Covid-19 Pandemic: Cardiovascular and Neurologic Impact, Early Signs and Symptoms

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Abstract

Background: Scientists and physicians continue their cutting-edge work on bacteria, parasites and viruses, responsible for 17 million deaths each year worldwide. The purpose of this article is to review our clinical experience with COVID-19 patients observed in the Cardiovascular Division of Pompidou Hospital (University of Paris, France) and the Department of Neurology of the Eastern Piedmont University (Novara, Italy), related with the impact on the cardiovascular, hematologic, neurologic systems and sense organs. Mathematical modeling for pandemic spreading control (protection versus hazards), proposed by Vienna University of Technology (Austria), are analyzed.

Methods: We sought to characterize the cardiovascular and neurologic manifestations, in patients with Covid-19. Special attention was given to initial signs and symptoms in order to facilitate early diagnosis and therapy. Indications of ECMO membrane oxygenation for cardio-respiratory support are evaluated, mathematical models for control pandemic spreading are described.

Results: Early neuro-sensorial symptoms like anosmia and dysgeusia are useful for diagnostic, patient’s isolation, and treatment. Angio-hematologic acro-ischemia syndrome and dermatological signs are mostly related with immune dysregulation, neoangiogenesis, vasculitis, and vessel thrombosis resulting from “cytokine storm syndrome”. Acro-ischemia syndrome allows diagnosis and early treatment of disseminated intravascular coagulation, with mortality risks for children and adults.

Conclusion: Covid-19 is a prothrombotic disease with a unique global lethality. Strong inflammatory response to viral infection severely affects cardiovascular, and neurologic systems, as well as respiratory, immune and hematologic systems. Fast identification of acro-ischemia syndrome permits the care of disseminated intravascular coagulation complications. Early symptoms like gustatory and olfactory loss are useful for Covid diagnostic. Mathematical models play a significant role in the understanding of COVID-19 transmission mechanisms, structures, and features. It should be of interest for policymakers and for explaining recurrent epidemic outbreaks.

Abbreviations: COVID : Corona Virus Disease; SARS CoV : Severe Acute Respiratory Syndrome Coronavirus; ECMO: Extra Corporeal Membrane Oxygenation; ACE: Angiotensin Converting Enzyme; RT-PCR : Real Time - Polymerase Chain Reaction; PEEP: Positive End-Expiratory Pressure; ARDS: Acute Respiratory Distress Syndrome; HFNC: High Flow Nasal Cannula; NIV: Non-Invasive Ventilation; L/min: Liter per Minute; HEPA: High Efficiency Particulate filter; CMR: Cardiac Magnetic Resonance; DVT: Deep Venous Thrombosis; MERS-CoV: Middle East Respiratory Syndrome Coronavirus; D-DIMER: Fibrin Degradation Product; ICU: Intensive Care Unit; DIC: Disseminated Intravascular Coagulation; CNS: Central Nervous System; CT: Computed Tomography; LU: Lung Ultrasound; SEIR Epidemiology model: Susceptible-Exposed-Infectious-Recovered
Background

COVID-19 caused by coronavirus 2 (SARS-CoV-2) has been declared a pandemic by the World Health Organization on March 11, 2020 [1]. After 3 months of critical clinical situation in Western Europe, the number of Covid cases, severity and mortality rates showed optimistic parameters. However, progressively coronavirus returns to Europe in September 2020 (Figure 1). As of 4 November 2020, COVID-19 has led to over 47.8 million confirmed infections and 1,220,224 deaths from coronavirus disease-2019 worldwide. Like most infections caused by members of the coronavirus family, SARS-CoV-2 manifests itself with upper respiratory tract infections and flu-like symptoms of varying severity. However, Covid-19 is unique in its ability to cause a multi-organ disease, with involvement of the cardiovascular and nervous systems. Coronavirus can send the body’s immune system into cataclysmic overdrive, and cause blood thrombosis that impede circulation to lungs, heart, brain, or limbs [2-4].

