β -Hydroxybutyrate Detection with Proton MR Spectroscopy in Children with Drug-Resistant Epilepsy on the Ketogenic Diet

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ABSTRACT

BACKGROUND AND PURPOSE: The ketogenic diet, including both classic and modified forms, is an alternative to antiepileptic medications used in the treatment of drug-resistant epilepsy. We sought to evaluate the utility of proton MR spectroscopy for the detection of β -hydroxybutyrate in a cohort of children with epilepsy treated with the ketogenic diet and to correlate brain parenchymal metabolite ratios obtained from spectroscopy with β -hydroxybutyrate serum concentrations.

MATERIALS AND METHODS: Twenty-three spectroscopic datasets acquired at a TE of 288 ms in children on the ketogenic diet were analyzed with LCModel using a modified basis set that included a simulated β -hydroxybutyrate resonance. Brain parenchymal metabolite ratios were calculated. Metabolite ratios were compared with serum β -hydroxybutyrate concentrations, and partial correlation coefficients were calculated using patient age as a covariate.

RESULTS: β -hydroxybutyrate blood levels were highly correlated to brain β -hydroxybutyrate levels, referenced as either choline, creatine, or *N*-acetylaspartate. They were inversely but more weakly associated with *N*-acetylaspartate, regardless of the ratio denominator. No strong concordance with lactate was demonstrated.

CONCLUSIONS: Clinical MR spectroscopy in pediatric patients on the ketogenic diet demonstrated measurable β -hydroxybutyrate, with a strong correlation to β -hydroxybutyrate blood levels. These findings may serve as an effective tool for noninvasive monitoring of ketosis in this population. An inverse correlation between serum β -hydroxybutyrate levels and brain tissue *N*-acetylaspartate suggests that altered amino acid handling contributes to the antiepileptogenic effect of the ketogenic diet.

ABBREVIATIONS: BHB = β -hydroxybutyrate; GABA = gamma-aminobutyric acid; KD = ketogenic diet; MAD = modified Atkins diet

The ketogenic diet (KD) is a high-fat, low-carbohydrate, low-protein diet developed in the 1920s as a treatment for epilepsy, in response to the observation that fasting decreased the incidence and severity of seizures.¹ Use of the KD was largely supplanted by the emergence of multiple highly effective antiepileptic medications throughout the subsequent decades.² More recently, there has been a resurgence of interest in the KD, especially for treatment of drug-resistant epilepsy in children. Clinical research has demonstrated that modified forms of the diet, including the modified Atkins diet (MAD)

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and modified ratio formulations of the KD, are also effective in controlling seizures.³

The KD restricts intake of carbohydrates and protein, driving the production of ketone bodies from fat by the liver. This alteration in energetic substrates results in increased serum levels of β -hydroxybutyrate (BHB) and, to a lesser degree, acetone and acetoacetate. Brain parenchymal concentrations of ketone bodies have been shown to increase as a function of serum concentrations,⁴ modulated by the duration of hyperketonemia. Prolonged ketosis results in upregulation of the monocarboxylic acid transporter family of proteins that governs the active transport of ketone bodies from serum, resulting in increased steady-state brain tissue concentrations.⁵ Ketone bodies serve as the primary oxidative substrate for neurons during ketosis.⁶

The mechanisms by which the KD exerts its antiepileptogenic effect are not definitely known. Theories that have been advanced include the following: an anticonvulsant effect of ketone bodies; a shift in brain amino acid handling favoring increased synthesis of the inhibitory neurotransmitter gamma-

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FIG 1. Voxel-of-interest (inner 8×8 matrix) and masked region (7×7 matrix) that retained spectra based on SNR, NAA, and adequate Cramér-Rao bounds. Across the brain (spectra labeled *A*–*F*, corresponding to labeled voxels in image at the upper right), 2 peaks comprising lactate and BHB are shown at 1.33 and 1.2 ppm, respectively, with lactate more prominent in voxels containing CSF in this patient with mitochondrial cytopathy. At the lower right, the composite spectrum from the masked region is shown, with the LCModel fit overlaid.

