Characteristics of COVID and post COVID polyneuropathies in adults and pediatrics: an Egyptian sample

Haidy Elshebawy1*, Mohamed Y. Ezzeldin2 and Eman Hassan Elzamarany3

Abstract

Background: The aim of this study is to describe the different forms of polyneuropathy associated with coronavirus disease 2019 (COVID-19) as a secondary neurological complication for (COVID-19) and the outcome from different therapeutic regimens in adults and pediatrics in first and second waves of the pandemic.

Case presentation: This study was conducted on 42 patients, they were divided into two groups, group (A) and group (B) in first and second waves respectively. Twenty-five patients presented by ascending weakness preceded by fever, dry cough and respiratory distress, electromyography (EMG) and nerve conduction (NC) studies done and confirmed the clinical diagnosis of demyelinating polyneuropathy. Eight patients presented by acute flaccid quadripareisis, more severe in upper limbs preceded by fever and diarrhea diagnosed as acute axonal polyneuropathy. Five patients presented by severe fatigue and progressive weakness of both lower and upper limbs, they developed fever and cough 10 days after the neurological symptoms. EMG and NC done and confirmed clinical diagnosis of polyneuropathy of demyelinating with secondary axonal picture. Four patients presented 30 to 40 days after their recovery form corona virus infection with gradual progressive weakness of both upper and lower limbs over 2 to 3 months duration, mainly the proximal muscles of lower limbs were affected with areflexia. EMG and NC done and confirmed the diagnosis of chronic inflammatory demyelinating polyneuropathy (CIDP).

Conclusion: We should gain a better understanding of the underlying pathophysiology and therapeutic options of polyneuropathies related to COVID-19, which will have an impact on the treatment of the COVID related respiratory failure presenting with neuropathy.

Keywords: Post COVID, Polyneuropathies, Adults, Pediatrics

Introduction

The novel coronavirus disease 2019 (COVID-19) pandemic is a world-shattering infection that affects all geographical areas. With the rapid increase in the numbers of cases of COVID-2 pandemic with its different waves, clinical presentation and complications are rising across the countries and the whole world since December 2019 [1].

Corona virus like many respiratory viruses exhibit respiratory as well as extra pulmonary symptoms, and known that it may include the nervous system and causing a variety of neurological manifestations through direct neural invasion or indirect virus-induced host immune responses [2].

COVID-19 associated neuromuscular disorders are the second most commonly encountered neurological complications, especially Gullian barre syndrome (GBS) [3]. Diagnosis and management of these complications will have an impact on the treatment of the severe acute respiratory syndrome coronavirus 2 (SARSCoV-2) related respiratory failure presenting with neuropathy [4].
aim of this study is to describe COVID-19 patients who presented or complicated with different forms of polyneuropathy in adults and pediatrics in first and second waves of the pandemic and their outcome with different therapeutic regimens.

**Methods**

Our study is a cross sectional descriptive study conducted on 42 patients presented to us in the Neurology outpatient clinics from multiple health care centers in Egypt by multiple clinical forms of polyneuropathy in period from March 2020 to end of February 2021. Patients recruited from the outpatient clinics were divided into two groups. Group (A) involved 23 patients recruited in the first wave of corona virus in the period from March 2020 till the end of September 2020 and group (B) involved 19 patients recruited in the 2nd wave in the period from October 2020 till February 2021. We included in our study patients who presented with new onset polyneuropathy in any age group following or complicating COVID 19 infection. Patients who have a previous history of polyneuropathy or have a chronic illness which may be associated with polyneuropathy eg. (DM, autoimmune disorders, HCV) were excluded. An informed written consent was taken from each patient. All data obtained from every patient were confidential and were not used outside the study. Our study was approved by ethical committee of the Department of Neurology, Faculty of Medicine (Cairo University).

All patients in our study were conducted for history taking, neurological examination, EMG and NC studies were carried out for all patients to confirm clinical diagnosis of polyneuropathy.

SARS-CoV-2 testing from nasal swab, computed tomography (Ct) chest and other laboratory investigations (CBC, ESR, CRP, ferritin and D-dimer); were done for all patients.