Keywords: Covid-19 pandemic, Cardio-vascular & Neuro-sensorial impact, Acro-ischemia Syndrome, Disseminated intravascular coagulation, Anosmia, Dysgeusia, Cytokine storm syndrome, Ecmo membrane oxygenation, Synchrotron radiation diffraction

The main goal of this article is to review our clinical experience observed with COVID-19 patients during the 2020 pandemic in the Cardiovascular Division of Pompidou Hospital (University of Paris, France) and the Department of Neurology of the Eastern Piedmont University (Novara, Italy). It analyses the impact on the cardiovascular, hematologic, neurological systems and initial signs and symptoms of Covid-19. Additionally, we present mathematical models proposed by Vienna University of Technology (Austria), for control pandemic spreading (protection versus hazards).

Covid-19, from SARS to Multiorgan Disease

Although Severe Acute Respiratory Syndrome (SARS) is at the core of the disease, covid-19 has revealed itself to be more than a simple viral pneumonia. SARS-CoV-2 disease seriously affects respiratory, cardiovascular, neurologic, and immune systems [5-7]. The three most common initial symptoms in COVID are fever, dry cough, and shortness of breath. Patients can also suffer from muscle pain, sore throat, nasal congestion, headache, loss of smell and taste and persistent hiccups. Current clinical experience is showing that it is important to monitoring blood oxygen saturation using pulse oximetry devices to indicate mask oxygen therapy. Non-invasive ventilation is recommended. Barotrauma in mechanical ventilation can be observed creating rupture of the alveolus [8,9]. Characteristics of COVID patients who become seriously ill are different of classic “pneumonia” or ARDS (Acute Respiratory Distress Syndrome). Coronavirus causes prolonged and progressive hypoxia, desaturation, leading to organ failure and fatal outcome. Thrombolytic therapy in COVID-19 patients with ARDS may be of benefit due to the unusually high incidence of pulmonary embolism and pulmonary thrombosis, particularly microvascular thrombosis, which are thought to contribute significantly to hypoxemia. It may also ameliorate the effects of extravascular and intra-alveolar fibrin deposition described in ARDS [10-12].
Covid-19 patients can remain subjectively comfortable even when their oxygen saturation levels fall far below borderline ranges, becoming a “silent hypoxemia”, low oxygen saturation associated with hypopacnia can be observed in pulmonary artery embolism [13]. Thoracic CT angiography scan and lung ultrasound imaging provide information for COVID diagnose and follow-up [14,15].

**Cardiovascular and Hematological Impact**

Covid-19 can have fatal consequences for people with underlying cardiovascular disease [7,16,17]. Over 65 years-old, patients with coronary artery disease, arterial hypertension and cardiac arrhythmias are at more severe risk. More worrisome is the fact that COVID could cause heart damage in patients who did not have any previous problems [7]. It is reasonable to expect that significant cardiovascular complications from Covid-19 will occur in patients with severe symptoms due to the strong inflammatory response associated with this disease that affects the myocardium. The critical cases of Covid-19 are those which have reported respiratory failure, septic shock with evolution towards multiorgan failure. 

In a new study including 100 patients recently recovered from COVID-19 infection [18], cardiovascular magnetic resonance imaging (CMR) revealed cardiac involvement in 78 patients (78%) and ongoing myocardial inflammation in 60 patients (60%), independent of pre-existing conditions, severity and overall course of the acute illness, and time from the original diagnosis. These findings indicate the need for ongoing investigation of the long-term cardiovascular consequences of COVID-19.

