Renal, cardiac, neurological, cutaneous and coagulopathic long-term manifestations of COVID-19 after recovery; A review

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Abstract

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) caused the novel global coronavirus disease 2019 (COVID-19) disease outbreak. Its pathogenesis is mostly located in the respiratory tract. However, other organs are also affected. Hence, realising how such a complex disturbance affects patients after recovery is crucial. Regarding the significance of control of COVID-19-related complications after recovery, the current study was designed to review the cellular and molecular mechanisms linking COVID-19 to significant long-term signs including renal and cardiac complications, cutaneous and neurological manifestations, as well as blood coagulation disorders. This virus can directly influence on the cells through Angiotensin converting enzyme 2 (ACE-2) to induce cytokine storm. Acute release of Interleukin-1 (IL1), IL6 and plasminogen activator inhibitor 1 (PAI-1) to be related to elevating risk of heart failure. Also, inflammatory cytokines like IL-8 and Tumour necrosis factor-α cause the secretion of von Willebrand factor (VWF) from human endothelial cells and then VWF binds to Neutrophil extracellular traps to induce thrombosis. On the other hand, the virus can damage the blood–brain barrier by increasing its permeability and subsequently enters into the central nervous system and the systemic circulation. Furthermore, SARS-induced ACE2-deficiency decreases [des-Arg9]-bradykinin (desArg9-BK) degradation in kidneys to induce inflammation, thrombotic problems, fibrosis and necrosis. Notably, the angiotensin II-angiotensin II type 1 receptor binding causes an increase in aldosterone and mineralocorticoid receptors on the surface of dendritic cells cells, leading to recalling macrophage and monocyte into inflammatory sites of skin. In conclusion, all the pathways play a key role in the pathogenesis of these disturbances. Nevertheless, more investigations are necessary to determine more pathogenetic mechanisms of the virus.

Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) caused the novel global coronavirus disease 2019 (COVID-19) disease outbreak [1, 2]. This new virus is highly contagious as well as is a deadly virus [3]. As individuals with the virus are elevating worldwide, the complications and some disturbances caused by COVID-19 have widely increased [4]. Although, the most frequent signs of COVID-19 are respiratory symptoms [5], but, there are documents of COVID-19-correlated symptoms including blood coagulation disorders, renal and cardiac complications as well as cutaneous and neurological manifestations (Fig. 1) [3, 6–8]. As we know, Angiotensin converting enzyme 2 (ACE-2) is the receptor for coronaviruses [9] and the spike (S) protein of virus attaches it on the cell surface of some organs (e.g. cardiovascular system, kidney, lung, gastrointestinal tract (GI), skin and brain). After cleavage by transmembrane serine protease 2 (TMPRSS2), the virus enters the cell [10].

Upon viral entry, the innate immune system is recruited to fight the infection. Meanwhile, SARS-CoV-2 replicates in the pulmonary and endothelial cells of the pulmonary arteries and after destroying the cells, releases to the circulation. Macrophages and dendritic cells (DCs) endocytose the virus and present virus antigens to T helper type 1 (Th1). Further stimulation of Th1 gradually activates B cells to secrete neutralising antibodies and CD8 + T cells to destroy virus-infected cells. This stimulates the secretion of pro-inflammatory cytokines like Interlukine-6 (IL-6), IL-21, IL-18 and Tumour necrosis factor-α (TNF-α) and activates the...
nuclear factor kappa B (NF-κB) pathway, causing a systemic hyperinflammatory state that may involve other organs [11, 12]. Along with systemic hyperinflammatory, other cellular and molecular mechanisms may influence on vital organs. So, realising how such a complex disturbance affects patients after recovery is crucial. Regarding the significance of control of COVID-19-related complications after recovery, the current review was designed to study the cellular and molecular mechanisms linking COVID-19 to significant long-term signs including renal, cardiac, cutaneous and neurological complications, as well as blood coagulation disorders. To the best our knowledge, there are no comprehensive review to explore these events related to many long-term complications of COVID-19.

