Neurophysiological Monitoring in the Endovascular Therapy of Aneurysms

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BACKGROUND PURPOSE: Endovascular aneurysm therapy has associated risks of ischemic complications. We undertook this study to evaluate the efficacy of neurophysiological monitoring (NPM) techniques in the detection of ischemic changes that may be seen during endovascular treatment of cerebral aneurysms.

METHODS: Thirty-five patients underwent NPM during endovascular treatment of cerebral aneurysms. The patients underwent a total of 50 endovascular procedures, including balloon test occlusion (19 patients), GDC embolization (22 patients), and permanent vessel occlusion (nine patients). NPM included electroencephalography, somatosensory evoked potentials, and/or brain stem auditory evoked potentials, depending on the location of the aneurysm.

RESULTS: NPM changes were seen in nine (26%) of 35 patients and altered the management in five (14%) of 35 patients. In three of the five cases, NPM changes were observed without corresponding neurologic physical examination changes after balloon test occlusion (performed while the patients were under general anesthesia in two cases). In the two other cases in which NPM changes altered management, ischemia was detected at the time of intra-aneurysmal therapy while the patients were under general anesthesia. Overall, 18 of 35 patients underwent a total of 19 balloon test occlusion procedures. Of the 17 remaining patients, 13 underwent aneurysm coiling, two were not treated because of inability to safely place coils, and two were treated for distal aneurysms. Two patients developed transient neurologic deficits without concurrent NPM changes, representing false-negative NPM test results.

CONCLUSION: NPM is a valuable adjunct to endovascular treatment of cerebral aneurysms. Our study suggests that these monitoring techniques may reduce ischemic complications and can be used to help guide therapeutic decisions.
bolization, and basilar artery thrombolysis. The purpose of our study was to assess the efficacy of multimodality NPM in detecting significant ischemia during endovascular treatment of cerebral aneurysms at the time of intra-aneurysmal GDC coil embolization or permanent vessel occlusion.

Methods

The study was a retrospective evaluation of patients treated during a 24-month period ending in June 1999. Thirty-five patients ranging in age from 33 to 81 years (mean age, 59.9 years) who underwent NPM during endovascular treatment of cerebral aneurysms were included. At our institution, personnel for NPM are available for nearly all elective and emergency cases. Many of the patients treated in this study were treated on an emergency basis. At the time of the study, only one physician was actively involved in NPM; currently, two are actively involved. For this reason, two patients who received endovascular therapy during the study period did not undergo NPM because it was unavailable at the time of treatment. They were therefore excluded from the study.

Of the 35 patients included in the study, 33 were deemed to be poor surgical candidates because of age, aneurysm size/morphology, or preexisting medical conditions and two had requested endovascular aneurysm treatment. Eleven of 35 patients had ruptured aneurysms, whereas 24 of 35 had unruptured aneurysms. The demographic characteristics of these patients are summarized in Table 1.

All GDC embolization procedures were performed with the patients under general anesthesia to minimize patient motion. Conscious sedation was used whenever possible for patients who required balloon test occlusion of the parent vessel before definitive treatment, to perform neurologic examination testing in addition to NPM. Patients who tolerated the temporary balloon occlusion were often then placed under general anesthesia, for either GDC embolization or permanent vessel occlusion, and underwent continuous NPM. For four patients, conscious sedation was not possible during balloon test occlusion and general anesthesia was used. Therefore, only NPM (without neurologic physical examination) was performed in these cases.

Electrophysiological Monitoring

NPM included EEG, somatosensory evoked potentials (SSEPs), and/or brain stem auditory evoked potentials (BAEPs), based on the location of the aneurysm and vascular territory at risk (Table 1). In cases in which the internal carotid artery and/or middle cerebral artery vascular territory was at risk, NPM was accomplished by using a combination of EEG and SSEP testing (performed after bilateral median nerve stimulation). Anterior cerebral artery territory aneurysm treatment (especially anterior communicating artery) was monitored with SSEP testing by using bilateral median and posterior tibial nerve stimulation. Vertebral and basilar artery vascular supply required a combination of median nerve-generated cortical SSEPs and BAEPs, which were acquired after bilateral independent stimulation.

In elective cases, preoperative EEG, SSEP, and/or BAEP testing was performed in addition to a thorough neurologic examination during a clinic visit. No adverse effects were observed in those patients who underwent preoperative evoked potential testing. In emergency cases, pre-procedural baseline monitoring was performed while the patient was on the angiography table and a limited clinical neurologic examination was also performed of conscious patients.

