Postoperative Cognitive Dysfunction: An Updated Review

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Abstract

Post operative Cognitive Dysfunction is a condition that follows immediately after a major surgery or any other serious health ailment. It shows significant clinical and social impact on the patient that may often result in either death or impairment, disabling the individual to enter into the job market. While 30% to 40% of postoperative patients, particularly elders are affected immediately after getting discharged from the hospital, 5% to 12% of patients are affected after three months.

The origin and the reasons for POCD are not clearly known and there are no universally acceptable evidence based management strategies to cope with this emergency. There is no consensus among experts across the globe on how to treat these kinds of disorders and the definition for the Post Operative Cognitive Dysfunction is missing in the existing literature, leading to further confusion.

This article attains importance as tries to analyze the most recent theories of pathogenesis and the risks that may follow Post operative Cognitive Dysfunction by reviewing articles published on this subjects for the past 60 years, starting from 1955 to 2014 in PubMed and Google Scholar and suggests relevant measures on how to cope with it.

Introduction

Postoperative cognitive dysfunction (POCD) is an important condition that may follow surgery [1] or other procedures [2] and has significant clinical and social impacts. It is associated with increased risk of mortality, premature departure from the labor market and social dependency [3]. It affects a significant percentage of the postoperative patients, of all age groups [4,5], particularly the elderly [6], with an estimated incidence of 30-40% at hospital discharge and 5-12% at 3 months for noncardiac surgeries [7]. The pathogenesis is not fully understood and effective, evidence-based management strategies are not currently described. There are no universally accepted guidelines for the diagnosis and treatment of POCD, and there have been few consensus statements on the subject. For example, there was a consensus statement published in Annals of Thoracic Surgery in 1995 which highlighted the recommended battery of neuropsychological testing to assess for POCD [8], although this has not been adopted consistently in the subsequent literature. The definition of POCD is neither included in the Diagnostic and Statistical Manual (DSM-V) nor in International Classification of Diseases [ICD-10], and the clinical awareness of anesthesiologists is still incomplete. In this review, we shed light on the most recent theories of pathogenesis and risk factors, experimental and clinical treatments, and recommendations for clinicians based on the available data.

Methods

A comprehensive search was done using PubMed and Google Scholar. Articles were extracted over a period of 60 years from 1955 through November 2014. However, most are dated from 1998 onward. We also selected and examined appropriate references from the gathered articles. Key words included, but were not limited to the following: POCD review article, POCD in the elderly, middle aged, pediatric population, assessment of cognitive impairment/POCD, pathophysiology of POCD, management of POCD, anesthetics and POCD, anesthetics and Alzheimer’s disease, general versus regional anesthesia and POCD, POCD and depth of anesthesia, pharmacology and POCD, Isoflurane, sevoflurane, desflurane anesthesia and POCD; Ketamine, propofol, benzodiazepines and POCD, biomarkers and POCD, neuroinflammation and POCD, Total Intravenous Anesthesia [TIVA] and POCD, dexmedetomidine and POCD.

Awareness and assessment of the problem

As early as 1955 and 1957 postoperative neurocognitive changes were reported to occur after noncardiac and cardiac surgeries, respectively [9,10]. Investigational activity regarding this topic culminated with publication of large multicenter prospective randomized trial results by the International Study of PostOperative Cognitive Dysfunction group (ISPOCD), some of which included long term follow-ups [11]. Since then, interest in this topic has continued in the clinical scientific perioperative community as evidenced by a steady stream of insightful papers.

Currently there is no standardized definition of POCD; the closest term in ICD-10 is “mild cognitive impairment”. POCD is simply a deviation from normal cognition; however the definition of the extent of deviation varies among studies. Furthermore, the cognitive changes in POCD are subtle and must be confirmed by a battery of neuropsychological tests. Hence the diagnosis cannot be made on purely clinical grounds. The same domains that are assessed in dementia are also assessed in POCD. They include learning and
memory, language, executive function, complex attention, perceptual-motor function and social cognition [12].

