Short- and long-term outcome and predictors in an international cohort of patients with neuro COVID-19

Ettore Beghi, MD; Raimund Helbok, MD; Serefnur Oztturk, MD; Omer Karadas, MD; Vitalie Lisnic, MD, PhD; Oxana Grosu, MD, Ph.D; Tibor Kovács, MD, PhD; Levente Dobronyi, MD; Daniel Bereczki, MD, PhD; Maria Sofia Cotelli, MD; Marinella Turla, MD; Eugenia Irene Davidescu, MD, PhD; Bogdan Ovidiu Popescu, MD, PhD; Franco Valzania, MD; Franceso Cavallieri, MD; Hanno Ulmer, MBA, PhD; Luis F Maia, MD, PhD; Anne Hege Amodt, MD, PhD; Carmel Armon, MD; Waldemer Brola, MD; Gryb Victoria, MD, PhD; Anis Riahi, MD; Ingomar Krehan MD; Tim von Oertzen, MD; Mohammed A Azab, MD; Michael Crean, Mr; Maria Lolich, PhD; Maria João Lima, MD; Johann Sellner, MD; Julian Perneczky, MD; Tom Jenkins, MD, PhD; Sara Meoni, MD, PhD; Elisa Bianchi, MD; Elena Moro, MD, PhD; Claudio L.A. Bassetti, MD on behalf of the ENERGY Study Group (*).

(*) ENERGY Study Group: Onur Ural & Iskender Kara, Konya, Turkey; Bilgin Ozturk, Ankara, Turkey; Mihail Gavriliuc & Olesea Odainic, Chisinau, Moldova; Patrizia Civelli & Marta Bianchi, Esine-Brescia, Italy; Teodora Bunea & Georgiana Sandu, Bucharest, Romania; Giulia Toschi, Reggio Emilia, Italy; Vanessa Oliveira & Alexandre Dias, Porto, Portugal; Simon Jung & Robert Hoepner, Bern, Switzerland; Marion Boldingh, Oslo, Norway; Netta Agajany & Sharon Wolfson, Zerifin, Israel; Lipowski Michał, Konskie, Poland; Lesiv Marjiana, Ivano-Frankivsk, Ukraine; Hajer Derbali, Tunis, Tunisia; Mafalda Seabra, São João, Brazil; Vanessa Carvalho, Matosinhos, Portugal; Heidi Øyen Flemmen, MD, Skien, Norway; Clarissa Lin Yasuda, Campinas, SP, Brazil; Pille Taba, Tartu, Estonia; Osama Yassin, Alexandria, Egypt; Gordana Kiteva-Trenchevska, Skopje, Macedonia.

Corresponding Author:

Ettore Beghi, MD

Laboratory of Neurological Disorders, Istituto di Ricerche Farmacologiche Mario Negri IRCCS, Via Mario Negri 2, 20156 Milan, Italy. Tel. +39 02 39014542. Fax +39 02 39001916.

e-mail: ettore.beghi@marionegri.it

This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the <u>Version of Record</u>. Please cite this article as <u>doi:</u> 10.1111/ENE.15293

https://orcid.org/0000-0003-2542-0469

Manuscript word count: 3202

Running head: Outcome of Neuro-COVID and predictors

Key-words: COVID-19, neurological disorders, outcome, predictors, SARS-CoV-2.

DR. ETTORE BEGHI (Orcid ID : 0000-0003-2542-0469)
DR. RAIMUND HELBOK (Orcid ID : 0000-0001-5682-0145)
DR. TIBOR KOVÁCS (Orcid ID : 0000-0002-8603-8848)
PROF. DÁNIEL BERECZKI (Orcid ID : 0000-0002-8374-0500)
DR. TIM J VON OERTZEN (Orcid ID : 0000-0003-2164-7842)
DR. MARIA JOÃO LIMA (Orcid ID : 0000-0002-0198-1766)
PROF. JOHANN SELLNER (Orcid ID : 0000-0001-8749-5533)
PROF. ELENA MORO (Orcid ID : 0000-0002-7968-5908)
PROF. CLAUDIO L BASSETTI (Orcid ID : 0000-0002-4535-0245)

Article type : Original Article

Short- and long-term outcome and predictors in an international cohort of patients with neuro COVID-19

Ettore Beghi, MD; Raimund Helbok, MD; Serefnur Oztturk, MD; Omer Karadas, MD; Vitalie Lisnic, MD, PhD; Oxana Grosu, MD, Ph.D; Tibor Kovács, MD, PhD; Levente Dobronyi, MD; Daniel Bereczki, MD, PhD; Maria Sofia Cotelli, MD; Marinella Turla, MD; Eugenia Irene Davidescu, MD, PhD; Bogdan Ovidiu Popescu, MD, PhD; Franco Valzania, MD; Franceso Cavallieri, MD; Hanno Ulmer, MBA, PhD; Luis F Maia, MD, PhD; Anne Hege Amodt, MD, PhD; Carmel Armon, MD; Waldemer Brola, MD; Gryb Victoria, MD, PhD; Anis Riahi, MD; Ingomar Krehan MD; Tim von Oertzen, MD; Mohammed A Azab, MD; Michael Crean, Mr; Maria Lolich, PhD; Maria João Lima, MD; Johann Sellner, MD; Julian Perneczky, MD; Tom Jenkins, MD, PhD; Sara Meoni, MD, PhD; Elisa Bianchi, MD; Elena Moro, MD, PhD; Claudio L.A. Bassetti, MD on behalf of the ENERGY Study Group (*). (*) ENERGY Study Group: Onur Ural & Iskender Kara, Konya, Turkey; Bilgin Ozturk, Ankara, Turkey; Mihail Gavriliuc & Olesea Odainic, Chisinau, Moldova; Patrizia Civelli & Marta Bianchi, Esine-Brescia, Italy; Teodora Bunea & Georgiana Sandu, Bucharest, Romania; Giulia Toschi, Reggio Emilia, Italy; Vanessa Oliveira & Alexandre Dias, Porto, Portugal; Simon Jung & Robert Hoepner, Bern, Switzerland; Marion Boldingh, Oslo, Norway; Netta Agajany & Sharon Wolfson, Zerifin, Israel; Lipowski Michał, Konskie, Poland; Lesiv Marjiana, Ivano-Frankivsk, Ukraine; Hajer Derbali, Tunis, Tunisia; Mafalda Seabra, São João, Brazil; Vanessa Carvalho, Matosinhos, Portugal; Heidi Øyen Flemmen, MD, Skien, Norway; Clarissa Lin Yasuda, Campinas, SP, Brazil; Pille Taba, Tartu, Estonia; Osama Yassin, Alexandria, Egypt; Gordana Kiteva-Trenchevska, Skopje, Macedonia.

Corresponding Author:

Ettore Beghi, MD Laboratory of Neurological Disorders, Istituto di Ricerche Farmacologiche Mario Negri IRCCS, Via Mario Negri 2, 20156 Milan, Italy. Tel. +39 02 39014542. Fax +39 02 39001916. e-mail: ettore.beghi@marionegri.it https://orcid.org/0000-0003-2542-0469

Manuscript word count: 3222

Running head: Outcome of Neuro-COVID and predictors

Key-words: COVID-19, neurological disorders, outcome, predictors, SARS-CoV-2.

Abstract

Background Despite the increasing number of reports on the spectrum of neurological manifestations of COVID-19 (neuro-COVID), few studies have assessed short and long-term outcome of the disease.

Methods This is a cohort study enrolling adult patients with neuro-COVID seen in neurological consultation. Data were collected prospectively or retrospectively in the EAN NEuro-covid ReGistrY. The outcome at discharge was measured using the modified Rankin Scale (mRS) and defined as: "stable/improved" if mRS score was equal or lower than pre-morbid score; "worse" if the score was higher than pre-morbid score. Status at 6 months was also recorded. Demographic and clinical variables were assessed as predictors of outcome at discharge and 6 months. **Results** From July 2020 to March 2021, 971 patients from 19 countries were included. 810 (83.4%) were hospitalized. 432 (53.3%) were discharged with worse functional status. Older age, stupor/coma, stroke and ICU admission were predictors of worse outcome at discharge. 132 (16.3%) died in hospital. Older age, cancer, cardiovascular complications, refractory shock, stupor/coma and ICU admission and degree of functional impairment at discharge were predictors of worse outcome. 65/221 hospitalized patients (29.4%) and 10/32 non-hospitalized patients (24.4%) experienced persisting neurological symptoms/signs. 10/262 patients (3.8%) developed new neurological complaints during the 6 months of follow-up.

Conclusions Neuro-COVID is a severe disease associated with worse functional status at discharge, particularly in older subjects and those with comorbidities and acute complications of infection.

INTRODUCTION

The spectrum of Coronavirus Disease 2019 (COVID-19) includes several neurological manifestations that, when present, are associated with higher severity and worse outcome.¹⁻⁴ However, neurological symptoms, signs and diagnoses in patients with COVID-19 (neuro-COVID) vary according to the target populations, setting (in- vs. outpatients), diagnostic criteria, and the background of those in charge of data collection.⁵ At present, there are only few publications with follow-up, mainly from single center studies⁶ or non-hospitalized patients,⁷ or based on self-reports,⁸ electronic databases,^{9,10} small samples,¹¹ or with short follow-up,^{12,13} or high attrition rates.¹⁴ Thus, available evidence is insufficient to define the full spectrum of neuro-COVID and verify how patient's profile (demographics, baseline clinical features) and acute manifestations of infection predict the outcome of the disease.

On this background, an international registry of patients with COVID-19 and neurological symptoms, signs or diagnoses was established for a better understanding of the disease spectrum, along with risk factors, comorbidities and outcome.¹⁵ The advantage of such a registry is the investigation of a large sample of patients from various countries, from which data on neuro-COVID are collected using uniform diagnostic criteria and standardized methods. The aims of this study were: 1: To compare the outcome of neuro-COVID at hospital discharge and at six months with patient's profile (comorbidities, general and neurological findings during the acute phase) and find outcome predictors; 2: To illustrate the demographic and clinical features of in-patients and outpatients with neuro-COVID from different countries; 3. To define incidence and types of new neurological manifestations after the acute phase.