aminobutyric acid (GABA); a direct consequence of glucose restriction; or an effect mediated by altered polyunsaturated fatty acid metabolism.⁷ Whatever the mechanism, steady-state blood levels of BHB have been shown to correlate with the degree of seizure control.⁸

Proton MR spectroscopy (¹H-MR spectroscopy) is a clinically available imaging technique by which tissue concentrations of metabolites can be noninvasively evaluated. Anecdotally, we have observed a resonance near 1.2 ppm on long-TE (288 ms) spectra in patients treated with the KD, tentatively attributed to BHB. There have been rare anecdotal reports of BHB detection by ¹H-MR spectroscopy in the brain parenchyma of children treated with the KD,^{9,10} with only 4 patients reported to date. We sought to evaluate the utility of ¹H-MR spectroscopy for the routine detection of BHB in a cohort of children treated with the KD for drug-resistant epilepsy and to correlate ¹H-MR spectroscopy metabolite ratios with BHB serum concentrations.

MATERIALS AND METHODS

Following institutional review board approval, we retrospectively identified 247 patients treated with the classic KD, a modifiedratio KD, or the MAD at our institution between 2006 and 2014. The classic KD used a ratio of 4:1, indicating a target dietary intake of 4 g of fat per gram of combined carbohydrate and protein. Patients were shifted to a modified-ratio formulation of the KD depending on clinical course and seizure response. The MAD, which limits carbohydrate intake to a maximum of 20 g per day, was used in some patients to increase tolerability and compliance.

Of the 247 patients, 38 underwent \geq 1 ¹H-MR spectroscopy during their treatment, for a total of 49 MR spectroscopy datasets. Clinical data on patient age, epilepsy and seizure type, diet type, medications, serum BHB concentration, and lag in days between

blood testing and MR spectroscopy were recorded after retrospective chart review.

Data were further reduced to 23 spectra from 18 total patients in whom the lag between blood BHB sampling and MR spectroscopy was <30 days. Medication histories were screened to exclude medications—including phenytoin and phenobarbital, known to contain propylene glycol or 1,2-propanediol—that generate a doublet resonance on ¹H-MR spectroscopy at 1.1 ppm, which could potentially confound results.

MR spectroscopy data were acquired axially for all subjects on a 3T Trio magnet (Siemens, Erlangen, Germany) with a multivoxel point-resolved spectroscopy sequence acquisition centered on the basal ganglia (16×16 matrix with a VOI of 8×8 , TE = 288, TR = 1700, slice thickness = 15 mm, NEX = 2, time of acquisition = 5 minutes 16 seconds). Slab location was assigned in a standardized fashion per our clinical MR spectroscopy protocol, with generally equivalent anatomic sampling across patients (see Fig 1 for a sample slab position). Twenty-two of 23 MR spectroscopy examinations were performed with sedation, using either nitrous oxide/sevoflurane or propofol for induction and sevoflurane or propofol for maintenance.

Data were processed using LCModel (Version 6.3; http:// www.lcmodel.com/) to generate phase and frequency corrected spectra.¹¹ Spectral arrays were masked to exclude the outer row within the VOI, as well as any individual spectrum having *N*-acetylaspartate signal Cramér-Rao lower bounds of \geq 15, a typical criterion for spectral quality. The remaining voxels were summed to create a single composite spectrum for each subject.

The composite spectra were reprocessed in LCModel with the addition of a simulated doublet centered at 1.2 ppm, correspond-

ing to the dominant BHB resonance produced by its methyl group $(-CH_3)$ (see Fig 1 for a sample fitted summed spectrum). The additional low-amplitude multiplet resonances of BHB at 2.3–2.4 ppm and 4.1 ppm were not modeled.

Results were expressed as a ratio of tissue BHB concentration to commonly assessed brain metabolite concentrations, including NAA, creatine/phosphocreatine (Cr), choline-containing compounds (Cho), and lactate. Derived metabolite ratios were correlated to serum BHB concentrations, corrected for age in months at the time of MR spectroscopy acquisition.

