Heart failure in COVID-19 pandemic: Challenging management issues

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Abstract
Over these 12 months of coronavirus disease 2019 (COVID-19) pandemic, the patients of heart failure (HF) showed higher risk of severe disease and increased mortality. Their evaluation and treatment when additionally affected with COVID-19 posed many new challenges, as one condition can potentiate the other and there is considerable overlap of their presenting features. The pandemic also impacted the existing health care systems with changed priorities. Known HF patients becoming worse after acquiring the COVID-19 infection or the new development of acute cardiac complications in persons without any prior heart disease- both are novel therapeutic issues which have no perfect answers for want of adequate data and randomised trials. Newer suggestions are emerging in the management of HF with concomitant COVID-19 from experience from various sources. We have to keep our minds open and learn every day.

Keywords: heart failure; coronavirus; COVID-19; management; SARS-CoV-2

Introduction
Coronavirus disease 2019 (COVID-19) pandemic is about to complete one-year of its emergence into our lives. First of all, it jeopardised the existing health care system including the cardiac services with changed priorities all over the globe. It is now clear that this viral disease affects multiple systems with frequent cardiac manifestations with enhanced degree of morbidity and mortality. The treatment of ischemic syndromes, decisions on revascularisation procedures and cardiac surgeries, management of patients with heart failure including the heart transplant recipients and many other cardiac care issues became more challenging in these pandemic months.

Zhou et al found that 23% of their 191 patients of COVID-19 inpatients had heart failure, which was as
high as 52% among the non-survivors [1]. Inciardi and Chen also reported similar trends of heart failure incidence among the Covid-19 patients [2, 3]. Currently there are no randomised trials to elucidate rationale therapies and HF management algorithms, due to the acuteness of the pandemic and lack of prior experience with COVID-19 infections. Lack of clarity or uniformity prompted us to review available literature on the impact of COVID-19 on heart failure management.

Possible mechanisms of cardiac dysfunction and heart failure
Heart Failure is common enough to be encountered at different stages in the course of the COVID-19 pandemic. There are multiple ways by which a prior HF patient may worsen, or a new onset HF can occur in a healthy individual.

1. Acute cardiac injury
Elevated cardiac troponins or new ECG or echocardiographic abnormalities are considered to indicate acute cardiac injury. Wang D et al showed that as many as 7.2% of all COVID-19 patients (22% of those requiring ICU admission) presented with acute cardiac injury [4]. However, National Health Commission of China reported that as many as 12% of patients, without prior CV disease or any cardiac arrest can show evidence of acute cardiac injury [5]. A level of >99 percentile URL hs-cTnl was noted in 46% of non-survivors versus 1% among the survivors in Wuhan study [1]. Those with higher levels of troponins had more complications and higher mortality. The need for mechanical ventilation and circulatory support was higher in them [6-8]. Studies varied in their definition of cardiac injury and timing of the sampling, based on cTn levels; it is suggested to have serial measurements and report their data in a universally accepted template designed by Sandoval et al [9].

2. Generalised inflammation and acute myocarditis
In advanced stages, there can be a state of cytokine storm with increased IL-3, IL-6 and IL-7, granulocyte-colony stimulating factor, interferon-gamma inducible protein-10, monocyte chemoattractant protein 1, macrophage inflammatory protein 1-alfa and tumour necrosis factor alfa and ferritin. The inflammatory status and production of cytokines secondary to infection increases blood viscosity and coagulopathy. Endothelial dysfunction, fluid and electrolyte imbalances add up resulting in decompensation of the sub-clinical heart failure [10, 11].

Acute myocarditis can manifest as a part of generalised inflammatory state or independent of it. In sporadic cases, histopathological confirmation showed low-grade inflammation with non-specific myocardial changes and low or absent myocyte necrosis [12-14]. Among those with elevated TnT or ECG abnormalities suggestive of acute cardiac injury, a small percentage of patients presented with florid symptoms of myocarditis with markedly raised inflammatory markers or MRI evidence for inflammation, most often, with a rapidly progressive down-hill course. Tomasoni et al and Luetkens et al observed generalised myocardial oedema in absence of LGE at cardiac MRI and these patients had normal echocardiogram findings and elevated biomarkers of cardiac injury [15, 16].

While some of those presenting with myocarditis were a component of the cytokine storm with multi-organ dysfunction, a few were isolated myocarditis or presented with associated respiratory failure. Most of such cases succumbed fatally after a period of circulatory shock and arrhythmias [17, 18].

