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Brief Report

Case Reports of Implantable Cardiac Device Physiologic Sensor Changes in Subjects with Coronavirus Disease-2019 Infection

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ABSTRACT

The severe acute respiratory syndrome novel coronavirus-2 pandemic has established a new set of challenges to health care delivery. Remotely monitored physiologic sensors on implantable cardiac devices can provide insight into the differential diagnosis of dyspnea in the heart failure population. We report on a unique pattern of sensor deviations that seem to occur specifically with severe acute respiratory syndrome novel coronavirus-2 infection. (J Cardiac Fail 2021;27:373–378)

The severe acute respiratory syndrome novel coronavirus-2 (SARS-CoV-2) pandemic has created evolving models of health care delivery with an increased reliance on telehealth.1 Individuals with underlying cardiovascular disease are at increased risk of complications from SARS-CoV-2 infection.2 As such, clinicians have been moving to virtual visits in these high-risk populations to avoid unnecessary exposure within the health care system. Infection with SARS-CoV-2 has a broad range of symptoms, with cough and shortness of breath among the top presenting symptoms.3 Given reports that only 48% of patients exhibit fever at the time of admission, the differentiation between congestive heart failure and SARS-CoV-2 infection becomes more difficult on a virtual platform.3

Many patients with heart failure due to reduced ejection fraction have implantable cardiac defibrillators and/or cardiac resynchronization therapy devices with physiologic sensors that are remotely monitored. HeartLogic is a multiparameter heart failure index and alert algorithm that has been validated in predicting impending heart failure decompensations.4 The algorithm utilizes sensors that collect information on heart sounds (S1 and S3), respiration, thoracic impedance, heart rate, and activity level to create a daily index value. An alert to the clinician is automatically triggered when the index value crosses a configurable alert threshold. Sensor data are downloaded and transmitted remotely, with clinicians able to access individual sensor data via an online remote patient monitoring system.

The identification of a unique signature of sensor patterns that differentiates heart failure from possible infection with SARS-CoV-2 may allow for a safer and more effective triage, testing, and treatment in these high-risk patients. We describe here the pattern of HeartLogic physiologic sensor changes in 3 patients with SARS-CoV-2 infection. Analysis of these sensors demonstrates a pattern with SARS-CoV-2 infection that is distinct from heart failure. Changes in sensor values were compared to a 60-day baseline average, 90 days before symptom onset.

Case Studies

Case 1

A 62-year-old Caucasian man with ischemic cardiomyopathy, a left ventricular ejection fraction of 30%, New York Heart Association functional class I, with comorbidities of hypertension and hyperlipidemia, who was implanted with a Boston Scientific RESONATE device for primary prevention. On March 13, he developed fever, chills, myalgia, anorexia, and severe fatigue. He was tested for SARS-CoV-2 on March 16, which ultimately returned as positive.

As seen in Fig. 1, his symptom onset coincided with a substantial increase in the respiratory rate (from 20 to 34 bpm), rapid shallow breathing index (RBSI; from 12.5 to 35.5 br/min/ V), nighttime heart rate (from 75 to 106 bpm), and temperature (from 93.6˚F to 98.2˚F), with a sudden...
decrease in his activity level (from 2.2 to 0.3 hours). Interestingly, a significant increase in thoracic impedance (from 46.7 to 54.0 Ω), S1 (from 4.8 to 7.0 mG) and S3 heart sounds (from 0.85 to 1.42 mG) was also noted with infection. This increase in thoracic impedance and S1 heart sound is atypical for decompensated heart failure, because thoracic impedance and S1 typically decrease as vascular congestion and fluid overload increase.5–7

Fig. 1. Case 1. HeartLogic tracings depicting a sharp increase in respiratory rate, RSBI, nighttime heart rate, S1, S3, temperature and thoracic impedance with onset of severe acute respiratory syndrome novel coronavirus-2 symptoms (purple vertical line). Major symptom resolution (green vertical line) was associated with recovering sensor trends. Maximum and minimum values are indicated with red dots. A baseline was calculated for each sensor trend (tan shaded interval). RSBI, rapid shallow breathing index.
In our cases, infection with SARS-CoV-2 created a unique pattern of sensor changes that was distinct from uncomplicated heart failure. Thoracic impedance, known to increase in volume overload, \(^5,^6\) increased in all 3 cases. Interestingly, this increase in thoracic impedance also differs from the pattern seen with pneumonia, which has been noted to contribute to false-positive results with intrathoracic impedance monitoring devices for heart failure. \(^8\) The precise reason as to why the thoracic impedance increases with SARS-CoV-2 infection is not well-understood. However, this finding is consistent with findings from Heggermont et al, \(^10\) who described a similar phenomenon of increasing thoracic impedance in a hospitalized patient with SARS-CoV-2 infection. Furthermore, in all of our cases, S1 heart sound increased with infection, in contrast with the decrease expected in decompensated heart failure. \(^7\) An increasing heart rate and respiratory trends in conjunction with increasing thoracic impedance and S1 would argue against heart failure and points to an alternate diagnosis.

**Discussion**

In the era of telemedicine, data derived from physiologic sensors can be extremely useful to clinicians in triaging and treating patients with heart failure complaining of dyspnea. Sensor data are monitored remotely and can provide insight into differential diagnoses without necessitating in-person contact. This technology holds the promise to decrease the spread of infection in the outpatient clinic setting by allowing clinicians to evaluate changes in physiologic sensors before patient presentation.

In our cases, infection with SARS-CoV-2 created a unique pattern of sensor changes that was distinct from uncomplicated heart failure. Thoracic impedance, known to decrease in volume overload, \(^5,^6\) increased in all 3 cases. Interestingly, this increase in thoracic impedance also differs from the pattern seen with pneumonia, which has been noted to contribute to false-positive results with intrathoracic impedance monitoring devices for heart failure. \(^8\) The precise reason as to why the thoracic impedance increases with SARS-CoV-2 infection is not well-understood. However, this finding is consistent with findings from Heggermont et al, \(^10\) who described a similar phenomenon of increasing thoracic impedance in a hospitalized patient with SARS-CoV-2 infection. Furthermore, in all of our cases, S1 heart sound increased with infection, in contrast with the decrease expected in decompensated heart failure. \(^7\) An increasing heart rate and respiratory trends in conjunction with increasing thoracic impedance and S1 would argue against heart failure and points to an alternate diagnosis.
Our data suggest that a unique set of sensor trends is identifiable in the patient who is infected with SARS-CoV-2. Using this pattern of individual sensor changes, there is the potential that alerts configurable for specific respiratory conditions could be developed. However, this report is a retrospective observational review of 3
cases. Further studies are needed to determine if the trends seen with these 3 cases remain consistent in larger sample sizes.

With the recent literature focusing on the use of telemedicine in the patient with advanced heart failure, remotely monitored physiologic sensor data have become increasingly used.
in patient evaluation. Given the somewhat broad differential in the patient with heart failure presenting with dyspnea, these distinctive trends have the potential to be extremely useful to clinicians. The identification of patterns that are atypical for heart failure before patient presentation may expedite triage and decrease potential exposure to SARS-CoV-2.

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