Disorders of Consciousness Associated With COVID-19: A Prospective, Multimodal Study of Recovery and Brain Connectivity

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Abstract

Background and Objectives: In patients with severe coronavirus disease 2019 (COVID-19), disorders of consciousness (COVID-DoC) have emerged as a serious complication. The prognosis and pathophysiology of COVID-DoC remain unclear, complicating decisions about continuing life-sustaining treatment. We describe the natural history of COVID-DoC and investigate its associated brain connectivity profile.

Methods: In a prospective, longitudinal study, we screened consecutive patients with COVID-19 at our institution. We enrolled critically ill adult patients with a DoC unexplained by sedation or structural brain injury, and who were planned to undergo a brain MRI. We performed resting state functional MRI and diffusion MRI to evaluate functional and structural connectivity, as compared to healthy controls and patients with DoC resulting from severe traumatic brain injury (TBI). We assessed the recovery of consciousness (command-following) and functional outcomes (Glasgow Outcome Scale Extended [GOSE] and the Disability Rating Scale [DRS]) at hospital discharge, three
months post-discharge, and six months post-discharge. We also explored whether clinical variables were associated with recovery from COVID-DoC.

Results: After screening 1,105 patients with COVID-19, we enrolled twelve with COVID-DoC. The median age was 63.5 years [interquartile range 55-76.3]. Excluding one who died shortly after enrollment, all of the remaining eleven patients recovered consciousness, after 0-25 days (median 7 [5-14.5]) following the cessation of continuous intravenous sedation. At discharge, all surviving patients remained dependent – median GOSE 3 [1-3], median DRS 23 [16-30]. However ultimately, except for two patients with severe polyneuropathy, all returned home with normal cognition and minimal disability – at three months, median GOSE 3 [3-3], median DRS 7 [5-13]; at six months, median GOSE 4 [4-5], median DRS 3 [3-5]. Ten patients with COVID-DoC underwent advanced neuroimaging; functional and structural brain connectivity in COVID-DoC was diminished compared to healthy controls, and structural connectivity was comparable to patients with severe TBI.

Discussion: Patients who survived invariably recovered consciousness after COVID-DoC. Though disability was common following hospitalization, functional status improved over the ensuing months. While future research is necessary, these prospective findings inform the prognosis and pathophysiology of COVID-DoC.

Introduction

Months into the coronavirus disease 2019 (COVID-19) pandemic, neurologic manifestations of the disease were recognized. Impaired consciousness was observed in 1-20% of patients with COVID-19, mostly in patients with severe infection and comorbidities. It soon became evident that these disorders of consciousness in severe COVID-19 (COVID-DoC) may be prolonged, carrying an unclear prognosis for neurologic recovery. This uncertainty has had profound implications. For families and surrogates already unsure about whether to continue intensive medical care for their loved ones, COVID-DoC has often prompted discussions about withdrawing life-sustaining treatment. To assist in these challenging decisions, some institutions have built dedicated services for deliberating the probability of neurologic recovery (i.e., “Coma Boards”) and have utilized ethics consultations. However, the uncertain prognosis of COVID-DoC raises the alarming possibility of error in decisions about life-sustaining treatment – continuation for patients with little chance of meaningful recovery, or withdrawal for patients who would have otherwise recovered.

Data on recovery from prolonged COVID-DoC are slowly emerging. In July 2020, we described a patient with COVID-DoC, who recovered consciousness ~40 days after sedation was discontinued. In November 2020, another group reported a patient with
COVID-DoC who recovered consciousness after two months.\textsuperscript{10} A case series in December 2020 subsequently described six patients with COVID-DoC who similarly recovered consciousness after 8-31 days.\textsuperscript{11} While these reports indicate that recovery of consciousness after COVID-DoC is possible, their susceptibility to selection bias makes the likelihood of recovery uncertain. Recovery from COVID-DoC has not been evaluated in a prospective cohort, and longer-term functional outcomes, which are crucial to guiding discussions of prognosis, have not been assessed.

