs • RVLS, FAC_{RV}, and TAPSE were associated with higher mortality (area under the curve 0.87, 0.72 and 0.67, respectively)
D'Alto et al. [16]	<ul style="list-style-type: none"> • COVID 19-induced ARDS is associated with early RV-PA uncoupling. 25 (67%) of the 37 invasively ventilated patients died. Non-survivors had nearly 3 times higher D-dimer plasma levels ($p < 0.001$) as well as higher PAPS and lower TAPSE ($p < 0.001$). Reduction of TAPSE/PASP was identified as an independent predictor of mortality. In non-survivors LV size was normal, and LVEF was $\geq 50\%$
Szekely et al. [39]	<ul style="list-style-type: none"> • RV dilation and dysfunction were the most frequent ECHO abnormalities (detectable in 39% of patients) with a 5 times higher incidence than the reduction of LVEF. RV dysfunction correlated with troponin levels. • During follow-up, 50% of patients showing clinical deterioration revealed also further worsening of RV parameters, whereas the LV function remains normal in all those patients
Schott et al. [63]	<ul style="list-style-type: none"> • ECHO revealed RV dilation and increased RV/LV ratio in the majority of hospitalized patients (74% and 82%, respectively), whereas reduction of LVEF was detected only in 3% of the patients
Rivinius et al. [64]	<ul style="list-style-type: none"> • In mechanically ventilated heart transplant recipients with COVID-19, the mortality of can reach 87% and appeared associated with RV dysfunction accompanied by increased PAP, whereas none of them showed LVEF reduction
Caravita et al. [65]	<ul style="list-style-type: none"> • The prevalence of PH (mean PAP ≥ 25 mmHg) was 4 times higher in ventilated COVID-19 patients with ARDS compared to those without the need for mechanical ventilation (76% vs. 19%, $p < 0.001$)
Doyen et al. [66]	<ul style="list-style-type: none"> • Patients with cardiac injury (increased hs-TnI) experienced 3.6 times more RV than LV systolic dysfunction (47% and 13%, respectively)

RV right ventricle, ICU intensive care unit, ECHO echocardiography, PAPs systolic pulmonary arterial pressure, LV left ventricle, EF ejection fraction, RVLS right ventricular longitudinal strain, FAC_{RV} right ventricular fractional area change, TAPSE tricuspid annulus peak systolic excursion, PH pulmonary hypertension, PAP pulmonary arterial pressure, ARDS acute respiratory distress syndrome, hs-TnI high-sensitive troponin I

and its presence is associated with high mortality [45]. The mechanical ventilation in patients with ARDS can reduce the venous return due to the positive intrathoracic pressure and increase the RV afterload by lung volume changes which can markedly alter the pulmonary vascular resistance (PVR) [45]. The increased PVR leads to RV-pulmonary arterial

(PA) uncoupling with subsequent development and/or progression of RV dysfunction [37]. These side effects of ventilation can be exacerbated by the use of high positive end-expiratory pressure (PEEP) and were observed particularly often in CARDS accompanied by pulmonary small vessel thrombosis [15, 46]. This also explains why decreased PEEP

Fig. 1 Pathophysiologic role of heart–lung interactions for the development and progression of life-threatening RV dysfunction during SARS-CoV-2 infection. The bold red arrows indicate the major direct impact of the virus on the lung tissue. The red dotted arrows indicate other possible direct cardiovascular damages of the virus with prognostic relevance



appeared independently associated with improved survival of patients with severe COVID-19 [15].

RV dilation and dysfunction also affect the LV loading and ejection by ventricular interdependence. The resulting reduction of the cardiac output, which impairs the blood supply to vital organs, can further aggravate the critical condition of the patient by exposing the overloaded RV to ischemia. In CARDS all this is amplified by the pulmonary microangiopathy with small vessel thrombosis which explains the particularly high incidence of cardiorespiratory failure in critically ill COVID-19 patients [2, 19].

A possible cause of COVID-19-related cardiac dysfunction is also the development of myocarditis which is a recognized severe complication of the virus infection [2]. The exact pathophysiology of SARS-CoV-2-associated myocarditis remains still not clearly definable. Direct virus-induced myocardial injury is possible since cardiomyocytes express a high level of ACE2 [47]. However, isolation of the virus from myocardial tissue was reported only in few autopsy studies and pathological reports have also described inflammatory infiltrates without myocardial evidence of SARS-CoV-2 [20, 48]. Virus-host interactions with participation of both innate and acquired immune responses, as well as autoimmune mediated mechanisms also contribute to SARS-CoV-2 myocardial injury. In a clinical study without availability of myocardial biopsies, although 12.5% of the investigated COVID-19 patients had cardiac abnormalities that could indicate myocarditis, the changes of cardiac hs-TnI over time and the absence of typical signs on ECG and TTE have suggested that myocardial injury might have been more likely related to indirect deleterious effects resulting from hyperactivation of inflammation and/or consequences of altered heart–lung interaction rather than direct virus-mediated cell damage [49]. Thus, although signs of infection can be detected in myocardial tissues, cardiac involvement

in COVID-19 is possibly more integrated with systemic disorders. This supports the current recommendation to be rather reluctant to make use of cardiac biopsy for COVID-19 patients with suspected myocarditis because of the high risks of spreading the virus.