**Viral Myocarditis**

1% to 5% of all patients with acute viral infections may involve the myocardium [19]. The incidence of myocarditis is between 10 to 22 cases per 100,000 persons, it was observed in many viruses including enterovirus coxsackie B, parvovirus B-19, herpesvirus, influenza virus, and Covid-19 [20]. Viral myocarditis symptoms can range from nonspecific fatigue and shortness of breath to more aggressive symptoms that mimic acute coronary syndrome. After the initial acute phase, remission of symptoms can be observed or the viral infection may persist, leading to a persistent autoimmune-mediated inflammatory process with continuing symptoms of heart failure. Pulmonary myocarditis associated with influenza A (H1N1) virus were observed in previously healthy patients [21]. Treatments include drugs, cardiac bio-assist [22,23], mechanical circulatory assist devices or heart transplantation [24].

**Hematology**

Clinical patient’s outcome allows to consider COVID-19 as a new hematologic disease, thrombosis would be favored by the vascular attacks related to the Covid-19 infection. Curative anticoagulation in these patients prevents endothelial lesions. Angiopoietin-2 as a marker of endothelial activation is a good predictor factor for intensive care unit (ICU) admission [25-28]. Pulmonary artery and vein thrombo-embolism doubled among ICU patients. Endotheliopathy, the activation of coagulation and the inflammatory syndrome contribute to increasing the risk of thrombosis, causes seem to be multifactorial. In obese patients, there is also impaired ventilation which can affect inflammation and organs perfusion. Elevated levels of D-dimers and clotting factors are known risk factors for pulmonary embolism in these cases [10]. COVID-19 seems to be an additional risk factor for Deep Vein Thrombosis (DVT). The prevalence of DVT in hospitalized patients with SRAS-CoV-2 infection is high and is associated with adverse outcomes [29].

**Pathological Findings**

Autopsies around the world show that the lungs of the COVID-19 had hundreds of micro clots inside the pulmonary vasculature, not observed in classic pneumonia. In some cases, migrate causing myocardial ischemia or brain stroke. In 38 autopsies analyzed in North Italy [30], the predominant pattern of lung lesions in COVID-19 patients was diffuse alveolar damage, as previously described for the other two coronavirus that infect humans, SARS-CoV and MERS-CoV. Hyaline membrane formation and pneumocyte atypical hyperplasia were frequently found. The main relevant finding was the presence of platelet-fibrin thrombi in small arterial vessels; this important observation fits into the clinical context of coagulopathy which dominates in these patients and which is one of the main targets of therapy [10-13].

Extensive histopathological analysis [31] showed that SARS-CoV-2 infection can result in diverse, multi-organ pathology, the most significant being in the lungs (diffuse alveolar damage in its different phases, microthrombi, bronchopneumonia, necrotizing bronchiolitis, viral pneumonia), heart (lymphocytic myocarditis), kidney (acute tubular injury), central nervous system (microthrombi, ischemic necrosis, acute hemorrhagic infarction, congestion, and vascular edema), lymph nodes (hemophagocytosis and histiocytosis), bone marrow (hemophagocytosis) and vasculature (deep vein thrombosis).

**Role of ECMO in Covid-19 infection**

Extracorporeal Membrane Oxygenation has been proposed in the treatment of severe pulmonary and cardiac compromise in COVID-19 [32]. ECMO is an extracorporeal respiratory assistance system, its indications concern patients with severe acute respiratory distress syndrome (ARDS) for which all other approaches have failed. There are 2 indications for ECMO: A) can serve as pure respiratory assistance, the blood is then taken from the right side at the level of the inferior vena cava and is re-injected into the upper vena cava. The blood passes through an artificial membrane, which allows complete extracorporeal respiration with oxygenation and decarboxylation of the blood. B) ECMO machine acts as an artificial oxygenator.
heart-lung, with additional cardiac support. The blood is reinjected this time, not into the jugular vein, but into the aorta [33,34]. The care under ECMO can last several weeks. For influenza ARDS patients who are managed every winter, the ECMO is used over an average of 2-3 weeks. For Covid-19 patients, some of them need 1 month of care (Figure 2).

