Neurology and Covid-19: Present and future

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Abstract

We’re witnessing a pandemic due to COVID-19, which has generated a strong impact in Public Health, Health Services and an economic level, still to be determined. It’s a recently discovered respiratory virus that has the capacity to produce neurological symptoms, either indirectly or directly. Much has been written about neurological symptoms, mainly by no-neurologist, and the frequency and type of neurological manifestations must be precisely defined in both, acute and medium-long term phases. After carrying out an extensive bibliographic review, the objective of this paper is to analyze what has been described so far and how it can be improved to know exactly the type and frequency of neurological manifestations related to COVID-19.

Introduction

Back in 1896, Sir William Osler said that humankind is facing three major problems: fever, famine and warfare, being fever the worst of them. Nowadays, we find ourselves fighting against a pandemic that has directly threatened Public Health, Sanitary Services and Economy. Causing a vast amount of infections and deaths, as well the need to develop several measurements such as social distancing or confinement, always seeking the common good. Something similar took place at the end of the Great War: the 1918 flu pandemic, wrongly named “the Spanish Flu”, from which there are almost no survivors left.

Our current pandemic started at the end of 2019 in Wuhan, China [1], rapidly acquiring the epidemic status and, short after, the pandemic one. It affected the areas of China, Southern Europe- specially Italy and Spain-, U.S.A., and Latin America, with a special virulence. Few are the places which escape this virus’ grip. To this day, 1st of June 2020, the number of infected rises to 6.171.341, with 372.166 deaths and 2.642.568 recoveries.

A beta-coronavirus is responsible for this pandemic. It was unknown until the recent events and it was named Covid-19 (SARS-Cov-2), encapsulated RNA virus, with a molecular mass that oscillates between 20-32 KB, genetic diversity and with the ability of carrying out recombination and thus, causing cross-species transmission. It presents a high contagiousness index [2], higher than the flu’s and the one of the SARS-Cov-1. However, the current virus is less lethal than the last one. The virus is transmitted through airways, within 1 to 2 meters, and its survival improves in closed spaces, where aerosols can be created, and it can survive in fomites for 72 hours. It has been isolated in saliva, pulmonary secretions, urine and, rarely, in cerebrospinal fluid. Several asymptomatic patients are produced, and only a small percentage of them develops the disease, being older the ones at a higher risk [3]. The disease is divided in three stages, I or initial, II, with respiratory issues, and III, which is related with the cytokine storm; the ability of producing neurological manifestations along three stages.

The pathogenic mechanism by which the disease can reach the nervous system is being studied. Recent studies point to haematogenous dissemination and through the olfactory bulb, where receptor ACE-2 [4,5] and retrograde transsynaptic transport [6] play an important role. As well as this, there’s anecdotal evidence of neurotropism [7] caused by the virus. That determines the cytokine release syndrome [8], that tends to be produced in stage III of the disease, to alter the blood-brain barrier’s permeability, allowing the flow of molecules and cells, causing immune response at the nervous system’s level and start inflammatory immune mechanisms that can be responsible for neurological disorders [9] and later manifestations (Figure 1).

Few diseases have generated this high number of publications and in such a short period of time, regarding neurological manifestations. Covid-19 has the properties of neuroinvasion (penetrate the nervous system), neurotropism (affect glial cells and neurons) and neurovirulence (produce a neurological pathology). It is based on several criteria, including: biological plausibility, neurological damage that it can produce in other species, animal models of nervous system’s invasion and neurological complications by other coronaviruses that Covid-19 patients have developed [10]. There are general series that define those manifestations. However, few of them have been done by neurologists.

Our main goal is to carry out a bibliographic review of those manifestations, allowing us to catalogue and define with a better accuracy the Covid-19 related neurological manifestations.

Material and methods

In order to carry out our project, we have done a bibliographic research, using only those English-written articles from November 2019 to 2020 which are the most representative ones, for our analysis, and describe with the best quality the neurological manifestations caused by infection of beta-coronavirus Covid-19.