The pathophysiology of COVID-DoC remains similarly unclear. DoC caused by other etiologies of brain injury, whether traumatic, anoxic, or cerebrovascular, are characterized by diminished neural connectivity – both functional connectivity as measured with resting state functional magnetic resonance imaging (rs-fMRI)\textsuperscript{12–14}, and structural connectivity as measured with diffusion MRI (d-MRI)\textsuperscript{15–18}. Whether COVID-DoC is characterized by similar connectivity disruptions is unknown. We reported intact functional connectivity in one patient with COVID-DoC, but these findings may not be generalizable.\textsuperscript{9} Preliminary d-MRI studies have found diminished white matter integrity in patients with COVID-19.\textsuperscript{10,19} Investigating brain connectivity in a larger cohort of patients with COVID-DoC, and comparing their connectivity to that of patients with other types of DoC, may shed light on the pathophysiology of this condition.

We launched a prospective, longitudinal, multimodal study to characterize long-term recovery from COVID-DoC and evaluate its brain connectivity profile (ClinicalTrials.gov NCT04476589). Given the urgent need for information to guide discussions of prognosis and decisions about life-sustaining treatment, we provide data from an initial patient cohort, additionally addressing short-term recovery. We aim to (1) identify the demographic and clinical features of patients with COVID-DoC, (2) describe the recovery of consciousness and longer-term function after COVID-DoC, (3) evaluate
functional network connectivity in COVID-DoC as compared to healthy controls, (4) evaluate structural connectivity in COVID-DoC as compared to healthy controls, and as compared to patients with DoC due to severe traumatic brain injury (TBI) – a condition known to disrupt white matter tracts\textsuperscript{15,16,18} – and (5) explore variables potentially associated with recovery from COVID-DoC.
Methods

Patients

Between July 2020 and March 2021, we consecutively screened all patients admitted or transferred to any intensive care unit at Massachusetts General Hospital (given that reported instances of COVID-DoC occurred in critically ill patients). Inclusion criteria were: (1) age 18 or greater, (2) positive severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) assay; (3) DoC (coma, vegetative state or minimally conscious state, based on behavioral examinations in the medical record, per the Aspen Neurobehavioral Workgroup criteria\textsuperscript{20}); and (4) the treating team’s intention to investigate the etiology of the patient’s DoC with a clinical brain MRI, to which research sequences could be appended. Patients were excluded for doses of sedation or structural brain injury on computed tomography sufficient to explain the patient’s DoC, as determined both by the treating team and a team of investigators trained in neurocritical care (DF and BLE). Patients were not excluded for neuroimaging findings associated with COVID-19 (specifically, microhemorrhages and leukoencephalopathy\textsuperscript{21,22}), given their uncertain effects on consciousness.

Once patients were stable for transport, clinical MRI sequences as well as rs-fMRI and d-MRI were acquired. Although sedation during the scan was minimized when possible, some patients received sedatives for safety, comfort and immobility. Though the clinical applications of functional MRI remain uncertain, guidelines have begun to suggest its judicious use in DoC diagnosis and prognostication.\textsuperscript{23,24} Therefore, rs-fMRI data were rapidly processed and reported to the clinical teams and patient families in accordance with an Institutional Review Board-approved protocol.
Neuroimaging

MRI data were acquired with a 32-channel head coil on a 3T Skyra MRI scanner (Siemens Healthcare, Germany). The neuroradiologists’ interpretation of the clinical sequences were reviewed. Reports of microhemorrhages (or microthrombi, which are often indistinguishable) and leukoencephalopathy, neuroimaging findings associated with COVID-19\textsuperscript{21,22}, were included in the analyses below.

Rs-fMRI measures spontaneous fluctuations in the blood-oxygen-level-dependent (BOLD) signal, a proxy for neuronal activity, and evaluates whether such fluctuations correlate across spatially disparate brain regions, a proxy for network connectivity.\textsuperscript{25} The rs-fMRI sequence was obtained with an echo time of 30.3 ms, a repetition time of 1250 ms, and simultaneous multi-slice acquisition (SMS=4) to optimize spatial and temporal resolution (parameters and analytic code provided at www.github.com/ComaRecoveryLab/COVID-19_rsfMRI).\textsuperscript{9} Images were normalized to Montreal Neurological Institute (MNI) space, denoised, and processed with the CONN toolbox software\textsuperscript{26}, using previously described parameters.\textsuperscript{27}