Acro Ischemia Syndrome

The basement membrane of the skin is equal to the renal membrane, the ocular structures, etc., and it receives the same type of aggression by immunoglobulins, immunocomplexes and all inflammatory cascades [35]. Acro-ischemia syndrome can be observed in Covid-19 children and adults requiring ICU hospitalization. This syndrome includes toe and/or finger cyanosis, skin bulla and dry gangrene. D-dimer, fibrinogen, and fibrinogen degradation product (FDP) are frequently elevated, increasing progressively when Covid exacerbated (Figure 3). Most of these patients are diagnosed with disseminated intravascular coagulation (DIC), receiving anticoagulation therapy (low molecular weight heparin). The association of respiratory and cardiovascular complications leads to a fatal prognosis [36]. Therefore, the existence of hypercoagulation status in critical Covid patients should be closely monitored.
Figure 3: Acro-ischemia syndrome. Cutaneous small vessel vasculitis secondary to COVID-19 infection. Presentation signs include cyanosis on the fingers/toes, sole and heel, skin bulla and dry gangrene. In these cases, D-dimer, fibrinogen, and fibrinogen degradation product (FDP) are significantly elevated, this is related with disseminated intravascular coagulation (DIC).

Dermatological lesions are observed in the face, hands and feet, like “frostbite”, including redness at the level of the fingers, nose and ears, swelling, temporary pain which corresponds to what it is observed in winter sports. In addition to this pseudo-frostbite, other skin manifestations have been identified, such as the sudden onset of persistent, sometimes painful redness, and urticarial eruptions (Figure 4A-4D). The pathophysiology of these lesions is unclear but may include immune dysregulation, vasculitis, vessel thrombosis and neoangiogenesis [37,38].
Figure 4A-4D: Initial dermatological signs of COVID-19. Small vessel vasculitis: hives (urticaria), generalized eruptions (rashes), redness, perniosis (childblains), violaceous macules, non-necrotic purpura.

Impact on Neurological System

Neurological involvement in Covid-19 [39,40], with expression in the most serious patients, can be divided into three categories:

a) Symptoms of central neurological concern (such as headache, dizziness, syncope, consciousness alteration, ataxia)

b) Symptoms of neuro-peripheral origin (hypo-ageusia, hyposmia, neuralgia, peripheral neuropathies)
c) Symptoms of peripheral-muscular damage, often associated with the most critical stages of the disease, with liver and kidney suffering

To support the clinical suspicion of neurological involvement, it is already known that human coronaviruses can spread from the respiratory tract to the central nervous system (CNS) through transneuronal and hematogenous routes, resulting in encephalitis and other neurological complications. Moreover, has been hypothesized that Covid-19, similarly to other coronavirus can spread via a synapse-connected route to the medullary cardiorespiratory center from the mechanoreceptors and chemoreceptors in the lung and lower respiratory airways triggering the acute respiratory failure in infected patients [41]. Another important observation is that HCoV-OC43 RNA, a kind of human coronaviruses, could be detected for at least a year in the CNS of infected mice. Hence If the SARS-CoV-2 is latent in CNS for a long-time, neurological disease can manifest as possible late complications of the cured patients [42].

A wide range of neurologic manifestations of SARS-CoV-2 infection have been recognized, and evidence of their severity and persistence is increasing. Neurologic manifestations were analyzed in 509 consecutive patients admitted with confirmed Covid-19 within a hospital network (Chicago, Illinois) [43]. Neurological manifestations were present at Covid-19 onset in 215 (42%), at hospitalization in 319 (62%), and at any time during the disease course in 419 patients (82%). The most frequent neurologic manifestations were myalgias (44%), headaches (37%), encephalopathy (31%), dizziness (29%), dysgeusia (15%), and anosmia (11%). Strokes, movement disorders, motor and sensory deficits, ataxia, and seizures were uncommon (0.2 to 1.4% of patients each). Severe respiratory disease requiring mechanical ventilation occurred in 134 patients (26%). Independent risk factors for developing any neurologic manifestation were severe Covid-19 and younger age. Of all patients, 362 (71%) had a favorable functional outcome at discharge. However, encephalopathy was independently associated with worse functional outcome and higher mortality within 30 days of hospitalization. In summary, neurologic manifestations occur in most hospitalized Covid-19 patients. Encephalopathy was associated with increased morbidity and mortality, independent of respiratory disease severity.