A Nicolet Viking IV Electrodiagnostic System (Nicolet Instrument Corporation, Madison, WI) was used to obtain the evoked potentials. Standard surface EEG electrodes were used and placed, by using collodion (except for Erb’s point), accord-
Table 2 lists the patients for whom NPM or physical examination changes were identified and describes when these changes altered our treatment decisions. NPM changes were observed for nine (26%) of 35 patients and altered management for five (14%) of 35 patients. For three of the five patients, NPM changes were observed without corresponding neurologic physical examination changes after balloon test occlusion (performed with the patient under general anesthesia in two of three cases). For the
other two of five patients for whom NPM changes altered management, ischemia was detected at the time of intra-aneurysmal therapy while patients were under general anesthesia.

NPM changes indicating cerebral ischemia altered treatment decisions for five patients as follows: 1) permanent vessel occlusion was delayed for 6 weeks to allow for maturation of an extracranial-to-intracranial bypass graft (patient 5), 2) permanent vessel occlusion was performed in the right vertebral artery rather than the basilar artery (patient 11), 3) GDCs were not deployed in an aneurysm because of ischemic changes that occurred with coil placement (patient 26) (Fig 1), 4) mean arterial pressure was increased during superselective catheterization to increase cerebral perfusion (patient 7), and 5) perma-
A. Lateral projection angiogram of the left internal carotid artery shows the aneurysm arising in the supraclinoid segment shortly after the takeoff of the ophthalmic artery. The aneurysm was thought to be unfavorable for GDC embolization, and balloon test occlusion was thus performed in anticipation of permanent vessel occlusion. Because of the patient’s physical condition, the procedure could be performed only with the patient under general anesthesia.

B. Baseline cerebral SSEPs after bilateral median nerve stimulation. Note the baseline asymmetry, with the right median nerve SSEP (arrow) being smaller in amplitude than the left. Top tracing, left median nerve stimulation (ie, right brain); bottom tracing, right median nerve stimulation (ie, left brain) (arrow).

C. Left internal carotid artery balloon test occlusion was performed, resulting in gradual amplitude reduction of the left cerebral (right median nerve stimulation) SSEP over 5 min and precipitous decrease in the 6th min. The SSEP obtained 6 min after balloon occlusion shows a nearly complete loss of the left cerebral SSEP (arrow). Top tracing, left median nerve stimulation (ie, right brain); bottom tracing, right median nerve stimulation (ie, left brain) (arrow).

D. Balloon was deflated immediately after the SSEP tracing shown in 2C. The cerebral SSEP returned to baseline amplitude levels after 1 min (arrow). Top tracing, left median nerve stimulation (ie, right brain); bottom tracing, right median nerve stimulation (ie, left brain) (arrow).

tent vessel occlusion was not performed in wide neck aneurysms (patient 27) (Fig 2). The NPM changes resolved and returned to normal or nearly normal in all five cases, and none of these patients had permanent neurologic deficits.

Among the four patients for whom NPM changes did not alter treatment decisions, the reasons were as follows: 1) aneurysm rupture (in two cases) required continuation of the procedure despite evidence of cerebral ischemia (patients 9 and 29); 2) a small clot was seen in the M1 segment of the middle cerebral artery during ipsilateral internal carotid artery aneurysm coiling, but thrombolysis was not attempted because of aneurysm perforation that had occurred during aneurysm coiling (patient 4); and 3) diffuse NPM changes due to cerebral vasospasm were identified with successful aneurysm coiling, and the decision was made to treat the vasospasm systemically (patient 30).

None of the patients in this series had both positive NPM and physical examination changes. Two patients did, however, develop transient neurologic deficits without concurrent NPM changes at the time of the procedure: 1) facial droop during balloon test occlusion (patient 10), and 2) new quadrantanopia after sodium amytal in the appropriate vascular territory (patient 33). The transient deficits resolved completely within 10 min. These were classified as false-negative NPM results, even though the affected cortical territory did not correspond to the cerebral distribution tested by the evoked potentials.

Six (11.5%) delayed ischemic complications in patients with no NPM changes occurred after therapy, at 15 hr to 5 days after treatment, among our 52 procedures. In five cases, radiologic follow-up showed these infarcts to be embolic (generally small cortical lesions) and they were therefore not considered false-negative responses to NPM. In one case involving a fusiform aneurysm of the basilar artery, a complication due to hypoperfusion was initially suspected but was also later shown to be embolic in nature: a left posterior inferior cerebellar artery territory infarction occurred after the patient discontinued his regimen of warfarin anticoagulation.

