Intraoperative Neurophysiological Monitoring for Spinal Cord Tumor Surgery: Comparison of Motor and Somatosensory Evoked Potentials According to Tumor Types

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Objective  To identify which combination of motor evoked potentials (MEPs) and somatosensory evoked potentials (SEPs) is most reliable for postoperative motor deterioration during spinal cord tumor surgery, according to anatomical and pathologic type.

Methods  MEPs and SEPs were monitored in patients who underwent spinal cord tumor surgery between November 2012 and August 2016. Muscle strength was examined in all patients before surgery, within 48 hours postoperatively and 4 weeks later. We analyzed sensitivity, specificity, positive and negative predictive values of each significant change in SEPs and MEPs.

Results  The overall sensitivity and specificity of SEPs or MEPs were 100% and 61.3%, respectively. The intraoperative MEP monitoring alone showed both higher sensitivity (67.9%) and specificity (83.2%) than SEP monitoring alone for postoperative motor deterioration. Two patients with persistent motor deterioration had significant changes only in SEPs. There are no significant differences in reliabilities between anatomical types, except with hemangioma, where SEPs were more specific than MEPs for postoperative motor deterioration. Both overall positive and negative predictive values of MEPs were higher than the predictive values of SEPs. However, the positive predictive value was higher by the dual monitoring of MEPs and SEPs, compared to MEPs alone.

Conclusion  For spinal cord tumor surgery, combined MEP and SEP monitoring showed the highest sensitivity for the postoperative motor deterioration. Although MEPs are more specific than SEPs in most types of spinal cord tumor surgery, SEPs should still be monitored, especially in hemangioma surgery.

Keywords  Spinal cord neoplasm, Intraoperative neurophysiological monitoring, Motor evoked potentials, Somatosensory evoked potentials, Postoperative complications
INTRODUCTION

Improvements in surgical techniques increasingly lead to more aggressive tumor resections. However, even for the most skillful surgeons, spine surgery has an inherent risk of postoperative neurological deterioration [1]. Intraoperative neurophysiologic monitoring (IONM) enables the surgeon to proceed with more aggressive maneuvers for near total removal of the tumor [2].

There have been various studies on effective monitoring, based on the clinical importance of IONM for intramedullary spinal cord tumor surgery. However, a specific guideline on IONM techniques has yet to be established. Recently, Scibilia et al. [3] reviewed several previous studies on different combinations of IONM techniques (such as somatosensory evoked potentials [SEPs], motor evoked potentials [MEPs], direct wave, free-running electromyography), and emphasized the multimodal IONM for safe spinal cord tumor surgery. There are controversies concerning certain IONM combination techniques, but they found combined motor and somatosensory evoked potential monitoring is most commonly used for spinal cord tumor surgeries.

A follow-up study compared certain pathologic types of spinal cord tumors to identify which type easily affects the IONM alarm criteria [4]. Considering the signal pathways of MEPs and SEPs, the vulnerability to the tracts would differ by anatomical and pathologic types.

Theoretically, using combined MEPs and SEPs for all spinal cord tumor surgeries would be the most sensitive for postoperative neurologic complications. However, considering cost-effectiveness and time constraints, establishing a standard guideline would be helpful. Such a guideline should suggest each combination of evoked potentials, according to the different types of spinal cord tumors.

There have been many studies on IONM and spine surgery, but studies particularly focusing on spinal cord tumor cases are not common. Furthermore, very few studies have categorized spinal cord tumors by anatomical and pathologic type, or examined the reliability of IONM for each spinal cord tumor type.

To fill this gap in research, our study analyzes IONM cases of spinal cord tumor resection and identifies which combination of MEPs and SEPs is most reliable during spinal cord tumor surgery. We identify which combination is the most reliable by anatomical and pathologic type, which thereby enables us to better predict the postoperative motor deterioration.

