

# The use of nutritional supplements to induce ketosis and reduce symptoms associated with keto-induction: A narrative review

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**Background.** Adaptation to a ketogenic diet (keto-induction) can cause unpleasant symptoms, and this can reduce tolerability of the diet. Several methods have been suggested as useful for encouraging entry into nutritional ketosis (NK) and reducing symptoms of keto-induction. This paper reviews the scientific literature on the effects of these methods on time-to-NK and on symptoms during the keto-induction phase.

**Methods.** PubMed, Science Direct, CINAHL, MEDLINE, Alt Health Watch, Food Science Source and EBSCO Psychology and Behavioural Sciences Collection electronic databases were searched online. Various purported ketogenic supplements were searched along with the terms “ketogenic diet”, “ketogenic”, “ketosis” and ketonaemia (/ ketonemia). Additionally, author names and reference lists were used for further search of the selected papers for related references. **Results.** The only available evidence, from one mouse study, suggests that leucine doesn’t significantly increase beta-hydroxybutyrate (BOHB). Animal studies indicate that the short chain fatty acids acetic acid and butyric acid, increase ketone body concentrations. However, only one study has been performed in humans. This demonstrated that butyric acid is more ketogenic than either leucine or an 8-chain monoglyceride. Medium-chain triglycerides (MCTs) increase BOHB in a linear, dose-dependent manner, and promote both ketonaemia and ketogenesis. Exogenous ketones promote ketonaemia but may inhibit ketogenesis. **Conclusions.** Despite the clear ketogenic effect of MCTs, it is unclear whether they independently improve time to NK and reduce symptoms of keto-induction. There is limited research on the potential for other supplements to improve time to NK and reduce symptoms of keto-induction. Few studies have specifically evaluated symptoms and adverse effects of a ketogenic diet during the induction phase. Those that have typically were not designed to evaluate these variables as primary outcomes, and thus, more research is required to elucidate the role that supplementation might play in encouraging ketogenesis, improve time to NK, and reduce symptoms associated with keto-induction.

**The use of nutritional supplements to induce ketosis and reduce symptoms associated with keto-induction. A narrative review**

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3

4 **ABSTRACT**

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18 fatty acids acetic acid and butyric acid, increase ketone body concentrations. However, only one  
19 study has been performed in humans. This demonstrated that butyric acid is more ketogenic than  
20 either leucine or an 8-chain monoglyceride. Medium-chain triglycerides (MCTs) increase BOHB  
21 in a linear, dose-dependent manner, and promote both ketonaemia and ketogenesis. Exogenous  
22 ketones promote ketonaemia but may inhibit ketogenesis.

23 **Conclusions.** Despite the clear ketogenic effect of MCTs, it is unclear whether they  
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26 keto-induction. Few studies have specifically evaluated symptoms and adverse effects of a  
27 ketogenic diet during the induction phase. Those that have typically were not designed to  
28 evaluate these variables as primary outcomes, and thus, more research is required to elucidate the  
29 role that supplementation might play in encouraging ketogenesis, improve time to NK, and  
30 reduce symptoms associated with keto-induction.

