Cerebral Metabolic Alterations in Rats with Diabetic Ketoacidosis: Effects of Treatment with Insulin and Intravenous Fluids and Effects of Bumetanide

Running title: MRS assessment of cerebral metabolism in rats with DKA

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Objective: Cerebral edema is a life-threatening complication of diabetic ketoacidosis (DKA) in children. Recent data suggest cerebral hypoperfusion and activation of cerebral ion transporters may be involved, but data describing cerebral metabolic alterations during DKA are lacking.

Research design and methods: We evaluated 50 juvenile rats with DKA and 21 normal control rats using proton and phosphorus magnetic resonance spectroscopy (MRS). MRS measured cerebral intracellular pH and ratios of metabolites including ATP/inorganic phosphate (Pi), phosphocreatine (PCr)/Pi, N-acetyl aspartate (NAA)/creatine (Cr), and lactate/Cr, before and during DKA treatment. We determined the effects of treatment with insulin and intravenous saline ± bumetanide, an inhibitor of Na-K-2Cl cotransport, using analysis of covariance with a 2 X 2 factorial study design.

Results: Cerebral intracellular pH was decreased during DKA compared to control (mean [± SE] difference = -0.13±0.03, p<0.001), and lactate/Cr was elevated (mean difference = 0.09±0.02, p<0.001). DKA rats had lower ATP/Pi and NAA/Cr (mean differences -0.32±0.10, p=0.003 and -0.14±0.04, p<0.001, respectively) compared to controls, but PCr/Pi was not significantly decreased. During two hours treatment with insulin/saline, ATP/Pi, PCr/Pi and NAA/Cr declined significantly, despite an increase in intracellular pH. Bumetanide treatment increased ATP/Pi and PCr/Pi, and ameliorated the declines in these values with insulin/saline treatment.

Conclusions: These data demonstrate that cerebral metabolism is significantly compromised during DKA, and further deterioration occurs during early DKA treatment, consistent with possible effects of cerebral hypoperfusion and “reperfusion injury”. Treatment with bumetanide may help to diminish the adverse effects of initial treatment with insulin/saline.
Cerebral injury resulting from diabetic ketoacidosis (DKA) is the most frequent diabetes-related cause of death in children. The cause of this complication has been a subject of much debate and is not well understood. Recent data from studies using an animal model demonstrate that DKA is associated with reduced cerebral blood flow and with brain cell swelling. (1; 2) The same studies suggest that activation of cerebral microvascular endothelial ion transport (particularly Na-K-Cl cotransport) during DKA contributes to brain cell swelling in this setting, as it does in animal models of stroke. During DKA treatment with insulin and intravenous fluids, cerebral blood flow rises and “vasogenic” edema develops. (3; 4) These data suggest that cerebral injury resulting from DKA may be similar to hypoxic/ischemic brain injury resulting from stroke or other causes, and raise the question of whether bumetanide, an inhibitor of Na-K-Cl cotransport, may be helpful in reducing cerebral injury in the setting of DKA.

Children who develop DKA-related cerebral injury often present with normal mental status or only mild mental status abnormalities. After several hours of DKA treatment, however, these patients have a decline in mental status, often followed by loss of consciousness and sometimes by clinical signs of increased intracranial pressure. Although a small percentage of children have clinically-apparent cerebral injury at presentation of DKA, prior to treatment, neurological decline during DKA treatment is more common. (5-7) The reason(s) that children may worsen during treatment with insulin and intravenous fluids are not clear. Initial hypotheses focused on the role of fluid infusion and osmotic change in causing brain cell swelling. (8; 9) More recent data suggest instead that reperfusion injury or related mechanisms may be more likely. (1-3)

Magnetic resonance spectroscopy (MRS) provides a non-invasive method for evaluating cerebral metabolism. Proton MRS can be used to measure concentrations of lactate in the brain, as well as the relative concentrations of N-acetyl aspartate (NAA) and creatine, a measure thought to be indicative of neuronal health.(10; 11) In stroke and other forms of hypoxic/ischemic brain injury, brain lactate is elevated and NAA/Cr is reduced. Phosphorus MRS can be used to measure cerebral intracellular pH and concentrations of high-energy phosphates, which are typically decreased during hypoxia/ischemia. (12-15)

