Cerebrovascular Response to Propofol, Fentanyl, and Midazolam in Moderate/Severe Traumatic Brain Injury: A Scoping Systematic Review of the Human and Animal Literature

Logan Froese,1,* Joshua Dian,2 Carleen Batson,3 Alwyn Gomez,2,3 Bertram Unger MD,4 and Frederick A. Zeiler1–3,5,6

Abstract
Intravenous propofol, fentanyl, and midazolam are utilized commonly in critical care for metabolic suppression and anesthesia. The impact of propofol, fentanyl, and midazolam on cerebrovasculature and cerebral blood flow (CBF) is unclear in traumatic brain injury (TBI) and may carry important implications, as care is shifting to focus on cerebrovascular reactivity monitoring/directed therapies. The aim of this study was to perform a scoping review of the literature on the cerebrovascular/CBF effects of propofol, fentanyl, and midazolam in human patients with moderate/severe TBI and animal models with TBI. A search of MEDLINE, BIOSIS, EMBASE, Global Health, SCOPUS, and the Cochrane Library from inception to May 2020 was performed. All articles were included pertaining to the administration of propofol, fentanyl, and midazolam, in which the impact on CBF/cerebral vasculature was recorded. We identified 14 studies: 8 that evaluated propofol, 5 that evaluated fentanyl, and 2 that evaluated midazolam. All studies suffered from significant limitations, including: small sample size, and heterogeneous design and measurement techniques. In general, there was no significant change seen in CBF/cerebrovascular response to administration of propofol, fentanyl, or midazolam during experiments where PCO2 and mean arterial pressure (MAP) were controlled. This review highlights the current knowledge gap surrounding the impact of commonly utilized sedative drugs in TBI care. This work supports the need for dedicated studies, both experimental and human-based, evaluating the impact of these drugs on CBF and cerebrovascular reactivity/response in TBI.

Keywords: brain injury; cerebral blood flow; cerebrovascular response; fentanyl; midazolam; propofol

Introduction
Intravenous anesthesia is used universally within care for patients with severe brain injury for its neuroprotective properties.1 Its use is not limited to its ability to moderate cerebral metabolism; it also provides a more stable cerebral physiology in the presence of the severe trauma.1,2 Despite large-scale use of intravenous anesthetic agents, the impact that these commonly employed drugs have on various aspects of cerebral physiology in critical care patients, especially those with a traumatic brain injury (TBI), is largely unknown. This is in spite of their widespread adoption and recommendation through consensus-based guidelines for the management of moderate/severe TBI.3–5

1Biomedical Engineering, Faculty of Engineering, 7Centre on Aging, University of Manitoba, Winnipeg, Manitoba, Canada.
2Section of Neurosurgery, Department of Surgery, 6Division of Anesthesia, Department of Medicine, Addenbrooke’s Hospital, University of Cambridge, Cambridge, United Kingdom.
3Department of Anatomy and Cell Science, 4Section of Critical Care, Department of Medicine, Rady Faculty of Health Sciences, University of Manitoba, Winnipeg, Manitoba, Canada.

*Address correspondence to: Logan Froese, BSc, GF231 Health Sciences Centre, Biomedical Engineering, Faculty of Engineering, University of Manitoba, Winnipeg, Manitoba R3A 1R9, Canada, E-mail: log.froese@gmail.com

© Logan Froese et al., 2020; Published by Mary Ann Liebert, Inc. This Open Access article is distributed under the terms of the Creative Commons License (http://creativecommons.org/licenses/by/4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly credited.
Of particular interest is the impact on cerebral blood flow (CBF) and cerebrovascular reactivity of such sedative agents in TBI care, as current clinical guidelines focus on improving cerebral perfusion, CBF, and end-organ nutrient delivery.\textsuperscript{3,6–9} The body of literature surrounding the link between impaired cerebrovascular reactivity and poor patient outcome after TBI is growing,\textsuperscript{10–14} with data suggesting that in modern TBI care much of the ongoing cerebral physiological insult seen is dominated by impaired cerebrovascular reactivity.\textsuperscript{9,12,13,15} Further, cerebrovascular reactivity-based individual cerebral physiological targets, such as optimal cerebral perfusion pressure (CPP\textsubscript{opt})\textsuperscript{8,16–18} or individual intracranial pressure (iICP) thresholds,\textsuperscript{19,20} are emerging as novel methods to personalize treatment in TBI. Understanding the effects these commonly employed sedative agents have on CBF/cerebrovascular reactivity in the patient with severe TBI is a pivotal step in advancing personalized care.

The goal of this study was to perform a systematically conducted scoping review of all available literature on the impact of three commonly employed sedative agents used in moderate/severe TBI care (i.e., propofol, fentanyl, and midazolam) on cerebrovascular responsiveness/CBF response in human patients with moderate/severe TBI and animal TBI models.

Methods
A systematic review of the available literature was conducted using the methodology outlined in the Cochrane Handbook for Systematic Reviews of Interventions.\textsuperscript{21} The data were reported in line with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA).\textsuperscript{22} Supplementary Table S1 provides the PRISMA checklist. The review questions and search strategy were decided upon by the supervisor (F.A.Z.) and primary author (L.F.).

Ethical considerations
All articles are from previously published journals and have been vetted by their respective journals.

Search question, population, and inclusion and exclusion criteria
The question posed for systematic review was: “What is the effect of exogenous systemically administered propofol, fentanyl, or midazolam on the cerebrovascular response/CBF in human patients with moderate/severe TBI and animal models with TBI?” All studies, prospective and retrospective, of any size, based on humans and animals were included.

The primary outcome measure was the impact on CBF or the cerebrovascular responsiveness as documented by any objective means of CBF/cerebrovascular reactivity assessment, including continuous measures and neuroimaging-based or blood sampling-based techniques.

All original studies, whether prospective or retrospective, of all sizes, of any human age category or animal TBI model design, with the use of propofol/fentanyl/midazolam, and with formal documentation of cerebrovascular response/CBF during administration were eligible for inclusion in this review. Exclusion criteria were as follows: mild TBI literature, non-TBI human literature, being a non-English language study, or conducting CBF mediation with a substance other than propofol/fentanyl/midazolam.

Search strategy
MEDLINE, BIOSIS, EMBASE, Global Health, SCOPUS, and the Cochrane Library from inception to May 2020 were searched using individualized search strategies for each database. The search strategy for MEDLINE can be found in Supplementary Table S2, and a similar search strategy was used for the other databases. Finally, the reference lists of reviewed articles on the cerebral blood vessels/CBF response to propofol, fentanyl, and midazolam were examined to ensure no references were left out.

