Exogenous Ketones as Therapeutic Signaling Molecules in High-Stress Occupations: Implications for Mitigating Oxidative Stress and Mitochondrial Dysfunction in Future Research

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ABSTRACT: High-stress occupations (ie, firefighters, military personnel, police officers, etc.) are often plagued by cardiometabolic diseases induced by exposure to chronic stressors. Interrupted sleep cycles, poor dietary patterns, lack of physical activity, and smoke exposure along with simultaneous psychological stressors promote chronic low-grade inflammation and excessive oxidative stress. Collectively, these data suggest that practical interventions which might mitigate the underlying pathologies of these cardiometabolic diseases are warranted. Ketones, specifically R-βHB, modulates intracellular signaling cascades such as the cellular redox ratios of NAD+/NADH, the activity of NAD dependent deacetylases SIRT1 and SIRT3, and promotes a robust mitochondrial environment which favors reductions in oxidative stress and inflammation. To date, the literature examining R-βHB as a signaling metabolite has mostly been performed from endogenous R-βHB production achieved through nutritional ketosis or cell culture and mouse models using exogenous R-βHB. To the authors knowledge, only 1 study has attempted to report on the effects of exogenous ketones and the mitigation of oxidative stress/inflammation. Therefore, the scope of this review is to detail the mechanisms of R-βHB as a signaling metabolite and the role that exogenous ketones might play in mitigating diseases in individuals serving in high-stress occupations.

KEYWORDS: Ketosis, cardiovascular disease, nutrition, metabolism, inflammation, oxidative stress, firefighters, police officers, soldiers

Introduction

Men and women serving in high-stress occupations (HSO; military soldiers, police personnel, firefighters, paramedics, etc.) are exposed to a variety of acute and chronic physiological and psychological stressors. Indeed, HSO present a unique model for studying metabolic dysregulation provided these individuals often demonstrate components of metabolic syndrome while also conducting occupational tasks (ie, short sprints, victim search and rescue, fire suppression, long endurance runs, etc.) that mimic the physical and cognitive demands of athletes. However, chronic exposure to factors such as disturbed sleep patterns, nutrient poor diets, exposure to micro particulates, post-traumatic stress disorder, and lack of physical activity all potentially result in the prolonged activation of the hypothalamic pituitary adrenal (HPA) and sympathetic (SA) axes. Activation of the HPA and SA axes results in the “fight or flight” response and while these physiological responses (ie, increased cortisol, epinephrine, and norepinephrine levels) are acutely beneficial for immediate stressors, persistent activation of these axes have been linked to chronic low-grade inflammation, excessive oxidative stress (OS), and an elevated risk for developing several cardiometabolic diseases.1 In support of these data, firefighters suffer more fatalities from events related to cardiometabolic diseases, such as sudden heart attacks, than from firefighting.2

Moreover, metabolic syndrome is largely prevalent among military soldiers and law enforcement personnel as approximately 7% of law enforcement deaths are attributed to heart infarctions.3 Subsequent investigations demonstrated that police are also at an increased risk for cardiovascular morbidity compared to the general population4,5 and observational data revealed that 51% of members in the military service are too overweight (body mass index > 25.0) to be considered occupationally ready to complete their military-specific job requirements.6 Collectively these data suggest that interventions are needed which might alleviate the detrimental effects associated with serving in an HSO. Currently, the American Heart Association and American College of Sports Medicine recommend that individuals working to prevent and mitigate cardiometabolic diseases should aim to burn more calories than consumed, follow a dietary approach to stop hypertension, participate in 75 to 150 min a week of moderate-to-vigorous intensity aerobic exercise, and eliminate exposure to smoke.7 While the authors do not refute these recommendations as each would dramatically improve the health and well-being of an individual serving in a HSO, currently the data suggest poor adherence to both dietary8,9 and exercise10-13 interventions. Provided these findings, alternative and practical interventions which could be implemented in a HSO are warranted.
Inducing a state of ketosis ($\beta$-HB $\geq 0.5$ mM)$^{14}$ represents 1 intervention for combating cardiometabolic diseases and simultaneously complimenting the rigorous physical and mental exertion required by individuals in HSO. Ketone bodies acetooacetate, acetoacetate, and $\beta$-hydroxybutyrate (2 enantiomers; $S$- and $R$-$\beta$HB) have re-emerged in recent years for their purported properties as signaling metabolites, provision as an alternative fuel source during exercise, and characteristics which induce a range of pleiotropic effects. More specifically, it is the ketone body $R$-$\beta$HB which has attracted sport scientists and clinicians alike. Produced in the liver during periods of carbohydrate restriction, $R$-$\beta$HB is accumulating evidence as a metabolite which positively affects components of metabolic syndrome,$^{15}$ acts as an alternative and efficient metabolic fuel substrate compared to either glucose or fatty acids,$^{16}$ hampers OS via intracellular signaling events,$^{17}$ and augments training adaptations.$^{18,19}$ which might potentially benefit performance.

