TITLE:

THE CEREBRAL METABOLIC RATE FOR OXYGEN IS NOT DEPENDENT ON CEREBRAL OXYGEN DELIVERY IN CRITICALLY

ILL PATIENTS

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Introduction: Normally, systemic oxygen consumption ( $\mathrm{VO}_2$ ) is not altered by changes in systemic oxygen ( $\mathrm{O}_2$ ) delivery, except at very low levels of 0,2 delivery ( $\mathrm{DO}_2$ ). However, in ARDS, and 2below a critical threshold in anesthetized man,  $\mathrm{VO}_2$  does appear to be flow-limited. The relationship between regional 0,2 consumption and delivery has not been studied in critically ill patients. We performed studies of cerebral blood flow (CBF) and the cerebral metabolic rate for 0,4 (CMRO $_2$ ) in six critically ill patients to determine whether systematic changes in cerebral oxygen delivery (CDO $_2$ ) would produce corresponding changes in CMRO $_2$ .

Methőds: After approval by the Clinical Research Practices Committee, written consent was obtained. CBF was determined by the clearance of intravenous 133Xenon (133Xe) measured by 16 gamma detectors. End-tidal 133Xe concentration measured to correct for recirculation. CBF was calculated by the  ${\rm CBF}_{15}$  technique, corrected for temperature and hematocrit. Mean global CBF was determined by averaging CBF from all detectors. each patient a right jugular bulb catheter was placed. An initial measurement of CBF was then made. After this, CBF was increased or decreased so that CMRO, was calculated for each patient at a lower and higher CDO<sub>2</sub> (CBF CaO<sub>2</sub>). CBF was altered by increasing or decreasing PaCO<sub>2</sub> by changing mechanical ventilatory rate so as to result in the safest change in systemic pH. CBF measurement was repeated 30 minutes after changing the ventilatory rate. One minute after injection of 133Xe, systemic blood pressure was recorded, and samples were temic blood pressure was recorded, and samples were drawn for measurement of arterial and jugular venous partial pressures of O<sub>2</sub> and carbon dioxide, O<sub>2</sub> saturation (SaO<sub>2</sub>, SjvO<sub>2</sub>), and hemoglobin (Hb). From these data were calculated arterial and jugular venous O<sub>2</sub> contents (CaO<sub>2</sub>, CjvO<sub>2</sub>), arteriovenous O<sub>2</sub> content difference (Ca-jvDO<sub>2</sub>), CMRO<sub>2</sub> (CBF Ca-jvDO<sub>2</sub>), and CDO<sub>2</sub>. Mean CMRO<sub>2</sub> and CDO<sub>2</sub> data were compared using paired two-tailed <u>t</u> tests, significant at p<.05. significant at p<.05.

Results: At a mean lower CDO<sub>2</sub> of 3.8±1.8 (SD) ml/100g/min, CMRO<sub>2</sub> was 1.2±0.6 ml/100g/min. At a mean higher CDO<sub>2</sub> of 5.1±2.5 ml/100g/min, CMRO<sub>2</sub> was 1.4±0.9 ml/100g/min. While the difference in CDO<sub>2</sub> was significant, the difference in CMRO<sub>2</sub> was not. Mean arterial pressure, SaO<sub>2</sub>, hemoglobin, and temperature were not significantly different between data points. Before CDO<sub>2</sub> was experimentally altered, CDO<sub>2</sub> averaged 4.4±1.9 ml/100g/min, and CMRO<sub>2</sub> averaged 1.3±0.5 ml/100g/min (normal: CDO<sub>2</sub> = 7=10 ml/100g/min; CMRO<sub>2</sub> = 3-3.5 ml/100g/min). The figure shows data for six subjects, two of whom were studied on two different days. Patients 1, 2, and 3 were septic, and patients 4, 5, and 6 had closed head injuries without intracranial hypertension. Triangles indicate CDO<sub>2</sub> and

CMRO, obtained before alteration of CDO. Circles indicate  ${\rm CDO}_2$  and  ${\rm CMRO}_2$  obtained after alteration of CDO,.

<u>Discussion</u>: O<sub>2</sub> consumption is not normally limited by O<sub>2</sub> delivery, except at very low levels of O<sub>2</sub> delivery. Changes in O<sub>2</sub> consumption, systemic or cerebral, are primary determinants of systemic or cerebral blood flow. In some critically ill patients this relationship is abnormal, and oxygen delivery becomes a primary determinant of oxygen consumption. In this series of critically ill patients, changes in CDO<sub>2</sub> did not change CMRO<sub>2</sub>. Before experimental alteration of CDO<sub>2</sub>, both CDO<sub>2</sub> and CMRO<sub>2</sub> were approximately half normal. The failure of CNRO<sub>2</sub> to change in response to alteration of CDO<sub>2</sub> süggests that reduced utilization was not due to a primary CBF reduction. Efforts to improve CMRO<sub>2</sub>, and inferentially neurologic outcome, by increasing CDO<sub>2</sub> may be ineffective in some critically ill patients. However, in two of the head-injured patients, an increase in CDO<sub>2</sub> increased CMRO<sub>2</sub>. In such cases an increase in CBF could be beneficial, or hyperventilation could further impair O<sub>2</sub> utilization. More studies are needed to define subsets of patients in which an increase in CDO<sub>2</sub> produces an increase in CMRO<sub>2</sub>. It is also necessary to determine the most effective means of increasing CDO<sub>2</sub>, and to determine whether this improves neurological outcome.

## References:

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