Case Report

Electroencephalographic evidence of organic alteration in a patient with SARS-CoV2 induced delirium

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Abstract

We have recently described, by means of quantified EEG (qEEG), a specific pattern in COVID-19 patients released from intensive care unit. In this work, we describe a patient showing an atypical delirium with insomnia, memory disturbances, tempo-spatial disorientation and significant language skills impairment, while maintaining a good level of attention and alertness. Three months later, the patient recovered his cognitive and behavioral premorbid state. Electroencephalographic evidence of organic alteration in a patient with SARS-CoV2 induced delirium

Introduction

Neurological complications in COVID-19 infected patients have been extensively reported. CNS affections include encephalitis, toxic encephalopathy, ageusia and anosmia, headache or acute cerebrovascular disease and delirium [1-5]. The mechanisms of CNS infection by CoV2 are still debated and it has been proposed a direct invasion through blood-brain barrier, a neuronal pathway, hypoxia damage, immune-response mediated injury or angiotensin-converter enzyme 2, among others [2,6,7]. COVID-19 patients suffering severe respiratory compromise are prone to develop neuropsychiatric symptoms [8]. Among these, one of the most frequent is the acute confusional syndrome or delirium [up to 28%] [5].

We have recently described, by means of quantified EEG (qEEG), a specific pattern in COVID-19 patients released from intensive care unit (ICU), showing brain alteration associated with neuropsychiatric symptoms [4].

In this work, we describe a patient with an atypical delirium and its evolution, highly correlated with alterations in qEEG, demonstrating an organic brain affection induced by COVID-19.

Case presentation

Clinical case

A 55-year-old male physician in active, with premorbid conditions of essential tremor without treatment and moderate alcoholic consumption, was directly admitted at intensive care unit (ICU) by acute severe respiratory syndrome (SARS-CoV-2), with a 86% minimum capillary oxygen saturation (SaO2). The patient was intubated under sedo-analgesia (fentanyl, midazolam and dexmedetomidine) and treated with tocilizumab (3 doses), hydroxychloroquine (12 days), lopinavir/ritonavir (14 days) and levofloxacin (15 days). Fifteen days later was released to conventional hospitalization area. The patient developed a critically illness polyneuropathy, later resolved by rehabilitation. After five days of clinical stability, the patient developed significant cognitive and behavioral changes. Neurological examination did not uncover any further focal neurological deficits. He developed insomnia and an atypical delirium with memory disturbances, tempo-spatial disorientation and significant language skills impairment, while maintaining a good level of attention and alertness. The patient was treated with high doses of antipsychotics, initially aripiprazole (10 mg/day) and quetiapine (100 mg/day), later replaced by risperidone (6 mg/day) when vascular damage was discarded. Imaging studies such as computed tomography and magnetic resonance imaging were normal. EEG (EEGcont) was nearly physiologic in a first de visu inspection, although qEEG showed a mild encephalopathy (see below). HIV and syphilis serology resulted negative, vitamin complex B and thyroid function were normal. The patient was delivered from hospital one month after admission.

Three months later, the patient was re-evaluated and a new EEG was performed (EEGcont). He showed a complete cognitive recovery and had returned to his job as physician. Only a residual fatigability remained (after climbing stairs 1-2 floors).

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Quantification of EEG

EEGs’ records were performed using a 32-channel digital system (EEG32U, NeuroWorks, XLTek®, Oakville, Canada) with 19 electrodes placed according to a 10–20 international system. Recordings were performed at 512 Hz sampling rate, with a filter bandwidth of 0.5 to 70 Hz and notch-filter of 50 Hz. Electrode impedances were usually below 15 kΩ. Artifact-free periods (excluding electro-oculogram, muscular or movement in awake patient) were selected and exported in ASCII file to be quantified. The algorithm used has been previously published [4,9]. Briefly, areas under the power spectrum, obtained by fast Fourier transform, were computed for classical EEG bands defined as delta (0.5–4.0 Hz), theta (4.0–8.0 Hz), alpha (8.0 – 13.0 Hz) and beta (13.0–30.0 Hz). Shannon Spectral entropy (SSE) and Pearson’s correlation coefficient (ρ) for pair of electrodes were also calculated. One second windowing 10% superimposed were used to compute all of these measurements. Exported records were between 120 and 300 s, which allowed a minimum of 130 and a maximum of 330 windows to be computed. Numerical analysis of EEG recordings was performed with custom-made Matlab® R2019 software (MathWorks, Natic, MA, USA).

Statistical comparisons between groups were performed using the Student’s t-test for data with normal distributions. Normality was evaluated using the Kolmogorov–Smirnov test. The Mann–Whitney Rank sum test. SigmaStat® 3.5 software (SigmaStat, Point Richmond, CA, USA) and Matlab® were employed for statistical analysis.

The significance level was set at p=0.05. Results are shown as the mean ± SEM, except where otherwise indicated.