MATERIALS AND METHODS

Subjects

Between November 2012 and August 2016, we monitored intraoperative MEPs or SEPs during 123 spinal cord tumor operations. Extensively experienced spinal surgeons performed all of these operations. We conducted a prospective analysis on the patient data, which included IONM records, medical records, operative narratives, anesthesia records, and outpatient clinical records. Preoperatively, 59 patients were neurologically intact and 64 patients had motor deteriorations. Table 1 lists the baseline characteristics of the patients.

Anesthesia

Rocuronium bromide Esmeron 50–150 mg (Han Wha Pharma Co. Ltd., Seoul, Korea) was administered intravenously as a short-acting muscle relaxant to facilitate endotracheal intubation. No paralytic agents were subsequently administered.

General anesthesia was induced by total intravenous anesthesia with remifentanil Ultiva (GSK, Middlesex, UK), propofol Fresofo1 (Fresenius Kabi, Seoul, Korea), or midazolam Vascam (Hana Pharm, Seoul, Korea) were used in several combinations to initiate and maintain the general anesthesia. During anesthesia, body temperature, direct radial artery pressure, pulse rate, oxygen saturation, and end-tidal carbon dioxide concentration were continuously monitored. All patients were kept normothermic and normotensive.

IONM monitoring techniques

A single technician performed IONM using the Cascade (Cadwell Industries Inc., Kennewick, WA, USA). In 110 out of 123 patients, both MEPs and SEPs were monitored. In the other 13 patients, only MEPs were monitored. Median SEPs were monitored in 38 patients, and tibial SEPs were monitored in 104 cases.

Somatosensory evoked potentials

SEPs were elicited by electrical stimulation of the median nerve at the wrist and the posterior tibial nerve at the
ankle (intensity 40 mA, duration 0.2 ms, with a repetition rate of 5 Hz). We recorded SEPs from needle electrodes placed on the scalp at C3 (right median nerve), C4 (left median nerve), and Cz (right or left tibial nerve) referenced to Fpz according to the 10-20 international electroencephalography (EEG) system.

### Motor evoked potentials

We obtained motor evoked potentials by multipulse transcranial electric stimulations using the Cascade electrical stimulator (Cadwell Industries Inc.). We recorded transcranial electric motor evoked potentials bilaterally from the abductor pollicis brevis muscles in upper extremities, and tibialis anterior muscles in the lower extremities using a pair of needle electrodes inserted 3 cm apart in each muscle. Needle electrodes delivered short trains of 6 square-wave stimuli of 0.5 ms duration, with an interstimulus interval of 5 ms. The needles delivered up to 2 Hz of repetition rate and were placed at C1 and C2, according to the 10-20 international EEG system. A C1/C2 montage is preferable for right-extremity MEPs, while C2/C1 is preferentially used to elicit left-extremity MEPs. To elicit lower-extremity MEPs, we used a Cz/Fz montage, which produces less intense muscle twitching. We gradually escalated the intensity of the stimulus by 50 mV increments (from 50 mV to a maximum of 400 mV) until MEP amplitudes were maximized above a minimum of 10 mV.

### Alarm criteria for evoked potentials

We performed neurophysiological monitoring throughout the surgical procedures. We obtained baseline EPs...
after at least 60 minutes after intubation and awaited the muscle relaxant effect to fade.

We continuously monitored values of N20 latencies of median SEPs, P37 latencies of tibial SEPs, and peak-to-peak amplitudes of MEPs. We considered a more than 50% reduction in amplitude, compared to the baseline MEPs, as a significant change [5,6]. We defined the prolongation of N20 or P37 latencies of more than 10% from the baseline SEPs as a significant change [7]. If any significant EP changes occurred, the surgeons were promptly informed and the surgical procedures were stopped temporarily until the values of EPs returned to normal. However, if no signal reversal occurred even after the surgical correction or after removing the spinal implant placed immediately prior to signal change, surgeons considered procedure cessation.