Study selection
Using two reviewers (L.F. and J.D.), a two-step review of all articles returned by our search strategies was performed. First, the reviewers independently screened all titles and abstracts of the returned articles to decide whether they met the inclusion criteria. Second, full text of the chosen articles was assessed to confirm whether the articles met the inclusion criteria and that the primary outcome of CBF/cerebrovascular response to propofol, fentanyl, and midazolam was documented. Any discrepancies between the two reviewers were resolved by a third party (F.A.Z.).

Data collection
Data were extracted from the selected articles and stored in multiple electronic databases to ensure data integrity.

Human studies
Data fields included the following: number of patients/animals, type of study, patient/model characteristics,
the goal of the study, dose of anesthetic administered, type of anesthetic administered, technique of CBF/vasculature assessment, CBF/cerebral vasculature response to drug, other outcomes, and general conclusions.

Bias assessment
Given the goal of this review was to provide a comprehensive scoping overview of the available literature, a formal bias assessment was not conducted.

Statistical analysis
A meta-analysis was not performed in this study because of the heterogeneity of model types, study designs, and data.

Results
Search results and study characteristics
The results of the search strategy across all databases and reference sections of articles are summarized in Figure 1. Overall, a total of 9896 articles were identified, all from the databases searched. A total of 4534 were removed because of duplication of references, leaving 5362 to review. By applying the inclusion/exclusion criteria to the title and abstract of these articles, we identified 400 articles that fit these criteria. One article was added from reference sections of pertinent review articles, leaving a total of 401 articles to review. The portable document formats (PDFs) of these 401 were then gathered. Applying the inclusion/exclusion criteria to these PDFs, only 14 articles were found eligible for inclusion in the systematic review.

Within the 14 TBI studies identified, there were 10 human TBI studies, and 4 animal TBI model studies. In the 10 human TBI studies, all the patients suffered a moderate/severe TBI, with human patients having a Glasgow Coma Scale (GCS) score of 12 or less on presentation. All studies measured CBF response to propofol, fentanyl, midazolam, and other agents: 5 used arterio-jugular differences of oxygen (AVDO2),10,23–26 2 studies used a Xenon133 diffusion technique,27,28 1 study used laser speckle imaging,29 1 study used radio-labeled microspheres,30 4 studies used transcranial-Doppler flow velocity,10,26,28,31 and 4 studies used CPP/PO2 as a surrogate for CBF.35 There were 3 studies that evaluated cerebrovascular reactivity/responsiveness, as measured by response of CBF to CO2 reactivity25,26 or a variety of other methods that used CBF and CBF velocity (CBFv).28 Regarding specific sedative agent studies, there were 8 studies that used propofol (2 of which used rat models29,31), 5 studies that used fentanyl (1 used rats36 and 1 used cats30), and 2 studies that used midazolam. The characteristics of the studies can be found in Table 1, Table 2, and Supplementary Table S3.

Propofol, fentanyl, and midazolam impact on objectively measured CBF
The following subsections provide a narrative summary of the impact of propofol, fentanyl, and midazolam administration on objectively measured cerebrovascular response/CFB in human patients followed by a brief summary of the four animal model studies. A summary of main study results can be found in Table 2, with more details for the interested reader in Supplementary Table S3. Of note, the following sections describe the trends presented in the parent articles. In all the human studies but one,34 partial pressure of carbon dioxide (PCO2) levels were either maintained or accounted for in cerebral response. PO2 was controlled in all studies through constant ventilation parameters. MAP was maintained at a constant level for most of these human studies, except for three studies where MAP was changed due to the sedative agent.25,27,32

Propofol. Within the six studies10,23,27,28,33,34 that evaluated propofol and CBF in human patients with TBI, most had a non-significant change in CBF. However, one study had a trend toward decrease to regional CBF when measured through a Xenon133 diffusion technique. Although it should be noted that there was also a significant drop in CPP and MAP, which could account for the decrease in CBF seen.27 Also, in this study individual patient responses were measured, demonstrating that most patients had a drop in CBF by at least 10 mL/100 g/min; further, in one patient cerebrovascular resistance (CVR; measured by CPP/CFB) was found to increase by 90% from baseline values (other patients had a limited response).

Two other studies displayed a non-significant response in CBF to propofol. Using transcranial-Doppler (TCD) to measure middle cerebral artery velocity (MCAv; which is a surrogate measure of CBF), these studies found the MCAv trended toward a decrease during propofol administration.10,28 In contrast to this CBFv change, CBF measured through AVDO2 methods demonstrated little response to propofol infusions.10 MAP, PCO2, and PO2 were relatively constant throughout, in both studies.

