SARS-CoV-2 interacts with renin-angiotensin system: impact on the central nervous system in elderly patients

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Abstract  SARS-CoV-2 is a recently identified coronavirus that causes the current pandemic disease known as COVID-19. SARS-CoV-2 uses angiotensin-converting enzyme 2 (ACE2) as a receptor, suggesting that the initial steps of SARS-CoV-2 infection may have an impact on the renin-angiotensin system (RAS). Several processes are influenced by RAS in the brain. The neurological symptoms observed in COVID-19 patients, including reduced olfaction, meningitis, ischemic stroke, cerebral thrombosis, and delirium, could be associated with RAS imbalance. In this review, we focus on the potential role of disturbances in the RAS as a cause for central nervous system sequelae of SARS-CoV-2 infection in elderly patients.

Keywords  COVID-19 · SARS-CoV-2 · Central nervous system · Renin-angiotensin system · Elderly

Introduction  Coronaviruses are positive single-strand RNA viruses, 80–220 nm in size, pleomorphic although often spherical, and enveloped with crown-shaped glycoprotein spikes, that comprise three genera: alphacoronavirus, betacoronavirus, and gammacoronavirus [1]. SARS-CoV-2, a betacoronavirus, has caused a global pandemic that started in 2019 [2]. While the pathogenesis of SARS-CoV-2 remains under study, common findings with SARS-CoV-1 and MERS-CoV pathogenesis may offer insights into SARS-CoV-2 pathogenesis. Most coronaviruses are largely associated with respiratory infections. SARS-CoV-2 infection results in a series of symptoms comprising fever, pulmonary insufficiency, dry cough, myalgia, headache, and intestinal dysfunction [3]. The complications and loss of function through the affected organs are particularly exacerbated in patients with co-morbidities [4]. Reports of a wide range of neurologic symptoms including stroke [5–7], viral presence in the cerebrospinal fluid (CSF) [8], and brain tissues from autopsies [9–11] introduced a neuroinvasiveness potential of SARS-CoV-2. It is increasingly evident that SARS-CoV-2 is not only neurotropic but also associated with a much broader spectrum of acute and atypical neurological syndromes and manifestations than prior infections, particularly those involving β-coronaviruses [12–17].
The genome of SARS-CoV-2 is 79.5% similar to previous SARS-CoV [18–20]. The most important structural proteins are the spike (S), envelope (E), matrix (M), and nucleocapsid (N) [19, 21]. Despite similarities in their genomes, a remarkable difference is the longer length of the S glycoprotein present in SARS-CoV-2 as compared to other coronaviruses [21]. It has been hypothesized that the higher transmissibility of SARS-CoV-2 is due to this difference in the S protein [21].

Angiotensin-converting enzyme 2 (ACE2) serves as a receptor for SARS-CoV-2 entry into susceptible cells of multiple organs [22]. The ubiquitous expression of ACE2 in multiple cell types allows SARS-CoV-2 to infect different organs including the nasopharynx, lungs, lymph nodes, small intestine, stomach, spleen, kidney, and brain leading to multiple organ damage [23]. The trimeric S glycoprotein interacts with the human ACE2 to allow for viral entry into host cells by viral membrane fusion. The affinity of the interaction between SARS-CoV-2 S glycoprotein and ACE2 is 10 to 20-fold higher than SARS-CoV, and this increased affinity is critical for the neuroinvasiveness of SARS-CoV-2 [19, 24, 25]. Viral fusion is helped by prior proteolytic cleavage of S by the transmembrane protease serine 2 (TMPRSS2), or by cathepsin B and L [22]. Hence, TMPRSS2 mediates spike protein activation and promotes SARS-CoV-2 entry via direct fusion, thereby subverting entry through endocytosis [26]. Moreover, TMPRSS2 or TMPRSS4 can generate circulating ACE2 by cleavage of the membrane-bound protein in vascular endothelial cells. Once ACE2 is shed, SARS-CoV-2 entry may occur by TMPRSS2 and TMPRSS4-mediated endocytosis [27].

Furin is a pro-protein convertase present in multiple tissues, including the brain. At the intracellular level, furin cleaves the viral S protein to a mature form, thus reinforcing its receptor binding and membrane fusion capabilities, therefore contributing to the multi-organ involvement particularly where ACE2 expression level is low [28].