EMG and nerve conduction studies were done using Nihon Kohden Neuropack M1 MEP-9200 EMG machine, Japan. Posterior tibial, common peroneal, superficial peroneal, motor and sensory ulnar nerves were examined for NCS, bilaterally. Both Facial nerves were added for NC protocol of study if there were clinical facial signs. EMG examination was carried out for both tibialis anterior, vastus medialis, deltoid, biceps brachii and first dorsal interosseous muscles.

**Clinical presentations**

Twenty-five patients presented by ascending weakness affecting both lower and upper limbs with severe hypotonia and areflexia after manifestations of fever, dry cough and headache by (5–10) days. Three of them developed respiratory distress. EMG and NC done for all of them fulfilling the criteria of acute demyelinating polyneuropathy, at least three of the following were found: prolonged DLs (two or more nerves, not at entrapment sites) DL > 115% of ULN (for normal CMAP amplitudes), DL > 125% of ULN (for CMAP amplitudes < LLN), CV slowing (two or more nerves, not across entrapment sites) CV < 90% of LLN (for CMAP amplitudes > 50% LLN) CV < 80% of LLN (for CMAP amplitudes < 50% LLN), Prolonged F responses (one or more nerves) > 125% of ULN; or absent F responses, in addition to conduction block and/ or temporal dispersion may be found. EMG examination showed MUP’s of reduced recruitment and no at rest activities. PCR was positive for COVID in nasopharyneal swap in all of them. All of them received eight sessions of plasma exchange, twenty of them improved with good outcome and discharged on physiotherapy and five partially improved and need intravenous immunoglobulins (IVIG) for 5 days with better outcome and discharged.

Eight patients presented by acute flaccid quadripareisis, more severe in upper limbs mainly distal with areflexia, preceded by fever and diarrhea 1 week before the neurological manifestations, MRI cervical spine was free for all of them, EMG and NC showed picture of acute axonal polyneuropathy in form of reduced amplitudes of CMAPs (compound motor action potentials) and SNAPs (sensory nerve action potentials) and slightly reduced conduction velocities not reaching the demyelinating range, MUP’s of reduced recruitment and resting activities in the form of positive sharp waves and fibrillation potentials may be found in EMG. Also, PCR from nasal swap and other laboratory data confirm COVID diagnosis. Five of them received one cycle of IVIG for 5 days with marked improvement and the others need 2 cycles of IVIG with partial recovery.

Five patients presented by severe fatigue and progressive weakness of both lower and upper limbs with areflexia and hypoesthesia distally, they developed fever and cough 10 days after the neurological symptoms. EMG and NC were done and the electrophysiological findings of mixed demyelinating and axonal polyneuropathy picture were found. Those patients started treatment in form of 6 sessions of plasma exchange in three of them and the other two received 5 days of 1 gm methylprednisolone with marked improvement for all of them, when the respiratory symptoms manifested, laboratory tests and PCR done and confirmed COVID-19 diagnosis.

Four patients presented 30 to 40 days after their recovery form COVID-19 infection evidenced by 2 negative nasopharyngeal swaps with 48 h in between and discharged from hospital by gradual progressive weakness of both upper and lower limbs over 2 to 3 months duration mainly proximal muscles are affected more
in lower limbs with areflexia. Two of them have bilateral lower motor neuron (LMN) facial and neck muscle weakness, and the other two patients had numbness in both lower limbs. EMG and NC were done and confirmed the diagnosis of chronic inflammatory demyelinating polyneuropathy (CIDP) for all of them fulfilling the electrophysiological criteria in form of markedly prolonged distal latencies (>130% of ULN), markedly slowed conduction velocities (<75% of the LLN), and markedly prolonged or absent late responses (>130% of ULN), there is usually conduction block and/or temporal dispersion. The EMG examination showed MUP's of reduced recruitment and no resting activities. All patients received 5 days of 1 gm methylprednisolone with marked improvement in one of them and minimal improvement in other three patients, with gradual withdrawal of steroids was planned to all of them.

With the start of 2nd wave of COVID 19 in Egypt, we added 19 patients to our study, 11 of them were in pediatric age group aged from 2 to 10 years old presented by picture of acute motor paralysis with ascending weakness affecting both lower and upper limbs with severe hypotonia and areflexia and 4 of them developed bilateral LMN facial. The condition preceded by gastrointestinal manifestations in 7 of them with vomiting, diarrhea with high grade fever. The other 4 had fever with cough preceding the weakness. EMG and NC done to all of them and revealed the typical picture of acute inflammatory demyelinating polyneuropathy (AIDP). All of them received IVIG with good response.