PATIENTS AND METHODS

A multinational registry of patients with neuro-COVID was activated in May 2020 by the European Academy of Neurology (EAN) to provide epidemiologic data on neurological signs and symptoms in patients with COVID-19 infection reported by neurologists in outpatient services, emergency rooms, and hospital departments (the EAN NEuro-covid ReGistrY, ENERGY). Details on ENERGY structure and organization have been published.¹⁵ Briefly, all neurologists participating in the registry were asked to record neurologic symptoms, signs and diagnoses in clinically or laboratory-confirmed COVID-19 patients in an electronic case record form (e-CRF) (**Supplementary Appendix 1**). Data was collected prospectively or retrospectively and included patient's demographics and lifestyle habits, comorbidities, date of first symptoms of infection, hospital and intensive care unit (ICU) admission, incident general and neurological manifestations during the acute phase, diagnostic tests and outcome. Each variable was reported as "Yes", "No" or

"Unknown". In addition, for each documented neurological manifestation, the local investigator was asked to indicate whether or not it was associated with COVID-19.

All adult patients with symptoms and/or signs and/or diseases requiring neurological consultation were eligible for inclusion. A guide is included in the e-CRF (**Supplementary Appendix 2**) to define each variable and facilitate data collection in the e-CRF at study entry and during follow-up. Registration and follow-up of eligible patients is ongoing.

All registered patients were followed through telephone contacts at 6 and 12 months. At each contact, the modified Rankin score (mRS) was assigned and new neurological manifestations were noted; for patients who died, date of death and, if performed, autopsy are noted. As the mRS is reliable even when applied by telephone,¹⁶ in addition to follow-up we measured functional disability at baseline enquiring patients or caregivers on their pre-morbid functional status. Descriptive statistics were performed on all variables collected during the acute phase in the entire sample and comparing hospitalized and non-hospitalized patients, and prospective and retrospective observations. The outcome of the infection, in terms of functional impairment, was defined as: "stable/improved" if mRS at discharge was equal as or lower than the baseline score; "worse" if mRS score at discharge was higher than the baseline score. Stable/improved and worse outcome were also assessed in patients who deceased during hospital stay compared to those discharged alive.

Neurological symptoms, signs and diagnoses persisting at six months were listed. Demographic and clinical profile of patients with new neurological manifestations occurring during follow-up was illustrated. The same methods were used to assess the effect of variables collected during the acute phase or at discharge on the 6-month outcome.

The association of all variables included in the registry with outcome (worse vs. stable/improved) and status (dead vs. alive) at hospital discharge was evaluated using univariable logistic regression models. Variables identified as statistically significant in univariable models were included in multivariable models, and a stepwise selection (with p<0.05 as criteria for entering and removing effects) was applied to identify variables most strongly associated with the outcome and status at discharge. Results of univariable and multivariable logistic regression models are presented as odds ratios (OR) and adjusted odds ratios (adj.OR) with 95% confidence intervals (CI). Significance was set at the 5% level (0.05).

For demographic and lifestyle variables, mRS and outcome, the number of missing data was reported in the tables, and indicated as "unknown" or missing. For all other variables, "unknown" values were grouped with "No". For neurological findings, the categories "present, not COVID associated" and "present, likely COVID associated" were combined. Data presented as numbers

with percentages or as means with standard deviations or medians and ranges were calculated only in subjects with the corresponding values.

The study was approved by the ethics committees of all participating sites and informed consent was obtained from all eligible patients in line with each participating country's legal requirements.

RÉSULTS

As of March 31st 2021, 1,004 patients were enrolled. COVID-19 infection was not laboratoryconfirmed in 33 cases, which were excluded from further analyses. The final sample included 971 patients from 19 countries (Europe 14; Asia 2; Africa 2; South America 1) (**Table 1**). A flow-chart of the study is illustrated in Figure 1.

With few exceptions, there were no major differences in the general characteristics of prospective compared to retrospective cases (**Supplementary Table 1**). There were 497 men and 466 women (plus eight intersex or unknown) aged 16 to 101 years (median 63; interquartile range, IQR 48-74). One or more comorbidities were present in 619 cases (63.75%). The most frequent was hypertension (52.0%), followed by cardiovascular disease (29.8%) and diabetes (22.0%). History of transient ischemic attacks (TIAs) or stroke (154 cases, 15.9%), dementia (86 cases, 8.9%) and Parkinson's disease (35 cases, 3.6%) were the commonest neurological comorbidities.

810 patients (83.4%) were hospitalized. Compared to non-hospitalized patients, hospitalized patients were older, preferably men, with one or more baseline comorbidities, and with functional impairment at baseline (**Table 1**). Functional disability at admission (mRS 2+) was present in 34.1% of hospitalized patients and 10.6% of non-hospitalized cases. Hospitalized patients experienced more systemic COVID-19 complications and had more neurological manifestations during the acute phase. The most common neurological complaints/manifestations during the acute phase included headache (38.2% in hospitalized and 52.6.9% in non-hospitalized patients), cognitive impairment (31.6% vs. 20.8%), stroke (30.1% vs. 3.9%), delirium (26.7% vs. 10.4%), hyposmia/hypogeusia (24.6% vs. 58.4%), sleep disorders (14.8% vs. 25.4%), myalgias (24.9% vs. 51.9%), and stupor/coma (14.7% vs. 2.6%). Two hundred and 24 patients (27.6%) were admitted to the ICU.

At discharge, the proportion of hospitalized patients with functional impairment (mRS 2+) increased to 62.3% (vs. 14.5% of non-hospitalized subjects). 432 hospitalized patients (53.3%) were discharged with a worse functional status as compared to admission (**Table 2A**). Compared to patients who improved or were stable, patients with worse outcome were older, had more non-neurological (hypertension, cardiovascular and renal diseases) and neurological comorbidities (history of TIAs or stroke, dementia) and presented more systemic complications during the acute

phase. Stroke was the most common neurological manifestation in patients with worse outcome (40.7%) followed by cognitive impairment (34.0%), headache (31.0%), stupor/coma (23.6%), and myalgia (20.8%). In contrast, in patients with stable/improved outcome the commonest manifestations were, in decreasing order, headache (48.2%), hyposmia/hypogeusia (32.3%), myalgias (30.5%), vertigo (23.4%), and cognitive impairment (20.7%).

132 patients died in hospital (**Table 2B**). Compared to those discharged alive, patients who died were older (median age at COVID-19 onset 76 years, IQR 67-85), with more comorbidities (hypertension, cardiovascular and renal diseases, cancer and, among neurological diseases, stroke, dementia and Parkinson's disease). Stupor/coma, stroke and cognitive disturbances were the commonest neurological manifestations/complaints in patients who died in hospital (59.1%, 47.7% and, 49.2%, respectively) along with dysexecutive syndrome (23.5%) and hypoactive delirium (24.2%). Almost all deceased patients presented systemic COVID-19 complications (predominantly pneumonia, 84.9% of cases), 48.5% were admitted in ICU and 36.4% required mechanical ventilation. Refractory shock occurred in 27.3% of in-hospital deaths.

The variable resulting most highly associated with worse outcome was refractory shock (OR 30.6; 95% CI 4.2-224.5) (Table 3A). Increasing age also predicted worse outcome (OR 1.04 for each additional year; 95% CI 1.03-1.05). Among neurological manifestations, stupor/coma (OR 6.2; 95% CI 3.6-10.8) and stroke (OR 3.1; 95% CI 2.2-4.3) showed the highest risk for worse outcome. The need for mechanical ventilation (OR 8.1; 95% CI 4.35-15.0) and ICU admission (OR 5.8; 95% CI 3.9-8.6) indicated worse outcome. Older age, stupor/coma, stroke and ICU admission were confirmed as predictors of worse outcome at discharge in multivariable model, while syncope and dystonia were predictors of stable/improved outcome. In univariable models, stupor/coma carried the highest death risk (OR 21.8; 95% CI 13.6-34.8), followed by cognitive impairment (3.05; 95% CI 2.1-4.5), hypoactive delirium (OR 2.5; 95% CI 1.6-4.0) and stroke (OR 2.4: 95% CI 1.7-3.56). Use of mechanical ventilation (OR 5.1; 95% CI 3.3-7.9), pneumonia (OR 4.3; 95% CI 2.6-7.0), ICU admission (OR 3.0; 95% CI 2.05-4.4), cardiovascular complications (OR 3.0; 95% CI 1.9-4.6) and renal insufficiency (OR 2.9; 95% CI 1.7-5.1) also predicted in-hospital mortality. Among preexisting comorbidities, hypertension (OR 3.8; 95% CI 2.4-6.1), chronic kidney diseases (OR 3.6; 95% CI 2.2-5.8. cardiovascular diseases (OR 2.5; 95% 1.7-3.7) and cancer (OR 2.5; 95% CI 1.5-4.3) were the variables most highly associated with in-hospital death. Among pre-existing neurological comorbidities, stroke (OR 2.6; 95% CI 1.7-4.0) and Parkinson's disease (OR 2.6; 95% CI 1.02-6.5) carried the highest risk. Pre-morbid mRS score was significantly associated with inhospital mortality, showing an increasing risk with the increase of the disability score (Table 3B). Older age, cancer, cardiovascular complications, refractory shock, stupor/coma and ICU admission

were predictors of death in the multivariable model, while hyposmia/hypogeusia predicted a lower risk of death.

At the time of data collection, a total of 269 patients (224 hospitalized and 45 non-hospitalized) had been followed for six months. Of them, 262 had laboratory-confirmed COVID-19 infection. This sample included 131 men and 130 women (unknown sex in 1) aged 19 through 91 years (IQR 47-71) (**Supplementary Table 2**). 199 patients (76.0%) had neurological manifestations during the acute phase of the COVID-19 infection, predominantly headache (40.8%), hyposmia/hypogeusia (34.7%), myalgia (29.1%), delirium (25.2%), cognitive impairment (23.3%), stroke sequelae (21.0%) and sleep disorders (17.1%). A mild-to-severe functional impairment (mRS 2+) was present in 48 patients (18.9%) before the onset of symptoms, in 133 patients (53.8%) at the end of the acute phase of the infection, and in 118 patients (46.1%) at the 6-month follow-up. Almost all the variables associated with worse outcome at discharge were negative prognostic predictors for outcome at 6 months (**Table 4**). Experiencing stroke or ataxia during the acute phase (OR 8.5; 95% CI 2.8-26.1 and, respectively, OR 6.9; 95% CI 1.2-40.7), ICU admission (OR 3.6; 95% CI 1.5-8.7) were confirmed as predictors of worse outcome at 6 months, along with functional impairment at discharge. In contrast, history of stroke was associated with stable/improved outcome (OR 0.3; 95% CI 0.1-0.9).

