

# Propofol Reduces Microelectrode-Recording Artefacts caused by Parkinsonian Tremor during Deep Brain Stimulation

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## Abstract

A 51-year-old male with medically refractory Parkinson's disease was scheduled for bilateral deep brain stimulation (DBS). During microelectrode recordings (MERs) of right side DBS, the patient developed severe sustained whole-body tremors causing severe artefacts in MER. The right side DBS electrode was inserted with suboptimal MER. For the creation of left burr hole, propofol infusion at a rate of 20 µg/kg/min, was used and soon after, all tremor activity ceased. Propofol infusion was continued during left side MER. With the absence of tremors, left subthalamic nucleus spike activity was better identified and neurological testing could take place. At 6 months after DBS, the patient's symptoms had improved significantly without the need for levodopa.

## Keywords

- ▶ a tremor effect
- ▶ deep brain stimulation
- ▶ Parkinson's disease
- ▶ propofol

## Introduction

The anesthetic goals for deep brain stimulation (DBS) are to ensure patient comfort for insertion of electrodes through burr holes while avoiding all drugs that interfere with microelectrode recordings (MERs) and macrostimulation. There are few studies which examine the effects of anesthetic drugs on MER during deep brain nucleus (DBN) localisation. This has led some neurosurgical teams to totally avoid sedation, especially propofol, during DBS. We would like to report a case, however, where the anti tremor effect of propofol was used to facilitate MER.

The patient gave written permission for the authors to publish the report.

## Case Report

A 51-year-old male with medically refractory Parkinson's disease (PD) was scheduled for bilateral DBS under conscious sedation. Despite taking levodopa of 125 mg and amantadine of 100 mg twice daily, he suffered severe resting tremor in

all limbs affecting his daily activities. The parkinsonian drugs were withheld on the day of surgery. He had no other significant medical history.

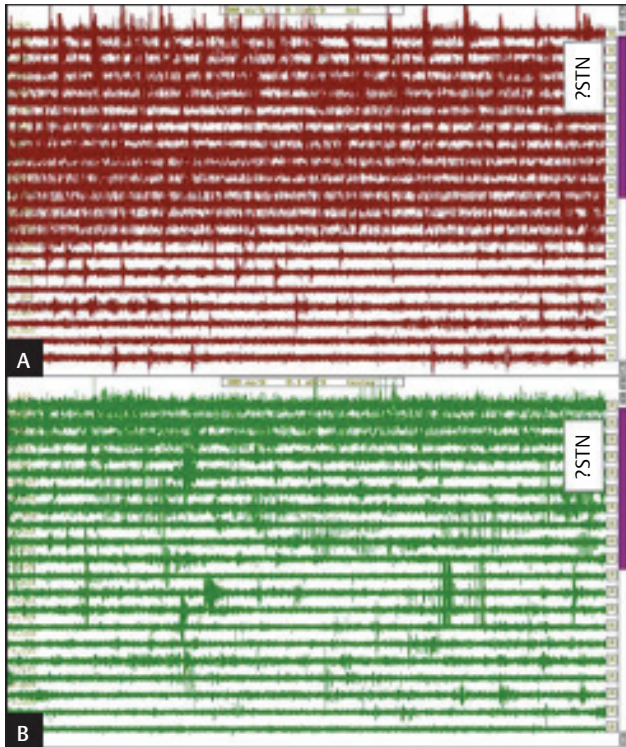
A Leksell stereotactic frame was placed on the patient using a local anesthetic to pin sites before computed tomography scan of the head and transferred to the operation room. No premedication was given. The standard monitoring, including electrocardiogram, pulse oximetry, and non invasive blood pressure monitoring were used. Propofol infusion (30–50 µg/kg/min) was used for sedation during the creation of first burr hole on the right side. Twenty minutes before MER, propofol was stopped in accordance with our institution's protocol, to avoid possible interference with MER. During MER, the patient developed severe sustained whole-body tremors leading to violent shaking of the Leksell head frame and operating table. The tremors caused so many artefacts in the MER that they prevented identification of the inferior border of the right subthalamic nucleus (STN) (→ Fig. 1). MER was suboptimal despite 45 min of mapping with repeated trajectories. The final placement of the right