**Neurologic examination**

We evaluated the motor status of each patient before surgery, 48 hours after surgery, and then 4 weeks later. We measured the strengths of 10 key muscles of the International Standards for Neurological Classification of Spinal Cord Injury. We evaluated muscle strength using the Medical Research Council (MRC) scale range of 0 to 5. The total score ranged from 0 to 50 points on each side. We considered any motor reduction scoring with 1 point or more, compared to the preoperative evaluation, as ‘postoperative neurologic motor deterioration.’ We considered any postoperative weakness observed within 48 hours after surgery, which subsequently recovered after 4 weeks, as a ‘transient motor deterioration.’ If the decreased motor score at 48 hours did not recover in 4 weeks, we considered the patient to have ‘persistent postoperative motor deterioration.’

We conducted statistical analyses to reveal the correlations between the results of intraoperative changes of SEPs or MEPs and the motor status (intact, transient or persistent motor deterioration). We conducted the analyses according to each anatomical or pathologic type of spinal cord tumor.

**Statistical analysis**

We used an independent t-test to compare the mean age, height and sex between the patients with motor deterioration (transient and persistent) and the patients without motor deterioration. We calculated the sensitivity, specificity, positive and negative predictive values in each type of spinal cord tumor by anatomical lesion and pathology. We analyzed this data using SPSS ver. 20.1 (IBM, Armonk, NY, USA), with p-values of <0.05 considered statistically significant.

**RESULTS**

**Baseline characteristics of the subjects**

We enrolled a total of 123 spinal cord tumor patients in this study. Fourteen patients had shown motor deterioration, 8 had motor improvement, 101 had no motor change postoperatively, and 9 of postoperative motor deterioration patients had preoperative motor deterioration already (Table 2).

Table 1 displays the baseline demographic and clinical characteristics of the population. There were 53 male patients (43.1%) and 70 female patients (56.9%). The mean age of the patients was 48.3 years. Table 1 also compares the group that was neurologically intact postoperatively (group A) to the group with postoperative motor deteriorations (group B). There were no statistically significant differences in age and sex.

We classified the types of tumors by anatomy; there were 79 intradural extramedullary (IDEM) tumors; 31 intramedullary (IM) tumors; 11 extradural (ED) tumors; and 2 cases of IDEM combined with ED tumors. We also categorized the patients according to pathology: there were 51 schwannomas, 25 meningiomas, 15 ependymomas, 6 hemangiomas, 5 neurofibromas, 5 cystic lesions, 4 metastatic tumors, and 14 ‘others’ (Table 1).

**Postoperative motor deteriorations**

Among 123 patients, 109 patients showed no postoperative motor deterioration. Of the 14 patients with postoperative motor deterioration, 7 patients had significant

| Table 2. Preoperative and postoperative motor change of patients |
|---------------------------------------------------------------|
| Postoperative | Postoperative | Postoperative |
| motor improvement | no motor change | motor deterioration |
| (n=8) | (n=101) | (n=14) |
| Preoperative motor score | | | |
| 100 (n=59) | 0 | 54 | 5 |
| <100 (n=64) | 8 | 47 | 9 |
changes in both MEPs and SEPs, 5 patients had only MEP changes, and 2 patients had only SEP changes (Table 3).

Ten patients showed persistent motor deteriorations. We observed both MEP and SEP changes in 4 patients, MEP changes alone in 4 patients, and SEP changes alone in 2 patients (Fig. 1).

**Combinations of intraoperative neurophysiologic monitoring**
We monitored all patients in this study with intraoperative MEPs or SEPs. Both intraoperative MEPs and SEPs had no significant changes during surgery in 73 patients (59.4%). In 15 cases (12.2%), only amplitude of MEPs significantly decreased without significant changes of SEPs. In 23 cases, only latencies of SEPs were significantly prolonged (18.7%). We observed both significant changes in MEPs and SEPs in 12 cases (9.8%).