### 31 **Introduction**

32 Very-low-carbohydrate ketogenic diets (VLCKDs) are becoming increasingly popular for  
33 mainstream and athletic use for a range of outcomes including weight-loss and maintenance,<sup>1</sup>  
34 improved satiety and a reduction in hunger.<sup>2-4</sup> The diet also offers specific benefits for health  
35 conditions ranging from neurological disorders, obesity, and diabetes and other conditions on the  
36 spectrum of metabolic syndrome, and offers potential for the adjunct treatment of various  
37 cancers.<sup>5-15</sup> Ketogenic diets elicit a state of ketosis known as ‘nutritional ketosis’ (NK), a state of  
38 hyperketonaemia distinct from pathological ketosis such as diabetic ketoacidosis (DKA).<sup>16</sup>  
39 Ketosis refers to the production of ketone bodies, derived from fats (and some amino acids) for  
40 use as an alternative fuel in times of fasting or drastic carbohydrate restriction. A restriction of  
41 carbohydrate, either by fasting or by restricting dietary carbohydrate, results in reduced insulin  
42 levels, thereby reducing lipogenesis (the creation of fats) and fat accumulation. When glycogen  
43 reserves become insufficient to supply the glucose necessary for normal  $\beta$ -oxidation of fat, via  
44 the provision of oxaloacetate in the Krebs cycle, acetyl-CoA is then used instead in the  
45 biosynthesis of ketone bodies via acetoacetyl-CoA and  $\beta$ -hydroxy- $\beta$ -methylglutaryl-CoA<sup>17</sup> to  
46 ensure provision of fuel to the Central Nervous System (CNS), which usually relies on glucose.  
47 The process of ketogenesis further allows coenzymes to be freed to ensure continued fatty-acid  $\beta$ -  
48 oxidation.<sup>17</sup> To elicit this carbohydrate restriction, while also providing sufficient alternate fuel to  
49 ensure sustainability of the diet, i.e. in comparison to fasting to achieve ketosis, VLCKDs have  
50 been used to encourage ketosis. Early research on KDs focussed on children with epilepsy and

51 for this purpose, a VLCKD typically consists of a 3:1 to 4:1 ratio of lipid to non-lipid. This  
52 treatment for epilepsy was pioneered at Johns Hopkins University Hospital,<sup>18,19</sup> and is referred to  
53 as a ‘classic’ or ‘standard’ ketogenic diet.

54 Ketogenic diets are now more commonly applied with differing definitions of what constitutes a  
55 ketogenic diet. Both low-energy diets and VLCKDs with fewer than 50 g of carbohydrate per day  
56 typically result in BOHB levels of  $\geq 0.5$  mmol.L<sup>-1</sup>.<sup>20</sup> This threshold has been used as a cut-off  
57 point for entry into ketosis by Guerci and colleagues,<sup>21</sup> and is commonly applied as a marker for  
58 entry into NK in the nutrition field, as compared to the typically higher levels expected in the  
59 medical field to elicit beneficial effects for seizure control in epileptic children.<sup>22</sup>

60 Adaptation to a VLCKD, or keto-induction, and the achievement of NK, from a standard, higher  
61 carbohydrate diet, can cause various unpleasant effects.<sup>23</sup> Symptoms of keto-induction are  
62 predominantly constipation, headache, halitosis, muscle cramps, diarrhoea, and general weakness  
63 and rash.<sup>24,25</sup> These occur because of increased urinary sodium, potassium and water loss in  
64 response to lowered insulin levels,<sup>26-29</sup> greatest between days 1-4 of a fast or ketogenic diet,<sup>26</sup> and  
65 transient reductions in glucose provision to the brain, observed to occur on days 1-3, with blood  
66 glucose normalising after day four.<sup>30</sup> Constipation may result from reduced food volume or  
67 reduced fibre intake, although this finding could be due to the groups that have been studied,  
68 which have included children with disabilities, who commonly experience constipation due to  
69 immobility.<sup>25</sup>

70 These symptoms are often referred to in the mainstream and grey literature as ‘keto-flu’ but are  
71 not well illustrated in the scientific literature. For example, a Google search returns over 22,000  
72 results for the term “keto-flu,” but the same term searched in MEDLINE Complete, CINAHL  
73 Complete, Alt HealthWatch, Food Science Source, SPORT Discus with Full Text, Psychology,  
74 and the EBSCO Behavioural Sciences Collection returns no results. Several studies have  
75 described adverse effects during ketogenic diets but to our knowledge, no studies have

76 specifically described symptoms of keto-induction in the short time between commencing a  
77 ketogenic diet and the achievement of NK.

78 Adverse effects resulting from a VLCKD are likely to reduce compliance and tolerability,<sup>31</sup> and  
79 therefore the effectiveness of these diets as clinical interventions.

