Deceptive Slanders of Cardiovascular Pathology in Covid-19 Ethnic Minorities

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Abstract
Reviewing the Covid-19 literature over the past six months we can identify pervasive endothelial dysfunction disorders, whether in the form of comorbid conditions e.g. arterial hypertension, mediated by pathogens, e.g., thromboembolic syndrome and/or iatrogenic due to inadequate therapies, e.g., ventilators, vasopressors. The conclusion is unless you are young and slim and Caucasian, we cannot cure you! Indeed, it is unacceptable in the twenty-first century to involve racial or ethnic assumptions in science without providing substantial evidence, especially in renowned journals. The question is, do we realize the extent of the psychological damage we are causing to individuals belonging to these ethnic minorities, to their wives, children, and friends? We aim through the present work to correct this erroneous thought, as well as to expose our visions concerning the management of Covid-19, which unfortunately became a politico-mediatic subject and remains without an effective solution.

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Introduction
In a post-mortem study of 10 African American patients with COVID-19, Fox and colleagues claimed key pathological states in this demographic, e.g. right ventricular (RV) dilatation [1]. In addition to the lack of significant statistics of substantial clinical data, e.g., echocardiography, ECG, ...etc., they have mentioned normal cardiac dimensions (3.4 cm and 3.6 cm) and presented a normal heart figure (Figure. C3), showing a well trabeculated right ventricular free wall, which is the most vulnerable part that cannot withstand intracardiac pressure. We should emphasize that the right-heart side is a highly compliant circuit, in which the claimed acute ventricular dilatation, which is a rapidly fatal hemodynamic condition, most commonly occurs due to right ventricular outflow tract (RVOT) surgery, e.g., excessive resection of the moderator band in the tetralogy of Fallot.

Thereafter, in less than a month, the authors published a new series, in which they had faded their exclusive identification of cardiac risk factors in the African American population and maintained claiming RV dilatation in Covid-19 [2]. It is important to precise that all these patients presented endothelial dysfunction comorbidities, linked to stressful socioeconomic status [3]. All studies on ethnic minorities lack comparative data with Caucasian groups, subjected to similar environmental conditions. Genetics rather than ethnicity, is the most rational Vector in Science, for example, difficulties of ventilators weaning in Noonan or Down syndrome. We stitch wounds for better healing, which is an endothelial function process. We know how to avoid sunrays in dark skin or absorbable stitches in Down syndrome to avoid undesired results. But we don’t add corrosives and complain about keloids (fibrosis). Only endothelial function tests in multi-ethnic infants can refute or confirm these claims. The ugly fact is that we are still blowing up the most important circulatory driving force that controls hemodynamics and microcirculation: the heaven’s door of cellular biology and tissue oxygenation of vital organs, heart included. Surgically, we can bypass the right or left ventricle but with the permission of the almighty respiratory pump [4].

As a reminder, the endothelium is the precursor of the cardiovascular system, depends on the shear stress of the bloodstream to maintain its functions like vascular tone, coagulation, angiogenesis, apoptosis, diabetes, atherosclerosis, immune system, inflammatory response, nitric monoxide synthesis, etc.

Endothelial shear stress (ESS) controls vasculogenesis, cardiogenesis, embryogenesis, organogenesis through the angiogenesis-apoptosis interdependency process, from the 8th day of gestation until death. An imbalanced angiogenesis-apoptosis interdependency can induce irreversible cellular damages like Eisenmenger syndrome, Cor pulmonale, or heart failure: cardiomyocytes apoptosis, compensated by angiogenic hypertrophy or fibrotic dilatation [5]. In antenatal, the right ventricle (RV) is the main trigger of ESS and moderator of fetal development even in severe cardiomyopathies, e.g., hypoplastic left heart syndrome. It distributes blood flow to the left ventricle (LV) through the foramen ovale, to pulmonary and collapsed lungs circuit (=10%), and the descending aorta through the ductus arteriosus. In the postnatal period, with shunts closures, the respiratory pump becomes a key circulatory driving force to deal with the massive blood volume at the right-heart side (≥80%). It becomes the main trigger of ESS to continue cardiovascular remodelling e.g., increasing LV mass in maintaining low remodelling at the right-heart side (RV/LV ratio)
mass \approx 1/6). It squeezes the pulmonary parenchyma in an accordion-like manner, releasing plenty of endothelial mediators to drop the pulmonary vascular resistance (PVR), to improve hemodynamics as well as tissue oxygenation with first breath after birth. By controlling the pulmonary afterload, the respiratory pump controls RV preload and cardiac output (Frank-Starling law), helped with other influential forces like the muscle pumps, gravity, atmospheric pressure [6].