**Statistical analysis**

The collected data was reviewed for completeness and logical consistency. Pre-coded data will be entered on the computer using Microsoft Office Excel software program for Windows 2019. Data was transferred to the Statistical Package for the Social Sciences (SPSS version 26) (IBM Corp., Armonk, NY, USA) to be statistically analyzed. Quantitative variables were described as mean, standard deviation, median, IQR. They were compared using Mann–Whitney U test, where p-value is significant if less than 0.05. Qualitative variables were described as frequency and percentage. They were compared using Chi-square test or Fisher’s exact test accordingly where p-value is significant if less than 0.05. Exact test was used instead when the expected frequency is less than 5

**Results**

Out of 42 patients with confirmed COVID-19, 23 (54.8%) in the first wave in the period from March to end of September 2020 and 19 (45.2%) patients in the second wave from October 2020 to end February 2021, 22 males (52.4%) and 20 females (48.6%), 28 adults (66.7%) their ages range from (35–55 years) and 14 in the pediatric group(33.3%), their ages range from (2–10 years). Clinical characteristics of studied patients illustrated in Table 1.

Regarding the comparison between the two waves of the pandemic, there is a significant difference between the 2 groups regarding the age with P value 0.00. Also, there is a significant difference regarding the mean age in the first wave comparing it with second wave, with P value 0.000, but there was no significant difference

| Table 1 | Clinical characteristics of studied patients |
|---------|-------------------------------------------|
| **Neuropathy** | **Frequency** |
| AIDP (motor and sensory) | 25 (59.5%) |
| AMSAN | 8 (19%) |
| Demyelinating with secondary axonal | 5 (11.9%) |
| CIDP like | 4 (9.5%) |
| **CT chest** | **Frequency** |
| Normal | 2 (4.8%) |
| Interstitial pneumonia | 3 (7.1%) |
| Interstitial pneumonia without parenchymal opacity | 4 (9.5%) |
| Bilateral GGO | 33 (78.6%) |
| **PCR** | **Frequency** |
| Positive | 36 (85.7%) |
| **CBC** | **Frequency** |
| Lymphopenia | 30 (71.4%) |
| Elevated | 31 (73.8%) |
| **CRP** | **Frequency** |
| Elevated | 31 (73.8%) |
| **Outcome** | **Frequency** |
| Full Improvement | 31 (73.8%) |
| No improvement and needed second treatment to improve | 8 (19%) |
| Minimal improvement | 3 (7.1%) |
| Total | 42 (100%) |

AIDP acute inflammatory demyelinating polyneuropathy, AMSAN acute motor sensory axonal neuropathy, CIDP chronic inflammatory demyelinating polyneuropathy, CT computed tomography, GGO ground glass opacity, CBC complete blood count, CRP C reactive protein
between both groups regarding the sex with P value 0.757 (Table 2).

There was no significant difference between both waves regarding the type of polyneuropathy (P value 0.193) and the outcome of the recovered patients (P value 0.084) (Table 3).

Regarding the comparison between adults and pediatric groups, we found that the picture of neuropathy affecting the pediatric mainly is the AIDP rather than different types of neuropathy affecting the adults group with a significant difference between the two groups (P value 0.003) (Table 4), but there was no significant difference regarding the outcome of the neuropathy in treatment between the two groups (P value 0.129) (Table 5).

**Discussion**

Coronaviruses are not primarily neurotropic viruses and their primary targets are respiratory and cardiovascular systems. The virus is attached to host cells through the Angiotensin-converting enzyme 2 (ACE-2) receptors leading to internalization and subsequent viral replication. This receptor is also found in glial cells in the Central Nervous System (CNS) and spinal neurons. The virus can invade peripheral nerves and lead to retrograde transfer via synapse mediated route to CNS [5, 6].