Functional network connectivity was quantified in two ways. Given its well-established association with DoC and neurologic recovery, we first assessed connectivity between nodes of the default mode network (DMN) – i.e., intra-network connectivity – measured as the average correlation between each pair of nodes, as done previously.\textsuperscript{13,14,28} Because a negative correlation between resting state networks may similarly carry prognostic value\textsuperscript{14}, we also assessed connectivity between the DMN and salience network (SN) – i.e., inter-network connectivity – measured as the average correlation between each DMN node and each SN node. Previously described DMN and SN nodes\textsuperscript{29} were used as regions of interest, including 10 DMN nodes and 23 SN nodes (10 mm spheres for cortical nodes and 4 mm spheres for subcortical nodes). For
qualitative visualization in figures, rs-fMRI data are presented as seed-to-voxel maps, with seeds defined as four principal nodes of the DMN – the medial prefrontal cortex, the posterior cingulate cortex, and the bilateral inferior parietal lobules – as done previously.\textsuperscript{9,27}

D-MRI evaluates the integrity of axonal pathways by measuring the diffusion of water along white matter tracts. We acquired high-angular-resolution diffusion imaging, which uses a multidirectional model to optimize resolution of crossing and branching axons\textsuperscript{15,30}, and processed images with FSL (FMRIB, UK), with parameters described previously.\textsuperscript{31} Fractional anisotropy (FA), which represents the directionality of water diffusion in each voxel of the brain (with higher values suggestive of greater white matter integrity, or structural connectivity), has been associated with neurologic recovery after brain injury.\textsuperscript{17} We therefore first measured the average FA of white matter across the whole brain. Second, given evidence of brainstem dysfunction in COVID-DoC\textsuperscript{19,32}, we measured the average FA within the brainstem. See eMethods, http://links.lww.com/WNL/B660 for details.

The imaging analysis was conducted after clinical data were abstracted from the medical record, to avoid the possibility that the connectivity results could bias the interpretation of the clinical data. Though the investigators conducting the imaging analysis were not blinded to the clinical data, the analysis procedure described above was uniform across subjects and required no manual adjustment or subjective interpretation.

We compared functional and structural connectivity of patients with COVID-DoC to that of fourteen healthy control subjects, recruited and scanned with an identical MRI protocol as part of a separate research study approved by the Institutional Review Board (ClinicalTrials.gov NCT03504709). Healthy control subjects had no history of
neurological/psychiatric disease, diabetes, hypertension, heart disease, or kidney disease. One healthy control did not complete the d-MRI sequence. To determine if patients with COVID-DoC exhibit similar connectivity impairments to other etiologies of DoC, we compared FA values from the COVID-DoC cohort to that of eighteen patients with DoC due to severe TBI, recruited and scanned with an identical d-MRI sequence as part of a separate study (eTable 1, http://links.lww.com/WNL/B660). Functional connectivity was not compared due to differences in the rs-fMRI sequences. Ages of the healthy control subjects ranged from 22-52 (median 32.5, interquartile range [IQR] 28.3-37.3), and the ages of the TBI cohort ranged from 19-51 (median 27, IQR 22.5-32.8).

Outcome Assessment

The primary outcome measures established on clinicaltrials.gov (NCT04476589) were the Glasgow Outcome Scale Extended (GOSE; where lower values indicate higher levels of disability) and the Disability Rating Scale (DRS; where higher values indicate higher levels of disability). These measures were assessed at hospital discharge via evaluations by physical and occupational therapists (either measured directly by therapists, or in cases where therapists did not complete these measures, estimated based on documentation of patient function). These measures were assessed again prospectively at three and six months (+/- 4 days) following hospital discharge based on patient interviews, with an interpreter when necessary. If patients were still admitted to an inpatient rehabilitation facility at the time of these post-discharge assessments, scores were also informed by records from physical and occupational therapists at the facility.

Here we also focus on earlier metrics of recovery. We evaluated whether consciousness was recovered (as indicated by chart documentation of command-
following after COVID-DoC), time to recover consciousness (TTRC; measured as the number of days from the cessation of continuous intravenous sedatives [propofol, midazolam, hydromorphone, fentanyl, or ketamine] to the first documented report of command-following), hospital length of stay, discharge disposition, and mortality. Of note, because the criteria for enrollment included the minimally conscious state, two patients already demonstrated intermittent command-following by the time of enrollment; for these two patients, the day of consciousness recovery occurred prior to enrollment.

