Assessment of Cerebrovascular Autoregulation in Head-Injured Patients
A Validation Study

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Background and Purpose—Cerebrovascular autoregulation is frequently measured in head-injured patients. We attempted to validate 4 bedside methods used for assessment of autoregulation.

Methods—PET was performed at a cerebral perfusion pressure (CPP) of 70 and 90 mm Hg in 20 patients. Cerebral blood flow (CBF) and cerebral metabolic rate for oxygen (CMR O2) were determined at each CPP level. Patients were sedated with propofol and fentanyl. Norepinephrine was used to control CPP. During PET scanning, transcranial Doppler (TCD) flow velocity in the middle cerebral artery was monitored, and the arterio-jugular oxygen content difference (AJD O2) was measured at each CPP. Autoregulation was determined as the static rate of autoregulation based on PET (SROR PET) and TCD (SROR TCD) data, based on changes in AJD O2, and with 2 indexes based on the relationship between slow waves of CPP and flow velocity (mean velocity index, Mx) and between arterial blood pressure and intracranial pressure (pressure reactivity index, PRx)

Results—We found significant correlations between SROR PET and SROR TCD ($r^2=0.32$; $P<0.01$) and between SROR PET and PRx ($r^2=0.31$; $P<0.05$). There were no significant associations between PET data and autoregulation as assessed by changes in AJD O2. Global CMR O2 was significantly lower at the higher CPP ($P<0.01$).

Conclusions—Despite some variability, SROR TCD and PRx may provide useful approximations of autoregulation in head-injured patients. At least with our methods, CMR O2 changes with the increase in CPP; hence, flow-metabolism coupling may affect the results of autoregulation testing. (Stroke. 2003;34:2404-2409.)

Key Words: autoregulation ▪ brain injuries ▪ monitoring, physiologic ▪ tomography, emission computed

Cerebrovascular autoregulation is defined as the brain’s ability to keep cerebral blood flow (CBF) relatively constant despite changes in cerebral perfusion pressure (CPP). This mechanism is frequently disturbed after head injury, even when the injury is only mild,1,2 and poor autoregulation after head injury is associated with unfavorable outcome.3–5 suggesting that this mechanism is a powerful protector of the injured brain against perfusion pressure–related secondary insults. It has also been suggested that targeting CPP according to an index of autoregulation might allow determination of an individual optimal CPP after head injury.6 Therefore, determination of autoregulation is of clinical interest in patients with traumatic brain injury.

Determination of autoregulation depends on an accurate assessment of CBF, which can be difficult. There are many methods available to measure CBF in head-injured patients, but many bedside methods do not measure CBF but instead monitor a surrogate marker considered to be proportional to CBF.7 In clinical practice, transcranial Doppler (TCD) is commonly used for dynamic8 and static9 measurements of autoregulation, although some investigators have used arterio-jugular oxygen content difference (AJD O2)10 or methods based on waveform analysis.4,11 When autoregulation is determined at the bedside, the assumption is made that the cerebral metabolic rate for oxygen (CMR O2) does not change during autoregulation testing. The validity of this assumption is critical, because changes in CMR O2 would mean that the measurement would reflect both autoregulation and flow-metabolism coupling. Measurement of static autoregulation requires a change in CPP, and vasoactive drugs such as norepinephrine are used to achieve this change. However, such drugs may influence cerebral metabolism, especially in head-injured patients, in whom the blood-brain barrier is potentially damaged. In view of these shortcomings, it is...
Twenty patients were investigated. Mean patient age was 33 ± 15 years; scans were performed 2.7 ± 1 days after injury, and median admission GCS was 6.5. Individual patient data are shown in Table 1. All patients were intubated and mechanically ventilated, sedated with propofol (2 to 5 mg · kg$^{-1}$ · h$^{-1}$) and fentanyl (1 to 2 μg · kg$^{-1}$ · h$^{-1}$), and paralyzed with atracurium. Infusion rates of these drugs were not changed during scanning. Patients had variable degrees of therapy for intracranial hypertension, including sedation, moderate hyperventilation (PaCO$_2$ 34°C to 37°C), surgical evacuation of space-occupying lesions, and barbiturate infusions (2 patients). However, no patient had received mannitol or hypertonic saline within the 6 hours preceding the study. All patients required catecholamines to maintain baseline CPP. Mean arterial pressure (MAP) and ICP were monitored by use of standard kits for invasive blood pressure monitoring (Baxter Healthcare Corp, CardioVascular Group) and intraparenchymal pressure transducers (Codman MicroSensors ICP Transducer, Codman & Shurtleff Inc).

