Low-dose adenosine-induced transient asystole during intracranial aneurysm surgery

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ABSTRACT

Background: Few studies have evaluated the adenosine dose that induces cardiac arrest during intracranial aneurysm surgery. We present our experiences with adenosine-induced transient asystole (AiTA) during intracranial aneurysm surgery and dosage recommendations.

Methods: We retrospectively reviewed the medical records of all patients who underwent intracranial aneurysm surgery between July 2016 and December 2018. Patients who experienced AiTA during intracranial aneurysm surgery were included in the study.

Results: Our study included nine intracranial aneurysm surgeries performed in eight patients. Thirteen episodes of AiTA were reported. Five of these were performed to facilitate bleeding control due to intraoperative aneurysm rupture (IAR), and adenosine doses were 9 mg (0.20 mg/kg), 12 mg (0.25 mg/kg), 12 mg (0.26 mg/kg), 18 mg (0.34 mg/kg), and 18 mg (0.39 mg/kg), resulted in transient asystole for 12, 14, 9, 44, and 18 s, respectively. For episodes without IAR, adenosine doses ranging from 6 to 18 mg (0.11–0.39 mg/kg) caused asystole for 8–33 s. In five episodes without IAR, low-dose adenosine (lower than 0.2 mg/kg) was used and caused asystole ranging from 8 to 12 s. Postoperatively, two patients had elevated cardiac troponin T levels but normal electrocardiograms.

Conclusion: AiTA can facilitate the clipping of intracranial aneurysms at low-risk of serious cardiac complications. An adenosine dose of 0.2–0.4 mg/kg is safe and effective in both IAR and non IAR situations. In non IAR cases, we propose that low-dose AiTA is an option to facilitate aneurysm clipping. A starting dose of 6 mg or 0.1–0.2 mg/kg can adequately induce brief asystole by softening the aneurysmal sac during clip application.

Keywords: Adenosine, Cerebral aneurysm, Induced transient asystole, Subarachnoid hemorrhage, Surgery

INTRODUCTION

During the intraoperative aneurysm rupture (IAR), the proximal control and cardiac standstill have been used to facilitate bleeding control. Several transient cardiac standstill techniques have been evolved including deep hypothermia with circulatory arrest on cardiopulmonary bypass, rapid ventricular pacing, and intraoperative adenosine-induced transient asystole (AiTA).

AiTA was proved to be an effective and safe technique to minimize bleeding caused by IAR. Recently, AiTA was shown to be useful as an alternative technique for the decompression of aneurysms during clip application. Several studies have demonstrated the utility of AiTA in facilitating intracranial
aneurysm clip ligation and its safety regarding perioperative and postoperative outcomes. There was no report on AiTA-related death. One serious complication was reported that the patient required closed cardiac chest compression due to prolonged extreme hypotension (>5.5 min) after the rapid redosing of adenosine due to IAR. After 3 min, the spontaneous circulation was returned, and chest compression was stopped.

The previous studies have used various starting doses of adenosine. However, few have explored its optimal dosage. We present a case series of using AiTA and determine the appropriate dose of adenosine for IAR and without IAR (non IAR) situations during intracranial aneurysm surgery.

MATERIALS AND METHODS

This study was approved by our institutional review board. We retrospectively reviewed the medical records, radiographic characteristics, anesthetic records, and operative records of all patients who underwent intracranial aneurysm surgery between July 2016 and December 2018. Patients who received AiTA were included in the study.

The anesthesiologists performed general anesthesia. The anesthetic management included arterial catheterization and standard monitoring, and external pacing-defibrillator pads were prepared. The patient was inducted with fentanyl and propofol, and cisatracurium was used to facilitate endotracheal intubation. Anesthesia was maintained with a propofol infusion and/or inhaled sevoflurane (<0.5 minimum alveolar concentration) and a fentanyl bolus. A bolus dose of IV nicardipine was used to control hypertension. The dose of adenosine was administered in an antecubital vein as an intravenous bolus followed by a rapid saline infusion. Repeat doses were administered after circulation returned to baseline values. The decision to use adenosine was made by the surgeon.

The following data were collected: sex, age, comorbidities, height, weight, aneurysm location, size, rupture status, dose of adenosine, reason for adenosine use, hemodynamics after the adenosine bolus, success of the clipping facilitated by AiTA, surgeon satisfaction, and postoperative complications. For cardiovascular complications, the electrocardiograms (EKGs) and cardiac enzymes were reviewed.

The descriptive data are presented as the mean ± SD for normally distributed data, as the median (range) for continuous variables, and as the count (percentage) for categorical variables.

For each patient, a scattergram of the adenosine dose and the duration of asystole was constructed. The relationship between the dose of adenosine (mg per ideal body weight [IBW] in kg) and the duration of asystole was examined by a linear regression analysis.