Several other molecules have been reported as SARS-CoV-2 cellular receptors and proteases that mediate viral entry [29, 30]. For epithelial cells, these proteases may assist viral entry through non-ACE2-mediated routes such as a pathway involving the CD147-spike protein and CD26, which is expressed ubiquitously [31, 32].

ACE2 is a key component of the renin-angiotensin system (RAS), which plays an essential role in the homeostatic regulation of blood pressure, electrolyte, and fluid balance, as well as in the regulation of vital organ function by the renal and cardiovascular system [33].

The SARS-CoV-2-ACE2 interaction links viral pathogenesis to the function of the renin-angiotensin system (RAS). In the brain, there are a circulating RAS and a local one. The former exerts its effect in circumventricular organs that lack the blood–brain barrier (BBB) and project to nuclei in the hypothalamus and medulla. The brain RAS synthesize de novo all components, but independently of the circulatory RAS [34].

The brain RAS may actively participate in the modulation of neurotransmitter release [35–37], and therefore, it is thought to control blood pressure and regulate metabolism [38, 39]. Besides its role in normal organ development and function, RAS appears to be involved in age-associated organ dysfunction by promoting pathophysiological processes of various age-related disorders. These include heart failure and other cardiovascular diseases, diabetes, cancer, chronic kidney disease, osteoporosis, and dementia [40]. In line with these observations, Benigni et al. reported that normative aging may be delayed by inhibiting RAS [41], probably through the decrease of oxidative stress and upregulation of prosurvival genes [42].

Thus, SARS-CoV-2 infection and age-associated dysregulation of the RAS may contribute to adverse clinical outcomes. In this review, we focus on the potential role of disturbances in RAS as a cause for central nervous system sequelae of SARS-CoV-2 infection in elderly patients.
may impair multiple systemic cellular, tissue, and organ regulatory mechanisms, resulting in the varied symptomatology of COVID-19.

It appears that the brain may constitute a replicative niche for SARS-CoV-2 [44]. Examination of brain tissues from post-mortem SARS-CoV-2-infected patients revealed the presence of virus in brain capillaries, endothelial cells, pericytes, and neurons [9, 45, 46]. SARS-CoV-2 was also found in areas of the cardiorespiratory center and medulla, suggesting that infection of these brain areas may lead to or contribute to respiratory failure in COVID-19 patients [47]. Moreover, SARS-CoV-2 was detected in the cerebrospinal fluid (CSF) of patients with acute neurological symptoms, like seizures or encephalitis in conjunction with magnetic resonance image findings on the condition [8, 48]. Furthermore, viral antigens were detected in the CSF of COVID-19 patients [8, 49]. Analysis of CSF also revealed the presence of SARS-CoV-2-specific antibodies, which hypothesized that auto-antigenicity may underlie post-infectious autoimmune demyelinated pathology of the brain in COVID-19 patients [50, 51].

Infection of the CNS by SARS-CoV-2 is possible due to ACE2 expression in neurovasculature, choroid plexus, ventricles, and substantia nigra, as well as in astrocytes, oligodendrocytes, and neurons, but not in microglia [52]. Nevertheless, ACE2 expression is relatively high in some neurovascular unit components, particularly in brain pericytes; these cells are derived from neural crest stem cells and are physically linking endothelial and astrocytic cells, thus promoting its maturation and production of basement membrane components [53]. Using cortical organoids as a model, pericyte-like cells (PLCs) are permissive of infection with authentic SARS-CoV-2 and have been proposed to serve as viral “replication hubs,” able to spread the virus to astrocytes and mediating inflammatory type I interferon transcriptional responses [54].

The structural changes in cerebral small vessels of patients with COVID-19 and consequent neurological symptoms have been associated with direct viral damage of infected brain endothelial cells mediated by SARS-CoV-2’s main protease [55]. Indeed, virus progeny release into the CNS has been associated with intracranial hypertension and edema that further contribute to increasing its neuroinvasiveness [56, 57].

Studies performed in vitro using human brain organoids and neurons suggest that SARS-CoV-2 infection promotes cell proliferation, metabolic processes, and organelle fission, suggesting that the brain is a site of replication for SARS-CoV-2 through a mechanism that involves reduction of interferon-driven gene activation. The observed reduction in interferon-mediated responses following SARS-CoV-2 infection may also result from signaling emanating by infected neurons that promote the death of neighboring cells [44].