De visu EEGEnc showed a near normal background, with presence of a low-frequent medium-voltage sharp waves at left temporal lobe. Diagnosis of a mild encephalopathy was performed by qEEG (see below). The EEGCont performed three months later, showed a quite similar background without irritative activity at left temporal lobe. Nevertheless, the numerical analysis of EEGEnc showed more interesting and no self-evident results. Spectral analysis revealed a generalized excess of delta bands, especially at anterior regions, with a posterior dominant rhythm at 8.5 Hz and a mean amplitude of 17.1 ± 1.7 μV. On the contrary, the numerical analysis EEGEnc showed very different results. In fact, the posterior dominant rhythm was 9.0 Hz, with the same amplitude (21.1 ± 1.3 μV, not significant, Mann-Whitney on ranks). In both cases, amplitude of EEG was normal and not reduced, as described in de visu analysis [10]. It’s was quite surprising that not only the power of different bands were different between both recordings, with excess of delta band, mainly at fronto-temporal regions, but it was observed that distribution of frequencies was different, with an increase in the peak frequency of alpha band (Figure 1 A-B). Moreover, the pattern of EEG, defined as the percentage of different bands by lobe, resulted highly different. In fact, recovery is associated with a significant increase in theta and alpha bands of both hemispheres, together with beta band at frontal lobes (Figure 1 C-E).

Nonetheless, not only cerebral rhythms were changed during recovery, but also the synchronization was modified (Table 1). Interestingly, during the encephalopathy the patient presented an excessive synchronization in left hemisphere and right temporal lobe (Table 1).

Another interesting fact described in encephalopathic COVID-19 patients is the relative increase in SSE. We have observed this fact for all the scalp regions in our patient and its decrease with recovery (Table 2).

Discussion

We have analyzed the scalp EEG performed in a patient during acute COVID-19 illness and three months later after clinical resolution and we have observed very relevant changes in qEEG among both periods that can be helpful to explain pathophysiology and can be used to diagnosis. Reports about EEG in COVID-19 patients are performed by de visu and different findings have been described but without specific features [10-12]. Nevertheless, we have described the EEG structure of encephalopathy in COVID-19 patients released from ICU after severe illness that clearly differs from other kinds of encephalopathy [4], showing specific features.

During encephalopathy, background EEG was scarcely pathological and the mild left temporal irritative activity observed cannot be responsible of the delirium. However, the brain activity was significantly impaired, as we observed in qEEG, e.g. the cortical bands (alpha and beta [9]) were severely decreased, mainly at temporal and parieto-occipital lobes, together with an increase of theta activity, besides; delta activity was increased in temporal lobes, mainly in the right. Moreover, these modifications were associated with changes in synchronization and structure of band composition across the scalp, as indicated by SSE. The relative normality of frontal lobes, together a most severe impairment of temporal lobes, in the context of abnormal synchronization in the

Table 1. Comparison for Pearson’s coefficient (ρ) for EEGEnc and EEGCont

| Location     | Total     | Frontal   | Parieto-occipital | Temporal |
|--------------|-----------|-----------|-------------------|----------|
| Left hemisphere | Encephalopathy | 0.514 ± 0.003 | 0.525 ± 0.007 | 0.495 ± 0.007 | 0.468 ± 0.004 |
|               | Control   | 0.497 ± 0.003 | 0.498 ± 0.003 | 0.482 ± 0.006 | 0.610 ± 0.008 |
|               | Probability | 0.001     | 0.001             | n.s      | 0.001      |
| Right hemisphere | Encephalopathy | 0.515 ± 0.003 | 0.526 ± 0.007 | 0.495 ± 0.007 | 0.468 ± 0.004 |
|               | Control   | 0.508 ± 0.005 | 0.532 ± 0.012 | 0.510 ± 0.009 | 0.643 ± 0.011 |
|               | Probability | 0.05       | n.s               | n.s      | 0.001      |

Table 2. Comparison Shannon spectral entropy (SSE) for EEGEnc and EEGCont

| Location     | Frontal     | Parieto-occipital | Temporal |
|--------------|-------------|-------------------|----------|
| Left hemisphere | Encephalopathy | 4.3 ± 0.03 | 4.60 ± 0.03 | 4.91 ± 0.04 |
|               | Control     | 4.17 ± 0.03 | 4.23 ± 0.04 | 4.28 ± 0.04 |
|               | Probability | 0.001     | 0.001             | 0.001      |
| Right hemisphere | Encephalopathy | 4.30 ± 0.03 | 4.33 ± 0.02 | 4.61 ± 0.02 |
|               | Control     | 4.08 ± 0.03 | 4.23 ± 0.04 | 4.19 ± 0.03 |
|               | Probability | 0.001     | 0.01             | 0.001      |
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**Figure 1.** Numerical comparison between qEEG. (a) Mean spectra by channels of the left hemisphere and (b) right hemisphere. Dashed lines represent values from EEGenc and continuous lines EEGcont. Comparison of values for bands at (c) frontal, (d) parieto-occipital and (e) temporal lobes. Note: Filled circle=left encephalopathy; empty triangle=left control; red square=right encephalopathy and green diamond=right control. Blue asterisks=statistically significant difference for left hemisphere and red ones for right hemisphere. *p<0.05; **p<0.01; ***p<0.001, Student t-test.

The left hemisphere can explain the anomalous features of the delirium presented by the patient [5,13,14].

We hypothesize that change affecting specific cerebral structures can be responsible of the different symptomatology described.

Considering that mild changes in synchronization or brain rhythmicity cannot be observed *de viso*, we encourage the use of objective measurements of bioelectrical variables to define more specific associations between electrical brain activity and symptoms in COVID-19 patients.

**Conclusion**

This is the first report of objective changes of bioelectrical brain activity during COVID-19 induced encephalopathy and after resolution. Reversible changes observed in EEG structure and connectivity can explain the anomalous delirium suffered by the patient.

**Conflicts of interest**

None.
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