**Renal manifestations**

ACE2 plays a vital role in regulating the cardiovascular and renal systems via Renin-Angiotensin-Aldosterone System (RAAS). Renal juxtaglomerular cells begin to release renin in response to some conditions, including reduced renal perfusion, hypotension, ischaemia, sodium diuresis and sympathetic stimulation, which converts liver-derived angiotensinogen to angiotensin I (ANGI) [13]. Subsequently, ANGI is converted by ACE to ANGII, which in turn is further metabolised by ACE-2 into ANG I–7. Meanwhile, ANGI can also be metabolised by neutral endopeptidase to ANG I–7. ANG I–7 and its receptor, i.e. Mas, acts as vaso-dilator, anti-inflammatory and anti-coagulant agents, whereas ANGII and its receptor, i.e. Angiotensin II type 1 receptor (AT1R), are vasoconstrictor, pro-inflammatory, pro-coagulant and fibrotic effectors [14]. Thus, in patients with cardiorenal diseases, ANGII can control fibrosis, hypertrophy and prooxidative hormone, which leads to an increase in salt and water retention [15, 16]. In kidneys, ACE2 is expressed on proximal tubule epithelial cells (PTECs), podocytes and endothelial cells [17]. Although the TMPRSS2 expression is low on PTECs, high levels of other proteases such as cathepsin and glutamyl aminotransferase, in these

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**Fig. 1.** COVID-19 long-term symptoms. SAE, systemic arterial embolism; MI, myocardial infarction.
cells may help the virus entry [11]. SARS-CoV-2 moves to the kidneys through the glomerular capillaries and arterioles and then enters the glomerular epithelial cells to infect the podocytes. In this way, SARS-CoV-2 enters tubular fluid and reach PTECs [13, 18]. The internalisation of ACE2 receptor induced by SARS-CoV-2 leads to ACE2 deficiency on cells, which shifts the ANG1–7–Mas pathway to angiotensin II-angiotensin II type 1 receptor (ANGII-AT1R) and induces a proinflammatory state [11].

Furthermore, ACE2 regulates the Kallikrein Kinin System. In this regards, after producing kinin from bradykininogenins, this metabolize acts on both B1 and B2 receptors to induce inflammation and increase vascular permeability [14, 19, 20]. The most important agonist of B1 receptor is a bradykinin metabolite, i.e., [des-Arg9]-bradykinin (desArg9-BK), which is cleaved by ACE2. As regards SARS-induced ACE2-deficiency, it should said that not only it increases ANGII, but also decreases desArg9-BK degradation [14]. Also, Kallikrein can trigger the coagulation imbalance through activating factor 12 and plasmin.

The mentioned renal changes can exacerbate systemic inflammation, thrombotic problems, fibrosis and necrosis mediated by COVID-19 in kidneys, resulting in renal damages [14].

The severity of these injuries varies from proteinuria to acute kidney injury (AKI) in ~5–70% patients [11, 14, 21]. However, a recent meta-analysis on 18 043 patients showed that 32.6% of patients admitted to the intensive care unit (ICU) suffered from AKI [22], the incidence of AKI has been higher in non-survivors, ranging from 25–50% [23, 24]. Based on a systematic study on 193 patients with COVID-19, the corresponding figures for proteinuria, haematuria, urea nitrogen levels and creatinine levels were 44–65%, 27–44%, 14% and 10–14% [21]. Theoretically, Transient Receptor Potential Canonical Channel 6, as an ion channel in podocytes, can directly be activated by ANGII, which leads to proteinuria [13, 25]. The most common COVID-19-related renal complications are including acute tubular injury, collapsing glomerulopathy, thrombotic microangiopathy, complement activation and IL-6-induced renal vascular permeability [13, 17]. Therefore, the kidney injury leads to: (i) hypoxia-induced acute tubular necrosis with high level of creatinine, (ii) collapsing focal segmental glomerulosclerosis together with haematuria, which are common in patients with APOL1 alleles, (iii) direct viral tropism of renal tissue with high levels of micro- or macro-albuminuria and (iv) hemodynamic changes mediated by endothelitis along with electrolyte imbalance [26]. Furthermore, the infiltration of some immune cells like lymphocytes into the renal interstitium can cause cells to release more pro-inflammatory cytokines and lymphocyte-derived toxic particles (e.g. perforin), causing more kidney injuries [13]. It is worth noting that rhabdomyolysis and acute cardiorenal syndrome are also associated with kidney failure [26]. In addition, some drug-induced nephrotoxicity may also occur in COVID-19 cases; for example, remdesivir can be nephrotoxic through damaging the mitochondria in the epithelial cells of the renal tubules [27]. According to a systematic review, renovascular complications can be seen in COVID-19 cases with/without a pre-existing renal disorder. Hence, all COVID-19 patients should be checked for their renal functions [28].