While the ketogenic diet in past years has arguably been the most common method for elevating $S$- and $R$-$\beta$HB levels and attaining nutritional ketosis, the ketogenic diet is also one of the more rigorous diets in terms of compliance requirements (carbohydrate intake $<50$ g a day or $<10\%$ total kcals$^{14}$). However, over the last decade the development of isolated ketone bodies for human consumption has produced various commercially available forms of ketone salts and ketone esters. These exogenous ketones (EK) induce an acute state of ketosis without the limitations imposed by the ketogenic diet. Provided that some of the health benefits (ie, decreased OS, inflammation, skeletal muscle catabolism, and increased cardiac muscle energetics) observed in subjects following a ketogenic diet are potentially a direct or indirect result of elevated $R$-$\beta$HB acting as a signaling metabolite, it is then reasonable to speculate that EK potentially extend the benefits of nutritional ketosis or even afford some of the purported properties of nutritional ketosis without severely limiting carbohydrate intake.

As such, the scope of this review is to discuss the mechanistic data of ketone bodies and highlight the potential field implications, if any, EK might offer individuals serving in HSO. The intricate details of ketogenesis and ketolysis exceed the scope of this article and readers are directed to a comprehensive review on ketone metabolism.$^{16}$ However, a brief overview is provided for the reader. During periods of low glucose availability and subsequent elevated glucagon-to-insulin ratios, free fatty acids are liberated from adipose tissue and circulated to hepatic mitochondria. Following several sequential reactions within the mitochondrial matrix, the ketone body acetoacetate is synthesized and either transported into circulation, converted to acetone and excreted, or the primary pathway is a reduction to $S$- and $R$-$\beta$HB. Currently, the specific effects of $S$-$\beta$HB as a signaling metabolite and in various metabolic roles are poorly understood. $S$-$\beta$HB is a minor biological intermediate during $\beta$-oxidation, a cellular antioxidant and scavenger of reactive oxygen species (ROS),$^{20}$ and 1 study found that $S$-$\beta$HB might influence the pro-inflammatory pathway NOD-, LRR- and pyrin domain-containing protein-3 (NLRP3).$^{21}$ However, Stubbs et al. demonstrated that $S$-$\beta$HB remained elevated in circulation longer than $R$-$\beta$HB likely due to differences in metabolic rates. These data support the work of earlier experimental studies in rats$^{22}$ and suggest that $S$-$\beta$HB is a poor oxidative fuel compared to $R$-$\beta$HB.$^{23}$ While only speculative, due to $S$-$\beta$HB low oxidative rates and longer elevations in the circulatory system, there is potential that $S$-$\beta$HB serves a more prominent role in redox signaling than currently thought. However, provided the continued discrepancies regarding the intracellular and systemic mechanisms of $S$-$\beta$HB, the rest of the present review will focus solely on $R$-$\beta$HB.

Unlike nutritional ketosis which is a result of inadequate glucose availability and an upregulation of the ketogenesis pathway, an alternative method for achieving acute ketosis is via EK. Currently, EK are available in either a racemic salt mixture (ie, containing both the $S$- and $R$-$\beta$HB enantiomer in a 50:50 ratio) or several different ketone ester forms. While ketone salts are a practical option for the general public due to greater availability and lower cost compared to the ester, a recent systematic review showed that ketone salts lack an ergogenic effect regarding physical performance and poorly elevate circulating $R$-$\beta$HB levels ($\leq 1.0$ mM).$^{24}$ It is worth noting that simply elevating $R$-$\beta$HB levels via EK will not independently improve physical performance.$^{24}$ Muscles seem to display an $R$-$\beta$HB saturation point between 0.8 and 2.0 mM$^{25}$ which likely evolved as a protective mechanism for sparing circulating $R$-$\beta$HB for extrahepatic tissues. Overall these data suggest a ceiling effect in skeletal muscle where further increases in $R$-$\beta$HB levels might offer no additional ergogenic benefits and potentially put an individual at risk for metabolic acidosis should ketone levels start to exceed 10.0 mM.$^{26}$ Still, although ketone salts technically achieve a state of ketosis, the lack of findings have led to speculations by several investigators of a hypothesized threshold ($\geq 2.0$ mM) for physiological benefits that extend beyond merely performance.$^{24,27-29}$ Circulating $R$-$\beta$HB of 2.0 mM or greater is commonly observed among ketone ester studies,$^{23,30-34}$ specifically those using the ketone monoester. Additionally, when ergogenic effects exist, improvements to performance tend to only be observed when $R$-$\beta$HB surpasses this hypothesized threshold.$^{30}$ It is reasonable to then speculate that other beneficial properties (eg, reduced OS and inflammation) might also occur when $R$-$\beta$HB levels are elevated beyond this threshold, such as is observed during ketogenic diet interventions. For more detailed information on