Discussion

To our knowledge, this study represents the first series describing the routine use of NPM during endovascular treatment of cerebral aneurysms. NPM including EEG and evoked potential is particularly valuable when the patient must be placed under general anesthesia for balloon test occlusion or for GDC aneurysm embolization because concurrent neurologic examination is not possible. Alternative approaches to balloon test occlusion have been published (1). These include the use of xenon CT to assess cerebral perfusion after a clinical neurologic examination with the balloon inflated. This has the disadvantage of requiring the patient to be transported to the CT scanner, where the balloon is again inflated in a blinded fashion.

Neurophysiological techniques can directly assess the functional state of specific cerebral regions and are indirect measures of local ischemia, which can be continuously monitored during endovascular aneurysm treatment. Several studies have documented the correlation between the NPM changes and cerebral ischemia. In patients undergoing carotid endarterec-
found that major EEG changes occurred with rCBF <10 mL/100 g/min, with the critical level defined as 15 mL/100 g/min. Primate models show that SSEPs are maintained at levels of rCBF >16 mL/100 g/min but are absent at levels of rCBF <12 mL/100 g/min, with sharp reductions in the cortical SSEP amplitude (50% of baseline level) at rCBF levels between 14 and 16 mL/100 g/min (3, 5, 19, 20). Central conduction time is also prolonged with cerebral ischemia at an rCBF threshold of approximately 15 mL/100 g/min (21, 22). Still other primate studies have shown that cerebral infarction occurred at an rCBF threshold of 12 mL/100 g/min maintained for >2 hr (23, 24). These findings suggest that a 50% reduction in amplitude of the SSEP or a prolonged central conduction time >10 ms corresponds to an rCBF of 14 to 16 mL/100 g/min and is indicative of ischemia and possible progression to infarction. The usefulness of NPM has also been shown during surgical treatment of cerebral aneurysms (25).

This study evaluates the use of NPM to detect ischemia and alter management decisions during endovascular treatment of cerebral aneurysms; as such, it does not focus on clinical outcomes. Most of the patients in this study were deemed poor surgical candidates. Clinical outcomes were primarily influenced by the patient’s baseline state, including preexisting medical condition, by atherosclerotic disease in the cerebrovascular circulation, and by complication of the primary disease, such as severe subarachnoid hemorrhage and/or vasospasm. NPM changes altered treatment decisions for 14% of our patients (five of 35 patients). None of these five patients developed postprocedural neurologic deficits due to ischemia. In another 11% of the patients (four of 35 patients), treatment decisions were not altered despite NPM changes because an aneurysm rupture had occurred (three cases), requiring further GDC embolization of the aneurysm, or because diffuse NPM changes due to severe cerebral vasospasm were present.

The six ischemic events that resulted in permanent mild or moderate neurologic deficits in the study population were delayed complications occurring 15 hr to 5 days after therapy. Pelz et al (26) reported a 28% incidence of thromboembolic complications after GDC embolization, with a 5% incidence of permanent neurologic deficits. These are recognized complications for both GDC embolization and parent artery occlusion occurring after the procedure and do not diminish the potential value of NPM, which detects ischemic changes during the procedure.

We advocate a multimodality approach to NPM. The type of NPM is determined by the location of the aneurysm and its associated vascular territory, and the setup is critical. A complete description of the neurophysiological monitoring techniques is beyond the scope of this article but is described in a review article by Lopez (27). We use a combination of EEG, SSEPs, and BAEPs in the neuroangiography suite. The combination used is dependent on the vascular territory at risk and the patient’s baseline neurologic examination. Each technique has advantages and disadvantages, and it is best to use complementary modalities in combination.

