COVID-19: A PERSONALIZED CARDIOMETABOLIC APPROACH FOR REDUCING COMPLICATIONS AND COSTS.
THE ROLE OF AGING BEYOND TOPICS

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Abstract: COVID 19 is much more than an infectious disease by SARS-CoV-2 followed by a disproportionate immune response. An older age, diabetes and history of cardiovascular disease, especially hypertension, but also chronic heart failure and coronary artery disease among others, are between the most important risk factors. In addition, during the hospitalization both hyperglycaemia and heart failure are frequent. Less frequent are acute coronary syndrome, arrhythmias and stroke. Accordingly, not all prolonged stays or even deaths are due directly to SARS-CoV-2. To our knowledge, this is the first review, focusing both on cardiovascular and metabolic aspects of this dreadful disease, in an integrated and personalized way, following the guidelines of the Cardiometabolic Health/Medicine. Therefore, current personalized aspects such as ACEIs and ARBs, the place of statins and the most appropriate management of heart failure in diabetics are analysed. Aging, better than old age, as a dynamic process, is also considered in this review for the first time in the literature, and not only as a risk factor attributed to cardiovascular and non-cardiovascular comorbidities. Immunosenescence is also approached to build healthier elders, so they can resist present and future infectious diseases, and not only in epidemics or pandemics. In addition, to do this we must start knowing the molecular mechanisms that underlying Aging process in general, and immunosenescence in particular. Surprisingly, the endoplasmic reticulum stress and autophagy are implicated in both process. Finally, with a training in all the aspects covered in this review, not only the hospital stay, complications and costs of this frightening disease in high-risk population should be reduced. Likely, this paper will open a gate to the future for open-minded physicians.

Key words: Metabolic Syndrome, cardiometabolic syndrome, cardiometabolic risk, cardiometabolic health, atherothrombotic disease, endothelial dysfunction.

Introduction: historical perspective, original focus and pandemic expansion. Why a Cardiometabolic approach is necessary?

Coronavirus (CoV) is a subfamily of virus with positive single-stranded RNA belonging to the coronaviridae family of the order of Nidovirales; they are responsible for diseases in animals (AcCoV) but also in humans (HCoV). Other Nidovirales, like arteriviridae (1), are responsible for diseases only in animals, and the most important is the equine viral arteritis (2).

Until now, CoV had caused illnesses in human beings ranging from the common cold, the most frequent, to more severe diseases such as SARS (Severe Acute Respiratory Syndrome) and MERS (Middle East Respiratory Syndrome). In 2003, SARS-CoV involved 32 countries, with 8422 confirmed cases and 916 (10.87%) casualties from November 2002 to August 2003 (3). In 2012, the MERS-CoV MERS-CoV spread over 27 states, causing 2496 cases and 868 casualties (34%). The most affected patients produced an in Saudi Arabia with 124 cases and a mortality rate as high as 40%. Isolated cases were subsequently diagnosed in other areas, the last one in 2015 (4).

A new coronavirus is responsible for the current pandemic with the initial focus in Wuhan (Hubei, China), in December 2019 (5). The latter expansion to South Korea, Spain and Italy took place at the end of January 2020. Therefore, on 30th January 2020, the World Health Organisation (WHO) declared the COVID-19 (Corona Virus Disease 2019) as an epidemic (6). On 11th February 2020, the WHO named this new disease as COVID-19 and the new virus as SARS-CoV2 (7). Finally, on 12th March 2020, the WHO announced the COVID-19 outbreak as a pandemic.

When we finished writing this paper, on 16th April 2020, 21:08 GMT, there were 210 countries from the 5 continents, 2,173,168 confirmed cases and 144,949 deaths. The average global Case Fatality Rate (CFR) is around 5%, depending on the age, and is slightly higher in males than females (6).

In financial terms, this pandemic is triggering a major worldwide economic meltdown with the stock market flash crash. On 9th April 2020 the International Monetary Fund, a United Nations (UN) specialized agency, predicted that COVID-19 will trigger the worst economic fallout since the Great Depression of the early 1930s shortly after stock market crash of Black Tuesday on 29th October 1929.

Facing such health and economic events, why is a cardiometabolic approach necessary here and now? Recently, five reviews (8-12) about cardiovascular aspects of this terrible disease, at the moment ignored, are opened in order to focus on a personalized cardiometabolic approach necessary here and now.
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Table 1
Summary of five cardiovascular-related studies in COVID-19 patients

| Author          | Journal / year | Diseases studied                                   | Conclusions                                                                 | Recommendations                                      |
|-----------------|----------------|----------------------------------------------------|----------------------------------------------------------------------------|-----------------------------------------------------|
| Driggin E et al | JACC 2020      | Patient risk assessment: infection and cardiovascular issues | The COVID-19 has substantially increased the difficulty of treating patients with severe emergent CVD | Individualized diagnosis and treatment measures tailored to specific local epidemic situations should be developed |
| Guo T et al     | JAMA Cardiol 2020 | Association of underlying CVD and myocardial injury with fatal outcomes in COVID-19 | Myocardial injury is associated with cardiac dysfunction, arrhythmias and fatal outcome of COVID-19. The prognosis of patients with underlying CVD but without myocardial injury is relatively favourable. | Aggressive treatment may be considered for patients at high risk of myocardial injury |
| Zheng YY et al  | Nat Rev Cardiol 2020 | Underlying CVD                                    | COVID-19 patients with underlying CVD have an adverse prognosis | Particular attention should be given to CV protection during treatment for COVID-19 |
| Madjid M et al  | JAMA Cardiol 2020 | Underlying CVD and/or cardiac risk factors         | COVID-19 is associated with a high inflammatory burden that can induce vascular inflammation, myocarditis, and cardiac arrhythmias. | CV risk factors and conditions should be controlled per evidence-based guidelines |
| Han Y et al     | Circulation AHA 2020 | Patient risk assessment of both infection and CV issues | Patient risk assessment of both infection and CV issues. Consider the prevention and control of COVID-19 transmission as the highest priority | Individualized diagnosis and treatment measures tailored to specific local epidemic situations should be developed |

CV = cardiovascular, CVD = cardiovascular diseases

disease had been published, but none are about an integrated cardiometabolic approach (See Table 1). Our questions are, why?, and beyond, would it be useful?.

The term “Cardiometabolic Medicine” (CMM), synonym of “Cardiometabolic Health” (CMH), was launched officially in October 2006 in Boston, with the celebration of the first congress, a meeting point for physicians from different specialties, biologists, biochemists, pharmacists, cardiovascular bioengineers and researchers in all these fields. The term “cardiometabolic” had previously used by Pescatello in 1999 for obesity (13), by Sowers in 2001 for “CM Syndrome” (14), and by Vasudevan & Ballantyne in 2005 (15) for “CM Risk» in two phases. Firstly, the term was used by Khan, Buse, Ferramini and Stern in a symposium whose proceedings were never published. Fortunately, their conclusions were collected and valued as of great interest by other authors (Vasudevan & Ballantyne); they published them that same year (15) laying the groundwork for a new enlightened medicine. This medicine, the CMM, is much more eagerly, dynamic, open that the classical cardiovascular medicine, and at the same time much more predictive, preventive and anticipatory. To topping it all off, the CMM is the prototype of the modern “Precision Medicine”, on which the American government spent so many millions in research during the last decade (16).