Also, the cytokine release syndrome (CRS), caused by an exacerbated recruitment and activation of macrophages, neutrophils, and natural killer cells (NK) in response to SARS-CoV-2 infection result in releasing

---

**Table 2** Comparisons between both waves regarding demographics

| Sex   | Wave | Male | Female | P  | Age group | P  | Total |
|-------|------|------|--------|----|-----------|----|--------|
|       | 1st wave | 13 (56.5%) | 10 (43.5%) | 0.757 | 20 (87%) | 3 (13%) | 0.003 | 23 (100%) |
|       | 2nd wave | 9 (47.4%) | 10 (52.6%) | | | | | 19 (100%) |

| Wave | n | Mean | SD   | P   |
|------|----|------|------|-----|
| Age (in years) | 1st wave | 23 | 41.17 | 15.52 | 0.000 |
| | 2nd wave | 19 | 19.63 | 18.43 | |

n number, SD standard deviation

**Table 3** Comparisons between both waves regarding the type of neuropathy and outcome

| Neuropathy type | Wave | AIDP | AMSAN | Demyelinating with secondary axonal | CIDP like | P   |
|----------------|------|------|-------|-----------------------------------|----------|-----|
|                | 1st wave | 11 (47.8%) | 5 (21.7%) | 3 (13.0%) | 4 (17.4%) | 0.193 |
|                | 2nd wave | 14 (73.7%) | 3 (15.8%) | 2 (10.5%) | 0 (0.0%) | |

AIDP acute inflammatory demyelinating polyneuropathy, AMSAN acute motor sensory axonal neuropathy, CIDP chronic inflammatory demyelinating polyneuropathy

| Outcome | Wave | Full improvement | Minimal improvement | No improvement and needed second treatment to improve | P   |
|---------|------|------------------|---------------------|-----------------------------------------------|-----|
|         | 1st wave | 14 (60.9%) | 3 (13.0%) | 6 (26.1%) | 0.084 |
|         | 2nd wave | 17 (89.5%) | 0 (0.0%) | 2 (10.5%) | |

**Table 4** Comparison between the adult and pediatric groups according to type of neuropathy

| Neuropathy type | Total | P   |
|----------------|-------|-----|
| AIDP | AMSAN | Demyelinating with secondary axonal | CIDP like |
| Adult | 11 (39.3%) | 8 (28.6%) | 5 (17.9%) | 4 (14.3%) | 28 (100%) | 0.003 |
| Pediatric | 14 (100%) | 0 (0.0%) | 0 (0.0%) | 0 (0.0%) | 14 (100%) | |

AIDP acute inflammatory demyelinating polyneuropathy, AMSAN acute motor sensory axonal neuropathy, CIDP chronic inflammatory demyelinating polyneuropathy
many activated leukocytes during the inflammatory phase can lead to extensive tissue damage, including the peripheral nervous system, and appears to correlate with COVID-19 severity [7, 8].

COVID-19 may not directly invade nerves, its roots or the anterior horn cells as seen in polio virus or West Nile virus leading to their damage. Even the cerebrospinal fluid (CSF) and the polymerase chain reaction (PCR) for coronavirus in multiple reported cases of COVID-19 related GBS has been negative [9]. It is likely to be a post infectious or a para-infectious complication resulting from an aberrant immune response which can be considered a second mechanism explaining GBS in COVID-19 by production of antibodies against ganglioside components of the peripheral nerves, owing to molecular mimicry with surface antigens of the infectious pathogen [10].

Our study was conducted on 42 patients, included the first and second waves of the pandemic, through the period from March 2020 till the end of February 2021. We found that COVID and post COVID polyneuropathies with its different types and forms are of the common neurological complications of COVID virus infection that may affect and threaten the health if not taken in consideration.

In our study we reported that more than half of our patients (59.9%) presented with typical pattern of AIDP which was against Yaranagula and Koduri who reported that more than half of the patients in the their cohort had paraparesis [11]. The paraparetic variant is an uncommon form of GBS and constitutes about 5–10% of all GBS cases [12].

In the current study, most of our patients presented with demyelinating neuropathy rather than the axonopathy which was in accordance with Yaranagula and Koduri who reported that more than half of the patients in the their cohort had paraparesis [11]. The paraparetic variant is an uncommon form of GBS and constitutes about 5–10% of all GBS cases [12].

