KETONE BODY THERAPY:

From the Ketogenic Diet to the Oral Administration of Ketone Ester

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Conflict-of-Interest statements:

Dr. Hashim is the recipient of a patent involving the triglyceride of \(\beta\)-hydroxybutyrate as a food supplement for use in disorders characterized by impairment of glucose utilization by the brain.

Dr. VanItallie is a minority shareholder in a company that markets a product that yields medium-chain fatty acids.
Abstract

Ketone bodies (KBs), acetoacetate and β-hydroxybutyrate, were considered harmful metabolic by-products when discovered in the mid-19th century in urine of patients with diabetic ketoacidosis. It took physicians many years to realize KBs are normal metabolites synthesized by the liver and exported into the systemic circulation to serve as an energy source for most extrahepatic tissues. Studies have shown that the brain (which normally uses glucose for energy) can readily utilize KBs as an alternative fuel. Even when there is diminished glucose utilization in cognition-critical brain areas, as may occur early in Alzheimer’s disease, there is preliminary evidence that these same areas remain capable of metabolizing KBs. Because the ketogenic diet (KD) is difficult to prepare and follow, and effectiveness of KB treatment in certain patients may be enhanced by raising plasma KB levels to ≥2 mM, KB esters, such as 1,3-butanediol monoester of β-hydroxybutyrate and glyceryl-tris-3-hydroxybutyrate, have been devised. When administered orally in controlled dosages, these esters can produce plasma KB levels comparable to those achieved by the most rigorous KD, thus providing a safe, convenient, and versatile new approach to the study and potential treatment of a variety of diseases, including epilepsy, Alzheimer’s, and Parkinson’s.
Ketone bodies: ugly duckling or swan?

Acetoacetate (AcAc) and β-hydroxybutyrate (βHB) are collectively known as ketone bodies (KBs). KBs have been dubbed “metabolism’s ugly duckling” because, in the mid-19th century, they were first discovered in large quantities in the urine of patients succumbing to diabetic ketoacidosis. Thus, it is not surprising that physicians of the era considered KBs to be toxic by-products of impaired carbohydrate metabolism. It took almost half a century for medical scientists to understand that KBs are normal metabolites manufactured by the liver in increasing amounts when dietary sources of carbohydrate and glucogenic amino acids are in short supply (1). Unfortunately, some physicians still fail to distinguish between the safe “physiological” hyperketonemia that occurs in healthy individuals during fasting or adherence to a ketogenic diet, and the pathologic, out-of-control hyperketonemia associated with insulin-deficient diabetes.

When Owen et al (2) reported that, during a prolonged fast, KBs can provide 60% or more of the brain’s daily energy requirement (thereby sparing ~80g/d of glucose that otherwise would have been derived largely from breakdown of the body’s limited protein stores), it was finally acknowledged that—as in Hans Christian Andersen’s 1843 fairy tale—the creature first thought to be an ugly duckling was turning out to be an emerging swan. It became evident that the

Although some consider acetone to be an authentic member of the KB family, its importance for purposes of this review is minimal.
ketogenic response to starvation is an indispensable metabolic adaptation designed by nature to preserve strength and prolong life during times when food is unavailable (3).

It is now known that (in nondiabetic individuals), owing to the blood’s efficient buffering capacity, plasma KB levels can increase to 6-8 mM during a prolonged fast without giving rise to clinically hazardous acidosis (4).

**Physiology of ketogenesis**

Four physiologic facts lie at the root of the ketogenic adaptation: [i] the body’s small reserve supply of preformed carbohydrate (largely as glycogen); [ii] the body’s limited protein stores; [iii] the relative plenitude in human adipose tissue of stored triglyceride (triacylglycerol [TAG]); and [iv] the inability of long-chain fatty acids (≥ C12) to cross the blood-brain barrier (BBB). Given these considerations, the evolutionary advantage of having a TAG-derived metabolite capable of crossing the BBB and nourishing the brain during times when food is unavailable is self-evident.