MRS provides an ideal method to evaluate the cerebral metabolic alterations associated with DKA and the changes in cerebral metabolic state that occur during DKA treatment with insulin and intravenous fluids. We undertook the current study to characterize these metabolic changes and to determine the effects of treatment with bumetanide, an inhibitor of Na-K-2Cl cotransport in the blood-brain barrier and astrocytes, as well as many other cell types, on these metabolic alterations. In analogy with ischemia/reperfusion injury, we hypothesized that DKA would be associated with metabolic abnormalities similar to those of hypoxic/ischemic brain injury, and that these abnormalities would worsen during initial DKA treatment, as normal cerebral perfusion is re-established. Further, we hypothesized that bumetanide treatment would result in improvements in cerebral metabolic state.

RESEARCH DESIGN AND METHODS

Overview: A sequence of two experiments was performed. The first experiment compared metabolic measures in 50 rats with DKA to 21 normal control rats. DKA rats were then randomized to one of the four treatment combinations from a 2 X 2 factorial experiment designed to assess the treatment
effects of insulin/saline and bumetanide. Twelve rats were treated with bumetanide only, 11 with insulin/saline only, 14 with both and 13 DKA control rats were treated with neither. To assess whether estimates of the bumetanide-only effect could be confounded with the small volume of saline fluid used to deliver bumetanide intravenously, five of the 13 DKA control rats were intentionally treated with a small volume of saline and compared with the remaining control rats.

**Induction of DKA** - 71 four-week old Sprague Dawley rats (150 g, Charles River Laboratories, Wilmington, MA) were given an intraperitoneal injection of streptozotocin (STZ, 150 mg/kg, n=50) or STZ vehicle, as described previously.(1) Rats were given unlimited access to D10W (water with 10% dextrose, Fisher Scientific, Santa Clara, CA) in the first 24-hour period after STZ injection to prevent hypoglycemia and then were subsequently allowed unlimited access to tap water and standard rat chow. Rats were weighed daily. Urine glucose and ketoacids (acetoacetate) were measured using Multistix urinalysis strips (BAYER, Fisher Scientific, Santa Clara, CA) as described previously.(1) Rats in the DKA group had urine glucose and acetoacetate concentrations ≥ 110 mmol/L and 15,680 µmol/L, respectively. To induce a level of dehydration similar to severe human DKA and ensure acidosis, rats were deprived of drinking water 24 hours before imaging. This study was conducted in accordance with the Animal Use and Care Guidelines issued by the National Institutes of Health using a protocol approved by the Animal Use and Care Committee at University of California Davis.

**Magnetic resonance imaging** - MRS measurements were performed in anesthetized rats in a horizontal bore magnet (Oxford Instruments, Oxford, UK) using a two channel Biospec system (Bruker Biospin, Billerica, MA) running ParaVision software. A double tuned 1H/X Litz coil (Doty Scientific, Columbia, SC) was used where X is tunable for 23Na or 31P. Field homogeneity was optimized by localized shimming on 1H over a 9x9x9mm voxel of interest (VOI). The VOI was positioned inside the brain and selected to encompass as much of the cortex as possible. After shimming, 31P and 1H MRS data were acquired.

**31P MRS** - An 8x8x8mm VOI was centered on the shimmed volume. Spectra were acquired in 43.2 min intervals using Bruker software for image selected in vivo spectroscopy (ISIS), (TR=4 sec, number of signals averaged=80, line broadening=25 Hz). Intracellular pH (pH\textsubscript{i}) was calculated from the chemical shift of the inorganic phosphate (Pi) peak relative to the phosphocreatine (PCr) peak, using the equation pH\textsubscript{i} = 6.7 + log [(shift-3.186) / (5.691-shift)].(16) PCr, β-ATP, and Pi peaks were integrated using NUTS software (Acorn NMR, Livermore, CA) and presented as ratios (ATP/Pi, PCr/Pi).