At 6 months, 65/221 hospitalized patients (29.4%) and 10/41 non-hospitalized patients (24.4%) experienced persisting neurological symptoms/signs, the commonest being hemiparesis/plegia (11 patients), cognitive impairment (10 cases), anosmia/ageusia (10 cases), para/tetraparesis (6 cases) and fatigue (5 cases) (**Figure 2**).

Ten patients (3.8%) developed new neurological complications during follow-up. Two were not hospitalized during the acute phase. The general characteristics of those patients are illustrated in **Supplementary Table 3**. Incident neurological manifestations varied in type and severity. The majority of patients had one or more comorbidities and complications of COVID-19 (mostly pneumonia). Three patients had severe functional impairment at six months. Those patients developed vertical diplopia and, respectively, status epilepticus and recurrent stroke during follow-up. New neurological complaints were more severe in patients with sequelae at hospital discharge.

DISCUSSION

This is the largest international cohort study including 6-month follow up in adult patients with neuro-COVID seen by neurologists. We found that neurological complications are highly prevalent and have a dramatic impact on outcome of hospitalized patients. Further strengthening the relevance of neurological involvement, we found a 76% persistence of neurological involvement with mildto-severe functional impact in 68%.

At admission, one or more comorbidities were present in 63.7% of cases and functional disability was documented in 34.1%; 51.6% of patients experienced systemic complications of SARS-CoV-2 infection, 83.4% were hospitalized, 23.4% were admitted to the ICU, 56.1% had worsening of their functional abilities at discharge, and 16.7% died while in hospital. Stupor or coma, ICU admission and stroke carried a worse outcome at discharge whereas history of cancer, development of cardiovascular complications and refractory shock were associated with increased mortality. Older age and coma were negative prognostic predictors (increased functional disability and death) but did not predict worse outcome at 6-months among survivors at discharge. ICU admission was a negative prognostic factor both at discharge and at 6 months.

At 6 months, 28.6% of patients still presented persistent neurological sequelae of the acute phase, the commonest being focal or generalized motor weakness and cognitive impairment. The development of stroke or ataxia, ICU admission, and functional impairment at discharge were predictive of worse 6-month outcome. These findings support the dispute that only the severity of the acute COVID-19 spectrum and some neurological complications, rather than older age, the presence of comorbidities, and the baseline functional impairment, are significant long-term prognostic predictors.

A number of neurological symptoms or signs (hyposmia/hypogeusia, syncope, dystonia, history of stroke) were associated with stable/improved outcome. However, for some of them (hyposmia/hypogeusia, syncope, dystonia) interview bias is a possible explanation (as more severe cases were perhaps unable to report those symptoms). The protective role of history of stroke cannot be easily interpreted. Although the mechanisms of previous strokes might have been different from COVID-19's mechanism of action, a coincidental finding cannot be excluded. Another study assessed incident neurological symptoms, signs and diagnoses in 4,491 hospitalized patients seen in neurological consultation.¹⁷ In that study, 88% of patients had new neurological manifestations. The most common were toxic/metabolic encephalopathy (51%), stroke (14%), seizures (12%), and hypoxic/ischemic brain injury (11%). In line with us, those patients were older, more severely ill, and less likely to be discharged home.

Our findings differ from other reports. In a large retrospective cohort (N=236,379) using data from an electronic health records network,⁹ the estimated incidence of neurological or psychiatric diagnoses at 6 months following the acute phase of COVID-19 was 33.6% (first diagnosis, 12.8%). The commonest neurological diseases were, in decreasing order, stroke (2.7%), dementia (0.7%), and parkinsonism (0.1%). Our higher rates can be explained by the source of our cases (80%)

hospitalized) and by patients seen in neurological consultation. However, in line with us, the incidence of stroke and dementia were significantly higher in patients with more severe disease. In a prospective study of 4,182 incident cases of COVID-19 who self-reported their symptoms using a mobile application, 558 participants (13.3%) experienced symptoms lasting \geq 28 days, 189 (4.5%) for \geq 8 weeks, and 95 (2.3%) for \geq 12 weeks.⁸ The commonest were fatigue, headache, dyspnea, and anosmia and were more frequent with increasing age, BMI and female sex. The presence of more than five symptoms during the first week of illness was associated with prolonged complaints during follow-up. This is in line with us and suggests that the higher severity of the disease is the consequence of a multisystem involvement by the virus, as shown by others.¹⁸ Post-hospital persistent symptoms (including memory loss [34%], concentration and sleep disorders [28% and 30%]) were reported during phone calls by 279 patients who had COVID-19.⁶ Although some study limitations (single center, inclusion of patients without neurological complaints, high attrition rate) can explain the differences with our findings, the frequent report of cognitive impairment and sleep disorders indicated similarities.

In a study including 1,733 of 2,469 discharged patients with a median follow-up of 6 months, fatigue or muscle weakness (63%) and sleep difficulties (26%) were the commonest persistent symptoms.¹³ Twenty-four percent of cases reported median 6-minute walking distance less than the lower limit of the normal range. Compared to our study, these higher rates might be explained by high attrition (736 patients, perhaps the least severe cases, did not attend follow-up appointments). In a population-based cohort study including non-hospitalized subjects, 938 subjects were invited to participate in a postal survey and 48% responded. Although the interviewees reported reduction of symptoms 1.5-6 months after the acute phase, 16% manifested persisting dyspnea, 12% dysosmia, and 10% dysgeusia.⁷ The differences between this study and ours are reflected, on one side, by our longer follow-up and, on the other side, on the possible underascertainment of non-neurological manifestations in our study or underascertainment of neurological complaints in that study. Sequelae at 6 months were reported in a prospective cohort study by 32.8% of 177 adults recovering from COVID-19.¹¹ The commonest persistent symptoms included fatigue and loss of smell or taste. The lower prevalence of sequelae in our study might reflect the focus on neurological manifestations.

Our study has strengths and limitations. The major strength is the large sample, which includes data from different countries and settings. Another strength is the accurate search and diagnostic assessment of neurological manifestations. All patients were examined by a neurologist and, to optimize inter-rater agreement, diagnoses were guided by standard definitions. Although each

neurological manifestation was investigated based on the findings available at the time of the interview, the e.CRF included precise questions and clinical assessment of the patient was to be completed according to a detailed check-list (Supplementary Appendix 1). The major limitation of our study is the lack of a population base. We tried to define the reference population to estimate incidence and prevalence of the various neurological manifestations. However, the differing catchment areas served by the participating sites did not consent precise calculations. Another important limitation is the focus on neurological manifestations. Eligible patients were those seen in neurological consultation. Although we did our best to collect information on all comorbidities with impact on patients' health and the major complications of COVID-19, our investigation of the full spectrum of the disease has been incomplete. Then, diagnostic accuracy was not always high as a more detailed assessment of registered patients (results of neuropsychological and imaging tests, treatments) was not required to maintain data collection least time consuming, given the emergency context in which neurological consultation was performed. We also opted not to collect data on treatments as they were rarely supported by evidence-based recommendations. Finally, the use of mRS at discharge to predict functional disability at home or in residential or rehabilitation settings could be debated.

In conclusion, in a multinational cohort of patients with neuro-COVID undergoing structured neurological consultation, we found a severe disease in a high proportion of patients. The presence of severe infection with complications predicted worse outcome at discharge, persistence of functional disability, and a number of sequelae at 6 months follow-up, some of those occurring after the remission or stabilization of the acute phase of the disease. Patients with neurological manifestations during the acute phase of COVID-19 infection should be carefully monitored to prevent the occurrence of long-term complications and premature mortality.

ACKNOWLEDGMENTS

The authors are indebted with Sherry H-Y Chou MD and Molly McNett PhD for their valuable intellectual support to the organization of the ENERGY registry, with Lalit Kaltenbach for the preparation of the web-database, with Ms. Anja Sander for her assistance during the activation and conduct of the ENERGY registry, and all patients participating patients.

CONFLICT OF INTEREST

Dr. Beghi reports grants from Italian Ministry of Health, grants from SOBI, personal fees from Arvelle Therapeutics, grants from American ALS Association, outside the submitted work. Dr. Moro reports personal fes from Medtronic, personal fees from Abbott, grants from Boston, outside the submitted work. Dr. Cavallieri reports personal fees from Zambon, outside the submitted work. Dr. Kiteva-Trenchevska reports personal fees from Roche, personal fees from Pliva, personal fees from Medis, outside the submitted work. Dr. Aamodt reports research grants foutside the submitted work from Medtronic and Boehringer Ingelheim and personal fees outside the submitted work from Bayer, Boehringer Ingelheim, Roche, Allergan, Novartis and Teva. Dr. von Oertzen reports personal fees from Liva Nova, grants from Merck, personal fees from Indivior Austria GmbH, personal fees and non-financial support from gtec GmbH Austria, grants, personal fees and non-financial support from Boehringer-Ingelheim, personal fees from Philips, personal fees from UCB Pharma, personal fees from Almirall, grants and personal fees from Eisai, personal feees from Arvelle Threapeuticos, personal fees from GW Pharma, personal fees from Zogenix GmbH, personal fees from Angelini Pharma Österreiche, personal fees from Novartis Pharma GmbH, outside the submitted work; and he is co-chair of the Communication Committee, scientific panel for epilepsy, and Covid taskforce, all of the European Academy of Neurology (EAN), president of the Österreichische Gesellschaft für Epileptologie (Austrian ILAE chapter), and president of the upper Austrian MS society. The other authors have nothing to disclose.

AUTHOR CONTRIBUTIONS

Concept & Design: Ettore Beghi, Raimuld Helbok, Elena Moro, Claudio Bassetti. Acquisition, analysis or interpretation of data; all Authors Drafting of manuscript: Ettore Beghi Critical revision of the manuscript for important intellectual content: [Main authors] Statistical analysis: Elisa Bianchi Obtained funding: Claudio Bassetti Supervision: Main Authors Acquisition of data and critical review of manuscript: ENERGY Study Group

FUNDING

The study was supported by the European Academy of Neurology.