In a 70kg man of normal body composition, the amount of fuel reserves in the form of TAG is approximately 12kg. Muscle protein is about 6kg, while the carbohydrate reserves (glycogen) in liver and muscle are ~100g and ~400g respectively (5). Glucose is the brain’s usual fuel source. After an overnight fast, owing to increased glucagon secretion and diminished insulin release, amplified mobilization of free fatty acids (FFA) from adipose tissue is associated with their increased utilization by muscle and enhanced hepatic ketogenesis. However, at this early stage of carbohydrate privation, while plasma KBs are still low, the brain remains heavily dependent on glucose.
During total caloric starvation, the only source of new glucose is that synthesized from the glycerol released from adipose tissue together with FFA, and from glucogenic amino acids derived from breakdown of stored protein. With continued starvation, gluconeogenesis is curtailed, and the liver shifts acetyl-CoA to KB synthesis (see below). During glucose scarcity, the astrocytes also may contribute to KB formation. Astrocytes in culture have been shown to produce KBs from fatty acids (6) and from leucine (7). The mechanism by which the astrocytes synthesize KBs is very similar to that of cultured hepatocytes. In a review of KB synthesis in the brain, it was suggested that production of KBs by astrocytes contributes to the survival of neurons subjected to hypoxia (8). Most studies of astrocyte ketogenesis come from cell culture experiments, and the extent of KB formation by astrocytes in vivo remains to be determined. Nevertheless, the major determinants of cerebral KB metabolism are the prevailing plasma KB concentrations and availability of suitable monocarboxylic acid transporter (MCT) isoforms (9).

Studies based on positron emission tomography (PET) imaging in rats found a seven- to eight-fold enhancement of brain uptake of ketones during a ketogenic diet or fasting (10).

**The brain's high energy requirement**

Usually, the brain obtains its fuel mainly from glucose/pyruvate-derived substrate, which is almost completely oxidized in the mitochondria, generating CO₂, water and high energy phosphate bonds (principally ATP). The brain is responsible for ~20% of the body’s total resting energy expenditure; yet, it represents only about 2% of adult body weight. The brain metabolizes ~100-120 grams of glucose per day under conditions of normal glucose availability. Studies have shown that most of the glucose-derived energy entering the brain is used to
maintain pre- and post-synaptic ion gradients required for neurotransmission, and for maintenance of the resting potential of neurons (11).

When glucose is in short supply, KBs serve as the brain’s principal alternative fuel. However, the brain can only use them in quantity if their levels in the plasma substantially exceed default concentrations (≤ 0.2 mM). In the postabsorptive state, for example in the morning upon awakening, there exists a mild degree of transient hyperketonemia, with plasma ketone levels of 0.1-0.3 mM. These concentrations drop precipitously after ingestion of a mixed meal, only to rise again in the next postabsorptive state. In diabetic ketoacidosis, plasma concentration of KB can exceed 25 mM (12).

The liver forms KB but lacks the enzymes to use them as energy substrates. Transfer of AcAc and βHB across cell membranes (including those of neurons) is enabled by monocarboxylate transporters (MCTs). In the mitochondrial matrix, βHB is converted to AcAc by βHB dehydrogenase, and the resulting AcAc, together with any AcAc that has entered the matrix as such, are then transformed to AcAc-CoA by oxoacid-CoA transferase. AcAc-CoA is then converted to acetyl-CoA by acetoacetyl-CoA thiolase, with the resulting acetyl CoA units entering the Krebs (tricarboxylic acid [TCA]) cycle. In the cycle, they undergo oxidative degradation, with reduction of the electron carriers NAD+ (nicotinamide adenine dinucleotide) and FAD (flavine adenine dinucleotide) to NADH and FADH2. NADH and FADH2 donate electrons to the protein Complexes I and II of the electron transport chain (ETC). Energy derived from the transfer of electrons along the ETC to oxygen (O2) is used by the electron transport system to pump protons (H+) into the mitochondrial intermembrane space, thereby generating a gradient across the inner mitochondrial membrane (proton motive force [pmf])
that provides energy to regenerate ATP from ADP and P\. The role of mitochondrial dysfunction in neuronal degeneration has been reviewed by Schon and Manfredi (13).

**KB: Source of energy for brain, heart and muscle**

There is evidence that the whole brain uses energy from KBs as a function of the blood (plasma) concentration, as shown in Table 1.