**1H MRS** - A 7x7x7mm VOI was centered on the shimmed volume. Acquisition of 1H MRS data was initiated immediately after 31P MRS acquisition was completed. Spectra were collected in 2.8 min intervals, but to improve the signal-to-noise ratio for 1H measurements, two 2.8min files were added together for final analysis. Data were acquired using Bruker software for point-resolved spectroscopy (PRESS) (TR=6974ms, number of signals averaged=20, line broadening=2Hz) with chemical-shift-selective (CHESS) pulses and dephasing gradients to suppress water. Cerebral 1H metabolite peaks were identified according to their chemical shifts(17): N-acetylaspartate (NAA, 2.02 ppm), creatine and phosphocreatine (Cr, 3.0 ppm), lactate (1.38 ppm), and beta-hydroxy butyrate (βOHB, 1.15 ppm). With a TE of 132 ms, the lipid peak was suppressed, and the lactate and βOHB appeared as inverted peaks. The NAA, Cr, lactate and βOHB peaks were integrated.
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using NUTS software and presented as ratios (NAA/Cr, lactate/Cr, βOHB/Cr).

Experimental treatments - Saline/Insulin Infusion: For treatment with saline and insulin, rats were infused via cannulated femoral vein with Regular insulin at 1.5 U/kg/hr (Humulin, Lilly & Company, Indianapolis, IN) and 0.9% NaCl at 80 ml/Kg/hr for one hour, followed by infusion with insulin and saline at 1.5 U/kg/hr and 40 ml/Kg/hr, respectively, for the remainder of the 2 hour experiment. These rates were determined by comparisons of human vs. rat metabolic rate, body surface area and percentage dehydration during DKA. In initial studies, these rates of infusion resulted in biochemical changes during DKA treatment (decline in serum glucose and urea nitrogen concentrations and resolution of acidosis) at rates similar to those observed in children with DKA.

Bumetanide Treatments: For experiments designed to evaluate the effects of bumetanide on cerebral metabolite concentrations, bumetanide (30 mg/kg) was administered in one of two ways. For rats not receiving intravenous infusion of saline/insulin, bumetanide was injected into a femoral vein cannula (0.8 cc total volume) immediately before the start of imaging as described previously.(1) For rats treated with saline/insulin infusion, bumetanide was given via femoral vein cannula at the start of the saline/insulin infusion. Bumetanide (ICN Biomedicals, Costa Mesa, CA) was prepared as described previously.(1)

Animal preparation for imaging - Prior to imaging, rats were anesthetized using Na pentobarbital (IP 65 mg/kg). Body temperature was monitored via rectal probe (Cole-Parmer Instruments, Vernon Hills, IL) and a heating pad with circulating water (Gaymar Inc., Orchard Park, NY) maintained body temperature at 36.8-37.0°C throughout surgery and brain imaging. The femoral artery and vein were cannulated for blood sampling and for drug and treatment infusion, respectively. Rats were subjected to tracheal intubation and ventilated (Harvard Small Animal Ventilator, Holliston, MA) throughout surgery and imaging. Ventilation was done to offset the tendency toward respiratory depression in the anesthetized rats and thereby ensure that the animal model closely mimicked human DKA. Blood samples were analyzed for pCO\textsubscript{2} and pH immediately after intubation, and the respiratory rate and tidal volume adjusted with the goal of maintaining pCO\textsubscript{2} levels within the range expected for a normal physiological response to the degree of acidosis.(18)

Blood Chemistry - Blood samples were withdrawn from the femoral artery cannula before and hourly during imaging, and from the abdominal aorta, after imaging, at the conclusion of the experiment. We measured serum electrolyte concentrations, pH, blood urea nitrogen and glucose concentrations using an I-STAT Portable Clinical Analyzer (I-STAT; Sensor Devices, Waukesha, WI, U.S.A.).