The two cases in our study in which neurologic deficits were identified based on physical examination without corresponding NPM emphasize that NPM should not serve as a substitute but rather a complement to concurrent clinical neurologic testing whenever possible. We speculate, however, that NPM changes lagged behind the physical examination findings and would have subsequently become evident had our management not been altered. Unfortunately, the NPM literature does not yet address the issue of temporal correlation between onset of NPM changes and onset of neurologic deficits. In our study, several cases showed neurophysiological changes immediately after balloon inflation or attempted coil deployment. The literature also does not provide a definitive answer regarding how long NPM changes can persist before ischemia progresses to infarction. One study evaluated temporary occlusion in aneurysm surgery (28). In this study, the mean occlusion time was 20.3 min for the patients with middle cerebral artery aneurysms and 15.8 min for the patients with internal carotid artery aneurysms. The SSEP disappeared during occlusion in 42 of 97 patients (30 middle cerebral and 12 internal carotid arteries). All except three eventually returned to baseline after recirculation, and none of the 39 patients experienced postoperative sequelae. The time period from the start of occlusion until the complete loss of the SSEP averaged 8.6 min among these 39 patients, and the occlusion time from total SSEP loss until recirculation averaged 12 min. For the remaining three of the 42 patients (all three of whom had internal carotid artery aneurysms), however, the SSEP did not recover after recirculation, and all three patients experienced postoperative sequelae.

One limitation of NPM is that it may be relatively insensitive in detecting ischemic changes in some vascular territories, such as in the cerebellum or posterior cerebral artery territories, because NPM monitors the integrity of only specific sensory pathways and is dependant on somatosensory cortex and afferent pathways to generate the evoked response. SSEPs that use median and posterior tibial nerve stimulation are useful in assessing the functional state of the middle and anterior cerebral artery territories (by detecting impairment along the respective somatosensory pathways). BAEP monitoring detects functional changes along the auditory brain stem pathways. This is most often caused by a brain stem insult, which could result from vertebrobasilar ischemia. However, ischemia in the cerebellum or posterior cerebral artery territories could still be missed. EEG provides a more global assessment of cerebral ischemia but cannot be used to monitor the posterior fossa. It may be used to detect ischemia in the posterior cerebral artery territory, but its sensitivity is limited. Although the sensitivity of EEG could potentially be improved by increasing the number of electrodes used for monitoring, only a limited number of
electrodes can be placed on the patient’s head (usually six) because the procedure is performed under fluoroscopy and more electrodes would obscure optimal working projections for endovascular treatment. In addition, the most commonly used monitoring equipment accepts only a limited number of electrode inputs.

Other technical limitations include confounding anesthesia-related effects, which may mimic cerebral ischemia. The effects of specific anesthetic agents are discussed in detail in the review article by Lopez (27). These effects can generally be distinguished from true ischemia by recognizing typical waveform characteristics (eg, bilateral rather than unilateral waveform changes) and by recognizing changes either in the type or dose of anesthetic agent used. Finally, evoked potential changes cannot be monitored at the time of coil release because of the nature of the electrolytic coil detachment, which interferes with the acquisition of the evoked potential. However, evoked potentials can be measured when the decision to detach the coil is made and can be monitored afterward. Also, with the advent of more rapidly detaching electrolytic coils, this drawback has been further diminished. Despite these limitations, we think that our experience shows the value of routine NPM during endovascular treatment of cerebral aneurysms, with NPM changes detected for 26% of patients treated and with NPM changes altering treatment decisions for 14%.

NPM is performed under the direction of one of the neurologists in the Department of Neurology and Neurologic Sciences. The neurologist supervising the NPM will typically examine the patient preoperatively and also perform a physical examination. The NPM team is composed of one neurodiagnostic technologist, who was originally trained as an EEG and evoked potential technologist and later received specialized training in intraoperative monitoring, and a neurologist trained in clinical neurophysiology and preferably intraoperative monitoring. An EEG/evoked potential machine is used to record the electrophysiological signals during the case. It is usual for the technologist to remain in the room for the duration of the case and for the neurologist to be present at the beginning of the procedure, during the time of balloon occlusion, and at the end of the case. The cost incurred for NPM includes an hourly hospital technical and physician professional fee. At our institution, this has been estimated to be approximately 10% of the cost of the neurointerventional procedure. NPM does not significantly prolong the interventional procedure because most of the patient electrode setup and the preoperative neurologic examination have been performed before the patient arrives in the radiologic suite. In addition, the time required to obtain baseline evoked potentials and EEG is approximately 5 min. NPM performed during the endovascular procedure usually does not require that the embolization procedure be interrupted.

In conclusion, NPM is a valuable adjunct to endovascular treatment of cerebral aneurysms. NPM may be particularly useful in situations in which neurologic examination is not possible (such as when the patient is under general anesthesia) or when a patient’s condition (such as obtunded subarachnoid hemorrhage) precludes neurologic testing. Our study suggests that these monitoring techniques may reduce ischemic complications and can be used to help guide therapeutic decisions.

References