The accurate assessment of POCD is difficult with a variety of contributing elements [13]. The most often mentioned are variability of examiners, different definitions of POCD, interval between test sessions, mood changes and anxiety, pain, sleep-deprivation, pharmacologic effects, failure of some patients to complete the studies, language and cultural problems, timing of postoperative assessment, differences in the tests used, surrounding environments, dissimilar exclusion criteria, and statistical design. Most well designed studies include a baseline assessment one week before; they follow up at one week and at three months after surgery; occasionally follow-ups as long as several years after surgery are reported. Large sample size, inclusion of non-surgical control groups, and using full set of neuropsychological tests are acknowledged as adding validity to study design. The Mini-Mental Status Examination (MMSE) is useful as a surrogate for clinical but not investigational purposes. Though it has sensitivity of 87% and specificity of 82% and is less time consuming than neuropsychological testing [14], it lacks sensitivity for detecting mild dementia [15].

Several investigator groups have searched for biomarkers of POCD for diagnostic or prognostic purposes. For example, Apolipoprotein E4 is a known marker for neurodegenerative diseases, but was not found useful for assessment of POCD [16]. In a pilot study, Price et al. evaluated whether preoperative MRI neuroimaging might predict POCD in non-demented patients. They found that reduced hippocampal/entorhinal volume predicted memory change; moreover, the degree of leukoaraiosis and lacunar volume predicted decrements in executive function [17]. Similarly, increased levels of IL-6 and S-100B correlated well with the incidence of POCD [18]. Other markers that have been studied include tau protein, malonaldehyde, melatonin level, neutrophil-lymphocyte ratio, and aspartic acid level, some with promising results [19-24]. However, further study is needed to prospectively validate how useful these biomarkers might be to identify high risk patients or those who might benefit from prophylactic or therapeutic intervention.

Pathophysiology, etiology and risk factors

Much of the available work suggests that causative factors of POCD could include the surgical experience [including the effect of hospitalization – recently subsumed under the term “post-hospital syndrome” [25] or potentially neurotoxic effects of anesthesia [26]. Therefore, in some studies, POCD is explained neither by surgery nor by anesthesia alone. Indeed, postoperative cognitive improvements relative to preoperative function has even been reported after surgery and anesthesia [27, 28]. This suggests that other factors are involved in the pathogenesis of POCD such as the patient’s own pathological state and subsequent systemic inflammatory conditions. For example, a Japanese prospective study [29], showed that preoperative existence of low grey matter volume and white matter lesions on MRI were associated with higher incidence of POCD after elective cardiac surgery. In a study using standardized assessment tools and good statistical design, Evered et al. compared the occurrence of POCD after CABG [on-pump: general anesthesia], coronary angiography [sedation only], and total hip arthroplasty [spinal and light general anesthesia]; the incidence of POCD after 3 months was 16%, 21% and 16% respectively. Surprisingly, the least invasive procedure performed only with sedation was associated with the highest incidence of POCD. This suggests that neither the intensity of surgical or procedural intervention nor the type of anesthesia alone can predict the occurrence of POCD [30].

It has been proposed that the surgical experience might induce POCD through surgery-induced systemic inflammation via activation of the immune system and release of proinflammatory cytokines [e.g. IL-1β, IL-6, TNF]. The latter are thought to violate the integrity of blood brain barrier and induce inflammation in the hippocampus, an area known to mediate memory and learning [31,32]. This response is even more exaggerated in the diseased brain [33-35]. General anesthetics could potentially lead to memory impairment through hyperphosphorylation of tau-protein, an important protein involved in the pathogenesis of Alzheimer’s disease. Tau hyperphosphorylation has been attributed to anesthesia-induced hypothermia [36,37]. However, a recent study showed that sevoflurane induced memory deficits even in normothermic mice [38]. Moreover, earlier studies attributed neuroinflammation to surgery only [39,40], but a more recent study has suggested anesthetic mediation as well. For example repeated exposures to anesthetics have more influence on learning, memory, and function than a single exposure [41]. Therefore, POCD or at least the components of POCD relating to memory function may also be generated through a direct effect of anesthetics on brain cells and circuits.