Overall, although COVID-19-related renal histopathological changes involve both parenchyma and the interstitium of kidney, glomerular injuries are milder than interstitial ones [29, 30]. These changes have already been described in a study by Faour et al. [13]. However, some questions still remain in this area that need further study.

Cardiac manifestations

It has been shown that about 7.3% of COVID-19 patients suffer from heart pulsation as a primary sign [31]. The most prevalent COVID-19-related cardiovascular manifestations have been cardiac injury and myocardial injury, with 35.29% and 29.41% of cases respectively [32]. COVID-19-induced hypoxia and myocardial damage can create cardiac electrical dysregulation and arrhythmias, which is common in 30%–50% of ICU cases [33]. Moreover, about 20%–30% of hospitalised cases have had an increase in troponin I (cTnI) levels which is related to myocardial involvement [31, 34]. In cardiac tissues, ACE-2 is highly expressed in venous and arterial smooth muscle cells (SMC), endothelial cells and cardiac fibroblasts [35, 36]. It has been reported that COVID-19 can directly affect cardiomyocytes through their ACE-2 to induce inflammatory responses and cytokine storm, as aforementioned [35].

It has been offered that there is a connection between acute myocardial damages caused by COVID-19 infection and ACE-2. In fact, by shifting RAAS toward ANGII/AT1R, the level of inflammatory cytokines such as IL4, IL10 and IL6 are increased, which in turn activates T-cells in peripheral blood of COVID-19 patients. The overactivation of T cells leads to a rise in Th17 and cytotoxic CD8 T-cells [37]. In other words, when ANGII binds to AT1 receptor, atherosclerosis, hypertrophy, fibrosis, proliferation and vasoconstriction are increased, while, natriuresis and diuresis are decreased [38]. On the contrary, if ANGII binds directly to AT2 receptor or converts to ANGIII and then binds to AT2 receptor or converts to ANG (1–7) receptor (MasR), atherosclerosis, hypertrophy, fibrosis, proliferation and vasoconstriction are diminished and natriuresis and diuresis are elevated [38].

The myocardial damage may occur during COVID-19-associated ‘cytokine storm’ that is followed by a Th1/Th2 imbalance and can exacerbate respiratory distress syndrome, hypoxaemia, shock and/or hypertension. Moreover, in COVID-19 patients, some factors including GCSF, IL10, IL7, IL2, IP10, MCPI, MIP1A and TNFα are elevated in inflammatory response [39–42]. It has been indicated that acute release of IL1, IL6 and plasminogen activator inhibitor 1 (PAI-1) were related to elevating risk of heart failure [43]. The cytokines caused the release of reactive oxygen species, superoxide anion and endogenous nitric oxide so that all of them could injure myocardial cells [44]. C-reactive protein (CRP) is an important marker in systemic inflammatory and several infections. CRP helps in stimulation of atherosclerosis and instability of atherosclerotic plaque [43]. In addition, a high prevalence of COVID-19-mediated inflammatory stress can also cause arrhythmias through inducing electrolyte and hemodynamic disorder [35, 45, 46]. In COVID-19 patients, strong interferon-mediated responses can contribute to myocardial dysfunction, especially in protective adaptive immunity by interferons [47]. It has also been recommended that severe COVID-19 patients with higher N-terminal pro-brain natriuretic peptide (NT-proBNP) levels were older patients who had high levels of markers of systemic inflammatory. NT-proBNP levels presented as an independent risk factor of in-hospital survival rate in severe COVID-19 patients. So that, COVID-19 patients with higher NT-proBNP (above 88.64 pg/ml) levels had a lower survival rate [48].

Transforming growth factor-β1 (TGF-β1) acts as important mediator in the process of tissue fibrosis and leads to scarring by activating its downstream small mother against decapentaplegic (Smad) pathway [49]. In this direction, SARS-CoV-2 activates
TGF-β signalling by the smad pathway to stimulating lung fibrosis. In addition, smad pathway activated by COVID-19 is a prevalent pathway of interstitial fibrosis development in the myocardium [47]. In fact, TGF-β signalling in COVID-19 process is activated by both the non-canonical and the canonical signalling pathways [50]. A papain-like protein as well as a nucleocapsid protein along with smad3 activate the canonical and the non-canonical pathways, so, TGF-β is upregulated [50].