Impact of ventilatory support on heart–lung interactions

Although the entrance portal for SARS-CoV-2 is inhalational, and alveolar infiltrates are commonly detectable, the CARDS usually includes a major pulmonary vascular insult (endothelial injury with highly activated coagulation cascade) with particularly severe impairment of heart–lung interaction that may also require specific treatment approaches than usually applied for ARDS. The large disparities in mortality rates across different ICUs may also indicate that the approach to ventilatory management could indeed affect patient outcome [50].

The hemodynamic effects of mechanical ventilation (MV) are mainly induced by changes in pleural pressure and transpulmonary pressure (TP). MV can reduce the venous return due to the positive intra-thoracic pressure and increase the RV afterload by increasing the TP [46]. Tidal forces and PEEP increase PVR in direct proportion to their effects on mean airway pressure [46]. The particularly recommended lung protective ventilation strategies in CARDS refer mainly to PEEP, which, by modifying both stress and strain, contributes to the applied mechanical power [50, 51]. Whereas optimal PEEP is beneficial by acting as a counterforce against superimposed pressure, thereby preventing atelectasis, and too low PEEP can induce atelectotrauma, high PEEP levels cause overinflation of the normal alveoli and compression of intraalveolar vessels, which lead to high PVR and

thereby to increased RV afterload [43, 51]. RV dysfunction can therefore evolve as a deleterious consequence of MV. Being an early sign of RV overloading, RV dilatation during the course of MV requires particular attention.

COVID-19 causes quite unique lung injury, and the necessary respiratory support approaches depend largely on the underlying pathophysiology, which varies in accordance with the stage of disease. In the early stage of CARDS, although minute ventilation is characteristically high, many patients are not overtly dyspneic despite very poor oxygenation [50, 52]. In a simplified model, these patients are assigned to “type-L” disease characterized by low lung elastance (high compliance), lower lung weight (estimated by CT), low ventilation/perfusion mismatch, and low response to PEEP [50, 53, 54]. Due to their good lung compliance, these patients accept larger tidal volumes than those usually prescribed for ARDS, without increasing the risk of ventilator-induced lung injury (VILI) [50]. Whereas for many patients, the disease may stabilize at this stage, some may progress to a more characteristic clinical picture of CARDS defined as “type-H,” with extensive lung consolidations, low compliance, higher lung weight, high right-left shunt, and high PEEP response [50, 54]. Between the 2 extremes, there are inter-mediate stages in which the type-L and type-H characteristics may overlap. Despite this over-lapping, it may be helpful to categorize patients as having either L or H phenotype. However, because of the severe widespread endothelial damage that promotes thrombogenesis in small pulmonary vessels and impairs the pulmonary vasoregulation (i.e., failure to develop a normal pulmonary vasoconstriction in response to hypoxia), the resulting ventilation-perfusion mismatch with life-threatening hypoxemia, becomes the key problem already in early CARDS stages [50].

Noninvasive respiratory support (e.g., high-flow nasal O₂, continuous positive airway pressure [CPAP], noninvasive ventilation [NIV]) providing preconditions for the lowest possible mechanical load on the lungs may stabilize the course in mild cases [50, 54]. The support must aim to minimize the volume and TP because high TP during inspiration and large tidal volumes will increase the lung stress and strain [50]. However, if the respiratory drive is not reduced by noninvasive support, the high drive and persistently strong spontaneous inspiratory efforts will result in patient “self-induced lung injury” [SILI] (mainly as a consequence of an excessive global and regional lung stress) followed by pulmonary transvascular pressure rise and fluid leakage with progressive deterioration of lung and RV function [50, 51, 55]. Timely intubation with effective sedation may interrupt this cycle and in patients with compliant lungs targeting higher tidal volumes and lower PEEP (8–10 cm H₂O) will be appropriate [50]. If lung edema increases in type-L patients (because of the disease itself and/or SILI), the shrinking lung (“shrinking baby lung”) shrinks further with a parallel decrease in respiratory system compliance, and

converts progressively into a type-H phenotype [50, 51]. In this advanced state, it is advisable to apply a more conventional lung-protective strategy: PEEP \leq 15 cm H₂O, lower tidal volume (6 mL/kg), and prone positioning while minimizing O₂ consumption [50]. Concentrating the entire ventilation workload on an already overstrained baby lung accentuates its potential for progressive lung injury. The forces of mechanical power (a function of strain, stress, and respiratory rate) are primarily applied to the extracellular matrix to which both the epithelial and endothelial cells are anchored [51]. The strain of these two cell populations contributes through a cytokine release to a further recruitment of inflammatory cells. This inflammatory reaction can severely alter the reaction to and tolerance of repeated cycles of high stress and strain [51]. Therefore, it can be necessary to consider also extracorporeal membrane oxygenation (ECMO) for some carefully selected patients with severe CARDS [50, 54, 56].