Finally, the three remaining studies demonstrated a non-significant CBF response to intravenous propofol
FIG. 1. PRISMA flow diagram. PRISMA, preferred reporting in systematic reviews and meta-analysis.
| References | No. patients/animals | Study type | Article location | Mean age | Patient/Animal characteristics | Primary and secondary goal of study |
|------------|---------------------|------------|-----------------|----------|-------------------------------|----------------------------------|
| **Human studies** | | | | | | |
| Lee et al.28 | 28 patients | Prospective cohort study | Journal | 33±13 years | TBI patients with GCS <7 | Primary: Assess influence of CO2 reactivity, pressure autoregulation, and metabolic suppression reactivity after head injury Secondary: Compare hemisphere response in CBF velocity |
| Steiner et al.10 | 10 patients | Prospective cohort study | Journal | 35±12 years | TBI patients with GCS score ≤12, 7 men and 3 women, with evacuated mass lesion in 8, diffuse injury in 2, large bilateral lesions in 5, and 7 had a craniectomy | Primary: Effect of propofol plasma concentration on pressure autoregulation |
| James et al.24 | 8 patients | Prospective randomized unblinded single crossover observational pilot study | Journal | Not mentioned | 4 patients with TBI, 3 with subarachnoid hemorrhage and one with intracerebral hemorrhage with median GCS score of 6.1 and acute physiology and chronic health evaluation 2 scores of 13.5 | Primary: Effects of dexmedetomidine and propofol on cerebral physiology in acute brain injury patients |
| Johnston et al.23 | 10 patients | Prospective cohort study | Journal | 21–53 years | TBI patients with GCS score 3–9, 8 patients with evacuated mass lesion and 2 with diffuse injury 2 | Primary: Assess the effect of propofol on cerebral oxygenation and metabolism in head-injured patients Secondary: Use propofol to achieve EEG burst suppression and evaluate overall effects on ischemic burden |
| Pinaud et al.27 | 10 patients | Prospective cohort study | Journal | 14–40 years | TBI patients with GCS score ≤6 | Primary: Effects of propofol on cerebral hemodynamics and metabolism in TBI patients Primary: Compare the cerebral microdialysis effects of propofol vs midazolam in TBI patients |
| Tanguy et al.33 | 30 patients | Retrospective cohort study | Journal | 35±18 years | TBI patients with acute physiological score 2–4 with mean GCS score 5 | Primary: Assess the effects of sufentanil, fentanyl, and alfentanil on cerebral hemodynamics |
| Albanese et al.24 | 6 patients | Randomized unblinded crossover study | Journal | 20–44 years | TBI male patients with GCS score 4–8 | Primary: Evaluate the cerebral hemodynamic effects of morphine and fentanyl in TBI patients Secondary: Correlation of morphine and fentanyl to cerebral autoregulation |
| de Nadal et al.26 | 30 patients | Randomized crossover study | Journal | 30±13 years | TBI patients with GCS score ≤8 | Primary: Evaluate the effects of fentanyl in TBI patients Secondary: Effect of midazolam on ICP and CPP in TBI patients Secondary: Evaluate cerebral damage by CPP increase |
| de Nadal et al.25 | 30 patients | Prospective cohort study | Journal | 30.2±13.2 years | TBI patients with GCS score ≤8 | |
| Papazian et al.32 | 12 patients | Prospective cohort study | Journal | 14–44 years | TBI patient with GCS score ≤6 | |
| **Animal studies** | | | | | | |
| Feuerstein et al.29 | 28 rats | Four-arm study | Journal | Not applicable | Male Wistar rats initially anesthetized with isoflurane, TBI method not mentioned | Primary: Evaluation of different methods to detect CBF and tissue deterioration after TBI Primary: Effects of propofol and isoflurane on cerebral hemodynamics during hypothermic conditions |
| Kahveci et al.31 | 16 rats | Two-arm study | Journal | Not applicable | Female Wistar rats with hypothermia and TBI caused from accelerated impact | |
| Bedell et al.30 | 17 cats | Two-arm study | Journal | Not applicable | Cats initially anesthetized with isoflurane and nitrous oxide, then TBI was induced with a fluid percussion injury | Primary: Influence of fentanyl on CBF during hypotension after TBI |
| Statler et al.36 | 51 rats | Two-arm study | Journal | Not applicable | Sprague-Dawley rats initially anesthetized with nitrous oxide and isoflurane, TBI was induced with control cortical impact | Primary: Evaluate the effects of isoflurane and fentanyl in TBI rats Secondary: Lesion volumes after TBI in rats |

CBF, cerebral blood flow; CPP, cerebral perfusion pressure; EEG, electroencephalogram; GCS, Glasgow Coma Scale; TBI, traumatic brain injury.
### Table 2. Sedation Treatment and Cerebrovascular Response: Summary of Study Details

| References                      | medication and dose | CBF/Cerebrovascular response | Limitations                                                                 | Conclusions                                                                 |
|--------------------------------|---------------------|-----------------------------|----------------------------------------------------------------------------|-----------------------------------------------------------------------------|
| **Human studies**              |                     |                             |                                                                            |                                                                             |
| Lee et al.²⁸                    | Propofol: 1 mg/kg   | Metabolic reactivity was induced through propofol burst suppression  | MAP during burst suppression was maintained with phenylephrine, which may interfere with CBF | Propofol through metabolic suppression decreased CBFv                        |
|                                |                     | -CPP increase by 5% (p<0.01) |                                                                            |                                                                             |
|                                |                     | -SjVO₂ increase by 3% (p<0.01) |                                                                            |                                                                             |
|                                |                     | -MAP was constant            |                                                                            |                                                                             |
|                                |                     | -MCAv in most models decrease by 30%, CBF also demonstrated a decrease but was not significant |                                                                            |                                                                             |
|                                |                     | -Trend to deteriorate vasoreactivity with 20% of patients having reduced response |                                                                            |                                                                             |
|                                |                     | Pco₂ and P0₂ levels were controlled through ventilation |                                                                            |                                                                             |
| Steiner et al.¹⁰               | Propofol: 3-4 mg/kg/h | Propofol                      | MAP was maintained with norepinephrine, which may interfere with CBF         | Propofol decreases MCAv, which is a surrogate measure of CBFv                |
|                                |                     | -Higher doses decreased MCAv by 8% |                                                                            |                                                                             |
|                                |                     | -Little change to CPP or AVDO₂ |                                                                            |                                                                             |
|                                |                     | -MAP remained relatively constant |                                                                            |                                                                             |
|                                |                     | -Static rate of autoregulation on average decreased from 56±36 to 28±35% but increased in some patients |                                                                            |                                                                             |
|                                |                     | Pco₂ and P0₂ levels were controlled through ventilation |                                                                            |                                                                             |
| James et al.³⁴                 | Propofol: 25.5 µg/kg/min | Propofol                      | Based on limited number of patients                                         | Propofol demonstrated little to no effect on CBF derived from the ICP/P0₂ comparison |
|                                | Dexmedetomine: 0.54 µg/kg/h | -Slight decrease in ICP and no change in Pbto₂ although both lacked statistical significance |                                                                            |                                                                             |
|                                |                     | -CPP increased during and fell after injection by about 6% |                                                                            |                                                                             |
|                                |                     | -Lactate/Pyruvate ratio increased drastically after injection |                                                                            |                                                                             |
|                                |                     | -CBF had minimal changes, based on limited response in ICP and P0₂ |                                                                            |                                                                             |
|                                |                     | Dexmedetomine                  |                                                                            |                                                                             |
|                                |                     | -A slight increase in ICP with no change in Pbto₂ though both lacked statistical significance |                                                                            |                                                                             |
|                                |                     | -CPP fell slightly by 2%      |                                                                            |                                                                             |
|                                |                     | -Lactate/Pyruvate ratio increased drastically after injection |                                                                            |                                                                             |
|                                |                     | -CBF had minimal changes, based on limited response in ICP and P0₂ |                                                                            |                                                                             |
|                                |                     | Pco₂ and P0₂ levels were controlled through ventilation |                                                                            |                                                                             |
|                                |                     | From ICP and CPP, MAP can be assumed to be near constant |                                                                            |                                                                             |
| Johnston et al.²³              | Propofol: 3-4 mg/kg/h | Propofol                      | ICP, CPP, and MAP can be assumed to be near constant                        | Propofol demonstrates a slight non-significant increase to CBF in the setting of TBI |
|                                |                     | -AVD0₂, ICP, and P0₂ all slightly decreased as compared with baseline values |                                                                            |                                                                             |
|                                |                     | -P0₂ and Pbto₂ slightly increased |                                                                            |                                                                             |
|                                |                     | -CPP and lactate/pyruvate ratio had little variation |                                                                            |                                                                             |
|                                |                     | -All changes were not significant |                                                                            |                                                                             |
| Pinaud et al.²⁷                | Propofol: 2 mg/kg then 150 µg/kg/min (3-5 µg/mL) | Propofol                      | Large variation within individual patients was not accounted for            | Propofol caused a varying decrease in rCBF in all patients; this was associated with a decrease in ICP and CPP |
|                                |                     | -rCBF decrease by 25% (p<0.01) |                                                                            |                                                                             |
|                                |                     | -iCBF decreased by 18% (p<0.001), this decrease was then inverted after propofol infusion ceased |                                                                            |                                                                             |
|                                |                     | -CPP dropped by 28% (p<0.001) |                                                                            |                                                                             |
|                                |                     | -AVD0₂ decreased by 6% but was not significant |                                                                            |                                                                             |
|                                |                     | -CVR increased then decrease as a result from propofol although this was not significant apart from 1 patient |                                                                            |                                                                             |
|                                |                     | -P0₂ remained constant at 33±2 mm Hg |                                                                            |                                                                             |