### 80 *Time to ketosis*

81 There is a paucity of research that identifies specific time points to nutritional ketosis. In a study  
82 comparing fasted ketogenic protocols to a more gradual initiation of a ketogenic diet, Bergqvist  
83 and colleagues observed that participants fasting, achieved mean levels of  $\geq 0.5$  mmol.L<sup>-1</sup> BOHB,  
84 on the day following initiation of the diet, whereas those on a 1:1 ketogenic diet (by weight)  
85 achieved the same level two days after initiation of the diet.<sup>32</sup> Other studies have measured either  
86 tangentially or directly, the achievement of ‘ketosis’ but have not specifically identified the time  
87 at which a level of  $\geq 0.5$  mmol.L<sup>-1</sup> was achieved. Berry-Kravis and colleagues observed a mean  
88 time to ketosis (urinary  $>80$ mg/dl) of 42 hours.<sup>33</sup> Wirrell and colleagues have demonstrated a  
89 mean time to ketosis of 33 and 58 hours for any trace of urinary ketones or ‘good ketosis’ (of  
90  $>0.8$  mmol.L<sup>-1</sup>) respectively.<sup>34</sup> Wusthoff et al. recorded two cases of adults with prolonged  
91 nonconvulsive status epilepticus in which ‘stable ketosis’ was achieved after 8 and 10 days  
92 respectively, 3.6 and  $>1.6$  mmol.L<sup>-1</sup>,<sup>35</sup> but the definition for ketosis was not mentioned and we  
93 cannot extrapolate the time to NK as defined above. Strzelczyk et al. suggested ketosis as the  
94 presence of urinary ketones some 3.5 days after initiation of a ketogenic diet but at that time  
95 participants had achieved serum BOHB of 3.6 mmol.L<sup>-1</sup>.<sup>36</sup>

96 Hoorn and colleagues observed no difference between fasted and non-fasted ketogenic protocols  
97 for time to ketosis, without specifically describing their definitions for ketosis or the time to  
98 ketosis itself.<sup>37,38</sup> So, while the achievement of ketosis has been described in the medical  
99 literature, there is little consistency in the measurement or definition for ketosis in these papers.

100 There have been several methods suggested to reduce symptoms of keto-induction (the so-called  
101 ‘keto-flu’) and to reduce the time taken to achieve NK, including the ketogenic amino acid  
102 leucine, short chain fatty acids, medium chain fatty acids, and exogenous ketones.

103 This paper reviews the available scientific literature relevant to improvements in time to ketosis  
104 and symptoms of keto-induction, resulting from nutritional supplementation.

## 105 **Methods**

106 PubMed, Science Direct, CINAHL, MEDLINE, Alt Health Watch, Food Science Source and  
107 EBSCO Psychology and Behavioural Sciences Collection electronic databases were searched  
108 online. Various purported ketogenic supplements were searched along with the terms “ketogenic  
109 diet”, “ketogenic”, “ketosis” and ketonaemia (/ ketonemia). Additionally, author names and  
110 reference lists were used for further search of the selected papers for related references. There is a  
111 paucity of studies on time to NK and mitigation of symptoms of keto-induction. As data related to  
112 the effects of various supplements on time to induction of ketosis and on symptoms of keto-  
113 induction are limited, and there is a lack of homogeneity between study objectives, outcomes, and  
114 measures, a narrative review style was chosen.