**Reliability of intraoperative neurophysiologic monitoring**
After comparing the patients’ motor status at 48 hours and at one month after surgery with the intraoperative EPs data, we calculated four measurements of reliability for each category of tumor by anatomy or pathology. Table 4 illustrates these calculations.

Overall, for MEP changes, sensitivity was 67.9%, specificity was 83.2%, positive predictive value was 48.7%, and negative predictive value was 91.7%. For SEP changes, sensitivity was 57.1%, specificity was 73.9%, positive predictive value was 34.0%, and negative predictive value was 88.0%. For both MEP and SEP changes, sensitivity was 25.0%, specificity was 95.8%, positive predictive value was 58.3%, and negative predictive value was 84.4%. For MEP or SEP changes, sensitivity was 100%, specificity was 61.3%, positive predictive value was 37.8%, and negative predictive value was 100% (Table 4).

As shown by this data, both overall positive and negative predictive values of MEPs were higher than the predictive values of SEPs. However, the positive predictive value was higher by the dual monitoring of MEPs and SEPs compared to MEPs alone.

**Analysis by anatomical type**
We calculated the reliabilities by anatomical type. Table 5 illustrates these calculations.

In ED tumors, for MEP changes, sensitivity was 66.7% and specificity was 100%. For SEP changes, sensitivity was 66.7% and specificity was 80.0%. There was no significant difference between sensitivity and specificity of MEP and SEP. However, in IM tumors, for MEP changes, sensitivity was 70.8% and specificity was 81.0%. For SEP changes, sensitivity were 54.2%, specificity was 71.4%.

### Table 3. Patients’ data with postoperative motor deteriorations after spinal cord tumor surgery

| Case no. | Age (yr)/sex | Pathologic diagnosis | Anatomical type | Level of lesion | EPs change | Motor deterioration |
|----------|--------------|----------------------|----------------|----------------|------------|-------------------|
| 1        | 75/F         | Schwannoma           | IDEM           | C3-4           | SEPs       | Persistent        |
| 2        | 24/M         | Anaplastic astrocytoma | IM            | C2-T1          | SEPs       | Persistent        |
| 3        | 61/M         | Ependymoma           | IM            | C5-7           | MEPs       | Persistent        |
| 4        | 54/M         | Ependymoma           | IM            | C2-4           | MEPs and SEPs | Persistent      |
| 5        | 34/M         | Ependymoma           | ED            | T11-12         | MEPs and SEPs | Persistent      |
| 6        | 44/F         | Ependymoma           | IM            | C2-T2          | MEPs       | Persistent        |
| 7        | 56/F         | Cavernous hemangioma | IM            | T11-12         | MEPs and SEPs | Persistent      |
| 8        | 43/M         | Cavernous hemangioma | IM            | T6             | MEPs and SEPs | Persistent      |
| 9        | 58/M         | Ependymoma           | IM            | C2-7           | MEPs       | Persistent        |
| 10       | 24/F         | Ependymoma           | IM            | T10-11         | MEPs       | Persistent        |
| 11       | 44/F         | Ependymoma           | IM            | C2-T4          | MEPs and SEPs | Transient      |
| 12       | 49/F         | Ependymoma           | IM            | T7-8           | MEPs and SEPs | Transient      |
| 13       | 39/M         | Ependymoma           | IM            | C4-7           | MEPs       | Transient        |
| 14       | 29/F         | Cystic glioma        | IM            | C2-5           | MEPs and SEPs | Transient      |

MEPs, motor evoked potentials; SEPs, somatosensory evoked potentials; IDEM, intradural extramedullary; IM, intramedullary; ED, extradural.
We can conclude from this data the specificity and sensitivity of MEPs for the postoperative motor deterioration was superior to SEPs in intramedullary tumors. Both positive and negative predictive values of MEPs in intramedullary tumor surgery were higher than with SEPs (Table 5).

