Cerebral pressure autoregulation and carbon dioxide reactivity during propofol-induced EEG suppression

B. F. Matta, A. M. Lam, S. Strebel and T. S. Mayberg

Summary
We studied cerebral pressure autoregulation and carbon dioxide reactivity during propofol-induced electrical silence of the electroencephalogram (EEG) in 10 patients. Anaesthesia was induced with propofol 2.5 mg kg⁻¹, fentanyl 3 μg kg⁻¹ and vecuronium 0.1 mg kg⁻¹, and a propofol infusion of 250–300 μg kg⁻¹ min⁻¹ was used to induce EEG silence. Cerebral pressure autoregulation was tested by increasing mean arterial pressure (MAP) by 24 (SEM 5) mm Hg from baseline with an infusion of phenylephrine and simultaneously recording middle cerebral artery blood flow velocity (v(mca)) using transcranial Doppler. Carbon dioxide reactivity was tested by varying Pₐₐ₆ₐ in a dose-dependent manner. As typified by the barbiturates, this depression is associated with prolonged awakening which is undesirable in neurological patients. Propofol has similar cerebrovascular effects as thiopentone [9] and because of its pharmacokinetic profile, it may be suitable for this purpose. If propofol is to be used for cerebral protection, in high enough doses to produce EEG suppression, it is important to know the effects of complete EEG suppression on cerebral pressure autoregulation and carbon dioxide reactivity. The purpose of this study was to determine if cerebral carbon dioxide reactivity and pressure autoregulation remain intact during propofol-induced isoelectric EEG.

Patients and methods
After obtaining local Ethics Committee approval and informed written consent, we examined carbon dioxide reactivity and pressure autoregulation during propofol-induced isoelectric EEG in 10 patients (ASA I or II, mean age 37 (range 23–44) yr, weight 76 (SD 10) kg) undergoing anaesthesia for non-neurosurgical procedures. Patients who had cardiovascular, respiratory or neurological disease, and those receiving psychotropic drugs were excluded. After commencing routine monitoring (ECG, non-invasive arterial pressure and pulse oximetry), anaesthesia was induced with propofol 2.5 mg kg⁻¹ and fentanyl 3 μg kg⁻¹. Neuromuscular block was produced with vecuronium 0.1 mg kg⁻¹ and the lungs ventilated with an air-oxygen mixture to maintain normocapnia (Pₐₐ₆ₐ 5.3 (SD 1) kPa). Anaesthesia was maintained with a propofol infusion of either 250 or 300 μg kg⁻¹ min⁻¹ to produce isoelectricity of the EEG and the rate of infusion remained unchanged for the duration of the study. Repeated doses of vecuronium were used to maintain neuromuscular block. A radial arterial catheter was then inserted for continuous monitoring of mean arterial pressure (MAP) and repeated sampling for measurement of blood-gas tensions.

Key words
Anaesthetics i.v., propofol. Brain, blood flow.

Cerebral pressure autoregulation and vasoreactivity to carbon dioxide, two homeostatic mechanisms important for the control of cerebral blood flow (CBF), are impaired by pathology and some drugs used commonly during anaesthesia [1–5]. With the exception of ketamine, most commonly used i.v. anaesthetic agents depress also cerebral metabolism in a dose-dependent manner. As typified by the barbiturates, this depression is associated with a reduction in cerebral metabolic requirements for oxygen (CMRO₂) and CBF, which stabilize when electroencephalographic (EEG) activity becomes isoelectric [6, 7]. Barbiturate-induced cortical suppression has been advocated as a method of cerebral protection during temporary occlusion of any vessel that provides nutrients to the central nervous system [8]. Although such use is well documented, it is associated with prolonged awakening which is undesirable in neurological patients. Propofol has similar cerebrovascular effects as thiopentone [9] and because of its pharmacokinetic profile, it may be suitable for this purpose. If propofol is to be used for cerebral protection, in high enough doses to produce EEG suppression, it is important to know the effects of complete EEG suppression on cerebral pressure autoregulation and carbon dioxide reactivity. The purpose of this study was to determine if cerebral carbon dioxide reactivity and pressure autoregulation remain intact during propofol-induced isoelectric EEG.