**R-$\beta$HB as a Signaling Metabolite**

**Endogenous versus exogenous ketones**

The intricate details of ketogenesis and ketolysis exceed the scope of this article and readers are directed to a comprehensive review on ketone metabolism.$^{16}$ However, a brief overview is provided for the reader. During periods of low glucose availability and subsequent elevated glucagon-to-insulin ratios, free fatty acids are liberated from adipose tissue and circulated to hepatic mitochondria. Following several sequential reactions within the mitochondrial matrix, the ketone body acetoacetate...
the characteristics of EK and nutritional ketosis, the reader is directed to a more comprehensive review.35

**Stress, R-βHB, and the mitochondria**

When considering the stress-induced redox implications for HSO, it is important to elucidate what is meant by the term “stress.” Stress is widely associated with negative aspects and outcomes.36 However, over 100 years ago, work from Yerkes and Dodson37 demonstrated a positive effect of moderate amounts of physiological arousal (ie, stress) on physical performance which was represented by the inverted U relationship between physiological arousal and performance.38 Interestingly, the concept of the U shape relationship between arousal and performance can also be applied to various aspects of cardiometabolic health, and not only to physical performance.36 For example, while OS is associated with the progression of numerous chronic diseases, acute exposure to ROS is potentially beneficial, as moderate ROS exposure serves as a trigger to up-regulate antioxidant defense mechanisms and initiate signaling for mitochondrial biogenesis.39-41 Thus, the same U shape was represented by the inverted U relationship between physiological arousal and performance.38 Interestingly, the concept of the U shape relationship between arousal and performance can also be applied to various aspects of cardiometabolic health, and not only to physical performance.36

Interestingly, the concept of the U shape relationship between arousal and performance can also be applied to various aspects of cardiometabolic health, and not only to physical performance.36 For example, while OS is associated with the progression of numerous chronic diseases, acute exposure to ROS is potentially beneficial, as moderate ROS exposure serves as a trigger to up-regulate antioxidant defense mechanisms and initiate signaling for mitochondrial biogenesis.39-41 Thus, the same U shape relationship between stress and performance can also be applied to redox balance and the hermetic stress response. In other words, excessive exposure to ROS leads to OS and chronic low-grade inflammation, while acute/moderate exposure to ROS is beneficial for facilitating favorable mitochondrial adaptations known as mitohormesis.42

HSO regularly encounter numerous chronic stressors (eg, heat exposure, intense physical exertion, disrupted sleep, etc.). Additionally, a unique aspect of HSO is the exposure to concurrent psychological and physiological stressors (ie, dual stress challenges) which have been shown to elicit greater activation of the HPA and SA axes compared to that noted from a single stressor alone. The subsequent impact is exacerbated concentrations of epinephrine, norepinephrine, cortisol, as well as markers of OS and inflammation.43,44 While acutely this response might serve beneficial physical functions, those working in HSO are at greater risk of being exposed to chronic inflammation and OS and subsequently, increased risk for cardiometabolic diseases.45 Although a direct cause and effect relationship between psychological stress and OS is currently lacking, extensive data demonstrate a strong relationship between stress, OS, mitochondrial dysfunction, and an elevated risk for developing numerous chronic cardio- and neurovascular diseases.46-48

OS is the result of an imbalance in the redox environment such that the production of ROS overwhelms antioxidant defense mechanisms resulting in oxidative damage to endogenous biomolecules. Exposure to environmental, physiological and psychological stress is known to induce OS through numerous mechanisms.49-51 It is well accepted that the mitochondria are common sources of ROS.49 An increased strain on the electron transport complex during mitochondrial respiration is likely to lead to oxygen/electron leakage from complex I and III leading to ROS, specifically formation of superoxide and hydroxyl radicals, and potentially OS.49 However, it is important to clarify that numerous enzyme systems are likely contributors to ROS as well, including (but not limited to) NADPH and xanthine oxidases as well as the endoplasm.50 Regardless where ROS originated, continual metabolic stress challenges the redox environment resulting in depleted antioxidative defense mechanisms and elevated OS, ultimately increasing risk for cardiometabolic dysfunction.