## 115 **Results**

### 116 **Leucine**

117 Leucine and lysine are solely ketogenic amino acids. Thus, they do not contribute to  
118 gluconeogenesis. Higher leucine (and isoleucine) concentrations result from a ketogenic diet and  
119 are related to reduced glutamate-to-GABA ratio and this might explain some of the anti-seizure  
120 activity of a ketogenic diet in epilepsy.<sup>39</sup> There appears to be a high affinity of kidney cells for  
121 ketogenesis from leucine.<sup>40</sup>

122 Progression of fasting increases the conversion of leucine to ketone bodies and peripheral tissue  
123 is catabolised to provide leucine for ketogenesis.<sup>41</sup> Leucine can also be degraded in rat astroglial  
124 cells to the ketone bodies, including BOHB, and when released by these cells, used by

125 neighbouring neurones as a fuel substrate.<sup>42</sup> Leucine also results in hepatic ketogenesis.<sup>43</sup>  
126 However, studies in mice have shown that while ingested L-leucine can reduce seizure activity  
127 similarly to a KD, it does not independently increase blood levels of BOHB.<sup>44</sup>

### 128 **Short chain fatty acids**

129 Short-chain fatty acids (SCFAs) have carbon chains between two and five in length. These fatty  
130 acids include acetic acid (C:2), propionic acid (C:3), butyric acid (C:4), and valeric acid (C:5).  
131 Short chain fatty acids, especially butyric acid, are used extensively as a fuel substrate by  
132 intestinal epithelial cells.<sup>45</sup> It is generally accepted that chain length affects the relative deposition  
133 of fatty acids into either lymph or the portal vein.<sup>46</sup> Those short-chain fatty acids that escape  
134 metabolism by epithelial cells are, therefore, primarily absorbed via the hepatic portal vein and  
135 do not require ‘bundling’ with micelles and chylomicrons for absorption.<sup>47</sup> The highest quantities  
136 of short-chain fatty acids have been observed in portal blood, followed by hepatic, and far less in  
137 peripheral blood.<sup>48</sup> Thus, they bypass the usual route of absorption (for the more common long-  
138 chain fatty acids) into the lymphatics and deposition into the bloodstream via the subclavian vein,  
139 and instead, are transported via the hepatic portal vein to the liver where they can be converted  
140 into the ketone bodies.<sup>49-51</sup>

### 141 *Acetic acid*

142 Acetic acid is a two-carbon SCFA. It comprises approximately 4-20% of vinegar. Vinegar has  
143 been demonstrated to improve postprandial insulin sensitivity in healthy and diabetic people and  
144 improve glycaemic responses to meals.<sup>52-54</sup> Urinary excretion of acetone (a ketone body) is  
145 increased in phloridzinised dogs and fasting rats after feeding with acetic acid.<sup>55</sup> Acetone is the  
146 spontaneous breakdown product of the ketone bodies acetoacetate and BOHB. Thus, it is likely  
147 that acetic acid is ketogenic, and has additional benefits for overall metabolic health, however, no  
148 research has been performed on acetic acid and its specific effects on the induction of ketosis or  
149 mitigation of keto-induction symptoms in humans. Interestingly, vinegar is commonly prescribed

150 as a ‘free food’ in ketogenic diet trials,<sup>56-58</sup> and may provide an under-recognized stimulus for  
151 ketogenesis.

### 152 ***Butyric acid***

153 Butyric acid (BTA) is a four-carbon, short-chain fatty acid found in the milk of ruminants and  
154 present in small amounts in many dairy foods. Most BTA in humans is produced by microbial  
155 intestinal fermentation of dietary fibre and resistant starch. Most of the butyric acid produced by  
156 this fermentation of starches is absorbed and used directly by colonocytes, with most of the  
157 remainder absorbed into the hepatic portal vein, and transported to the liver where it can be  
158 converted to ketone bodies.<sup>50,51</sup> A small amount is absorbed directly from the large colon and  
159 enters systemic circulation, to be used directly by peripheral tissue.<sup>50</sup> Butyrate exerts effects  
160 directly on the colonic mucosa, including inhibition of inflammation and carcinogenesis,  
161 decreasing oxidative stress, and promotion of satiety.<sup>59,60</sup> Thus, it serves an important role in  
162 preserving the health of the colon, microbiota, and may have other beneficial roles for general  
163 and systemic health. Animal studies on the ketogenic potential of butyrate are mixed. For  
164 example, silage butyrate content has been shown to provide no significant effect on subclinical  
165 ketosis in dairy cows,<sup>61</sup> however, sub-clinical ketosis is higher in those receiving silage higher in  
166 butyrate content.<sup>62</sup>