Furthermore, anesthetics may have a direct toxic effect on neural structures. For example, high concentrations of inhalational anesthetics can increase blood brain barrier permeability [42]. In a recent study [43], isoflurane disrupted hippocampal Neuregulin 1-ErbB4 in mice, which plays a key role in synaptic plasticity, and impairs learning and hippocampal long-term potentiation (LTP) [44]. Other studies in neonatal monkeys showed that prolonged ketamine infusion causes neurodegeneration and neuroapoptosis [45,46]; the same was found with isoflurane [47]. Dong et al. observed that sevoflurane induced apoptosis and elevated levels of B-site amyloid precursor protein-cleaving enzyme [48]. Taken together, these and many other findings lead to a consensus statement on the possible link between Alzheimer’s disease and anesthetics which states “there is sufficient evidence at multiple levels to warrant further and more definitive investigation of the onset and progression of Alzheimer’s disease and neurodegeneration after anesthesia and surgery” [49].

Currently, however, there remains no concrete evidence to support a direct causative relationship of anesthesia with regard to initiation or progression of Alzheimer’s disease. Interestingly, The ISPOCD group followed up patients from ISPOCD-1 and ISPOCD-2 for a median duration of 11 years without finding a significant relationship between anesthesia, POCD and dementia [50].

The prospect that general anesthesia (GA) may be neurotoxic led researchers to compare patients receiving GA with those having regional anesthesia [RA] or neuroaxial anesthesia. A review of seventeen clinical studies [51,52] failed to find a significant difference in the incidence of POCD after GA compared to RA. The main criticism of these studies is that RA is often combined with sedation, frequently deep sedation. Initially regional anesthesia without deep sedation [i.e. BIS>80 was shown to result in a substantially lower occurrence of delirium compared with [53]. Because delirium and POCD are linked, it was reasonable to assume a similar relationship for POCD. Therefore, Silbert et al. [2] prospectively compared the incidence of POCD after GA for Extracorporeal Shockwave Lithotripsy [ESWL] with the use of spinal anesthesia without sedation. Currently, however, there remains no concrete evidence to support a direct causative relationship of anesthesia with regard to initiation or progression of Alzheimer’s disease. Interestingly, The ISPOCD group followed up patients from ISPOCD-1 and ISPOCD-2 for a median duration of 11 years without finding a significant relationship between anesthesia, POCD and dementia [50].

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at 3 months the incidence was 19.6% and 6.8% for spinal and GA, respectively. The investigators could not conclude superiority of one technique over the other; clearly POCD could not be avoided through the use of RA without sedation. This and other findings after minimally invasive interventions point to a likely effect of hospitalization on patient functional status. Physician deconditioning has long been recognized as a consequence of hospital stays [54,55].