Numerous dietary interventions have been examined as potential methods to reduce OS including ketogenic diets, time restricted feeding, and caloric restriction.51-55 Indeed, the literature on viable methods of restricting calories is somewhat extensive involving both human and animal trials, demonstrating caloric restriction results in a favorable hermetic redox balance and greater resistance to stress as noted by increased antioxidant status, reduced OS, and increased lifespan.55-57 It has been widely speculated that some of the benefits associated with caloric restriction are attributed to endogenous production of ketone bodies.58-61 While more data are needed to clarify this relationship, metabolism of R-βHB has been shown to reduce OS59 and susceptibility to developing several chronic diseases.

Metabolically speaking, R-βHB is often cited as an efficient substrate for cellular work by providing a thermodynamic advantage due to the greater free energy released during ATP hydrolysis compared to either glucose or fat.62 This is supported by earlier work demonstrating ketone body infusion in perfused rat heart models suppressed glycolytic activity and improved the mitochondrial oxygen consumption efficiency.63 These findings are interesting as nutritional ketosis is often cited as a mechanism for increasing mitohormesis.64 When consuming a ketogenic diet, energy demands shift from glycolytic sources to longer chain fatty acids. The shift in substrate metabolism alters the FADH₂ to NADH ratio, effectively placing a greater demand on complex II and increasing the chance of mitochondrial ROS production.65,66 The modest increase in ROS production then acts as a signaling stimulus to induce hormetic adaptations.64 However, recent survey data suggests that individuals in HSO would depict low adherence (~10%)67 to a ketogenic diet. An argument could also be made that an intervention which further increased ROS generation in a population potentially suffering from mitochondrial dysfunction, (5-15 days)68 would likely not be an optimal intervention. It is then reasonable to speculate that in this model, chronic EK supplementation paired with a modest caloric restricted diet (~10%-20% total kcal reduction) might be advantageous provided ketones require less oxygen per mole of carbon to oxidize and therefore, reduce the strain on the electron transport complex and related antioxidant defense systems.

The oxidation of endogenous R-βHB has been shown to beneficially impact mitochondrial ROS production and reduce
OS susceptibility through a variety of proposed mechanisms.\textsuperscript{63} Endogenous \(R\)-\(\beta\)HB can function as a direct or indirect antioxidant by facilitating reduction reactions, donating electrons to ROS\textsuperscript{20} and by acting as a trigger to up-regulate mitochondrial signaling proteins.\textsuperscript{59} The oxidation of \(R\)-\(\beta\)HB changes cellular ratios of NAD\(^+\)/NADH which favors redox changes and results in the activation of signaling proteins such as NAD dependent deacetylases (SIRT1, SIRT3) that are known to increase the activity of antioxidants such as heme oxygenase 1, superoxide dismutases, and catalases through the activation of forkhead box O1 and 3 (FOXO1 & FOXO3).\textsuperscript{59} It has also been speculated that the mitochondrial consumption of \(R\)-\(\beta\)HB activates nuclear factor-erythroid 2-related factor-2 (NRF2) which serves as an additional mechanism for the up-regulation of endogenous antioxidants.\textsuperscript{59} Collectively, the potential for \(R\)-\(\beta\)HB to act as a direct and indirect antioxidant provides evidence that some of the metabolic benefits with caloric restriction may be attributed directly to \(R\)-\(\beta\)HB itself. These findings have important implications for EK and HSO who often suffer from OS induced from numerous occupational stressors. To date however, the body of literature surrounding \(R\)-\(\beta\)HB as a signaling metabolite in mitigating OS has largely been demonstrated using endogenous ketone bodies, as opposed to EK.