Two sets of PET scans were performed, each assessing CBF and CMRO$_2$. CPP was controlled with an infusion of noradrenaline that was adjusted to reach the desired CPP and then as necessary to keep CPP constant during scanning. The first scan (baseline) was carried out at a CPP of 69 ± 6 mm Hg; the second (intervention), at a CPP of 92 ± 4 mm Hg. Because of the change in CPP, ICP increased from 18 ± 6 to 19 ± 6 mm Hg (P < 0.01). Arterial partial pressure of CO$_2$ (PaCO$_2$) was measured at 5 time points during each scan, and PaCO$_2$ was kept stable by adjusting the ventilator as necessary. Variability of PaCO$_2$ during acquisition of CBF data for individual patients is shown in Table 1. The precision of PaCO$_2$ control is limited not only by the respiratory status of the patients but also by the precision of the blood gas analyzer. For all blood gas measurements, an AVL Omni blood gas analyzer (AVL Graz, A-8020 Austria) was used. The total precision of this analyzer according to NCCLS document EPS-5 is SD < 0.07 kPa (personal communication, J. Riegebaumer, AVL Graz, 2002). PaCO$_2$ during the baseline scan was 4.41 ± 0.33 kPa and during the intervention scan was 4.48 ± 0.34 kPa (P = 0.07). Patient temperature was kept constant with the use of a heating/cooling mattress at the level desired by the team responsible for the clinical management. Patient temperature at baseline and at intervention was 35.4 ± 0.7°C (P = 0.4).

**PET Methods**

The studies were undertaken on a General Electric Advance scanner (GE Medical Systems). The steady-state protocol used has been described in detail previously. Emission images were coregistered to spiral CT images obtained immediately after PET scanning. All emission data were normalized to Talairach space. CBF and CMRO$_2$ were calculated globally and from the middle cerebral artery (MCA) territory for each hemisphere separately. The MCA territory was defined using high specificity at the cost of sensitivity: All pixels included had a very high probability of being perfused by the MCA, although some pixels that may have been within the MCA territory in individual patients may have been excluded.18

**Transcranial Doppler**

Bilateral TCD (DWL Multidop X4, DWL Elektronische Systeme GmbH) of the MCA was performed throughout scanning with two 2-MHz probes held in place with a head rack. Phantom testing carried out in the PET scanner before the patient studies established that there were no image artifacts caused by the ultrasound probes and head rack.

**Determination of Autoregulation**

For comparison of PET- and TCD-based determinations of autoregulation, SROR was used. SROR is calculated as the percent change in cerebrovascular resistance divided by the percent change in CPP used to induce the change in resistance. Results are expressed as percentage, with 0% representing complete autoregulatory failure and 100% representing optimal autoregulation. The formula is shown in the Appendix, which is available online at http://stroke.ahajournals.org. For calculation of SROR based on PET data (SROR$_{PET}$), CBF from the MCA territory and CPP were used as inputs into the formula. SROR$_{TCD}$ was calculated using mean flow velocity (FVm) instead of CBF. For comparison of PET and TCD, only FVm and CPP acquired during the phase of the PET scan during which CBF data were acquired were used.
During both scans, \( \text{AJDo}_2 \) was calculated from paired arterial and jugular blood samples withdrawn immediately before the \( \text{H}_2\text{O} \) infusion was started and used to calculate the percent change in CBF. According to the Fick principle, if \( \text{CMR}_{\text{O}_2} \) is constant, then the percentage decrease in \( \text{AJDo}_2 \) will equal the percentage increase in CBF, which can be compared with the percent change in global CBF as measured by PET. The formula is shown in the Appendix.

Two methods based on waveform analysis were investigated. The first method calculates an index of pressure reactivity from the analysis of spontaneous slow waves of MAP and ICP. Average values of MAP and ICP were calculated for 6-second intervals and used to calculate a pressure reactivity index (PRx) every 60 seconds as the linear moving correlation coefficient between 40 consecutive intervals for the regression. Similarly, this stratification implies intact autoregulation efficiency across the interval of CBF values that we studied. For comparisons between PET data and PRx or Mx, the last 2 variables were averaged over the complete duration of scanning at the higher level of CPP (\( \approx 90 \text{ mm Hg} \)), because sampling over longer time periods improves the estimation of these parameters.\(^{21,22} \) The value at the higher CPP was chosen because these measures of dynamic autoregulation would clearly have been nonconcordant at the 2 CPP levels and a consideration of cerebral vascular physiology shows that dynamic measures at the higher CPP value more closely represent the SROR that interrogated autoregulatory efficiency across the interval of CBF values that we studied. For comparisons between PRx and PET CBF, global CBF was used, whereas MCA territory CBF was used for the comparison with Mx.