Depth of GA has been investigated as a contributing factor for POCD. Two studies suggested that deeper level of anesthesia monitored by Bispectral Index [BIS] to values between 30-40 result in better cognitive outcome than lighter values [i.e. 50-60] [56,57]. The main criticisms of this work are relatively small sample sizes and that assessment for POCD was done in the early postoperative period [4-6 weeks and 5 days, respectively]. In addition, only 3 neuropsychological tests were used for assessment in the first study instead of a full battery of tests. The CODA trial [Cognitive Dysfunction after Anesthesia] is a prospective randomized study of 921 patients. The patients were divided into a BIS-titrated group with a target of 40-60 versus BIS-blinded group. The BIS-titrated group had less delirium, less POCD and decreased anesthetic delivery than the BIS-blinded group, where BIS values were lower than in the BIS-guided group [58]. Another large prospective study found an association between low BIS values and higher incidence of delirium but not POCD [59]. More alarmingly, multiple studies have linked low BIS values with mortality [60,61]. A large retrospective study from Cleveland Clinic reported an association between a composite “triple low state” [low BIS, low blood pressure, and low Minimal Alveolar Concentration] and mortality [62]. On the other hand, investigators from Duke University could not confirm this association of mortality with the “triple low state” in a large retrospective study [63]. Although the “triple low state” studies include depth of anesthesia as one of the three components, it is unclear to what extent any one parameter contributed to the observed associations. Results from a pilot study from New Zealand and Australia, conducted in preparation for a larger clinical trial, indicate that “expected higher complication and mortality will be present with low BIS values” [64]. While the evidence for an association between depth of anesthesia and incidence of POCD is far from conclusive, currently available evidence seems to suggest that deeper levels of anesthesia may be associated with less than optimal outcome. Clinicians may well consider adjusting anesthetic dose to a level minimally capable of achieving the goals of anesthetic management, prioritized on surgical and patient factors. Interestingly, a recent survey of Australian anesthesiologists indicates that most anesthesiologists who utilize depth of anesthesia [DoA] electroencephalogram [EEG] monitors do so in an attempt to prevent awareness; the survey also suggests that practitioners may in the future look toward DoA monitoring as a means to improve recovery and minimize complications. Survey results showed that 62% of respondents thought that DoA monitors allowed them to use less anesthetic. This capability provided benefits that were thought to include “faster emergence, less morbidity, and improved hemodynamic stability” [65].

The type of GA has been examined as a contributing factor to POCD. Cai and co-workers conducted a large single center case-controlled study to investigate the association between Apolipoprotein E4 and POCD in an elderly population. They compared surgical patients receiving total intravenous anesthesia [TIVA] with others receiving general inhalational anesthesia [GIA]. They found that patients who received GIA had a significant decrease in postoperative MMSE scores on postoperative day 3 when compared to preoperative baseline; their postoperative MMSE scores were also lower than those of the TIVA group. In the GIA group, decreases in MMSE score were related to the presence of the Apo E4 allele. This association was not found in the TIVA group [66]; this finding suggests an interaction between type of anesthetic exposure and genetic predisposition in the development of POCD. In another study, type of anesthetic exposure, neuro-inflammation, and POCD were examined in patients specifically undergoing on-pump CABG. The TIVA group received propofol and fentanyl and the GIA group received sevoflurane and fentanyl. Neuro-inflammation was assessed with the serum biomarker protein S100B. The TIVA group had higher levels of S100B as well as lower MMSE and Montreal Cognitive Assessment [MoCa] scores when compared to the GIA group [67]. The differences in the results from these two studies are difficult to explain. Potential confounders include type of surgery, the patient population, or the individual risk factors of the patients [e.g. ApoE4 presence, etc.]. Furthermore, the investigator teams used different tools to detect POCD [MMSE alone and MMSE plus MoCA], and both evaluated their patients only early in the postoperative period. These studies highlight the difficulties that may arise from when standards for testing and timing of POCD evaluation are not adopted, as outlined in the consensus statement mentioned previously [8].

Old Age is considered a major risk factor for POCD and is universally commented on in the available literature. Strom et al summarized seven mechanisms to explain the higher incidence of POCD in older subjects [68]. This includes decreased brain volume, decreased density of the blood-brain barrier, decreased neurogenesis, decreased baseline cognition, decreased cognitive reserve, increased likelihood of inflammation, and cerebrovascular disease. Others risk factors for POCD include low level of education, preexisting cerebrovascular disease, noisy hospital environment, metabolic syndrome and systemic inflammatory states, duration of anesthesia, respiratory and infectious complications, male gender, genetic polymorphism, pain, longer hospital stay, more invasive complicated surgery, diabetes and hyperglycemia [11,52,58,69-74]. Hypoxia and hypotension are mechanisms that could induce cerebral ischemia. They were examined in ISPOCD1 study [11] and were not found to be significant risk factors for POCD. However, direct measurement of cerebral oxygen desaturation predicts POCD in cardiac [75] and noncardiac surgery and might be useful in selected cases [76,77]. For investigative work it is important to account for such risk factors in the development of one’s study design. Until effective preventative measures and therapeutic interventions have been identified, it is difficult to envision their use by practitioners in clinical decision making.