3. Endothelitis
In these last 12 months it was observed that there is sharp rise in cases of STEMI and when taken up for primary PCI, the operators were surprised to see no significant culprit lesion in most cases. Some of these could be explained as a type of endothelitis involving the microcirculation in COVID-19 infection. This could be an early stage of systemic inflammatory response to infection especially in those with pre-existing cardiovascular disease (CVD) [19].

4. Hypoxemia induced myocardial dysfunction
Pneumonia is a predominant manifestation of COVID-19 affection. Extensive pneumonia leads to hypoxemia and subsequent myocardial damage. There can be pulmonary hypotension and right heart dysfunction as well. Use of elevated positive end-expiratory pressure during mechanical ventilation also adds to the right ventricular afterload and wall-stress.
5. Stress-cardiomyopathy
Jabri et al studied the incidence of stress-cardiomyopathy (SCMP) among 1914 ACS patients with COVID test negative reports during the COVID pandemic compared to historical controls from 2018 and 2019. The incidence of SCMP was 7.8% vs 1.5%. The length of hospital-stay was longer but there was no difference in mortality or 30-day re-hospitalisation rates. Minhas documented a case of stress-cardiomyopathy in the setting of COVID-19, which he claims as the first ever such case in USA [29, 30]. These observations suggest that in a few cases, severe stress itself can produce cardiomyopathy with or without getting actual COVID-19 infection [20, 21].

6. Worsening of prior HF
Worsening heart failure in previously diagnosed patients is not an uncommon event, most requiring ICU admission. HF was observed in about 20-30% of the COVID-19 patients admitted to intensive care units, as reported in various centres [1-3]. Myocarditis, acute cardiac injury, cardiac arrhythmias, thrombotic events, fluid imbalances, plaque instability or disruption in guidelines directed medical therapies can all precipitate a decompensated state in a previously stabilised HF patient.

Type of presentations
It is better to consider presentation and prognosis of heart failure in three clinical scenarios-
(a) In those without any known prior cardiac affection,
(b) Those with prior stable heart failure who decompensated after COVID-19 infection,
(c) Those with HF preserved EF.

(a) No prior heart disease
Cardiac affection is not unique to the current COVID-19 pandemic. During the severe acute respiratory syndrome (SARS), Middle East respiratory syndrome (MERS) and Influenza epidemics in the past, acute cardiac injury was noted in a good number of patients resulting in the development of heart failure, arrhythmias, and sudden cardiac death [22]. Severe COVID-19 infection can affect the cardiovascular system even in those without prior heart disease and can manifest as acute cardiac injury, myocarditis, acute heart failure, myocardial infarction (Type 1/II), acute pulmonary embolism, cardiac arrhythmias and stress-induced cardiomyopathy [1].

(b) Prior HF present
Viral infections like Influenza were known to precipitate acute decompensation in heart failure patients [23]. In a large study from Spain, among 3080 positive RT-PCR patients 153 (4.9%) had CHF. Of these 98 (64.5%) had some degree of LV systolic dysfunction prior to diagnosis. The rest 54 had preserved or normalised LVEF (following treatment) or had valvular heart disease with no systolic dysfunction. The former had more CV risk factors and co-morbidities and were of advanced age. They showed higher troponins and pro-BNP, higher incidence of thrombotic events and higher mortality. The authors observed that withdrawal of GDMT was associated with precipitation of acute decompensation and higher mortality in those who had prior HF [24]. In a study of 6,439 patients those who had prior HF, had longer hospital stay, increased risk of mechanical ventilation and mortality compared those who never reported HF prior to their admission regardless of EF at admission [25].

Chatrath et al in a retrospective analysis of 134 hospitalised patients of chronic heart failure of which 40 tested positive for COVID-19 infection. Top co-morbidities were comparable between positive and negative groups. Among the patients who tested positive the mortality was almost 5 times greater at 50.0% vs 10.6% (RR 4.70). COVID-19 pneumonia and decompensated heart failure was the cause of death in 40% of them. Evidence for myocardial injury and need for Oxygen and treatment for super-added infection was higher in the positive group. The authors attributed the excess morbidity and mortality to excess of inflammatory and immunological pathways, that lead to cardiac dysfunction and renal hypoperfusion [26]. COVID-19 infection manifests in severe form in the elderly HF patients with increased mortality due to reduced immunity and general frailty and age-related disabilities.