Analyses were also performed on 3 exploratory datasets (one with a restricted sampling lag threshold of 14 days [n = 13], a second with an extended sampling lag threshold of 120 days [n = 31], and a third including all examinations regardless of sampling lag [n = 49]) to estimate the effect of sampling lag on the strength of correlations.

Statistical Analysis

Descriptive analyses were performed for all variables, with age at MR spectroscopy in months covaried for parametric Spearman correlations. Significance was set at P < .05, with the primary time point of interest being a lag of \leq 30 days. Other lag intervals were analyzed to explore the strength of the correlation, with reported values not corrected for multiple comparisons and/or autocorrelation owing to overlapping samples with decreasing lags. Analyses were performed in SPSS 19 (IBM, Armonk, New York).

RESULTS

Clinical data for the 18 included patients are summarized in Table 1. Nineteen datasets were obtained from patients on the KD, and 4 were from patients treated with the MAD. Mean blood serum BHB concentrations were 35.2 ± 25.9 mg/dL (range, 0.2-81.3 mg/dL). The mean lag time between blood sampling and MR spectroscopy was

Table 1: I	Patient clinical	l data includi	ng underlying	epilepsy	syndrome,	seizure typ	e, and
antiepile	ptic medicatio	ons					

Patient	Age (yr)	Epilepsy Syndrome ²¹⁻²³	Seizure Type	Medications
1	1	EE (Unclassified)	Т	LEV
2	2	Metabolic (Mito)	TS	ZNS/LTG/LEV
2	7	Metabolic (Mito)	TS/GTC	ZNS/LTG/LEV/RFM
2	8	Metabolic (Mito)	TS/MY/GTC	ZNS/LTG/LEV/RFM
3	9	MAE	MY/AT/GTC	VPA/CLB/RFM
4	7	Metabolic (Mito)	MY/RFL	TPM
4	10	Metabolic (Mito)	MY/RFL	TPM/ZNS
5	7	EE (MCD)	T/AT/TS	VGB/LTG
6	2	Structural (MCD)	Focal	ZNS/TPM
7	1	Structural (Aicardi)	ES	VGB
8	6	Genetic (SCN1A)	MY/AT	LEV/ETX/CLN
9	6	Metabolic (Mito)	MY/AT	LEV/LTG/ETX/CLN
10	6	MAE	T/MY/TS	LTG
11	0.5	Genetic (ATP1A3)	Focal/SE	LEV
11	1	Genetic (ATP1A3)	SUB/MY	LTG/CLB
12	6	EE (unclassified)	AT/T/GTC	CLB
13	1	EE (unclassified)	ES	VGB
14	2	Metabolic (Mito)	ES	TPM
15	4	EE (LGS)	AT/T/AA	LTG/LEV/ZNS
16	1	EE (Unclassified)	ES	ZNS/LEV
17	5	Genetic (DEAF 1)	T/AT/GTC	LEV
18	1	Metabolic (Mito)	ES	ZNS

Note:—"Epilepsy Syndrome": EE indicates epileptic encephalopathy; Mito, primary mitochondrial disease; MAE, myoclonic astatic epilepsy; MCD, malformation of cortical development; LGS, Lennox-Gastaut Syndrome; "Seizure Type": T, tonic; TS, tonic spasms; GTC, generalized tonic-clonic; MY, myoclonic; AT, atonic; RFL, reflexive seizure; ES, epileptic spasms; SE, status epilepticus; AA, atypical absence; SUB, subclinical seizures; "Medications": LEV, levetiracetam; ZNS, zonisamide; LTG, lamotrigine; RFM, rufinamide; CLB, clobazam; TPM, topiramate; VGB, vigabatrin; ETX, ethosuximide; CLN, clonazepam; VPA, valproic acid. 11.1 ± 9.7 days (range, 0–28 days).

Included spectra (n = 23) were acquired at a mean patient age of 54.0 ± 37.3 months (range, 1–129 months). In terms of spectral fit parameters, the mean SNR was 25.7 ± 6.3 (range, 12– 36), and the mean full width at half maximum was 0.04 ± 0.01 (range, 0.02– 0.06), with mean NAA Cramér-Rao lower bounds of 1.9 ± 0.6 (range, 1–3).