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Cerebral blood flow velocity was measured by insonating the right middle cerebral artery (MCA) using a 2-MHz transcranial Doppler probe (Multidop by DWL, Sipplingen, Germany) through the temporal window. This method has been reported previously [10]. Briefly, the probe was secured in position so that the angle of insonation remained constant throughout the study. Doppler signals from the right MCA were identified and measured at a depth of 45–50 mm. The shift in frequency spectra of the Doppler signals converted into time–mean peak flow velocity (vmca) were displayed on a video monitor. To reduce ventilatory effects, measurements were obtained only during end-expiration and averaged over 3–4 cycles. Brain electrical activity was monitored using a two-channel fronto–occipital montage (Spacelab, Redmond, WA, USA), using a module which displayed unprocessed EEG signals. In all patients, an infusion of 0.01% phenylephrine was used to maintain mean MAP within the normal range of 70–90 mm Hg.

After 5–15 min of isoelectric EEG, relatively stable arterial pressure and normocapnia, the infusion rate of phenylephrine was increased so as to elevate MAP by approximately 20 mm Hg from baseline and vmca was recorded simultaneously. The infusion rate of phenylephrine was then decreased and MAP allowed to return to baseline. Carbon dioxide reactivity was then tested by varying PaCO2 between 4.0 and 7.0 kPa and recording vmca simultaneously. Arterial PaCO2 was varied by altering minute ventilation, primarily by changing the rate so that inspiratory pressure remained relatively constant. A minimum of 5 min for stabilization (unchanged end-tidal carbon dioxide) was allowed at each PaCO2 before measurements were made. Three pairs of PaCO2–vmca data were obtained in each patient for determination of carbon dioxide reactivity.

The index of autoregulation (IOR) was defined as the ratio of percentage change in estimated cerebral vascular resistance to percentage change in MAP. Estimated cerebral vascular resistance (CVRe) was calculated using the equation CVRe = MAP/vmca, where MAP = mean arterial pressure at the time of vmca measurement. An unchanged vmca would theoretically occur if the percentage change in CVRe was equal to the percentage change in MAP. Thus an IOR of 1 implies perfect autoregulation and an IOR of 0 complete absence of autoregulation (IOR = %ΔCVRe/%ΔMAP). Based on a dynamic autoregulation study [11], we considered a 15% change in IOR to be clinically insignificant. For a power of 0.80, an α error of 0.05 and a β error of 0.20, the required number of patients to reject the null hypothesis was 10. The computed IOR and data recorded before and after the phenylephrine-induced increase in MAP were analysed using Student’s paired t test. P < 0.05 was considered statistically significant.

To determine carbon dioxide reactivity, regression lines were constructed for the paired PaCO2–vmca data for each patient. Relative vmca was expressed also as a percentage of vmca at a PaCO2 of 5.3 kPa for all patients. Carbon dioxide reactivity was expressed in both absolute (change in vmca per kPa change in PaCO2) and relative values (percentage change in vmca per kPa change in PaCO2). For percentage changes, vmca was normalized to vmca at a PaCO2 of 5.3 kPa.

### Results

The major findings of the study are shown in table 1 and figures 1–3. There was no significant change in patients’ body temperature, haemoglobin concentration or partial pressure of oxygen (PaO2) in arterial blood during the course of the study (table 1). There was no change in PaCO2 during the autoregulation part of the study and no change in MAP during the carbon dioxide reactivity part of the study. The autoregulation data are shown in table 1. An increase in MAP of mean 24 (SEM 5) mm Hg had no effect on vmca. IOR was 0.93 (0.02) and was not significantly different from 1, indicating near perfect autoregulation. A representative trace is shown in figure 1.

Linear regression analysis demonstrated a close relationship between vmca and PaCO2 with correlation coefficients greater than 0.90 in all instances (P < 0.001). Although absolute cerebral vasoreactivity to carbon dioxide derived by linear regression analysis was lower than reported awake values, relative cerebral vasoreactivity to carbon dioxide was within normal limits for all patients studied (8.5 (SEM 0.8) cm s–1 kPa–1 and 22 (2)% kPa–1, respectively). Absolute vmca vs PaCO2 for all paired data and the respective linear regression lines (slopes) are shown in figure 2. Figure 3 shows

<table>
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<tr>
<th>Table 1: Temperature and haemodynamic data during pressure autoregulation (mean (SEM)). **P &lt; 0.01 compared with baseline (Student’s paired t test)</th>
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<tr>
<td><strong>Haemoglobin (g dl−1)</strong></td>
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<td>Temperature (°C)</td>
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<td>MAP (mm Hg)</td>
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<td>Heart rate (beat min−1)</td>
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<td>PaCO2 (kPa)</td>
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<td>vmca (cm s−1)</td>
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**Figure 1** Representative trace of autoregulation during propofol-induced EEG silence showing mean arterial pressure (MAP), mean cerebral blood flow velocity (vmca) and unprocessed EEG activity. Because of flow–metabolism coupling, the onset of EEG silence is associated with a sudden decrease in vmca.
relative cerebral vasoreactivity (relative umca vs $P_{a\text{CO}_2}$). Normalization is necessary to allow pooling of data with different baseline values.