Showers of microemboli during cardiac surgery, especially during cardiopulmonary bypass, were thought to represent a mechanism of subtle ischemia and POCD [78]. MRI is a useful modality for detecting cerebral ischemia and microinfarcts caused by micro-emboli. Knipp et al. [79] prospectively examined 39 patients undergoing CABG by neuropsychological testing and MRI preoperatively and up to 3 years after the surgery. They found ischemic cerebral lesions in 51% of patients, but they failed to find an association between these lesions and POCD. Given the small population studied, further large scale studies are needed to confirm these findings and their clinical significance. To wit, studies using Transcranial Doppler [TCD] did not find a relationship between embolic load and occurrence of POCD [80,81]. Indeed, an association between the number of emboli and POCD was absent in both cardiac and orthopedic surgery [82,83] as well as coronary angiography procedures [84]. A meta-analysis by
Marasco et al. [85] failed to find a higher incidence of POCD after off-pump cardiac surgery [less emboli] versus on-pump cardiac surgery [more emboli], suggesting that microemboli do not represent a significant pathogenic factor.

Interestingly, a recent meta-analysis concluded that cardiac surgery results in postoperative cognitive improvement relative to pre-operative baseline testing. This suggests that improvement of myocardial ischemia may improve cognitive dysfunction [86]. Myocardial ischemia has emerged as an important potential risk factor for POCD. Zhu et al. [87] demonstrated that transient myocardial ischemia [induced by ligation of the left anterior descending artery for 30 minutes] caused in vivo cognitive dysfunction evidenced by impaired long term potentiation [LTP] and increased expression of inflammatory biomarkers. This effect has been attenuated by preconditioning with sevoflurane. LTP impairment did not occur after a sham procedure. Myocardial ischemia-reperfusion injury therefore likely plays a significant role in the occurrence of POCD after cardiac surgery.

**Therapeutic Interventions: Experimental and clinical trials**

Anti-inflammatory and neuroprotective therapies have been investigated for their potential to prevent POCD. In one animal study, neutralizing antibody to alarmin, a high-mobility group box 1 protein involved in neuro-inflammation, prevented the inflammatory response and decreased the incidence of memory deficits [88]. Similarly, resolvins are potent endogenous lipid mediators that are biosynthesized during the resolution phase of inflammation. Aspirin has been shown to trigger resolvin D1 production which in turn attenuated the effect of memory decline [89]. Minocycline mitigated isoflurane-induced cognitive impairment in aged rats [90]. Atorvastatin, an anti-inflammatory drug, attenuated in vivo reduction of the hippocampal-dependant fear response induced by surgery [91]. An animal study involving 344 rats suggests that amantadine attenuates learning and memory impairment after surgical intervention [92]. Dexmedetomidine attenuated isoflurane-induced neurocognitive impairment in developing neonatal rats [93].

In a recent clinical trial, magnesium failed to improve cognitive function after cardiac surgery [94]. Low dose bolus ketamine may attenuate POCD after cardiac surgery [95]. Two studies, one experimental and one clinical, demonstrated the role of intranasal insulin in maintenance of normoglycemia which lead to decreased tau hyperphosphorylation and improved cognitive outcome [96,97].