This CMM approach is not new, we saw it was born starting this 21st century, but we frequently forget, at the expense of the most classical cardiovascular term, actually dating from the 20th century. Despite the latter one, is still useful in certain specialties (Cardiology, Neurology and Vascular Surgery), it is seldom useful for defining prevention strategies. Therefore, CMM had allowed detection of causes and had predicted the consequences of a rise of cardiometabolic diseases across the world (especially in low- and middle-income countries) (17), and had proven to prevent and predict events (18), and at the same time, facilitating the cost analysis.

Bearing this in mind, it is quite likely, that if we have fewer complications, particularly severe ones (cardiac arrest, ventricular tachyarrhythmia, acute heart failure, acute coronary syndrome, haemorrhagic or massive ischaemic stroke), this integrated approach could cut down the elevated mortality in the highest risk group (cancer, COPD and oldest subjects with comorbidities), usually preceded by a multi-organ failure.

The SARS-CoV-2 as new human Coronavirus

Seven types of HCoV had been described. On one side, we have four genuinely human ones (229E, NL63, OC43 and KU1) which are responsible for almost 30% of common colds. On the other hand, three mutated variants of ACoV had been described (see below) causing zoonotic disease when infecting human beings (19). This happened also with the Nipah virus infection in 1998 (Malaysia) and 2004 (Bangladesh) from the so-called “fruit bat”. After this outbreak, as they did after latest MERS epidemic, the CEPI (Coalition for Epidemic Preparedness Innovations), worked successfully on vaccines to prevent future epidemics with both viruses.

In relation to mutated CoV they are included strangely enough as HCoV (20). In either case, unlike the first group of
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HCoV, they usually show a wide clinical range, with higher lethality in risk populations. Of course, we are speaking about SARS-CoV, MERS-CoV and SARS-CoV-2, responsible of SARS, MERS and COVID-19 respectively, three emerging CoV diseases in just two decades (21). MERS and SARS were probably originated from bats and then moving into other mammalian hosts (the Himalayan palm civet for SARS-CoV, and the dromedary camel for MERS-CoV) before jumping to humans. The exact role of bat and others mammalian hosts in this current pandemic is still under investigation.

Regarding ACoV, the main target is usually the digestive tract, and rarely can induce a mild bronchitis (22, 23).

Like all coronaviruses, SARS-CoV-2 is composed of RNA involved by a glycoprotein and lipid membrane. This is not a specific membrane composition of this subfamily of viruses, but of all membranes of living beings, including plants (these are surrounded by a cell wall). This new virus’ membrane contains moreover specific proteins of CoV named S, E, N and M proteins. Regarding the external is named ”S” protein, forms spikes (hence its name), and have two subunits (S1 and S2) that allows it, if recognized by the right receptor (see below), to penetrate into the cells. From what is known so far in COVID-19, the mutation especially affects it, making it especially harmful. The “E” protein is key to infect other cells. The “N” protein, a haemagglutinin-esterase protein that allows them to camouflage the genetic material. Finally, the “M” (membrane) protein plays a central role in virus assembly, interacting through both the transmembrane domain and endodomain, turning cellular membranes into workshops where virus and host factors come together to make new virus particles (24).

Even more interesting, taking into account that at least 50% of COVID-19 patients have hyperglycaemia on admission or in the first days of hospitalization (25), the transmembrane domain has a potential N-glycation site (26) that in COVID-19 would deserve further research. The reason is that other proteins (e.g., LDL, myelin-proteins, collagen, myosin) become more pathogenic when they bind non-enzymatically to glucose (27).

On the other hand, a host transmembrane serine protease, TMPRSS2, synthesized by the microvascular endothelium, promotes the entrance of SARS-CoV into the cells in two steps: firstly, TMPRSS2 acts on the S2 facilitating fusion of the virus to the cell membrane. Secondly TMPRSS2 activates the spike and cleaves the receptor that is ACE2 (Angiotensin-I Converting Enzyme type 2) (31), the homologue of ACE, a dipeptidyl-carboxypeptidase. ACE is the classical enzyme of RAS that cleaves angiotensin I to generate angiotensin II, which is the effector peptide of the system and exerts multiple biological functions. ACE is occasionally named incorrectly as ACE1, is simply ACE because history, including the scientific history, cannot be rewritten. ACE is the main enzyme of the RAS with low ACE2 levels in blood (28) but these levels increase in patients treated with ACEIs/ARBs, generally with hypertension (HTN) and/or diabetes mellitus (DM) (32).

At a local level, RAS, with all its components such as ACE, renin, angiotensinogen and AngII, was firstly reported in dog’s brain and later in other organs (28, 33). ACE in vascular cells is expressed in the endothelial cells (EC) and smooth muscle cells (SMC) independently; so, ACE in SMC, unlike the EC, is insulin-dependent (34) including the lymphatic system. On their behalf, its homologue ACE2 is expressed mainly locally (by the local RAS) with a low production by the systemic RAAS axis which include, unlike the local system, the aldosterone in the period of about 5.2 days. A single systematic review found no statistically significant difference in common symptoms between patients with severe or mild / moderate COVID-19 (30).

In an initial phase, the virus enters through the upper airway and may produce signs and symptoms like fever, malaise, myalgia and occasionally sore throat and sneezing. Afterwards, it descends into the lower airway and there it is responsible for a dry or barely productive cough. Not uncommonly, the virus colonizes the airway and patients remain asymptomatic for about two weeks; they can be contagious in this period, whilst this has not been documented in patients with common flu. Fever is the most common symptom, but is accompanied by generalized myalgia only in 44% of total affected, markedly different from the flu, where it is close to 100%. Other frequent symptoms are cough (present in about 75%), anosmia is a frequent symptom, up to 60% of confirmed cases in some series (30), headache and drowsiness (<10%), and confusion (<5%, especially in older patients). Occasionally in this phase the patient may have sore throat, ageusia and rhinorrhoea (<5-10%), and gastrointestinal reaction (<10%) (7, 25).

In a second phase, presumably preceded of a virus-receptor interaction (see below), it is possible a pulmonary involvement, but not always. If this happens, dyspnoea is more frequent in severe cases and, in some studies, is a marker of severe disease (30).

Dyspnoea usually coincides with the appearance of lung infiltrates in the chest X-ray and CT-scan.

In a third phase, by the seventh to the tenth day about 20% of patients can develop acute respiratory distress syndrome (ARDS) and 10% a multiple organ failure (7, 25).