Primary Analysis

We used two-sample t-tests to compare functional and structural connectivity measures between patients with COVID-DoC and healthy controls. Given that patients tended to be older than controls, and that connectivity metrics may change with age\textsuperscript{36,37}, we also used a linear regression model with terms for condition (patient versus control) and age, to determine if the main effect for condition remained statistically significant after controlling for age. We used two-sample t-tests to compare the d-MRI measures between patients with COVID-DoC and patients with TBI. We applied a Bonferroni correction to account for multiple comparisons. To ensure that t-test results were not driven by outliers, all comparisons were repeated with nonparametric Mann-Whitney \textit{U} tests.

Exploratory Analyses

We conducted exploratory analyses to determine if any clinical factors or biomarkers were associated with recovery of consciousness. Due to low variability in whether consciousness was recovered, we instead used TTRC as the outcome metric.
for these analyses. A Spearman correlation coefficient was computed between TTRC and several clinical variables, including age, medical comorbidities, COVID-19 treatment, degree of hypoxemia, renal dysfunction, liver dysfunction, and presence of microhemorrhages or leukoencephalopathy (see eMethods, http://links.lww.com/WNL/B660).

Though patients were excluded for the ongoing administration of sedatives sufficient to explain the DoC, the cumulative effect of previous sedatives may contribute to COVID-DoC. Thus, a Spearman correlation coefficient was computed between TTRC and the cumulative amounts of midazolam-equivalents, morphine-equivalents, propofol, ketamine, and dexmedetomidine administered from intubation to cessation of continuous intravenous sedation, and from cessation of continuous intravenous sedation to recovery of consciousness (for patients who received intravenous sedatives before transfer to our institution, values were estimated based on available reports). For each patient, we also evaluated whether the recovery of consciousness occurred before or after the estimated elimination of sedative medications (see eMethods, http://links.lww.com/WNL/B660).

We computed the Spearman correlation coefficients between the rs-fMRI/d-MRI metrics and TTRC, and between the rs-fMRI metrics and time from rs-fMRI to recovery of consciousness. For these neuroimaging metrics, we used two-sample t-tests to assess whether they differed between patients with and without microhemorrhages, and between patients with and without leukoencephalopathy. Given the exploratory nature of these analyses and our limited sample size, we did not correct them for multiple comparisons.

**Standard Protocol Approvals, Registrations, and Patient Consents**
The study protocol was approved by the Mass General Brigham Institutional Review Board. We obtained informed consent for research from all participants, or surrogate decision-makers for participants, in this study. Study information can be found at ClinicalTrials.gov (NCT04476589).

Data Availability

Anonymized data not published within this article will be made available by request from any qualified investigator.

Results

Patients and outcomes

We screened 1,105 consecutive patients admitted to the hospital with COVID-19. Patients who died in the hospital (n=98, 8.8%), had a negative SARS-CoV-2 test on confirmatory testing (n=22, 2.0%), were younger than 18 years old (n=9, 0.8%), or were not admitted to an intensive care unit (n=880, 79.6%) were excluded. Of the remaining 185 adult patients with COVID-19 admitted to any intensive care unit, we identified 33 (17.8%) with a DoC unexplained by sedative medications. Of those 33, four (12.1%) had a brain injury sufficient to explain the DoC (one with anoxia caused by cardiac arrest, two with large strokes and malignant edema, and one with aneurysmal subarachnoid hemorrhage and hydrocephalus), nine (27.3%) recovered from the DoC
prior to enrollment, four (12.1%) were too medically unstable for enrollment, three (9.1%) had a DoC attributed to renal failure and thus were not planned to undergo a brain MRI, and one was eligible but not enrolled.

We prospectively enrolled the remaining twelve patients (see eTable 2, http://links.lww.com/WNL/B660). Patient ages ranged from 33-82 (median 63.5 years, IQR 55-76.3). Five were male (42%). The distribution of race and ethnicity paralleled that of other patients hospitalized with COVID-19 across the United States. At the time of enrollment, two were comatose (17%), eight were in a vegetative state (67%), and two were in a minimally conscious state (17%). At admission, eleven carried a diagnosis of hypertension (92%), eight of obesity (67%), eight of diabetes (67%), seven of hyperlipidemia (58%), and five of asthma (42%). All patients were intubated for acute respiratory distress syndrome (ARDS) due to COVID-19 (nine severe [75%], two moderate [17%], one mild [8%]) and remained hospitalized for 27-65 days (median 49.5, IQR 36.8-62). Four patients had microhemorrhages, another three had leukoencephalopathy, and another two had both. Of the six patients with microhemorrhages, three had a mild burden (<5 microhemorrhages; patients 8, 10 and 11) and three a severe burden (>100 microhemorrhages; patients 1, 7 and 9); see Figure 1 and eFigure 1, http://links.lww.com/WNL/B660. Eleven patients underwent a clinical EEG – all demonstrated slowing, four demonstrated generalized rhythmic delta activity, and one demonstrated seizures (generalized periodic discharges up to 3 Hz) that resolved prior to enrollment.