Statistical Analysis - All statistical analyses were conducted using Version 9.1 of the SAS System for Windows. Two-sided testing was used for all study hypotheses, with P values less than 0.05 considered statistically significant and p values between 0.05 and 0.10 considered to represent a trend. Statistical analysis began with graphical and numerical summaries of the distributions of study outcomes and baseline measurements. When indicated, variables were log-transformed to ameliorate skewness and/or to stabilize variances across the groups under comparison. Group-specific (geometric) means and (geometric) standard deviations are reported for (log-transformed) baseline measures. At baseline, biochemical measures for some rats fell above or below the detection limits of measurement (BUN, log-transformed TCO\textsubscript{2} and serum glucose.
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concentrations). To account for this, maximum likelihood estimates of pre-treatment distribution parameters for these measures were produced, assuming Normal distributions.

For comparing DKA rats to normal control rats, Student’s independent groups t test was used for all outcomes, except those with some values outside the detection limits of measurement, which were analyzed using maximum likelihood estimates of regression models for heterogeneous limited dependent normally distributed variables (using SAS PROC QLIM). These models used a single independent variable that indicated group membership (1 “DKA rat” vs. 0 “Control rat”). The coefficient for this regressor was compared to the heterogeneity-robust standard error estimate to form the t-statistic used for testing between-group differences.

Student’s t test was used within the set of DKA control rats to compare the eight untreated rats with the five administered a small amount of saline. These comparisons were performed on baseline values and on change scores (from pre- to post-treatment). After establishing that these two subgroups did not have statistically significant differences on any comparison, the two subgroups were analyzed as a single group in subsequent analyses.

Within each of the four factorial treatment combinations, over-time changes in metabolite ratios were assessed using paired t-tests comparing pre- and post-treatment values. Between-groups comparisons on over-time changes were conducted using analysis of covariance models for a 2 X 2 factorial experiment, with main effects for the two binary treatment factors and, when indicated, heterogeneous error variance components. Baseline covariates were selected based on a preliminary stage of analysis that aimed to find a parsimonious set of predictors to improve model fit and the precision of estimates of main treatment effects. This set included serum pH for all study outcomes and, for outcomes where it improved model fit, an indicator for whether the baseline value of TCO₂ was outside the limit of detection. PCO₂ was also evaluated as a candidate variable for inclusion in the model, but was not found to improve model fit. Between-model comparisons for goodness-of-fit were performed using the Akaike Information Criterion and by comparing the root mean square error of the residuals. Analysis of residuals and influence statistics and fits of alternative regression models (with interaction terms) were used to verify that models for mean and covariance parameters were appropriately specified, were not unduly influenced by a small number of extreme observation and satisfied distributional assumptions needed for valid hypothesis testing. To accommodate the features of our final model, we used PROC MIXED to estimate model parameters, using restricted maximum likelihood estimates for variance components.

RESULTS

Biochemical values for DKA rats and normal control rats are summarized in Table 1. Using phosphorus MRS, we found that rats with DKA had significantly decreased cerebral intracellular pH compared with normal control rats (Figure 1). Peaks corresponding to the ketone body, βOHB, were readily detectable on proton MRS in DKA rats (mean βOHB/Cr ratio -0.16± 0.12), whereas no such peaks could be identified in normal control rats. Lactate to creatine (Cr) ratios were significantly increased on proton MRS in DKA rats (mean βOHB/Cr ratio -0.16± 0.12), whereas no such peaks could be identified in normal control rats. Lactate to creatine (Cr) ratios were significantly increased on proton MRS in DKA rats and N-acetyl aspartate (NAA)/Cr ratios were significantly decreased compared to control values. In phosphorus MRS, ATP/ inorganic phosphate (Pi) ratios were significantly decreased in DKA rats compared to normal controls, but PCr/Pi ratios were not significantly different between the two groups.
When rats with DKA were treated with intravenous insulin and saline, a deterioration in MRS measures of cerebral metabolism occurred (Figures 2 and 3). We observed a significant decrease in ATP/Pi, PCr/Pi and NAA/Cr ratios in rats treated with insulin and saline for 2 hours (Figures 4 A-C). These changes occurred despite improvements in cerebral intracellular pH (Figure 4 D), and decreased brain levels of βOHB (cerebral βOHB /Cr before treatment 0.12± 0.07 vs. 0.03± 0.03 after treatment, p<0.001), consistent with improvements in the ketoacidotic state. In contrast, rats with DKA who were left untreated for the same 2 hour period had no significant changes in ATP/Pi, PCr/Pi, NAA/Cr, cerebral intracellular pH (Figure 4 A-D) or intracerebral βOHB/Cr (0.17± 0.06 vs. 0.18± 0.11, p=0.84).