Regarding lung involvement, the main target in COVID-19, the SARS-CoV-2 induces a direct damage at this level, via the same receptor that SARS-CoV used in 2003, the carboxypeptidase ACE2 (Angiotensin-I Converting Enzyme type 2) (31), the homologue of ACE, a dipeptidyl-carboxypeptidase. ACE is the classical enzyme of RAS that cleaves angiotensin I to generate angiotensin II, which is a key effector peptide of the system and exerts multiple biological functions. ACE is occasionally named incorrectly as ACE1, is simply ACE because history, including the scientific history, cannot be rewritten. ACE is the main enzyme of the RAS with low ACE2 levels in blood (28) but these levels increase in patients treated with ACEIs/ARBs, generally with hypertension (HTN) and/or diabetes mellitus (DM) (32).

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equation. ACE2 is one of the type-I integral proteins, expressed fundamentally at the cell surface as an ectoenzyme and is found at heart, kidney, small intestine, lung (type II pneumocytes), pancreas and liver among others. The expression of ACE2 in the endothelium and vascular SMC opens an unexplored pathway in subjects with COVID-19 who develop acute coronary syndromes. It had been demonstrated that in patients treated with RAS-blockers, the risk of atherothrombosis decreases. This could be explained because by dampening local A-II and vice versa, an over activated ACE2 could produce the opposite effect (35); this does not happen in older people, because they have ACE2 expression reduced in all organs in which it has been described, but especially at the vascular level (36). At both the systemic and vascular level, ACE2 acts as a buffer, so that Angiotensin-II does not rise excessively when its receptor is exogenously blocked. Therefore, ACE2 is a key counter regulatory enzyme that degrades angiotensin II to angiotensin-(1–7), thereby attenuating its effects on vasoconstriction, sodium retention, and fibrosis. Although angiotensin II is the primary substrate of ACE2, that enzyme also cleaves angiotensin I to angiotensin-(1–9) and participates in the hydrolysis of other peptides (28). Both ACEs uses Zn²⁺ and Cl⁻ as cofactors for being metalloproteases (37). It could have therapeutic implications especially in people who use zinc as a nutraceutical (38).

The ACE2, but not the ACE is expressed in the type II pneumocytes (28) but it cannot be ruled out that endothelial ACE2 could collaborate coming from neighbouring microvascular endothelial cells as demonstrated in SARS (39). The true physiological significance of ACE in the pneumocyte is currently unknown.

After learning the important role played by ACE2 in SARS and COVID-19, it is strange that the CoV NL63-S, which also binds to ACE2, do not cause severe lung disease (40). This would lead us to investigate a co-receptor in COVID-19, a pathway that is not being considered.

There is no doubt that in COVID-19, the immunology plays a relevant role from the beginning. In the initial contagiousness, the older people are more vulnerable because of their immunosenescence (41). Afterwards, the immunology is responsible of the fever, inducing the virus directly a secretion of MIP1a/ CCL3 by the macrophage in the phase 1; this is responsible of a monophasic fever at hypothalamic level (42), that is resistant to non-steroidal anti-inflammatory drugs (NSADs) (e.g. ibuprofen, a cyclooxygenase inhibitor). In contrast, the fever induced by Tumoral Necrosis Factor alpha (TNFα) and Interleukin 1 (IL-1), the two usual endogenous pyrogens in flu and other viral infections, they do respond to NSAIDs (43).

Mainly in risk groups, by the seventh to the tenth day, some patients can develop a more severe phase; it has been correlated with chemokine secretion from endothelial cell (MCP-1) and the macrophage (CXCL10/IP-10). The massive knock-on (or pull effect) of chemokines attracts new macrophages, that are hyperactivated and release large amounts of cytokines, that can act locally or pass into the circulation. The more severe cases are related to a cytokine storm (commanded by IL6), related to a Macrophage Activation Syndrome (MAS), which is the cause not only of ADRS but occasionally, of myocarditis (see below), and in the most severe cases. Of a Multi-Organ Failure. This syndrome, initially described as Multiple Organ Dysfunction Syndrome (44), is not always lethal despite the impaired function of the liver, kidneys and brain (among other organs), but even when not lethal can leave disabilities in up to 15-20% of the patients.

**Insufficient response to the tested treatment. A change of strategy is urgent**

The genome of this mutated coronavirus, has been published recently (45) and it is the basis for the investigation of an adequate vaccine (46). Alignment of the full-length genome sequence showed that is a positive-sense RNA, 29903-bp (Watson–Crick base pairs) betacoronavirus. The closest relationship was with the bat SARS-like coronavirus strain (BatCov RaTG13), with 96% identity. The other rational therapy, convalescent plasma transfusion (47), are not available at present.

In the meantime, different therapies have been tried and their results may be considered as frustrating in high risk groups (6). Recently, two antiviral agents, remdesivir and ivermectin, are being tested at a great scale. Remdesivir, a classical antiviral agent, up to now not traded worldwide (48); ivermectin, a classical antiparasitic drug recently approved by the FDA for this epidemic because of its antiviral action (49).

Regarding to hydroxychloroquine / azithromycin (HC/AZ) and corticosteroids, two of the most tested therapies, we would like to point out a couple of comments related to their cardiometabolic aspects. Firstly, HC and AZ induce inhibition of cardiac hERG/IKr (rapid IK) potassium channel, inducing mild QT prolongation, with subsequent risk of ventricular arrhythmia and even cardiac arrest, thus needing QT monitoring. Secondly, high doses of corticosteroids not only produce hyperglycaemia in patients, even in not diabetics (50), but also raise blood pressure and in many cases lead to hypokalaemia, increasing the risk of arrhythmias. Other causes of non-diabetic hyperglycaemia, in many cases associated to corticosteroid administration would be high doses of loop diuretic, sepsis and very especially the enteral and parenteral nutrition (50, 51).

It is clear that we are still far from a useful vaccine and convalescent plasma transfusion. Furthermore, usual management of the disease is not enough, and two of the most commonly used treatments either have frequent side effects (corticosteroids) or have potentially severe adverse effects (HC/AZ). The question now is, what can we do pro-actively, in the meantime?
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What the truth does not hide: The COVID-19 is much more than an infectious disease and a dysfunctional immune response; it is also a cardiometabolic disease

Within the cardiometabolic (CM) aspects of COVID-19, we first looked at DM and HTN. Both of them, in addition to be among the main risk factors for pandemic mortality, are strictly related to age, weight gain and low physical activity. DM and HTN form part of three syndromes closely related but not synonymous, the Metabolic Syndrome (MS) (52), the CM Syndrome (14) and the CM Risk (53). MS is the core of the other two (Figure 1), and CM risk is the core of CMM/CMH born just a year later.

In the past, during the SARS and MERS outbreaks, the high prevalence of DM and HTN was attributed to the age (54) but no further studies were conducted.

