Case report: Remote neuromodulation with direct electrical stimulation of the brain, as evidenced by intra-operative EEG recordings during wide-awake neurosurgery

Real-time functional mapping of the brain with direct electrical stimulation (DES) is used to guide the resection of slow-growing infiltrative tumours during wide-awake surgery. The DES technique reduces the probability of resecting essential areas near or within the tumour (Duffau, 2005). During the neurosurgery, patients perform a number of neuropsychological tests while DES is applied both cortically and subcortically, in order to detect and thus preserve connectivity online. This technique is especially useful for critical white matter pathways. This functional mapping of the area near the tumour enables the removal of as much as non-eloquent infiltrated tissue as possible, while minimizing sequelae.

The local electrophysiological effects of DES have only been partially characterized. By using electrocorticography and an implanted grid on the surface of the grey matter for pre-surgical planning in drug-resistant epileptic patients, Matsumoto et al. (2004, 2007) observed “corticocortical” evoked potentials around 10–50 ms after the start of low-frequency DES (1 Hz, 10–12 mA) at a cortical site. The evoked potentials were measured in contiguous cortical areas whose centres were up to three centimetres apart and were linked by axons in the grey matter or in local U-shaped subcortical fibres. In contrast, the propagation of DES and its remote effects have not yet been investigated (Mandonnet et al., 2010; Szélényi et al., 2010).

Here, we report on a case of wide-awake surgery for a slow-growing, right frontal lobe tumour in a 33-year-old man. Intra-operative electroencephalographic (iEEG) recording was used to determine whether cortical or subcortical DES can have remote neuromodulatory effects on electrophysiological signals. The volume of the resection was 116.8 cm³. DES (60 Hz, with biphasic current: I = 2 mA, pulse duration = 1 ms) was applied at different locations during the recordings (Fig. 1a). iEEG signals were recorded at four scalp sites: three on the contralateral hemisphere (F3, C3 and O1) and one on the ipsilateral hemisphere (O2). An additional reference electrode was placed on the right mastoid. During the recording itself, the Biosemi system’s common-mode sense electrode served as the reference electrode. Electrophysiological signals were sampled at 2048 Hz and acquired with ActiView software. iEEG signals were detrended, and different stimulation periods were determined based on DES induced artefacts. For cortical DES, we obtained 23 periods with durations ranging from 1.3 to 10.2 s, separated by intervals of at least 2.3 s. For subcortical DES, we obtained nine periods with durations ranging from 3.3 to 11.6 s, separated by intervals of at least 5.1 s. Mean spectrograms were first determined (moving window: 500 ms; overlap: 90%, Hamming window: 1024 samples, nfft = 2048 samples) in order to detect noise (at 50 Hz, plus 60 Hz for DES artefacts). Mean power spectral densities (PSDs) were computed for the [0, 40] Hz frequency band, which clearly contained most of the iEEG signals.

When DES was applied to the right frontal lobe, the median frequency of the mean PSD computed for the [0, 40] Hz frequency band increased at C3 for both cortical and subcortical stimulation ($p = .045$ and $p = .006$, respectively), at F3 for cortical stimulation only ($p = .002$) and at O1 and O2 for subcortical stimulation only ($p = .004$ and $p = .026$, respectively). Furthermore, the power of the [20, 30] Hz frequency band also increased at C3, F3, O1, and O2 for subcortical DES (Fig. 1b and c, $p < .029$ in all cases). Interestingly, the increase in the median frequency, averaged over the different periods, at O1 was found to be progressive when DES was applied subcortically (see Fig. 1d, $p = .04$). In contrast, the power of the [10, 15] Hz frequency band decreased during DES ($p = .038$). After cortical DES, the power of a [0, 10] Hz frequency band increased at F3 ($p = .026$). A non-significant trend was observed for the [0, 10] Hz frequency band at F3 with subcortical DES ($p = .063$).

We observed significant changes in the frequency content at different iEEG sites during DES. Subcortical DES (i.e. stimulation of white matter pathways) led to neuromodulation at more sites than cortical DES. This may have been due to (i) better conduction and propagation following the direct stimulation of large, myelinated axons and (ii) the greater current intensity in subcortical DES. Further research will have to characterize these aspects more carefully and apply cortical and subcortical DES with identical current intensities. Most of the observed changes occurred immediately after the start of DES. However, for subcortical DES, the median frequency at O1 increased progressively during the first second of the stimulation train. This observation suggests that immediate changes and progressive changes have differing mechanisms. The O1 electrode was furthest from the stimulation site; when more neural elements within a network are involved, neuromodulation with DES may be delayed by intermediary electrophysiological interactions. For some electrodes, we observed that the power increased within the [20, 30] Hz frequency band (i.e. the beta band) with cortical or subcortical DES. At F3, we also observed an effect of increasing power on the [0, 10] Hz frequency band. At this stage, it is premature to link DES-associated electrophysiological modulations to particular cognitive aspects: a variety of tasks were performed during surgery, and clinical assessments are performed by a neuropsychologist in real-time to detect perturbations. It remains to be seen whether DES induces these effects indirectly (i.e. through behavioural changes) or directly.

Although we investigated evoked potentials in the time domain, the method needs to be modified because the frequency used for DES (60 Hz) and the propagation time of the evoked
potential (10–20 ms) are close enough to either mask the interesting signal or induce error in the determination of sequences of events. Accordingly, time-domain analyses are not yet possible; one would have to change the frequency of DES, which would require a specific scientific rationale and approval by an investigational review board since this change in DES is not part of routine clinical practice. On the basis of the present case report, we are developing a protocol for application to a series of patients undergoing wide-awake surgery. We expect a time-domain assessment of the future series to provide very useful, complementary information.

In conclusion, DES may induce some remote neuromodulatory effects and electrophysiological changes in both the contralesional and ipsilesional hemispheres.

Conflict of interest

None.

References


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