DATA AVAILABILITY STATEMENT

Data can only be shared with permission of individual countries.

REFERENCES

1) Chou SH, Beghi E, Helbok R, et al. Global Incidence of Neurological Manifestations Among Patients Hospitalized With COVID-19-A Report for the GCS-NeuroCOVID Consortium and the ENERGY Consortium. JAMA Netw Open. 2021;4(5):e2112131. doi: 10.1001/jamanetworkopen. 2021.12131.

2) Cagnazzo F, Arquizan C, Derraz I, et al. Neurological manifestations of patients infected with the SARS-CoV-2: a systematic review of the literature. J Neurol. 2020:1-10. doi: 10.1007/s00415-020-10285-9.

3) Favas TT, Dev P, Chaurasia RN, et al. Neurological manifestations of COVID-19: a systematic review and meta-analysis of proportions. Neurol Sci. 2020;41(12):3437-3470. doi: 10.1007/s10072-020-04801-y.

4) Yassin A, Nawaiseh M, Shaban A, et al. Neurological manifestations and complications of coronavirus disease 2019 (COVID-19): a systematic review and meta-analysis. BMC Neurol. 2021;21(1):138. doi: 10.1186/s12883-021-02161-4.

 Beghi E, Michael BD, Solomon T, Westenberg E, Winkler AS; COVID-19 Neuro Research Coalition. Approaches to understanding COVID-19 and its neurological associations. Ann Neurol. 2021 Apr 9. doi: 10.1002/ana.26076.

 Garrigues E, Janvier P, Kherabi Y, et al. Post-discharge persistent symptoms and health-related quality of life after hospitalization for COVID-19. J Infect. 2020;81(6):e4-e6. doi: 10.1016/j.jinf.2020.08.029.

7) Stavem K, Ghanima W, Olsen MK, Gilboe HM, Einvik G. Prevalence and Determinants of Fatigue after COVID-19 in Non-Hospitalized Subjects: A Population-Based Study. Int J Environ Res Public Health. 2021;18(4):2030. doi: 10.3390/ijerph18042030.

8) Sudre CH, Murray B, Varsavsky T, et al. Attributes and predictors of long COVID. Nat Med. 2021;27(4):626-631. doi: 10.1038/s41591-021-01292-y.

9) Taquet M, Geddes JR, Husain M, Luciano S, Harrison PJ. 6-month neurological and psychiatric outcomes in 236 379 survivors of COVID-19: a retrospective cohort study using electronic health records. Lancet Psychiatry. 2021:S2215-0366(21)00084-5. doi: 10.1016/S2215-0366(21)00084-5.

Daugherty SE, Guo Y, Heath K, et al. Risk of clinical sequelae after the acute phase of SARS-CoV-2 infection: retrospective cohort study. BMJ. 2021 May 19;373:n1098.

11) Logue JK, Franko NM, McCulloch DJ, et al. Sequelae in Adults at 6 Months After COVID-19 Infection. JAMA Netw Open. 2021;4(2):e210830. doi: 10.1001/jamanetworkopen.2021.0830. 12) Moreno-Pérez O, Merino E, Leon-Ramirez JM, et al. Post-acute COVID-19 syndrome. Incidence and risk factors: A Mediterranean cohort study. J Infect. 2021 Mar;82(3):378-383.

13) Chopra V, Flanders SA, O'Malley M, Malani AN, Prescott HC. Sixty-Day Outcomes Among Patients Hospitalized With COVID-19. Ann Intern Med. 2021 Apr;174(4):576-578.

14) Huang C, Huang L, Wang Y, et al. 6-month consequences of COVID-19 in patients discharged from hospital: a cohort study. Lancet. 2021;397(10270):220-232. doi: 10.1016/S0140-6736(20)32656-8.

15) Beghi E, Helbok R, Crean M, et al. The European Academy of Neurology COVID-19 registry (ENERGY): an international instrument for surveillance of neurological complications in patients with COVID-19. Eur J Neurol. 2020 Nov 21;10.1111/ene.14652. doi: 10.1111/ene.14652.
16) Savio K, Pietra GL, Oddone E, Reggiani M, Leone MA. Reliability of the modified Rankin Scale applied by telephone.Neurol Int. 2013 Feb 19;5(1):e2. doi: 10.4081/ni.2013.e2.

17) Frontera JA, Sabadia S, Lalchan R, et al. A Prospective Study of Neurologic Disorders in Hospitalized Patients With COVID-19 in New York City. Neurology. 2021 Jan 26;96(4):e575e586. doi: 10.1212/WNL.000000000010979.

18) Dennis A, Wamil M, Alberts J, et alCrooks M, Gabbay M, Brady M, Hishmeh L, Attree E, Heightman M, Banerjee R, Banerjee A, COVERSCAN study investigators. Multiorgan impairment in low-risk individuals with post-COVID-19 syndrome: a prospective, community-based study BMJ Open 2021 Mar 30;11(3):e048391.

Figure 1. Study flow-chart

Figure 2. List of neurological symptoms, signs and diseases persisting at 6 months

		OVID-19 l (n=971)(*)	Hospitaliz	zed (n=810)	Not hospi	talized (n=154)	p-valu
	<u>n</u>	<u>%</u>	n	%	n	%	P
Country							< 0.000
Austria	66	6.80	64	7.90	2	1.30	
Brazil	3	0.31	1	0.12	2	1.30	
Egypt	6	0.62	5	0.62	1	0.65	
Estonia	0	0.00	0	0.00	0	0.00	
France	22	2.27	19	2.35	3	1.95	
Hungary	101	10.40	85	10.49	16	10.39	
Israel	30	3.09	30	3.70	0	0.00	
Italy	165	16.99	96	11.85	69	44.81	
Macedonia	1	0.10	1	0.12	0	0.00	
Moldova	118	12.15	116	14.32	2	1.30	
Norway	50	5.15	39	4.81	11	7.14	
Poland	26	2.68	9	1.11	17	11.04	
Portugal	56	5.77	53	6.54	0	0.00	
Romania	84	8.65	84	10.37	0	0.00	
Russia	13	1.34	7	0.86	6	3.90	
Switzerland	42	4.33	22	2.72	19	12.34	
Tunisia	19	1.96	14	1.73	5	3.25	
Turkey	145	14.93	141	17.41	1	0.65	
Ukraine	24	2.47	24	2.96	0	0.00	
Sex							0.0033
Male	497	51.18	434	53.58	59	38.31	
Female	466	47.99	368	45.43	95	61.69	
Intersex	2	0.21	2	0.25	0	0.00	
Unknown	6	0.62	6	0.74	0	0.00	
Smoking							0.0949
Yes	122	12.56	109	13.46	13	8.44	
No	729	75.08	606	74.81	120	77.92	
Unknown	120	12.36	95	11.73	21	13.64	
Source of COVID-19							0.000
contact							< 0.000
Occupation	75	7.72	42	5.19	33	21.43	
Family member	168	17.30	119	14.69	46	29.87	
Social	86	8.86	66	8.15	18	11.69	
Travel	17	1.75	16	1.98	1	0.65	
Other	61	6.28	61	7.53	0	0.00	

Table 1. Confirmed COVID-19 cases, hospitalized and not hospitalized cases

	Unknown	564	58.08	506	62.47	56	36.36	
		Median	IOD	Median	IOD	Median	IOD	
		(n)	IQR	(n)	IQR	(n)	IQR	
	Age at COVID onset	63 (909)	48-74	66 (751)	52-76	48 (151)	34-61	< 0.0001
	BMI	25 (840)	23-28	26 (706)	23-29	24 (130)	22-28	0.0455
		n	%	n	%	n	%	
	Any comorbidity	619	63.75	565	69.75	49	31.82	< 0.0001
	Hypertension	505	52.01	464	57.28	36	23.38	< 0.0001
	Diabetes Type 1	8	0.82	8	0.99	0	0.00	0.2054
	Diabetes Type 2	206	21.22	191	23.58	14	9.09	< 0.0001
Ż	Cardiovascular disease	289	29.76	269	33.21	17	11.04	< 0.0001
	Chronic kidney disease	88	9.06	87	10.74	1	0.65	< 0.0001
	Chronic liver disease	37	3.81	37	4.57	0	0.00	0.0057
	Chronic pulmonary disease	93	9.58	86	10.62	6	3.90	0.0136
	Anemia	50	5.15	47	5.80	3	1.95	0.0389
	Cancer	85	8.75	80	9.88	4	2.60	0.0055
	Immunosuppressed state	50	5.15	44	5.43	6	3.90	0.3712
	Other non-	222	22.90	216		15	0.74	-0.0001
	neurological comorbidity	232	23.89	216	26.67	15	9.74	<0.0001
	Dementia	86	8.86	79	9.75	6	3.90	0.0275
	Parkinson's disease	35	3.60	21	2.59	14	9.09	0.0001
	Stroke: ICH, ischemic stroke, TIA	154	15.86	147	18.15	6	3.90	< 0.0001
	Multiple sclerosis	47	4.84	19	2.35	28	18.18	< 0.0001
	Motor neuron disease	4	0.41	3	0.37	1	0.65	0.6501
	Neuromuscular disorder	12	1.24	11	1.36	1	0.65	0.4395
	Neuropathy	34	3.50	31	3.83	3	1.95	0.2157
	Other neurological disease	99	10.20	82	10.12	16	10.39	0.8675
	COVID systemic complications	501	51.60	480	59.26	17	11.04	< 0.0001
	Dyspnea	503	51.80	453	55.93	46	29.87	< 0.0001
	Pneumonia	528	54.38	501	61.85	22	14.29	< 0.0001
	Cardiovascular	121	12.46	117	14.44	1	0.65	< 0.0001
	-				8.02			