(continued)
| References | medication and dose | CBF/Cerebrovascular response | Limitations | Conclusions |
|------------|---------------------|----------------------------|-------------|-------------|
| Tanguy et al.33 | Propofol: 1 mg/kg/h and increased by same increment with 5 mg/kg/h being max Midazolam: 0.03 mg/kg/h and increased by 0.01 mg/kg/h | Propofol - ICP of 19±12 mm Hg - P02 of 97±2% - P02 of 38±7 mm Hg - CPP of 73±11 mm Hg - MAP of 91±11 mm Hg - CBF had minimal changes, based on limited response in CPP and P02 | MAP was maintained with catecholamines, which may interfere with CBF. Therapeutic goals and sedation levels were independent from microdialysis biomarkers | Using the CPP/P02 to find CBF, it is indicated that propofol and midazolam are near identical in CBF effect, both demonstrating no significant response |
| Albanèse et al.24 | Sufentanil: 1 μg/kg then 0.005 μg/kg/min Alfentanil: 100 μg/kg then 0.7 μg/kg/min Fentanyl: 10 μg/kg then 0.075 μg/kg/min | Sufentanil, alfentanil, and fentanyl - Initial increase ICP (25%) then after 60 min ICP returned to baseline - CPP decreased by 41% (p < 0.05) - SvO2 remained relatively unchanged - Based on CPP/ SvO2, CBF was indicated to increase - Based on CMRO2/AVD02, CBF slightly decreased - P02 levels were maintained between 32 and 35 torr and CPP stayed between 27 to 37 mm Hg - P02 and P0 levels were controlled through ventilation | No difference was seen in the lactate/pyruvate ratio was seen | CBF was approximated or found from the MCA MAP was maintained with phentylephrine, which may interfere with CBF |
| de Nadal et al.26 | Morphine: 0.2 mg/kg Fentanyl: 2 μg/kg | Morphine - Slight increase in CBF (10%) with no change in MCAv - When comparing autoregulation there was little difference in MCAv; however for CBF, impaired autoregulation demonstrated lower overall response (7%) then intact autoregulation (13%) Fentanyl - Slight increase in CBF (10%) and a slight decrease in CBFv (10%) - There was little difference in impaired vs. intact autoregulation in CBF and CBFv response All changing in AVD02 were adjusted for P02 levels MAP remained relatively constant in all groups Slight decrease in CBF associated with ICP then increase to baseline | CBF was approximated or found from the MCA MAP was maintained with phentylephrine, which may interfere with CBF | Fentanyl showed little change in CBF in any group whether with intact or impaired autoregulation although the direct measurement by CBFv and 1/AVD02 contradicted in response Morphine showed a slight increase in CBF with no significant change in CBFv. Intact autoregulation had a higher response then impaired autoregulation in any group whether intact or impaired autoregulation although the direct measurement by CBFv and 1/AVD02 contradicted in response |
| de Nadal et al.25 | Fentanyl: 2 μg/kg | Fentanyl - ICP: Increased then slowly decreased in both the group with intact and impaired autoregulation - CPP: Moderately decreased by 6% - AVD02 initially decreased (11%) then returned to the baseline at 60 min but was not significant MAP showed a similar decrease as CPP All changing in AVD02 were adjusted for P02 levels | CBF was approximated or found from the MCA MAP was maintained with phentylephrine, which may interfere with CBF | Fentanyl showed a decrease in CPP with a small increase in 1/AVD02 as a surrogate measure for CBF; this was a small and nonsignificant increase |