Infection of the central nervous system by SARS-CoV-2 could be achieved by several routes: (i) the hematogenous route followed by a breakdown of the blood–brain barrier (BBB); (ii) through the blood-cerebrospinal fluid barrier (BCSFB); (iii) following retrograde axonal transport of SARS-CoV-2 virions and trans-synaptic viral spreading; and (iv) through entry to circumventricular organs.

Early during infection, SARS-CoV-2 enters the bloodstream after primary infection of type II alveolar epithelial cells in the airway and, to a lesser extent, after infection of enterocytes at the gastrointestinal tract. Both sites that support the initial phase of viral replication are characterized by the high expression levels of ACE2. Later during infection, bloodstream invasion by the virus may increase when endothelial cells of the BBB or BCSFB are infected and disrupted, allowing for paracellular transmigration of virions. This route of viral propagation involves intercellular adhesion molecule 1 (ICAM-1)-mediated transport that is upregulated by tumor necrosis factor-alpha (TNF-α) and matrix metalloproteinases activation, which promote destabilization or disruption of tight junctions of the BBB leading to BBB leakage [9, 23, 47, 58–62]. Potent activation of the immune system known as “cytokine storm” is a state characterized by prominent overproduction and release of numerous active soluble components, such as interferons –IFN-, chemokines, interleukins –IL-, and TNF-α. The discharge of excessive amounts of pro-inflammatory cytokines (i.e., IFN-I, IFN-II, IL-1β, IL-6, IL-12, IL-18, IL-33, TNF-α, TGF-β) and chemokines (i.e., CXCL-8, 10, CCL-2, CCL-3, and CCL-5) is responsible for an abnormal systemic inflammatory response, which afterward causes acute respiratory distress syndrome and organ failure [63], including an exacerbation of BBB breakdown and CNS dissemination [64–67].
Another route that SARS-CoV-2 may use to access CNS may be via the brain lymphatic drainage system. The presence of viral nucleotide sequences in macrophages and T lymphocytes distributed at the periphery of germinal centers, lymph nodes, and in peripheral blood suggests that SARS-CoV-2 could use a “Trojan horse”-type mechanism by infecting T lymphocytes, macrophages, and monocytes in the blood. Viremia in the circulation may be exacerbated by productive infection of peripheral endothelial cells [68–71].

Like other human coronaviruses, SARS-CoV-2 infection may occur via the BCSFB. The CSF circulation involves a local fluid exchange between blood, interstitial fluid, and CSF that occurs normally by directional and pulsatile flow through the brain [72]. Cells at the BCSFB may activate the expression of transcriptional factors (i.e., NF-kB) and metalloproteinases that can promote BCSFB permeability and immune cell trafficking, potentially leading to a neuroinflammatory environment [73].

SARS-CoV-2 may also reach the CNS through retrograde axonal transport and transneuronal spread from different nerves, allowing the virus to infect the brainstem and disseminate in the forebrain through neuroanatomically interconnected pathways [74]. SARS-CoV-2 spread through trans-synaptic transfer may involve exocytosis/endocytosis mechanisms or rapid axonal transport, which would move the virus along microtubules to the neuronal soma [75, 76]. The transneuronal pathway is one of the potential routes that would allow SARS-CoV-2 to enter the CNS through primary sensory neurons. In this scenario, SARS-CoV-2 could enter through the olfactory mucosa (causing anosmia), spread through the olfactory nerve, and reach the olfactory cortex [77, 78]. Likewise, the virus could also enter the CNS through neurons innervating exocrine tissues such as salivary and lacrimal glands, spread through the facial and glossopharyngeal nerves, and reach various brain stem nuclei. The SARS-CoV-2 infection could also spread from gustatory cells on taste buds (producing ageusia) and move retrogradely through nerves that end in the nucleus tractus solitarius of the brainstem [79]. Lastly, SARS-CoV-2 could also reach the brainstem through the vagus nerve through infection of terminals in the respiratory tract, allowing the virus to spread to other organs innervated by vagal terminals [80–82]. The nucleus of the solitary tract and the dorsal motor nucleus of the vagus nerve express ACE2 [83], but after SARS-CoV-2 infection, ACE2 expression decreases forcing toward ACE-Ang-II-AT1 receptor axis, dysregulation of anti-inflammatory response leading to a systemic inflammatory response that results in the elevation of pro-inflammatory cytokines, chemokines, acute phase proteins, complement, and modification of leukocyte profiles in blood with consequent disruption of BBB, and microglial activation which results in increased vascular permeability [84–86] (Fig. 1).