Data were sampled from the analog output of the hemodynamic monitors, processed through an analog-to-digital converter (DT 2814, Data Translation Marlboro), and stored on a computer using software developed in house.\(^{23} \) Sampling frequency was 30 Hz. Time-averaged means for MAP, ICP, and CPP (CPP=MAP−ICP) were calculated and stored every 6 seconds. In 1 patient, we were unable to insonate the left MCA; therefore, for comparisons between TCD and PET, data from 39 hemispheres were available. Because of unavailability of monitoring equipment, data from 17 patients were available for comparisons between PRx, Mx, and PET. In 2 patients, technical problems prevented us from collecting CMRO\(_2\) data; therefore, CMRO\(_2\) data from 18 patients were available for analysis. Data are presented as mean±SD unless otherwise indicated. Linear correlations and Bland Altman plots\(^{24} \) were used to assess associations and agreement of measurement methods as appropriate. Calculations were performed with SPSS 11.0 (SPSS Inc). A value of \( P<0.05 \) was considered statistically significant.

**Results**

PET and TCD data are presented in Table 2. TCD FVm correlated significantly with PET CBF in both hemispheres (CPP=70 mm Hg: left, \( r^2=0.24, P=0.03 \); right, \( r^2=0.33, P<0.01 \); pooled, \( r^2=0.23, P=0.002 \); CPP=90 mm Hg: left, \( r^2=0.33, P=0.01 \); right, \( r^2=0.36, P<0.01 \); pooled, \( r^2=0.34, P=0.0001 \); Figure 1). However, there was marked variability that limits the usefulness of TCD as an estimator of absolute CBF. The change in CBF correlated with the change in FVm (left, \( r^2=0.48, P=0.001 \); right, \( r^2=0.42, P<0.01 \)). There was a significant correlation between SROR\(_{\text{PET}}\) and SROR\(_{\text{TCD}}\) (left, \( r^2=0.53, P<0.001 \); right, \( r^2=0.32, P<0.01 \)), suggesting that SROR\(_{\text{TCD}}\) is a useful approximation of autoregulation within the MCA territory (Figure 2a). The Bland Altman plot

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<td>SROR(_{\text{PET}}), %</td>
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Data are presented as mean±SD. FVm values represent data collected during PET CBF measurements.

(Figure 2b) shows that SROR\(_{\text{TCD}}\) measurements are on average 30% lower than SROR\(_{\text{PET}}\) measurements, and again there is a large variability.

\( \text{AJDo}_2 \) at baseline was normal or low, making global ischemia unlikely. However, increases in CPP resulted in a significant fall in \( \text{AJDo}_2 \) (4.0±1.2 to 3.2±1.1 mL · dL\(^{-1}\); \( P<0.001 \)), implying poor flow-metabolism coupling. There was no significant relationship between the estimated percent change in CBF based on \( \text{AJDo}_2 \) and the percent change in global CBF determined by PET (\( P=0.6 \)).

PRx was significantly associated with global \( \text{SROR}_{\text{PET}}\) (\( r^2=0.31, P=0.02 \)), with the relationship very close for low values of SROR but less so for those >80% (Figure 3). There was no significant relationship between \( \text{SROR}_{\text{PET}}\) and Mx in either hemisphere.

The correlation between PET and TCD did not appear to be confounded by the presence of lesions within the MCA territory. When MCA territories were grouped as lesioned or nonlesioned, the association for each subset was not significantly different from that observed in the nonlesioned (79.5±12.5 versus 69.1±10.1 mL · 100 mL\(^{-1}\) · min\(^{-1}\); \( P=0.03 \)) regions of interest. The reduction in MCA territories containing lesions was not significantly different from that observed in nonlesioned MCA territories without a lesion.

**Discussion**

We found that some but not all side measurements provide useful estimates of autoregulation. However, we observed a large variability even when significant associations were present. CMRO\(_2\) was not identical at the 2 levels of CPP; thus, measurements may reflect not only autoregulation but also variable\(^{25} \) flow-metabolism coupling.