<table>
<thead>
<tr>
<th>Plasma KB Concentration (mM)</th>
<th>Proportion of Brain Energy Metabolism (%)</th>
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<tbody>
<tr>
<td>0.3-0.5 mM (12-24 hr fast)</td>
<td>3-5%</td>
</tr>
<tr>
<td>1.5 mM (2-3-day fast)</td>
<td>18%</td>
</tr>
<tr>
<td>5 mM (8-day fast)</td>
<td>60%</td>
</tr>
<tr>
<td>7 mM (≥ 20-day fast)</td>
<td>&gt;60%</td>
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In the human brain, the transport system for KBs (unlike that for glucose) remains relatively intact with advancing age. Certain monocarboxylic acid transporter (MCT) isoforms are well expressed in neurons (MCT2), astrocytes (MCT4) and brain capillaries (MCT1). When glucose utilization is impaired in neurodegenerative diseases, transport of KBs into the brain appears to be less affected and their utilization for energy by the brain mitochondria is not impeded by such factors as local insulin resistance that, by interfering with the neuronal fuel supply, may contribute to the progressive nerve cell damage observed in AD (1,5,14-16).

The central actions of βHB have been reviewed by Laeger et al. (17). These include its sources, its metabolism during starvation and cellular signaling, its effects on food intake, its role in ATP production, energy metabolism and thermogenesis, its neuroprotective effects, and its influence on pituitary hormone release. The authors cite studies indicating that
all the enzymes needed for KB oxidation, such as β-hydroxybutyrate dehydrogenase, 3-ketoacid CoA transferase, and acetyl-CoA thiolase, are present in the brain.

**Regulation of plasma KB concentrations**

In the first few days of a prolonged fast, while the body’s carbohydrate stores are being rapidly depleted, the liver accelerates its manufacture of KBs from FFA released in increasing amounts from adipocytes. In the absence of dietary carbohydrate, and as depletion of the body’s stored glycogen continues, the liver also increases its production of new glucose. Krebs cycle intermediates—notably oxaloacetate—are diverted to gluconeogenesis, which entails conversion in the liver of pyruvate derived from the carbon skeletons of glucogenic amino acids, to glucose. Glycerol released from adipocytes along with FFA is also converted to glucose in the liver.

At the same time, insulin production tends to wane as glucose availability diminishes. Reduced concentrations of circulating insulin result in attenuation of insulin’s inhibiting effect on FFA/glycerol release. At this point, because much of the limited supply of oxaloacetate is being used for gluconeogenesis, metabolism in the Krebs cycle of fatty acid-derived acetyl-CoA is slowed and the resulting accumulation of the 2-carbon units is then redirected to production of KBs for export into the systemic circulation.

To promote regeneration of oxaloacetate and thereby allow restoration of earlier levels of gluconeogenesis, the intrahepatic accumulation of acetyl Co-A apparently stimulates pyruvate carboxylase activity, resulting in conversion of more pyruvate to oxaloacetate—a key intermediate in both the Krebs cycle and the gluconeogenic process.
As the liver increases its KB output, the plasma total KB concentration rises gradually to 5-7 mM, or even slightly higher, depending in considerable part on the duration of the fast. In individuals whose islet beta-cells are intact and functional, an elevated plasma ketone concentration can directly stimulate the beta-cells to increase insulin secretion. However, it should be kept in mind that much of the evidence for hyperketonemia-induced enhancement of insulin release was obtained from dog studies in which infusions of KBs produced plasma KB concentrations of ~3 mM (21, 22). The relatively brief time frame in which the infusion experiments took place is very different from the slow rate at which metabolic changes occur during the development of fasting-induced hyperketonemia. During a prolonged fast, blood glucose plateaus at a lower-than-usual level, with an associated reduction in insulin release.

Nevertheless, a KB-generated negative feedback effect could explain the fall in arterial glucose concentration, the gradual increase—followed by a leveling off—of plasma FFA levels, and the stabilization of plasma KB observed over time in fasting individuals. Reducing the quantity of FFA released from adipocytes decreases FFA traffic through the liver. Reduction in rate of FFA entry into the liver would be expected to cause a decrease in hepatic KB formation— in effect, closing the negative feedback loop that prevents plasma KBs from rising to unsafe levels during starvation. Moreover, hyperketonemia per se may limit fatty acid release from adipose tissue (23). However, the presence of insulin is believed necessary for this effect (3).