Overall, various cellular and molecular mechanisms such as recognition of ACE-2 and receptor infection, cytokine storm and induction in several inflammatory responses are contributed in cardiovascular dysfunction, which can increase pulmonary vascular resistance, as well as result in pulmonary heart disease and hypertension.

Neurological complications

Besides the acute clinical manifestation of respiratory virus, understanding the neurological complications following COVID-19 illness will be crucial [51]. A tremendous number of clinical studies have reported that many of COVID-19 patients had at least one neurological manifestation [7]. In a 2022 meta-analysis, 2791 out of 18 258 COVID-19 patients had neurological symptoms with the mortality rate of 29.1% [52]. A scoping review on neurological manifestations in COVID-19 patients has revealed that the spectrum of these symptoms ranged from mild to severe [53]. The mild manifestations included olfactory and gustatory disorders, dizziness, headache, vomiting, malaise, fatigue and anorexia. Whereas, ischemic stroke, impaired consciousness, encephalopathy, cognition and memory impairments, meningitis, diplopia and ophthalmoalgia are regarded as severe symptoms [23, 54, 55].

Neurologic complications can also be divided into central nervous system (CNS) and peripheral nervous system (PNS) involvements. The CNS involvement comprises headache and dizziness, impaired consciousness, acute cerebrovascular disease, corticospinal tract signs, ataxia and seizure, meningitis, encephalitis and stroke. While, the PNS involvement includes hypoxia and dysgeusia, vision impairment, nerve pain and Guillain-Barre syndrome [56, 57].

Interestingly, sex and age differences have been reported to affect the severity of neurological manifestations. Differences in humoral and innate immune responses to viral infections between men and women as well as age-related expression of ACE2 are supposed to be involved [58, 59]. In addition, Heart disorders, diabetes and dyslipidemia have increased the risk of developing neurological complications in COVID-19 by 2-fold in a meta-analysis on 4401 patients [60]. [59]. Moreover, some neurological symptoms like Parkinsonism can be developed in cases that recovered from COVID-19 as shown on a clinical study. Thus, Parkinsonism is a post-COVID neurological symptom [61].

However, the aetiology for COVID-19 neurological consequences is not completely understood. Post- COVID-19 neurological disorders are shown in Figure 2. From the existent studies on the neuropathogenicity of COVID-19, two possible underlying mechanisms have been proposed: (1) Direct mechanisms and (2) Indirect mechanisms [7, 53]. In direct invasion, SARS-CoV-2 could affect the nervous system mainly through the neuronal pathways and to a lesser extent through the blood circulation pathways or lymphatic system [23, 54]. In terms of neuronal pathway, the virus mainly spreads via the transcribal route from the olfactory epithelium along with the olfactory nerve to the olfactory bulb in the nasal cavity and forebrain [62]. This latter route effectively makes it a channel among the nasal epithelium, the entire brain and cerebrospinal fluid and subsequently causes inflammation and demyelinating reactions [58, 59, 62, 63]. Another potential mechanism is the neuronal retrograde or anterograde transmission. In this route, the viruses invade the sensory or motor nerve endings and migrate by the motor proteins including dynein and kinesins [64]. These mentioned pathways can lead to dysfunction of the respiratory and/or gastrointestinal centres [65].

Accumulating evidence suggests that SARS-CoV-2 gains entry to the CNS through the ACE2 and TMPRSS2 receptors on several brain structures as well as via CD147 and Neurinol-1 (NRP1) receptors in sensory neuron [7, 23, 51, 53–59, 62–66]. Based on studies, both ACE2 and NRP1 are highly expressed in the brain. Although ACE2 is mostly expressed in circumventricular organs and endothelial vasculature, NRP1 is generally expressed in the hippocampus, endothelial cells, mural cells, perivascular macrophages and microglia [67, 68]. Following the dissemination of the coronaviruses in the nervous system, it can damage the blood–brain barrier by increasing its permeability and subsequently enter into the CNS and the systemic circulation [56, 57].

In addition to direct infection injury, the nervous system may be affected indirectly due to a severe systemic reaction in response to a viral infection outside the nervous system. In this regard, neuro-inflammation, hypoxia, coagulation disorders, dysregulated blood pressure (due to binding to ACE2 receptors) as well as altered glucose and lipid metabolism are considered as indirect possible mechanisms in neurological consequences of COVID-19 [23, 53–59, 62–65].