(c) Heart failure with preserved EF
This type of presentation in patients with COVID-19 disease is possible by direct viral infiltration, inflammation, or cardiac fibrosis or by unmasking of pre-existing HFpEF. Direct myocardial injury is evidenced by elevated biomarkers, cardiac
imaging, and autopsy reports. The release of pro-inflammatory cytokines like IL-1 and IL-6 favour the role of inflammation. At least 2 echocardiographic studies have shown that in nearly 90% of hospitalised patients the LVEF was normal but there could be RV dysfunction and LV diastolic dysfunction. This may be due to hypoxemic respiratory failure and thrombotic events [27]. Higher morbidity in the diabetic and elderly patients suggests possible metabolic injuries in the development of HFpEF [28].

Management of heart failure

Pneumonia and pulmonary oedema can be confused with each other and also may co-exist in the same patient. Early admission of all suspected cases to ICU has to be decided by a designated COVID physician without wastage of precious time as patients can deteriorate very rapidly. Evaluation includes extensive and repeated imaging studies and tests for the presence of the virus. Besides basic cardiac, pulmonary, renal, and hepatic evaluation tests like D-dimer, platelet count, serum ferritin, CRP, troponins, BNP etc are warranted. Evaluation of the thrombotic potential and the pro-inflammatory markers is of paramount importance in heart failure patients affected with COVID-19. Lymphopenia, acute kidney injury and elevated LDH, liver enzymes, CRP and serum ferritin, raised PT, troponins; creatinine kinase and D-dimer are predictors of cardiac affection and bad prognosis [1].

Like pre-COVID days the primary focus in acute decompensated heart failure is the enhancement of hemodynamic status through reduction of vascular congestion, improving the preload, after-load, and the ventricular contractility. Correction of hypoxemia, anaemia, infection, electrolytes disturbances and the fluid management are principal concerns. Judicious diuretic, inotropic and vasodilator use, CPAP/BIPAP timel y intubation enhances recovery from the decompensated state, irrespective of the COVID-19 status.

Pharmacotherapy of HF

Diuretics, RAS inhibitors, beta-blockers and MRAs continue to be the main drugs in the conventional treatment of HF even in COVID-19 patients.

1) Diuretics: Critical care physicians and nephrologists share the responsibility for proper fluid and electrolyte balance in each patient. Their inputs have to be respected while giving diuretics for HF management. Hypotension has to be investigated for any precipitating factor like low fluid intake, fever, use of NSAIDS, GI bleed or septic shock or any drug-effect. Invasive monitoring has to be decided on case to case basis by the team.

2) RAS inhibitors: Those who are on ACEI/ARBs/ARNI have to continue the previous medications unless hypotension or renal derangements warrant dose reduction or temporary stoppage. Based on a single study, initially it was debated whether ACEI/ARBs make the host cells susceptible to COVID-19 infection through an up-regulation of ACE2 [29]. Subsequent observations and analysis confirmed that there is no increase in adverse effects in patients on continued ACEI/ARBs and in fact such RAS inhibition was protective to lung inflammation at a later stage of COVID-19 infection [30]. Almost every professional body advised the continuation of ACEI/ARBs as before for hypertension and heart failure. Hypotension or worsening renal function should be the main indication for stoppage or reduction of dose of these cornerstone molecules [31-33].

3) Beta-blockers: Metoprolol, bisoprolol, carvedilol, or nebulol are the approved beta-blockers for heart failure; but they should be used with caution in presence of hemodynamic instability. Drugs like lopinavir/ritonavir or darunavir are also known to decrease heart rates and may cause hypotension and they have to be used judiciously. It is suggested that carvedilol could have unique anti-cytokine properties; but it not been recommended to prefer it over the other three [34].