(continued)
| References | medication and dose | CBF/Cerebrovascular response | Limitations | Conclusions |
|------------|---------------------|----------------------------|-------------|-------------|
| Papazian et al.\(^{32}\) | Midazolam: 0.15 mg/kg | Midazolam | CBF assumed through CMRO\(_2\) coupling | CBF had little change apart from the mentioned ICP <18 mm Hg, in which case midazolam caused a slight increase in CBF. Midazolam was assumed to have limited influence on autoregulation due to limited difference in ICP groups. |
| Animal studies | | | | |
| Feuerstein et al.\(^{29}\) | Isoflurane at 2% Propofol: 33 to 53 mg/kg/h | Isoflurane: -rCBF increased initially with injection then returned to baseline after 1 min (19.8 ± 27.2%) Propofol: -rCBF increased initially with injection then returned to baseline after 1 min (27.5 ± 38.2%) -Atrial diameter decrease of 50% where isoflurane had no response Blood gasses were maintained through ventilation | Limited number of subjects | There was little response in rCBF in both groups, with propofol demonstrating a constriction of cerebral pial vessels. |
| Kalveci et al.\(^{31}\) | Propofol: 12 mg/kg/h Isoflurane: 0.9 ± 0.04% | Propofol: -Decrease ICP from 50% (\(p < 0.01\)) -CPP increased by 10% -No significant change to PO\(_2\), CBF\(_v\), or MAP Isoflurane: -No significant effect on CBF\(_v\), ICP, or PO\(_2\) -MAP and CPP decrease over time by 30% Blood gasses were maintained through ventilation | Subjects were also in a hypothermic state, which influences CBF | Despite the limited result, it was indicated that propofol is the better choice in hypothermic conditions, with no response in CBF\(_v\). The limited CBF effects of isoflurane are exaggerated by hypothermia indicating that isoflurane either caused no change or an increase in CBF. |
| Bedell et al.\(^{30}\) | Isoflurane: 1-1.5% Fentanyl: 50 \(\mu\)g/kg/h | Isoflurane: -ICP increased -CPP decreased by 7% then returned to baseline -MAP, CBF, and CVR remain relatively constant Fentanyl: -ICP decreased then slightly increased -CPP decrease by 30% -MAP decreased from 30% -CBF decreased by 22% at 75 min then increased to baseline -CVR decreased by 28% EEG, ICP, PO\(_2\), PCO\(_2\), pH, and temperature were similar between groups | Surgery may influence CBF | In the presence of hypotension fentanyl demonstrated a prevention of CBF indicating the fentanyl may increase CBF, along with this there was a decrease in CVR indicating a vasoconstrictive effect. Isoflurane had little influence on cerebral vasculature. |
| Statler et al.\(^{36}\) | Isoflurane at 4% then reduced to 1% Fentanyl: 50 \(\mu\)g/mL then 50 \(\mu\)g/kg/h | Fentanyl: MAP was higher than isoflurane; however during infusion the MAP and CPP remained constant throughout the experiment CPP after 4 h was greater in fentanyl than isoflurane group by 10%, but both constant CBF was 2 to 3 times higher in isoflurane then fentanyl group PO\(_2\) and PCO\(_2\) were controlled by ventilation | Subjects also sedated with nitrous oxide | The increase in CPP by isoflurane indicates that CBF is increased in contrast to the fentanyl demonstrating only minor change in CPP and therefore demonstrated little effect to CBF. |

AVDO\(_2\), arterio-venous oxygen differences; CBF, cerebral blood flow; CBF\(_v\), cerebral blood flow velocity; CMRO\(_2\), cerebral metabolic rate of oxygen; CPP, cerebral perfusion pressure; CVR, cerebrovascular resistance; EEG, electroencephalogram; h, hour; ICP, intracranial pressure; MAP, mean arterial pressure; MCA, middle cerebral artery; MCA\(_v\), middle cerebral artery velocity; min, minute; mm Hg, millimeters of mercury; PbtO\(_2\), brain tissue oxygen tension; PO\(_2\), partial pressure of oxygen; rCBF, regional cerebral blood flow; sec, second; SvO\(_2\), jugular venous oxygen saturation; TBI, traumatic brain injury.
administration. However, they did demonstrate a trend toward a decrease in CPP with no change in PO2. Such CPP and PO2 responses may indicate a decrease in CBF, based on CPP/PO2 as a surrogate measure for CBF. CPP and MAP remained unchanged in these studies.

**Fentanyl.** Within the three studies that evaluated the CBF effects of fentanyl in patients with TBI, all three had a non-significant response to fentanyl. However, there was a trend toward a decrease in CBFv found through TCD, with this drop found to be similar in patients with intact and impaired autoregulation (autoregulation was measured by comparing response of CBF with CO2 reactivity). In contrast to the CBFv decrease seen in these studies, a trend toward a CBF increase was demonstrated with an increase in 1/AVDO2 (surrogate measure for CBF). The PCO2 in these studies was between 29 and 35 torr, and MAP remained relatively unchanged during CBFv measurements.

**Midazolam.** In the two studies that evaluated CBF and midazolam in patients with TBI, there was a non-significant response to midazolam in CPP, PO2, and CBF. Although in one study the CPP and PO2 values were slightly higher in the midazolam group, compared with the propofol group. The second study demonstrated that midazolam decreases MAP by over 15 mm Hg, with patients who had an ICP <18 mm Hg before infusion demonstrating a slight increase in CPP. No definitive conclusions regarding the CBF/cerebrovascular reactivity response of midazolam can be made at this time.

**Animal studies**

In the four animal studies two compared propofol with isoflurane and the other two compared fentanyl with isoflurane. In all studies PO2 and PCO2 remained constant in all models. In the two studies in which propofol was evaluated in rat models, MAP was relatively constant. Both studies demonstrated a decrease in CBFv (measured through TCD of the MCA) or a decrease in CBF measured through laser speckle imaging with propofol administration. One of these studies had ICP drastically decreasing from 18 ± 2 to 7 ± 1 mm Hg (CPP decrease of 10%), and the other demonstrated a constriction of pial cerebral vessels by 50% (through direct visualization of vessels).

In the two remaining animal studies, the effects of fentanyl on CBF was evaluated. Both studies found the fentanyl groups displayed lower CBF and CPP values compared with the isoflurane groups, although CPP did trend toward increasing with fentanyl administration. The one study with feline models found fentanyl decreased MAP from 120 to 80 mm Hg with a significant drop in CBF (measured through radiolabel microspheres) and a slight decrease in CVR (calculated from MAP/CBF). Whereas the other study with rodents found that the fentanyl group had a CBF value that was 2 to 3 times lower than that in the isoflurane group, although the technique used and true value of CBF were not indicated.

**Discussion**

Through this systematically conducted scoping review of the literature surrounding the impact of propofol, fentanyl, and midazolam on CBF/cerebrovascular response in human and animal TBI, we have identified a significant knowledge gap. Although 14 studies were identified, they all suffered from some significant limitations, which restricted our ability to derive concrete conclusions regarding the CBF/cerebrovascular effects of these sedative agents. However, some general trends were seen in these studies.