To the authors knowledge, only 1 study has attempted to examine the role of an EK on OS markers. In a randomized, crossover design, McAllister et al. supplemented firefighters for 7 days with either a placebo or racemic ketone salt mixture.\textsuperscript{69} Markers of OS (ie, reduced and oxidized glutathione, superoxide dismutase, catalase, total antioxidant capacity, and malondialdehyde) were collected at multiple time points pre- and post-exercise task with firefighters exercising in personal protective equipment. Their findings showed that exercising in personal protective equipment did result in OS but EK had no effect on this response. However, a glaring limitation to the investigation was the lack of blood \(R\)-\(\beta\)HB analysis throughout the study duration (ie, 7-day treatment periods). While the investigative team reported the supplementation duration as a limitation in the absent changes in OS markers and suggested longer supplementation trials, additional speculation would suggest that the null findings can also be attributed to the team’s incorporation of a racemic ketone salt supplement which arguably only slightly elevated \(R\)-\(\beta\)HB levels (~0.5 mM) and put the study’s subjects in the minimum threshold for achieving nutritional ketosis. The present team would then argue it was the non-meaningful rise in \(R\)-\(\beta\)HB levels that resulted in a lack of OS changes and perhaps not the study duration. More specifically, data reporting improvements to a subjects cardiometabolic profile are often observed following a prolonged ketogenic diet (≥3 weeks)\textsuperscript{36,80}, when \(R\)-\(\beta\)HB levels consistently approach the hypothesized threshold of ~2.0 mM. Provided these data and our speculation, if therapeutic properties do exist from EK supplementation, then they will likely be revealed through future studies incorporating ketone esters as opposed to racemic ketone salt blends. Indeed, it is the ketone monooester which routinely elevates \(R\)-\(\beta\)HB levels above 1.0 mM\textsuperscript{23,30,34,71} and is generally well tolerated compared to the gastrointestinal issues that accompany the acetocacetate diester.\textsuperscript{72}

It is difficult to elucidate whether the improvements in cardiometabolic markers are a result of the robust cellular environment induced by nutritional ketosis (eg, mitohormesis, modulation in cellular signaling, downregulation of the NLR3P pathway, etc.) and \(R\)-\(\beta\)HB is simply a byproduct of this environment, or if \(R\)-\(\beta\)HB is “truly” a signaling metabolite in the absence of these mitochondrial adaptations and adherence to a ketogenic diet. Alternatively, due to the nature of ketogenic diets, ingestion of lipids is substantially increased. A single high-fat feeding is known to induce OS acutely\textsuperscript{73} and it is possible that the mitochondrial antioxidant defense mechanisms upregulate in response to the dietary pattern to protect against lipid induced OS, not necessarily as a result of rising \(R\)-\(\beta\)HB levels. While it does not seem that \(R\)-\(\beta\)HB directly induces mitohormesis, mitochondrial health is paramount to \(R\)-\(\beta\)HB oxidation considering ketolytic enzymes are highest in type I fibers and located within the mitochondria. In vitro and mouse model studies have shown \(R\)-\(\beta\)HB to directly act as a modulator of signaling events by promoting the anti-atherogenic and anti-inflammatory throughout cells of the epithelial wall and immune system.\textsuperscript{74} Additionally, \(R\)-\(\beta\)HB has been shown to directly inhibit histone deacetylase enzymes in a dose-dependent manner and thus promote gene expression and enhanced modulation of the immune system and global reduction in OS.\textsuperscript{17,75} These data alone should warrant additional research into the potential therapeutic impact that EK might offer clinicians as current mechanistic data is scarce regarding the signaling potential of ketone bodies in human trials. An overview of \(R\)-\(\beta\)HB as a signaling metabolite within the mitochondria can be viewed in Figure 1.

**Practical implications**

Individuals serving in HSO are plagued by cardiometabolic diseases (ie, insulin resistance, atherosclerosis, obesity) which are exacerbated by the countless stressors encountered during a typical work cycle. While these diseases are multi-factorial in consideration of lifestyle factors, at the cellular level, cardiometabolic diseases tend to stem from chronic low-grade inflammation and excessive OS.\textsuperscript{76} The present review is not to suggest that EK might act in place of nutrition, exercise, or lifestyle interventions, but rather to direct future research efforts among clinicians and sport scientists alike into the therapeutic properties, if any, that EK supplementation might offer individuals serving in HSO or suffering with metabolic dysregulation.

Current in vitro and mouse data have demonstrated \(R\)-\(\beta\)HB to act as a powerful signaling molecule capable of hampering pro-inflammatory pathways. In addition, \(R\)-\(\beta\)HB seems to...
enhance the mitochondrial environment and thus improving the anti-oxidant capabilities and OS reduction properties of cells. R-βHB has also been shown to alter whole-body metabolism, consistently displaying a reduction in plasma glucose concentrations (10%-20%) as well as reductions in circulating free fatty acids. It has been proposed that the reduction observed in lipolysis can be partially explained due to an acute increase in insulin sensitivity which would highly favor metabolic health in a diseased population.