Finally, SARS-CoV-2 might enter the CNS through the highly vascularized circumventricular organs that express ACE2 such as the subfornical organ, or through the paraventricular nucleus of the hypothalamus, the nucleus of the solitary tract, and the rostral ventrolateral medulla that also express ACE2 [87]. These areas are expected to be permissive to SARS-CoV-2 infection and thus could undergo neurovascular damage as a consequence of infection as discussed previously [88, 89]. The circumventricular organs and the areas of the hypothalamus and brainstem proximal to the third and fourth ventricles lack BBB and allow communication between the blood, the CSF, and the brain parenchyma [90].

**Renin-angiotensin system (RAS) in the CNS during SARS-CoV-2 infection**

RAS is a central mechanism of regulation of blood pressure that involves multiple organs including the brain, where its action impacts cerebral vasodilation, neuroprotection, and cognition [91]. RAS initiates in the kidney with the synthesis of the protease renin. This enzyme produces angiotensin I (Ang-I) by cleavage of its substrate angiotensinogen produced in the liver. Ang-I is not biologically active and is transformed in Ang-II mainly by the action of the angiotensin-converting enzyme (ACE). Ang (1–7) is subsequently produced by the action of angiotensin-converting enzyme 2 (ACE2) on Ang-II. RAS function is dependent on the balance of two opposing pathways: One involves ACE and its product Ang-II that signals through seven transmembrane-spanning G protein-coupled AT1 or AT2 receptors (AT1R and AT2R, respectively) based on their selective affinity for peptide and non-peptide ligands. Ang-II mediates biological functions of the RAS system and controls
physiological responses in the renal system. The alternative pathway involves ACE2, which mediates the hydrolysis of Ang-II into Ang (1–7), which binds to the Mas receptor (MasR with the highest affinity. The first pathway results in vasoconstriction and is pro-inflammatory and pro-thrombotic, and fibrotic; the second pathway is vasorelaxant, natriuretic, anti-thrombotic, anti-inflammatory, and anti-fibrotic through the actions of Ang (1–7) [92, 93].

Interestingly, an alternative renin isoform known as renin-b has been reported, supporting the existence of an intracrine RAS [94]. Renin-b is an alternative renin isoform transcribed in the brain but not present in other tissues [95, 96]. Catalytically active renin-b lacks a signal peptide and, as an intracellular form, it has been proposed to regulate brain RAS rather than generating intracellular Ang-II [97]. Whether Ang-II can be generated as a result of renin-b activity in presynaptic neurons to be subsequently released in presynaptic terminals upon depolarization is still unclear [97]. If this were the case, however, this pathway would involve the migration of virions involving ICAM-1-mediated transport that is upregulated by TNF-α and MMPs activation, which promote destabilization or disruption of tight junctions of the BBB leading to BBB leakage that is exacerbated by the cytokine storm. Additionally, the presence of viral nucleotide sequences in macrophages and T lymphocytes distributed at the periphery of germinal centers, lymph nodes, and in peripheral blood suggests that SARS-CoV-2 could use a “Trojan horse”-type mechanism by infecting T cells, macrophages, and monocytes in the blood.
potential role of Ang-II as a neurotransmitter or a neuromodulator [98].

The role of the RAS and especially the implications of ACE2 activity in the brain have been explored only recently. However, the recently demonstrated influence of the Ang (1–7)/Mas pathway in neuronal plasticity suggests a role of ACE2 in CNS homeostasis [99].

SARS-CoV-2 entry into the brain activates microglia and induces astroglisis, increasing the secretion of proinflammatory cytokines (TNF-α, IL-6, and IL-1β) and prostaglandin E2, leading to chronic inflammation, neural hyperexcitability, and exacerbated neuron programmed cell death [84, 100].

Levels of Ang-II are increased in SARS-CoV-2 infected patients. Increased Ang-II may lead to exacerbated inflammatory responses observed in COVID-19 since Ang-II potently activates NF-κB in different cell types [101, 102]. An increase in activity of the Ang-II pathway would be more harmful in older patients because of its exacerbated inflammatory component [103]. Thus, activation of the Ang-II pathway may contribute to increased risk of mortality in older individuals with COVID-19 by exacerbating the impact of dysregulation of the immune system, which leads to hyper-inflammation [103]. Additionally, in patients that recover, peripheral inflammation during SARS-CoV-2 infection could have long-term consequences leading to CNS disorders such as neurodegenerative disease and dementia [104].