In COVID-19, in a population of 686 patients with mean age 48.9 years, 42.7% females, the composite endpoints that consisted of admission to the intensive care unit (ICU), invasive ventilation or death, were analysed. The most prevalent comorbidity was HTN followed by DM. After adjusting for age and smoking status, DM and HTN had similar hazard ratio of reaching to the composite endpoints, but they never overtook COPD nor cancer (55).

It is known that other RNA viruses such as enterovirus B, coxsackie (B1, B3 and B5) and human mastadenovirus C are implicated in autoimmune destruction of pancreatic islet β cells, which results in a typical autoimmune T1DM (56). The question that immediately can be raised is if SARS-CoV-2 (also a RNA virus) is able to cause also a typical autoimmune T1DM. This possibility has not been discarded, but it would be more likely a not autoimmune, atypical T1DM (57). It looks strange that no cases of this type of DM had been described, in more than a million and a half test-confirmed patients, while in SARS, many cases were described in less than one thousand infected patients (58). The most surprising fact about this study, was that the pancreas damage was correlated with the expression of ACE2 in the endocrine tissues of the pancreas, leaving no doubt about a cause and effect relationship.

In relation to the heart, both HTN and DM can predispose to myocardiopathy (59) and in presence of viral respiratory infections, like flu annually (60) or now in COVID-19 (61), can predispose to heart failure (HF). This is without doubt the most frequent cardiac complication of these patients, especially if we add anaemia and/or tachyarrhythmia (flutter and atrial fibrillation, episodes of supraventricular or ventricular tachycardia) both not uncommon in Intensive Care Units (62, 63).

In addition, SARS-CoV-2 can induce severe myocarditis with TnT and NT-proBNP as biomarkers (64-66). This is not a surprise, because in animal models of CoV, myocarditis was described almost 30 years ago, in 1992 (67); but in SARS, although creatine-kinase MB isoenzyme was a predictor for death (58) no study has demonstrated any case of well-documented myocarditis. COVID-19 patients who develop myocardial injury, usually DM and/or elderly patients, have a more acute presentation, with higher incidence of ADSR and more frequent need for assisted ventilation than those without myocardial injury (64).

The likely damage of the myocardium in COVID-19 may be, at least partially, from the release of cytokines and secondary activation of the NF-kB signalling pathway (68); actually, the virus cannot penetrate because this cell does not have a receptor for it (69). Although abundant ACE2 (SARS-CoV and SARS-CoV-2 receptor) immunostaining was found in the heart, unfortunately the authors did not specify if was in the endocardium and/or myocardium of the affected tissue (70).

In aged COVID-19 patients or with history of coronary artery disease (CAD) an acute coronary syndrome (ACS) can also be seen for plaque vulnerability in the presence of a pro-inflammatory state with cytokine release (71) but from the experience in animals, could it be plausible that any of them could be due to arteritis? As we already pointed out at the beginning of this paper, arteriviridae, of the order of Nidovirales (like the CoV), are responsible of equine viral arteritis (EVA) (72) a disease that has too many similarities with COVID-19. The main one is that both of them can present with interstitial pneumonia and the microscopic findings, included diffuse alveolar damage with exudates, lymphocytic inflammation and multinucleated giant cells, suggesting hyperactive macrophages, were seen alongside large atypical pneumocytes (51). In EVA’s necropsies, antigens can be demonstrated not only within the cytoplasm of the epithelial cells, but also within endothelial, macrophages and cardiac myocytes (51), something has not been proven in COVID-19, at least not yet.

It should also be considered in COVID-19 the suitable diagnosis and management of other vascular complications such as stroke and finger ischaemia, both frequent in the ICUs (73, 74).

Practical aspects of COVID-19 regarding the personalized aspect of the Cardiometabolic Medicine

If we look at Figure 1 and Table 2, we can observe that there are as many possible combinations as patients; that implies that treatments must be personalised. This had been advocated during last years by the most prestigious societies and institutes, such as the AHA, ADA, AACE, ESC, EASD and the NICE institute among others.

Among all these combinations, we have chosen the following three, because they are highly topical in the CMM. They can be applied to patients with COVID-19 in their acute phase of hospitalization or from the respiratory point of view, in a more stable phase (prehospitalization phase or when they are already convalescing at home).

The first aspect regarding personalised treatment is related to HTN and COVID-19, a debate about keeping or discontinuing
ACEIs and ARBs started. The European Medicines Agency (EMA), European Society Cardiology (ESC) and American College Cardiology (AMC) all agreed that they should be continued in well-controlled patients and also in patients with previous cardiovascular pathologies of various kinds (28, 32).

Whilst discussing the above, a new debate has emerged with force: whether to start ACEIs/ARBs in patients without another clinical indication or not. Based upon what we had learned from SARS, the biological plausibility of salutary effects of ACEIs/ARBs in those with COVID-19 is intriguing (28) (32).

Table 2
Cardiometabolic evaluation
(Excluding exhaustive metabolic and cardiovascular genetic evaluation)