It has been supposed that the neurotropic potential of the COVID-19 is closely related to the development of a systemic inflammatory response syndrome (SIRS) [58]. SIRS mediates the release of inflammatory cytokines (cytokine storm) and free radicals, which affect the central and peripheral nervous system. As a consequence of a cytokine storm [cytokines (e.g., IL-6, IL-1TNF-α, IL-1β, IFN-γ, IL-4, IL-10)], glial cells will be activated [53, 54]. This activation can induce a pro-inflammatory state which is positively correlated with the severity of neurological symptoms [69]. Encephalitis and encephalopathy have been related to a rise in the levels of proinflammatory cytokines and antioxi- dants based on two clinical studies [70, 71]. On the other hand, headache has less been associated with mortality since patients with headaches have experienced less cellular effectors related to cytokine storm, including D-dimer and ferritin [72].

Another detrimental effect of coronaviruses is hypoxia injury that caused by lung tissue lesions. This, in turn, leads to subsequent nervous system damage such as interstitial oedema and encephalopathy. Moreover, headache due to ischaemia, congestion and even coma can be observed [7]. Interestingly, hypoxia has been implicated in the tau hyperphosphorylation in Alzheimer’s. Furthermore, increased ACE expression in brain of Alzheimer’s patients suggests that COVID-19 may contribute the incidence of this neurodegenerative disease as well [73, 74].

A review of current published literatures has indicated that coagulation dysfunction was another indirect mechanism in case of neuroinvasive potential of SARS-CoV2. Coagulopathy, including elevated D-dimer levels, prolonged prothrombin time, decreased platelet counts, elevated PAI-1 and Von Willebrand factor (VWF) have been highlighted in severe cases [23, 53–59, 62–65]. It may render these patients prone to ischaemic strokes and other cerebrovascular events [54].

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To put these findings together, understanding the neurotropic characteristics of SARS-CoV-2 is substantial to individualise and prioritise the treatment protocols in the context of patients with COVID-19. However, more studies are needed to clarify the underlying mechanism of neuropathogenicity of the new coming virus.

**Cutaneous manifestations**

Cutaneous manifestations of COVID-19 are heterogeneous and occur in 1–20% of cases. Many studies classify them into five general categories, which are summarised in Table 1 [75, 76]. Overall, the spike protein of SARS-CoV-2 enters the skin through ACE2 receptors on the surface of epithelial cells, keratinocytes, endothelial vessels and eccrine glands that can directly damage them [77]. Upon virus entry, innate immunity is activated to prevent viral replication and plasmacytoid dendritic cells (pDCs) establish synapse with infected cells and upregulate toll-like receptor 7 (TLR7) and release the type-I interferons (IFN-I) [78, 79]. Indeed, TLRs sense pathogen-associated molecular patterns (PAMPs) such as single-stranded RNA, unmethylated double-stranded DNA (CpG), lipoproteins, flagellin and lipopolysaccharide that induce the secretion of inflammatory cytokines [80]. In chronic infections and autoimmune diseases, pDCs are constantly activated and participate in the pathogenesis through IFN-I overactivity [81]. Thus, the most skin lesions, including erythematous/vesicular/urticarial rashes, are those that are also common in other viral infections (e.g. *Herpesviridae*). Additionally, some lesions may either be exacerbated by co-infection/reactivation of SARS-CoV-2 with other viruses or by a drug-induced reaction; however, a combination of the two is also possible [75, 76, 82].

The other two types of eruptions are of vascular origin: (1) chilblain- or pernio-like and (2) ischaemic types such as livedoid racemosa, retiform purpura and acral ischaemia. A histopathological study shows the perivascular and perieccrine infiltration of T lymphocytes and pDCs in these lesions [78]. Chilblain-like is a late-onset eruption, thus PCR may be negative, that may exhibit in young people with mild and asymptomatic COVID-19 through IFN-I responses and an endothelial dysfunction. Since youngers have stronger innate immune systems; they can inhibit the virus without activating acquired immunity, resulting in negative antibody tests [83]. Nevertheless, the ischaemic lesions are related to hypercoagulability and complement deposition in elderly with severe COVID-19. In critical COVID-19, the impairment of IFN-I responses downregulates ACE2, shifting the RAAS toward ANGII. ANGII in turn stimulates T cells to increase the expression of AT1R [78, 84]. The ANGII-AT1R binding causes an increase in aldosterone and mineralocorticoid receptors on the surface of DC cells, leading to the production of pro-inflammatory cytokines (such as IL-6) and further recalling macrophage and monocyte into the inflammatory sites of skin [78]. Furthermore, in ischaemic lesions of patients with pulmonary...
thromboembolism, complement induces thrombotic microvascular injury through mannann binding lectin (MBL) pathways [85, 86]. In other words, the viral spike glycoproteins enter directly into ACE2-containing cells and through their glycan moiety interact with MBL, activating it to increase IL6, followed by platelet aggregation and complement deposition [85]. In Chilblain, however, an increase in IFN1 upregulates ACE2 and its vasoprotective mechanism via shifting the RAAS toward ANG1–7, resulting in only an impermanent acral vasospasm [78, 83]. Both vasospasm and thrombotic states can also mediate a local hypoxia in the cutaneous vasculature, which inhibit the regulatory T cells and their anti-thrombotic states can also mediate a local hypoxia in the cutaneous vasculature [87, 88].