167 In a recent study in humans, the effect of L-leucine, octanoyl-monoacylglycerol (O-MAG), a  
168 monoglyceride consisting of an 8-carbon fatty acid, L-carnitine, and butyric acid on acetoacetate  
169 and BOHB were studied. Both 2 g and 4 g of butyric acid were demonstrated to be more  
170 ketogenic than either 5 g of leucine, or 5 or 10g of O-MAG.<sup>63</sup>

### 171 **Medium Chain Triglycerides**

172 In medium chain triglycerides (MCTs) two-to-three of the fatty acid chains attached to the  
173 glycerol backbone are medium in length. These medium-chain fatty acids (MCFAs) are  
174 comprised of a 6–12 carbon chain. The MCTs are: caproic (C6), caprylic (C8), capric (C10) and

175 lauric acid (C12).<sup>64</sup> Similar to the short-chain fatty acids and unlike long-chain triglycerides  
176 (LCTs), MCTs do not require the actions of bile, nor micellar-chylomicron mediated absorption  
177 into the lymphatics and instead are diffused directly into the hepatic portal vein and preferentially  
178 converted into bio-available ketone bodies in the liver. Huttenlocher and colleagues first  
179 demonstrated that diets containing fewer calories from lipids than a ‘classic’ ketogenic diet—  
180 around 60%-75% of calories—can induce NK if they include a high proportion of medium chain  
181 triglycerides (MCTs).<sup>65</sup> A VLCKD with 60% of energy derived from MCTs, a three-fold greater  
182 intake of carbohydrate (18% vs. 6%) and a ~50% (7% vs. 10%) increase in protein compared to a  
183 standard ketogenic diet induces NK with no appreciable difference in BOHB levels.<sup>66</sup>

184 Dietary MCTs are also known to promote both ketonaemia and ketogenesis in animals<sup>67,68</sup> and  
185 humans with and without health conditions.<sup>69,70</sup> MCTs promote ketonaemia and ketogenesis  
186 (useful to reduce the risk of night-time hypoglycaemic coma) in those with carnitine  
187 palmitoyltransferase deficiency, a rare genetic condition which inhibits the ability to produce  
188 ketone bodies from long-chain fatty acids.<sup>71,72</sup> MCTs also increase BOHB when calorically dose-  
189 matched to either LCTs or carbohydrate in single feeding and non-ketogenic diet studies.<sup>73-76</sup>

190 When fed intravenously, MCTs increase ketogenesis when compared to both structurally similar  
191 fats<sup>77</sup> and LCTs.<sup>78,79</sup> However, ketogenesis is reduced by the simultaneous application of  
192 glucose.<sup>80</sup> It has been demonstrated by Sandstrom and colleagues that in a hypercaloric diet, there  
193 are increased BOHB levels observed with the application of MCTs that aren’t seen in a  
194 hypocaloric state.<sup>81</sup>

195 MCTs increase BOHB in a linear and dose-dependent fashion. For example, when eleven pre-  
196 term infants were fed formulas with either 25% or 50% of fat calories coming from MCTs for at  
197 least 96 hours (30 kcal/ml, around 50% calories from fat in total, 10% protein, 40%  
198 carbohydrate) the 50% MCT formula resulted in a mean plasma level of BOHB of  $0.14 \pm 0.03$   
199 mmol/L/, a nearly three-fold increase over the lower MCT formula ( $0.06 \pm 0.01$ ).<sup>82</sup>