Ang (1–7) can induce endothelial and neuronal nitric oxide (NO) synthase (eNOS) activity through Ang (1–7)-Mas, and bradykinin-NO [105] signaling, increasing NO production [106], which promotes vasodilatation and apoptosis reduction [107, 108]. Ang (1–7)/Mas receptor signaling also inhibits inducible NO synthase (iNOS) in glial cells and neurons, thus reducing tissue damage by peroxynitrite [109, 110]. In the brain, NO release is required for neural processes including locomotor activity, memory, and learning [111]. Under physiological conditions, NO activates Akt and cyclic AMP-responsive-element-binding protein (CREB), which are involved in survival pathways. Excess NO, on the other hand, leads to the formation of reactive nitrogen species which cause cell damage [112]. NO production is strongly associated with the activity of RAS, and in patients with COVID-19, some neurological signs could be associated with diminished physiological NO levels in the CNS [113].

Numerous reports have shown that the brain ACE2-Ang-(1–7)-MasR axis acted as a pivotal regulator of blood pressure, counteracting the pressor effect of ACE-Ang-II-AT1R [114]. Increased levels of oxidative stress and proinflammatory cytokines prompted to favor and maintain hypertension through activating redox signaling in the blood pressure regulatory centers. In this sense, in vitro and in vivo studies have revealed that inhibition of oxidative stress and inflammation may represent part of the underlying mechanisms for the antihypertensive effects of ACE2-Ang-(1–7)-MasR axis [115, 116] (Fig. 2).

In the bloodstream, the level of soluble ACE2 is normally lower than 17 mU/L [117, 118] but becomes elevated in different cardiovascular disorders [117–120]. Probably promoted by the SARS-CoV-2 infection, it is also raised among patients with severe COVID-19 [121], which could be more pronounced among older patients [86, 119, 122–124]. However, contradictory results have been reported regarding circulating ACE2 activity in SARS-CoV-2 infected patients that is ranging from elevated [125–131] to unchanged [132, 133] or even lowered circulating ACE2 levels [134]. Particularly in the brain, overactive RAS is associated with decreased cellular ACE2 level and activity during the development of neurogenic hypertension. Among patients, increased shedding of ACE2 takes place as indicated by its increased activity in their CSF samples. Such phenomenon involves the ADAM17, a member of the “A Disintegrin And Metalloproteases” (ADAM) family known to cleave a variety of membrane-anchored proteins, including ACE2. In the central nervous system, the AT1R on neurons mediates the ADAM17 upregulation upon brain RAS over-activation, leading to increased shedding of ACE2 and favoring the development of neurogenic hypertension and compromising the compensatory effect of Ang (1–7) [135]. However, the level of circulating ACE2 in the CSF in patients hospitalized with COVID-19 remains unknown.

Previous evidence obtained from SARS-CoV-1 patients indicates that SARS-CoV-2 infection can down-modulate ACE2 expression [136–139]. Considering that ACE2 mediates the Ang (1–7) production, it is expectable that SARS-CoV-2-mediated ACE2 downregulation will reduce Ang (1–7) production.
There is cumulative evidence that the ACE2-Ang (1–7)-MasR axis in the brain exerts mainly beneficial effects against hypertension [114], atherosclerosis [140] and antithrombotic activity [141]. The Mas receptor concentration is high not only in the brain structures associated with memory and cognition such as the hippocampus but also in the piriform cortex involved in smell being observed in neurons, astrocytes, and endothelial cells of cerebral resistance vessels [142–145]. As an endogenous constituent of the brain, Ang (1–7) is detected in the hypothalamus, medulla oblongata, and amygdale [146]. Consequently, when the ACE2 expression is decreasing after its SARS-CoV-2 interaction, a deleterious effect on the ACE2-Ang (1–7)-MasR axis becomes prominent with an impaired endothelial function in cerebral arteries and oxidative damage [147], difficulties in learning and memory [145, 148, 149], and reduced antioxidant and anti-inflammatory actions [146, 150, 151] [152].