**Analysis by pathologic type**

The specificity of MEPs for the postoperative motor deterioration was superior in schwannoma and meningioma. In ependymoma, the sensitivity of MEPs was superior to SEPs, but the specificities were the same. However, the specificity of SEPs was superior to MEPs for hemangioma, and the sensitivity was higher with SEPs alone than with both MEP and SEP monitoring.

In some tumor types, we were unable to calculate the reliabilities of each value of EP monitoring due to the absence of a motor deterioration case (Table 6). In ependymoma, the positive and negative predictive values were higher with MEPs than SEPs, and even higher than the dual monitoring of both MEPs and SEPs. In hemangioma, SEPs showed a 100% positive predictive value, and the negative predictive value was also higher with SEPs than MEPs.

**DISCUSSION**

In this study, we included only patients with spinal cord tumors, with a sample size of 123. The male to female ra-
Spinal cord tumors occur within or nearby the spinal cord, and account for 2%-4% of CNS tumors. The prevalence of certain types of tumors in our study reflected the type found in the general population. The three most common types of tumors in the United States and Japan are meningioma, schwannoma, and ependymoma, in descending order [8,9]. Among the cases included in this study, we also found these three to be the most common tumors, although with differing rates of abundance. Schwannoma was the most common (45.5% of cases), followed by meningioma (18.7%), and ependymoma (12.2%). Differences in abundance may have occurred because we only included cases for which IONM was carried out. In particular, tumors of the spinal nerve accounted for the highest percentage of cases. IONM is more frequently carried out in function-sparing operations, where there is a benign and slow-growing tumor [10].

Our study has some strong points. First, we analyzed the spinal cord tumors by two different categories, considering the anatomical and pathologic types. Second, in each type of tumor, we calculated and compared the value for reliability. Third, we were able to verify the superiority of SEPs in correlation with the postoperative motor deterioration in some tumor types.

We subdivided spinal cord tumors by anatomical site. Intradural extramedullary tumors are located outside the spinal cord but within the dural sheath. Intramedullary spinal cord tumors refer to a subgroup of intradural spinal tumors that arise from cells within the spinal cord, as opposed to adjacent structures, such as the nerve roots or meninges. Extradural tumors are located outside the dura mater lining [11-13].

Anatomically, MEPs monitor the corticospinal tract descending from motor cortex to the anterior horn of the spinal cord and peripheral nerve fibers [14]. The SEPs represent the function of the dorsal column-lemniscal pathway of the somatosensory system. The pathway consists of the somatosensory cortex, ventroposterior nuclei of thalamus, medial lemniscus, dorsal column, and peripheral nerve fibers [7]. Likewise, EPs’ neural pathways consist of the spinal cord of the intramedullary portion and the peripheral nerve fibers of the extramedullary portion. Thus, we suggested that intraoperative neurophysiological monitoring of spinal cord tumor surgery should consider the anatomical location of tumors.

We also categorized spinal cord tumor cases by pathologic type. Some tumors like schwannoma, hemangioma,

| Table 5. Reliabilities of each combination of intraoperative MEPs and SEPs for the postoperative motor deterioration by anatomical type of spinal cord tumor |
|---------------------------------------------------------------|
| Significant changes | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|----------------------|-----------------|-----------------|---------|---------|
| IDEM tumor (n=79)    |                 |                 |         |         |
| MEPs only            | NA              | 81.4            | NA      | 98.6    |
| SEPs only            | 100             | 73.3            | 4.2     | 100     |
| Both MEPs and SEPs   | NA              | 95.3            | NA      | 98.8    |
| Either MEPs or SEPs  | 100             | 59.3            | 2.8     | 100     |
| IM tumor (n=31)      |                 |                 |         |         |
| MEPs only            | 70.8            | 81.0            | 81.0    | 70.8    |
| SEPs only            | 54.2            | 71.4            | 68.4    | 57.7    |
| Both MEPs and SEPs   | 25.0            | 95.2            | 85.7    | 52.6    |
| Either MEPs or SEPs  | 100             | 57.1            | 72.7    | 100     |
| ED tumor (n=11)      |                 |                 |         |         |
| MEPs only            | 66.7            | 100             | 100     | 90.9    |
| SEPs only            | 66.7            | 80.0            | 50.0    | 88.9    |
| Both MEPs and SEPs   | 33.3            | 100             | 100     | 83.3    |
| Either MEPs or SEPs  | 100             | 80              | 60.0    | 100     |