The clinical neurological practices have, by necessity, changed dramatically during the pandemic spreading. Patients with acute, non-infectious, neurological disorders, that it is means, for almost all cases, patients with ischemic or hemorrhagic vascular events, are tending to postpone the access to the emergency care for several reasons, such as the contagion fear or the anxiety of subsequent isolation from the family unit, or to avoid a further overloading of national health service. However, the delay in access leads to an impossibility of early essential treatment with an inevitable worsening of the long-term outcome. On the other hand, the approach and needs of patients with chronic neurological diseases (e.g. epilepsy or multiple sclerosis) and of patients with neurodegenerative diseases (e.g. Alzheimer’s disease and other forms of progressive dementia, Parkinson’s disease and Amyotrophic Lateral Sclerosis) are inevitably different. In this worldwide situation, where neurological diseases do not stop only because a pandemic is ongoing, the switch to non-face-to-face care, is becoming mandatory to prevent the physical and psychological consequences that can be associated with this emergency [44]. Various telehealth technologies, in particular video teleconferencing between healthcare providers and patients, have been incorporated into the practice of neurology [45].

**Loss of Sense of Smell and Taste as Viral Infection Markers**

Early single reports and surveys have suggested that gustatory and olfactory loss may be early symptoms associated with COVID-19 infection also in the absence of other known symptoms of the disease. The first large Multicenter European Study [46] reported that in a population of 417 mild-to-moderate COVID-19 patients 85% and 88% had smell and taste dysfunctions respectively with a significant positive association between the two symptoms. Interesting the symptoms were persistent in 56% of patients over the days following the resolution of the infection general clinical manifestations. Smell and taste loss were reported in 68% and 71% of Covid-19-positive subjects in US. Anosmia, with or without dysgeusia, seems to manifest either early in the disease process or in patients with mild or no constitutional symptoms. According to the results of these studies the prevalence of olfactory and gustatory dysfunction seems lower in Asiatic than in European and American COVID-19 patients.

Post-viral anosmia is one of the leading causes of loss of sense of smell in adults, accounting for up to 40% cases of anosmia. Viruses that give rise to the common cold are well known to cause post-infectious loss, and over 200 different viruses are known to cause upper respiratory tract infections. Previously described coronaviruses are thought to account for 10-15% cases. It is therefore perhaps no surprise that the novel COVID-19 virus would also cause anosmia in infected patients. However, the traditional nasal cavity manifestations, such as nasal congestion or rhinorrhea are commonly absent in patients with COVID-19 suggesting a direct SARS-CoV-2 actions in the nervous system, especially in the olfactory pathway. Most studies suggest that the nasal cavity olfactory epithelium is the likely site of enhanced binding of SARS-CoV-2 and that multiple non-neuronal cell types present in the olfactory epithelium express two host receptors, ACE2 and TMPRSS2 proteases, that facilitate SARS-CoV-2 binding, replication, and accumulation. Hence the olfactory epithelium from the nasal cavity has been sug-
gested as the more appropriate tissue for detection of SARS-CoV-2 virus at the earliest stages. The olfactory receptor neurons may initiate rapid immune responses at early stages of the disease and viruses may first invade peripheral nerve terminals, and then gain access to the CNS via a synapse-connected route. The chance of brain infection through olfactory bulb during COVID-19 infection should be also considered. Clinical evidence of the association of COVID-19 infection and encephalopathy have been reported. Moreover, it is currently suspected that the neuroinvasive potential of SARS-CoV2 plays a key role in the respiratory failure of COVID-19 patients [47,48]. These preliminary observations suggest that research should be focused on additional aspects of SARS-CoV-2 actions in the olfactory pathway.