**Therapeutic uses of ketone bodies**

Traditionally, physicians have been taught to fear ketosis because the marked hyperketonemia that results from insulin deficiency can cause severe acidosis and death in
individuals with type 1 diabetes. Thus, in their description of the potential therapeutic uses of KBs, Veech et al. (14, 18) emphasize that, in marked contrast to the clinical picture in diabetic ketoacidosis, mild to moderate hyperketonemia (up to ~8mM) can materially prolong survival during periods of caloric starvation. As glucose availability diminishes, KBs manufactured in the liver from fatty acids mobilized from adipose tissue, become major sources of energy for muscle, heart and brain (18).

Veech et al. (14) described “clinical maneuvers” for readily increasing blood levels of βHB to 2-8mM—concentrations similar to those produced by starvation or various ketogenic diets. To achieve this objective, they recommended use of small synthetic, digestible KB polymers (including dimers), or esters of βHB administered orally at 100-150g/d in divided doses. The goals were to [i] obtain relatively high plasma KB levels which might enhance the clinical effectiveness of KB therapy in some cases; and [ii] provide a more efficient source of energy per unit oxygen consumed for the treatment of certain types of heart failure, and neurodegenerative diseases characterized by focal brain hypometabolism, such as Parkinson’s and Alzheimer’s. The authors also suggested that the ability of βHB to reduce nicotinamide adenine dinucleotide phosphate (NADP+) might be important in decreasing the oxidative damage associated with various kinds of metabolic stress (14).

**KBs are a “high-octane” fuel for the body**

The effect of adding insulin or KBs (4mM) to a buffer containing 10 mM of glucose in a perfused rat heart preparation was studied by Kashiwaya et al (24) and by Sato et al (25). The addition of either insulin or ketones increased the efficiency of the working heart (hydraulic
work/energy from O\textsubscript{2} consumed) by 25%. The addition of both insulin and KBs in combination increased heart efficiency by 36%. The authors concluded that moderate hyperketonemia (~4 mM) may compensate for defects in mitochondrial transduction associated with insulin deficiency, local glucoprivation, or mitochondrial senescence. Later work by the same group showed that moderate hyperketonemia following ingestion of the 1,3-butanediol monoester of βHB (ketone monoester [KME]) significantly improves endurance of rats on a treadmill and also the physical performance of competing University athletes (26).

**Alzheimer's disease**

**Possible triggering role of mitochondrial dysfunction**

Mitochondrial dysfunction has been implicated in the etiology of mild cognitive impairment (MCI) and Alzheimer's disease (27). Such dysfunction, which may be related to diminished energy production from mitochondrial glucose/pyruvate oxidation, potentiates the pathologic intraneuronal (and later extracellular) deposition of amyloid-β and hyperphosphorylated tau. The mechanism for the mitochondrial dysfunction is not certain. However, several possible explanations have been proposed and are discussed in recent reviews (28, 29). Manifestations of impaired mitochondrial function include a decrease in oxidative phosphorylation and ATP synthesis, increased superoxide anion production, evidence of oxidative damage, inhibition of mitochondrial pyruvate dehydrogenase complex (PDH) activity, and functional impairment in the mitochondrial electron transport chain (ETC), particularly involving cytochrome c oxidase. Magnetic resonance spectroscopy (MRS) has been used to access neuronal mitochondrial metabolism in healthy elderly and young volunteers (27). MRS studies of these two groups revealed that, in the aging subjects, there was a reduction in neuronal and glial mitochondrial
metabolism compared with the healthy young subjects. In a mouse model of Alzheimer’s disease, Chou et al (30) found that early dysregulation of the mitochondrial proteome precedes the development of plaque and tangle pathologies. A number of mitochondrial proteins were down-regulated in the cerebral cortices of these mice, notably in Complexes I and IV of the oxidative phosphorylation system. Other studies have provided strong evidence that the impaired glucose metabolism in certain parts of the brain, which is characteristic of AD, is related to mitochondrial dysfunction (31-37). In AD, changes in glucose metabolism in cognition-associated parts of the brain have been detected by PET imaging with 2-[\(^{18}\)F]fluoro-2-deoxyglucose (FDG), decades before the appearance of typical Alzheimer’s dementia (38). Four apparently normal individuals with FDG-PET evidence of reduced glucose utilization in cognition-related brain sites were followed for 9-19 years to the onset of clinical symptoms of dementia, and subsequently to post-mortem confirmation of the diagnosis of AD.