**Assessment of current anesthetics**

As a general rule, there is no clear superiority of any one the currently available anesthetics with respect to development of POCD. Bilotta et al reviewed 16 drugs [106], the incidence of POCD didn't differ between patients and controls in thiopental, ketamine, propofol and xenon, with conflicting results for lidocaine, ketamine and magnesium sulfate. Some studies suggest there may be a beneficial effect of desflurane over sevoflurane over isoflurane [107,108]. In a clinical study comparing spinal anesthesia plus desflurane, versus spinal anesthesia plus isoflurane versus spinal anesthesia alone, POCD incidence at one week was lowest in the desflurane group [109]. Repeated exposure of neonatal mice [3 times versus one time] didn’t induce neuroinflammation or cognitive dysfunction with desflurane or sevoflurane, respectively [110]. Benzodiazepines are known to affect concentration in elderly patients [111], however a clear relation between their serum concentration and POCD has not been proven [112].

Delirium is another condition that is often looked at in comparison to POCD. Dexmedetomidine, compared to midazolam, decreased delirium and facilitated early extubation in ICU mechanically ventilated patients [113]. Despite this finding, delirium and POCD are separate entities. A relationship has yet to be fully elucidated and properly studied, and it is still unclear if strategies that reduce delirium can also reduce POCD [114]. One study reviewed examined delirium in mechanically ventilated ICU patients as a predictor for long term cognitive impairment. The majority of patients’ evaluated were critically ill with diagnoses of sepsis and ARDS. The patients were evaluated with 9 neuropsychiatric tests administered by a neuropsychiatrist at 3 months and 12 months. They found that delirium was in fact an independent predictor of long term cognitive impairment and also noted that an increase in the duration of delirium also increased long term cognitive impairment [115]. This study suggests a relationship between delirium and cognitive impairment, but this is also another example of investigators using different definitions/evaluation tools for cognitive impairment. In another study that evaluated patients after cardiac surgery, the relationship between delirium and long term cognitive and functional outcomes was examined. In this study, patients diagnosed with delirium based off Diagnostic and Statistical Manual for Mental Disorders [DSM-IV] were given a questionnaire 1-1.5 years after having cardiac surgery. Sleep disturbance was the only category evaluated noted to have a statistically significant difference between patient’s with post operative delirium and without delirium. Mortality, readmission rates, memory/concentration, and dependency for activities of daily living [ADLs] all had a negative tendency in the delirium groups, but there was no statistically significant difference. Of note, this study was limited by a small sample size and the use of a mailed questionnaire [116]. The results of this study suggest a need for further study in this area to truly determine if a relationship between delirium and cognitive impairment exists. A Taiwanese study evaluated similar data in orthopedic surgery patients. They utilized a Comprehensive Geriatric Assessment pre-operatively and 24 hours post-operatively. This assessment looked at visual/hearing deficits, polypharmacy, depression, nutrition, comorbidity, pain, cognitive function, and ADLs. After discharge at 1, 3, 6, and 12 months, the patients received a phone call to assess their ability to perform ADLs. This study showed worsening of scores regarding independence with ADLs at 6 and 12 months in patients with delirium, as well as even worse scores in patients with delirium and cognitive impairment [117]. This study calls for early identification of cognitive impairment and the...
implementation of strategies to prevent delirium. Another study looked at neuroimaging in relation to both delirium and cognitive impairment. Patients that were diagnosed with delirium in the hospital were evaluated at 3 and 12 months post discharge. The investigators evaluated 5 indices: immediate memory, visuospatial, language, attention, and delayed memory. The patients also underwent an MRI scan at discharge and 3 months later. Specific areas of the brain were examined for fractional anisotropy which is a calculation done with diffusion tensor imaging to evaluate axonal integrity. The study showed that longer duration of delirium was associated with white matter disruption on imaging, and white matter disruption was associated with worsened cognitive outcomes at 12 months [118]. These studies all suggest that further study into the relationship between POCD and delirium needs to be investigated further. Unfortunately, these studies also highlight the need for standardization of evaluation tools for cognitive impairment if more progress into understanding POCD is hoped to be achieved.