**Discussion**

We have shown that during propofol-induced isoelectric EEG, both cerebral pressure autoregulation and carbon dioxide reactivity are intact. This suggests that both of these mechanisms are not dependent on cortical activity. The findings have important clinical implications as there is some experimental evidence which suggests that propofol has cerebral protective properties when used in high enough doses to induce EEG suppression [12].

There was no significant difference in patients' body temperature or haemoglobin concentration during the course of the study. Haemoglobin can affect flow velocity as haemodilution decreases viscosity and increases CBF and flow velocity [13]. Steady-state conditions were maintained during both parts of the study; MAP was maintained constant during the carbon dioxide reactivity part of the study, whereas $P_{a\text{CO}_2}$ was unchanged between normotension and phenylephrine-induced hypertension phases. The only other drug used was fentanyl, which has no direct cerebrovascular effects [14].

Using transcranial Doppler ultrasonography (TCD) it is possible to measure CBF velocity in a non-invasive continuous manner. However, CBF velocity is not a direct measure of CBF. Although techniques to measure CBF, such as radioactive xenon, are available, these are cumbersome and slow, require steady state and allow only a limited number of measurements. TCD recordings, on the other hand, may be obtained continuously. Although correlation between absolute flow velocity and CBF in any given population is poor, largely because of variation in MCA diameter, good correlation between relative changes in flow velocity and CBF has been demonstrated [15, 16]. Moreover, TCD studies on carbon dioxide reactivity indicated that this may be a particularly suitable tool for such investigations, as multiple paired measurements are taken and linear regression lines can be constructed more accurately than with a limited number of conventional blood flow measurements [15, 17, 18]. The validity of TCD rests on the assumption that the MCA is a conductance vessel and its diameter does not change significantly with changes in $P_{a\text{CO}_2}$. Changes in cerebral vascular resistance occur primarily by dilatation of arterioles and not the arteries of the circle of Willis [19]. Consequently, the MCA as a conductance vessel is unlikely to be affected by vasoactive agents. These assumptions have been confirmed recently by Giller and colleagues who observed during operation that the diameter of the proximal MCA does not dilate more than 4% during variations in $P_{a\text{CO}_2}$, or arterial pressure, nor does it change appreciably with systemic administration of nitroprusside or phenylephrine [20].

If we accept the assumption that the MCA region that TCD insonates does not change in diameter, then the change in blood flow velocity is proportional to changes in CBF. Indeed, Kirkham and co-workers have shown that carbon dioxide reactivity determined with TCD correlates well with carbon dioxide reactivity determined using direct methods of measuring CBF [18].

Normal umca varies from 35 to 90 cm s$^{-1}$ with a mean of approximately 60 cm s$^{-1}$ during the awake and resting states [22]. This range of umca probably reflects individual differences in MCA diameter, baseline CBF and the angle of insonation. The low absolute carbon dioxide reactivity seen during propofol-induced EEG silence can be explained by the cerebral vasoconstrictive effect of propofol and is consistent with results we reported previously during propofol anaesthesia [10]. Therefore, as baseline umca is low, the change per kPa change in $P_{a\text{CO}_2}$ in absolute numbers is similarly reduced. When this is normalized to a umca at a $P_{a\text{CO}_2}$ of 5.3 kPa, the relative slope expressed as a percentage approximates the awake value. Compared with the absolute change, the percentage change in umca with change in $P_{a\text{CO}_2}$ shows less dependence on baseline values, is therefore more appropriate for statistical analysis [23], and consequently is a more valid indicator of carbon dioxide reactivity.