200 While there is a paucity of research on the effect of MCTs on the time taken to achieve NK,  
201 MCTs are demonstrably ketogenic and thus, allow induction of NK with lower proportions of fat  
202 in the diet, than that used in ‘classic’ 3 or 4:1 lipid to non-lipid (or ‘ketogenic ratio’) protocols.  
203 When ‘classic’ ketogenic diets with a greater than 3:1 ratio of lipid to non-lipid are compared to  
204 MCT ketogenic diets with 60% of calories from MCT, NK can be achieved with a lower lipid  
205 intake. Huttenlocher first observed higher BOHB levels in children with epilepsy aged 2-9 years,  
206 at up to one month on an MCT ketogenic diet, and marginally lower after this time, when  
207 compared to a classic ketogenic diet, although these differences were not significant.<sup>83</sup> In a study  
208 of 55 children with severe epilepsy, Schwartz and colleagues found modified ketogenic diets,  
209 MCT ketogenic diets, and classic ketogenic diets to all be ‘ketogenic’ (inducing NK) with peak  
210 ketone body concentrations of approximately 1 mmol/L, 1.5 mmol/L and 4 mmol/L respectively,  
211 after three weeks on the differing ketogenic protocols.<sup>84</sup> Nine children were subsequently trialed  
212 on a second diet and profiled three weeks later. Cumulative results over 24 hours of metabolic  
213 testing demonstrate that expression of ketone bodies rises (in order) from a normal diet (little  
214 change) to a modified MCT diet, an MCT ketogenic diet, and the greatest rise in ketone bodies  
215 over 24 hours resulting from a classic (4:1) ketogenic diet. In a 12-month study, a classic  
216 ketogenic diet resulted in higher levels of BOHB (and acetoacetate) over all time periods (three,  
217 six, and 12 months) but this was only statistically significant at three and six months ( $p < 0.001$ ).<sup>85</sup>  
218 After ingestion of MCT at a dosage of 30g MCT/m<sup>2</sup> body surface area by nine children (in a  
219 study of seizure control), BOHB levels rose progressively after administration from a mean of 0.2  
220  $\pm$  0.1 mmol/L after an overnight fast to 1.05  $\pm$  0.3 mmol/L at 180 minutes. Participants reached  
221 NK on average at 30-60 min with most participants in NK by the 90<sup>th</sup> minute, but there was  
222 significant variation in BOHB between individuals.<sup>86</sup> With a lower dosage of 7.5 g of MCT taken  
223 three times per day after an acclimation period of 5 g MCT taken three times per day for one  
224 week, plasma BOHB was higher, yet not inducing NK.<sup>87</sup>

**225 Exogenous ketones**

226 Exogenous ketone supplements provide BOHB directly to the body without requiring ketogenesis  
227 and without concurrent elevations in free fatty acids.<sup>88</sup> They are considered to be a safe and  
228 effective way to increase ketone body concentrations.<sup>89</sup> Ketone supplements demonstrate promise  
229 as potential adjunct treatments for brain injury,<sup>90</sup> cancer,<sup>91,92</sup> Angelman syndrome,<sup>93</sup> for reducing  
230 inflammation by suppressing activation of the NLRP3 inflammasome,<sup>94</sup> and Alzheimer's  
231 disease.<sup>95</sup> Ketone supplements might also improve fueling during exercise, reduce lactate  
232 production, and improve performance due to glucose sparing,<sup>96</sup> and have positive effects on  
233 anxiety,<sup>95</sup> and mental performance and memory.<sup>95</sup>