Our study has 2 major methodological limitations. First, we use a very specific, albeit standard, clinical management strategy. This limits the transfer of our results to centers that use distinctly different management strategies. The second
important limitation is that, because of the precision of PET, which is strongly influenced by the low signal-to-noise ratio of the method, interpretation of results in patients with small changes in CBF, ie, good autoregulation, may be difficult. Despite reasonable overall agreement of data, measurements in patients with good autoregulation and therefore small changes in CBF are more likely to be influenced by noise. With our PET methods, the SD of repeated CBF measurements under constant physiology from the MCA territory in head injured patients is $1.7 \times 10^{-3}$ mL $\cdot$ 100 mL $\cdot$ min $\cdot$ 1 (unpublished data). This is illustrated in Figure 4, which shows that the agreement between PET and TCD measurements of SROR is better when patients with very low changes in CBF are excluded. This is further supported by Figure 3, which shows that the relationship between PET and PRx is close for low values of SROR but less so for high values of SROR. This suggests that a comparison of methods in patients with good autoregulation is affected not only by the sensitivity of the bedside methods to detect small changes in CBF but also by the precision and signal-to-noise ratio of PET. This also shows that it would be impossible to perform a validation in healthy volunteers with our methods.

Our data suggest that the percent change in AJD O$_2$ is not a reliable estimator of changes in CBF and therefore not a useful estimator of autoregulation. This may be due in part to the fact that the key assumption of constant CMRO$_2$ was not fulfilled. These changes in CMRO$_2$ associated with norepinephrine-induced CPP increases were often small but may be
particularly relevant with respect to estimates of autoregulation based on metabolism (eg, AIDO₂ measurements) as opposed to those based purely on an estimate of CBF (eg, SRORₚₑᵗ). We have not been able to validate Mx. There are several possible explanations for this. Earlier work has shown a close correlation between Mx and PRx,²⁶ which was not present in this group of patients. A selection bias could therefore have influenced our results. Alternatively, the number of investigated patients could be too low to establish this relationship. However, we suspect that our methods have prevented us from establishing a relationship between Mx and changes in PET CBF. Mx depends on spontaneous slow fluctuations of CPP of at least 5 mm Hg to elicit changes in FVm,²² whereas we tried to keep CPP as stable as possible and possibly suppressed the necessary slow waves to a relevant extent or created overly abrupt changes in CPP with some of the changes in the norepinephrine infusion rate. Our results suggest that an index based on MAP and ICP is more robust than one based on FVm and CPP. This is supported by data from another group that also used waveform analysis for quantification of autoregulation.²⁷

The decrease in CMRO₂ that we observed with the increase in CPP is unexpected. For CMRO₂, the SD for repeated scans is 1.3 μmol · 100 mL⁻¹ · min⁻¹ (unpublished data); therefore, the reductions, albeit small, are likely not to be due to the limited sensitivity or noise of the PET scan. We can only speculate on the reasons for this decrease in CMRO₂. One possible explanation is that the CMRO₂ changes are due to the clinical management, with propofol delivery increased at the higher CPP. Alternatively, it could be a specific effect of norepinephrine or of a disrupted blood-brain barrier. Regard-

Figure 3. Relationship between static rate of autoregulation measured by PET and PRx.

Figure 4. Effect of small changes in CBF. This Bland Altman plot illustrates the effect of small changes in CBF and their influence on the agreement between SROR measured with TCD and PET. Excluding CBF changes <1.7 mL · 100 mL⁻¹ · min⁻¹ improves the agreement considerably. Axes are scaled as in Figure 2b to enable direct comparison.
less of the underlying mechanism, this reduction in \( \text{CMRO}_2 \) is likely to lead to a reduction in CBF and thus an overestimation of autoregulation. The extent of the error will depend on the degree of flow-metabolism coupling, which may be disrupted to a variable degree after head injury.\(^{25,28} \) Bedside methods do not allow us to quantify this reliably.

In conclusion, the static rate of autoregulation calculated from TCD data and PRx provide useful information on autoregulation in head-injured patients. Studies grading autoregulation on that basis of changes in \( \text{AJDO}_2 \) need to be interpreted with caution. PRx seems to be a more robust estimator of autoregulation than Mx. More data are needed to validate Mx. At least when our methods are used, all measurements may be influenced by flow-metabolism coupling.\(^{20} \)

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References

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