Factors impeding glucose utilization by the brain may contribute to, or precipitate, AD neuropathology. This possibility is strengthened by evidence that diminished glucose utilization can be present well in advance of measurable cognitive decline (29).

Studies have shown that certain glucose transporters in the brain (GLUT 1 and GLUT 2) may be diminished significantly in the Alzheimer brain (34). In addition, there is evidence that the concentration of GLUT 3, the principal neuronal glucose transporter, is diminished in the brains of Alzheimer patients (39). A decrease in glucose transporters also correlates with abnormal hyperphosphorylation of tau in Alzheimer’s disease (40). Such GLUT deficiencies presumably contribute to the impaired glucose metabolism implicated in neuronal degeneration.
There is preliminary evidence that, unlike glucose, transport and metabolism of KBs are not diminished in the AD brain (41,42). This finding underlines the importance of developing a safe, simple, and reliable way to provide the brain with KBs as an alternative fuel to glucose. The subject of brain fuel metabolism in aging and AD has been extensively reviewed by Cunnane et al. (41). In a more recent communication, Castellano et al. reported that, at the same time a diminished brain glucose utilization in AD could be demonstrated, ketone uptake was unchanged (42).

In recent years, extensive evidence has accumulated suggesting that regional hypometabolism within the brain may be a root cause of cognitive decline in sporadic AD (15). For example, carriers of one copy of the APOE-ɛ4 allele (a situation which enhances risk of developing AD), exhibit abnormally low rates of glucose metabolism bilaterally in the posterior cingulate, parietal, temporal, and prefrontal cortex (15). Under normal conditions, the energy used by the adult human brain is derived almost exclusively from glucose (42,43). In individuals with an increased risk of developing AD, glucose hypometabolism (manifested by a reduced cerebral metabolic rate for glucose [CMRglu]) may occur in cognition-critical parts of the brain decades before symptoms of dementia become manifest, and may precede intra- and extra-neuronal deposition of abnormal proteins. These findings suggest that neuronal energy privation may be an important contributor to the decline in cognitive performance exhibited by patients with early AD. Early support for the concept that the Alzheimer brain may retain its ability to use ketone bodies for energy even when glucose utilization is impaired, was obtained by feeding a mildly ketogenic (0.5-0.8mM) MCTG (tricaprylin) to AD patients. Even at such relatively low plasma KB concentrations, a modest rise in cognitive performance occurred
transiently in a subset of the Alzheimer cohort under examination. Yet, despite the unspectacular nature of the improvement that occurred, the studies reviewed were well designed and the cognitive improvement measured following MCTG ingestion was statistically significant (15).

In a mouse model of AD, the feeding of a KME (comprised of D-β-hydroxybutyrate and R-1,3 butanediol) as 21.5% of dietary calories was associated with lessening in anxiety and improvement in performance on learning and memory tests. Moreover, the mice fed the KME exhibited reduced Aβ peptide deposition in the hippocampus and amygdala, and reduced levels of hyperphosphorylated tau deposits in the same areas and in the cortex (44).

**Histone acetylation and deacetylation**

During the past ten years, a number of studies have addressed the phenomenon of histone acetylation and deacetylation, and the role of these processes in cognitive impairment and Alzheimer’s disease. For example, degradation of histone acetylation is associated with age-dependent memory impairment in mice. In contrast, restoration of histone acetylation leads to recovery of cognitive performance (45). More recent studies suggest that there is an urgent need to develop additional selective histone deacetylase (HDAC) inhibitors (46).