Using age as a covariate, we calculated correlation coefficients between blood levels and brain metabolites. They are reported in Table 2. BHB blood levels were strongly correlated to brain BHB levels (calculated correlation coefficient for blood BHB-BHB/NAA was R = 0.81 with P < .001; calculated correlation coefficient for blood BHB-BHB/Cr was R = 0.80 with P < .001) and inversely but more weakly related to NAA levels, regardless of the ratio denominator (calculated correlation coefficient for blood BHB–NAA/Cr was R =-0.55 with P < .01; calculated correlation coefficient for blood BHB-NAA-Cho was R = -.44 with P < .04). No

Table 2: Partial correlations of blood BHB concentration (mg/dL) and brain metabolite ratios derived f	rom ¹ H-MRS, using patient age ir
months as a covariate	

Group	BHB/Cr	BHB/Cho	BHB/NAA	BHB/Lac	Cho/Cr	NAA/Cr	NAA/Cho	Lac/NAA	Lac/Cr
Lag $\leq 14 d (n = 13)$									
R	.871	.898	.886	.507	287	645	540	.376	.253
Significance	.000	.000	.000	.092	.367	.024	.070	.228	.428
Lag ≤30 d (n = 23)									
R	.803	.814	.807	.538	165	553	443	.421	.334
Significance	.000	.000	.000	.010	.463	.008	.039	.051	.128
Lag ≤120 d (n = 31)									
R	.790	.802	.781	.484	312	658	515	.523	.423
Significance	.000	.000	.000	.008	.099	.000	.004	.004	.022
All ($N = 49$)									
R	.579	.588	.584	.393	170	387	312	.290	.162
Significance	.000	.000	.000	.006	.254	.007	.033	.048	.278

Note:-Lac indicates lactate.



FIG 2. Scatterplot of blood serum BHB concentrations (milligrams/deciliter) versus brain BHB/ NAA and BHB/Cr ratios demonstrates the strong associations between measures.

strong concordance with lactate was demonstrated. BHB/NAA and BHB/Cr plots versus blood BHB concentration for individual examinations are shown in Fig 2.

Results were calculated for the exploratory datasets, also reported in Table 2. For the dataset with a threshold lag of 14 days, the calculated correlation coefficient for blood BHB–BHB/NAA was R = 0.89 with P < .001. For the dataset with a threshold lag of 120 days, the calculated correlation coefficient for blood BHB–BHB/NAA was R = 0.781 with P < .001. When the entire cohort was included, blood BHB–BHB/NAA demonstrated an R = 0.584 with P < .001.

DISCUSSION

Prior reports have demonstrated ketone bodies by ¹H-MR spectroscopy in small numbers of mainly adult patients during prolonged fasting,^{4,12} diabetes,^{13,14} and BHB supplementation¹⁵ and in patients treated with the KD.^{9,10,12,16,17} To date, elevated BHB levels have been anecdotally reported in a only 4 children using ¹H-MR spectroscopy.^{9,10} In our cohort, we demonstrated routine detection of BHB in pediatric patients treated with the KD or a modified KD using long-TE multivoxel ¹H-MR spectroscopy performed at 3T.

High concordance between steady-state brain parenchymal BHB ratios and serum BHB levels was observed using a lag threshold between blood sampling and MR spectroscopy of 30 days (Fig 2). The strength of this correlation progressively increased as the lag threshold between blood BHB sampling and MR spectroscopy decreased, suggesting that the concordance reflects a true physiologic relationship. Results persisted when the lag threshold increased to 120 days, with a similar but weaker relationship demonstrated when the entire cohort was included, implying that BHB ratios are relatively stable with time in patients on the KD. These findings suggest that ¹H-MR spectroscopy might be an effective tool for noninvasive monitoring of ketosis in children on the KD, particularly when performed in conjunction with routine follow-up MR imaging.