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Encephalopathy- fluctuating, reversible and global cerebral dysfunction- is more common between older patients, along with hypertension and other comorbidities, primarily obesity and with previous cognitive decline. They look like a vulnerable group, as well as those who take ECA suppressors. In Mao’s study, a 15% of severe Covid-19 patients, underwent consciousness’ alterations. Fitalov publishes [15] that encephalopathy’s origin is multifactorial, including toxic-metabolic causes, focusing on hypoxia, neurotoxic side effects of drugs and subclinical seizures.

Moniguchi, et al. [16] publish the first case of encephalitis linked with Covid-19, in a Japanese patient with inflamed cerebrospinal fluid and positive RCP. This is the first case where a direct threat to the nervous system by the virus is observed. A case of Covid-19 associated acute haemorrhagic necrotizing encephalopathy is described by Poyiadji, et al. [17], in a radiology journal. The disease is confirmed by nasopharynx RCP and injuries at bilateral anterior temporal, thalamus and insula levels are observed. The previous manifestations are related with the cytokine release syndrome. In both above described cases, even though they are anecdotal, a certain predisposition to produce limbic encephalitis is appreciated. Myelitis [18] is thought to have a connection with Covid-19.

Thromboembolic manifestations, specifically stroke [19], have been described with the disease, being more frequent in the late stages of it and seem to share a connection with D-dimer, hypercoagulability, cytokine release syndrome and the infection’s gravity. Five cases of young patients with large vessel stroke in New York, with no vascular risk factors history, are published in NEJM [20]. Moreover, Barrios-López, et al. [21] describe four stroke cases in severe Covid-19 infected patients. Mao’s description, with 214 hospitalised patients, several neurological manifestations are described: ischemic strokes, one haemorrhagic stroke and cerebral venous thrombosis.

Regarding peripheral nervous system’s manifestations, Sedaghat and Toscano [22,23] describe Guillain-Barré’s syndrome, which occurs two weeks after the initial stage of the infection. They found evidence of demyelination in the electromyography, suggesting an immunemediated origin. In addition, Gutiérrez-Ortiz, et al. [24] describe two cases of cranial polynuertis, compatible with Miller-Fisher’s syndrome, two weeks after infection. Muscle affection seems to be a common manifestation in the early stages of the disease, and tend to increase as the illness degenerates. Cases of rhabdomyolysis have been described [25].

In relation to the medium to long term neurological manifestations, Serrano Castro, et al. [26] state in their article, the possibility of producing an immune/inflammatory phenomenon on vulnerable patients with a neurodegenerative and psychiatric pathology that modifies those diseases’ normal course.

ALBACOVID register’s data, published in Neurology, by Romero-Sánchez, et al. [27] is the first hospital record in which the neurological manifestations are described by neurologists from two Spanish hospitals. It is a retrospective and hospital study with 814 positive Covid-19 patients, where the 50.4% develop neurological manifestations. The average age of the sample is 66.4 years, with 56% males. They develop myalgias (17.2%), cephalgia (14.1%), dizziness (6.1%), dysgeusia (6.2%) and hyposmia (4.9%). Consciousness’ level manifestations happen in a 19.6% of the patients, being more frequent in the graver and older ones. Myopathy (6.1%), dysautonomia (2.5%), stroke (1.7%), seizures and movement disorders (0.7%), one case of encephalitis, Guillain-
Barre syndrome and optic neuropathy. Neurological manifestations were related with the 4.2% of deaths. The authors suggest that in order to analyse the neurological manifestations due to Covid-19 infection, there must be a separation between the following levels: unspecific neurological symptoms (dizziness, headache and myalgia), neuro-psychiatric alterations (anxiety, insomnia, depression and psychosis), central nervous system’s alterations (direct viral infection, consciousness’ level alterations, stroke and seizures), peripheral nervous system’s alterations (cranial and peripheral neuropathies), myopathy and demyelinating disease.