**Recommendations**

Despite significant and copious research on the subject, Post-operative Cognitive Dysfunction remains poorly defined and poorly understood. While many potential explanations are suggested, a definitive pathophysiology has not been described, and a direct causal relationship has not been firmly established between the disease and any suggested insult. Furthermore, no definitive peri-operative or, more specifically, anesthetic strategy has been shown to definitively improve the incidence or severity of POCD.

Despite this, there are several intriguing possibilities both for future research and for application of studied practices to attenuate the problem. We strongly advocate pre-operative discussion with patients and their families regarding the possibility of post-operative cognitive dysfunction in at-risk groups.

Pre-operative discussion with the patient’s family is highly recommended to discuss the potential for POCD to occur. It is important to stress that there are various causes of this phenomenon, including the surgery itself, and also that the effect is temporary. This is especially important in elderly patients and/or patients with baseline dementia or cognitive deficits. A MMSE can be considered before and after surgery for patients that are at risk for POCD [elderly, patients with baseline dementia or cognitive deficits, etc.] in order to establish a baseline and monitor for the occurrence of POCD, respectively. The routine use of midazolam or other benzodiazepines except in populations that are felt to be at risk should be avoided unless strongly indicated. Meperidine should also be avoided. While it is unclear whether it definitely improves outcome, minimally invasive surgery or laparoscopic surgery should be advocated to decrease the extent of inflammatory response and subsequently reduce the chances of developing POCD.

Hypothermia in conjunction with inhalational anesthetics enhances tau phosphorylation in animal studies which leads to increased memory deficits. Hyperthermia is also recognized as harmful in cardiac and brain surgeries. Hence, maintaining normothermia is strongly advocated. Hyperglycemia appears to be associated with POCD, and worsens outcome in neurosurgery and cardiac surgery. Normoglycemia should be maintained.

There is conflicting evidence regarding depth of anesthesia and post-operative cognitive dysfunction. Some evidence suggests the potential for higher rates of POCD with deeper anesthesia, while more recent evidence seems to suggest that anesthesia titrated to BIS-scores between 40-60 improves the likelihood of POCD. It is recommended to avoid the use of anesthetics for non-anesthetic purposes, e.g., treating hypertension by increasing the dose of inhaled anesthetic. It is important to pay attention to anesthetic requirements on a patient by patient basis. DoA monitors may be used in the future to help providers use less anesthetics per case and therefore improve outcomes. Some studies suggest that for inhalational anesthetics, desflurane is superior to sevoflurane, and sevoflurane is superior to isoflurane in reducing POCD. Dexmedetomidine may be the best current sedating agent for the mechanically ventilated patient that could enhance earlier release and decrease the incidence of delirium, but it is still unclear if these strategies also decrease the incidence of POCD. Ketamine was shown to reduce POCD in cardiac patients in one study and further research is likely indicated here. Epidural analgesia may be better than parenteral analgesia during the in-hospital recovery period [119], but, surprisingly, spinal anesthesia was not better but actually worse than general anesthesia for reducing post-operative cognitive dysfunction.

Obviously, myocardial ischemia should be avoided in general for several reasons, but recently gained insight indicates that maintenance of normal cardiac perfusion may reduce the chance for POCD. Furthermore, monitoring of cerebral oxygenation may be useful in select patient populations, although this has not been shown to reduce the incidence of the disease.

While both anesthesia and surgery have been associated with POCD, there are other factors that appear to contribute as well. For example, prolonged hospital stays, sleep deprivation in the hospital, and postoperative pain may all contribute to POCD. Minimizing length-of-stay, carefully managing post-operative pain, and improving patient sleep-efforts may help with this disease. Implementation of fast-track policy in orthopedic surgery could decrease early POCD [120].

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