When bumetanide was added to the insulin and saline treatment, the mean ATP/Pi, PCr/Pi and NAA/Cr showed no significant change during treatment, rather than declining (Figures 4 A-C). Treatment of DKA rats with bumetanide alone, without insulin or saline, resulted in improvements in some metabolic measures. ATP/Pi rose significantly and there was a trend toward an increase in PCr/Pi (p=0.051). NAA/Cr levels, however, did not improve significantly in this group, nor did intracellular pH (Figure 4 D). Results of the ANCOVA analysis (Table 2) confirmed significant opposing effects of insulin/saline and bumetanide. While insulin/saline treatment worsened MRS metabolic measures despite improvements in intracellular pH, bumetanide treatment tended to improve MRS metabolic measures without significantly changing intracellular pH.

DISCUSSION
Case reports of cerebral edema and cerebral injury occurring during DKA in children often describe the child’s initial mental state as normal or nearly normal at the time of presentation with DKA. After several hours of treatment with insulin and intravenous fluids, however, a decline in mental status occurs, often with loss of consciousness, seizures or other substantial neurological abnormalities.(19-21) This decline in mental status occurs despite improvements in acidosis and hyperglycemia. Although clinically-apparent cerebral edema and cerebral injury can also occur before treatment of DKA, the more frequent occurrence of cerebral edema during DKA treatment suggests that some aspect of treatment may cause or enhance cerebral injury.

Our data demonstrate that cerebral intracellular pH is low during untreated DKA, cerebral lactate levels are high, and levels of high-energy phosphates are low, similar to cerebral ischemia. NAA/Cr ratios are also decreased, suggesting neuronal compromise or injury.(10-14; 22-25) Taken together with our previous results demonstrating that DKA diminishes CBF in this model(2), these data provide further evidence consistent with the hypothesis that cerebral hypoperfusion occurs in untreated DKA and may lead to cerebral injury. Similar findings have been observed in both human and animal studies of stroke and other ischemic brain injury, including declines in brain concentrations of high energy phosphates, elevated brain lactate concentrations and decreased NAA/Cr ratios.(12; 14; 15; 23-26) Additionally, our data provide the first evidence that during initial DKA treatment with insulin and intravenous saline, key aspects of the cerebral metabolic state worsen. High-energy phosphate levels decline further, as does the NAA/Cr ratio. These data suggest that the initial period of DKA treatment may lead to additional cerebral injury, possibly caused by reperfusion of previously hypoperfused cerebral tissues or some other aspect of treatment.

Data from previous studies suggest that hyperglycemia augments ischemic brain
Hyperglycemia results in increased brain lactate concentrations and reduced high energy phosphate concentrations following an ischemic insult. During ischemia and reperfusion, hyperglycemia is associated with greater and more prolonged intracellular acidosis. Our data correlate well with these findings and suggest that hyperglycemia may result in greater vulnerability of the brain to injury resulting from diminished perfusion.

Although osmotic fluctuations during DKA therapy have been suspected to cause cerebral edema, our data are more consistent with the effects of ischemia and reperfusion. Limited data from other studies suggest that osmotic fluctuations do not result in changes in cerebral high-energy phosphate levels. In addition, previous studies by our group have demonstrated high apparent diffusion coefficient (ADC) values measured by MR diffusion weighted imaging during DKA treatment in children, in contrast to the low ADC values observed in cerebral edema induced by osmotic fluctuations. These data suggest that declines in osmolality during DKA treatment are unlikely to be the main cause of cerebral injury.