| METABOLIC TESTS | NON-METABOLIC TESTS |
|-----------------|---------------------|
| Glycaemia a and A1C (%) Lipid profile: | Transthoracic Echocardiography: mandatory in suspected or diagnosed HTN, DM, or in patients ≥ 60, or stroke evaluation |
| - HDL-C and TG (Metabolic Syndrome) | Supra-aortic Trunks Ultrasound: mandatory in DM, ≥60 or history of CVD |
| - LDL-C (Cardiometabolic Risk) if FH is suspected request a genetic LDL/ApoB100 receptor study | Incipient Atherosclerosis Evaluation |
| - ApoB and non-HDL-C (New targets alternative to LDL-C) | Intima-media thickness (IMT) |
| Glomerular Filtration Rate: Chronic Kidney Disease-Epidemiology | Ankle-brachial index (ABI) |
| Collaboration (CKD-EPI) | Ambulatory Blood Pressure Monitoring: Mandatory in: non-controlled HTN, history of masked HTN, suspected “white coat” effect, suspected overtreatment. |
| Ionic balance: | 24h-Holter Monitoring (ECG): if any of the following: sinus arrhythmia, bradycardia, tachycardia, anaemia, hyperthyroidism, extrasystole, complete LBBB, short PR, prolonged corrected QT |
| Na/K | Arterial stiffness evaluation: In HTN and in the course of an Aging evaluation |
| Ca/P | - Pulse wave amplitude (PWA) |
| Albumin / Creatinine ratio (urine sample): Essential in DM and HTN or low/borderline GFR | - Pulse wave velocity (PWV) |
| Emergent CV Risk Factors: | Coronary Calcium Score (CAC): with CCT |
| High-sensitivity C-Reactive Protein | If ≥60 with no risk factors; ≥50 with 1 risk factor; ≥45 if T1DM, FH or cardiovascular family history of premature coronary disease |
| Homocysteine: if high request polymorphisms C 677T and A1298C of MTHFR gen | Cardiorespiratory fitness (CRF): VO2-Max |
| Lipoprotein (a): only in initial consultation | Direct (cycle ergometry) |
| Uric acid: is always recommendable because it is related with MS, CMS and CMR, but especially if gout history or diuretic treatment | Indirect (Balke Treadmill Test) |
| Ferritin: especially recommendable if family history of hyperferritinaemia or fatty liver disease | Endothelial Function Study |
| If above the range request a genetic HFE test for study of polymorphisms HD63 and C282Y | - microvascular: EndoPAT / DRT4-lasser |
| Mutation G51C of IRE (iron responsive element) of L-ferritin gen | - macrovascular: Celermajer test |
| Hormonal & Vitamins Study: | Especific imaging techniques |
| Hormones: | Cranial CT / RM: for cerebral small vessel disease |
| Women: β-estradiol / FSH / LH. | study, including «white matter lesions» and «lacunar infarcts» |
| Men & Women: TSH / testosterone / DHEA / IGF-1 | Cardio-RM: old necrosis / current ischemia / EF |
| Cortisol in 24 h-urine | Angio-CT: evaluation of a stenosis detected by doppler / possibility of stent |
| Vitamins: Vitamin D: essential for right management of Aging | Angio-MR: alternative to the CT in stenosis evaluation |
| - Folic acid and Vitamin B12 | |
| Hypercoagulability study | Noninvasive testing in myocardial ischemic evaluation (CAC Score >75th percentile) |
| D-dimer; Prothrombin G20210A Factor II mutation; Factor V Leyden; Proteins C and S; Anti-phospholipid autoantibodies (aPL): anticoagulants (aCL) / anti antiphosphatidylserine; Anti-beta2-glycoprotein I antibodies (anti apo-H); Lupus anticoagulant LA: evidence of a circulating LA sensitive PTT or DRVVT diluted Russell Viper Venom test | Stress echocardiography, on treadmill or bike or drugs (dobutamine) |
| Intima-media thickness (IMT) | Ergometry with submaximal Bruce protocol |
| Incipient Atherosclerosis Evaluation | 99m Tc-MIBI-SPECT if necessary: LBBB, females |
| Coronary Calcium Score (CAC): with CCT | Optional Tests: |
| If ≥60 with no risk factors; ≥50 with 1 risk factor; ≥45 if T1DM, FH or cardiovascular family history of premature coronary disease | Handgrip: essential in an Aging evaluation |
| Coronary Calcium Score (CAC): with CCT | Interventionist catheterism if required |

CT: Computed tomography; CCT: Cardiac computed tomography; CVD: Cerebrovascular diseases; DHEA: Dehydroepiandrosterone. DM: Diabetes mellitus. EF: Ejection fraction; FH: Familial hypercholesterolemia activity. FSH: Follicle stimulating hormone. GFR: Glomerular Filtration Rate. HFE (gene and protein implicated in HH: Hereditary Hemochromatosis); High FE2+: HTN: hypertension. IGF-1: Insulin-like growth factor-1. LH: Luteinising hormone. LBBB: Left bundle branch block. TG: Triglycerides. TSH: Thyroid-stimulating hormone; (1) In hospital setting, glycaemia, hsCRP and ferritin are not valuable in a basal cardiometabolic status. Must be repeated at 2-month follow-up; (2) LDL-P (particle) (nmol/L) better than LDL-C if a Nuclear Magnetic Resonance Spectroscopy is available; (3) If Ca is high or P is low: Request PTH (parathyroid hormone). Also request PTH, if GFR is<50; (4) As stress measure in Aging and as diagnostic tool in patients with suspected secondary HTN.
have reduced ACE2 expression and upregulation of angiotensin II (AII) and consequently its proinflammatory signalling way could be hyper-activated. The increase in ACE2 levels, and consequent reduction of AII, ACEIs/ARBs treatment would be especially beneficial for this age subgroup, in line with the evidence of a protective role of AII antagonism against sepsis-associated acute lung injury (22).

The third aspect would be the combination of T2DM and Heart failure (HF) (the most frequent cardiac complication in any of the phases of the disease), which is present in a high percentage of patients, especially those at higher risk. Excluding secondary DM (corticosteroids, diuretics, sepsis, enteral / parenteral nutrition), these patients would benefit from taking a SGLT2 inhibitors (Sodium-glucose co-transporter-2, or gliflozins) (78) when they are in a stable phase. In these patients, the risk of ketoacidosis is quite low, but it may happen in the hospitalized patient (myocardial or cerebral infarction, sepsis ...) and more likely, if they are being treated with these drugs.

Cardiometabolic syndrome (Sowers et al 2001): Gathering the classic components of the Metabolic Syndrome (Central obesity, hyperglycaemia, low HDL, hypertriglyceridaemia, pre-hypertension or hypertension) and adding endothelial dysfunction, oxidative stress, and inflammation and hypercoagulability biomarkers. Cardiometabolic Risk (Khan, Buse, Ferrannini and Stern, 2005): The Cardiovascular Risk maintains the Metabolic Syndrome as its core, and includes what the Cardiometabolic Syndrome had provided. It only keeps the inflammatory and hypercoagulability biomarkers, leaving the rest of biomarkers (Endothelial Dysfunction and Oxidative Stress) for research. Apart from the above, their main contribution was to add classic risk factors, not related to insulin resistance such as a family history of premature coronary disease (<55 in male / <65 in female), LDL-C and tobacco use.

On the other hand, SARS-Cov-2 particularly attacks DM patients (75), more frequently and with greater severity, being DM the cardiometabolic disease par excellence because they are often accompanied by other risk factors and atherothrombotic disease (ATD) (76). From lessons learned from experimental DM, we know that affected animals have a decreased vascular expression of ACE2, and it has been associated with an increased predisposition to ATD, coincidentally, as in elderly patients (77). This would lead us to extend the trials with RAS-blockers recommended by the NIA (36), if it were not for the fact that most of them are already on that treatment.

The second aspect should be regarding the use of statins, the star drug treatment of CMM/CMH in an outpatient context. This is a subject of debate in hospitalized patients except for those with previous history or recent stroke or ACS, and possibly in those with familiar hypercholesterolemia, but only if a prolonged benefit is presumed. Apart from this, the risk/benefit balance in patients with COVID-19 remains to be clarified.

Furthermore, in HF we should consider not only the association with DM, but also its type; i.e., if HF is acute or chronic, and if chronic, to consider if reduced or preserved ejection fraction (HFrEF or HfP EF), because they have different therapeutic approaches (79).

Synthesising all the above, it looks like we are facing a disease, which is much more than an infectious/immune respiratory disease: simultaneously, it is also a cardiometabolic disease. The respiratory side is well known and managed, with poor results in the risk group, exactly the one that has more cardiometabolic alterations. Regarding this group of disorders,
discussed in detail above, we do not know why they are always approached separately, when there are many advantages of their overall management, and it would have an unavoidable impact on morbidity, lethality and costs as demonstrated in patients followed up in outpatient clinic (27, 28).