**Mathematical Modeling to Control Epidemic Spreading: Protection Versus Hazards (Polytechnic University of Vienna, Austria)**

Mathematical models address the need for understanding the transmission dynamics and other significant factors of the disease that would aid policymakers to make accurate decisions and reduce the rate of transmission of the disease.

The multi-disciplinary fight against the global spreading of COVID-19 has a strong applied mathematics/engineering science component. The outbreak of COVID-19 put in evidence the huge positive potential of diligent and careful use of applied mathematics. In this context, stochastic transmission models [49], also in combination with digital twin approaches, have turned out to be extremely helpful for estimating the efficiency of different approaches for well-tailored partial or full lock-down scenarios. In countries such as Singapore or Austria, characterized by comparatively low mortality of these Asian or European regions, governmental lock-down decisions were in fact strongly supported by such mathematical approaches.

The reliability of non-modified classical epidemiological models, i.e. compartmental model of the SEIR-type, has turned out as very questionable. Such models have repeatedly delivered predictions which, just a few days or weeks later, turn out to lie far away from the recorded data on new infections or new deaths. The basic reason for this situation seems to be in the actual non-identifiability of the involved model parameters [50]. This calls not only for simple model structures focusing on the key characteristics of the pandemic spreading, but also for careful modification of the compartmental model-related differential equations themselves. These mathematical models play a significant role in the understanding of COVID-19 transmission mechanisms, structures, and features. This may pave the way to new coronavirus spreading models - namely to such models which might help us to successfully fight COVID-19, while still seeing a beautiful, “simple”, mathematical structure telling us analytically where the weaknesses of the virus could be [51-53].

**Bioengineering Developments for Diagnostic and Therapy**

**Non-Invasive Ventilation (NIV) in SARS**

High Flow Nasal Cannula (HFNC) and Facemask are in development for severe acute hypoxemic respiratory failure of COVID-19 patients [54]. Instead of mechanical ventilation, a recent clinical study [55] showed that HFNC, which targeted a flow ≥ 50 L/min, improves oxygenation, reduces minute ventilation, and work of breathing. This study (performed in 379 patients) demonstrated clinical benefits, reduced patients’ intubation, and subsequent invasive mechanical ventilation. Thus, HFNC seems to be as safe as standard oxygen in COVID-19 patients (Figure 5A).

**Figure 5A:** Face Mask for non-invasive nasal ventilation in severe acute hypoxemic respiratory failure of Covid patients.

The high flow rates of HFNC and Facemask are likely to increase virus aerosolization. The Helmet device can limit virus spread into the ambient air; it is a reusable single patient interface made of a clear plastic hood on a hard-plastic ring with a silicon-polyvinyl chloride soft collar. With an additional tool, the patient’s exhalate can be filtered by applying a high efficiency particulate (HEPA) filter at the Helmet outlet [56,57] (Figure 5B).
Therapeutic Plasma Exchange (TPE)

It is clinically estimated that 13% of Covid-19 cases are severe, and 6% are critical; the evolution can be acute respiratory distress syndrome (ARDS), sepsis and/or multiorgan failure. The response to fulminant COVID-19 infection is characterized by excessive immune dysregulation (cytokine storm), inflammation, hypercoagulable state, and endothelial dysfunction. Severe COVID-19 disease has been associated with lymphopenia and high levels of ferritin, C reactive protein (CRP), lactate dehydrogenase (LDH), D-dimer, and interleukin-6 (IL-6) [4,58,59].