In the current study, treatment with bumetanide, an inhibitor Na-K-2Cl cotransport, resulted in improvements in metabolic measures during untreated DKA, and amelioration of the declines in metabolic measures during initial DKA treatment. These data suggest a protective effect of bumetanide. Previous data from our group have demonstrated that untreated DKA is associated with reduced brain ADC values, suggesting brain cell swelling. Treatment in these studies resulted in an increase in ADC, suggesting reduced cell swelling. While elucidating the mechanisms underlying this effect of bumetanide will require further investigation, previous studies of the Na-K-Cl cotransporter in healthy and diseased brain provide some clues. The Na-K-Cl cotransporter is known to be present in cerebral microvascular endothelial cells (also called blood-brain barrier, BBB), astrocytes and neurons and to serve a number of functions, depending on cell type and prevailing physiological/pathophysiological conditions. These findings have been reviewed previously. Briefly, in healthy brain the BBB Na-K-Cl cotransporter (predominantly in the luminal membrane) is thought to participate in secretion of Na, Cl and water from blood into brain, accounting for up to 30% of brain interstitial fluid generation. During the early hours of ischemic stroke, factors including hypoxia, aglycemia and vasopressin stimulate activity...
of the BBB cotransporter leading to increased secretion of Na, Cl and water across the intact barrier from blood into brain. (42; 43; 45) Ischemic factors also stimulate astrocyte Na-K-Cl cotransport activity causing the cells to take up ions and water crossing the BBB and to swell (cytotoxic edema). As ischemia progresses, the endothelial cells themselves begin to swell by a process that is at least partially dependent on Na-K-Cl cotransporter activity. Ischemic stimulation of the cotransporter can also cause swelling of neurons, although there is some debate about the extent of neuronal swelling compared to astrocytes. In addition, increased Na-K-Cl cotransporter activity in GABAergic neurons causes elevation of intracellular [Cl] and thus increased efflux of Cl through GABA-activated Cl channels and depolarization of the cells. Neuronal cotransporter activity is high in immature neurons and appears to contribute to neonatal seizures.(41; 46; 47) In mature brain, the cotransporter may also contribute to seizures occurring after ischemia/reperfusion by increasing intracellular [Cl] and causing hyperexcitability of GABAergic neurons. Previous studies have also shown that elevation of intracellular [Na] stimulates Na/K ATPase activity, consequently increasing ATP consumption as long as ATP is available (48; 49) and thus inhibition of Na uptake pathways can decrease ATP consumption (50-53). The observed effects of bumetanide on metabolic parameters in the present study are consistent with the possibility that DKA-induced cerebral hypoxia/ischemia stimulates Na-K-Cl cotransporter-mediated Na influx (in BBB, astrocytes and/or neurons), elevating intracellular [Na] and stimulating Na/K ATPase activity and ATP consumption and that bumetanide reduces ATP consumption by reducing cotransporter-mediated Na influx. Interestingly, although ATP/Pi levels were significantly reduced in DKA rats, PCr/Pi levels in DKA rats were not significantly different from control values. These data initially appear counterintuitive because declines in PCr/Pi caused by cerebral ischemia typically are of greater magnitude than observed declines in ATP/Pi.(13; 14; 23; 54) Data from human studies of hyperglycemia, however, demonstrate that brain PCr concentrations increase during hyperglycemia.(55) A modest increase in ATP concentrations also occurs, but the increase in PCr is greater, resulting in an increase in the PCr/ATP ratio. The lack of a detectable difference between DKA rats and controls in PCr/Pi in the current study may therefore reflect higher baseline PCr levels in the DKA rats induced by hyperglycemia. The current study has some limitations. First, under conditions where brain high energy phosphate metabolism is near normal, Pi levels are commonly near the noise level obtained in our data. This is likely to have caused a relative increase in variability for parameters dependent on Pi, (intracellular pH, PCr/Pi and ATP/Pi) particularly under control conditions. This variability may have decreased our ability to detect differences of smaller magnitude between groups. In addition, because of the inherent limitations of mechanical ventilation in small animals, we were not always able to precisely adjust the animals’ pCO2 level to that expected for the degree of acidosis. For these reasons, we conducted a sub-analysis including the pCO2 level as a covariate in the model. Inclusion of pCO2 was not found to improve model fit, suggesting that differences in pCO2 level between the groups did not have a significant effect on the study outcomes. Finally, although our data suggest a beneficial effect of bumetanide, we investigated only the initial phase of DKA treatment. Whether bumetanide treatment results in decreased neurological injury later in the course of DKA treatment, or improved outcomes after recovery from DKA is not yet known.
In summary, our data demonstrate that DKA results in metabolic changes in the brain similar to those occurring in hypoxic/ischemic conditions. Furthermore, initial DKA treatment with insulin and intravenous saline, rather than resulting in improvements in the cerebral metabolic state, results in further deterioration despite recovery of intracellular pH. These data may help to explain the more frequent occurrence of DKA-related cerebral injury during DKA treatment, rather than at the time of presentation. Treatment with bumetanide to inhibit Na-K-2Cl cotransport results in improvements in cerebral metabolic measures, suggesting a protective effect. Our data suggest the need for further investigation of the effects of bumetanide during DKA treatment in children.

