Bispectral Index Guided Propofol and Remifentanil Anaesthesia: Evaluation of Drug Consumption and Immediate Recovery in a Ghanaian Population

Akwasi Antwi-Kusi1,*, Bright Ighodaro Obasuyi2

1Department of Anaesthesia and Intensive Care, Komfo Anokye Teaching Hospital, Kumasi, Ghana
2Department of Anaesthesiology, University of Port Harcourt Teaching Hospital, Port Harcourt, Nigeria

Email address:
antwikusi@yahoo.com (A. Antwi-Kusi)
*Corresponding author

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Abstract: General anaesthesia is a balance between the amount of anaesthetic drug(s) administered and the state of arousal of the patient. Different patients may need variable doses of anaesthetics to reach and maintain a surgical level of anaesthesia. Even though lack of autonomic response suggests sufficient analgesia during surgery, there is no such clear clinical sign for the assessment of the depth and sedation of anaesthesia especially in a paralysed patient. Bispectra index (BIS) monitor improves anaesthetic delivery by allowing clinicians to administer the minimal dose of agents required to achieve an adequate hypnotic state. In this study conducted at the Komfo Anokye Teaching Hospital, patients undergoing general anaesthesia received intravenous propofol and remifentanil under BIS monitor as the sole anaesthetic. At the end of the surgery the total amount of propofol and remifentanil consumed and the time it took for the patients to open the eye and respond to verbal commands were noted. Propofol consumption was calculated to beat an average rate of 5.44mg/kg/h and remifentanil at 13.68ug/kg/h to maintain a surgical level of anaesthesia. The mean time from stopping the anaesthetic agents until the patients opened the eyes and responded to verbal commands were 2.20 minutes and 2.70 minutes respectively. The study found that Propofol and remifentanil infusions at a rate of 5.44mg/kg/h and 13.68ug/kg/h respectively, under BIS monitor, effectively controls intraoperative responses and allow for rapid emergence from anaesthesia in a Ghanaian population.

Keywords: Total Intravenous Anaesthesia, Bispectra Index, Remifentanil, Propofol

1. Introduction

General anaesthesia is a balance between the amount of anaesthetic drug(s) administered and the state of arousal of the patient [1]. Different patients may need variable doses of anaesthetics to reach and maintain a surgical level of anaesthesia [2]. It has been established that patients vary widely in their responses to drugs. Genetic factors can account for 20% to 95% of patient variability [3]. Genetic polymorphisms for many drug- metabolising enzymes and drug targets [eg. Receptors] have been identified. There are also differences in drug administration, metabolism and excretion among races [2] and these differences are of importance to the anaesthesiologist.

It is therefore necessary to monitor the effect of anesthetic drugs on patients based on individual patient variations. Even though lack of autonomic response suggests sufficient analgesia during surgery, there is no such clear clinical sign available for the assessment of the depth and sedation of anaesthesia especially in a paralysed patient. Standard clinical practice is cautious and anaesthetists may err on the side of safety by administering larger than adequate drug doses in the hope of avoiding awareness [3]. This may lead to the administration of high doses of anaesthetic agents which will delay emergence and affect recovery from anaesthesia. Bispectra index (BIS) monitor improves anaesthetic delivery by allowing clinicians to administer the minimal dose of agents required to achieve an adequate hypnotic state.
It is one of several technologies used to monitor the depth of anaesthesia. Titrating the anaesthetic agents to specific bispectra index during general anaesthesia in adults and children over 1 year old allows the anaesthetist to adjust the amount of anaesthetic agent to the needs of the patient, possibly resulting in more rapid emergence from anaesthesia.

The non-invasive sensor has self-adhesive backing, much like a typical EEG pad. After wiping the patient's forehead and temple with alcohol and drying the skin to ensure that a quality signal will be obtained, you'll place the sensor across the patient's forehead and over either the left or right temple. The sensor then sends raw EEG information through the cable and converter to the BIS engine. This engine processes the EEG data according to an algorithm that combines select EEG features to produce a BIS index. This index is a number between 0 and 100 that is displayed on the monitor and reflects the patient's level of sedation. A BIS value between 40 and 60 indicates an appropriate level for general anaesthesia as recommended by the manufacturer. The essence of BIS is to process a complex signal (the EEG), analyse it and process the result into a single number [4, 5, 6]