In this sense, are we really optimizing the main targets during the hospitalisation? Do we give the value it really has? We are not speaking of LDL-cholesterol, as in the case of high CV risk patients followed as outpatients. In the presence of recent events where the LDL can be important, but even more important is the inflammation, to justify treatment with statins (80). Except in this last group, we speak of high and low blood pressure and/or high and low glycaemia, both type of decompensations are important in the two diseases. However, if we have to choose in the case of blood glucose, hypoglycaemia can have more serious consequences, especially in older patients, as the ADA stated in their last report (81).

Furthermore, we talk also of early detection and treatment of heart failure, arrhythmia or stroke that can go unnoticed in ICU patients with induced coma, and so many more aspects that should not be overlooked by monitoring only the lung disease when this is not the one that will always kill our patients.

**Common molecular aspects between cardiometabolic diseases and Aging. Do they embrace the immunosenesence also?**

One of the great advantages that CMM has over classic CV medicine, and that we did not say previously, is that CMM is seriously concerned with aging, both at clinical level (for healthy aging) and at research level.

In the same sense, whilst most researchers and clinicians interested in COVID-19 have attributed its greater aggressiveness to previous comorbidities (6), as convinced CMM supporters and of its interest in the aging process, we believe that despite our suspicions are correct, this is not enough. We will review the molecular aspects that underlie human aging and whether any of these mechanisms are involved in immunosenescence. If this is true, it will be easier in the future to deal with the whole process of aging, by slowing down the harmful processes and favouring the beneficial ones.

A modern theory of aging named «A Vascular Theory of Aging» (81), has rescued from oblivion two old aphorisms: «man has the age of his arteries» (Thomas Sydenham, 17th century) and «man has the age of his endothelium» (Rudolf Altshul) (82). Both were visionaries, but especially the latter when he published his book (in 1954) without even knowing the nitric oxide, the main endothelial product, discovered in the 1980s, leading Furchgott, Ignarro and Murad, to be recognized as the Nobel Prize in Physiology or Medicine in 1998.

Advances in the physiology in the last decade consider atherothrombosis and aging as an inflammatory disorder (81) but, which molecular aspects are common to both conditions?. There are many important molecular aspects involving them, but the principal one is the combined activation of what we consider as the «lethal triad». The three elements of this triad are: 1) the mTOR (mammalian target of rapamycin), 2) the NF-kB (Nuclear Factor Kappa-B) signalling pathways (83) on one side, and 3) the NOX enzymes (84) of the submembranous cytoplasm on the other side, which release more free radicals than the mitochondria itself.

This triad alters the lipidoma at three levels, (membranous, cytoplasmatic and circulating), as well as the so-called endoplasmatic reticulum stress. This is a key process to understand both the endothelial dysfunction (85) preceding atherothrombotic disease (ATD), and the dysfunction of the beta cell (86) in T2DM and in the aging process (87) in part by the endothelial dysfunction at microvascular level within of vascular theory of aging (81). Fortunately, autophagy and especially mitophagy (mitochondrial autophagy) (88) protect us from apoptosis (programmed cell death), as long as our genetics and epigenetics (influenced by healthy nutrition and regular physical activity) allow it.

Finally, regarding the immunosenescence, its three hallmarks are: 1) the reduction in the number of peripheral blood naïve T cells population; 2) a relative increase in the frequency of CD28 memory T-cells subset, and 3) a low-grade chronic inflammation that characterizes aging: “inflamm-aging” or “inflammaging” (89). Surprisingly, there are extensive data showing, moreover, that latent persistent human cytomegalovirus infection is also associated with age-related immune dysfunction in the T cells, which might enhance immunosenescence. In this way, in the case of COVID-19, in older groups, a virus could be predisposing to another virus (90). If we can still get even more surprised, both the endoplasmatic reticulum stress and autophagy are involved in the process of immunosenescence (91). In accordance with the above, could the immunosenescence be related to the special aggressiveness of COVID-19 in the elderly?

**Conclusions**

The coronavirus disease 2019 (COVID-19) outbreak is evolving rapidly worldwide. From a dangerous situation due to its respiratory facet, there is another less well-known one (at least up to now), the cardiometabolic facet that can precede, accompany or even kill us if we do not keep our eyes open.

The high lethality of SARS-COV-2 in subjects with previous hypertension, diabetes and atherothrombotic disease. In addition to several cardiovascular complications arising during the hospitalization, include severe arrhythmias, acute heart failure by myocarditis and stroke. Often forgotten in reported series, are an ample justification that we are dealing with a disease of a great cardiometabolic potential that we should not overlook.

Other cardiometabolic aspects, like the possible N-glycation of viral protein M in patients with hyperglycaemia will need further investigation. In addition, the role of serinprotease TMPRSS2, synthesized by the endothelium and involved
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in the interaction of the virus with its recipient, could be a potential target for the treatment. Its secretion is not the unique endothelial product, also ACE2, considered only of pulmonary origin could derive from the microvascular endothelium as demonstrated in SARS.

The role of convalescent plasma transfusion and recent antiviral agents such as ivermectin and remdesivir, in improving COVID-19 prognosis in high-risk patients remains to be demonstrated.

In the meantime, here we have learned that Cardiometabolic Medicine could help us to reduce, using an anticipative and integrated approach, the morbidity and mortality related to this current SARS-CoV-2 pandemic while reducing costs.

Finally, the molecular aspects of aging in general and in the immunosenescence in particular, will serve us better in the near future. Knowing what happens, the solution is a matter of time. With healthier old people, we will make it more difficult for viral infections in general, not only in the course of epidemics or pandemics. On the other hand, in these situations, the best way for prevention is to not let them happen and for this purpose, CEPI (Coalition for epidemic preparedness innovations) exists, let us help them with more resources.

Acknowledgment: The authors are indebted to Dr. Joseph Hogg, partner at Abbey House Medical Centre, for his critical review, helpful comments and careful English revision.

Conflict of interest: None declared.

Ethical Standards: Not applicable.