dialysis							
Coagulation disorder							
/ disseminated	45	4.63	44	5.43	1	0.65	0.0080
intravascular	45	4.05		5.45	1	0.05	0.0080
coagulation							
Refractory shock	38	3.91	37	4.57	0	0.00	0.0183
Extra-corporeal							
membrane	5	0.51	5	0.62	0	0.00	0.3176
oxygenation (ECMO)							
Mechanical	121	12.46	116	14.32	0	0.00	< 0.0001
ventilation	121	12.10	110	11.52	0	0.00	(0.0001
Neurological findings	747	76.93	633	78.15	107	69.48	0.0434
Headache	394	40.58	310	38.27	81	52.60	0.0010
Hyposmia /	291	29.97	199	24.57	90	58.44	< 0.0001
hypogeusia	-						
Dysautonomia	139	14.32	107	13.21	31	20.13	0.0274
Vertigo	194	19.98	159	19.63	32	20.78	0.5409
Myalgia	284	29.25	202	24.94	80	51.95	< 0.0001
Sleep disorders	161	16.58	120	14.81	39	25.39	0.0009
Cognitive impairment							
(including	288	29.66	256	31.60	32	20.78	0.0029
dysexecutive							
syndrome)							
 Hyperactive delirium	122	12.56	111	13.70	10	6.49	0.0163
Hypoactive delirium /	112	11.53	106	13.09	6	3.90	0.0007
acute encephalopathy	10.1	10.55	110	11.50		2 50	0.0001
Stupor / coma	124	12.77	119	14.69	4	2.60	< 0.0001
Syncope	51	5.25	46	5.68	5	3.25	0.1813
Seizures / status	81	8.34	76	9.38	5	3.25	0.0085
epilepticus							
Meningitis /	42	4.33	38	4.69	4	2.60	0.2087
encephalitis Stroke	253	26.06	244	30.12	6	3.90	< 0.0001
Tremor	233 68	7.00	60	50.12 7.41	8	5.19	0.2681
Chorea	1	0.10	1	0.12	0	0.00	0.6556
Dystonia	18	1.85	18	2.22	0	0.00	0.0562
Myoclonus	15	1.54	15	1.85	0	0.00	0.0302
Dyskinesia	12	1.24	9	1.11	3	1.95	0.4301
 Parkinsonism	26	2.68	21	2.59	5	3.25	0.4301
Ataxia	86	8.86	78	9.63	8	5.19	0.0573
Spinal cord disorder	38	3.91	36	4.44	2	1.30	0.0575
	50	5.71	50	т .тт	2	1.50	0.0001

Peripheral neuropathy	92	9.47	85	10.49	7	4.55	0.0150
Other neurological							
findings	121	12.46	106	13.09	15	9.74	0.1859
Hospital admission	810	83.42	810	100.00	0	0.00	< 0.000
ICU admission	227	23.38	224	27.65	0	0.00	< 0.000
Pre-morbid mRS							< 0.000
0	488	52.87	381	48.97	104	75.36	
1	153	16.58	132	16.97	20	14.49	
2	95	10.29	84	10.80	10	7.25	
3	96	10.40	90	11.57	4	2.90	
4	64	6.93	64	8.23	0	0.00	
5	27	2.93	27	3.47	0	0.00	
Missing	48		32		16		
Discharge mRS							< 0.000
0	264	28.12	170	21.49	94	64.83	
1	158	16.83	128	16.18	30	20.69	
2	116	12.35	101	12.77	15	10.34	
3	130	13.84	123	15.55	6	4.14	
4	88	9.37	87	11.00	0	0.00	
5	51	5.43	50	6.32	0	0.00	
6	132	14.06	132	16.69	0	0.00	
Missing	32		19		9		
Outcome							< 0.000
Worse (**)	448	49.12	432	56.10	14	10.07	
Stable/Improved (**)	464	50.88	338	43.90	125	89.93	
Not available	59		40		19		

BMI = Body mass index; **ICH** = Intracerebral haemorrhage; **ICU** = Intensive care unit; **mRS** = Modified Rankin Score; **TIA** = Transient ischemic attack.

(*) Setting was unknown in 7 cases: Portugal 3, Switzerland 1, Turkey 3

(**) Worse outcome: mRS score at discharge higher than pre-morbid mRS score; Stable/improved outcome:

mRS score at discharge equal or lower than pre-morbid mRS score.

		1	4				В	3		
		orse *) (n=432)		mproved *) (n=338)	p-value	Alive ((n=659)	Dead ((n=132)	p-value
	n	%	n	%		n	%	n	%	_
Country					< 0.0001					< 0.0001
Austria	35	8.10	22	6.51		52	7.89	10	7.58	
Brazil	0	0.00	1	0.30		1	0.15	0	0.00	
Egypt	0	0.00	1	0.30		5	0.76	0	0.00	
Estonia	0	0.00	0	0.00		0	0.00	0	0.00	
France	15	3.47	3	0.89		14	2.12	4	3.03	
Hungary	28	6.48	56	16.57		75	11.38	10	7.58	
Israel	8	1.85	22	6.51		24	3.64	6	4.55	
Italy	70	16.20	23	6.80		69	10.47	26	19.70	
Macedonia	0	0.00	1	0.30		1	0.15	0	0.00	
Moldova	97	22.45	19	5.62		89	13.51	27	20.45	
Norway	23	5.32	7	2.07		31	4.70	0	0.00	
Poland	4	0.93	3	0.89		7	1.06	0	0.00	
Portugal	35	8.10	18	5.33		44	6.68	9	6.82	
Romania	51	11.81	33	9.76		56	8.50	28	21.21	
Russia	2	0.46	5	1.48		7	1.06	0	0.00	
Switzerland	6	1.39	5	1.48		17	2.58	1	0.76	
Tunisia	10	2.31	4	1.18		11	1.67	3	2.27	
Turkey	48	11.11	92	27.22		132	20.03	8	6.06	
Ukraine	0	0.00	23	6.80		24	3.64	0	0.00	
Sex					0.1716					0.6373

 Table 2. Outcome at discharge (only confirmed and hospitalized COVID-19 cases)

Yes 40 9.26 67 19.82 100 15.17 8 6.06 No 336 77.78 240 71.01 492 74.66 101 76.52 Unknown 56 12.96 31 9.17 67 10.17 23 17.42 Source of COVID- 19 contact 0.0002 0.0002 1 0.0002 1 0.76 Social 13 3.01 25 7.40 39 5.92 1 0.76 Family member 50 11.57 61 18.05 103 15.63 12 9.09 Social 31 7.18 34 10.06 51 7.74 14 10.61 Travel 9 2.08 6 1.78 15 2.28 0 0.00 Other 444 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 65.97 196 57.99 413											
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Male	237	54.86	170	50.30		349	52.96	69	52.27	
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	Female	193	44.68	162	47.93		302	45.83	63	47.73	
Smoking <0.0001 0.000 Yes 40 9.26 67 19.82 100 15.17 8 6.06 No 336 77.78 240 71.01 492 74.66 101 76.52 Unknown 56 12.96 31 9.17 67 10.17 23 17.42 Source of COVID- 19 contact 0.0002 0.0002 1 0.76 1 0.76 Social 13 3.01 25 7.40 39 5.92 1 0.76 Family member 50 11.57 61 18.05 103 15.63 12 9.09 Social 31 7.18 34 10.06 51 7.74 14 10.61 Travel 9 2.08 6 1.78 15 2.28 0 0.00 Other 44 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 <td>Intersex</td> <td>1</td> <td>0.23</td> <td>1</td> <td>0.30</td> <td></td> <td>2</td> <td>0.30</td> <td>0</td> <td>0.00</td> <td></td>	Intersex	1	0.23	1	0.30		2	0.30	0	0.00	
Yes 40 9.26 67 19.82 100 15.17 8 6.06 No 336 77.78 240 71.01 492 74.66 101 76.52 Unknown 56 12.96 31 9.17 67 10.17 23 17.42 Source of COVID- 19 contact 0.0002 0.0002 1 0.0002 1 0.000 Social 13 3.01 25 7.40 39 5.92 1 0.76 Family member 50 11.57 61 18.05 103 15.63 12 9.09 Social 31 7.18 34 10.06 51 7.74 14 10.61 Travel 9 2.08 6 1.78 15 2.28 0 0.00 Other 444 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 65.97 196 57.99 413 <td>Unknown</td> <td>1</td> <td>0.23</td> <td>5</td> <td>1.48</td> <td></td> <td>6</td> <td>0.91</td> <td>0</td> <td>0.00</td> <td></td>	Unknown	1	0.23	5	1.48		6	0.91	0	0.00	
No 336 77.78 240 71.01 492 74.66 101 76.52 Unknown 56 12.96 31 9.17 67 10.17 23 17.42 Source of COVID- 19 contact	Smoking					< 0.0001					0.0027
Unknown 56 12.96 31 9.17 67 10.17 23 17.42 Source of COVID- 19 contact 0.0002	Yes	40	9.26	67	19.82		100	15.17	8	6.06	
Source of COVID- 19 contact 0.0002 colspan="6">colspan="6" colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6">colspan="6" colspan="6" colspa="6" colspa="6" colspan="6" colspan="6" colspan="6" colspan="6"	No	336	77.78	240	71.01		492	74.66	101	76.52	
0.0002 -	Unknown	56	12.96	31	9.17		67	10.17	23	17.42	
19 contact Occupation 13 3.01 25 7.40 39 5.92 1 0.76 Family member 50 11.57 61 18.05 103 15.63 12 9.09 Social 31 7.18 34 10.06 51 7.74 14 10.61 Travel 9 2.08 6 1.78 15 2.28 0 0.00 Other 44 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 65.97 196 57.99 413 62.67 83 62.88 Median IQR Median IQR Median IQR Median IQR $Age at COVID$ 70 (405) 60-79 58 (308) 43-71 <0.0001	Source of COVID-										
Family member 50 11.57 61 18.05 103 15.63 12 9.09 Social 31 7.18 34 10.06 51 7.74 14 10.61 Travel 9 2.08 6 1.78 15 2.28 0 0.00 Other 44 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 65.97 196 57.99 413 62.67 83 62.88 Median IQR Median IQR Median IQR Median IQR (n) 70 (405) 60-79 58 (308) 43-71 <0.0001	19 contact					0.0002					< 0.0001
Social 31 7.18 34 10.06 51 7.74 14 10.61 Travel 9 2.08 6 1.78 15 2.28 0 0.00 Other 44 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 65.97 196 57.99 413 62.67 83 62.88 Median IQR Median IQR Median IQR Median IQR Median (n) 10 10 57.99 413 62.67 83 62.88 62.88 Median IQR Median	Occupation	13	3.01	25	7.40		39	5.92	1	0.76	
Travel92.0861.78152.2800.00Other4410.19164.73385.772216.67Unknown28565.9719657.9941362.678362.88Median (n)IQRMedian (n)IQRMedian (n)IQRMedian (n)IQRMedian (n)IQRAge at COVID onset70 (405)60-7958 (308)43-71<0.0001	Family member	50	11.57	61	18.05		103	15.63	12	9.09	
Other 44 10.19 16 4.73 38 5.77 22 16.67 Unknown 285 65.97 196 57.99 413 62.67 83 62.88 Median (n) IQR Median (n) IQR <th< td=""><td>Social</td><td>31</td><td>7.18</td><td>34</td><td>10.06</td><td></td><td>51</td><td>7.74</td><td>14</td><td>10.61</td><td></td></th<>	Social	31	7.18	34	10.06		51	7.74	14	10.61	
Unknown 285 65.97 196 57.99 413 62.67 83 62.88 Median (n) IQR (n) Median (n) Median (n) </td <td>Travel</td> <td>9</td> <td>2.08</td> <td>6</td> <td>1.78</td> <td></td> <td>15</td> <td>2.28</td> <td>0</td> <td>0.00</td> <td></td>	Travel	9	2.08	6	1.78		15	2.28	0	0.00	
Median (n) IQR Median (n) IQR Median (n) IQR Median (n) IQR Median (n) IQR Age at COVID onset 70 (405) 60-79 58 (308) 43-71 <0.0001	Other	44	10.19	16	4.73		38	5.77	22	16.67	
IQR IQR IQR IQR IQR IQR IQR Age at COVID onset 70 (405) 60-79 58 (308) 43-71 <0.0001	Unknown	285	65.97	196	57.99		413	62.67	83	62.88	
n % n % n % n % Any comorbidity 335 77.55 202 59.76 <0.0001 63 (606) 49-73 76 (127) 67-85 <0 Hypertension 289 66.90 158 46.75 <0.0001 348 52.81 107 81.06 <0			IQR		IQR			IQR		IQR	
n % n % n % n % Any comorbidity 335 77.55 202 59.76 <0.0001	-	70 (405)	60-79	58 (308)	43-71	<0.0001	63 (606)	49-73	76 (127)	67-85	<0.0001
Any comorbidity 335 77.55 202 59.76 <0.0001	BMI	26 (379)	23-29	25 (302)	23-28	0.0135	26 (573)	23-28	25 (117)	22-29	0.6456
Hypertension 289 66.90 158 46.75 <0.0001 348 52.81 107 81.06 <0		n	%	n	%		n	%	n	%	
	Any comorbidity	335	77.55	202	59.76	< 0.0001	434	65.86	118	89.39	< 0.0001
Diabetes Type 1 3 0.69 5 1.48 0.2865 8 1.21 0 0.00 0.	Hypertension	289	66.90	158	46.75	< 0.0001	348	52.81	107	81.06	< 0.0001
	Diabetes Type 1	3	0.69	5	1.48	0.2865	8	1.21	0	0.00	0.2033