Recently, βHB was found to inhibit histone deacetylases 1, 3 and 4 at concentrations of 5.3, 2.4 and 4.5mM, respectively. Thus, millimolar concentrations of βHB appeared to increase histone acetylation via inhibition of histone deacetylases. Moreover, the same study provided evidence that βHB exerts a suppressive effect on oxidative stress (19). Inhibition of histone deacetylase also was shown in mice that were protected from methyl tetrahydropyridine-induced dopaminergic damage by feeding a triglyceride of βHB (20).
The human and rodent genome encodes for eleven HDAC proteins that are divided into four classes (HDAC I—IV). There is evidence that inhibition of HDACs 1–3 (Class I) reverses memory dysfunction in a mouse model of AD (47,48). Agents reported to inhibit HDAC include sodium butyrate, trichostatin A, suberoylanilide hydroxamic acid, and sodium phenylbutyrate. βHB also qualifies as an HDAC inhibitor (19,20). Most HDAC inhibitors influence the activities of the HDAC isoforms and classes nonselectively, and the term “pan-inhibitor” has been used to distinguish them from inhibitors that are class-selective or isoform-selective.

**Parkinson’s disease**

Although the pathogenesis of sporadic Parkinson’s disease remains unresolved, numerous studies suggest that—at the least—impairment of mitochondrial function involving the substantia nigra pars compacta (SNpc) plays an important contributory role (49-51). In 1983, Langston et al (52) reported that four persons developed marked parkinsonism after taking an illicit drug intravenously. The drug, 4-propyloxy-4-phenyl-N-methylpiperidine (MPPP), was a meperidine (Demerol®) analogue. A contaminant (and unwanted side product) resulting from apparently careless MPPP manufacture, 1-methyl-4-phenol-1, 2, 5, 6-tetrahydropyridine (MPTP) was found to be the likely culprit. It was the MPTP, after being oxidized in the brain to methylphenylpyridine (MPP+), that presumably caused selective destruction of dopaminergic neurons in the SNpc, giving rise to the human Parkinson’s disease-like syndrome described by Langston (52). Subsequently, MPTP has been used extensively to produce animal models of Parkinson’s disease.
Because a reduction in Complex I activity and impaired mitochondrial function had been reported in the brain and other tissues of patients with Parkinson’s disease (53,54), Tieu et al (55) reasoned that, inasmuch as the brain can utilize KB for energy via mitochondrial Complex II, KBs might protect against MPTP induction of parkinsonism in mice. Indeed, infusion of βHB into mice was found to confer protection against the dopaminergic neurodegeneration and motor deficits induced by MPTP.

In a tissue culture study of rat neurons, βHB protected hippocampal neurons from amyloid-beta (Aβ) 1-42 toxicity, and mesencephalic neurons from MPTP toxicity. These findings suggest that KBs have the potential of preventing, or possibly treating, both AD and PD (56). In a later recent study, Cheng et al (57) reported, in a rat model of PD, that a ketogenic diet protected dopaminergic neurons of the SNpc against the neurotoxicity of 6-hydroxydopamine (6-OHDA).

Recently, oral administration of glyceryl-tris-3-hydroxybutyrate (3GHB), the triglyceride of β-hydroxybutyrate, was found to exert an extended neuroprotective action against MPTP-induced neuronal destruction in the SNpc of mice. It was shown that 3GHB protects these neurons in a dose-dependent manner (20). The study’s authors suggested that this protection might be mediated via inhibition of HDAC. They concluded that this new KE (3GHB) represented a promising preventive and/or therapeutic strategy for a range of pathologic conditions affecting the brain, including PD and AD (20).

Another study in mice demonstrated that βHB inhibits HDAC in vitro and in vivo (19). The in vivo studies involved producing hyperketonemia (1.5 mM) in mice by means of a 24-hour fast, caloric restriction (0.6 mM), or infusion of buffered βHB (1.2 mM). A positive correlation was
observed between serum βHB level and histone acetylation, promoted by the KB-induced inhibition of HDAC. Treatment of mice with βHB also conferred significant protection against oxidative stress. Other studies indicate that KBs are protective against oxidative stress in neocortical neurons (58). They also help protect against the neuronal synaptic dysfunction induced by respiratory complex inhibitors (59).

In a 28-day outpatient study, the clinical effect of a “hyperketogenic” diet (hKD) (carbohydrate 2%; protein 8%; fat 90% of total calories) was studied in five patients with Parkinson’s disease (50). Unified Parkinson’s Disease Rating Scale (UPDRS) scores were determined at baseline and at weekly intervals. During adherence to the hKD, UPDRS scores improved in varying degrees in all five subjects.