234 Exogenous ketone supplements are available as either salts or esters of BOHB. Supplements  
235 containing ketone salts (KS) are some combination of sodium-, magnesium-, calcium or  
236 potassium-BOHB, and are available commercially from several companies under patent.<sup>97</sup>  
237 Ketone esters (KEs) at the time of writing, are only available for research, primarily as 1,3-  
238 butanediol monoester of BOHB<sup>89</sup> and thus, the animal and human research has mostly focused  
239 on the use of ketone esters. Both ketone esters and salts elevate BOHB to levels consistent with  
240 NK,<sup>98</sup> with ketone esters having greater effects on ketonaemia with ketone salts providing  
241 significantly higher reporting of gastrointestinal symptoms.<sup>99</sup> Ketone salts might provide a greater  
242 potential for long-term side effects if the inorganic ion load delivered is excessive for the  
243 individual.<sup>99</sup> Conversely, R-1,3-butanediol from ketone monoesters is readily metabolized in the  
244 liver to AcAc.<sup>100</sup> Clarke et al. detected no R-1,3-butanediol in the plasma of participants taking a  
245 ketone monoester supplement, except at the highest dosage of 714 mg/kg body weight, at which  
246 dose plasma R-1,3-butanediol was detectable at a level of  $\leq 1.0$  mmol/L and was undetectable 4  
247 hours later.<sup>100</sup>

248 At a dosage of 395 mg/kg bodyweight, KE increased BOHB in healthy volunteers from 0.2  
249 mmol/L ( $\pm 0.02$ ) at baseline to 3.3 mmol/L ( $\pm 0.2$ ) one hour later,<sup>101</sup> and from 0.16 mmol/L ( $\pm$

250 0.02) at baseline to 3.16 mmol/L ( $\pm$  0.14).<sup>102</sup> The same dose has been used to determine the effect  
251 on ketonaemia of KE taken with or without a meal. BOHB concentration (one-hour post-KE) was  
252 lower in those having taken a meal, but both groups achieved levels of ketonaemia consistent  
253 with NK; 2.1 mmol/L ( $\pm$  0.2) and 3.1 mmol/L ( $\pm$  0.1) respectively.<sup>103</sup> In a study using higher  
254 dosages (0.573 g/kg BW) in healthy male athletes performing an hour of bicycle exercise at 75%  
255 of maximal exercise intensity BOHB levels rose from 0.1 to 3.4 mmol/L ( $p < 0.01$ ) following  
256 ketone drinks.<sup>104</sup>

257 While it is clear that exogenous ketones increase serum BOHB, they are not ketogenic, and may,  
258 in fact, inhibit endogenous ketone production.<sup>105</sup> In other words, they promote ketonaemia but do  
259 not encourage the creation of ketone bodies in the liver. So, it is more accurate to say that  
260 exogenous ketones mimic the effects, many of which are positive, of NK, rather than inducing it.

## 261 **Conclusions**

262 It's unclear at this time whether an elevation in ketones over and above NK would mitigate the  
263 effects of keto-induction. It has, for example, been observed that mood is improved within the  
264 first two weeks of a diet irrespective of macronutrient composition,<sup>106</sup> and only one study, to our  
265 knowledge, has demonstrated a correlation between ketone levels and memory performance.<sup>107</sup>

266 With the exception of MCTs, there is limited research on the ketogenic potential of nutritional  
267 supplements, especially in human subjects. While the ketogenic amino acid leucine may not  
268 independently encourage ketogenesis to levels consistent with NK, more research is required, and  
269 the effect on time to NK and symptoms of keto-induction, particularly in a classic KD, are at this  
270 stage unknown.

271 Similarly, there is a paucity of research on the short-chain fatty acids and their effects on  
272 ketogenesis. Their mode of absorption and metabolism, like that of MCTs, but perhaps even more  
273 rapid, hints at a potential role for encouraging ketogenesis, and thus, the potential for improving  
274 time to NK and reducing symptoms of keto-induction.

275 There is a considerable amount of research demonstrating that MCTs promote both primary  
276 ketonaemia resulting from the conversion of medium chain fatty acids liberated from MCTs into  
277 bio-available ketone bodies, and longer-term ketogenesis by facilitating keto-adaptation.  
278 Expression of the ketone body BOHB is increased in a linear, dose-dependent manner in response  
279 to oral loads of MCT but it is unclear whether MCTs independently improve time to NK.  
280 Modified MCT ketogenic diets do not significantly hasten the induction of NK over a classic  
281 ketogenic diet with a minimum of three parts lipid to one part non-lipid, but they do allow NK to  
282 occur in diets containing greater amounts of non-lipid macronutrients.

