Case Report

Anesthetic Overdose Leading to Cardiac Arrest Diagnosed by End-Tidal Inhalant Concentration Analysis in a Dog

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A 5-year-old male-castrated Cocker Spaniel presented to the Veterinary Teaching Hospital of the University of Georgia for a total ear canal ablation. Premedication was with carprofen 2.2mg/kg SQ, hydromorphone 0.1mg/kg IM, diazepam 0.2mg/kg IM, and glycopyrrolate 0.01mg/kg IM. The patient was induced with lidocaine 2mg/kg IV and etomidate 1mg/kg IV and maintained with sevoflurane and a constant rate infusion consisting of lidocaine 0.05mg/kg/min. Before surgery start, the patient's systolic arterial blood pressure was 110mmHg, heart rate (HR) was 85 beats/min, respiratory rate was 8 breaths/min, end-tidal sevoflurane concentration was 3.2%, and end-tidal CO₂ (ETCO₂) was 23mmHg. As a scrub was being performed, the patient's HR abruptly dropped to 20 beats/min over the course of 2 minutes. His ETCO₂ simultaneously decreased to 16mmHg. At this time, cardiopulmonary arrest was diagnosed. After two minutes of resuscitation, a spontaneous heart beat was obtained and the patient was successfully recovered and discharged without further incident. The cardiac arrest in this case is most likely attributable to an overdose of inhalant anesthesia, which was diagnosed by an anesthetic inhalant concentration monitor. A gas analyzer may be a helpful contribution to the small animal practitioner, particularly those performing more lengthy or complex procedures.

1. Introduction

Anesthesia in pet dogs can be associated with intraoperative fatality due to the anesthesia. In people, the incidence of anesthetic-related arrest is approximately 1 in 40,000 patients [1]. In small animals, this value has been estimated at 2 in 1,000 [2], which represents a significantly greater risk than in people. In dogs undergoing anesthesia, successful return of spontaneous circulation occurs in approximately 50% of patients, indicating an anesthetic-related mortality rate of 1 in 1,000 [2]. The incidence of anesthetic arrest in small animals in general practice is unknown, but the incidence of perioperative fatality in dogs has been estimated at 2 in 100, which reflects a marked increase compared to a university setting [2, 3]. It is possible that this difference is due to increased expertise at an institution, increased vigilance, or greater monitoring technology and interventional capabilities.

A gas analyzer is a device that is equipped to measure end-tidal concentrations of various gasses and vapors, notably CO₂ and halogenated inhalant anesthetics [4]. A number of technologies can be used to determine this information, with the most simple being infra-red, whereby infra-red light of a specific wavelength is transmitted into a sample, and the amount of light sent out versus the amount of light received by a sensor indicates the concentration of the substance. The American Society of Anesthesiologists requires end-tidal CO₂ (ETCO₂) monitoring for people under anesthesia [5], but this recommendation has not been adopted by the American College of Veterinary Anesthesia. ETCO₂ monitoring has been well correlated with outcome during cardiopulmonary resuscitation in small animals [2]. Monitoring end-tidal inhalant anesthetics has been speculated as one possible explanation for an improvement in outcome with anesthetic arrests at a university [2, 6].

2. Case Presentation

A 5-year-old male-castrated Cocker Spaniel presented to the Veterinary Teaching Hospital of the University of Georgia for a total ear canal ablation (TECA) due to chronic unresolved otitis externa and media. Six months earlier, the same procedure had been performed on the left ear without complication.
On physical exam, the dog was bright, alert, and responsive and had a grade II/VI systolic heart murmur, severe otitis externa, and a normal temperature, pulse, and respiration. There were no significant abnormalities in the history except for the chronic otitis. Bloodwork, including a complete blood count, biochemical profile, and urinalysis revealed a mild neutrophilia with regenerative left shift and a 3+ proteinuria. The owners declined an echocardiogram.

The patient was premedicated with carprofen 2.2 mg/kg SQ, hydromorphone 0.1 mg/kg IM, diazepam 0.2 mg/kg IM, and glycopyrrolate 0.01 mg/kg IM. This was the same protocol he had received for his first TECA procedure 6 months before. After 30 minutes, an IV catheter was placed and the patient was induced with lidocaine 2 mg/kg IV and etomidate 1 mg/kg IV. Etomidate was used to minimize any cardiovascular depression associated with induction, in deference to the heart murmur which had not been fully worked up. The patient was maintained with sevoflurane in oxygen and a constant rate infusion (CRI) consisting of lidocaine 0.05 mg/kg/min. This was used to provide analgesia and allow for a lower concentration of inhalant anesthesia. The patient was instrumented with indirect blood pressure via doppler, an ECG, a pulse oximeter, and a capnograph/gas analyzer. Throughout the surgical prep, all vitals were stable.