Normalizing all umca values by expressing them as a percentage of umca at a $P_{a\text{CO}_2}$ of 5.3 kPa also allows comparisons with previously published
studies. As we measured umca and not CBF, we could only estimate cerebral vascular resistance (CVRe). This assumes that the changes in CBF correlate with changes in umca. This is not an unreasonable assumption as changes in CBF have been shown to correlate well with changes in umca [15, 16]. The index of autoregulation is then calculated by dividing the percentage change in CVRe by the percentage change in MAP. If the percentage change in CVRe is the same as the percentage change in MAP, no change in CBF results. Thus an autoregulation index of 1 indicates perfect autoregulation, and an IOR of 0 signifies complete absence of autoregulation. Mean IOR was 0.93 (SEM 0.02), indicating that good autoregulation was present during propofol-induced EEG silence. We considered a 15% deviation from IOR of 1 to be unimportant clinically and within normal limits, as dynamic autoregulation in normal volunteers has been shown to have a SD of 15% [11]. Power analysis ensured that 10 patients were sufficient to reject the null hypothesis with a power of 0.80. Based on this, we found the autoregulatory capacity to be intact during propofol-induced EEG suppression. Although we only examined autoregulation in one direction (increase in MAP), we chose a level of MAP in the middle of the autoregulatory curve. Therefore, there is no reason to assume that a decrease instead of an increase in MAP would have yielded different results.

A hallmark of normal cerebral vasculature is an intact autoregulatory response to changes in cerebral perfusion pressure (CPP). In the normotensive human, autoregulation is operative over a range of mean CPP of 50–150 mm Hg. Within this range, cerebrovascular resistance varies directly with MAP to maintain CBF constant. In the event of an abrupt change in CPP, blood flow initially changes correspondingly for a brief period (1–5 s) before the autoregulatory mechanism returns flow to control levels [11]. Some pathological states and pharmacological interventions can modify or abolish autoregulation. Moreover, there is evidence to suggest autoregulation is more susceptible to perturbation than carbon dioxide reactivity. Okuda and colleagues have demonstrated that in baboons made hypotensive by haemorrhage to an MAP beyond the lower limit of autoregulation (MAP of 40 mm Hg), and subsequently rendered normotensive, there was rebound cerebral hyperaemia (i.e. lost autoregulation) which responded to hypocapnia (i.e. intact carbon dioxide reactivity) [24]. In contrast, mild to moderate head injury often impairs autoregulation without affecting carbon dioxide reactivity [25]. In this study we investigated carbon dioxide reactivity and autoregulation while minimizing the effect of metabolic activity by rendering the EEG isoelectric. Cerebral blood flow and metabolism coupling is assumed generally to be intact during i.v. anaesthesia. It has been shown that i.v. agents such as the barbiturates produce cerebral vasoconstriction indirectly by reducing cerebral metabolism. Maximum reduction in CMRO₂ coincides with the development of electrical silence in the EEG, and additional administration of barbiturates results in no further reduction in CMRO₂ or CBF [26]. Thus a flat EEG is considered the point at which metabolic activity and CBF are depressed maximally and has been used as the end-point for cerebral protection. Although recent studies suggest that metabolic suppression may play only a small role in the overall scheme of cerebral protection [27], it continues to be used clinically as a method of cerebral protection and for reducing ICP [8, 28]. It seems logical that in situations of reduced oxygen delivery, keeping CMRO₂ to a minimum would increase the margin of safety in those patients at risk of cerebral ischaemia. The use of barbiturates for cerebral protection is associated with profound cardiovascular depression and prolonged awakening which may be undesirable in neurosurgical patients. Propofol, with similar cerebral vascular effects as thiopentone, has a relatively short duration of action and is useful for producing short periods of cerebral metabolic suppression without delayed awakening. Indeed, the protective effect of propofol has been demonstrated in an experimental ischaemic model [12]. The effect of suppressing cortical activity to the point of EEG silence with barbiturates has been shown not to significantly affect carbon dioxide reactivity in humans [29] or autoregulation in sheep [30]. We have now confirmed that high-dose propofol, sufficient to produce EEG silence, also preserved carbon dioxide reactivity and autoregulation. The preservation of these normal homeostatic mechanisms increases the safety of intraoperative manipulation of arterial pressure and carbon dioxide, at least in neurologically normal patients. Because of its pharmacokinetic profile, propofol may be preferable to thiopentone when EEG suppression is indicated clinically.

We conclude that in patients without neurological pathology, cerebral pressure autoregulation and carbon dioxide reactivity are intact during propofol-induced EEG silence. Whether this is true for neurosurgical patients with intracranial pathology requires investigation.

References

CO$_2$ reactivity and autoregulation

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