In the current study, many activated leukocytes during the inflammatory phase can lead to extensive tissue damage, including the peripheral nervous system, and appears to correlate with COVID-19 severity [7, 8].

Table 5 Comparison between the adult and pediatric groups according to outcome of neuropathy

| Outcome | Full Improvement | Minimal improvement | No improvement and needed second treatment to improve |
|---------|------------------|---------------------|-----------------------------------------------|
| Adult   | 18 (64.3%)       | 3 (10.7%)           | 7 (25%)                                       |
| Pediatric | 13 (92.9%)   | 0 (0.0%)            | 1 (7.1%)                                      |

Table 5 Comparison between the adult and pediatric groups according to outcome of neuropathy [11], as has been reported with Zika virus [13] and that agreed with our study.

In 2020, Virani and colleagues reported a 54-year-old male with confirmed COVID-19 and a history of fever, cough and *Clostridium difficile* colitis of recent onset developed difficulty breathing and weakness and diminished reflexes of arms and legs eventually diagnosed as GBS [10]. A 65 years old male was reported by Sedaghat and colleagues, with ascending quadraparesis, areflexia and bifacial palsy, after positive nasal swab and normal CSF, when EMG and nerve conductions were done on day 9 of weakness that revealed GBS variant-AMSAN [14].

El-Otmani and colleagues reported 70 years old female with quadriplegia, hypotonia and areflexia, that her MRI was normal but EMG and nerve conductions revealed axonal pattern of atypical GBS and diagnosed as ASAN, that it's CSF protein was 100 mg/dL and normal WBCs [15] which agreed with our study.

Camdessanchi and colleagues reported a 64 years old male with flaccid paralysis and parasthesia. On day 5 EMG and NCV were done that revealed polyneuropathy of mixed demyelinating pattern, the CSF protein was 54 mg/dL [16].

In 2020, two other studies done by Scheidl and colleagues and Padroni and colleagues reported two patients with paraparesis and areflexia, EMG/NCS revealed demyelinating pattern and diagnosed as GBS and CSF proteins are elevated [17, 18].

Ottaviani and colleagues reported 66 years old Female with difficulty in walking, paraparesis and areflexia then progressed to quadriparesis with facial weakness. On day 10 EMG/NCS revealed mixed axonal and demyelinating patterns and diagnosed as GBS. The CSF protein was 108 mg/dL, normal WBCs and serum antiglycolipid antibodies were absent [19], which agreed with our study.

Abdelnour and colleagues reported 69 years old male presented with bilateral lower limb weakness and areflexia, brain and spine MRIs were normal, EMG/NCS were not performed however he was diagnosed as GBS, inspite that the CSF was not performed [20].

Caamaño DS and colleagues reported a 61 years old male with bilateral facial weakness, unresponsive blink reflex on both eyes. Brain MRI was normal. EMG/NCS
were not performed. Diagnosed as atypical GBS variant with facial diplegia. The CSF protein was 44 mg/dL [21].

Toscano and colleagues reported 5 patients with GBS from three hospitals in northern Italy during the COVID-19 outbreak. In sum, GBS occurred 5 to 10 days after onset of COVID-19 symptoms, a typical interval. Clinical neurophysiology was consistent with axonal-type GBS in 3 cases and demyelinating-type in 2 patients. Post-Gadolinium MRI showed enhancement of caudal nerve roots in 2 patients and the facial nerve in one. All patients were treated with IVIG and Patients 1 and 3 received two cycles. A similar treatment was used in GBS patients during the MERS outbreak [22].

With the emergence of more cases of polyneuropathies temporally linked to SARS-CoV-2 infection, we can understand the improvement of such patients on steroids, plasma exchange and IVIG. We should gain a better understanding of the underlying pathophysiology and potential therapeutic options of GBS related to COVID-19.

Several therapeutic options are still under study, with the intent to stabilize the immune system in COVID-19 and either prevent or minimize the consequences of this storm, as reviewed by Diamanti and colleagues [23].

Since these neuropathies are treatable and they pose increased morbidity and mortality, physicians working with COVID-19 patients must be aware of this association.

In our study, we noticed different types of polynuropathies which vary according to age groups and also it may be the initial COVID symptoms between Group A and Group B, this could be due to the genetic change in the corona virus phenotype or change in the autoimmune response of the body to the virus in the second wave.