Of the 152 patients in an intensive care unit with COVID-19, but without a DoC, we identified twelve who were age-matched to the patients with COVID-DoC (+/- 5 years) and regained command-following within 24 hours of the cessation of continuous intravenous sedation. See eTable 3, http://links.lww.com/WNL/B660 for details. This
age-matched cohort had similar comorbidities and ARDS severity. Only one underwent a clinical brain MRI (to assess for anoxia after a brief cardiac arrest), which was normal, without evidence of microhemorrhages or leukoencephalopathy. A post-hoc two-sample t-test showed that patients with COVID-DoC spent more days under intravenous sedation than patients with severe COVID-19 but no DoC (COVID-DoC=21.6 [Standard deviation=7.3], COVID-no-DoC=12 [SD=5.5], t[22]=3.65, p<0.005).

Of the twelve patients with COVID-DoC, one died due to medical complications of COVID-19 shortly after enrollment, before the MRI scan was obtained. Another patient improved neurologically shortly after enrollment, and research sequences were not obtained. For the remaining ten patients who underwent an MRI, nine required sedation at the time of the scan to facilitate safety and comfort: two were on a low-dose sedative infusion, two received a single dose of intravenous sedative prior to the scan, and the remaining five were on an enteral regimen of intermittent sedatives to prevent iatrogenic opioid or benzodiazepine withdrawal. See eTable 2, http://links.lww.com/WNL/B660 for details of sedation at enrollment and during the MRI.

A summary of patient outcomes is presented in Figure 1, eFigure 1, http://links.lww.com/WNL/B660 and eTable 4, http://links.lww.com/WNL/B660. Excluding the patient who died shortly after enrollment, all of the remaining eleven patients recovered consciousness. Ten recovered consciousness while still hospitalized, and one after discharge. The TTRC after the cessation of continuous sedation ranged from 0 to 25 days (median 7, IQR 5-14.5). Four patients died during the hospitalization, all due to medical complications of COVID-19; no deaths were attributable to the withdrawal of life-sustaining treatment based on a poor neurologic prognosis. Two were discharged to a long-term acute care facility and six to an inpatient rehabilitation facility.
A summary of neurologic outcomes is presented in Figure 2. By discharge, GOSE scores ranged from 1-3 (median 3, IQR 1-3), and DRS scores ranged from 9-30 (median 23, IQR 16-30); all patients who survived demonstrated cognitive impairment and required significant assistance with activities of daily living at hospital discharge. All patients who survived to discharge (eight) were evaluated three months later, when GOSE scores ranged from 3-6 (median 3, IQR 3-3), and DRS scores ranged from 0-18 (median 7, IQR 5-13). By three months, three patients (those with the longest DoC) remained in an inpatient rehabilitation facility, requiring supervision and support for weakness and cognitive impairment, and five returned home with normal or nearly normal cognition and mild disability due to weakness and pain. All eight patients were again evaluated at six months post-discharge; six lived at home with normal cognition and minimal or no disability, and two, whose recovery was complicated by severe polyneuropathy, remained in an inpatient living facility for more constant support; GOSE scores ranged from 3-6 (median 4, IQR 4-5) and DRS scores ranged from 0-18 (median 3, IQR 3-5). See eTable 4, http://links.lww.com/WNL/B660 for details.

Neuroimaging results

Compared to healthy controls, patients with COVID-DoC exhibited significantly reduced (i.e., less positive) intra-network connectivity within the DMN (COVID-DoC=0.12 [SD=0.06], healthy control=0.23 [SD=0.07], t[22]=3.88, p<0.001), and significantly reduced (i.e., less negative) inter-network connectivity between the DMN and SN (COVID-DoC=0.04 [SD=0.03], healthy control=-0.003 [SD=0.04], t[22]=2.86, p<0.01); see Figure 3. Both findings remained significant after Bonferroni correction (p<0.025) and with Mann-Whitney U tests. After controlling for age, COVID-DoC...
remained significantly associated with reduced intra-network \((b=-0.15, \ p<0.01)\) and inter-network connectivity \((b=0.06, \ p<0.05)\).