An important aspect related to both noninvasive and invasive MV is the negative impact of hyperoxia on the pulmonary microcirculation. This is of particular importance due to the fact that the anatomical and pathophysiological features of COVID-19-related lung infection are different from classic ARDS [57]. Exposure to hyperoxia over several days/weeks can induce complex endothelial, epithelial, platelet, and immune cell responses in the lung resulting in diffuse alveolar damage [58]. Given the spatial heterogeneity of pulmonary involvement in COVID-19, it is likely that segments of the severely damaged and largely consolidated lung experience lesser exposure to hyperoxia, whereas segments that are not consolidated and well ventilated are more prone to VILI and injury induced by hyperoxia. The vascular and immunological features of CARDS may reflect, at least in part, changes ascribed to prolonged hyperoxia that are superimposed on the original COVID-19-related disease, or develop in parallel in the well-ventilated segments of the lung [58]. There is evidence for overlapping pathological effects upon the lung of SARS-CoV-2 infection and exposure to high concentrations of inhaled O₂ and hyperoxia may exacerbate the local cytopathic effects of the virus [58], therefore limiting the exposure to hyperoxia to the lowest levels that are required appeared advisable [58].

Relevance of right heart dysfunction in COVID-19

In a multicenter study, the most frequent ECHO abnormality in patients with COVID-19 was RV dilation with or without dysfunction [39]. In hospitalized patients, two different ECHO studies revealed RV dilation in 28% and 41%, respectively, and RV dysfunction was evidenced in 27% of the patients in both studies [59, 60]. In another study, RV dilation was present in 31% of the hospitalized COVID-19

patients with similar prevalence of major comorbidities, laboratory markers of myocardial injury or inflammation, and also no differences in LV size and EF compared with patients without RV dilation [61]. Severely ill patients can develop isolated RVF, more often associated with increased PVR derived from PE and/or pulmonary small vessel thrombosis [2, 19]. ECHO allowed the detection of elevated PAPs in up to 70% of COVID-19 patients [59]. In hospitalized patients with COVID-19, ECHO revealed also a discrepancy between the involvements of the right-sided and left-sided heart in the prognosis of the disease [62]. Thus, non-survivors displayed significantly enlarged right heart chambers, reduced RV function, and elevated PAPs, whereas the LV and left atrial (LA) size as well as the LVEF were in the normal range [62]. Lower RV longitudinal strain (RVLS), RV fractional area change (FAC_{RV}), and the lateral tricuspid annulus peak systolic excursion (TAPSE) as well as RV dilation were identified as powerful and independent predictors of higher mortality [61, 62]. In another study (100 COVID-19 patients), RV dilation and dysfunction were the most frequent ECHO abnormalities (detectable in 39 patients), whereas LVEF reduction occurred only in 8 patients [39]. Among the patients developing clinical deterioration during follow-up, 50% showed further deterioration of RV parameters, whereas the LV function remains normal in all those patients [39]. An ECHO study on hospitalized COVID-19 patients revealed RV dilation in 74% of them and the RV/LV ratio, which was increased in 82% of the patients, reached 0.9 ± 0.3 [63]. In that study, the LVEF was 60% (52–74%) and reduction of LVEF was detected in only 3% of the patients. Heart transplanted patients with COVID-19, who needed mechanical ventilation, showed a mortality of 87.5%, which appeared mainly associated with RV dysfunction (at simultaneously increased PAP), whereas none of them showed LVEF reduction [64]. The prevalence of PH (mean PAP ≥ 25 mmHg) was 4 times higher in ventilated COVID-19 patients with ARDS compared to those without the need for mechanical ventilation (76% vs. 19%) [65]. In another study, 29% of COVID-19 patients treated in the ICU had PAP values > 40 mmHg [19]. A previously normal RV (without pre-existing hypertrophy) will be barely able to generate such high pressures for several days, and therefore, such patients are at high risk to develop acute RV failure. In other studies, almost 50% of critically ill patients with COVID-19 revealed thrombotic events (including pulmonary microvascular occlusions) and about 22% experienced myocardial ischemia and/or AMI without signs of congestive HF [2]. In a study which included COVID-19 patients without previously known cardiomyopathy, ECHO parameters on LV function were not different between survivors and non-survivors [62].

Whereas, in general, LV failure is one of the most common etiologies of acute RV failure in the ICU, in critically

ill COVID-19 patients, pulmonary tissue damages and small vessel thrombosis with consequent hypoxemia and excessive increase in the resistance in the pulmonary circulation are the cause of both acute respiratory and RV failure. Patients with cardiac injury (increased hs-TnI) experienced 3.6 times more RV than LV systolic dysfunction (47% and 13%, respectively) [66]. Timely identification of patients at high risk for RV failure can be life-saving for them. Although there are no specific biomarkers for RV failure, markers of myocardial injury like hs-TnI and N-terminal proBNP can help in prognostication because they have shown rapid increase in non-survivors. ECG lacks sensitivity and once chest X-ray or CT reveal RV dysfunction; RVF is usually advanced and associated with high mortality [67]. Pulmonary artery catheters (PACs) and ECHO are the most reliable tools to diagnose RV failure and evaluate the treatment response in the ICU. Their combined use provides crucial information for timely detection of potentially life-threatening alterations in the pulmonary circulation and RV function, as well as for ensuring the best possible patient care, including evaluation of the responses to therapies aimed at reducing RV afterload, optimizing the intravascular volume status and enhancing the RV inotropy. They can also help optimizing mechanical ventilation in order to avoid ventilator-induced lung injury and/or reduce its possible negative effects on RV preload and afterload [9, 68].