The presentation of the neurological complications attacked the younger age in the second wave may be due to the eased of the global restrictions of closure, opening of the schools and nurseries, making the children in this age group more vulnerable for infection, or may the adults gained immunity from previous exposure to the first coronavirus season as they were more exposed for infection for the sake of keeping their works and the individual financial income.

Conclusions

We should gain a better understanding of the underlying pathophysiology and therapeutic options of polynuropathies related to COVID-19, which will have an impact on the treatment of the COVID related respiratory failure presenting with neuropathy.

Abbreviations

COVID-19: Coronavirus disease 2019; SARS-CoV-2: Severe acute respiratory syndrome coronavirus 2; DM: Diabetes mellitus; HCV: Hepatitis C virus; EMG: Electromyography; NC: Nerve conduction; CT: Computed tomography; CBC: Complete blood count; DL: Distal latency; ESR: Erythrocyte sedimentation rate; CRP: C reactive protein; ULN: Upper limit of normal; LLN: Lower limit of normal; CMAP: Compound motor action potentials; SNAPs: Sensory nerve action potentials; MRI: Magnetic resonance imaging; PCR: Polymerase chain reaction; IVIG: Intravenous immunoglobulin; MUP’s: Motor unit potentials; CIDP: Chronic inflammatory demyelinating polyneuropathy; ADP: Acute inflammatory demyelinating polyneuropathy; AMSAN: Acute motor sensory axonal neuropathy; MERS: Middle East Respiratory Syndrome; ACE-2: Angiotensin-converting enzyme 2; CNS: Central nervous system; GBS: Guillain Barre Syndrome; GGO: Ground glass opacity; IVIG: Intravenous immunoglobulins.

Acknowledgements

The authors acknowledge subjects for their participation and cooperation in this study.

Authors’ contributions

EHE: research idea, data acquisition, data analysis and interpretation, HME: data acquisition, data interpretation and manuscript writing and reviewing. All authors have read and approved the final manuscript.

Funding

This research received no specific grant from any funding agency in the public, commercial, or not-for-profit sectors.

Availability of data and materials

The datasets generated and/or analyzed during the current study are not publicly available due to the current Cairo University regulations and Egyptian legislation but are available from the corresponding author on reasonable request and after institutional approval.

Declarations

Ethics approval and consent to participate

An informed written consent was taken from each patient. All data obtained from every patient were confidential and were not used outside the study. The patients have rights to withdraw from the study at any time without giving any reason. All the cost of the investigations was afforded by the researcher. Our study was approved by ethical committee of the Department of Neurology, Faculty of Medicine, Cairo University on 12/7/2021.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Author details

1 Neurology Department, Faculty of Medicine, Cairo University, Cairo, Egypt.
2 Present Address: Neuropsychiatry Department, Faculty of Medicine, South Valley University, Qena, Egypt. 3 Neurophysiology Department, Faculty of Medicine, Cairo University, Cairo, Egypt.

Received: 28 August 2021 Accepted: 8 December 2021

Published online: 20 December 2021

References

1. World Health Organization, 2020, director-General’s opening remarks at the media briefing on COVID-19—11 March 2020. https://www.who.int/dg/speeches/detail/who-director-general-s-opening-remarks-at-the-media-briefing-on-covid-19—11-march-2020.

2. Desforges M, Le Coupanec A, Dubeau P, Bourgouin A, Lajoie L, Dubé M, et al. Human coronaviruses and other respiratory viruses: underestimated opportunistic pathogens of the central nervous system. Viruses. 2019;12(1):14. https://doi.org/10.3390/v12010014.
3. Zhao H, Shen D, Zhou H, Liu J. Guillain–Barré syndrome associated with SARS-CoV-2 infection: causality or coincidence. Lancet Neurol. 2020;19(5):583–4.

4. Ng Keekwong KC, Mehta PR, Shukla G, Mehta AR. COVID-19, SARS and MERS. A neurological perspective. J Clin Neurosci. 2020. https://doi.org/10.1016/j.jocn.2020.04.124.