Disturbed RV preload provokes cardiovascular disorders like in astronauts, professional scuba divers, sleep apnea, and serious complications in bedridden ventilated patients. The respiratory pump which is a low-pressure momentarily closed hydraulic circuit, due to the epiglottis effect, must deal with two types of fluids: the compressible Newtonian (air) and the incompressible non-Newtonian (blood); and a delicate alveolar system composed of two types of single-cell layers: the epithelium and the endothelium, to ensure gas exchanges. Also, the extra-alveolar and alveolar endothelial cells have different embryological origins and behavior, e.g., drawbacks of inhalational PAH therapies [7]. Furthermore, we have proved that the pulmonary endothelium, which is unexploited before birth, responds differently and more effectively to ESS stimuli, compared to the left-heart side endothelium [8]. Then we come with the ventilator’s invasion and neuromuscular blockades to transform the respiratory pump into a piston-like, closed, pressurized, caged, purulent, hydraulic circuit.

Maintenance of endogenous pulmonary endothelial mediators is mandatory to respiratory or more correctly metabolically, distressed patients, whatever the underlying pathology. Once the pulmonary production of ESS is compromised due to pathological conditions of the contractile structures of the respiratory pump e.g., pneumonia, edema, patients exhibit symptoms and signs like dyspnea, tachypnea, tachyarrhythmia, orthopnea, even with very mild hypoxia (SpO2≤94%), which is normally uncompromising for life. This means desperate requirements of endogenous pulmonary mediators for hemodynamics and metabolic improvements.

It is all about how to fully engage the respiratory pump and its influential forces, in particular the gravitational effect. For example, pulmonary ESS enhancement is the hallmark of physical exercise, exhibited without shortness of breath in marathon runners due to the "second wind" effect. In contrast, an overweight nonathlete runner exhibits shortness of breath and leans forward with hands-on knees and not in a recumbent position. On the other hand, a congestive heart failure patient, in a recumbent position exhibits nocturnal orthopneic dyspnea to improve hemodynamics. Or patients in severe cyanosis (SpO2≤80%), exhibit squating position without shortness of breath. This is exactly what lies behind the untold explanations of studies showing the advantages of noninvasive ventilation, low-dose neuroblockade: maintaining chest wall recoil; prone position, obesity: correlated with diaphragmatic compression by stagnant hepatosplanchnic venous capacity; tracheostomy: decreasing intervalveolar pressure by reducing airflow energy losses and tracheal dead space, etc.

Besides, vasopressors and the encounter of circulatory assist devices (CAD) with the circulatory system creates a vicious circle of endothelial dysfunction and momentum energy losses. As it’s known, boundary wall friction of bloodstream inside rigid narrow conduits of CAD provokes the postcardiotomy syndrome [9], post-hemodialysis pains [10]. Similarly, pressurized airflow inside ventilators rigid conduits but with different diameters promotes barotrauma and surinfection by the Venturi effect. We think it is time to have a deep breath and calm down, through the respiratory pump, and let it do the knocking on heaven’s door instead of our unfortunate patients. We didn’t need a well-performing heart pump (Figure 1) [11,12], or even heartbeats, ventilators, and pharmacological supports [13], to improve hemodynamics in fragile animal models, but endogenous pulmonary

![Figure 1: A: Presumed mechanism and passage of induced pulmonary eNOS. 1 = Pulmonary artery (PA); 2 = pulsatile catheter fitting PA trunk; 3 = right ventricle (RV) inlet-outlet compartments; 4 = infundibular site of pulmonary catheter insertion; 5 = arrows showing presumed passage of pulmonary eNOS (backward through coronary ostia and/or forward through systemic circulation); 6 = left ventricle (LV) inlet-outlet compartments; 7 = permanent ligation of the left anterior descending coronary artery distal to the second diagonal branch; 8 = interventricular septum; 9 = cardiorespiratory monitor; 10 = pneumatic driving force. 1 = pulmonary eNOS primarily induced at PA zone with catheter pulsation; II = pulmonary eNOS natural passage through the left heart circuit; III = presumed pulmonary eNOS involvement in myocardial recovery most probably through micrcirculation and/or the RV interseptal coronary network. B: Macroscopic reduction of the ischemic zone. Left panel figure showing dark infarcted myocardial after 50 min of ischemia; Right panel figure showing significant reduction of ischemic myocardial zone after 10 min of pulsation; 1 = left anterior descending coronary artery snuggler; 2 = infundibular site of the intrapulmonary pulsatile catheter insertion. C: Myocardial apoptosis by TUNEL technique. Upper panel, apoptotic index (AI) in both groups. AI in group P was significantly lower than that in group NP (P < 0.01); lower panel, representative figures from both groups showing apoptotic cells manifestations (red arrows): lower left, group P; lower right, group NP. D: Myocardial eNOS mRNA expression. RT-PCR results shown with statistics, in which myocardial eNOS expression was significantly higher in group P compared to group NP (P<0.01). Pulsatile group (P), Non-pulsatile group (NP).](Image)
endothelial mediators. This is not polemical or science fiction work, it is what we have developed over the past two decades with pioneering doctors and legendary heart surgeons, including the former president of a well-known Academy of Sciences.

References