In specialized centers, ECHO is indispensable for the management of refractory respiratory failure with and without additional isolated RV failure or cardiogenic shock via artificial cardiopulmonary support strategies [56, 69–72]. Considering that without the option of ECMO therapy the mortality rate of patients with COVID-19-associated ARDS can reach 85% [2], the recently reported inhospital mortality of 38% for ECMO-supported patients with CARDS [72], which is consistent with the mortality rates reported for non-COVID-19 ECMO-supported patients, revealed the existence of important survival benefits achievable by ECMO therapy for appropriately selected patients with COVID-19. However, because in severe cases of RV afterload mismatch the RV output can decrease substantially even in the setting of normal intrinsic RV contractility (i.e., “acute cor pulmonale”), in such situations, attempting to treat the underlying etiology of impaired gas exchange using V-V ECMO alone may not be sufficient because V-V ECMO recirculation is exacerbated by reduced RV ejection and TR [72]. Given the prognostic importance of the relatively high incidence of right heart involvement in COVID-19, close monitoring of RV size, geometry, and function in patients with refractory respiratory failure associated with hemodynamic instability, before overt RV failure develops, can substantially improve therapeutic decision-making [70]. ECHO can help with the selection of candidates for ECMO therapy, selection of the most appropriate support strategy (e.g., V-V ECMO plus catheter-mounted RV assist device [RVAD]) or surgical

RVAD plus gas exchanger), and is also necessary for weaning decision-making from the mechanical support, after cardio-pulmonary recovery [56, 69, 70, 72].

Assessment of the right heart by echocardiography

The frequently normal LVEF even in critically ill patients may have contributed to the initially restraint in use of ECHO for monitoring purposes in hospitalized patients with COVID-19. This is not surprising because for years, the assessment of the RV has lagged behind other ECHO examinations.

The main reasons for that were the historical lack of awareness of the pathophysiological and clinical importance of the RV function, the intense clinical focus of the left heart, and the paucity of normative data. More recently, RV function was found to be a powerful prognostic factor in HF and PH of various etiologies. Since mechanical circulatory support has become a well-established life-saving treatment for drug-refractory cardiorespiratory failure, assessment of RV size, geometry, and function has emerged as a crucial component of cardiac evaluation [69]. In patients with refractory respiratory failure associated with hemodynamic instability, TTE allows timely recognition of RV dysfunction (i.e., before overt RV failure develops) and could therefore spare some patients from additional ECMO therapy. Cardiologists already warned years ago that “it is imprudent to ignore the RV any longer” [73].

The increasing role of point-of-care (PoC) ECHO in the management of patients with undifferentiated shock in the emergency departments was also recently emphasized [74]. Given that pulmonary vessel injuries and blood flow alterations followed by RV failure are major causes of COVID-19-related deaths, it becomes increasingly difficult to follow the initial recommendation to avoid unnecessary cardiac imaging (including TTE) in order to reduce transmission of the virus and protect health care professionals because, without any ECHO examination, it cannot be established whether ECHO is indeed unnecessary in a given patient. Whereas the restraint use of trans-esophageal ECHO is fully justified by the high risk of the dispersion of droplets and aerosols, focused TTE protocols can provide a favorable benefit/risk balance in especially in patients with cardiovascular instability and/or suspicion of RV dysfunction [75, 76]. With the ongoing COVID-19 pandemic, the increasingly recommended limited TTE with focused protocols, in particularly bedside PoC-TTE, can be indeed a good solution which considers both clinical and epidemiological concerns related to the management of hospitalized patients with COVID-19 [2, 76]. PoCUS, primarily focused on the pulmonary manifestations, can be particularly useful in the triage and decision making for COVID-19 patients [75, 76].

Compared with the LV, the RV is less suited for quantitative ECHO-analysis, mainly because of its complex geometry with a limited number of well-defined anatomical landmarks and its variable position in the thorax. Because of the relevant impact of LV size and function on RV geometry and performance, the complex and different contraction-relaxation mechanism among the main segments of the RV, and the particularly high dependency of RV size, geometry, and function on RV loading conditions, even in the presence of unaltered myocardial contractility, collection of ECHO-data and especially their accurate interpretation is often a very challenging task [73, 77].

Main objectives of right heart ECHO in severe COVID-19

Detection and monitoring of right-sided heart dilation, RV dysfunction, tricuspid regurgitation (TR), and alterations of inferior vena cava size and blood flow are major goals of ECHO examinations in critically ill COVID-19 patients. In patients with RV dysfunction, it is also important to establish whether dysfunction is induced by pressure overloading or by impaired contractility or even by both.