References

1. Lavi E, Schwartz T, Jin YP, Fu L. Nidovirus infections: experimental model systems of human neurologic diseases. J Neuropathol Exp Neurol 1999;58:1197-1206.
2. Cruz F, Newton R, Sanchez A, Ireland J, Mughini-Gras L, Moreno MA, Fores P. Equine viral arteritis in breeding and sport horses in central Spain. Res Vet Sci 2017;115:88-91. DOI: 10.1016/j.rvsc.2017.01.022
3. Bell DM. World Health Organization Working Group on International and Community Transmission of SARS. Public health interventions and SARS spread, 2003. Emerg Infect Dis 2004;10:1900–1906.
4. Wernery U, Lau SK. Woc PC. Genomics and zoonotic infections: Middle East respiratory syndrome. Rev Tech Sci 2016;35:191-202.
5. Li Q, Guan X, Wu P, Wang X, Zhou L, Tong Y et al. Early Transmission Dynamics in Wuhan, China, of Novel Coronavirus-Infected Pneumonia. N Engl J Med 2020;382:1199-1207
6. Guo YR, Cao QD, Hong ZS, Tan YY, Chen SD, Jin HJ et al. The origin, transmission and clinical therapies on coronavirus disease 2019 (COVID-19) outbreak - an update on the status. Mil Med Res 2020;7:11. doi: 10.1186/s40779-020-00240-0
7. Moe SA, Alhowikan AM, Al-Khiawi T, Moe IM, Halebeki PM, Iqbal M et al. Novel coronavirus 2019-nCoV: prevalence, biological and clinical characteristics comparison with SARS-CoV and MERS-CoV. Eur Rev Med Pharmacol Sci. 2020;4:2012-2019.
8. Driggan E, Madhavan MV, Bikdeli B, Chuich T, Carvalan C, Hyder AA, Lazo-Porras M, Oni T et al. Understanding the rise of cardiometabolic diseases in low- and middle-income countries. Nat Med 2019;25, 1667–1679.
9. Ras G. Prevention or reversal of cardiometabolic diseases. J Clin Prev Cardiol 2018;7:22-28
10. Plowright RK, Parrish CR, McCallum H, Hudson PJ, Ko AI, Graham AL, Lloyd-Smith JO. Pathways to zoonotic spillover. Nat Rev Microbiol 2017;15:502-510
11. Pyrc K, Berkhout B, van der Hoek L. Identification of new human coronaviruses. Expert Rev Anti Infect Ther 2007;5:245-253
12. Guan N. Three Emerging Coronaviruses in Two Theories of The Story of SARS, MERS, and Now COVID-19. Am J Clin Pathol 2020;153:420-421.
13. Imai Y, Kuba K, Penninger JM. The discovery of angiotensin-converting enzyme 2 and its role in acute lung injury in mice. Exp Physiol 2008;93: 543–548.
14. Chan JF, To K, Tse H, Jin DY, Yuen KY. Interspecies transmission and emergence of novel viruses: lessons from bats and birds. Trends Microbiol 2013;21:544-555
15. Neuman BV, Kiss G, Kunding AH, Bheila D, Baksh MF, Connolly S et al. A structural analysis of M protein in coronavirus assembly and morphology. J Struct Biol 2011;174: 11–22. doi: 10.1016/j.jsb.2010.11.021.
16. Huang C, Wang Y, Li X, Ren L, Zhao J, Hu Y et al. Clinical features of patients infected with 2019 novel coronavirus in Wuhan, China. Lancet 2020;395:497-506. doi: 10.1016/S0140-6736(20)30183-5
17. Mourir S, Talbot PJ. Sequence analysis of the membrane protein gene of human coronavirus OC43 and evidence for O-glycosylation. J Gen Virol 1992;73:2731-2736.
18. Welsh KJ, Kirkman MS, Sacks DB. Role of Glycated Proteins in the Diagnosis and Management of Diabetes: Research Gaps and Future Directions. Diabetes Care. 2016;39:1299–1306
19. Vaduganathan M, Vardeny O, Michel T, McMurray JJJ, Pfeffer MA, Solomon SD, Renin-Angiotensin-Aldosterone System Inhibitors in Patients with Covid-19. N Engl J Med March 2020;381:382
20. Asselta R, Paraboschi EM, Mantovani A, Daga S, ACE2 and TMPRSS2 variants and expression as candidates to sex and country differences in COVID-19 severity in Italy. MedRxiv, 2020.DOI: 10.1101/2020.03.30.20047878
21. Michelen M, Jones N, Stavropoulou C, on behalf of the Oxford COVID-19 Evidence Service Team Centre for Evidence-Based Medicine, Nuffield Department of Primary Care Health Sciences, University of Oxford (2020) In patients of COVID-19, what are the symptoms and clinical characteristics of mild and moderate cases? April 1, 2020. https://www.cernb.net/covid-19-in-patients-of-covid-19-what-are-the-symptoms-and-clinical-features-of-mild-and-moderate-case/
22. Hamming I, Timens W, Bulthuis ML, Lely AT, Navis G, van Goor H. Tissue distribution of ACE2 protein, the functional receptor for SARS coronavirus. A first step in understanding SARS pathogenesis. J Pathol 2004;203:631-637
23. Bivishi Ch, Maddux TM, Merssari FH. Coronavirus Disease 2019 (COVID-19) Infection and Renin Angiotensin System Blockers JAMA Cardiol. Published online April 3, 2020. doi:10.1001/jamacardio.2020.1288
24. Ballantyne CM. Cardiometabolic risk assessment: an approach to the management of patients with severe cardiovascular disease. JAMA Cardiol. Published online April 3, 2020. doi:10.1001/jamacardio.2020.1288
25. Hooper NM, Turner AJ. Understanding the rise of cardiometabolic diseases in low- and middle-income countries. Nat Med 2019;25, 1667–1679.
syndrome coronavirus receptors, ACE2 and CD209L in different organ derived microvascular endothelial cells. Zhonghua Yi Xue Za Zhi 2007;27:83-83
40. Hofmann H, Pyrc K, van der Hoek L, Geier M, Berkhout B, Pöhlmann S. Human coronavirus NL63 employs the severe acute respiratory syndrome coronavirus receptor for cellular entry. Proc Natl Acad Sci USA 2005;102:7988-7993
41. Tchkonia T, Zhu Y, Deursen JV, Campisi J, Kirkland JL. Cellular senescence and the senescent secretory phenotype: therapeutic opportunities. J Clin Invest 2013;123: 966-972
42. Mikiya FJ, Sancibrian M, Myers RD. Fever induced by macrophage inflammatory protein-1 (MIP-1) in rats: hypothalamic sites of action. Brain Res Bull 1991;27:701-6
43. Dinarello CA. The history of fever, leukocytic pyrogen and interleukin-1. Temperature (Austin) 2015;2:8-16. doi: 10.1002/2338940.201510708
44. Baue AE. Multiple, Progressive, or Sequential Systems Failure. A Syndrome of the 1970s Arch Surg 1975;109:779-781
45. Lu R, Zhao X, Li J, Niu P, Yang B, Wu H, et al. Genomic characterisation and epidemiology of 2019 novel coronavirus: implications for virus origins and receptor binding. Lancet 2020;22:395-565-574