Epilepsy

The anticonvulsant effect of fasting has been known for centuries (1). The ketogenic diet (KD) for the treatment of epilepsy, which mimics the metabolic effects of fasting, was first conceived in 1921 by Wilder (60). In terms of energy distribution, the original KD was 90% fat, ~8% protein and ~2% carbohydrate.

The very-high-fat, very-low carbohydrate, low-protein KD can produce rises in plasma LDL cholesterol, uric acid and free fatty acids. Occasionally, the KD may be associated with an increased incidence of nephrolithiasis and other serious complications (1). Some of these adverse effects can be prevented by guarding against chronic dehydration. Hyperlipidemia can be avoided in most cases by boosting the proportion in the diet of polyunsaturated (ω6 and ω3) and monounsaturated fatty acids (61). Also, incorporation of medium-chain triglyceride (MCTG)
into the KD may be helpful in formulating more tolerable ketogenic regimens for the long-term treatment of drug-resistant epilepsy (62-65).

KD’s have also been found therapeutically effective in approximately two-thirds of 104 patients with infantile spasm (66). In another study, at one to three months after the initiation of the KD in 26 patients with infantile spasm, 46% had a greater than 90% reduction in symptoms (67).

The mechanism responsible for the beneficial effect of the KD in epilepsy is not known. Several explanations have been proposed: [i] reduction in neural excitability; [ii] changes in energy availability; [iii] direct anti-convulsion action. Another mechanism for the anti-seizure action of the KD, suggested by Yudkoff et al (68), pertains to decreased availability of excitatory neurotransmitters (aspartate and glutamate), and increased availability of the inhibitory neurotransmitters (GABA), via stimulation of glutamic acid decarboxylase, which, in turn, increases GABA production from glutamate. Many studies have contributed in a variety of ways to our understanding of the beneficial effect of KDs on epilepsy (60,62,69-76). However, despite the abundance of hypotheses, the basis for the anti-seizure action of KBs remains unclear.

Because the new KEs (see Fig. 1) can elevate plasma KBs to concentrations comparable to those achieved during prolonged adherence to a KD, without concurrent need to change the composition of the habitual diet, it should now be possible to determine conclusively whether hyperketonemia has an anti-seizure effect in epileptic patients independent of any associated dietary change.

A recent study of brain metabolism in normal Wistar rats fed a KME (1,3- butanediol monoester of βHB) may provide a possible explanation for the anti-epileptic effect of KDs.
Animals fed KME as 28% of daily calories for 14 days had their brain metabolites measured after removal of their brains by freeze-blowing. The KME-supplemented animals had elevated blood KB levels in the 3.5 mM range, and had a two-fold decrease in food intake despite lowered plasma glucose, insulin and leptin. The authors attributed the diminished food intake to increased malonyl-CoA and uncoupling proteins 4 and 5. Feeding the KME diet resulted in a significant decrease in both L-glutamate and GABA. This observation provides additional support for the notion that the anti-epileptic effect of KDs may result from the reduction in the excitatory amino acid, glutamate, associated with their use (77).

The anticonvulsant effect of sustained hyperketonemia has also been studied in a rat model of central nervous system (CNS) oxygen toxicity seizures (78). In an attempt to mimic the sustained therapeutic hyperketonemia (~7 mM) that can be achieved by means of a strict KD, a single oral dose (10g/kg) of a KE (R,S 1,3-butanediol acetoacetate diester) was administered to rats over a 30-minute period before placing them in a seizure-inducing hyperbaric oxygen chamber. The KE treatment was associated with a substantial delay in occurrence of the CNS oxygen toxicity-induced seizures. Ingestion of the KE resulted in rapid and sustained elevations of βHB (>3 mM), AcAc (>3 mM) and acetone (~0.7 mM). The KE had no effect on blood glucose, and the ketonemia was induced despite the fact that the rats had been fed a standard carbohydrate-containing diet.

**Ketone esters**

Conversion of ketone bodies (KB) to ketone esters (KE) eliminates KB acidity, making the KEs suitable vehicles for the delivery of KBs to the blood circulation via the gastrointestinal route.
Ingestion of KE can directly increase plasma KBs to levels within the range achieved during fasting. The degree of KB elevation attained is readily controlled by the dose size (Fig.1).