RESULTS

We identified 26 patients who underwent 28 intracranial aneurysm surgeries in our institutions between July 2016 and December 2018. Of these 26 patients, eight who underwent nine intracranial aneurysm surgeries experienced 13 episodes of AiTA. These patients included six females and two males with a mean age of 61.6 ± 5.4 (range 55–72) years. The patient data and aneurysm characteristics are summarized in [Table 1]. The median maximum diameter of the aneurysm was 5.1 (range 2.0–8.9) mm. The location of the aneurysm was the middle cerebral artery in one patient, the anterior communicating artery in two patients, and the posterior communicating artery (Pcom) in six patients.

The details of the patients and episodes of AiTA are shown in [Table 2]. All of the patients had hypertension. Seven patients preoperatively had sinus rhythm. Only one patient (patient #1) had an EKG that revealed premature atrial contractions (PACs). None of the patients were previously diagnosed with cardiac and pulmonary disease.

The majority of cases were emergency surgery due to ruptured aneurysms. The patient #3 harboring bilateral Pcom aneurysms presented with a subarachnoid hemorrhage. We performed two separate clipping surgeries. The first clipping was for the ruptured side, while the second operation was an elective clipping for the contralateral unruptured aneurysm.

Of these 13 episodes of AiTA, five episodes of AiTA were used during an IAR to facilitate bleeding control. The other eight episodes of AiTA without IAR (Non IAR) were used as an alternative to temporary clipping.

The dose of adenosine ranged from 3 to 18 mg (0.06–0.39 mg/kg IBW). The duration of asystole ranged from 0 to 44 s. Four patients required a second dose of adenosine (patient #1, #4, #6, and #7), one for further dissection or clip adjustment (patient #6), and three to facilitate bleeding.

| Table 1: Demographic data of eight patients with nine aneurysm characteristics. |
|---------------------------------------------|--------------|
| Sex (M/F) | 2/6 |
| Age (year), mean±SD | 61.6±5.4 |
| Presentation (unruptured/ruptured) (n) | 2/7 |
| Size (mm), median (range) | 5.1 (2.0–8.9) |
| Location (n) | |
| Middle cerebral artery | 1 |
| Anterior communicating artery | 2 |
| Posterior communicating artery | 6 |
control due to IAR (patient #1, #4, and #7). One patient with an adenosine dose of 3 mg (0.06 mg/kg IBW) did not achieve asystole. However, the systolic blood pressure decreased from baseline for 10 s, thereby allowing for a successful aneurysm clipping. Intraoperative ruptures did not occur in any case during adenosine-induced flow arrest.

For those with IAR (patient #1, #3, #4, and #7), the dose of adenosine was 9, 12, 12, 18, and 18 mg (0.20–0.39 mg/kg IBW), and the duration of asystole was 12, 9, 14, 18, and 44 s, respectively.

Low-dose adenosine (lower than 0.2 mg/kg) was used in six episodes of non IAR. Adenosine applied at 3 mg did not induce asystole, whereas four episodes treated with 6 mg of adenosine (0.11–0.13 mg/kg) and one episode treated with 12 mg (0.18 mg/kg) caused asystole for 8–12 s. Clip application was successfully performed, and additional doses of adenosine were not required. Low-dose AiTA was used in one episode to control bleeding during IAR; in this case, adenosine (9 mg, 0.2 mg/kg) induced 12 s of asystole.

Scattergrams of the adenosine dose (mg/kg IBW) and the duration of asystole (s) are shown in [Figure 1]. We found that there was a relationship between the adenosine dose and the duration of asystole ($r = 0.74$), and a linear regression analysis of each adenosine dose showed a significant relationship with the duration of asystole ($R^2 = 0.55$).

Postoperatively, two patients had elevated cardiac troponin T levels, but none of their EKGs showed any changes. Atrial fibrillation (AF) occurred after adenosine dosing in one patient who had preoperative PACs. Amiodarone was initiated due to the degree of hypotension in that patient. However, the patient’s postoperative cardiac enzyme level was normal, and the AF changed to a sinus rhythm in the intensive care unit. In the other patients, the degree of hypotension or bradycardia resulting from AiTA was transient and resolved to normal baseline values.

**DISCUSSION**

During intracranial aneurysm surgery, AiTA was initially used in cases where temporary clipping was infeasible and to facilitate bleeding control if IAR occurred.\(^{[8,10,14,17]}\)
recently, AiTA has been used as an alternative to temporary clipping in patients without IAR. [2,5,9,11,13,15] There are no specific indications for adenosine use, but several authors have provided expert opinion on its use, such as in a large or deep-seated aneurysms in narrow corridors where temporary clip ligation is difficult or impossible, synergy with temporary clipping, and to improve visualization of adjacent perforating arteries. [7]

Many studies have demonstrated the safety of this technique with regard to perioperative cardiac morbidity. [2,5,11,12,20] However, no study has reported a difference in the dose recommendation between the use of AiTA for IAR versus without IAR, and few studies have evaluated the optimal dose of adenosine.