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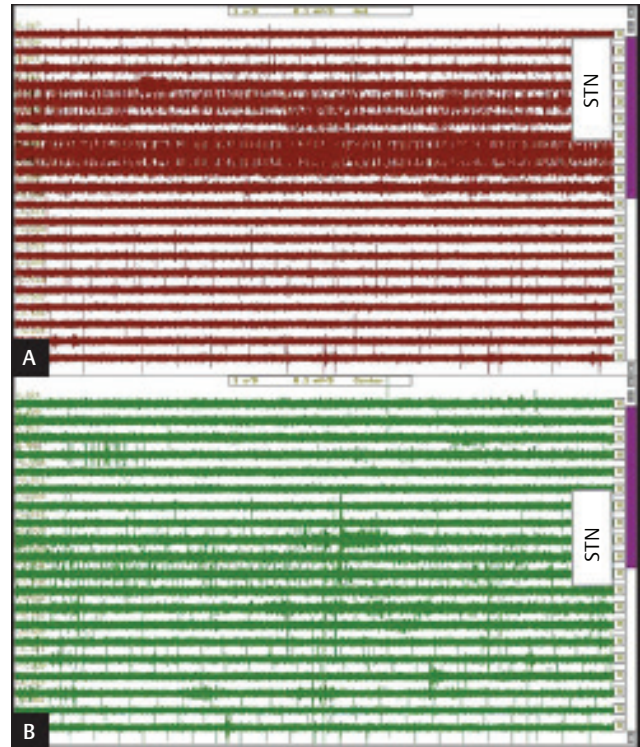
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**Fig. 1** (A) Right hemispheric anterior microelectrode shows excessive tremor artefacts throughout the entire length of the recording. The superior subthalamic nucleus is detected 1.5 mm above target. Inferior border of subthalamic nucleus is not detected, (B) Right hemispheric center microelectrode shows excessive tremor artefacts throughout the entire length of the recording. The superior subthalamic nucleus is detected 1 mm above target. Inferior border of subthalamic nucleus is not detected.



**Fig. 2** (A) Left hemispheric anterior microelectrode shows fewer tremor artefacts throughout the entire length of the recording. The superior subthalamic nucleus is detected 4 mm above target. Inferior border of subthalamic nucleus is 2 mm above target. (B) Left hemispheric center microelectrode shows very few tremor artefacts throughout the entire length of the recording. The superior subthalamic nucleus is detected 0.5 mm below target. Inferior border of subthalamic nucleus is 3.5 mm below target.

DBS electrode was based on the neurological examination with macrostimulation of the electrode.

Propofol infusion at a rate of 30 to 50  $\mu\text{g}/\text{kg}/\text{min}$  was recommenced for the creation of the left burr hole, and soon after, all tremor activity ceased. A decision was made to continue propofol during MER. With propofol infusion at a rate of 20  $\mu\text{g}/\text{kg}/\text{min}$ , there was no recurrence of tremor activity, and the patient obeyed verbal commands and spoke fluently. With the absence of tremor artefact in MER, left STN spike activity was better identified ( $\rightarrow$  Fig. 2). Then, propofol was stopped to allow the tremor resurface which it did within 7 min after cessation of propofol. Macrostimulation and neurological testing took place for the final placement of the left DBS electrode. Following this, general anesthesia was instituted for stage 2-tunneling and insertion of the generator.

At 6-months post-DBS, the patient's symptoms had improved significantly, and he was weaned off levodopa.

His simplified Movement Disorders Societies-Unified Parkinson's Disease Rating Scale III, a rating score used by our neurologists, was markedly reduced from 18.5 to 6.5 (stimulation-on; medication off state), equivalent to 65% reduction in symptoms ( $\rightarrow$  Table 1).