In our cohort, seizure response dictated varying ratios of fat to carbohydrates and protein on the KD. Some patients required the MAD for compliance or tolerability concerns. The specific formulation of the diet did not appear to alter the correlation of brain BHB ratios to serum BHB levels, though we had a relative paucity of children on the MAD. That the specific diet type did not appear to alter the observed correlation suggests that brain BHB levels may be correlated with serum BHB concentrations independent of the dietary mechanism used to achieve ketosis, be it increased fat ratio or carbohydrate restriction. Further evaluation of a cohort with larger subgroups of diet type would be useful to confirm this observation.

We demonstrated an inverse relationship between blood levels of BHB and brain NAA ratios, independent of patient age. This is in contrast to a prior animal study in rats with diabetic ketoacidosis, which found no significant dif-

ference in NAA/Cr ratios between study and control animals using ¹H-MR spectroscopy performed at 7T with a TE of 132 ms.¹⁸ Our finding may reflect in vivo changes in amino acid handling that occur with prolonged ketosis in the KD.7,19 Increased acetylcoenzyme A as a by-product of ketone body metabolism in the brain drives increased tricarboxylic acid cycling, resulting in increased conversion of cytosolic aspartate to glutamate. Aspartate serves as the precursor molecule for NAA, while glutamate is converted to the inhibitory neurotransmitter GABA. Thus, the observed decrease in NAA may serve as a marker for an increase in GABA production, one of the proposed mechanisms of seizure control in the KD. Further studies using an MR spectroscopy acquisition specifically tailored to interrogate brain GABA concentrations would be instructive, with specific evaluation of the relationship between serum BHB concentrations and both brain GABA ratios and the degree of seizure control.

Consistent with past studies using infusion samples, lactate ratios did not show significant correlation to BHB serum levels.⁴ This finding has the caveat that anything other than elevated lactate exists near the noise floor, which can preclude accurate association assessment. In 5/18 patients, mitochondrial cytopathy was the etiology of their drug-resistant epilepsy, contributing further uncertainty to ascribing lactate levels to dietary effects.

Of note, the ¹H-MR spectroscopy spectra included in our analyses were acquired with a TE of 288 ms, optimized for the detection of lactate. J-coupling for BHB is 6.3 Hz, suggesting that TEs of 160 ms (1/J) or 320 ms (2/J) would be more appropriate for optimization of BHB detection.¹⁰ Furthermore, this TE selection limits the evaluation of additional ketosis-related metabolites, including acetone and acetoacetate. Both metabolites have generally lower brain concentrations⁴ and shorter T2 relaxation times¹⁰ than BHB. Future work including short-echo data would be helpful to assess these metabolites.

An additional limitation of our methodology was inclusion of gray and white matter and CSF within the VOI. Tissue segmentation was not attempted, and the differential contributions of different tissue types to the composite spectra were not quantified. The presence of visually identifiable BHB peaks in parenchymal voxels suggests that the ketogenic diet metabolites are broadly distributed across tissue types, consistent with the work of Pan et al.⁴ Future work would benefit from specific analysis of the contributions of different tissue types to the overall summed spectrum.

Our described methodology, preprocessing and summing a multivoxel dataset to increase the signal-to-noise ratio, allowed routine detection of BHB in brain parenchyma. Further work using same-day blood sampling; integrated tissue-type segmentation using an approach like localization by adiabatic selective refocusing,²⁰ which minimizes chemical shift artifacts; absolute metabolite quantification; as well as evaluation of longitudinal changes will be helpful to extend and refine these results.

CONCLUSIONS

We demonstrated routine detection of a brain parenchymal BHB using conventional ¹H-MR spectroscopy in the brains of pediatric patients with epilepsy treated with the KD. We further demonstrated a strong correlation between brain tissue BHB ratios and serum BHB concentrations, apparently independent of specific diet formulation. These findings suggest that ¹H-MR spectroscopy may be a useful noninvasive method of interrogating brain tissue BHB levels in patients on the KD or a modified KD protocol. The observation of an inverse correlation between brain tissue NAA levels and serum BHB levels may reflect altered amino acid handling induced by the KD, consistent with one of the proposed antiepileptic mechanisms of the KD.

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