We are currently facing an increase in available literature about the nervous system’s damage, caused by Covid-19 infection. Nevertheless, few of them are written by neurologists [28] and virtually none are a population study that analyses the neurological manifestations in a global manner. It is important to define which are the essential mechanisms of that affection. The cytokine release syndrome appears to play an important role, contributing to increasing the blood-brain barrier’s permeability, causing molecules and cells to flow into the nervous system. Hypoxia is also important, as well as the direct injury mechanism by the virus, like HIV- from which we have anecdotal data. It can reach the nervous system through the blood vessels, using the “Trojan’s horse” mechanism. This event’s likelihood increases with the cytokine storm, in the gravest scenarios of the disease. Furthermore, the retrograde transynaptic transport through the olfactory bulb is also an essential mechanism. There is an report, where the RM signal seems to intensify in the frontobasal lobe and fade with Covid-19 recovery [29]. Respiratory system’s chemoceptors/mecanoreceptors can influence in the Covid-19 related type 2 respiratory failure, where the ACE2 receptors play an important role. A recent neuropathologic study with 18 Covid-19 patients [30], shows that the primal changes are caused by hypoxia and that there are no viral particles in those brains. As a result, hypoxia and cytokine storm acquire a lead role in this disease’s neurological manifestations.

Recently, “NeuroCovid Staging” [31] is being used as a better way of understanding physiopathology and the neurological manifestations produced (Figure 2).

- **Stage I.** The virus unites the ACE2 in the olfactory and gustatory neuroepithelia. Cytokine storm is mild. Symptoms are dyseguesia and hyposmia.
- **Stage II.** A strong immune response is produced, with a ferritin, reactive C protein and D dimer increase. That leads to, on one hand, a hypercoagulability that provokes strokes-, and on the other hand, an immune response and vasculitis. As a result, there is damage on the cranial nerves, peripheral nerve and muscles.
- **Stage III.** The cytokine storm damages the blood-brain barrier, favouring molecules and cells flow, as well as the virus. Consequently, there is brain damage and edema, leading to encephalopathy and seizures. An increased level of angiotensin favours hypertension and brain haemorrhage.

Just like it was observed in the 1918 Flu Pandemic, regarding Von Economo’s encephalitis; and currently, with the HIV-related cognitive impairment- that even though it has zero viral blood presence, it causes an inflammatory response, maintained by HIV, in the nervous system- it is necessary to define the medium to long term neurological manifestations present in a specially vulnerable population, such as those with neurodegenerative diseases caused by Covid-19. The SARS-Cov1 epidemic favoured Parkinson’s disease and multiple sclerosis development [32,33]. The high ACE-2 content in the neurons and the glial cells can determine its influence on developing medium to long term neurodegenerative and age-related diseases. Therefore, evidence shows the importance of neurologists monitoring Covid-19 patients.

Table 1 shows the Covid-19 related neurological manifestations described in literature.

This pandemic has caused a reorganisation in the Sanitary Services- as a whole- and in the Neurology Services-in particular [34,35].
Finally, we can conclude the following about the Covid-19 related neurological manifestations:
- Covid-19 related neurological manifestations and their frequency are still a subject of study.
- The most common neurological manifestations are cephalalgia, dizziness and myalgia.
- Anosmia might be an indicator of the disease.
- Encephalopathy, stroke and seizures are becoming more frequent. They are all related to the cytokine storm, where hypoxia acquires a critical role. Encephalopathy is the most frequent one. This has been observed in 18 patients.
- Encephalitis and polyradiculopathy are infrequent. A certain level of tropism could cause limbic encephalitis, due to the increase in the tempo-hematoencephalic barrier’s fragility.
- “Neurocovid Staging” seems to be very adequate for having a better understanding of physiopathology and neurological manifestations.
- It is necessary to increase the number of population studies in order to get to know the essential mechanisms that cause the Covid-19 related neurological pathologies.
- It might be a priority to increase the number of pathological autopsies, in order to get to know the essential mechanisms that cause the Covid-19 related neurological pathologies.

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