First, in the studies identified propofol had a tendency to decrease CBF and CBFv. This has been previously described in healthy patients. However, it must be acknowledged that with the reduction in CPP seen in some of these studies with propofol, this alone may account for the CBF reductions. Further, some of the propofol studies estimated CBF using the 1/AVDO2 method, which is predicated on a relatively constant cerebral metabolic rate of oxygen (CMRO2). This may be the case in healthy patient populations, but likely does not hold true in the setting of TBI, where both regional and global changes in CMRO2 may fluctuate. Further, literature exists suggesting propofol may alter flow-metabolism coupling, further muddying the interpretation of CBF using the 1/AVDO2 technique. As such, no conclusive comments regarding the impact of propofol in CBF can be made at this time in patients with TBI, highlighting the need for future work.

Second, a decrease in CPP after fentanyl was seen in these TBI studies; this has been commented on in past review articles. Along with this, there was a
limited response in CBF in the three TBI studies with
$\text{PCO}_2$ being constant through the studies, indicating
that fentanyl has little influence on CBF in the setting
of ventilatory and cardiovascular support/control seen
during treatment in an intensive care unit (ICU). In
the animal models there was a decrease in CBF seen
with fentanyl administration, compared with isoflu-
urane, although the true influence of response is hard
to identify. In one of the animal studies the decrease
in CBF occurred with a concurrent decrease in
MAP.30 Thus, as with propofol, we are limited in
the conclusions that can be made, although there ap-
pears to be a no significant impact on CBF.

Third, midazolam was only evaluated in two stud-
ies with patients with TBI where CBF was objectively
assessed and did not appear to have any significant
impact on CBF. In one study there was a significant
decrease in MAP from 89 to 71 mm Hg with a non-
significant response to CBF.32 In healthy patients,
midazolam has been documented to decrease CBF and increase in CPP.2 Based on the studies identified,
it appears that in the setting of cardiorespiratory
control in the ICU, midazolam does not appear
to significantly impact CBF, although it must be
acknowledged that further work is required in this
area.

Finally, there was a limited response in CVR to ad-
ministration of sedative agents. For example, propofol
was found to have limited effects on CVR (CPP/CBF),
with one patient having a significant response in
CVR.27 Similarly, there was in one animal study that
analyzed cerebral pial vessel response to propofol
through direct visualization; vessels constricted by
50% as compared with baseline diameter.29 Whereas,
fentanyl found a trend toward a decrease in CVR in
one animal study, from 1.68 ± 0.46 to 1.21 ± 0.58 (as esti-

mated through CPP/CBF).30

Limitations
As mentioned above, the identified literature car-
ries significant limitations, which hinder our ability
to make conclusive statements regarding the CBF/
cerebrovascular response of propofol, fentanyl, and
midazolam in moderate/severe TBI. First, the litera-
ture body is low in number, consisting mainly of
small case series with limited sample sizes. As well,
many studies only demonstrated a weak non-
significant response, which could be influenced by
publication bias, therefore only trends may be com-
mented on. Second, the studies were heterogeneous
in nature, with different dosing and co-administration
of medications. Further, some patients were on vaso-
pressor drugs to support MAP and CPP during the
recorded CBF response. These drugs have known ce-
rebral vasoconstrictive properties and may therefore
have confounded the results. Third, most studies
employed the 1/AVD02 method for CBF estimation.
This method estimated CBF under the assumption
of relatively fixed CMRO2. This may be the case in
non-TBI patient populations but does not hold true
in the setting of moderate/severe TBI. Similarly, pro-
opofol is known to impact flow-metabolism coupling
in the brain and systemic blood pressure changes
could have caused the CBF response in many of these
studies.

These outlined limitations of the CBF measure-
ment technique further limit our ability to interpret if
these agents have a true impact on CBF. As well, CBFv
methods to evaluate MCAv make the assumption
that medium/large vessel changes in CBFv reflect
downstream CBF/cerebrovascular responses. Finally,
there is a lack of recorded high temporal physiology
responses of each drug with respect to CBF, relying
mainly on serological information for CBF estimation.
Thus, the true temporal CBF/cerebrovascular response
to these sedative agents in moderate/severe TBI re-


Future directions
It is clear from this review that knowledge of the
impact of commonly administered sedative agents on
CBF/cerebrovascular response in TBI is limited. As
such, we believe this review both highlights the knowl-
edge gap and provides evidence to support further
work in this area. Future investigations would benefit
from both experimental animal TBI models and in vivo
human studies in TBI. Both types of research re-
quire the use of continuous high temporal frequency
CBF/cerebrovascular reactivity measurement tech-
niques. These data would need to be time-linked to
medication dosing information, to provide the optimal
platform for exploring the temporal impact of such
sedation agents on CBF/cerebrovascular reactivity.
A multi-modal cerebral physiological monitoring
approach would be preferred, employing ICP, brain tis-
sue oxygen tension (PbtO2), thermal diffusion CBF,


neuromyoximetry, and cerebral microdialysis.
Similarly, objective assessments of sedation depth,
such as via processed electroencephalogram (EEG)
data, may remove the uncertainty around individual
There were a limited number of articles objectively documenting the CBF/cerebrovascular response of propofol, fentanyl, and midazolam in human patients with moderate/severe TBI and in animal TBI models. All studies suffered from significant limitations and small sample sizes, limiting the conclusions that can be drawn. In general, none of the agents had a significant impact on estimated CBF in the TBI populations described. This review highlights a significant knowledge gap present regarding the CBF/cerebrovascular response of these sedative agents in moderate/severe TBI, emphasizing the need for future dedicated experimental and human studies.

**Funding Information**

This work was supported by funding from a University of Manitoba UMG GFT grant, and the University of Manitoba URGP grant programs.

F.A.Z. receives research support from the Manitoba Public Insurance (MPI) Neuroscience/TBI Research Endowment, the United States National Institutes of Health (NIH) through the National Institute of Neurological Disorders and Stroke (NINDS) (NIH grant #R03NS114335-01), the Canadian Institutes of Health Research (CIHR) (CIHR grant #432061), the Canada Foundation for Innovation (CFI) (CFI grant #38583), Research Manitoba, the University of Manitoba VPRI Research Investment Fund (RIF), the University of Manitoba Centre on Aging, the University of Manitoba Rudy Falk Clinician-Scientist Professorship, and the Health Sciences Centre Foundation Winnipeg. L.F. is supported through a University of Manitoba – Department of Surgery GFT Research Grant, and the University of Manitoba Office of Research Services (ORS) – University Research Grant Program (URGP). C.B. is supported through a University of Manitoba Centre on Aging Fellowship Grant. A.G. is supported through the University of Manitoba Clinician Investigator Program.

**Author Disclosure Statement**

No competing financial interests exist.

