Australian and New Zealand College of Anaesthetists (ANZCA)

Guidelines on Monitoring During Anaesthesia

Background Paper

PURPOSE

Monitoring is an integral part of observation and recording contributing to the management of anaesthesia and outcomes. In the perioperative period clinical changes can occur rapidly and at times unexpectedly, and consequently it is fundamental that such changes are detected early to facilitate management. In general, the term ‘monitoring’ is used to imply the use of electronic monitoring equipment; however other equipment such as a sphygmomanometer may meet the standard in some circumstances.

With advances in technology and monitoring equipment it is timely to review standards of monitoring. PS18 was last reviewed in 2008 and was republished in 2013. In line with A01 Policy for the Development and Review of Professional Documents, the title of PS18 has been changed from “Recommendations” to “Guidelines”. “Guidelines” offer advice on clinical and non-clinical aspects of the practice of anaesthesia and perioperative medicine, reflecting expert consensus and supported by other evidence when available.

SCOPE

PS18 is intended to apply wherever anaesthesia is administered and includes general anaesthesia, sedation, and major regional analgesia. While general anaesthesia and major regional analgesia is performed by anaesthetists, conscious sedation is administered by a wide group of practitioners. The guidelines are primarily intended for anaesthetists; however, any practitioner responsible for monitoring sedated patients should follow them.

The purpose of this guideline is to inform practitioners of the standards and to guide them in the use of monitoring aimed at achieving optimal clinical management and optimising patient safety and quality care.

BACKGROUND

There have been significant changes over time in the ability to observe and record, either manually or by computerised methods, physiological parameters, and this ability has led to advances in anaesthesia techniques. With the advances in anaesthesia and the development of more potent and titratable agents comes the benefit of producing very rapid onset of effects, but also the disadvantage of rapid onset of other (unwanted) effects. Increasing patient comorbidities and sometimes complex procedural requirements adds to the complexity of care.
Historically, rapid changes in physiology necessitated the anaesthetist to be in close physical contact to the patient, with constant observation, palpation of peripheral pulses, continual auscultation with precordial stethoscopes, and visualisation of pupils, amongst other things. Increasing sophistication of equipment has allowed the anaesthetist to be removed from direct physical contact and has provided the opportunity for improved quantification of parameters, as well as enabling alarm parameters to be set, and recording of data. This in turn contributes to improved quality care and outcomes.

Vigilance and situational awareness cannot be replaced by monitoring equipment and the purpose of the equipment is to confirm changes in clinical status, but also to signal changes earlier. Consequently, monitoring is essential to management of anaesthesia with specific regard to optimising outcomes.

The accompanying guidelines recommend that circulation, ventilation and oxygenation are monitored as a minimum and that other monitors should be added as required.

**ISSUES**

**Pulse Oximetry**
While monitoring of oxygen and use of pulse oximetry is regarded as mandatory for all cases the monitoring of other variables is strongly encouraged. Where, for any reason, other monitors are not used, they must be readily available should they become required.

**Electrocardiography**
It is recognised that ideally every patient should be monitored using an ECG and it is strongly encouraged; however, it is also acknowledged that there are situations where ECG monitoring may not be feasible. The omission of ECG monitoring in these circumstance must be clinically justifiable.

**Capnography**
Discussions as to whether capnography should be mandated for all cases indicated that there remains strong support for all cases in which general anaesthesia is administered. For sedation techniques that do not require airway instrumentation capnography is optional and should be determined by the patient’s clinical requirement and depth of sedation, and not the availability (or lack thereof) of suitable equipment. The transition between general anaesthesia and sedation and their definitions can be found in PS09 Guidelines on Sedation and/or Analgesia for Diagnostic and Interventional Medical, Dental or Surgical Procedures.

**Blood pressure**
Blood pressure monitoring is considered essential for the vast majority of cases. It is recognised however that in some situations this may be inappropriate (e.g. prior to induction of an extremely agitated patient), or impractical (emergency management of a life threatening airway condition, extremely brief paediatric procedure). In all such cases blood pressure monitoring should be initiated if and when circumstances permit.

**Neuromuscular blockade monitoring**
This is an emerging area of standardisation due to the awareness of the risk of residual curarisation. Guidelines such as the AAGBI (2015) provide excellent background and have moved towards mandating assessment of NMB; however the AAGBI Appendix does not consider suxamethonium and emergency cases if a NMB monitor is not available. A comprehensive review of non-depolarising neuromuscular blockade and reversal in 2017 concludes that “Objective measurement (a train-of-four ratio greater than 0.90) is the only
method to determine appropriate timing of tracheal extubation and ensure normal muscle function and patient safety\(^1\). Therefore, quantitative monitoring is recommended to assess depth of blockade prior to reversal and assessment of adequacy of reversal. Quantitative assessment is essential since tactile assessment is subjective and cannot detect fade at a TOF ratio of greater than 0.4. Only quantitative monitors can assess adequacy of reversal. Best available evidence suggests that a train-of-four ratio of >0.9 should be achieved prior to extubation following the use of non-depolarising neuromuscular blockade.