Diabetes Type 2	112	25.93	71	21.01	0.1114	149	22.61	40	30.30	0.0585
Cardiovascular disease	165	38.19	92	27.22	0.0014	195	29.59	68	51.52	< 0.0001
Chronic kidney disease	57	13.19	28	8.28	0.0309	54	8.19	32	24.24	< 0.0001
Chronic liver disease	14	3.24	20	5.92	0.0728	30	4.55	7	5.30	0.7093
Chronic pulmonary disease	47	10.88	35	10.36	0.8148	68	10.32	15	11.36	0.7207
Anemia	22	5.09	20	5.92	0.6171	34	5.16	13	9.85	0.0375
Cancer	46	10.65	29	8.58	0.3368	53	8.04	24	18.18	0.0003
Immunosuppressed state	20	4.63	19	5.62	0.5334	33	5.01	8	6.06	0.6184
Other non-										
neurological	145	33.56	63	18.64	< 0.0001	153	23.22	60	45.45	< 0.0001
comorbidity										
Dementia	49	11.34	27	7.99	0.1214	53	8.04	26	19.70	< 0.0001
Parkinson's disease	14	3.24	7	2.07	0.3227	14	2.12	7	5.30	0.0381
Stroke: ICH, ischemic stroke, TIA	98	22.69	44	13.02	0.0006	102	15.48	43	32.58	<0.0001
Multiple sclerosis	3	0.69	13	3.85	0.0023	18	2.73	1	0.76	0.1764
Motor neuron disease	0	0.00	2	0.59	0.1094	3	0.46	0	0.00	0.4374
Neuromuscular	5	1.16	4	1.18	0.9734	9	1.37	1	0.76	0.5681

disorder

Neuropathy	15	3.47	13	3.85	0.7833	26	3.95	5	3.79	0.9322
Other neurological disease	38	8.80	37	10.95	0.3179	60	9.10	17	12.88	0.1818
COVID systemic complications	312	72.22	151	44.67	<0.0001	350	53.11	122	92.42	<0.0001
Dyspnea	270	62.50	163	48.22	< 0.0001	335	50.83	105	79.55	< 0.0001
Pneumonia	306	70.83	172	50.89	< 0.0001	374	56.75	112	84.85	< 0.0001
Cardiovascular	76	17.59	39	11.54	0.0193	79	11.99	38	28.79	< 0.0001
Renal insufficiency / dialysis	46	10.65	17	5.03	0.0048	42	6.37	22	16.67	<0.0001
Coagulation disorder / disseminated intravascular coagulation	28	6.48	13	3.85	0.106	33	5.01	9	6.82	0.3971
Refractory shock	36	8.33	1	0.30	< 0.0001	1	0.15	36	27.27	<0.0001
Extra-corporeal membrane oxygenation (ECMO)	4	0.93	0	0.00	0.0761	4	0.61	0	0.00	0.3695
Mechanical ventilation	99	22.92	12	3.55	<0.0001	66	10.02	48	36.36	<0.0001
Neurological findings	377	87.27	230	68.05	<0.0001	498	75.57	125	94.70	<0.0001

Headache	134	31.02	163	48.22	< 0.0001	279	42.34	23	17.42	< 0.0001
Hyposmia /	79	18.29	109	32.25	< 0.0001	184	27.92	7	5.30	<0.0001
hypogeusia	19	10.29	107	52.25	<0.0001	104	21.92	1	5.50	<0.0001
Dysautonomia	50	11.57	54	15.98	0.0761	93	14.11	13	9.85	0.1893
Vertigo	72	16.67	79	23.37	0.02	140	21.24	14	10.61	0.0048
Myalgia	90	20.83	103	30.47	0.0022	184	27.92	12	9.09	< 0.0001
Sleep disorders	63	14.58	51	15.09	0.8446	105	15.93	14	10.61	0.1181
Cognitive										
impairment										
(including	157	36.34	86	25.44	0.0012	183	27.77	67	50.76	< 0.0001
dysexecutive										
syndrome)										
Hyperactive	76	17.59	32	9.47	0.0013	86	13.05	23	17.42	0.1833
delirium	10	11.05	52	5.17	0.0010	00	10.00	25	17.12	0.1000
Hypoactive										
delirium / acute	76	17.59	29	8.58	0.0003	74	11.23	32	24.24	< 0.0001
encephalopathy										
Stupor / coma	102	23.61	16	4.73	< 0.0001	41	6.22	78	59.09	< 0.0001
Syncope	14	3.24	30	8.88	0.0008	41	6.22	4	3.03	0.1485
Seizures / status	37	9.50	20	11.24	0.2136	62	0.41	14	10.61	0.6600
epilepticus	57	8.56	38	11.24	0.2130	62	9.41	14	10.01	0.6699
Meningitis /	19	4.40	19	5.62	0.4368	33	5.01	5	3.79	0.5498
encephalitis	17	1.10	17	5.02	0.1000	55	5.01	5	5.17	0.0170
Stroke	176	40.74	62	18.34	< 0.0001	180	27.31	63	47.73	< 0.0001
Tremor	24	5.56	32	9.47	0.038	54	8.19	5	3.79	0.0786

Chorea	0	0.00	1	0.30	0.2579	1	0.15	0	0.00	0.6543
Dystonia	2	0.46	15	4.44	0.0002	17	2.58	1	0.76	0.2001
Myoclonus	8	1.85	7	2.07	0.8271	12	1.82	3	2.27	0.7283
Dyskinesia	2	0.46	7	2.07	0.0394	7	1.06	2	1.52	0.6543
Parkinsonism	9	2.08	12	3.55	0.2149	20	3.03	1	0.76	0.1374
Ataxia	34	7.87	43	12.72	0.026	72	10.93	6	4.55	0.0248
Spinal cord	19	4.40	16	4.73	0.8244	34	5.16	2	1.52	0.0667
disorder	19	4.40	10	4.75	0.8244	54	5.10	2	1.52	0.0007
Peripheral	49	11.34	26	7.69	0.09	73	11.08	7	5.30	0.0466
neuropathy	77	11.54	20	1.09	0.07	15	11.00	7	5.50	0.0400
Other neurological	62	14.25	37	10.95	0.1613	88	12.25	16	10.10	0.7021
findings	62	14.35	57	10.95	0.1015	88	13.35	16	12.12	0.7021
ICU admission	180	41.67	37	10.95	< 0.0001	157	23.82	64	48.48	< 0.0001
Pre-morbid mRS					0.6792					< 0.0001
0	206	48.02	163	48.95		331	52.29	38	29.46	
1	78	18.18	53	15.92		118	18.64	13	10.08	
2	50	11.66	32	9.61		64	10.11	18	13.95	
3	48	11.19	41	12.31		66	10.43	23	17.83	
4	35	8.16	29	8.71		39	6.16	25	19.38	
5	12	2.80	15	4.50		15	2.37	12	9.30	
Missing	3		5			26		3		
Discharge mRS					< 0.0001					< 0.0001
0	0	0.00	170	50.30		170	25.80	0	0.00	
				10.01		100	10.42	0	0.00	
1	58	13.43	62	18.34		128	19.42	0	0.00	
1 2	58 57	13.43 13.19	62 38	18.34 11.24		128	19.42 15.33	0	0.00	

4	61	14.12	22	6.51	87	13.20	0	0.00
5	36	8.33	11	3.25	50	7.59	0	0.00
6	132	30.56	0	0.00	0	0.00	132	100.00

(*) Worse outcome: mRS score at discharge higher than pre-morbid mRS score; Stable/improved outcome: mRS score at discharge equal or lower than premorbid mRS score.