## Discussion

In this case, we observed a reversible anti-Parkinsonian tremor effect associated with the administration of low dose propofol,

facilitating MER, which might be helpful in many other DBS cases with a similar intraoperative problem. Spurious movement or tremor can interfere with MER during DBS. Tremor is the most common cardinal symptom in PD, which is found in three-quarter of PD patients.<sup>1</sup> As expected, tremor-related artefacts interfering MER are commonly encountered clinically. In addition, the typical target nucleus (e.g., STN) is only a few millimetres in diameter, precise placement of DBS electrode, a crucial step to achieve desired therapeutic effect, is technically challenging, especially in a patient with severe tremor.<sup>2</sup> DBS electrode placement is a blind procedure; the risks of intracranial hemorrhage or injury to vital structures are correlated to the number of microelectrode trajectories.<sup>3</sup> We would like to use this case as a discussion point to review the current understandings on the anti-Parkinsonian effect of propofol, as well as to discuss the concerns about propofol interferences on MER.

### Antitremor Effect of Propofol

Tremor pathophysiology involves complex anatomical pathways between basal ganglia, cerebellar circuits and motor cortex ( $\rightarrow$  Fig. 3). Tremor is exhibited as there is increased cerebellothalamocortical circuit activity. For example, dopaminergic depletion of the pallidum in PD or gamma-aminobutyric acid (GABA)-ergic dysfunction of the cerebellar dentate nucleus and brain stem in essential tremor (ET) cause hyperactivity of cerebellothalamocortical circuit, resulting in clinical tremor in both diseases.<sup>4</sup>



MER. It implied the effective dose for anti-Parkinsonian tremor is much lower than the effective dose for sedation, indicating the differential sensitivity of DBN (subcortical neurons) and cortical neurons towards anesthetic agents. It is also reasonable to question whether other anesthetic agents (GABA agonists) possess the same effect. A previous case report described ketamine abolished tremor and dyskinesia in a PD patient, which postulated to be through N-methyl-D-aspartate receptor activation.<sup>11</sup> Further study is required to evaluate the underlying mechanism, dose-response relationship and whether other anesthetics exert similar effects in the tremor pathway. On the other hand, propofol can also cause myoclonus; however, the dose-response relationship between myoclonus and the anti-tremor effect is yet to be determined.

### Concerns of Propofol during Microelectrode Recording

The common methods of DBN localisation include neuronavigation, MER and macrostimulation techniques.

High fidelity magnetic resonance imaging enables generation of highly accurate coordinates to grossly locate the DBN, and placement of DBS electrode is further finely adjusted according to the MER and microstimulation.

MER is typically started at 10 to 15 mm above the targeted nucleus. The electrode is advanced by a microdrive and look for occurrence and disappearance of STN-specific bursting pattern of electrophysiological activity to determine the border of STN.

The length of the STN, the presence of movement-responsive neurons, and distance from the STN border to adjacent structures are used as MER criteria to determine the best trajectory for permanent electrode implantation. Since the electrophysiological activities of DBN are very small (50–200  $\mu$ V) and sensitive to anesthetic agents, anesthetic suppression of MER during placement of electrodes becomes one of the major concerns affecting the accuracy of mapping during DBS.<sup>2</sup>

Macrostimulation, referring to an *in vivo* test stimulation of deep brain electrode, physical examination is used to check for side effects due to overstimulation of surrounding nucleus and to assess the clinical effect of DBS. A fully awake patient is required to participate neurological examination during macrostimulation. A previous study has showed adjustment of electrode placement after macrostimulation were required in 17%–87% of STN-DBS patients with average target adjustments of 1 to 4 mm.<sup>2</sup> Thus, a highly controlled anesthetics is desirable to achieve the best procedural benefit.