RAS also regulates stress and anxiety responses. Transgenic mice with increased ACE2 expression display behaviors consistent with reduced anxiety levels [153]. This outcome was reversed when mice were treated with a Mas receptor antagonist, suggesting the Ang (1–7)/Mas axis in the regulation of anxiety. In addition, overexpression of ACE2 in mice reduces levels of cortisol and proopiomelanocortin in plasma, indicating that ACE2 may modulate basal anxiety levels through actions on the hypothalamic–pituitary–adrenal (HPA) axis [154–157]. SARS-CoV-2-mediated shedding of ACE2 may favor anxiety as well as depressive symptoms. Expectedly, disturbance of the HPA and the subsequent hypocortisolism seen in SARS-CoV-1 patients has been associated with increased anxiety, depression, and post-traumatic stress disorder [158]. In the RAS pathway, a stress response system similar to the HPA axis is triggered by Ang-II — as a stress hormone — binding to AT receptors located in the HPA axis, hippocampus, and prefrontal cortex [159]. Such HPA response to stress is desensitized using ACE inhibitors [160].

An increasingly large body of evidence suggests that the ACE2/Ang (1–7)/Mas receptor pathway results in the generation of protective mediators in numerous neuropsychiatric pathologies and stress disorders [110, 145, 154, 161]. The anti-inflammatory and antithrombotic effect of the activity of the ACE2/
Ang (1–7)/Mas receptor pathway in the CNS reduces both oxidative stress and apoptosis [162, 163]. Experiments in mice revealed that the administration of Ang (1–7) reduced not only the level of Ang-II but also hormones associated with stress response such as norepinephrine, dopamine, and serotonin (87).

**Age-related changes in the brain RAS**

RAS is imbalanced during aging due to changes in levels of its components. Changes in RAS during the aging result in abnormal levels of inflammation, oxidative stress, and cell death that promote chronic age-related disorders. Several studies have shown that Ang-II mediates premature senescence [164, 165]. Benigni et al. have shown increased longevity by downregulation of AT1R, which attenuated oxidative stress and promoted expression of pro-survival genes [41, 166].

Age-dependent variations in RAS components are differently controlled in circulating and brain RAS [167]. The mitochondrial and nuclear AT1R levels increased significantly with age [168, 169] as opposed to decreased AT2R expression [170] which achieves its maximal expression in developing fetal tissues but decreases later reaching lower levels in adulthood [171]. Similarly, age-related alterations in the distribution of RAS have been documented in the brain. A progressive decrease in the expression of AT2Rs and mRNA/protein expression of other protective RAS receptors accompanied by elevated Ang-II and AT1R levels with aging was shown in the substantia nigra [172]. Similarly, protective Ang (1–7)/MasR axis expression may be decreased in the brain of aged rats [143, 173] (Fig. 3).

Mitochondrial dysfunction has a critical role in cellular aging [174]. A functional mitochondrial angiotensin system has been identified that exhibits distributional changes in their RAS receptor levels with aging such as a decrease in AT2R and an increase in AT1R density [168]. Mitochondria in human skeletal muscle cells and monocytes as well as mouse cardiac myocytes, renal tubular cells, neuronal cells, vascular endothelial cells, and hepatocytes express high levels of AT1R with aging, which increases mitochondrial ROS levels, leading to diminished mitochondrial integrity, lowered ATP generation, and further overproduction of ROS, a prominent molecular mediator of aging [175]. Increased ROS levels lead to oxidation of mitochondrial macromolecules and DNA damage, both linked to cellular senescence and apoptosis [175, 176]. Excessive ROS production promotes the uncoupling of endothelial NO synthase, which in turn reduces NO availability and enhances ROS production. Under physiological conditions, the Ang-II capability to propitiate oxidative stress is firmly regulated. At variance, uncontrolled Ang-II-dependent ROS generation takes place as a consequence of age-associated activation of RAS. This increased Ang-II level is associated with cellular senescence depicted by cardinal markers such as telomere shortening and cell cycle arrest, which are reversed by losartan [177]. Consistent with these observations, over-activation of the Ang-II/AT1R/NADPH-oxidase complex (NOX) axis in the brain results in increased oxidative stress and cellular dysfunction. Oxidative imbalance in this system is associated with diminished levels of sirtuin-3 (SIRT3) which suppress the pro-oxidative RAS axis [178]. Additionally, the uncoupling of endothelial NOS by superoxide anions may occur, thus decreasing NO availability and NOS activity with aging [179].

Upregulated AT1R but reduced AT2R expression impairs the counterbalance mechanism of the RAS in the aging brain while oxidative stress, neuroinflammation, and increased neuron vulnerability continue [180, 181]. It has been suggested that RAS imbalance leads to age-related pro-inflammatory changes known as “inflammaging,” observed in several tissues during aging [180].