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REFERENCES


### Table 1: Biochemical Values in Normal Control Rats and in Rats with DKA before and after Two Hours of Treatment with Insulin and Saline Infusion

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Control (n=21)</th>
<th>DKA before infusion (n=49)</th>
<th>DKA after 2 hr. saline/insulin infusion (n=11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose (mmol/L)</td>
<td>7.9 (0.9)</td>
<td>35.7 (9.1)</td>
<td>17.3 (8.4)</td>
</tr>
<tr>
<td>BUN (mmol/L)</td>
<td>3.9 (1.1)</td>
<td>34.3 (13.2)</td>
<td>25.3 (10.7)</td>
</tr>
<tr>
<td>pH</td>
<td>7.42 (0.06)</td>
<td>7.10 (0.23)</td>
<td>7.20 (0.13)</td>
</tr>
<tr>
<td>Total CO$_2$ (mmol/L)</td>
<td>27 (1)</td>
<td>7 (2)</td>
<td>12 (1)</td>
</tr>
</tbody>
</table>

* NOTE: DKA pre-infusion values represent pooled values for all DKA treatment groups. Post-infusion values include rats in the standard (insulin/saline) DKA treatment group only. Among DKA rats before infusion, 20 of 49 had glucose measurements above the detection limit (DL) of 38.5 mmol/L, 9 of 49 had BUN measurements above the DL of 50 mmol/L and 18 of 48 had a Total CO$_2$ measurement below the DL of 5 mmol/L. Pretreatment parameters for BUN, Glucose and Total CO$_2$ are maximum likelihood estimates, as described in the methods section. Values shown are means with SD in parentheses for glucose, BUN and pH. Values shown for total CO$_2$ are geometric means with geometric SD in parentheses.

### Table 2: Individual Effects of Insulin/Saline and Bumetanide on Cerebral Metabolites during DKA: Regression Adjusted Main Effects of each Treatment

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Adjusted Main effect of Insulin/Saline* (SE)</th>
<th>p</th>
<th>Adjusted Main effect of Bumetanide* (SE)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATP/Pi</td>
<td>-0.34 (0.13)</td>
<td>0.02</td>
<td>0.38 (0.15)</td>
<td>0.01</td>
</tr>
<tr>
<td>PCR/Pi</td>
<td>-0.29 (0.10)</td>
<td>0.007</td>
<td>0.31 (0.13)</td>
<td>0.03</td>
</tr>
<tr>
<td>NAA/Cr</td>
<td>-0.08 (0.03)</td>
<td>0.020</td>
<td>-0.001 (0.03)</td>
<td>0.97</td>
</tr>
<tr>
<td>Intracellular pH</td>
<td>0.13 (0.04)</td>
<td>0.005</td>
<td>-0.002 (0.04)</td>
<td>0.72</td>
</tr>
</tbody>
</table>

* Values are adjusted mean effect of changes within rat groups as defined in Methods. SE values are shown in parentheses. ATP/Pi and PCR/Pi were analyzed as log transformed values.
FIGURE LEGENDS

Figure 1. Cerebral metabolites measured by 1H and 31P MRS in DKA and control rats. A) ATP:Pi ratios B) PCr:Pi ratios C) NAA:Creatine ratios D) Intracellular pH E) Lactate:Creatine ratios. All values are means with 95% confidence intervals; figure A and B are geometric means and figure C, D and E are arithmetic means. n=20 for control and 41 for DKA rats in figure C and D; n=20 for control and 44 for DKA rats in figure A, B and E. * Values are significantly different from control group, p-values <0.005.