Management of other systems and co-morbidities

Management of critically ill patient of HF in the setting of COVID-19 infection in the high definition units is truly a multidisciplinary teamwork calling the services of physicians of other specialities, the critical care physician being the co-ordinator. Pulmonary disease is the predominant affection in many COVID-19 patients. The pulmonologist, ICU
specialist and anaesthesiologist play crucial role in decision making. Hypoxic respiratory failure and hypoxic cardiac injury can coexist. HF with pulmonary oedema can predispose or worsen the pulmonary complications and lead to enhancement of intensity of mechanical respiratory support. The aim is to achieve SpO2 >95% (In COPD >90%). Initially standard oxygen therapy, venturi mask or mask with reservoir bag is used. When high flow oxygen does not help, non-invasive ventilation and prone positioning may be considered. Next escalation step is endo-tracheal intubation and invasive mechanical ventilation with lung protective strategies like low tidal volume (6-8 ml/kg ideal body weight) and low-level airway platform pressures < 30 cm H2O [10, 35]. Acute kidney injury can be seen in about one-third of the ICU admitted patients. Most of them may need renal replacement therapy of which ultra-filtration may be best for diuretic resistant HF patients [36]. Hemofiltration can effectively control inflammatory response, reduce the offending factors in the blood and minimise the tissue injury in this critical stage of cytokine storm in COVID-19 affection [37].

Anti-viral drugs, immune-modulatory agents and steroids have to be used based on the current evidence and local institutional protocols as there are some controversies and newer updates are constantly pouring in. It is important to remember that some antiviral agents can cause myocardial dysfunction, QTc prolongation and cardiotoxicity as well.

**Use of MCS and ECMO in cardiogenic shock**

After oxygenation is taken care, intra-aortic balloon pump can be the initial choice for patients in borderline shock states. If they do not provide adequate support, micro-axial systems like Impella may be tried which adequately unload the failing ventricle and reduce cardiac inflammation. Next in the upgradation is veno-venous extracorporeal membrane oxygenation (ECMO) which is both expensive and not available in most centres. In presence of decreased CO and depressed LV contractility one may prefer the Veno-arterial ECMO and additional respiratory failue warrants a hybrid ECMO. Zheng et al cautioned about the potential for reduction of some classes of lymphocytes and inflammatory activation in patients on ECMO in COVID-19 patients [38-41].

**Advanced therapies**

**Heart transplant program**

ACC published the poll results of impact of the COVID-19 pandemic on advanced heart failure and heart transplantation, which was conducted in August, 2020 involving 47 adult program-physicians from 12 countries. Nearly 80% was altered their approach to the program priorities. One in five stopped the program at some time or became more selective. There was overall decrease in heart transplant volumes. COVID-19 related mortality in heart transplant recipients was 25% [42, 43].

Managing the heart transplant recipients can be quite challenging. In heart transplant recipients, immunosuppressive drugs can cause lymphopenia which may be further enhanced by COVID-19 infection. ISHLT guidelines suggested their temporary with-holding in in sever presentations of COVID-19 infection. The anti-viral agents can interfere with action of cyclosporine. It is also possible to misdiagnose COVID-myocarditis to be cardiac transplant rejection. Gene-profiling or donor-derived cell free DNA test can differentiate the two entities [44].

**Left ventricular assist devices**

Patients on left ventricular assist devices (LVADs) are prone for various infections and they have to be guarded against COVID-19 as well. The device may need tele-remote programming. In those on long term LVADs can have elevated cytokines even without COVID-19 affection. This has a bearing in evaluation of a COVID-19 positive patient. Increased potential for thrombosis and appropriate anticoagulation may pose additional challenges in these patients. These patients if contract COVID-19 disease can be at higher risk for right heart failure due to hypoxia and probable high output syndrome [44].

**Development of updated guidelines**

The recent publication of position paper by Chinese Heart Failure association and the European Society of Cardiology is praise-worthy exercise to put most of the existing data in the right prospective and formulate some suggestions for the management of current issues in management of HF in the COVID-19 pandemic. Emergence of guidelines has to wait until more experience is shared through standard publications involving larger number of patients [44].
For chronic management of discharged patients, tele-consultations and physician-guided patient self-management programs have been implemented at some centres to reduce hospital visits and admissions. Implantable sensors for remote PA pressure monitoring can be of particular benefit in these unprecedented times.

Conclusions
The 12-month old COVID-19 pandemic posed unprecedented challenges to the practice of public health and clinical medicine; HF management is no exception. Known HF patients becoming worse after acquiring the COVID-19 infection or the new development of acute cardiac complications in persons without any prior heart disease- both are novel therapeutic issues which have no perfect answers for want of adequate data and randomised trials. Many current practices and expert recommendations in the management of HF with concomitant COVID-19 are bound to be modified in the days ahead with further data and experience. We have to update ourselves every day.

Conflicts of interest
Authors have no conflict of interest.

References


