Cannabidiol – A Role for COVID-19?

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Abstract
Although the lung is considered as the primary organ affected by SARS-CoV-2, responsible for tissue and organ hypoxia, cerebrovascular complications such as stroke and neurological impairment such as Guillain-Barré syndrome have been repeatedly reported in COVID-19 patients. Many drugs that are currently administered to patients are experimental treatments which poorly cross the blood-brain-barrier. Cannabidiol (CBD) is known since long for reaching high brain levels and for its favourable safety profile. It has been shown to be neuroprotective and anti-inflammatory and has prevented sequelae in various hypoxic-ischemic disease models as well as in animal models of demyelination. In the past, it has been demonstrated that CBD can be safely co-administered with a number of other medications. CBD may be able to prevent and attenuate complications and long-term damages also in COVID-19 patients.

Keywords: Anosmia, Anxiety, Cannabidiol, COVID-19, Guillain-Barré syndrome, Hypoxia, Neuroinflammation, Neuroprotection, SARS-CoV-2, Stroke

Abbreviations: CBD: Cannabidiol; CVD: Cerebrovascular Disease; EAE: Experimental Autoimmune Encephalomyelitis; MOG: Myelin Oligodendrocyte Glycoprotein; TMEV: Theilers Murine Encephalitis Virus

Short Communication

Although the most challenging symptoms of COVID-19 have been related to acute respiratory distress, an increasing number of publications point to the infection of the vascular endothelial cells by SARS-CoV-2, subsequent endotheliitis, thromboembolic complications and impairment of the central and peripheral nervous system among other organs [1-7]. Most worrying are cerebrovascular events and polyneuroradiculitis, both occurring primarily in elderly patients.

Whereas cerebro- and other vascular complications likely result from a global inflammatory response and a hypercoagulable state induced by the infection with SARS-CoV-2, coronaviruses may disseminate to the central nervous system (CNS) through haematogenous spread, retrograde neuronal dissemination, or direct invasion of the olfactory epithelium. In a very recent publication, the Environmental Neurology Specialty Group of the World Federation of Neurology (ENSG-WFN) concludes that “respiratory failure, the lethal manifestation of COVID-19, responsible for 215,461 deaths worldwide, is probably neurogenic in origin and may result from the viral invasion of cranial nerve I, progressing into rhinencephalon and brainstem respiratory centers” [5]. This is supported by recent autopic findings of SARS-CoV-2 in olfactory pathways and brainstem [8]. A sudden dysfunction of smell and taste (up to complete anosmia and ageusia) often precede other symptoms and - most intriguingly - often in the absence of nasal congestion or rhinorrhoea seems to emerge as an early characteristic of SARS-CoV-2 infections; dysosmia has been observed in 70% to 98% of the patients, and more often in women than men [9,10]. Meanwhile, a wide range of additional neurological and neuropsychiatric symptoms and disorders have been described that include gait alteration, dizziness, seizures, headache, nausea and vomiting, cognitive impairment, inattention, irritability, agitation, confusion,
disorientation, anxiety, post-traumatic stress, sleep disturbances, depressed mood and even suicidality [11,12]. Although most patients seem to have recovered without experiencing mental illness, in a longer-term potential aftereffects may not be neglected [13,14]. Aftereffects of the pandemic are likely affecting also the general population [15].

Among many other signs and symptoms, cerebrovascular events have also been observed. Serious cerebrovascular disease (CVD) such as stroke, sinus thrombosis and cerebral haemorrhage have been reported primarily in patients who were significantly older (71-6 ± 15-7 years vs 52-1 ± 15-3 years), with more concomitant disorders and who were more likely to present with severe COVID-19 (84-6% vs. 39-9% [16]; in larger samples, ischemic stroke has occurred with an incidence of 1% to 6% [17], 2.4% [3], 2.5% [18], and 5.8% [16]. Individual case reports and case series also demonstrate that cerebrovascular events occur primarily in patients aged around 60 years and older [19-22], although ischemic stroke was occasionally observed in previously healthy, younger COVID-19 patients who were 36 years, 45 and 52 years old [23-25]. The average of time from SARS-CoV-2 infection to onset of CVD was about 12 days [16]. At present, data are still insufficient for assessing long-term sequelae of intermediary brain hypoxia during COVID-19.

Another manifestation which has been increasingly diagnosed with COVID-19 is the Guillain-Barré syndrome (GBS). GBS and its variant, the Miller-Fisher syndrome, is an autoimmune disorder of the peripheral nervous system, often triggered by acute infections. It manifests as an inflammatory autoimmune polyradiculoneuritis which has been related to a previous bacterial or viral infection (incidence ~1/100,000); it induces the production of antibodies which react with the myelin sheath of the peripheral nerves, resulting in demyelination and/or axonal injury [26]. By mid of May 2020, at least 21 cases related to COVID-19 have been reported with a median age of 61 years. Fourteen of these cases occurred in men; this contrasts to reports from the past, where women were more frequently affected [27]. GBS occurred shortly, after a median of 10 days (range 3 to 29 days) after the onset of symptoms [28-37]. This includes a case of polyneuritis cranialis in a 39 year old man [38].