BMI = Body mass index; **ICH** = Intracerebral haemorrhage; **ICU** = Intensive care unit; **mRS** = Modified Rankin Score; **TIA** = Transient ischemic attack.

Table 3. Predictors of outcome at discharge (N = 971)

A. Worse outcome vs. Stable/improved outcome (*) B. Dead vs. Alive at discharge Univariable model Multivariable model Univariable model

This article is protected by copyright. All rights reserved

Multivariable model

	OR	95% CI	p-value	adj. OR	95% CI	p-value	OR	95% CI	p-value	adj. OR	95% CI	p-value
Sex			0.1432						0.9611			
Male	1 (ref.)						1 (ref.)					
Female	0.86	0.64-1.14					1.05	0.73-1.54				
Intersex/Unknown	0.24	0.05-1.20					ne	ne				
Smoking			0.0001						0.0039			
Yes	0.43	0.28-0.65					0.39	0.18-0.83				
No	1 (ref.)	-					1 (ref.)	-				
Unknown	1.29	0.81-2.06					1.67	0.99-2.81				
Source of COVID-19 contact			0.0007						0.0471			
Occupation	0.53	0.35-0.80					0.50	0.27-0.94				
Family member	0.33	0.16-0.67					0.11	0.02-0.81				
Social	0.59	0.35-0.99					1.18	0.63-2.21				
Travel	0.97	0.34-2.76					ne	ne				
Other/Unknown	1 (ref.)	-					1 (ref.)	-				
Age at admission (1 year	1.04	1 02 1 05	< 0.0001	1.03	1.02-1.04	< 0.0001	1.00	1.05.1.09	< 0.0001	1.05	1.03-1.07	<0.0001
increase)	1.04	1.03-1.05	<0.0001	1.05	1.02-1.04	<0.0001	1.06	1.05-1.08	<0.0001	1.05	1.05-1.07	<0.0001
BMI (1 unit increase)	1.01	0.99-1.03	0.2869				0.99	0.97-1.02	0.5206			
Non neurological												
comorbidities												
Hypertension	2.30	1.72-3.09	< 0.0001				3.82	2.41-6.07	< 0.0001			
Diabetes Type 1/Type 2	1.25	0.90-1.74	0.1878				1.39	0.92-2.10	0.1171			
Cardiovascular disease	1.65	1.21-2.25	0.0014				2.53	1.73-3.70	< 0.0001			
Chronic kidney disease	1.68	1.05-2.71	0.0324				3.59	2.21-5.83	< 0.0001			
Chronic liver disease	0.53	0.27-1.07	0.0770				1.17	0.50-2.73	0.7095			
Chronic pulmonary disease	1.06	0.67-1.68	0.8154				1.11	0.61-2.02	0.7208			
Anemia	0.85	0.46-1.59	0.6174				2.01	1.03-3.92	0.0409			

Cancer	1.27	0.78-2.07	0.3345	2.54	1.51-4.29	0.0005	2.57	1.23-5.34	0.0117
Immunosuppressed state	0.82	0.43-1.55	0.5341	1.22	0.55-2.71	0.6189			
Neurological comorbidities									<u> </u>
Dementia	1.47	0.90-2.41	0.1232	2.81	1.68-4.68	< 0.0001			<u> </u>
Parkinson's disease	1.58	0.63-3.97	0.3267	2.58	1.02-6.52	0.0451			
Stroke: ICH, ischemic	1.96	1.33-2.89	0.0007	2.64	1.73-4.02	< 0.0001			
stroke, TIA	1.90	1.55-2.89	0.0007	2.64	1.75-4.02	<0.0001			
Multiple sclerosis	0.18	0.05-0.62	0.0068	0.27	0.04-2.05	0.2068			
Motor neuron disease	ne	ne	0.9796	ne	ne	0.9837			
Neuromuscular disorder	0.97	0.26-3.67	0.9734	0.55	0.07-4.39	0.5737			
Neuropathy	0.90	0.42-1.92	0.7822	0.96	0.36-2.54	0.9330			
COVID systemic									
complications									
Dyspnea	1.79	1.34-2.39	< 0.0001	3.76	2.40-5.90	< 0.0001			
Pneumonia	2.34	1.74-3.16	< 0.0001	4.27	2.59-7.04	< 0.0001			
Cardiovascular	1.64	1.08-2.48	0.0201	2.97	1.90-4.63	< 0.0001	2.08	1.07-4.06	0.0311
Renal insufficiency / dialysis	2.25	1.27-4.00	0.0058	2.94	1.69-5.11	0.0001			
Coagulation disorder /									
disseminated intravascular	1.73	0.88-3.40	0.1099	1.39	0.65-2.98	0.3986			
coagulation									
Refractory shock	30.63	4.18-224.56	0.0008	ne	ne	< 0.0001	44.72	5.68-352.5	0.0003
Extra-corporeal membrane	ne	ne	0.9818	ne	ne	0.9877			
oxygenation (ECMO)									
Mechanical ventilation	8.08	4.35-14.99	<0.0001	5.13	3.31-7.94	< 0.0001			
Neurological findings									
Headache	0.48	0.36-0.65	<0.0001	0.29	0.18-0.46	< 0.0001			
Hyposmia / hypogeusia	0.47	0.34-0.66	<0.0001	0.15	0.07-0.32	< 0.0001	0.12	0.04-0.40	0.0006

Dysautonomia	0.69	0.46-1.04	0.0772				0.67	0.36-1.23	0.1919				
Vertigo	0.66	0.46-0.94	0.0205				0.44	0.25-0.79	0.0059				
Myalgia	0.60	0.43-0.83	0.0023				0.26	0.14-0.48	< 0.0001				
Sleep disorders	0.96	0.64-1.43	0.8444				0.63	0.35-1.13	0.1209				
Cognitive impairment													
(including dysexecutive	1.67	1.22-2.29	0.0013				2.68	1.83-3.93	< 0.0001				
syndrome)													
Hyperactive delirium	2.04	1.31-3.17	0.0015				1.40	0.85-2.33	0.1846				
Hypoactive delirium / acute	2.28	1.44-3.58	0.0004				2.53	1.59-4.03	< 0.0001				
encephalopathy	2.28	1.44-5.56	0.0004				2.33	1.39-4.03	<0.0001				
Stupor / coma	6.22	3.59-10.77	< 0.0001	12.01	4.35-33.11	< 0.0001	21.77	13.62-34.81	< 0.0001	22.77	12.1-42.86	< 0.0001	
Syncope	0.34	0.18-0.66	0.0013	0.10	0.03-0.31	< 0.0001	0.47	0.17-1.34	0.1576				
Seizures / status epilepticus	0.74	0.46-1.19	0.2149				1.14	0.62-2.11	0.6701				
Meningitis / encephalitis	0.77	0.40-1.48	0.4379				0.75	0.29-1.95	0.5514				
Stroke	3.06	2.19-4.28	< 0.0001	2.89	1.88-4.44	< 0.0001	2.43	1.66-3.56	< 0.0001				
Tremor	0.56	0.33-0.98	0.0402				0.44	0.17-1.13	0.0865				
Chorea	ne	ne	0.9780				ne	ne	0.9858				
Dystonia	0.10	0.02-0.44	0.0024	0.02	0.00-0.14	< 0.0001	0.29	0.04-2.19	0.2288				
Myoclonus	0.89	0.32-2.48	0.8263				1.25	0.35-4.51	0.7288				
Dyskinesia	0.22	0.05-1.07	0.0600				1.43	0.30-6.98	0.6555				
Parkinsonism	0.58	0.24-1.39	0.2202				0.24	0.03-1.83	0.1704				
Ataxia	0.59	0.37-0.94	0.0273				0.39	0.17-0.91	0.0301				
Spinal cord disorder	0.93	0.47-1.83	0.8238				0.28	0.07-1.19	0.0853				
Peripheral neuropathy	1.54	0.93-2.53	0.0919				0.45	0.20-1.00	0.0499				
ICU admission	5.81	3.93-8.59	< 0.0001	5.62	3.54-8.95	< 0.0001	3.01	2.05-4.43	< 0.0001	2.17	1.18-4.00	0.0130	
Pre-morbid mRS			0.6842						< 0.0001				
0	1 (ref.)	-					1 (ref.)	-					

1	1.16	0.77-1.75	0.96	0.49-1.86
2	1.24	0.76-2.02	2.45	1.32-4.56
3	0.93	0.58-1.47	3.04	1.70-5.43
4	0.96	0.56-1.63	5.58	3.06-10.22
5	0.63	0.29-1.39	6.97	3.04-15.98

(*) Worse outcome: mRS score at discharge higher than pre-morbid mRS score; Stable/improved outcome: mRS score at discharge equal or lower than premorbid mRS score.

BMI = Body mass index; **ICH** = Intracerebral haemorrhage; **ICU** = Intensive care unit; **mRS** = Modified Rankin Score; **TIA** = Transient ischemic attack.