At present, there are two techniques for adenosine administration in AiTA, with each requiring different dosages of adenosine: the dose-escalation technique and the dose estimation technique. In the dose-escalation technique, 6–12 mg of adenosine is initially used, and then more doses are added in a titrated manner. In the dose estimation technique, a single dose of adenosine is administered to achieve a predictable asystolic time. In dose-response studies, dosages ranging from 0.24 to 0.42 mg/kg IBW were recommended to achieve 30–60 s of profound hypotension and bradycardia. [2,8] Meling et al. [15] reported the use of a relatively higher single dose of adenosine during surgical aneurysm clipping (range 18–45 mg); nevertheless, the duration of asystole was similar to that in our study (5–15 s). Our institution prefers the dose estimation technique, but we use a relatively lower dose of adenosine than has been reported in other studies. We have found that dose-response characteristics show both intra- and interindividual variability.

In cases of IAR, it is useful to maintain asystole until temporary clips can be placed on the parent vessels or permanent clips can be placed on the aneurysm’s neck. Thus, using AiTA might require a long asystolic duration. The findings of this study support the dosages of adenosine reported in previous studies. [2,6] An adenosine dose of 0.2–0.4 mg/kg can induce asystole for up to 44 s, which is sufficient for bleeding control, without requiring an additional dose. Since the efficacy of repeated adenosine doses on asystolic duration is unpredictable, a higher estimated initial dose is most appropriate. In our study, we used a relatively lower dose than has been reported in other studies but still achieved similar asystolic times. The observed difference in dose-response times might be partly due to the race or underlying diseases of the patients, the site of injection (central versus peripheral), and interactions with other drugs. [1,4,6,22,24] The use of low-dose AiTA during IAR could induce brief asystole that is adequate for proximal control and minimizing blood loss.

The purpose of using AiTA in non IAR cases is to prevent IAR during permanent clip application and minimize blood loss if IAR occurs. Intarakhao et al. [11] found that using AiTA as an alternative to temporary clipping could minimize the temporary clipping time without increasing the risk of IAR. Our findings demonstrate that low-dose adenosine (lower than 0.2 mg/kg) caused 8–12 s of asystole, which can adequately facilitate the placement of a permanent clip. A patient who received 3 mg of adenosine had brief hypotension, but the clipping was successful, and no IAR occurred. Thus, low-dose AiTA can be considered as an option for non IAR cases because profound systemic hypotension itself without asystole could facilitate safe clipping as well. Guinn et al. reviewed the use of adenosine in patients who had electively clipped aneurysms. Although the minimum total dose of adenosine was 3 mg, the patients in this case series experienced some degree of asystole or bradycardia. The aneurysm decompression was satisfactory, and the clipping procedure was successful. [9] These findings support the notion that both transient hypotension and relative asystole are important factors that facilitate the application of the clip to softened aneurysms. Thus, we propose that low-dose AiTA can be used during permanent clip application without IAR. This method could prevent IAR and minimize the cardiac complications that might occur as a result of prolonged asystole.

In our study, one patient who had preoperative PACs developed AF after adenosine dosing. Pre-existing cardiac conduction abnormalities might have been the cause of this patient’s cardiac arrhythmia, and the cardiac asystole was also longer in this patient than that observed in other patients. Therefore, clinicians should be aware of whether the patient has any conditions including severe reactive airway disease, severe coronary artery disease, and pre-existing cardiac connection abnormalities [7] that could be relative contraindications for adenosine before administering adenosine, and a small initial dose might be required.

The previous studies [2,5,9,11,13,18] have also reported relatively low incidences of transient cardiac arrhythmia and AF that resolved spontaneously and not related to the dose of adenosine. Postoperatively, elevated troponin T levels could also occur; however, it is unclear whether this finding is related to adenosine. Transthoracic echocardiography may be useful for evaluating impaired myocardial contractility or regional wall motion abnormalities.

Regarding the study limitations, the study design was retrospective, and it included a relatively small number of patients. In addition, the duration of asystole was recorded manually. The details about the duration of hypotension and precise heart rate after the bolus of adenosine are lacking, which might have been beneficial in our study. Finally,
each surgeon needs a different working time, so the request for a repeat dose of adenosine might be dependent on the individual.

CONCLUSION
AiTA can facilitate the clipping of intracranial aneurysms and has a low incidence of serious complications. When using AiTA during IAR, we suggest using an estimated dose of 0.2–0.4 mg/kg to provide long asystole to facilitate bleeding control and that additional doses of adenosine should be minimized. Under non IAR conditions, a low dose of adenosine (6 mg or 0.1–0.2 mg/kg) can be considered as an option to facilitate clip application. However, further studies with larger numbers of patients are needed to refine our understanding of the use of adenosine during aneurysm surgery.

Declaration of patient consent
Patient's consent not required as patients identity is not disclosed or compromised.

Financial support and sponsorship
Nil.

Conflicts of interest
There are no conflicts of interest.

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**How to cite this article:** Intarakhao P, Thirawat P, Tewarittrueangsri A, Pojasupawun S. Low-dose adenosine-induced transient asystole during intracranial aneurysm surgery. Surg Neurol Int 2020;11:235.