MEPs, motor evoked potentials; SEPs somatosensory evoked potentials; PPV, positive predictive value; NPV, negative predictive value; IDEM, intradural extramedullary; IM; intramedullary; ED, extradural; NA, not available.
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Table 6. Reliabilities of each combination of intraoperative MEPs and SEPs for the postoperative motor deterioration by pathologic type of spinal cord tumor

| Significant changes | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|---------------------|-----------------|-----------------|---------|---------|
| Schwannoma (n=51)   |                 |                 |         |         |
| MEPs only           | NA              | 85.2            | NA      | 97.9    |
| SEPs only           | 100             | 74.1            | 6.7     | 100     |
| Both MEPs and SEPs  | NA              | 96.3            | NA      | 98.1    |
| Either MEPs or SEPs | 100             | 63.0            | 4.8     | 100     |
| Meningioma (n=25)   |                 |                 |         |         |
| MEPs only           | NA              | 80.0            | NA      | 100     |
| SEPs only           | NA              | 68.0            | NA      | 100     |
| Both MEPs and SEPs  | NA              | 96.0            | NA      | 100     |
| Either MEPs or SEPs | NA              | 52.0            | NA      | 100     |
| Ependymoma (n=15)   |                 |                 |         |         |
| MEPs only           | 76.5            | 62.5            | 81.3    | 55.6    |
| SEPs only           | 47.1            | 62.5            | 72.7    | 35.7    |
| Both MEPs and SEPs  | 23.5            | 87.5            | 80.0    | 35.0    |
| Either MEPs or SEPs | 100             | 37.5            | 77.3    | 100     |
| Hemangioma (n=6)    |                 |                 |         |         |
| MEPs only           | 66.7            | 75.0            | 80.0    | 60.0    |
| SEPs only           | 66.7            | 100             | 100     | 66.7    |
| Both MEPs and SEPs  | 33.3            | 100             | 100     | 50.0    |
| Either MEPs or SEPs | 100             | 75.0            | 85.7    | 100     |
| Neurofibroma (n=5)  |                 |                 |         |         |
| MEPs only           | NA              | 80.0            | NA      | 100     |
| SEPs only           | NA              | 80.0            | NA      | 100     |
| Both MEPs and SEPs  | NA              | 100             | NA      | 100     |
| Either MEPs or SEPs | NA              | 60.0            | NA      | 100     |
| Cystic lesions (n=5)|                 |                 |         |         |
| MEPs only           | 66.7            | 100             | 100     | 80.0    |
| SEPs only           | 66.7            | 100             | 100     | 80.0    |
| Both MEPs and SEPs  | 33.3            | 100             | 100     | 66.7    |
| Either MEPs or SEPs | 100             | 100             | 100     | 100     |
| Metastatic tumors (n=4)|             |                 |         |         |
| MEPs only           | NA              | 100             | NA      | 100     |
| SEPs only           | NA              | 100             | NA      | 100     |
| Both MEPs and SEPs  | NA              | 100             | NA      | 100     |
| Either MEPs or SEPs | NA              | 100             | NA      | 100     |

MEPs, motor evoked potentials; SEPs, somatosensory evoked potentials; PPV, positive predictive value; NPV, negative predictive value; NA, not available.