Therapeutic plasma exchange has been used for the management of severe infections such as 2009 H1N1 influenza A, sepsis, and multiorgan failure with a trend towards improved survival. In patients with COVID-19 pneumonia, high risk of thrombosis became a current issue, and D-dimer levels indicating fibrin degradation products (FDPs) in the plasma were found as a predictor for mortality [60,61]. Although unfractionated heparin (UFH) and low-molecular-weight heparin (LMWH) decrease the production of FDPs by inhibiting factors Xa and II, they cannot contribute to metabolization of existing FDPs. Furthermore, FDPs cannot be filtered by known cytokine filters because of their molecular weight (minimum 240 kDa). FDPs can be removed by therapeutic plasma exchange. TPE might be proposed as supportive/adjunct therapy for the management of COVID-19 with severe pneumonia, cytokine storm, and multiorgan failure [62,63].

Structures of Coronavirus Proteins by Synchrotron Radiation Diffraction (Polytechnic University of Ancona, Italy)

Some determinations of 3D structures at atomic level of proteins and complexes, useful for discovery of drugs and vaccines, were performed by using the diffraction of X-Ray Synchrotron Radiation [64]. Synchrotron light is created when centripetal acceleration is applied to an electron beam (Figure 6A). The electrons inside a heated metal are excited enough to escape from the surface in a process known as thermionic emission, then they are directed by the linear accelerator or LINAC. This fits into the Booster ring, which adopts magnetic fields to force the electrons to travel in a circle. Microwaves are used to add even more energy to the electrons. The radiation is produced in the Booster ring, having a multi-sided shape (similar an octagon). A charged particle in motion subjected to a magnetic field perpendicular to its trajectory experiences a force. If the field is constant, the particle describes a circular motion (Figure 6B).
A series of electromagnetic devices (dipole and wave) are positioned around the storage ring, causing the beam to curve or wave in a winding path. Every time the beam passes through a corner with these magnets, the electrons lose energy, which is released in the form of light. These magnets are adjustable. If the intensity of the magnetic field increases, the forces in the electron stream increase, creating tighter curves along its path. This change in the curve produces a modification in the wavelength of the emitted radiation. A tight curve produces short wavelength radiation, like X-rays. While smooth curves produce longer wavelength radiation, such as infrared (Protein crystallography at BESSY II: A mighty tool for the search of anti-viral agents).

The crystal structure of SARS-COV-2 main protease has been determined [65,66]. Such an enzyme constitutes a possible target of antiviral drugs because it is essential for processing the polyproteins which are translated from the viral DNA. Then the 3D structure at the atomic level of the main protease complexed with an α-ketoamide inhibitor was determined including positive pharmacokinetic results obtained in CD-1 mice. The structure of SARS-CoV-2 papain-like protease was also investigated to study mechanism and inhibition [67].

The technique was used at the Advanced Photon Source of the Argonne National Laboratory (Chicago, USA) to determine the crystal structure of the SARS-CoV-2 receptor binding domain (RBD) in complex with the human cell receptor, namely angiotensin-converting enzyme 2 (ACE 2). A similar experiment was performed at the Synchrotron Research Facility (Shanghai, China).

The crystal structure of baicalein-bound SARS-CoV-2 3CLpro was investigated, demonstrating the 3CLpro reversibly inhibited by a small molecule, or even a fragment of one, derived from TCM, providing an ideal lead for the design of new and versatile classes of inhibitors against 3CLpros [68].

The crystal structure of the C-terminal domain of SARS-CoV-2 (SARS-CoV-2-CTD) spike (S) protein in complex with human ACE2 (hACE2) was also investigated, which reveals a hACE2-binding mode similar overall to that observed for SARS-CoV. The results shed light on the viral pathogenesis, providing important structural information to develop therapeutic countermeasures against the virus [69].

The crystal structure of the receptor-binding domain (RBD) of the spike protein of SARS-CoV-2 bound to the cell receptor ACE2 was also investigated, identifying residues in the SARS-CoV-2 RBD that are essential for ACE2 binding and helping the future identification of cross-reactive antibodies [70].

The above reported results of the different experiments constitute a fundamental target for antiviral strategies, together with the use of neutralizing monoclonal antibodies.