After preparing the patient for surgery, he was moved to the operating room, and intermittent positive-pressure ventilation (IPPV) provided by a mechanical ventilator was begun to deliver 12 breaths a minute with a peak inspiratory pressure of 14 cm H2O. At that time, his systolic arterial blood pressure (SAP) was 110 mmHg (normal >90 mmHg), heart rate (HR) was 85 beats/min, respiratory rate (RR) was 8 breaths/min, temperature was 98.2°F, end-tidal sevoflurane concentration was 3.2%, SpO2 was 99%, and ETCO2 was 23 mmHg (normal: 35–45 mmHg). As a surgical scrub was being performed, the patient’s HR abruptly dropped to 20 beats/min over the course of 2 minutes. His ETCO2 simultaneously abruptly decreased to 16 mmHg.

At this time, cardiopulmonary arrest (CPA) was diagnosed. The patient’s sevoflurane and lidocaine were discontinued, and he was given atropine 0.05 mg/kg IV and epinephrine 0.01 mg/kg IV, and external chest compressions were initiated. After two minutes of resuscitation, a spontaneous heart beat was obtained. The patient was moved out of the operating room and given mannitol 0.5 g/kg IV over 10 minutes to treat suspected cerebral edema from ischemia during the CPA. Fifty minutes after the CPA event, the patient was extubated. Over the next two hours, the patient regained his palpable reflex and then his menace reflex. Surgery was rescheduled for two weeks later, at which time no significant complications were encountered. An echocardiogram was performed prior to this later procedure and mild mitral regurgitation was diagnosed.

### 3. Discussion

The cardiac arrest in this case is most likely attributable to an overdose of inhalant anesthesia. Although the patient did have some underlying cardiac disease, the arterial blood pressure and heart rate were stable throughout surgical prep. No arrhythmias consistent with myocardial disease or ischemia were detected. Also, the patient responded rapidly to treatment with external chest compressions, atropine, and epinephrine. If the cardiac arrest was due to cardiac failure, it would be expected that the patient would have been less responsive to the cardiac resuscitation measures. Furthermore, the patient underwent anesthesia two weeks later without incident and had an echocardiogram which was unremarkable at that time.

Other possible causes include excessive vagal tone, a machine fault (such as a closed pop-off valve), or overdose of a different drug. There was no event surrounding the CPA which may precipitate vagal hypertonus, such as the manipulation of the eye or viscer. No machine fault was detected at the time, nor afterwards when the machine was evaluated thoroughly. No other drugs were being given with the exception of lidocaine. Lidocaine is a negative inotrope and can depress the cardiac output at toxic doses. However, the dose of lidocaine was checked and the syringe pump administering the CRI was also confirmed to be administering a proper concentration.

Anesthetics have a fairly narrow therapeutic index (TI), which is the LD50/ED50. Sevoflurane has a TI of 2.6 in rats [7] and propofol has a TI of 3.1 in rats [8]. Therefore, overdose with inhalant agents is relatively easy. The ED50 of inhalant anesthetics is the minimum alveolar concentration (MAC), the point at which 50% of patients will not move in response to a standard painful stimulus. The MAC for sevoflurane is approximately 2.1% [9]. For a surgical plane of anesthesia, typically 1.2–1.5 times the MAC is necessary (i.e., 2.5–3.2% for sevoflurane) [10]. However, this requirement will be decreased by other anesthetic and analgesic drugs. In this patient, the time of CPA was 2 hours after premedication, so it is unlikely that the premedicants were still exerting a significant MAC-sparing effect. However, the patient was concurrently receiving lidocaine.

Lidocaine is a fast sodium channel blocker which has potent analgesia and sedative properties in dogs. It profoundly decreases the MAC requirement in dogs, by up to 23% at the dose given to this patient [11]. Therefore, it would be expected that the patient could have been maintained at an end-tidal sevoflurane concentration between 1.9 and 2.4%. An end-tidal concentration of 3.2%, which was the concentration at the time of CPA, represents a 33% higher concentration than the highest which would be necessary to keep the patient adequately anesthetized for surgery. At the time of CPA there was no surgical stimulation. Typically, surgical stimulation partially counters the cardiodepressant effects of the inhalant anesthetics. Thus, this patient was at a higher concentration of sevoflurane than that required for surgery, was not actively receiving surgical stimulation, and was on a lidocaine CRI which may have been blunting any cardiovascular stimulus he had. This particular synergy of events likely led to the CPA event.

This case was unusual because the cause of CPA was very likely inhalant anesthesia overdose, and the concentration of inhalant at the time of CPA was known. Although end-tidal gas analyzers are expensive (ranging in price from $2,800 to $15,000), they are useful for documenting the
concentration of inhalant in the patient which helps guide clinical decision making and is the only way to definitively diagnose inhalant anesthesia overdose or underdose. There has been speculation that the presence of a gas analyzer dramatically improves the ability to manage equine cases [6], and this may also be true in small animals. A gas analyzer may therefore be a helpful contribution to the small animal practitioner, particularly those performing more lengthy or complex procedures. This case also highlights the potential risk with administering drugs as a CRI which significantly decreases the MAC, such as lidocaine, morphine, and/or ketamine [12]. Although balanced anesthesia with a variety of drugs each acting to reduce the dose of any one drug is generally preferable, in inexperienced hands these techniques may be dangerous. The clinician should always be thoroughly comfortable with the anesthesia technique employed and understand the effects of drugs on the inhalant anesthetic requirement.

References