ECHO-assessment of right heart anatomy

The apical 4-chamber (A4C) view is critical for assessment of right heart size and geometry, but the measurements rely on good visualization of the RV free wall and lateral tricuspid annulus. If apical images are poor, subcostal windows may provide similar views of the RV, but the difficulties in defining the anteroposterior plane of the RV from this window can induce misinterpretations in the evaluation of RV size [78]. Parasternal views allow measurements of RV outflow tract (RVOT) dimensions but the RVOT should be measured in both parasternal long-axis and short-axis views, in order to avoid measuring errors generated by patient position during the acquisition of RVOT images or rotation of the RVOT out of the section plane [77, 78]. During a limited bedside TTE, an A4C view is usually sufficient for assessment of right heart anatomy (including the TA) and the relationship between the size of right and left heart cavities, as well as for assessment of IVS and IAS position.

ECHO-assessment of right ventricular function

Assessment of RV function is more challenging than the assessment of its anatomical characteristics, and the interpretation of functional parameters for the RV is more demanding than for the LV [70]. RV pump performance depends mainly on myocardial contractility and loading conditions (preload and afterload) and is also influenced by heart rhythm, valve function, ventricular interdependence, and synchrony of ventricular contraction [73, 74].

Mechanical and timing heterogeneity even in normal RVs make the assessment of systolic function particularly challenging. For objective reasons (e.g., complex RV shape necessitating geometric assumptions for volume calculations, limited imaging window, limited definition of end-systolic RV endocardial surface), RVEF calculations from 2D-ECHO-derived measurements are less reliable and therefore not anymore recommended for clinical use [78, 79]. Nevertheless, 2D-ECHO is useful for assessment of RV function because it allows easy measurements of FAC_{RV} and TAPSE, which provide similar information to RVEF [80–82]. TAPSE and FAC_{RV} require no geometric assumptions, and TAPSE can be obtained even with poor image quality [77, 78]. During a limited bedside TTE, an A4C view (preferably RV-focused view) is sufficient also for assessment of RV systolic function because both FAC_{RV} and TAPSE are measured in this view. An additional subcostal view can provide useful data for assessment of TR.

There is evidence that severe COVID-19 ARDS can be associated with a specific phenotype of RV radial impairment with sparing of longitudinal function and TAPSE may become therefore less useful for detection of RV dysfunction [78]. In one study, although reduced FAC_{RV} was present in about 70% of critically ill COVID-19 patients, TAPSE remained normal in nearly 80% of these patients [83]. However, in another study, patients with COVID-19-induced ARDS, those who survived, had significantly higher TAPSE values ($p < 0.001$) compared to those who died in the ICU [16]. A limitation of RVEF, FAC_{RV} , and TAPSE is the fact that due to their load dependency, they can change without changes in RV myocardial contractility. Thus, they will decrease with increasing PVR, even if myocardial contractility remains unaltered [81, 84, 85]. Additionally, TR can induce RVEF, FAC_{RV} , and TAPSE changes which can be misleading for assessment of RV contractile function and estimation of RV myocardial contractility [77–79, 81, 85–87]. TR can induce overestimations of RV contractile function by facilitating the RV free wall transversal (inward) and longitudinal motion which will increase the measured TAPSE, FAC_{RV} , and EF, correspondingly to the increased blood volume leaving the RV in systole [85].

Alongside with the FAC_{RV} , also the RVOT velocity–time integral ($RVOT_{VTI}$) can be useful for detection of RV dysfunction in hospitalized patients with COVID-19 [83]. Thus, reduced $RVOT_{VTI}$ (i.e., < 19 cm) was present in about 85% of critically ill COVID-19 patients, whereas the RV free wall strain was reduced in only about 30% of the patients [83].

The major challenge in assessing the RV involvement in the prognosis of COVID-19 is the appropriate incorporation of ECHO measurements into an integrative evaluation of the right-sided heart in connection with the RV myocardial contractile abilities and the impact of heart–lung interactions on RV size, geometry, and pump performance [88]. There

is no individual ECHO-parameter, irrespective of whether derived from conventional ECHO, 3D-ECHO, tissue Doppler, or deformation (strain and strain rate) imaging, which can alone enable the distinction between high afterload-induced and impaired contractility-induced RV dysfunction (Table 2).

Distinction between right ventricular dysfunction with and without impaired myocardial contractility

A crucial question which arises particularly in patients with severe COVID-19 is whether RV dysfunction is induced by pressure overloading (afterload mismatch) or by impaired contractility, or by both.

Assessment of RV-PA coupling, using the FAC_{RV}/RV systolic pressure (RVSP) and TAPSE/RVSP as indices of RV contractile function and prognosis [89], could therefore be useful in assessing and monitoring of RV function in patients with COVID-19 [16]. Using a standard cut off for normal function of 1.0, $FAC_{RV}/RVSP$ identified 86% of hospitalized patients with COVID-19 as having RV-PA uncoupling [83]. Significant uncoupling ($FAC_{RV}/RVSP < 0.6$) was detectable in 50% of ICU-treated patients with COVID-19 [83]. In patients with COVID-induced ARDS, the mortality rate within the first 20 days was twice as high in those with a TAPSE/PASP ratio < 0.635 , and reduction of TAPSE/PASP was identified as an independent predictor of mortality [16].

RV myocardial contractile abilities are also reflected by the relationship between RV pressure load and alteration of its geometry and size. Integrative approaches using parameter combinations which consider also the RV afterload, appeared particularly useful for evaluation of RV function [85, 90, 91].