and neurofibroma are usually benign forms, while other tumors like astrocytoma, multiple myeloma, small round cell tumor are malignant forms. In the case of benign spinal cord tumors, the radical resection of the tumor is usually done for curative purposes [15,16]. On the other hand, in the case of malignant spinal cord tumors, tumor resection is usually performed for palliative management [17,18]. Moreover, the range of invasion in surgical procedure would be different according to the pathologic and anatomical type of tumor.
For postoperative neurologic deterioration, the alert criteria for intraoperative MEP changes vary from 50% to 75% in amplitude reduction [4]. Some studies included amplitude criterion in addition to the latency criterion for SEP monitoring [4,19,20]. The specific alarm criteria in our spine hospital are a more than 50% reduction in MEP amplitude, and a more than 10% of N20 or P37 latency prolongation for SEPs. As the alarm criteria vary with each individual study, precise comparing of each EP value for reliability is not easy. Nevertheless, we conclude the sensitivity and specificity of MEPs for postoperative weakness was superior to that of SEPs in IONM during spine tumor surgery. Thus, we should consider why we perform intraoperative SEPs monitoring.

In this study, 2 out of 14 cases with motor deterioration showed no significant changes in MEPs, and only SEPs showed significant changes. Therefore, it seems worthwhile to perform SEP monitoring to increase the detection rate for postoperative motor deterioration, and decrease the number of false negatives [21,22]. Further, as the stimulations for MEPs make muscles contract and patients move, intraoperative MEP monitoring is usually performed only periodically when there is a major procedure. On the other hand, SEPs can be monitored relatively continuously, since the stimulations can be applied in distal area with relatively small intensity, compared to MEPs. In addition, across all cases, SEPs may be more reliable for some categories of tumor types in our study, although MEPs showed superior sensitivity and specificity in general spine surgery [15,23]. Although we are presenting a 100% positive predictive value of SEPs in hemangioma surgery, there may be a bias due to the small sample size. Further study with a larger sample size is needed.

SEP monitoring is known to be affected by blood flow. When the blood flow to neural tissue drops below 18 mL/100 g/min, the SEP values start to change. Specifically, the SEP amplitudes decrease and the SEP latencies prolong. Indeed, when the blood flow to neural tissue drops below 15 mL/100 g/min, SEP loss occurs [24,25]. However, if the SEP loss begins just before the critical threshold of persistent neurologic damage, intraoperative SEP loss can be an early warning sign for neurologic damage [21]. Thus, intraoperative SEP monitoring would be more important than MEP’s in hemangioma, a highly vascularized tumor, which can lead to more blood loss during a removal operation than other types of tumors.

There are a few limitations of this study, which include the small number of patients enrolled in this study, and lack of long-term follow-up to assess neurological deteriorations. In addition, this study did not consider sensory problems as another neurologic deterioration. We only took into account SEP latencies, with no consideration of the changes in amplitudes. Further studies with a greater number of cases for each pathologic and anatomical type would bring more concise results.

In conclusion, dual IONM of both MEPs and SEPs in all cases would be ideal. However, if one must be selected due to cost or time, the selection must depend on the case at hand.

This study reconfirms the conclusion of previous studies that the highest reliability can be achieved by performing both MEPs and SEPs simultaneously. In our previous study, we reported only SEP changes predicted persistent motor deterioration in spinal cord tumor surgery, and a combined MEP/SEP monitoring showed high sensitivity and specificity in spinal deformity surgery [20]. To follow up on our previous study, we now make findings on the different sensitivities and specificities of MEPs and SEPs observed for each type of spinal cord tumor by anatomical and pathological type. This information is useful for selecting appropriate monitoring modalities for each type of surgery. Specifically, SEPs are less reliable as an overall indicator of neural deteriorations, when compared to MEPs in most types of spinal cord tumor surgeries. However, SEPs would be superior in some types of tumors, especially in hemangioma.

CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

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