Moreover, the humoral immunity dysregulation induced by COVID-19 can produce autoantibodies (including antiphospholipid antibodies), which may cause skin lesions through inducing an autoimmune reaction mediated by molecular mimicry, immune complex-triggered inflammation and prothrombotic state [83]. In one study, the injection of purified IgG from COVID-19 cases into thrombotic mice could accelerate thrombus formation [89]. This is probably the mechanism by which multisystem inflammatory syndrome develops cutaneous and organ damages in children after 4–6 weeks of COVID-19. More importantly, the activation of antibody-secreting cells needs several weeks, thus autoantibody-mediated lesions usually are a late-onset outcome [90].

Overall, the reviewed studies show that there is a link between skin manifestations and COVID-19, but whether SARS-CoV-2 is the cause of other underlying conditions is still unclear and needs further investigation.

**Coagulopathic disorders**

COVID-19 patients are also at emerging risk of thrombosis disorder. Two meta-analysis studies showed that the rates of the deep vein thrombosis, pulmonary embolism and arterial thrombus were 19.8%, 18.9% and 3.7%, respectively [91, 92]. Moreover, thrombogenesis was highly associated with the mortality so that 50% of non-survivors have shown a procoagulant state, while only 7% of survivors have presented this symptom [93]. Strikingly, thrombosis is a part of the innate immune reaction to pathogens that is called immunothrombosis [94, 95]. Neutrophil activation with neutrophil extracellular traps (NETOSIS), endothelial cell lesion and activation, as well as platelet activation and aggregation, altogether with coagulation protease activation, take a part in the process of immunothrombosis, particularly in lung [94, 96]. In addition, PAMPs, damage-associated molecular patterns (DAMPs), along with extracellular histones and DNA, have recently been considered in immunothrombosis [94, 97, 98]. Cell-free DNA and extracellular histones are the important parts of NETs which are secreted from dead cells, and they increase the inflammation and induce the thrombosis via damaging fibrinolysis.
raising thrombin production and stimulating platelet activation [94, 98, 99]. Based on clinical studies, the severity of COVID-19 is a key factor to increase the level of NETs in blood, tracheal aspirates and different organs of patients [100–102].

It has been offered that high fibrin degradation product levels (called D-dimer), long prothrombin time (PT), activated partial thromboplastin times, reduction in platelet counts, raised levels of fibrinogen, are commonly occurred in COVID-19 patients during thrombosis [94]. A considerable elevation of factors v, viii, and VWF has also been indicated in intense COVID-19 patients. Furthermore, high plasma levels of VWF antigen and pro-peptide are revealed in severe COVID-19 which they are indicative of endothelial stimulation as well as injury by secretion of VWF from Weibel-Plade bodies [94, 103]. In a study in 2020, Goshua et al. evaluated COVID-19 associated coagulopathy. They indicated that P-selectin and VWF antigen (markers of endothelial cell and platelet activation), as well as soluble thrombomodulin (a marker of endothelial cell activation) were significantly increased in ICU patients in comparison with non-ICU patients. In addition, they reported mortality to be significantly related to soluble thrombomodulin and VWF antigen among all patients [103].