Compared to healthy controls, patients with COVID-DoC exhibited reduced whole brain FA \((\text{COVID-DoC}=0.51 \ [SD=0.02], \ \text{healthy control}=0.59 \ [SD=0.02], \ t[21]=9.06, \ p<1 \times 10^{-8})\) and brainstem FA \((\text{COVID-DoC}=0.53 \ [SD=0.04], \ \text{healthy control}=0.59 \ [SD=0.02], \ t[21]=4.27, \ p<0.001)\); see Figure 4. Both findings remained significant after Bonferroni correction \((p<0.0125)\) and with Mann-Whitney \(U\) tests. After controlling for age, COVID-DoC remained significantly associated with reduced whole brain FA \((b=-0.09, \ p<0.05)\) and brainstem FA \((b=-0.06, \ p<0.005)\). Whole brain FA and brainstem FA in COVID-DoC were comparable to that of patients with severe TBI \((p>0.05)\). See eTable 5, http://links.lww.com/WNL/B660 for details and eTable 6, http://links.lww.com/WNL/B660 for FA within specific white matter tracts. Though the limited sample and heterogenous sedation regimens precluded formal statistical testing, there was no appreciable trend towards lower connectivity values in patients on higher amounts of sedation during the MRI (eFigure 2, eFigure 3, http://links.lww.com/WNL/B660).

Exploratory results

The clinical variables assessed did not demonstrate a significant correlation with TTRC. Of them, the presence of renal dysfunction at the time of intravenous sedation cessation demonstrated the strongest trend, where patients with renal dysfunction trended towards a longer TTRC \((\text{Spearman } r[9]=0.61, \ p=0.06)\). Neither intra-network functional connectivity, inter-network functional connectivity, whole brain FA nor brainstem FA correlated with TTRC or time from MRI to recovery of consciousness \((p>0.05)\). These neuroimaging metrics did not differ between patients with \((n=6)\) and
without \( (n=5) \) microhemorrhages, or between patients with \( (n=5) \) and without \( (n=6) \) leukoencephalopathy \( (p>0.05) \), nor were they correlated with the degree of hypoxemia \( (p>0.05) \). Patients with a severe burden of microhemorrhages trended towards a longer TTRC (26, 6, and 4 days) as compared to patients with a mild burden of microhemorrhages (6, 0 and 0 days).

Of the eleven patients who recovered consciousness, seven did so while feasibly still under the effect of sedation, based on the timing and estimated half-lives of sedatives they received. The other four patients, even with conservative half-life estimates for critically ill patients with prolonged exposures, did not recover consciousness for 2-20 days following the elimination of all sedatives. See eTable 7, http://links.lww.com/WNL/B660 for details of cumulative patient sedation.
Discussion

Here we report findings from a cohort of patients with COVID-DoC – the first prospective study of this condition. Notably, all patients who survived severe COVID-19 recovered consciousness. This observation corroborates prior reports that consciousness recovery from COVID-DoC is possible,\(^9\)\(^–\)\(^11\) and indicates that such recovery is highly likely. Moreover, enrolling patients prospectively, soon after patients developed an unexplained DoC, revealed that the weeks- or months-long DoC described in previous reports is not common in COVID-DoC, with 50% of patients regaining consciousness within a week after continuous sedation was discontinued.