The anesthetic effects on MER are complex. Several factors can modify the effect of anesthetics, including the site of target nuclei (STN, globus pallidus internus [GPI], ventromedial thalamic nucleus), the disease state (PD, ET and dystonia), the severity of the disease (variable degree of neuronal depletion). Following the report that propofol infusions at 50  $\mu$ g/kg/min significantly decreased STN neuronal activity (-23.2%),<sup>12</sup>

**Table 2** Review of patient outcomes following deep brain stimulation using microelectrode recording technique under general anesthesia

Study	Study design	Sample size	Anesthetic agents	UDPRS-III score reduction	LEDD reduction	Conclusion
Kim et al <sup>19</sup>	Comparison study	8	Propofol + remifentanyl (one side) LA (the contralateral side)	67%	N/A	No significant difference in the mean firing rate between the left and the right side MERs
Fluchere et al <sup>14</sup>	CS	213	Sevoflurane	61% (1 y) 37% (5 y)	46% (1 y) 49% (5 years)	STN stimulation performed under controlled GA is efficient and has similar short- and long-term motor effects to local anesthesia
Harries et al <sup>15</sup>	CS	82	Isoflurane + N <sub>2</sub> O (26 patients) Propofol + remifentanyl (56 patients)	22.89 (1 y)	58.1% (1 y)	Satisfactory MER of the STN were able to obtain under GA
Lin et al <sup>16</sup>	CS	10	Desflurane (0.5-1 MAC)	5.42% (6 mo)	N/A	Typical neuronal firing patterns of the STN and substantia nigra reticulata were able to observe in all patients
Hertel et al <sup>17</sup>	CS	9	Propofol (0.1-0.2 mg/kg/min) Remifentanyl	24%	N/A	All patients had satisfactory MER and STN
Maltête et al <sup>18</sup>	Case-control study	30 (15:15)	Propofol (TCl: 0.8-2 ng/mL)	N/A	N/A	Both GA and LA group has markedly improved the parkinsonian motor disability score The GA group has higher residual parkinsonian motor score than LA group

Abbreviations: CS, case-series; N/A, not available; GA, general anesthesia; LA, local anesthesia; LEDD, levodopa equivalent dose; UDPRS, Unified Parkinson's Disease Rating Scale; TCl, target controlled infusion; MERs, microelectrode recordings; STN, subthalamic nucleus; MAC, minimum alveolar concentration.

many neurosurgical teams decided to avoid propofol altogether during DBS. There was also similar report about anesthetic suppression (propofol) on MER during GPi-DBS in dystonia patients, where the anesthetic suppression was more pronounced in dystonia patients than PD patients.<sup>13</sup>

In contrast to these reports, there are several case series reporting successful MER and motor outcomes in PD patients undergoing STN-DBS with general anesthesia (propofol or volatile anesthetics) (►Table 2).<sup>14-18</sup> Interestingly, all these studies found MER were not affected by controlled general anesthesia and were able to detect bursting STN pattern in all study cases. Although one of the typical features of STN (widening of background noise baseline) were lost during general anesthesia, the overall motor outcomes and symptoms improvement were comparable between general anesthesia and historical controlled data under local anesthesia.<sup>14-18</sup> While anesthetic suppression is a genuine phenomenon; but under controlled general anesthesia condition, presumably a higher dose of anesthetics were used, MER and patient outcomes were no clinical differences based on non-randomized data. The avoidance of propofol or other anesthetic agents on MER in all DBS patients appears to be over-concerned, especially in some patients who might benefit from adequate sedation during this prolonged procedure.

## Conclusion

We present an interesting DBS case where low dose propofol infusion suppressed Parkinsonian tremor, facilitated MER and the successful placement of DBS electrodes. Further studies are needed to investigate the dose-response relationship of propofol and if other drugs can facilitate MER during DBS.

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### Conflict of interest

None.