Platelets release VWF cumulative in α-granules just till activation. ADAMTS-13 (a disintegrin and metalloproteinase with a thrombospondin type 1 motif, member 13) is expressed by hepatic stellate and endothelial cells [104, 105]. ADAMTS-13 controls the activation of VWF via excision of ultra-large VWF multimers (> 10,000 KDa) which are released from endothelial cells into active high molecular weight multimers (< 10,000 KDa) under stress states [106]. Intense deficiency of ADAMTS-13 results in accumulation of ultra-large VWF multimers and causes microvascular thrombosis, thrombocytopenia and thrombotic thrombocytopenic purpura (TTP) [107]. A mechanism that notably contributes in ADAMTS-13 deficiency is linked to the antiphospholipid antibody production within SARS-CoV-2 infection [108–111]. Antiphospholipid antibodies have been incompatibility announced in all patients of COVID-19 [108, 110, 111].

Despite, the prolongation of aPTT [111], the patients with antiphospholipid syndrome have unusual ADAMTS-13 plasmatic activity, so that, an increased risk of thrombosis is seen [112]. Antiphospholipid antibodies during active SARS-CoV-2 infection maybe attach the spacer domain of ADAMTS-13 which can intervene in the identification and proteolysis of VWF [113]. In this regard, studies reported that COVID-19 nonsurvivors have shown higher VWF and lower ADAMTS13 compared with survivors [114, 115].

Overall it can be seen that TLRs-3 and −7 are activated by the viral RNA to induce the NF-κB Pathway and the interferon regulatory factors, that they raise the production and release of pro-inflammatory cytokines [94]. This pro-inflammatory condition has been seen in several viral diseases such as influenza and SARS-CoV-2 [113]. The elevation of these pro-inflammatory cytokines such as IL-1, IL-6 and TNFα can promote thrombosis [103]. So that, inflammatory cytokines (IL-8 and TNF-α) cause the secretion of VWF from human endothelial cells and then VWF binds to NETs to induce thrombosis [113]. Moreover, proinflammatory situation is influential on haemostasis by blocking of fibrinolysis [113]. Also increased levels of myeloperoxidase as a NET-linked marker have been demonstrated in COVID-19 patients [94]. More importantly, the interaction among complement activation, inflammation and the coagulation cascade is seen to be crucial to understanding the COVID-19 pathophysiology and is involved to triggering disseminated intravascular coagulation [116]. Membrane attack complex (C5b-9) and C3a are both responsible for platelet activation. Moreover, C5a increases cellular and plasma TF expression [117]. Moreover, it has been established that considerable hypoxaemia in COVID-19 patients is caused in prothrombotic conditions through stimulation of endothelial synthesis of procoagulants, upregulation of plasminogen activator inhibitor and releasing tissue factor and VWF as well [113].

On the other hand, a large number of patients with nonvalvular atrial fibrillation who use direct oral anticoagulants (DOACs) (to prevent systemic embolism and stroke) may be treated with antiviral drugs (lupinavird/ritonavir, varonadin), cloquine or hydroxychloroquine, antibiotics and tocilizumab for the treatment of severe respiratory syndrome caused by COVID-19 [118]. Since both DOACs and antiviral drugs are substrates of cytochrome P450-based metabolic pathways; the simultaneous use of these two drugs can greatly elevate the plasma level of DOACs and lead to an increased risk of uncontrolled bleeding and thrombotic complications [118, 119].

In conclusion, hypercoagulability, as a noticeable feature of COVID-19, can cause thrombotic vascular events. Several mechanisms may be involved in the occurrence of the thrombosis, such as inflammatory storm, renin angiotensin system dysregulation and uncontrolled inflammation-mediated endothelial injury. However, to get a better understanding of COVID-19, more studies are needed.

Conclusions

Regarding the significance of control of COVID-19-related complications after recovery, the current study has summarised to review the cellular and molecular mechanisms linking COVID-19 to some long-term symptoms including renal, cardiac, cutaneous, neurological and coagulopathic complications. As mentioned earlier, this virus can directly influence on the cells through ACE-2 to induce cytokine storm to increase the risk of heart failure and thrombosis. On the other hand, the virus can damage the blood–brain barrier by increasing its permeability and subsequently enters into the CNS and the systemic circulation. Furthermore, SARS-induced ACE2-deficiency decreases desArg9-BK degradation in kidneys to induce inflammation, thrombotic problems, fibrosis and necrosis. Notably, the ANGII-AT1R binding causes an increase in aldosterone and mineralocorticoid receptors on the surface of DC cells, leading to recalling macrophage and monocyte into inflammatory sites of skin. All the pathways play a key role in the pathogenesis of these disturbances and physicians should be aware in their attention to all these complications and take precision steps for control, screening and even cure. Nevertheless, more investigations are necessary to determine more pathogenetic mechanisms of the virus.

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