Feedback received during the pilot phase revealed that interpretations of the available evidence varied with strong proponents advocating for mandatory use of quantitative neuromuscular function monitors while others were more moderate. Both benefits and cost implications were considered. The recommendations in the accompanying guidelines acknowledge this debate and fall short of mandating their use. However, quantitative monitoring of neuromuscular functions is recommended with a view to improving the management of neuromuscular blockade.

**Other monitors**

The emergence of new monitoring modalities requires that their role and validity be carefully assessed prior to firm recommendations being made. Cerebral oximetry (using near infrared spectroscopy) is an example of such where although it is receiving widespread attention in a number of clinical applications, it is not yet clear that it will reliably alert to cerebral hypoperfusion in certain circumstances (e.g. beach chair or sitting positions). As further data emerges the role of these monitors will be clarified.

Temperature monitoring has historically been for the purpose of managing hypothermia. However, with the emergence of forced air warming and contact warming devices the potential for hyperthermia and its consequences has been recognised. As a result, whenever active warming devices are being used it is recommended that temperature be monitored to avoid hyperthermia. Specification of use for extended cases was avoided due to the many other factors that may be relevant other than purely time alone. Each case needs to be assessed on its merits and risks.

**Presence of the anaesthetist**

An anaesthetist must be constantly present whilst providing anaesthesia during the procedure. It is recognised however, that on occasions, exceptional circumstances may arise requiring the anaesthetist to leave theatre or procedure room for brief periods. Such absences should occur only if they are unavoidable, in which case a handover to another anaesthetist in accordance with PS53 Statement on the Handover Responsibilities of the Anaesthetist is required. If another anaesthetist is not available and the situation is within the scope of another suitably skilled practitioner, then delegation of observation, including recording of observations must be to a practitioner who is judged to be competent for the task and capable of responding to any significant perturbation either directly or by summoning assistance. The term ‘practitioner’ is defined by the regulatory authorities in Australia and New Zealand and in the context of this document the term refers to an individual who is practising within their scope of practice as determined by the health care institution in which the case is being undertaken.

**SUMMARY**

Monitoring is an integral part of observation and recording contributing to the management of anaesthesia and outcomes. PS18 was revised based on the above advice from the
document development group. The goal of this revised document is to support uniform standards for the use of monitoring in conjunction with clinical observation.

RELATED ANZCA DOCUMENTS

A01 Policy for the Development and Review of Professional Documents
PS02 Statement on Credentialing and Defining the Scope of Clinical Practice in Anaesthesia
PS03 Guidelines for the Management of Major Regional Analgesia
PS06 Recommendations on the Recording of an Episode of Anaesthesia Care
PS07 Recommendations for the Pre-Anesthesia Consultation
PS09 Guidelines on Sedation and/or Analgesia for Diagnostic and Interventional Medical, Dental or Surgical Procedures
PS53 Statement on the Handover Responsibilities of the Anaesthetist
PS55 Recommendations on Minimum Facilities for Safe Administration of Anaesthesia in Operating Suites and other Anaesthetising Locations
ANZCA Handbook for Training and Accreditation
ANZCA Handbook for Training and Accreditation in the Affiliated Training Regions

REFERENCES

1. Brull, SJ and Kopman, AF Current Status of Neuromuscular Reversal and Monitoring, Challenges and Opportunities. *Anesthesiology*. 2017; 126:00–00

FURTHER READING


DOCUMENT DEVELOPMENT

ANZCA Safety and Quality Committee acted as the document development group for the 2015 review, led by:
Professor David A Scott, FANZCA, Councilor
Dr Peter Roessler, FANZCA, Director of Professional Affairs (Professional Documents)

In addition, the following were consulted:
ANZCA regional and national committees
Faculty of Pain Medicine Board, national and regional committees
ANZCA Safety and Quality Committee
ANZCA Trainee Committee
ANZCA Special Interest Groups

Professional documents of the Australian and New Zealand College of Anaesthetists (ANZCA) are intended to apply wherever anaesthesia is administered and perioperative medicine practised within Australia and New Zealand. It is the responsibility of each practitioner to have express regard to the particular circumstances of each case, and the application of these ANZCA documents in each case. It is recognised that there may be exceptional situations (for example, some emergencies) in which the interests of patients override the requirement for compliance with some or all of these ANZCA documents. Each document is prepared in the context of the entire body of the College's professional documents, and should be interpreted in this way.

ANZCA professional documents are reviewed from time to time, and it is the responsibility of each practitioner to ensure that he or she has obtained the current version which is available from the College website (www.anzca.edu.au). The professional documents have been prepared having regard to the information available at the time of their preparation, and practitioners should therefore take into account any information that may have been published or has become available subsequently.

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