## References

- Hughes AJ, Daniel SE, Blankson S, Lees AJ. A clinicopathologic study of 100 cases of Parkinson's disease. *Arch Neurol* 1993;50:140-148
- Benabid AL, Chabardes S, Mitrofanis J, Pollak P. Deep brain stimulation of the subthalamic nucleus for the treatment of Parkinson's disease. *Lancet Neurol* 2009;8:67-81
- Gross RE, Krack P, Rodriguez-Oroz MC, Rezai AR, Benabid AL. Electrophysiological mapping for the implantation of deep brain stimulators for Parkinson's disease and tremor. *Mov Disord* 2006;21 Suppl 14:S259-S283
- Helmich RC, Toni I, Deuschl G, Bloem BR. The pathophysiology of essential tremor and Parkinson's tremor. *Curr Neurol Neurosci Rep* 2013;13:378
- Deuschl G, Raethjen J, Hellriegel H, Elble R. Treatment of patients with essential tremor. *Lancet Neurol* 2011;10:148-161
- Filion M, Tremblay L. Abnormal spontaneous activity of globus pallidus neurons in monkeys with MPTP-induced Parkinsonism. *Brain Res* 1991;547:142-151
- Vila M, Levy R, Herrero MT, Ruberg M, Faucheux B, Obeso JA, et al. Consequences of nigrostriatal denervation on the functioning of the basal ganglia in human and nonhuman primates: *An in situ* hybridization study of cytochrome oxidase subunit I mRNA. *J Neurosci* 1997;17:765-773
- Ihara M, Tomimoto H, Ishizu K, Yoshida H, Sawamoto N, Hashikawa K, et al. Association of vascular Parkinsonism with impaired neuronal integrity in the striatum. *J Neural Transm (Vienna)* 2007;114:577-584
- Hall SD, Prokic EJ, McAllister CJ, Ronnqvist KC, Williams AC, Yamawaki N, et al. GABA-mediated changes in inter-hemispheric beta frequency activity in early-stage Parkinson's disease. *Neuroscience* 2014;281:68-76
- Anderson BJ, Marks PV, Futter ME. Propofol - Contrasting effects in movement disorders. *Br J Neurosurg* 1994;8:387-8
- Wright JJ, Goodnight PD, McEvoy MD. The utility of ketamine for the preoperative management of a patient with Parkinson's disease. *Anesth Analg* 2009;108:980-982
- Raz A, Eimerl D, Zaidel A, Bergman H, Israel Z. Propofol decreases neuronal population spiking activity in the subthalamic nucleus of Parkinsonian patients. *Anesth Analg* 2010;111:1285-1289
- Sanghera MK, Grossman RG, Kalhorn CG, Hamilton WJ, Ondo WG, Jankovic, et al. Basal ganglia neuronal discharge in primary and secondary dystonia in patients undergoing pallidotomy. *Neurosurgery* 2003;52:1358-1370
- Fluchere F, Witjas T, Eusebio A, Bruder N, Giorgi R, Leveque M, et al. Controlled general anaesthesia for subthalamic nucleus stimulation in Parkinson's disease. *J Neurol Neurosurg Psychiatry* 2014;85:1167-1173
- Harries AM, Kausar J, Roberts SA, Mocroft AP, Hodson JA, Pall HS, et al. Deep brain stimulation of the subthalamic nucleus for advanced Parkinson disease using general anesthesia: Long-term results. *J Neurosurg* 2012;116:107-113
- Lin SH, Chen TY, Lin SZ, Shyr MH, Chou YC, Hsieh WA, et al. Subthalamic deep brain stimulation after anesthetic inhalation in Parkinson disease: A preliminary study. *J Neurosurg* 2008;109:238-244
- Hertel F, Ziechner M, Weimar I, Gemmar P, Noll B, Bettag M, et al. Implantation of electrodes for deep brain stimulation of the subthalamic nucleus in advanced Parkinson's disease with the aid of intraoperative microrecording under general anesthesia. *Neurosurgery* 2006;59:E1138
- Maltête D, Navarro S, Welter ML, Roche S, Bonnet AM, Houeto JL, et al. Subthalamic stimulation in Parkinson disease: With or without anesthesia? *Arch Neurol* 2004;61:390-392
- Kim W, Song IH, Lim YH, Kim MR, Kim YE, Hwang JH, et al. Influence of propofol and fentanyl on deep brain stimulation of the subthalamic nucleus. *J Korean Med Sci* 2014;29:1278-1286