An easy approach to assess this relationship by ECHO is the calculation of the “RV load-adaptation index” (LAI_{RV}), a composite variable based on the relationship between RV load and cavity dilation, also taking the RA pressure into account (i.e., ratio between the systolic mean ΔP_{RV-RA} and RV end-diastolic volume per long-axis length) [85, 91]. Using the TR velocity–time integral (VTI_{TR}) instead of ΔP_{RV-RA} and the RV end-diastolic area (A_{ED}) instead of the RV end-diastolic volume (EDV), the LAI_{RV} defined accordingly as $LAI_{RV} = VTI_{TR} \cdot L_{ED}/A_{ED}$ becomes an easy calculable dimensionless index (Fig. 2) obtainable from a single ECHO view (i.e. A4C). A small RV area relative to long-axis length (unaltered size and geometry) in patients with high RV systolic pressure and relatively low RA pressure (i.e., high VTI_{TR}) yields a high LAI_{RV} which indicates good adaptation to load (suggesting a good RV contractility) and the potential of the RV to improve its performance after reduction of loading conditions. A large A_{ED} relative to the long-axis length (spherical dilation) in patients with a relatively low VTI_{TR} yields a low LAI_{RV} indicating poor adaptation to load

Table 2 Simple parameters and indices obtainable by bedside limited transthoracic echocardiography for assessment of the right heart in patients with high resistance in the pulmonary circulation [56, 63, 68, 70, 77, 81, 82, 85, 86, 88, 91]

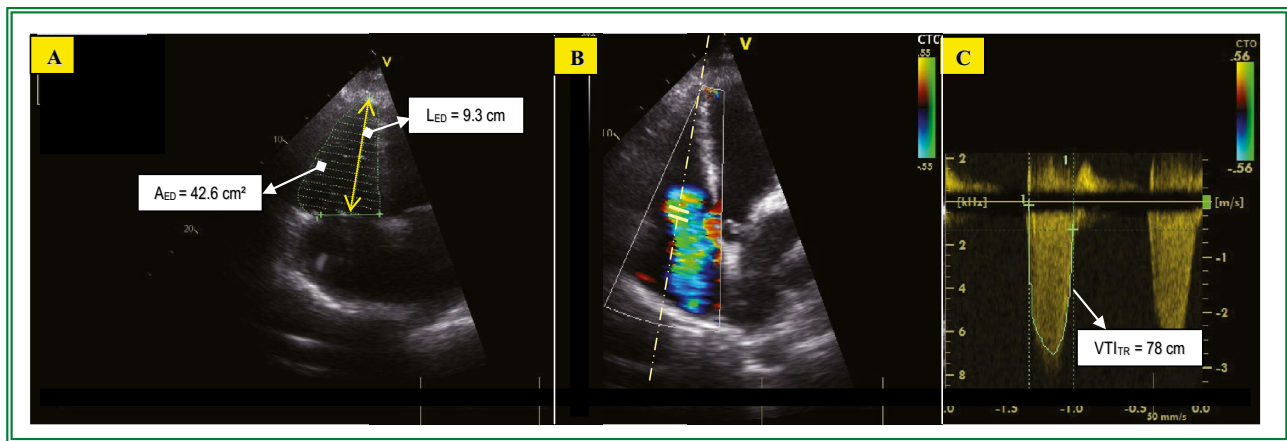
Parameters and indices	RV dysfunction due to high resistance in the pulmonary circulation	Preserved RV contractile function	Reduced RV contractile function
RVEDA, RVEDA	<ul style="list-style-type: none"> • Increased 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
S/L RV axis ratio	<ul style="list-style-type: none"> • Increased (> 0.57) 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
RA size	<ul style="list-style-type: none"> • Increased 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
Atrial septum	<ul style="list-style-type: none"> • Shift to left 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
TR	<ul style="list-style-type: none"> • New appearance or aggravation of preexistent TR 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
Ventricular septum	<ul style="list-style-type: none"> • Shift to left 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
RVEDD/LVEDD	<ul style="list-style-type: none"> • Increased (> 0.65 up to ≥ 1.0) 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
FAC_{RV}	<ul style="list-style-type: none"> • Reduced (< 0.35%) 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
TAPSE	<ul style="list-style-type: none"> • Reduced (< 1.6 mm) 	<ul style="list-style-type: none"> • No specific change* 	<ul style="list-style-type: none"> • No specific change*
SV	<ul style="list-style-type: none"> • Progressive reduction 	<ul style="list-style-type: none"> • Reduced only at excessively high afterload 	<ul style="list-style-type: none"> • Reduced already at moderate afterload increase
ΔP_{RV-RA}^{**}	<ul style="list-style-type: none"> • Initially progressive increase, then progressive reduction to low values 	<ul style="list-style-type: none"> • Increased (> 35 mmHg) 	<ul style="list-style-type: none"> • Reduced (no defined cut-off value)
FAC_{RV}/RVSP and TAPSE/RVSP (RV-PA coupling)	<ul style="list-style-type: none"> • Reduced (FAC_{RV}/RVSP < 1; TAPSE/RVSP < 0.635) <p>The initial decrease is mainly due to the RVSP increase, followed by further decrease also due to SV reduction</p> <p>Later, despite the additional reduction of myocardial contractility both FAC_{RV} and TAPSE reduction can be counterbalanced by the increasing TR which reduces the diagnostic relevance of the ratios</p>	<ul style="list-style-type: none"> • Reduction is likely in patients with only mild to moderately reduced FAC_{RV} and/or TAPSE in the presence of excessively high RVSP 	<ul style="list-style-type: none"> • Reduced in patients with relevantly reduced FAC_{RV} and/or TAPSE in the presence of only moderately increased RVSP
LAI_{RV}^a	<ul style="list-style-type: none"> • Initially progressive increase (preserved adaptability to load), then progressive reduction to low values (afterload mismatch) 	<ul style="list-style-type: none"> • High (> 18) 	<ul style="list-style-type: none"> • Low (< 15) indicates the necessity for temporary mechanical RV support