Vascular cognitive impairment is a consequence of cerebrovascular disease characterized by brain dysfunction and cognitive loss [182]. Aging and hypertension are major risk factors for vascular cognitive impairment. Aging and hypertension are associated with poor blood flow with consequent hypoperfusion and hypoxia. These conditions generate a pro-oxidative and pro-inflammatory milieu in the brain, which may promote neuronal death, thus contributing to cognitive impairment [183]. In this scenario, AT1R over-activity stimulates vasoconstriction and increases oxidative stress, inflammation, and susceptibility to ischemia. On the other hand, ACE2 overexpression and activity of the Ang (1–7)/MasR axis of RAS counteract the impact of AT1R activity, reducing both inducible NOS and the production of
pro-inflammatory cytokines in the brain and promoting angiogenesis [184] (Fig. 3).

CNS and post-acute sequelae of SARS-CoV-2 infection

“Long COVID” syndrome includes multiple post-acute sequelae of SARS-CoV-2 infection [185]. Numerous patients manifest prolonged multisystem compromise with significant disability [186]. After 6 months of SARS-CoV-2 infection, 76% of patients have at least one of the following symptoms: fatigue/muscle weakness, difficulty sleeping, hair loss, anosmia, and mobility difficulties [187].

Neuropsychiatry and movement abnormalities are reported post-infection [188–190]. Among these, the lack of movement coordination, falls, gait shuffling, and confusion was detected in individuals testing negative for SARS-CoV-2 for several days [5, 191]. Consistent with these observed abnormalities, demyelinating lesions of the CNS driven by inflammation arising from activation of glial cells are also a sequela of COVID-19 [192, 193].

An inflammatory mechanism is considered the most plausible association between SARS-CoV-2 brain infection and neurological dysfunction. Encephalitis as the central neuroinflammatory sequelae SARS-CoV-2, especially in older adults, has been documented in neuroimaging, electrophysiological, and laboratory studies [50, 194].

Fig. 3 RAS starts with the synthesis of protease renin in the kidney, which produces Ang-I from angiotensinogen in the liver. Ang-I is transformed in Ang-II by the action of ACE. Secondly, Ang (1–7) is produced by the action of ACE2 on Ang-II. RAS function is dependent on the balance of two opposing pathways. One formed by Ang-II, and the AT1R (inflammatory way), and the other comprises Ang (1–7) that intervenes in its actions by binding to Mas receptor and Ang-II/AT2R (anti-inflammatory way). The aging process unbalances the RAS activation promoting abnormal levels of inflammation, oxidative stress with concomitant cell death. AT1R and Ang-II levels are upregulated during aging. In contrast to the counterbalanced components, AT2R and Ang (1–7)/Mas axis expression appeared to be decreed. This process could be exacerbated during SARS-CoV-2 infection and explain the accelerated neurodegenerative manifestation observed in aging phenotypes of COVID-19 patients.
Although pathophysiological mechanisms of SARS-CoV-2 infection are still largely unknown, the neurological sequelae observed after SARS-CoV-2 infection may suggest infection-driven damage to glial cells and neurons that express ACE2. Furthermore, the ACE2 substrate (Ang-II) is synthesized by ACE activity, which might be inhibited by endogenous serum albumin present in the blood and potentially many tissues, but not present in the CNS [195], and it is also tightly coordinated by genotype-dependent expression and secretion mechanisms [196]. Consequently, in the CNS location, a high ACE activity, and consequently of Ang-II level, is expectable, thus highlighting the role of ACE2 as its cellular down expression would result in a more severe dysregulation of local RAS.

Sequelae may also be due to the disruption of bidirectional interactions between the immune and nervous systems, which could lead to a pro-inflammatory, hypercoagulable, and hypoxemic state in the brain. Such an environment is expected to drive neurological syndromes including demyelinating and movement disorders, degenerative dementias, encephalopathies, and also neuropsychiatric and unusual cognitive disorders [197]. Consistent with this hypothesis, increased coagulopathy, vasculopathy, neuroinflammation, and immune dysregulation are expected to underlie the increase in the risk of stroke during the COVID-19 [198].