It is difficult to suggest EK as a practical intervention to individuals serving in HSO due to the current cost of the ketone esters (~ US$33.00 per serving). However, our team has worked extensively with HSO (military, firefighters, police, and correctional emergency response teams) and the incorporation of EK supplements (ie, ketone salts) as a potential ergogenic aid has become common to observe. Consequently, it is apparent that officers, lieutenants, and some nutritionists working with these individuals have applied data from ketogenic diets interchangeably with EK. A thorough review of the convergence and divergence of these 2 distinct metabolic states was recently published and outlines our similar concerns regarding merging these studies without disentangling the methods utilized to induce nutritional ketosis. However, until further data are presented regarding EK and human trials, these practices will likely continue. Therefore, interested individuals are directed towards supplementing with the ketone esters, as opposed to the ketone salts. The present evidence on ketone salts suggests a negligible to weak effect on physical and cognitive performance, as well as lack of potential to mitigate OS. In contrast, ketone esters have demonstrated some benefits as an ergogenic aid for both physical and cognitive performance, although OS and inflammatory data are missing. In earlier studies, Veech et al. described a protocol of administering divided, oral doses (100-150 g/day) of R-βHB in ester form throughout the day. It was speculated that maintaining elevated plasma R-βHB levels through the day would result in dramatic reductions to NADP+ and subsequently mitigate OS and cellular metabolic stress. Finally, it should be noted this is not an exhaustive list of the purported properties that ketone esters might offer. Present hypotheses and investigations are ongoing, attempting to provide insight into the role of ketone bodies on appetite regulation, muscle recovery, protection against respiratory viral infections, sparing of muscle protein during prolonged exercise or during states of muscle wasting, and alleviation from type II diabetes mellitus symptoms. To date, the most recent collection of EK studies have overwhelmingly focused on the potential substrate (glucose, glycogen, and fatty acids) sparing effects and improvements to physical or cognitive performance when taken acutely and prior to exercise. While the data is not conclusive on EK in

Figure 1. Pleiotropic effects of R-BHB as a signaling molecule within the mitochondria. The mitochondria are often only known as the “powerhouse” of a cell due to the reliance on the mitochondria for adenosine triphosphate production. However, the mitochondria serve an important focal point in improving cellular health and metabolic flexibility. During periods of caloric or carbohydrate restriction or when ingested as a supplement, an elevation in R-BHB can contribute to the overall health of the mitochondria. R-BHB has been shown to mitigate oxidative stress, upregulate antioxidant enzymes and anti-inflammatory transcription factors, and decrease the oxidation of glucose and fatty acids. These effects make R-BHB a favorable metabolite for future projects with an aim to mitigate cardiometabolic diseases in high-stress occupations. R-BHB, R-β-hydroxybutyrate; NLRP3, NOD-, LRR- and pyrin domain-containing protein-3; NAD, nicotinamide adenine dinucleotide; FOXO1 and 3, forkhead box O1 and 3; NRF2, nuclear factor-erythroid 2-related factor-2.
this role, it would appear thatEK are a more promising application when incorporated as a training and recovery aid and likely supplemented over a longer duration.

**Conclusion**

Interest into the therapeutic potential of ketone bodies increased when ketogenic diets reemerged as a potential intervention for treating patients suffering from epilepsy in the 1990s and again in the early 2000s from the pioneering work of Dr. Veech whom demonstrated the capabilities of ketone bodies to alleviate underlying symptoms of metabolic diseased states. Current speculation suggests that it would be incorrect to assume thatEK mimic the robust mitochondrial environment induced from adhering to a ketogenic diet. However, data are limited in human trials, specifically as it relates to the effects of ketone bodies on inflammation and OS markers. If any mitoprotective properties are elicited from EK supplementation, they will likely be observed when plasma R-[3]-βHB levels near the hypothesized ~2-0.0 mM. Currently it is unknown if EK offer HSO an advantage, but until further data are provided, the effects of nutritional ketosis achieved from a ketogenic diet and acute ketosis induced from EK will continue to be conflated.

**Author Contributions**

H.S.W and M.J.M wrote the manuscript. All authors contributed equally to the manuscript. All authors read and approved the final manuscript.

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