**Supplementary Material**

Supplementary Table S1
Supplementary Table S2
Supplementary Table S3

**References**

1. Khandelwal, A., Bithal, P.K., and Rath, G.P. (2019). Anesthetic considerations for extracranial injuries in patients with associated brain trauma. J. Anaesthesiol. Clin. Pharmacol. 35, 302–311.

2. Oddo, M., Crippa, I.A., Mehta, S., Menon, D., Payen, J.-F., Taccone, F.S., and Citerio, G. (2016). Optimizing sedation in patients with acute brain injury. Crit. Care 20, 128.

3. Carney, N., Totten, L.A., Johnston, A.J., Chatfield, D.A., Czosnyka, M., Coleman, M.R., Coles, J.P., Gupta, A.K., Pickard, J.D., and Menon, D.K. (2003). The effects of large-dose propofol on cerebrovascular pressure autoregulation in head-injured patients. Anesth. Analg. 97, 572–576. table of contents.

4. Zeiler, F.A., Lee, J.K., Smielewski, P., Czosnyka, M., and Brady, K. (2018). Validation of intracranial pressure-derived cerebrovascular reactivity indices against the lower limit of autoregulation. Part II: experimental model of arterial hypotension. J. Neurotrauma 35, 2812–2819.

5. Steiner, L.A., Johnston, A.J., Chatfield, D.A., Czosnyka, M., Coleman, M.R., Coles, J.P., Gupta, A.K., Pickard, J.D., and Menon, D.K. (2003). The effects of large-dose propofol on cerebrovascular pressure autoregulation in head-injured patients. Anesth. Analg. 97, 572–576. table of contents.

6. Budohoski, K.P., Czosnyka, M., Kirkpatrick, P.J., Smielewski, P., Steiner, L.A., and Pickard, J.D. (2013). Clinical relevance of cerebral
autoregulation following subarachnoid haemorrhage. Nat. Rev. Neur. 9, 152–163.

12. Czosnyka, M., Smielewski, P., Kirkpatrick, P., Laing, R.J., Menon, D., and Pickard, J.D. (1997). Continuous assessment of the cerebral vasomotor reactivity in head injury. Neurosurgery 41, 11–19.

13. Zeiler, F.A., Ercole, A., Beqiri, E., Smielewski, P., Theilin, E.P., Cabeleira, M., Stocchetti, N., Vargiolu, A., Vilcinis, R., Wolf, S., and Younsi, A. (2019). Association between cerebrovascular reactivity monitoring and mortality is preserved when adjusting for baseline admission characteristics in adult traumatic brain injury: a CENTER-TBI Study. J. Neurotrauma 37, 1233–1241.

14. Zeiler, F.A., Ercole, A., Beqiri, E., Cabeleira, M., Aries, M., Zoerle, T., Malhotra, A.K., Schweitzer, J.B., Fox, J.L., Fabian, T.C., and Proctor, K.G. (2019). Predicting trauma mortality using Cerebral Autoregulation Monitoring System: a CENTER-TBI analysis. Acta Neurochir. Suppl. 140, 409–410.

15. Budohoski, K.P., Czosnyka, M., Smielewski, P., Menon, D.K., Cabeleira, M., Stocchetti, N., Kondziella, D., Koskinen, L.O., Meyfroidt, G., Moeller, K., Pickard, J.D., Laing, R.J., Menon, D., and Pickard, J.D. (1997). Continuous assessment of the cerebral vasomotor reactivity in head injury. Neurosurgery 41, 11–19.

16. Higgins, J.P.T., Thomas, J., Chandler, J., Cumpston, M., Li, T., Page, M.J., and Welch, V.A. (eds.). (2020). Cochrane Handbook for Systematic Reviews of Interventions, version 6.1 (updated September 2020). Cochrane. www.training.cochrane.org/handbook (Last accessed January 5, 2020).

17. Withers, D., Liberti, A., and Tetzlaff, J. (2009). Preferred reporting items for systematic reviews and meta-analysis: the PRISMA statement. Ann. Intern. Med. 151, 264–269.

18. Johnston, A.J., Steiner, L.A., Catchafield, D.A., Coleman, M.R., Coles, J.P., Alrawi, P.G., Menon, D.K., and Gupta, A.K. (2003). Effects of propofol on cerebral oxygenation and metabolism after head injury. Br. J. Anaesth. 91, 781–786.

19. Sufentanil, fentanyl, and alfentanil in head trauma patients: a study on cerebral hemodynamics. Crit. Care Med. 27, 407–411.

20. de Nadal, M., Sahuquillo, J., Pedraza, S., Garnacho, A., and Gancedo, V.A. (1998). Effects on intracranial pressure of fentanyl in severe head injured patients. Acta Neurochir. Suppl. 71, 10–12.

21. Higgins, J.P.T., Thomas, J., Chandler, J., Cumpston, M., Li, T., Page, M.J., and Welch, V.A. (eds.). (2020). Cochrane Handbook for Systematic Reviews of Interventions, version 6.1 (updated September 2020). Cochrane. www.training.cochrane.org/handbook (Last accessed January 5, 2020).

22. Moher, D., Liberati, A., and Tetzlaff, J. (2009). Preferred reporting items for systematic reviews and meta-analysis: the PRISMA statement. Ann. Intern. Med. 151, 264–269.

23. Johnston, A.J., Steiner, L.A., Catchafield, D.A., Coleman, M.R., Coles, J.P., Alrawi, P.G., Menon, D.K., and Gupta, A.K. (2003). Effects of propofol on cerebral oxygenation and metabolism after head injury. Br. J. Anaesth. 91, 781–786.

24. Fentanyl and alfentanil in head trauma patients: a study on cerebral hemodynamics. Crit. Care Med. 27, 407–411.

25. de Nadal, M., Ausina, A., Sahuquillo, J., Pedraza, S., Garnacho, A., and Gancedo, V.A. (1998). Effects on intracranial pressure of fentanyl in severe head injured patients. Acta Neurochir. Suppl. 71, 10–12.

26. Nadal, M., de Muren, F., Poca, M.A., Sahuquillo, J., Garnacho, A., and Rossello, J. (2000). Cerebral hemodynamic effects of morphine and fentanyl in patients with severe head injury: absence of correlation to cerebral autoregulation. Anesthesiol. J. Am. Soc. Anesthesiol. 92, 11–11.