Discussion

Covid-19 is a prothrombotic disease with a unique global lethality. Strong inflammatory response to viral infection severely affects the respiratory, cardiovascular, immune, and hematologic systems. The coronavirus can send the body’s immune system into cataclysmic overdrive, resulting from “cytokine storm syndrome” [71].

Early signs and symptoms associated with COVID-19 infection are skin vasculitis lesions represented by acro-ischemia syndrome of paramount importance for diagnostic in child and adults. These skin complications can be localized in toes, heel and sole of the foot, face, ears, nose, hands, fingers, they can be dangerous signs of disseminated intravascular coagulation with mortality risks [35-38]. They are mostly related with immune dysregulation, neoangiogenesis, vasculitis, and vessel thrombosis. Fast identification of acro-ischemia syndrome permits the care of disseminated intravascular coagulation complications. Furthermore, neuro-sensorial symptoms like olfactory and gustatory loss (anosmia, dysgeusia) are also initial symptoms useful for covid diagnosis and patients’ isolation [46-48]. The goal is avoiding epidemic spreading.

Clinical evolution of patients allows to consider COVID-19 as a new hematologic disease [72]. Curative anticoagulation in these patients prevents endothelial lesions. Angiopoietin-2 as a marker of endothelial activation is a good predictor factor for intensive care unit admission of COVID-19 patients [25-27]. Coronavirus antibodies can disappear after 2-3 months of disease. Assessment of antibodies levels after recovery showed that patients cured of SARS-CoV-2 are only immunized for several weeks, thus antibodies remain in patient’s serum for at least 40 days after the onset of signs. These antibodies persist active and dynamic to prevent reinfection for at least three months in 90% of cases [73]. This rate begins to decline after twenty to thirty days, most subjects tested lose 50% of active antibodies after six months. Afterwards lymphocytes develop and take in charge immunity, the so-called “memory” cells in turn would be ready to produce antibodies in the event of recontamination. Further research should clarify this subject. Exosomes could be a therapeutic option in coronavirus disease [74,71].

Mathematical models address the need for understanding the transmission dynamics and other significant factors of the disease that would aid policymakers to make accurate decisions and reduce the rate of transmission of the disease. Synchrotron radiation might play a role for diagnostic and therapy. Aftermath of Covid-19 is for the moment unknown; cardiovascular, hematological, neurological, respiratory, and immune systems are greatly altered [76-78]. Future translational research should be concentrated in this challenge.
Abbreviations

COVID : Corona Virus Disease
SARS CoV Syndrome Coronavirus
ECMO : Extra Corporeal Membrane Oxygenation
ACE : Angiotensin Converting Enzyme
RT-PCR : Real Time - Polymerase Chain Reaction
PEEP : Positive End-Expiratory Pressure
ARDS : Acute Respiratory Distress Syndrome
HFNC : High Flow Nasal Cannula
NIV : Non-Invasive Ventilation
L/min : Liter per Minute
HEPA filter : High Efficiency Particulate Air filter
CMR : Cardiac Magnetic Resonance
DVT : Deep Venous Thrombosis
MERS-CoV Syndrome Coronavirus
D-DIMER : Fibrin Degradation Product
ICU : Intensive Care Unit
DIC Coagulation
CNS : Central Nervous System
CT : Computed Tomography
LU : Lung Ultrasound
SEIR Epidemiology model : Susceptible-Exposed-Infectious-Recovered

Authors contributions

Conceived and designed the study: JCC, DM, FR. Search of information and database creation: JCC, LM, CH, BZ, MR. Analyzed the data: JCC, LM, CH, CL. Clinical and epidemiological explorations: JCC, DM, LM, CL. Contributed documentation & illustrations: JCC, LM, BZ, MR, CL. Wrote the paper: JCC LM CH DM BZ MR FR. Compiled the data: DM, CH, FR.

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