Figure 2. 1H MR spectra obtained before DKA treatment and at the end of 2 hours treatment with insulin/saline. [Peak label abbreviations are: phosphocreatine+creatine – Cr, N-acetylaspartate – NAA, lactate – Lact, ßhydroxybutarate – ßOHB]

Figure 3. 31P MR spectra obtained before DKA treatment and at the end of 2 hours treatment with insulin/saline. [Peak label abbreviations are: inorganic phosphate – Pi, phosphocreatine – PCr, γATP – γ, αATP – α, βATP – β]

Figure 4 A. ATP:Pi ratios in DKA rats pre- and post- saline/insulin infusion with and without bumetanide. ATP to Pi ratios were measured by 31P MRS as described in RESEARCH DESIGN AND METHODS. Bumetanide alone and saline/insulin with and without bumetanide treatments started immediately after baseline measurements (0 hr). All values are geometric means with 95% confidence intervals, n= 12, 10, 10, and 12 for no treatment, bumetanide alone, saline/insulin, saline/insulin with bumetanide treatment groups, respectively. * Within group comparisons: p < 0.02 for comparison of pre/post treatment values in bumetanide alone group and insulin/fluid group. Refer to table 2 for the overall effects of insulin/saline and bumetanide. Note: Data were analyzed with log transformed values.

B. PCr:Pi ratios in DKA rats pre- and post- saline/insulin infusion with and without bumetanide. PCr to Pi ratios were measured by 31P MRS as described in RESEARCH DESIGN AND METHODS. Bumetanide alone and saline/insulin with and without bumetanide treatments started immediately after baseline measurements (0 hr). All values are geometric means with 95% confidence intervals, n= 12, 10, 10, and 12 for no treatment, bumetanide alone, saline/insulin, saline/insulin with bumetanide treatment groups, respectively. * Within group comparisons: p =0.01 for comparison of pre/post treatment values in insulin/fluid group. p=0.051 for comparison of pre/post treatment values in bumetanide alone group. Refer to table 2 for the overall effects of insulin/saline and bumetanide. Note: Data were analyzed with log transformed values.

C. NAA:Creatine ratios in DKA rats pre- and post- saline/insulin infusion with and without bumetanide. NAA to creatine ratios were measured by 1H MRS as described in RESEARCH DESIGN AND METHODS. Bumetanide alone and saline/insulin with and without bumetanide treatments started immediately after baseline measurements (0 hr). All values are arithmetic means with 95% confidence intervals, n= 13, 9, 9, and 10 for no treatment, bumetanide alone, saline/insulin, saline/insulin with bumetanide treatment groups respectively. * Within group comparisons: p =0.03 for comparison of pre/post treatment values in insulin/fluid group. Refer to table 2 for the overall effects of insulin/saline and bumetanide.
D. Intracellular pH in DKA rats pre- and post- saline/insulin infusion with and without bumetanide. Intracellular pH values were determined by 31P MRS as described in RESEARCH DESIGN AND METHODS. Bumetanide alone and saline/insulin with and without bumetanide treatments started immediately after baseline measurements (0 hr). All values are arithmetic means with 95% confidence intervals, n= 12, 10, 10, and 12 for no treatment, bumetanide alone, saline/insulin, saline/insulin with bumetanide treatment groups, respectively. * Within group comparisons: p < 0.05 for comparison of pre/post treatment values in insulin/fluid group and insulin/fluid + bumetanide group. Refer to table 2 for the overall effects of insulin/saline and bumetanide.

Figure 1
Figure 2

Figure 3
Figure 4