Total intravenous anaesthesia (TIVA) involves the use of intravenous agents for induction and maintenance of anaesthesia. The advantage of TIVA under BIS guidance include rapid emergence from anaesthesia and better recovery characteristics. It has however been reported by Ortolani et al. that recovery of Senegalese African Blacks from Intravenous Anaesthesia with Propofol and Remifentanil is slower than that of Caucasians. Our experience on recovery characteristics in Ghanaians after TIVA was however different from that reported by Ortolani [7]. Our hypothesis was that West Africans require a lower dose of propofol and remifentanil for total intravenous anaesthesia and that if the normal standard dose is given to Ghanaians and for that matter Africans, it will lead to delayed awakening. There has not been any study on total intravenous anaesthesia using propofol and remifentanil under BIS guidance in Ghana. Our study therefore sought to find the consumption of propofol and remifentanil used for TIVA under BIS guidance among Ghanaians and the characteristics of their recovery.

2. Methodology

After institutional ethics approval and patients consent, male and female ASA physical status class I and II who were coming for various elective surgeries under general anaesthesia during the study period were recruited for this research (convenient sampling). The number of patients recruited was restricted to the available research materials like BIS electrodes and the medications available. Exclusion criteria included patients with neurologic disease, patients taking medications that affect the central nervous system, age below 18 or above 70 years, body mass index (BMI) > 35 and history of previous adverse reaction to any of the drugs to be used for the research. All the patients were fasted over night prior to the surgery. No premedication was given. Patients taking medications like anti-hypertensives that would not affect the anaesthesia were allowed to take their medications on the morning of surgery. An 18G intravenous cannula was inserted into a forearm vein and 500ml of lactated ringers solution was set up. This was followed by base line monitoring which included oxygen saturation, heart rate, electrocardiogram (ECG), non invasive blood pressure and the bispectral index of the electroencephalogram (BIS A-200 monitor), Aspect Medical System Inc.

Pre-oxygenation was done with oxygen flowing at 5L/min for three (3) minutes. Fentanyl 2µg/kg was given for preemptive analgesia. Induction of anaesthesia was done with Propofol 2mg/kg and titrated till loss of verbal response and eyelash reflex. After the patient was induced to sleep, test ventilation was done and when mask ventilation was satisfactory, the patient was given Rocuronium 0.5mg/kg to facilitate tracheal intubation and to maintain neuromuscular block for the surgery. The patient was tracheally intubated and the cuff of the endotracheal tube inflated. The patient was ventilated with oxygen and air with F\textsubscript{2}O\textsubscript{2} of 0.4%. Anaesthesia was then maintained with Propofol (Diprivan) 2% by Astra Zeneca at an initial rate of 6mg/kg/h and Remifentanil (Ultiva) by Glaxo Smithkline at 5µg/kg body weight through the use of a syringe pump.

Mean arterial pressure (MAP), Heart rate (HR) Blood pressure (BP) and bispectra index (BIS) were used to evaluate the depth of anaesthesia. Throughout surgery the propofol infusion was adjusted to maintain a BIS value of between 40 and 60. Remifentanil rate was also adjusted according to the autonomic response: at a pulse rate and non invasive systolic blood pressure greater than 20% of baseline value respectively, the dose of remifentanil was increased to abolish the autonomic response. All patients were also given 1.25g novalgin (Novamin sulphate) infused over 5 minutes at 30 minutes before the end of surgery for post operative analgesia. Approximately fifteen minutes before the end of surgery, the propofol infusion was stopped and the Remifentanil infusion adjusted according to the patient’s autonomic response. After the last stitch the remifentanil infusion was also stopped. The skin incision was infiltrated with 20ml of 0.5% ropivacaine for post operative analgesia. A nerve stimulator was used to assess neuromuscular function. Patients were extubated when neuromuscular function was adequate and the patient could open the eyes and obey command. No reversal agent was given for neuromuscular block. At the end of the surgery, the total amount of propofol and remifentanil consumed as well as the time taken for the patient to open the eyes spontaneously and respond to verbal command were noted. Any complications noted intra and post operatively were also noted and documented.

3. Results

30 patients were enrolled in this study with a mean age of 40.67 and a standard deviation (SD) of 12.01. There were 18 males and 12 females. The mean weight of the patients was
The mean duration of the surgeries was 69.30 minutes with a SD of 19.25 and the mean duration of anaesthesia was 79.57 minutes with a SD of 24.38. Table 1 is a summary of the recovery characteristics and amount of drugs consumed. The average amount of propofol consumed was 413.67 mg and that of remifentanil was 1.04 mg. The average consumption rate of propofol was calculated to be 5.44 mg/kg/h and that of remifentanil was calculated to be 13.68 ug/kg/h. This is lower than the dosage described in standard text books. The time from the end of anaesthesia to eye opening was 2.20 minutes and the time from end of anaesthesia to verbal response was 3.70 minutes. The patients spent an average of 37.7 minutes in the recovery ward.