27. Pinaud, M., Lalasue, J.N., Chetanneau, A., Fauchoux, N., Menegalli, D., and Souron, R. (1990). Effects of propofol on cerebral hemodynamics and metabolism in patients with brain trauma. Anesthesiology 73, 404–409.

28. Lee, J.H., Kelly, D.F., Oertel, M., McArthur, D.L., Glenn, T.C., Vespa, P., Boscardin, W.J., and Martin, N.A. (2001). Carbon dioxide reactivity, pressure autoregulation, and metabolic suppression reactivity after head injury: a transthalamic Doppler study. J. Neurosurg. 95, 222–232.

29. Feuerstein, D., Takagaki, M., Gramer, M., Manning, A., Endepols, H., Vollmar, S., Yoshimine, T., Strong, A.J., Graf, R., and Backes, H. (2014). Detecting tissue deterioration after brain injury: regional blood flow level versus capacity to raise blood flow. J. Cereb. Blood Flow Metab. 34, 1117–1127.

30. Bedell, E.A., DeWitt, D.S., and Prough, D.S. (1998). Fentanyl infusion preserves cerebral blood flow during decreased arterial blood pressure after traumatic brain injury in cats. J. Neurotrauma 15, 985–992.

31. Kahveci, F.S., Kahveci, N., Alkan, T., Goren, B., Korfall, E., and O’zlik, K. (2001). Propofol versus sufentanil anesthesia under hypothermic conditions: effects on intracranial pressure and local cerebral blood flow after diffuse traumatic brain injury in the rat. Surg. Neurol. 56, 206–214.

32. Papazian, L., Albanese, J., Thirion, X., Perrin, G., Durbec, O., and Martin, C. (1993). Effect of bolus doses of midazolam on intracranial pressure and cerebral perfusion pressure in patients with severe head injury. Br. J. Anaesth. 71, 267–271.

33. Tanguy, M., Seguin, P., Laviolle, B., Bleichner, J-P., Morandi, X., and Malledant, Y. (2012). Cerebral microdialysis effects of propofol versus midazolam in severe traumatic brain injury. J. Neurotrauma 29, 1105–1110.

34. James, M.L., Olson, D.M., and Graffagnino, C. (2012). A pilot study of cerebral and haemodynamic physiological changes during sedation with dexmedetomidine or propofol in patients with acute brain injury. Anaesth. Intensive Care 40, 949–957.

35. Heilbrun PM, Jorgensen PB, and Boysen G. (1972). Relationships between cerebral perfusion pressure and regional cerebral blood flow in patients with severe neurological disorders. Stroke 3 181–195.

36. Statler, K.D., Kochanek, P.M., Dixon, C.E., Alexander, H.L., Warner, D.S., Clark, R.S., Wnisierski, S.R., Graham, S.H., Jenkins, L.W., Marion, D.W., and Safar, P.J. (2000). Isoflurane improves long-term neurologic outcome versus fentanyl after traumatic brain injury in rats. J. Neurotrauma 17, 1119–1189.

37. Sahuquillo, J., Poca, M.A., Ausina, A., Baguena, M., Garcia, R.M., and Rubio, E. (1999). Arterio- jugular differences of oxygen (AVD02) for bedside assessment of CO2-reactivity and autoregulation in the acute phase of severe head injury. Acta Neurochir. (Wien) 138, 435–442.

38. Chong, K.Y., and Gelb, A.W. (1994). Cerebrovascular and cerebral metabolic effects of commonly used anaesthetics. Ann. Acad. Med. Singapore 23, 145–149.

39. Klein, K.U., Fukui, K., Schramm, P., Stadie, A., Fischer, G., Werner, C., Oertel, J., and Engelhard, K. (2011). Human cerebral microcirculation and oxygen saturation during propofol-induced reduction of bispectral index. Br. J. Anaesth. 107, 735–741.

40. Oshima, T., Karasawa, F., and Satoh, T. (2002). Effects of propofol on cerebral blood flow and the metabolic rate of oxygen in humans. Acta Anaesthesiol. Scand. 46, 831–835.
41. Slupe, A.M., and Kirsch, J.R. (2018). Effects of anesthesia on cerebral blood flow, metabolism, and neuroprotection. J. Cereb. Blood Flow Metab. 38, 2192–2208.

42. Maas, A.I.R., Menon, D.K., Steyerberg, E.W., Citerio, G., Lecky, F., Manley, G.T., Hill, S., Legrand, V., Sorgner, A., and CENTER-TBI Participants and Investigators. (2015). Collaborative European NeuroTrauma Effectiveness Research in Traumatic Brain Injury (CENTER-TBI): a prospective longitudinal observational study. Neurosurgery 76, 67–80.

43. Klein, S.P., De Sloovere, V., Meyfroidt, G., and Depreitere, B. (2019). Autoregulation assessment by direct visualisation of pial arterial blood flow in the piglet brain. Sci. Rep. 9, 13333.

44. Bernard, F., Gallagher, C., Griesdale, D., Kramer, A., Sekhon, M., and Zeiler, F.A. (2020). The CAnadian High-Resolution Traumatic Brain Injury (CAHR-TBI) Research Collaborative. Can. J. Neurol. Sci. J. Can. Sci. Neurol. 47, 551–556.

Cite this article as: Froese L, Dian J, Batson C, Gomez A, Bertram Unger B, Zeiler FA (2020) Cerebrovascular response to propofol, fentanyl, and midazolam in moderate/severe traumatic brain injury: A scoping systematic review of the human and animal literature, Neurotrauma Reports 1:1, 100–112, DOI:10.1089/neur.2020.0040.

Abbreviations Used

- AVDO₂ = arterio-jugular differences of oxygen
- CBF = cerebral blood flow
- CBFv = CBF velocity
- CMRO₂ = cerebral metabolic rate of oxygen
- CPPopt = optimal cerebral perfusion pressure
- CVR = cerebrovascular resistance
- EEG = electroencephalogram
- GCS = Glasgow Coma Scale
- ICU = intensive care unit
- iICP = individual intracranial pressure
- MAP = mean arterial pressure
- MCA = middle cerebral artery
- MCAv = middle cerebral artery velocity
- PbtO₂ = brain tissue oxygen tension
- PCO₂ = partial pressure of carbon dioxide
- PDF = portable document format
- rCBF = regional cerebral blood flow
- SvO₂ = jugular venous oxygen saturation
- TBI = traumatic brain injury
- TCD = transcranial-Doppler