Damage to endothelial cells arising from direct SARS-CoV-2 infection likely drives COVID-19-associated coagulopathy that underlies specific forms of neurological dysfunction. Damaged endothelial cells produce excess nitric oxide, promoting platelet and leukocyte adhesion, followed by the migration of inflammatory cells [199]. Release of tissue factor (also known as coagulation factor III or tissue thromboplastin) after endothelial cell damage, especially in the brain, activates α-thrombin, the final serine protease in the coagulation cascade [200]. The active thrombin disrupts BBB after cleaving protease-activated receptors known as PAR on endothelial cells thus gaining access to the CNS and impairing oxygen exchange. Once there, it can cleave to activate PARs on microglia/astrocytes for neuroinflammation and neurons to form neurofibrillary tangles [200]. Of note, microglial nodules formed after phagocytosis of hypoxic neurons were found in the cerebellum of COVID-19 patients [201]. Consistent with this knowledge, COVID-19 patients frequently present with microvascular ischemic and hemorrhagic parenchymal injury, microglial activation, and neuroinflammation [190, 199, 202]. Even mild respiratory SARS-CoV-2 infection without neuroinvasion may cause a multi-cellular dysregulation in the brain, including white matter microglial reactivity, abnormalities in neural precursor cell populations, reduction in hippocampal neurogenesis, depletion of myelinating oligodendrocytes, and myelin loss. Such anomalies are related to neurological symptoms, including impairment in attention, concentration, speed of information processing, and memory, as part of the long-COVID cognitive syndrome [203].

SARS-CoV-2 infection may directly impact mechanisms of disease in age-associated neurological disorders such as Alzheimer’s disease (AD) and Parkinson’s disease (PD). Vascular dysfunction plays a central role in AD and PD [204]. Expression of the apoE4 allele — a major risk factor for AD and chronic traumatic encephalopathy — in glial cells is associated with more severe neurodegenerative symptoms observed in older COVID-19 patients [205]. Moreover, ACE is upregulated in the limbic regions of the brain of patients with AD [206].

In addition to the direct involvement of neurovascular damage arising from increased inflammation, some of the post-acute neurological sequelae associated with SARS-CoV-2 infection could be due to dysregulation of RAS. Hyperinflammation has been reported more frequently in older patients with SARS-CoV-2 infection and is related to dysregulated immune and RAS in older adults that may underlie the observed increased risk of mortality and post-acute neurological sequelae in this demographic group [103].

The RAS plays a central role in the regulation of vascular function. Both the RAS and the vascular system are impacted by COVID-19 pathogenesis [207]. As discussed above, SARS-CoV-2 infection may exacerbate age-associated oxidative stress, which may compromise genome integrity in neurons and potentially other cells in the CNS, promoting neurodegenerative processes [208]. Studies that blocked the Ang-II receptor demonstrated that disruption of the blood–brain barrier is associated with RAS activation, dysfunction of the neurovascular unit, cognitive impairment, and dementia [209].
Although a strong association between SARS-CoV-2 and peripheral neuromyopathy as Guillain-Barré syndrome (GBS) is lacking, SARS-CoV-2 infection may be an occasional trigger for GBS. Previous studies showed that SARS-CoV-2-infected patients share neurological symptoms similar to those previously described in patients who experienced non-SARS-CoV-2 post-infection GBS [210, 211]. Moreover, peripheral neuropathy was most frequent among older adults [212].

Overall, a lower global cognitive performance score that is positively associated with severity of respiratory disease in COVID-19 patients suggests that brain injury could be due to reduced oxygen and subclinical neuroinflammation, both common in brain aging [213]. Taken together, this evidence may help explain the accelerated age-associated phenotypes observed in COVID-19 patients with neurodegenerative manifestations.

**Concluding remarks**

In addition to the respiratory tract, SARS-CoV-2 can also enter the CNS where it binds to cell receptors, including ACE2, expressed in several brain areas and both neuronal and non-neuronal cell types. The appearance or accelerated progression of neurodegenerative disease is manifested once the virus enters the CNS, promoting neuroinflammation, coagulopathies and hemorrhages, barrier dysfunction, and neuronal death. The varied range and severity of acute neurological COVID-19 syndromes as well as the consequences of sustained, viral-mediated neural dysregulation of peripheral systems may be a unique feature of SARS-CoV-2 infection. A disturbance of the bidirectional interactions of the nervous system and the RAS may accelerate aging through the generation of a pro-inflammatory state that can promote long-term neurological syndromes.

**Author contribution** Jorge Quarleri and Maria Victoria Delpino: Both have contributed equally to conceptualization, methodology, writing, review, and editing.

**Declarations**

**Conflict of interest** The authors declare no competing interests.
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