Two KEs are known to be under current study: (a) 1,3-butanediol monoester of βHB (ketone monoester [KME]) (77,79-84); (b) glyceryl-tris-3-hydroxybutyrate (3GHB) (17,85,86). Studies have demonstrated that orally or intravenously administered 1,3-butanediol or glycerol esters of βHB are safe and well tolerated in animals (80,86), and that the orally administered 1,3-butanediol monoester is also safe and well tolerated in humans (79).

Like other fatty acid esters, KEs described herein are hydrolyzed in the intestine into ketoacids and the esterifying polyol (1,3-butanediol or glycerol). Early studies on polyols such as 1,2-, 1,4- and 2,3-butanediols revealed that they had varying degrees of toxicity. In contrast, 1,3-butanediol was found to be non-toxic when fed to rats and dogs (87). When 1,3-butanediol was fed \textit{ad libitum} to rats for 43 days as a replacement for carbohydrate (which was added to a high-fat diet at 23.4% of daily calories), it was shown that 1,3-butanediol was readily metabolized in a manner similar to ethanol, with subsequent conversion to βHB, and eventually (at the peripheral tissue level) to AcAc (88). A similar study in rats later confirmed the conversion of 1,3-butanediol to βHB when it was added as a replacement of up to 20% of dietary carbohydrate energy (89).

Desrochers et al (81,82) synthesized R,S 1,3-butanediol acetoacetate monoesters and diesters as totally or partially water-soluble compounds that could replace emulsions of long-chain TAG for total parenteral nutrition. In a follow-up study, continuous intravenous administration to pigs of R,S 1,3-butanediol acetoacetate esters in amounts providing up to
30% of the hourly energy requirement resulted in their complete utilization, leading to plasma concentrations of 1,3-butanediol of 0.1 mM, and total KBs of 0.5 mM. In contrast, when the esters were given to pigs as intragastric boluses at 15% of daily calories, the blood 1,3-butanediol and KB levels were 2-3 mM and 5 mM respectively (82).

Various investigators have used the term “therapeutic ketosis”—a term that implies achievement of plasma KB levels in the 2-7 mM range—comparable to concentrations found in subjects maintained on various KDs, or in those undergoing a fast. Such degrees of hyperketonemia have been readily achieved by KE administration in rats, mice, pigs, and humans (17, 23, 54-59).

Summary

The advent of the 1,3-butanediol and glycerol esters of AcAc and βHB has made feasible oral administration of KEs as food supplements capable of providing an alternative fuel source (namely, KBs) for cognition-critical parts of the brain that, for various reasons, are manifesting impairment of glucose utilization. However, such impairment does not necessarily extend to the utilization of KBs during aging, and in certain types of early neurodegenerative disease. Given the high energy requirement of the brain and its critical dependence on the delivery of a constant supply of fuel, the consequences of leaving such an energy shortfall untreated can be dire. When the brain’s energy supply is insufficient to meet its metabolic needs, the neurons that work hardest—especially those concerned with memory and cognition—are among the first to exhibit functional incapacity (e.g. impairment of memory and cognitive performance). At the molecular level, neuronal energy deprivation is associated with impaired mitochondrial
function, with reduction in the efficiency of the electron transport chain (ETC), overproduction of reactive oxygen species (ROS), and intraneuronal (followed by extraneuronal) accumulation of deposits of amyloid-beta (Aβ) oligomers and (later) polymers, and hyperphosphorylated tau. As energy privation continues and worsens, fuel-deprived brain cells (particularly neurons that function at a high synaptic activity level) exhibit a drop in cellular energy followed by an increase in intracellular Na\(^+\) and Ca\(^{2+}\), excessive release of neurotransmitters, and apoptosis.

If the foregoing scenario is credible, it would seem critically important to test whether the hyperketonemia readily achievable by ingestion of an FDA-approved KE can prevent or delay the occurrence of neuronal energy privation (and its pathologic consequences) in individuals in whom preclinical AD or PD can be diagnosed.

It is also crucial to determine whether KE treatment \textit{per se} is effective in the prevention/control of epileptic seizures.

References


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Fig. 1. Changes in circulating D-β-hydroxybutyrate and acetoacetate concentrations for 24 hours following ingestion of a single dose of the ketone monoester. Note that concentrations reflect dose size. Reproduced from Clarke et al. (79).