While patients were typically of advanced age, with comorbidities such as hypertension, diabetes, and asthma, such demographics characterize patients susceptible to severe COVID-19 in general\(^40\), and may not indicate a susceptibility to COVID-DoC in particular (an age-matched cohort of patients with severe COVID-19 but no DoC had similar comorbidities). Similarly, while there was a high prevalence of structural neuroimaging findings associated with COVID-19 – 82% of patients demonstrated microhemorrhages and/or leukoencephalopathy – it is unclear if these findings cause, or predispose to, COVID-DoC. Given that the prevalence of these findings in severe COVID-19 is uncertain, and that these findings may be attributable to
alternative etiologies (e.g., cerebral amyloid angiopathy, coagulopathy from critical illness, or leukoaraiosis), it is challenging to confirm their association with COVID-DoC. We note that the severity of microhemorrhages and leukoencephalopathy varied between patients, and while patients with numerous microhemorrhages tended to demonstrate a longer TTRC than patients with few microhemorrhages, there was no clear link between severity and recovery: some patients with severe findings had brief COVID-DoC with favorable outcomes (e.g., patient 7 in eFigure 1, http://links.lww.com/WNL/B660), while other patients with no such findings had prolonged COVID-DoC with protracted disability (e.g., patient 3 in eFigure 1, http://links.lww.com/WNL/B660). Ultimately, given that some patients with COVID-DoC exhibited neither microhemorrhages nor leukoencephalopathy, we can conclude that such findings are not necessary for COVID-DoC.

Some have speculated that COVID-DoC may result from the high doses of sedation required to treat severe COVID-19. Patients with COVID-DoC were indeed sedated longer than those with severe COVID-19 but no DoC, but sedatives alone may not necessarily explain the differences between these two small cohorts, given potential confounds such as illness severity. While some patients with COVID-DoC may have been impacted by lingering sedatives, others remained unresponsive long after sedatives were likely eliminated. Thus, while sedation may contribute to COVID-DoC, our findings suggest that sedation alone is not sufficient to explain all instances of COVID-DoC.

The heterogeneity of clinical characteristics observed in this patient cohort suggests that the etiology of COVID-DoC is likely multifactorial. Structural brain injury, medical causes of encephalopathy, systemic inflammation (as suggested by a recent study), hypoxemia and the lingering effect of sedation may all play a role, to varying
degrees, in COVID-DoC. Though we found no variables consistently associated with TTRC, we note that there was a trend towards an association with renal failure, a known complication of COVID-19 that can cause encephalopathy.\textsuperscript{42}

Functional connectivity, both within and between networks, was reduced in COVID-DoC as compared to healthy controls. White matter integrity was also reduced in COVID-DoC as compared to healthy controls. Moreover, white matter integrity was comparable between patients with COVID-DoC and patients with DoC caused by severe TBI, a condition known to cause axonal injury.\textsuperscript{15,16,18} Just as diminished functional and structural connectivity has been observed in other DoCs, disruption of brain network integrity may similarly explain impaired consciousness in COVID-DoC.

The cause of diminished brain connectivity in COVID-DoC is unknown. The fact that microhemorrhages and leukoencephalopathy were observed in many patients raises the possibility that these findings represent overt manifestations of neurologic injury present across all patients with COVID-DoC. Whether such neurologic injury reflects systemic inflammation, toxic-metabolic insults, the effects of prolonged intubation and intensive care, direct viral infection, hypoxemia, or a combination of these and other factors, remains a question for future research.

Though the recovery of consciousness is a critical consideration in prognostication, the prospects for longer-term functional recovery are similarly important. All patients who survived the hospitalization required constant support at discharge, but by three months after discharge over 60% had returned home. By six months, 75% lived at home with normal cognition and minimal disability (due to persistent weakness or pain); the other 25% (two patients) remained in inpatient facilities due to ongoing medical illness.
This study has several limitations. Though among the largest to date, this cohort of patients with COVID-DoC remains small, due to the challenges of enrolling and obtaining advanced neuroimaging for critically ill patients with COVID-19. As such, it is possible that we failed to capture the full spectrum of potential neurologic outcomes. Similarly, while we found no significant association between TTRC and several clinical variables, this study may have been underpowered for detecting such associations. The clinical judgment inherent to the definition of COVID-DoC may lead to heterogeneity within this patient cohort; future studies are needed to develop uniform and objective diagnostic criteria. It is possible that sedatives reduced functional connectivity in patients with COVID-DoC, though the low dose sedatives that patients received have a limited impact on functional connectivity\textsuperscript{43}, and sedation and connectivity were not appreciably associated. Because the control subjects were enrolled in a separate study, we were unable to ensure that they were matched to the patients with COVID-DoC, raising the possibility of confounds; all comparisons, however, remained significant after accounting for differences in age. Comparing the MRI characteristics of patients with COVID-DoC to patients with COVID-19 but no DoC may be an informative topic of future investigation; given the potential risks to patients and staff, at the time of this study it was not feasible to obtain research scans on patients with severe COVID-19 who did not otherwise warrant the scans clinically. Finally, clinical teams were not blinded to the neuroimaging findings; while we cannot exclude the possibility that patient care was subtly influenced, life-sustaining treatment was never withdrawn for a poor neurologic prognosis, despite diminished brain connectivity. Ultimately, larger prospective cohorts are necessary to further elucidate the natural history of COVID-DoC, more precisely characterize its neural connectivity profile, and identify prognostic biomarkers.
Among the countless tragedies of the COVID-19 pandemic, COVID-DoC has presented unique and profound challenges. Rendering patients unable to communicate, and with uncertain prospects for recovery, COVID-DoC has forced families and clinicians to decide whether to continue life-sustaining treatment with little data available for guidance. This study indicates that patients who survive the medical complications of severe COVID-19 are highly likely to regain consciousness. Though rates of disability are high immediately following hospital discharge, such disability is mostly due to the sequelae of prolonged medical illness, and patients frequently improve substantially in the ensuing months. Moreover, COVID-DoC appears to be associated with loss of functional and structural brain connectivity, findings common to other DoCs caused by brain injury. While future research is necessary to further characterize COVID-DoC, these findings help inform the prognosis and pathophysiology of COVID-DoC, a condition fraught with uncertainty.