Table 1. Recovery characteristics of patients and amount of drugs consumed.

|                          | Mean     | S.D     | Number |
|--------------------------|----------|---------|--------|
| End of anaesthesia to eye opening | 2.20 minutes | 0.71    | 30     |
| End of anaesthesia to verbal response | 3.70 minutes | 1.44    | 30     |
| Amount of propofol consumed | 413.67mg | 120.19  | 30     |
| Amount of remifentanil consumed | 1.04mg   | 0.36    | 30     |
| Infusion rate of propofol | 5.44 mg/kg/h |        |        |
| Infusion rate of remifentanil | 13.68ug/kg/h |       |        |

S.D- Standard deviation

The main complication encountered was hypotension which was defined as systolic blood pressure falling below more than 20mmHg. When this happened, the infusion rate of the ringers lactate was increased and the anesthetic agents reduced. None of the patients reported any experience of intra-operative awareness (table 2).

Table 2. Complications.

| Complications                   | Incidence |
|---------------------------------|-----------|
| Bradycardia                     | 6.66%     |
| Post operative nausea and vomiting | 6.66%     |
| Hypotension                     | 13.32%    |
| Awareness                       | 0.00%     |

4. Discussion

Our results show that intravenous infusions of remifentanil at a rate of 13.68 ug/kg/h and propofol 5.44 mg/kg/h administered in a TIVA technique under BIS guidance is an effective regimen in Ghanaians, allowing for rapid recovery following anaesthesia. The dose of propofol administered in this study was smaller than the dose typically given for the maintenance of TIVA using propofol. Differences in sensitivity to anaesthetic agents among different races of people have been established [8, 9, 10]. There are also differences in sensitivity towards other medications. These differences may be due to both genetic factors and lifestyle. These lifestyles include nutrition, health, physical exercise etc. For example, Critchley et al found that West Africans show reduced metabolic activation of paracetamol [11]. Iron et al also demonstrated an increased sensitivity to alcohol in a Niger population possibly because of polymorphism of human alcohol dehydrogenase [12]. Due to these differences in sensitivity to drugs among races it is imperative to tailor your anaesthetic drugs to the needs of your patient. In reality this does not happen. Anaesthesia providers will always determine drug dosages as recommended by the manufacturer and standard textbooks. This is partly due to the unavailability of BIS monitor in developing countries to monitor the depth of anaesthesia. Because of this, there is always the fear of under dosing the patient with its attendant risk of awareness under anaesthesia. In order to avoid this problem anaesthesia providers tend to give standard doses of medications to their patients. This may lead to an overdose in certain group of patients due to genetic variations. In the case of total intravenous anaesthesia, this could lead to delayed awakening. A typical example is the research conducted by Ortolani et al, where a group of Black Senegalese Africans and Caucasians were given the same dosages of propofol and remifentanil in a total intravenous anaesthesia technique [7]. At the end of the surgery, they reported delayed awakening in the Black Senegalese patients even though the two groups received the same amount of medications. This delay is attributed to genetic polymorphism. The use of BIS monitor allows you to give the dose that is enough to put your patient to sleep, avoid awareness and also avoid overdose based on the individual patient [13, 14].

Our research did not show any differences in the awakening characteristics between males and females. There is abundant information in the literature on the concept of ‘personalized medicine’ in which a knowledge of genetic factors- guided prescribing, tailored to the individual, is popularly considered to be an inevitable consequence of completion of the International Human Genome Project [15, 16]. For example Lindi et al has established that race contribute to variability in dosing requirements for warfarin in anticoagulation with African Americans requiring higher doses and Asians requiring lower doses than whites. This difference was attributed to Whites having more of the variant enzyme CYP2C9 than in African Americans [15].
5. Conclusion

Propofol and remifentanil infusions at a rate 5.44mg/kg/h and 13.68ug/kg/h respectively under BIS monitor effectively controls intraoperative responses and allow for rapid emergence from anaesthesia in a Ghanaian population.

Limitations

The main limitation of this study is that it is a fairly small study based on patients recruited from one hospital. To enhance reliability of the results, the study size would need to be much higher.

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