Table 4. Predictors of outcome at 6-months (N = 262)

	Worse outcome vs. Stable/improved outcome (*)											
	U	nivariable mo	del	Mu	ltivariable n	odel						
	OR	95% CI	p-value	adj. OR	95% CI	p-value						
Sex			0.9979									
Male	1 (ref.)											
Female	1.02	0.62-1.67										
Intersex/Unknown	ne	ne										
Smoking			0.5947									
Yes	0.80	0.35-1.84										
No	1 (ref.)	-										
Unknown	1.63	0.52-5.14										
Source of COVID-19 contact			0.0008									
Occupation	0.23	0.10-0.50										
Family member	0.36	0.17-0.78										
Social	0.36	0.13-1.02										
Travel	0.79	0.23-2.71										
Other/Unknown	1 (ref.)	-										
Age at admission (1 year increase)	1.03	1.02-1.05	0.0001									
BMI (1 unit increase)	0.99	0.99-1.00	0.6458									
Non neurological												
comorbidities												
Hypertension	1.96	1.19-3.23	0.0085									
Diabetes Type 1/Type 2	1.26	0.66-2.40	0.4771									
Cardiovascular disease	2.25	1.22-4.15	0.0095									
Chronic kidney disease	7.88	1.75-35.45	0.0071									

Chronic liver disease	0.76	0.17-3-45	0.7181			
Chronic pulmonary disease	0.95	0.45-2.01	0.8851			
Anemia	2.08	0.51-8.52	0.3070			
Cancer	0.47	0.19-1.15	0.0996			
Immunosuppressed state	1.02	0.29-3.60	0.9796			
Neurological comorbidities						
Dementia	0.32	0.08-1.22	0.0955			
Parkinson's disease	0.67	0.11-4-09	0.6664			
Stroke: ICH, ischemic stroke,	2.09	1.01-4.31	0.0475	0.27	0.08-0.91	0.0302
TIA	2.08	1.01-4.51	0.0475	0.27	0.08-0.91	0.0302
Multiple sclerosis	0.25	0.03-2.25	0.2153			
Motor neuron disease	ne	ne	-			
Neuromuscular disorder	1.54	0.25-9.36	0.6410			
Neuropathy	0.33	0.03-3.25	0.3443			
COVID-19 systemic						
complications						
Dyspnea	1.13	0.69-1.86	0.6225			
Pneumonia	1.34	0.81-2.20	0.2501			
Cardiovascular	3.84	1.37-10.76	0.0105			
Renal insufficiency / dialysis	4.19	1.35-13.01	0.0131			
Coagulation disorder /						
disseminated intravascular	2.83	0.73-10.91	0.1317			
coagulation						
Refractory shock	ne	ne	0.9907			
Extra-corporeal membrane	1.02	0.14-7.33	0.9872			
oxygenation (ECMO)	1.02	0.14-7.33	0.9012			
Mechanical ventilation	2.77	1.30-5.88	0.0081			

Neurological	findings
--------------	----------

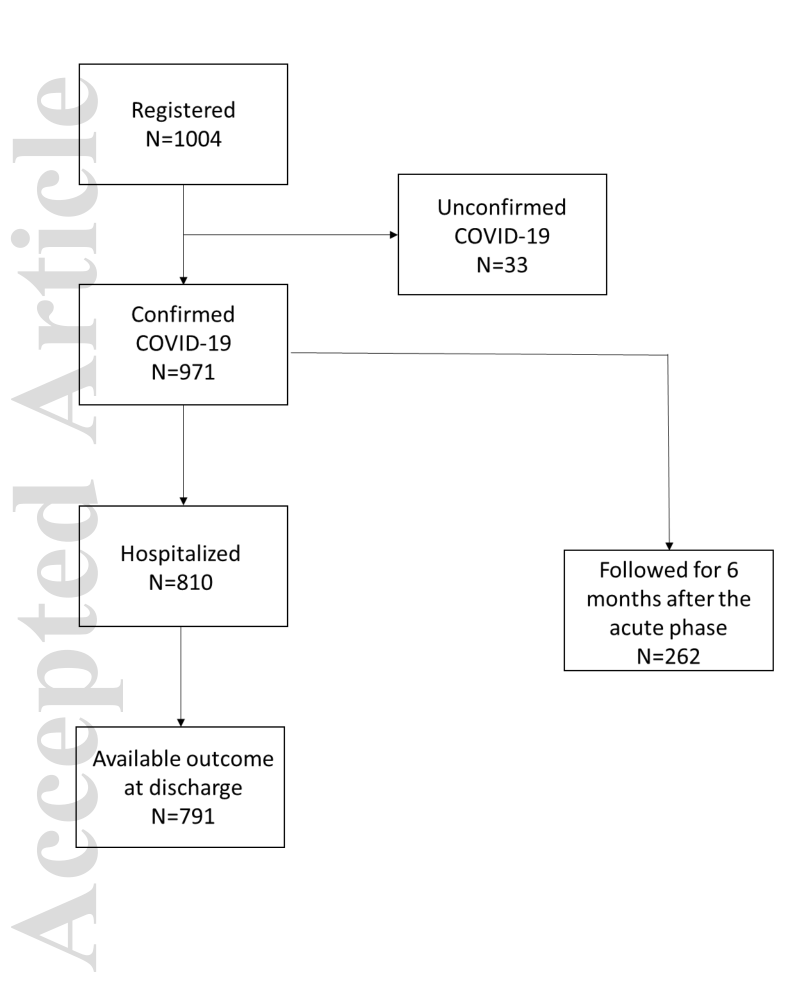
Inticle

i teur orogreur innungs						
Headache	0.71	0.43-1.18	0.1925			
Hyposmia / hypogeusia	0.52	0.30-0.88	0.0152			
Dysautonomia	1.22	0.53-2.84	0.6393			
Vertigo	1.27	0.66-2.44	0.4711			
Myalgia	0.95	0.55-1.63	0.8451			
Sleep disorders	1.14	0.59-2.18	0.1209			
Cognitive impairment						
(including dysexecutive	2.09	1.17-3.74	0.0125			
syndrome)						
Hyperactive delirium	1.65	0.83-3.28	0.1542			
Hypoactive delirium / acute	3.07	1.24-7.60	0.0151			
encephalopathy	5.07	1.24-7.00	0.0131			
Stupor / coma	5.43	1.17-25.32	0.0311			
Syncope	0.67	0.11-4.09	0.6664			
Seizures / status epilepticus	0.74	0.30-1.83	0.5196			
Meningitis / encephalitis	1.37	0.30-6.23	0.6869			
Stroke	6.35	3.02-13.36	< 0.0001	8.51	2.77-26.13	0.0007
Tremor	1.82	0.52-6.39	0.3475			
Chorea	ne	ne	0.8997			
Dystonia	ne	ne	0.9906			
Myoclonus	1.02	0.06-16.43	0.9910			
Dyskinesia	ne	ne	0.9907			
Parkinsonism	ne	ne	0.9790			
Ataxia	6.03	1.31-27.78	0.0212	6.94	1.18-40.68	0.0180
Spinal cord disorder	1.72	0.40-7.36	0.4635			
Peripheral neuropathy	1.84	0.81-4.20	0.1462			

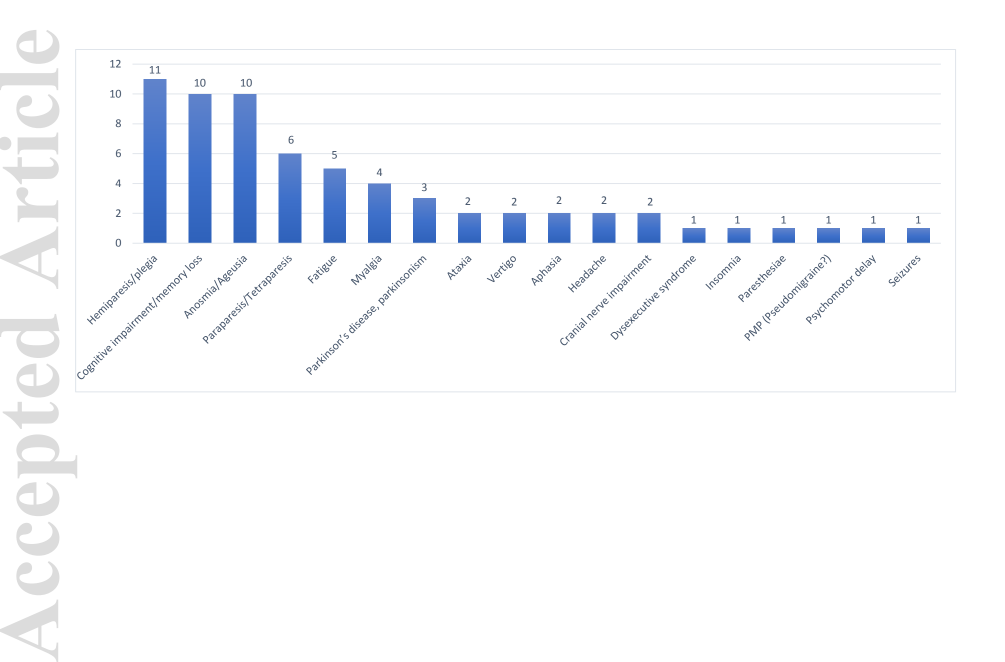
ICU admission	6.39	3.19-12.79	< 0.0001	3.59	1.49-8.66	0.0017
Pre-morbid mRS			0.8153			
0	1 (ref.)	-				
1	1.01	0.53-1.92				
2	0.65	0.23-1-79				
3	0.82	0.30-2.24				
4	0.37	0.07-1.97				
5	0.69	0.15-3.20				
Discharge mRS			< 0.0001			< 0.0001
0	1 (ref.)	-		1 (ref)	-	
1	11.60	3.96-33.97		6.71	2.03-22.15	
2	19.33	6.34-58.98		12.96	3.60-46.66	
3	39.44	12.44-125.0		21.46	5.92-77.73	
4	28.35	8.55-93.98		19.38	4.87-77.14	
5	50.26	10.64-237.4		23.64	4.32-129.3	

(*) Worse outcome: mRS score at discharge higher than pre-morbid mRS score; Stable/improved outcome: mRS score at discharge equal or lower than premorbid mRS score.

BMI = Body mass index; **ICH** = Intracerebral haemorrhage; **ICU** = Intensive care unit; **mRS** = Modified Rankin Score; **TIA** = Transient ischemic attack.



ene_15293_f2.docx



MANAGE-PD

Tool for Making Informed Decisions to Aid Timely Management of Parkinson's Disease

MANAGE-PD allows you to:

- Identify PD patients inadequately controlled on oral medications
- Determine which patients with PD may be adequately controlled on their current treatment regimen or may require changes to their treatment regimen



Scan the QR code to access to the web

abbvie

Click here to access to the web

MANAGE-PD is an AbbVie Inc. registered Medical Device. It is a collaborative research and development effort between AbbVie Medical Affairs and Health Economics and Outcomes, the Parkinson's Foundation and an international panel of Movement Disorder Specialists.

©2022 AbbVie Inc. All rights reserved. The Parkinson's Foundation logo is the sole property of the Parkinson's Foundation used with written permission. Any use of the Parkinson's Foundation name or logo without Foundation permission is prohibited. All content in https://www.managepd.eu/is intended only for informational use by healthcare professionals and is not offered as or intended to be medical advice for any particular patient. This information is not intended for patients. Only a healthcare professional exercising independent clinical judgement can make decisions regarding appropriate patient care and treatment options considering the unique characteristics of each patient.

PD: Parkinson's Disease