Supplement - http://links.lww.com/WNL/B660
References


**Figure Legends**

**Figure 1. Neuroimaging features and recovery of patients with COVID-DoC.**

Patients are listed by descending COVID-DoC duration. The patient who died shortly after enrollment (patient 12) is not depicted. Patients with microhemorrhages or leukoencephalopathy on a structural brain MRI are marked with a black circle; those without findings are marked with a white circle. Intra-network functional connectivity of the default mode network, as measured with resting state functional MRI (rs-fMRI), is represented by colored circles, where red represents the average intra-network default mode network connectivity of healthy controls ($Z=0.23$) and purple represents absent connectivity ($Z=0$). White matter integrity, measured as whole brain fractional anisotropy (FA) with diffusion MRI (d-MRI), is represented by colored circles, where red represents the average whole brain FA of healthy controls (0.59) and purple represents the lowest FA measured among patients (0.45). Recovery of consciousness is depicted relative to the day of intubation. The duration of intubation is depicted in dark blue (terminating with tracheostomy or extubation), and the duration of intravenous sedation (including propofol, midazolam, hydromorphone, or ketamine) in light blue. The duration of unresponsiveness, starting with the cessation of intravenous sedation, and ending with the first documentation of command-following (represented by a red circle), is depicted in red. The timing of the brain MRI is depicted as gray arrows. Outcomes at hospital discharge are depicted in the right-hand column, including acute hospitalization length of stay (LOS) and disposition.
Figure 2. Longer-term functional outcomes for patients with COVID-DoC. The Glasgow Outcome Scale Extended (GOSE; A) and Disability Rating Scale (DRS; B) scores for each patient are shown (higher scores on GOSE, and lower scores on DRS, reflect less disability). Functional outcomes are depicted at hospital discharge, 3 months post-discharge, and 6 months post-discharge.
Figure 3. Resting state functional connectivity for patients with COVID-DoC and healthy controls. Connectivity of the default mode network is depicted as group-wise seed-to-voxel maps, generated with seeds at four nodes – the medial prefrontal cortex, posterior cingulate cortex, and bilateral inferior parietal lobules – for healthy controls (A) and patients with COVID-DoC (B). Warmer colors represent positive correlations (reflecting intra-network connectivity) and cooler colors represent negative correlations (reflecting inter-network connectivity). Patients with COVID-DoC demonstrated less positive intra-network connectivity (C) and less negative inter-network connectivity (D) than healthy controls. ** = p < 0.01. *** = p < 0.001.
Figure 4. Diffusion MRI metrics for patients with COVID-DoC and healthy controls.

White matter integrity within a white matter skeleton is depicted as a three-dimensional group-wise fractional anisotropy (FA) map, with warmer colors representing higher FA and thus higher white matter integrity, for healthy controls (A) and patients with COVID-DoC (B). Whole brain (C) and brainstem (D) FA values of patients with COVID-DoC were low compared to healthy controls, but comparable to patients with severe traumatic brain injury (TBI). *** $= p < 0.001$. **** $= p < 1 \times 10^{-8}$. 
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