ECHO echocardiography; *RV* right ventricle; *RVEDA* and *RVEDA* RV end-diastolic diameter and end-diastolic area, respectively; *S/L* short/long axis ratio; *SV* stroke volume; *RA* right atrium; *TR* tricuspid regurgitation; *LVEDD* left ventricular end-diastolic diameter; *FAC_{RV}* fractional area change; *TAPSE* tricuspid annulus peak systolic excursion; ΔP_{RV-RA} pressure gradient between *RV* and *RA* during systole; *RVSP* RV systolic pressure; *RV-PA* right ventricular-pulmonary artery; *LAI_{RV}* RV load adaptation index

*Parameter alteration indicates RV alteration which could be caused by both high afterload and decreased myocardial contractility (etiopathogenetic differentiation not possible)

**Useful for differentiation between patients with and without reduced RV contractile function

^aThe LAI_{RV} can be useful to predict therapy-refractory RVF in both pre- and post-capillary PH



$$LAI_{RV} = \frac{VTI_{TR} \text{ (cm)} \cdot L_{ED} \text{ (cm)}}{A_{ED} \text{ (cm}^2\text{)}} = \frac{78 \text{ cm} \cdot 9.3 \text{ cm}}{42.6 \text{ cm}^2} = 17.0$$

Fig. 2 Calculation of the right ventricular load adaptation index (LAI_{RV}). **a** Measurement of the right ventricular (RV) end-diastolic area (A_{ED}) and long axis lengths in the apical 4 chamber view. **b** and **c** Measurement of the tricuspid regurgitation velocity–time integral (VTI_{TR}) using the continuous wave Doppler. The LAI_{RV} value of 17.0 is reduced and indicates that in this patient with pulmonary

arterial hypertension (pulmonary arterial systolic and mean pressure: 48 mmHg and 37 mmHg, respectively), the RV dilation is more pronounced than one would expect on the basis of its present afterload. The limited adaptation possibilities to the increased afterload are exceeded, and a reduction of pulmonary vascular resistance is highly needed in order to prevent further aggravation of RV dysfunction

(suggesting a reduced RV myocardial contractility). LAI_{RV} values < 15 indicate severely impaired adaptability to load being usually insufficient to prevent RV failure even after normalization of PVR [77]. Good adaptability to high afterload does not exclude a reduced RV pump function which will occur during afterload mismatch. The LAI_{RV} allows distinguishing between pressure overloading-induced and impaired contractility-induced RV dysfunction, as well as between patients with similar FAC_{RV} , but different RV myocardial inotropy [77]. In PAH, the LAI_{RV} can reach values of 25 even in patients with relevant RV dysfunction, suggesting that in the vast majority of PAH patients, the cause of RV failure is the excessive pressure overload and not a relevantly impaired RV contractility [77, 85, 92]. This explains why RV failure is more likely and more faster reversible than LV failure of similar severity and also the reverse remodeling with normalization of RV function in the overwhelming majority of patients with pre-capillary PH-induced severe RV dysfunction, after they underwent lung transplantation [77, 85]. Therefore, temporary mechanical support of the RV can be taken into consideration in life-threatening COVID-19 [72, 93]. In such cases, differentiation between primary afterload-induced and primary impaired contractility-induced RV failure can be crucial for a successful treatment.