Other encephalopathies that have been observed include severely impaired consciousness [7,29], meningoencephalitis [40], meningitis with convulsions [41], and hemorrhagic necrotizing encephalopathy [42]. All patients were aged above 55 years, except a 24 years old male with meningitis and a positive SARS-CoV-2 test in the cerebrospinal fluid [41]. Less threatening neurological and neuropsychiatric impairments which occurred in patients with COVID-19 are agitation (69%), confusion (65%), dysexecutive syndrome (33%), asthena (63.3%), ataxia (69%), as well as fatigue (26%-51%) [9,22]. In addition, 36% had myalgia, and increased creatine kinase (CK) was present in 33% of patients [5]. Overall, neurological involvement varies widely between about 15% to 21% to 36% [2,3,22,43], but was more common in patients with more severe disease (45.5%) than in non-severe patients (30.2%) [3]. Frequency is very much dependent on the population characteristics and can precede other symptoms by days [16].

Recent publications stressed that many drugs that are currently administered as experimental treatments for COVID-19 poorly cross the blood-brain barrier; among them monodonal antibodies, (hydroxy-)chloroquine, favipiravir, lopinavir + ritonavir or remdesivir [44], some of them with nervous system side effects that need to be carefully balanced against expected benefits [1]. In contrast to, cannabidiol (CBD) is well supported and known to penetrate in lipophilic brain tissue; this may open the way for CBD as a supportive treatment in inflammatory CNS infections. The potential of CBD and other cannabinoids for the prevention and treatment of tissue damages caused by severe hypoxia induced ischemic stroke has been repeatedly reviewed [45-47]; it has also been demonstrated in a number of various animal models of cardiac ischemia-reperfusion injury [48-50], renal ischemia-reperfusion injury [51-52], and newborn hypoxic ischemic encephalopathy [53-61]. In addition to protective effects in newborn hypoxic-ischemic encephalopathy, an impressive number of studies in adult animals has demonstrated that CBD can reduce hypoxia-induced brain damages when given either as prevention or as treatment up to about three days after occlusion [55,62-65], protecting animals from long-term sequelae [66]. Nonetheless, confirmation in human subjects is still lacking and no respective studies seem to be ongoing (www.clinicaltrials.gov).

In addition, CBD treatment has also demonstrated myelin protective effects in a wide array of different demyelinating disease models in animals. One of them studied hypomyelination induced by hypoxia / ischemia in the immature brain. When newborn rats were exposed to hypoxic-ischemic injury, the subsequent treatment with CBD (1mg s.c./kg) completely preserved function and myelination, suggesting a general neuroprotective effect [56]. In another model, experimental autoimmune encephalomyelitis (EAE) was induced in mice by injection of myelin oligodendrocyte glycoprotein (MOG35-55 peptide) as antigen; mice progressively developed EAE disease, mimicking the inflammatory lesions on the myelin sheath in multiple sclerosis. The disease typically follows the course of a progressive degeneration, with visible signs of pathology consisting of flaccidity of the tail and loss of motion of the hind legs. It was found that intraperitoneal administration of CBD (20mg ip./kg) delayed the onset of disease and significantly attenuated clinical signs of EAE, proliferation of MOG-specific T-cells, and also significantly reduced inflammatory cytokine levels of IFNγ and IL-17. A dose of 20mg/kg body weight of CBD translates to 1.6mg/kg when converted to human equivalent dose [67]. Furthermore, CBD reduced autoimmune encephalomyelitis also in a model of adoptively transferred EAE. When splenocytes and lymph nodes from mice with actively induced EAE were cultured in the presence of MOG35-55 and IL-12, and inoculated intraperitoneally in recipient mice, EAE is adoptively transferred to recipients (at-EAE); CBD proved to
be effective also in this model [68]. Using the same model, another group of researchers could demonstrate that treatment with CBD (10mg i.p./kg per day), starting from the occurrence of the first signs of disease (14th day) until when mice have been sacrificed on the 28th day from EAE-induction, can avoid programmed cell death in the spinal cord of animals affected [69]. Finally, a model which is widely used for studying the chronic, demyelinating process in multiple sclerosis, is the Theiler’s murine encephalitis virus model (TMEV-induced demyelinating disease, TMEV-IDD), a naturally occurring single-stranded RNA-virus. In this model, mice were infected intracerebrally with TMEV and then treated with CBD (5mg i.p./kg) or the vehicle alone, once daily for up to 10 days post-infection to analyze the long-term effects. CBD completely blocked TMEV-induced release of the adhesion molecule VCAM-1 which is mainly expressed by endothelial cells and that binds selectively to VLA-4 expressed by monocytes and lymphocytes; this significantly reduced the expression of the chemokines CCL2 and CCL5 transcripts and of IL-1β. CBD treatment for 10 days in the acute phase of the disease attenuated neurodegeneration and demyelination in the chronic phase of TMEV-IDD, completely restoring both horizontal and vertical motor activity when on day 80, motor function was evaluated [70].

In summary, although animal models do not necessarily correlate one-to-one to the therapeutic situation in man, it may be worth to try CBD in COVID-19 patients early when they start showing neurological symptoms. CBD reaches high concentrations in brain, has neuroprotective and anti-inflammatory properties, is well tolerated and can be combined with many other medications. Furthermore, it potentially limits stress, anxiety and other aftereffects.

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Conflict of Interest
Author declare no conflict of interest.

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