5. Baig AM, Khaleeq A, Ali U, Syeda H. Evidence of the COVID-19 virus targeting the CNS: tissue distribution, host-virus interaction, and proposed neurotropic mechanisms. ACS Chem Neurosci. 2020;11(7):995–8. https://doi.org/10.1021/acschemneuro.0c00122.

6. Natali S, Oliveira V, Calabresi P, Maia LF, Pisani A. Does SARS-CoV-2 invade the brain. Translational lessons from animal models. Eur J Neurol. 2020. https://doi.org/10.1111/ene.14277.

7. Moore JB, June CH. Cytokine release syndrome in severe COVID-19. Science. 2020;368(6490):473–4.

8. Jamilloux Y, Henry T, Belot A, Viel S, Faust M, Jammal T, et al. Should we stimulate or suppress immune responses in COVID-19, cytokine and anti-cytokine interventions. Autoimmun Rev. 2020;19(7):105267.

9. Ye Q, Wang B, Mao J. The pathogenesis and treatment of the Cytokine Storm in COVID-19. J Infect. 2020;80(6):607–13. https://doi.org/10.1016/j.jinf.2020.03.037 (PMID: 32283152; PMCID: PMC7194613).

10. Viriani A, Rabold E, Hainson T, Haag A, Elufayr R, Chema T, et al. Guillain–Barré Syndrome associated with SARS-CoV-2 infection. IDCases. 2020;20:e00771.

11. Yaranagula SD, Koduri VKC. Spectrum of acute neuropathy associated with COVID-19: a clinical and electrophysiological study of 13 patients from a single center. Int J Infect Dis. 2021;110:314–9.

12. Leonhard SE, Mandarakas MR, Gondim FAA, Bateman K, Ferreira ML, Corblath DR, et al. Diagnosis and management of Guillain–Barré syndrome in ten steps. Nat Rev Neurol. 2019;15(11):671–83. https://doi.org/10.1038/s41582-019-0250-9 (PMID: 31541214; PMCID: PMC6821638).

13. da Silva IRF, Frontera JA, Bispo de Filippis AM, Nascimento O. Neurologic complications associated with the Zika virus in Brazilian adults. JAMA Neurol. 2017;74(10):1190–8.

14. Sedaghat Z, Karmi N. Guillain–Barre syndrome associated with COVID-19 infection: a case report. J Clin Neurosci. 2020;76:235–5.

15. El Otmani H, El Moutawakil B, Rafai MA, Benna N, Kettani C, Soussi M, et al. COVID-19 and Guillain–Barré syndrome: more than a coincidence! Rev Neurol. 2020;176(6):518–9.

16. Camdessanche JP, Morel J, Poizot B, Paul S, Tholance Y, Botelho-Nevers E. COVID-19 may induce Guillain–Barré syndrome. Rev Neurol. 2020;176(6):516–8.

17. Scheidt E, Canseco DO, Hadji-Naumov A, Bereznai B, et al. Guillain–Barre syndrome during SARS-CoV-2 pandemic: a case report and review of recent literature. J Peripher Nerv Syst. 2020. https://doi.org/10.1111/jns.12382.

18. Padroni M, Mastrangelo V, Ascoli GM, Pavolucci L, Rumelehe S, Piscaglia M, et al. Guillain–Barre syndrome following COVID-19: new infection, old complication. J Neurol. 2020;24:1–3.

19. Ottaviani D, Bosco F, Tranquillini E, Gapeni I, Pedrotti G, Cozzio S, et al. Early Guillain–Barré syndrome in coronavirus disease 2019 (COVID-19): a case report from an Italian COVID-hospital. Neurol Sci. 2020;41(6):1351–4.

20. Abdelnour L, Abdalla ME, Babiker S. COVID 19 infection presenting as motor peripheral neuropathy. J Formos Med Assoc. 2020;119(6):1119–20.

21. Toscano G, Palmerini F, Ravaglia S, et al. Guillain–Barré syndrome associated with SARS-CoV-2. J Clin Neurosci. 2020. https://doi.org/10.1016/j.jocn.2020.05.016.

22. Diamanti AP, Rosado MM, Pioli C, Sesti G, Bruno LB. Cytokine release syndrome in COVID-19 patients, a new scenario for an old concern: the fragile balance between infections and autoimmunity. Int J Mol Sci. 2020;21(9):E3330.