Bedside acquisition of essential ECHO data

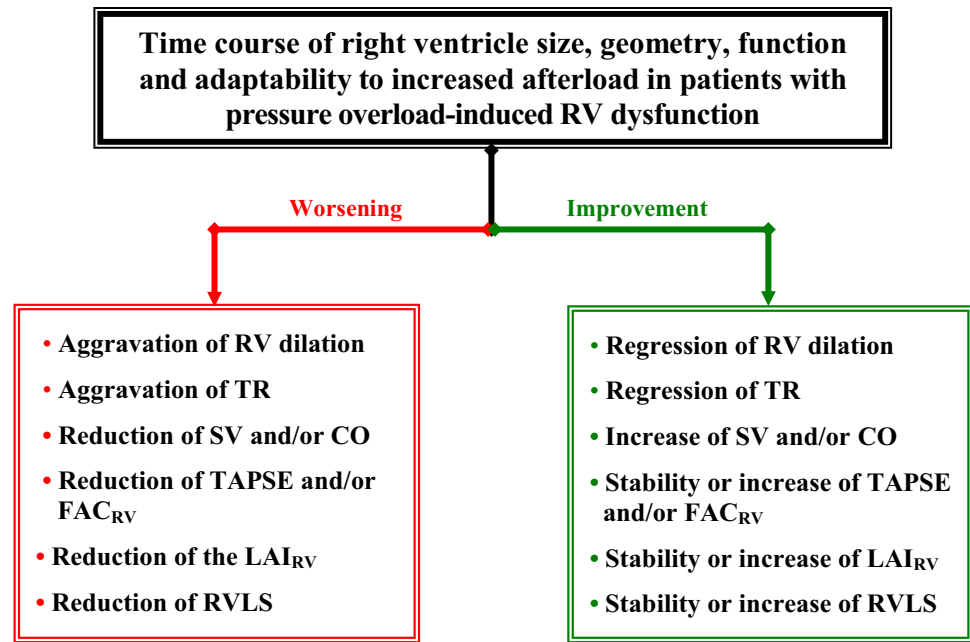
Considering the prognostic importance of right heart involvement in COVID-19, focusing more attention on the

RV during critical care ECHO can help in managing and monitoring patients with confirmed infection. Handheld laptop-based equipment is usually sufficient for the acquisition of essential ECHO data. An A4C view allows answering nearly all the major questions concerning the involvement of the right heart in COVID-19. An additional A5C view (easily acquirable from the A4C) plus a parasternal long-axis view enable the measurements necessary for calculation of the SV. In patients with relevant TR, a subcostal view can provide additional information for the severity of TR. Whereas the necessary TTE measurements are not time consuming, their interpretation remains more often the greater challenge. Although in most patients with severe COVID-19, TTE can provide the information needed, there are situations (e.g., patients with acute circulatory failure), in which critical care goal-directed TEE will become an invaluable diagnostic tool for managing such critically ill patients [94].

Conclusions and further challenges

Extensive pulmonary microangiopathy with small vessel thrombosis associated with particularly severe hypoxemic respiratory failure, high resistance in the pulmonary circulation, and life-threatening HF is a distinctive feature of COVID-19 and also a main cause for the particularly high mortality associated with the disease. Due to the direct alterations and damages of the pulmonary circulation, the pathophysiologic role of the heart–lung interactions becomes

Fig. 3 Major signs of either right ventricular improvement or further deterioration revealed by echocardiographic monitoring of patients with increased resistance in the pulmonary circulation. RV right ventricle, TR tricuspid valve regurgitation, SV stroke volume, CO cardiac output, LAI_{RV} right ventricular load adaptation index, TAPSE tricuspid annulus peak systolic excursion, FAC_{RV} RV fractional area change, RVLS right ventricular longitudinal strain



mainly involved in the development and progression of severe RV dysfunction. By contrast, severe reduction of LV pump function occurred more rarely (especially in patients without preexisting cardiovascular diseases). The close correlations between cardiac troponine and D-dimer levels with both PAPs and RV systolic dysfunction plus the correlation between D-dimer levels and RV dilation underline the pathogenetic significance of the COVID-19-associated pulmonary microvascular thrombosis-induced RV afterload increase. The high afterload-induced increase in RV wall tension associated with RV myocardial damage and dysfunction appeared to be the major cause for the COVID-19-related acute HF and troponin increase is mainly reflecting RV afterload increase.

Being the most impacted side of the heart in COVID-19 and because RV dilatation and dysfunction were often identified as the most prominent ECHO abnormalities, the right heart deserves proper consideration. Along with biological markers like hs-TnI and D-dimers, ECHO can help anticipating a worsening of the patient's condition. Bedside limited TTE focused on the right heart is a useful diagnostic tool in hospitalized COVID-19 patients, particularly in those with suspicion of RV dysfunction or PH and becomes necessary for monitoring of critically ill patients. Even hand-held laptop-based equipment can be often sufficient for acquisition of the strictly necessary ECHO data and it appears advisable to further promote the routine use of this time- and cost-effective diagnostic tool in the management of symptomatic COVID-19 infected patients.

The particular pathophysiology of COVID-19 places the RV at higher risk of failure, and therefore, one key problem remains the timely recognition and severity grading of right

heart involvement and the prediction of RV failure. In this regard, monitoring of RV and RA size, search for TR and assessment of its progress, as well as monitoring of the SV and the pressure gradient between the RV and RA (ΔP_{RV-RA}) are particularly useful (Fig. 3). Assessment of RV-PA coupling using the $FAC/RVSP$, plus evaluation of RV adaptability to pressure loading using the LAI_{RV} , can also allow prognostic statements to be made. Given the growing evidence of therapeutic successes achieved in highly selected COVID-19 patients with life-threatening refractory respiratory or cardiorespiratory failure using ECMO, whereas patients who are inappropriately selected stand a much lower chance of survival, the role of bedside TTE in identifying suitable patients and selection of the appropriate type of ECMO will also continue to grow.

Full implementation of bedside limited TTE focused on the right heart, which is currently still under-used, could facilitate more personalized management and treatment of hospitalized patients and can contribute towards reducing the high mortality associated with SARS-CoV-2 infection.

Declarations

Conflict of interest The authors declare no competing interests.

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