283 There has, however, been little research performed on the application of MCTs to classic  
284 ketogenic diets and whether, if applied, they would; a) improve time to NK, b) result in  
285 significantly higher levels of BOHB, and c) significantly reduce symptoms of keto-induction. It is  
286 also unknown if, in the context of a ketogenic diet, MCTs provide additional benefits, for  
287 example for physical and mental performance and mood.

288 Exogenous ketones are unlikely to be ketogenic per se, and may inhibit ketogenesis, however, the  
289 rapid and substantial elevation of BOHB offers potential to mitigate effects of keto-induction, and  
290 thus, could play a role in improving adherence to a ketogenic diet. Newport et al. have reported  
291 improvements in mood and cognitive performance resulting from ketone ester treatment over 20-  
292 months in an Alzheimer's Disease case. In this case, cognitive performance tracked plasma  
293 BOHB concentrations. In a direct, dose-matched comparison, Kesl and colleagues evaluated the  
294 effects of ketone esters, salts, MCTs, and MCT + KS on blood BOHB in Sprague-Dawley rats at  
295 a dose of 5 g/kg. At 0, 30, and 60 min and 4, 8, and 12 hrs post administration (by intragastric  
296 gavage) KS + MCT and MCT supplementation rapidly elevated and sustained significant BOHB  
297 elevation compared to control for the duration of the 4-week study. Ketone salts did not  
298 significantly elevate BOHB at any time point tested compared to controls. Ketone ester  
299 supplements significantly elevated BOHB levels for the duration of the 4-week study. This

300 further demonstrates, albeit, in non-human subjects, the superiority of KE to KS for elevating  
301 BOHB, and the utility of MCT for the same purpose, but is likely to be limited in applicability to  
302 health and performance as we have seen demonstrable increases in BOHB, consistent with NK  
303 levels with supplementation of KS in humans.<sup>98,99</sup> Research performed on exogenous ketone  
304 supplements is, at this time, highly preliminary, and has been predominantly performed using  
305 animal subjects. Further clinical research is required to translate the potential benefits seen in  
306 these studies, to human models of disease and disorder.

307 This review was limited by a dearth of studies demonstrating the effect of supplementation on  
308 specific time points to NK in humans, and on symptoms of keto-induction. While studies have  
309 described symptoms arising from a ketogenic diet, few studies have specifically evaluated  
310 symptoms and adverse effects of a ketogenic diet during the induction phase, and the studies that  
311 have been performed typically have not been designed to evaluate these as primary outcomes,  
312 and thus, our conclusions are extrapolated from a variety of sources. There is also little consensus  
313 on whether greater levels of BOHB (over and above NK threshold) are, in fact, associated with  
314 fewer symptoms of 'keto-flu', nor for that matter with improved outcomes but as previously  
315 noted, Newport and colleagues have observed a linear correlation between mood and cognition,  
316 and BOHB levels.<sup>108</sup> Adverse effects associated with the induction of NK might cause increased  
317 drop-out rates and preclude some of the positive effects for those that would otherwise benefit  
318 from a VLCKD. For example, Yancy and colleagues noted an 8% overall dropout rate due to  
319 difficulties adhering to an LCHF diet, with a further 5% withdrawing from their study due to  
320 adverse effects.<sup>24</sup> High attrition rates due to tolerability and gastrointestinal side effects have also  
321 been noted in childhood epilepsy research utilising VLCKDs.<sup>7,38</sup>

322 Preliminary research suggests that increased BOHB levels and a faster time-to-NK might  
323 improve the acceptability of the KD and improve compliance rates, but more research is required  
324 to understand the role that supplementation could play in encouraging ketogenesis, improving

325 time to NK, reducing symptoms associated with keto-induction, and the effect this might have on  
326 improving adherence to, and outcomes from a VLCKD.

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