46. Lurie N, Saville M, Hatchett R, Halton J. Developing Covid-19 Vaccines in Pandemic Speed. N Engl J Med March 30, 2020; DOI:10.1056/NEJMp2005630
47. Roback JD, Guarnier J. Convalescent plasma to treat COVID-19: Possibilities and challenges. JAMA Mar 27, 2020; DOI:10.1001/jama.2020.4940
48. Wang, M, Can R, Zhang J, Yang X, Liu J, Xu M, Shi Z, Hu Z, Zhong W, Xiao G. Remdesivir and chloroquine effectively inhibit the recently emerged novel coronavirus (2019-nCoV) in vitro. Cell Res 2020;Feb 2; [e-pub]. https://doi.org/10.1038/s41422-020-0282-0
49. Caly L, Druce JD, Catton MG, Jans DA, Wagstaff KM. The FDA-approved Drug Ivermectin inhibits the replication of SARS-CoV-2 in vitro. Antiviral Res 204787, 2020. doi:10.1016/j.antiviral.2020.104787
50. Corsino L, Dhatatarya K, Umpierrez G. Management of Diabetes and Hyperglycemia in Hospitalized Patients. www.endotext.org, October 1, 2017.
51. Gosmanov AR, Umpierrez GE. Management of Hyperglycemia during Enteral and Parenteral Nutrition Therapy. Curr Diab Rep 2013;13: 155–162
52. Hanefeld M, Leonhardt W. Das metabolische Syndrom. Dt Gesundheitswesen 1991;36:545-551
53. Vasudevan AR, Ballantyne CM. Cardiometabolic risk assessment: an approach to the prevention of cardiovascular disease and diabetes mellitus. Clin Cornerstone 2005;7:7-16.
54. Alnaimi KH, Abedi GR, Midlyce CM, Alkhamis A, Alsaqer T, Almoudi A et al. Diabetes Mellitus, Hypertension, and Death among 32 Patients with MERS-CoV Infection, Saudi Arabia. Emerg Infect Dis 2020;26:166–168.
55. Guan W, Liang W, Zhong S, et al. Clinical characteristics of coronavirus disease 2019 in China. N Engl J Med 2020;382:1708-1720. doi: 10.1056/nejmoa2002032
56. Yang JK, Lin SS, Ji XJ, Guo LM. Binding of SARS coronavirus to its receptor damages islets and causes acute diabetes. Acta Diabetol 2010;47:193-9
57. Gradman AH, Rutschiha F, COVID-19 Illness and Heart Failure: A Missing Link? JACC. Heart Failure, 2020;DOI: 10.1016/j.jchf.2020.03.004
58. Rawal G, Kumar Y, Yadav S, Amrita Singh. Anemia in Intensive Care: A Review of Vasudevan AR, Ballantyne CM. Cardiometabolic risk assessment: an approach to the prevention of cardiovascular disease and diabetes mellitus. Clin Cornerstone 2005;7:7-16.
59. Alnaimi KH, Abedi GR, Midlyce CM, Alkhamis A, Alsaqer T, Almoudi A et al. Diabetes Mellitus, Hypertension, and Death among 32 Patients with MERS-CoV Infection, Saudi Arabia. Emerg Infect Dis 2020;26:166–168.
60. Guan W, Liang W, Zhong S, et al. Clinical characteristics of coronavirus disease 2019 in China. N Engl J Med 2020;382:1708-1720. doi: 10.1056/nejmoa2002032
61. Yang JK, Lin SS, Ji XJ, Guo LM. Binding of SARS coronavirus to its receptor damages islets and causes acute diabetes. Acta Diabetol 2010;47:193-9
62. Gradman AH, Rutschiha F, COVID-19 Illness and Heart Failure: A Missing Link? JACC. Heart Failure, 2020;DOI: 10.1016/j.jchf.2020.03.004
63. Rawal G, Kumar Y, Yadav S, Amrita Singh. Anemia in Intensive Care: A Review of Vasudevan AR, Ballantyne CM. Cardiometabolic risk assessment: an approach to the prevention of cardiovascular disease and diabetes mellitus. Clin Cornerstone 2005;7:7-16.
64. Alnaimi KH, Abedi GR, Midlyce CM, Alkhamis A, Alsaqer T, Almoudi A et al. Diabetes Mellitus, Hypertension, and Death among 32 Patients with MERS-CoV Infection, Saudi Arabia. Emerg Infect Dis 2020;26:166–168.
65. Guan W, Liang W, Zhong S, et al. Clinical characteristics of coronavirus disease 2019 in China. N Engl J Med 2020;382:1708-1720. doi: 10.1056/nejmoa2002032
66. Edwards S, Small JD, Geratz JD, Alexander LR, Baric RS. An Experimental Model for Myocarditis and Congestive Heart Failure after Rabbit Coronavirus Infection. J of Inf Dis 1992;165:154-40
67. Maier HK, Schips TG, Wietelmann A, Krüger M, Brunner C, Sauter M, et al. Cardiomyocyte-specific IkB kinase (IKK)/NF-kB activation induces reversible inflammatory cardiomyopathy and heart failure. Proc Natl Acad Sci USA 2012;109:11794-11799
68. Schaufeleberger M, Drexler H, Schieffer E, Wedelberg K. Angiotensin-converting enzyme gene expression in skeletal muscle in patients with chronic heart failure. J Card Fail 1998;4:185-191
69. Yang JK, Lin SS, Ji XJ, Guo LM. Binding of SARS coronavirus to its receptor damages islets and causes acute diabetes. Acta Diabetol 2010;47:193-9
70. Toussilouts D, Antoniadis C, Koulamoulis N, Stefanidis C. Pro-inflammatory cytokines in acute coronary syndromes: from bench to bedside. Cytokine Growth Factor Rev 2016;27:225-233
71. Del Piero F. Equine Viral Arteritis. Vet et Pathol 2000;37:287-296. https://doi.org/10.1354/vp.37-4-287
72. Minahan RE, Bhardwaj A, Williams MA. Critical monitoring of future for cardiovascular disease. New Horiz 1997;5:406-21.1997
73. Labay G, Mostul CJ, Almoudi A et al. The Hoorn Study, on the development of recurrent disease, in patients who test positive for SARS-CoV-2. PLoS One 2020;15:e0229785
74. Landry G, Mostul CJ, Almoudi A et al. The Hoorn Study, on the development of recurrent disease, in patients who test positive for SARS-CoV-2. PLoS One 2020;15:e0229785
75. Sano M. A new class of drugs for heart failure: SGLT2 inhibitors reduce sympathetic overactivity. J Cardio 2018;71:471-476
76. Hanefeld M, Leonhardt W. Das metabolische Syndrom. Dt Gesundheitswesen 1981;36:545-551
77. Tiikellis C, Pickering R, Tsorotes D, Du XJ, Kiriazis H, Nguyen-Huu TP et al. Interaction of diabetes and ACE2 in the pathogenesis of cardiovascular disease in experimental diabetes. Clin Sci (Lond) 2012;123:519-529.
78. Bloomgarden ZT. Diabetes and COVID-19. J Diabetes 2020;12:347-348 doi: 10.1111/1753-0407.13027.
79. Zeng JH, Liu Y, Yuan J, Wang F, Wu W, Li J et al. First Case of COVID-19 Infection with Fulminant Myocarditis Complication: Case Report and Insights. Preprints 2020;2020030180 doi: 10.20944/preprints2020030180v1