Coronary Microvascular Injury in the COVID-19 Patient: The Need for Imaging Studies in Post-COVID-19 Cardiovascular Syndrome

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The COVID-19 pandemic has generated numerous questions about the pathophysiology of the acute phase of this disease, and as time goes by, also of the chronic phase, known as post-COVID-19 syndrome [1-4]. Although the neurological phenotype (post-COVID 19 neurological syndrome) is the most debated [1,2], the evidence points to the fact that the main manifestations of post-COVID 19 patients, regardless of disease severity, are dyspnea, fatigue, tachycardia, and difficulties in performing activities of daily living [5]. This raises alarms about the presence of persistent target organ injury since the acute phase of COVID-19, or whether previous inflammation may have decoupled the mechanisms that some organs perform in the face of chronic disorders. There is evidence that has reported the compromise of the endothelium and myocardium during the acute phase of the disease, with the presence of previous heart disease being considered a risk factor for severe disease and death [6].

Interestingly, Drakos, et al. [7] evaluated coronary microvascular disease in COVID-19 patients by cardiovascular magnetic resonance imaging, showing that patients who had COVID-19 had significantly reduced global myocardial perfusion reserve (2.73 [2.10-4.15 - 11] vs. 4.82 [3.70-6.68], p = 0.005), significantly increased coronary sinus flow at rest (1.78 ml/min [1.19-2.23 ml/min] vs. 1.14 ml/min [0.91-1.32 ml/min], p = 0.048), and reduced coronary sinus flow during stress activity (3.33 ml/min [2.76-4.20 ml/min] vs. 5.32 ml/min [3.66-5.52 ml/min], p = 0.05), compared to controls [7]. Based on the above, the authors concluded that there is cardiac microvascular injury in COVID-19 patients, which may trigger major cardiovascular events in the post-COVID-19 phase, and this is one of the reasons that would explain the persistence of fatigue and dyspnea during this phase [7].

Another study that supports the persistence of endothelial dysfunction, but using another vascular parameter (flow-mediated dilation), is that of Ambrosino, et al. [8], who carried out a case-control study with 133 patients where they found that a significantly lower flow-mediated dilation in post-COVID 19 patients as compared to controls (3.2% ± 2.6...
vs. 6.4% ± 4.1 p < 0.001), confirmed when stratifying the study population according to age and major clinical variables. Among cases, females exhibited significantly higher flow-mediated dilation values as compared to males (6.1% ± 2.9 vs. 2.5% ± 1.9, p < 0.001). Among post-COVID 19 patients, flow-mediated dilation showed a direct correlation with arterial oxygen tension (0.247, p = 0.004), forced expiratory volume in 1s (0.436, p < 0.001), forced vital capacity (0.406, p < 0.001), and diffusing capacity for carbon monoxide (0.280, p = 0.008). In conclusion, post-COVID-19 syndrome was a major and independent predictor of flow-mediated dilation values (β = -0.427, p < 0.001) [8]. This allows us to understand that although the study investigated pulmonary vascular dynamics and some respiratory parameters, the endothelial damage could be the same and can be correlated with the mechanism of lung-heart circulation [7-9].

It is valid then, even to raise the question of whether there is silent myocarditis in the patient with asymptomatic COVID-19, which through compensatory mechanisms does not develop symptoms, but leaves a trace in the vascular and cardiac territory? Clearly, in the patient with cardiovascular risk factors, we should inquire more quickly through translational research in the vascular and cardiac territory? Clearly, in the patient with heart failure) or death, or re-evaluate therapeutic plans based on the findings and the mechanisms of action of the most appropriate drugs for each case. Prospective multicenter studies are needed to define cutoff scores for hemodynamic and structural parameters to control the burden of cardiovascular disease that is expected in this group of patients.

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Conflicts of Interest

None.

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