Since 1921, ketogenic diet (KD) therapy has been a recognized effective tool in the treatment of epilepsy. Interest in diet therapy waned following the introduction of antiepileptic drugs (AEDs), until the 1990s, when studies and clinical trials emerged demonstrating its efficacy in drug-resistant patients and particular pediatric epilepsy syndromes (1–3). Indeed, there is growing evidence of diet efficacy in the management of glucose transporter type 1 deficiency syndrome, pyruvate dehydrogenase deficiency, infantile spasms, Lennox-Gastaut syndrome, Dravet syndrome, Angelman syndrome, and myoclonic-astatic epilepsy (4–9). Many patients diagnosed with these syndromes are now reaching adulthood, while others are being diagnosed de novo as adults with recent advances in genetic testing. In the management of drug-resistant epilepsy (seizures resistant to two or more AEDs), patients have a less than 5% chance of seizure freedom with additional drugs and may not be surgical candidates due to a generalized epilepsy, multifocal nature, or nonresectable seizure focus (10). Emerging evidence suggests that KDs can offer seizure reduction and seizure freedom in a subset of these patients, not only children but now adults as well.

**Ketogenic Diet Variants**
The classic KD is the original diet introduced into practice in the 1920s. This high-fat, low-carbohydrate diet induces ketone body production through fat metabolism with the goal of mimicking a starvation state without depriving the body of necessary calories to sustain growth and development (11,12). It is typically composed of a 4:1 ratio of fat (in grams) to protein plus carbohydrates (in grams), thus shifting the predominant caloric source from carbohydrate to fat. Lower ratios of 3:1, 2:1, or 1:1 (referred to as a modified KD) can be used depending on age, individual tolerability, levels of ketosis, and protein requirements (13). To increase flexibility and palatability, more relaxed variant forms (Figure) include the modified Atkins diet (MAD), the low glycemic index treatment (LGIT), and the KD combined with medium-chain triglyceride (MCT) oil. The MAD, introduced in 2003, typically employs a net 10 to 20 g carbohydrate per day limit, which is roughly equivalent to a ratio of 1–2:1 of fat to protein plus carbohydrates (14, 15). The MAD does not require calculating and weighing of food portions or an initial hospital stay for implementation (12, 16). The LGIT recommends 40 to 60 g of carbohydrates daily with the selection of foods with glycemic indices <50 and ~60% of dietary energy derived from fat and 20 to 30 percent from protein (17). The MCT variant KD uses medium-chain fatty acids provided in coconut and/or palm kernel oil as a diet supplement. It provides an option for individuals with carnitine deficiency, as carnitine is required for processing the long-chain fatty acids supplied by the classic KD, and allows for greater carbohydrate and protein intake than even a lower-ratio classic KD (18), which can improve compliance.

**Ketogenic Diets in the Management of Chronic Epilepsy in Adults**
The demand for KD access in adults comes from children already on KD therapy transitioning into adulthood, patients with adult-onset epilepsy refractory to medications or wishing to limit AEDs due to side-effect burden, and adults with chronic epilepsy who were not offered a KD as children and are appropriate candidates. These patients are motivated to achieve seizure control; secondary goals include improved overall quality of life and employment potential, independence in the activities of daily living, the ability to obtain driving privileges, and reduced risk of injury, sudden unexplained death in epilepsy, and AED burden. Thus, evidence supporting diet efficacy (≥50% reduction in seizure frequency) in adults is needed.

A 2011 review of dietary treatments in adolescents and adults pooled data from seven studies of the classic KD to show that 49% of 206 patients had ≥50% seizure reduction and, of these, 13% were seizure free (19). A 2015 meta-analysis reviewing ketogenic dietary treatments in adults from 12
Ketogenic Diet Use in the Management of Refractory Status Epilepticus in Adults

Status epilepticus (prolonged seizure lasting longer than 5 minutes or recurrent seizures without return to baseline between seizures) that continues despite appropriate first- and second-line AEDs is classified as refractory status epilepticus (RSE). If status epilepticus continues or recurs 24 hours or more after the initiation of treatment with anesthetic agents to induce burst or seizure suppression, patients are diagnosed with super-refractory status epilepticus (SRSE) (29). Several case reports and case series demonstrate the successful use of KD therapy for the management of RSE and SRSE. The first report of KD use for SRSE in an adult was published in France (30). Subsequently, a 4:1 ratio KD was introduced to two adult patients with SRSE at the University of Pennsylvania after 20 and 101 days of status epilepticus, respectively, and resulted in seizure cessation after 6 and 11 days, respectively (31). Similar cases of successful resolution of status epilepticus in adults within 2 weeks with the classic KD and 4 days with LGIT have also been reported (32–34). A case series of 10 adults with SRSE lasting a median of 21.5 days who were treated with a KD (nine with a 4:1 ratio KD and one with a 3:1 ratio KD) at four medical centers showed successful cessation of status epilepticus in 100% of patients who achieved ketosis (9 of 10 adults) at a median of 3 days (range = 1–31 days) (35). In the most recent and largest phase I/II clinical trial of 15 adult patients treated with 4:1 ratio KD therapy (14 of whom completed therapy) after a median of 10 days of SRSE at four medical centers, 11 (79% of patients who completed KD therapy, 73% of all patients enrolled) achieved resolution of seizures in a median of 5 days (range = 0–10 days) (36).

As both RSE and SRSE carry high rates of morbidity and mortality (37), KD therapy offers a needed adjunct strategy for management. It has the potential advantages of working rapidly and synergistically with other concurrent treatments; is relatively easy to start, monitor, and maintain in the controlled intensive care unit setting with close follow-up; does not contribute to hemodynamic instability seen with anesthetic agents; and could potentially reduce the need for prolonged

Diet adherence and compliance remain significant barriers to successful implementation of ketogenic diets and an adequate assessment of efficacy. The aforementioned meta-analysis of 11 studies of KDs in adults found a combined adherence rate of 45% for all KD types, 38% for the classic KD, and 56% for the MAD (20). A subsequent study of the classic KD noted that 39% of subjects had a ≥50% seizure reduction (9/23) and 8% had ≥90% seizure reduction (2/23); a subsequent MAD study noted that 31% of subjects had ≥50% seizure reduction (4/13) (21, 22). In the largest observational study of 101 adult patients naïve to diet therapy who subsequently started the MAD, 39% had ≥50% seizure reduction and 22% became seizure free following 3 months of treatment (23). Thus, the classic KD reduced seizures by ≥50% in 22 to 70 percent of patients and by ≥90% in up to 52% of patients, while the MAD reduced seizures by ≥50% in 12 to 67 percent of patients and by ≥90% in up to 67% of patients (21, 23).

Dietary treatment in patients with drug-resistant epilepsy has several advantages. Dietary treatments can be rapidly initiated, and the beneficial effects can be seen almost immediately. They can be used safely and effectively for most types of epilepsy and in patients of all ages worldwide (25). Several studies in adults have also reported benefits beyond seizure control, including improvements in arousal, mood, alertness, energy, and concentration (22, 26). Other studies show that diets are effective in treating comorbid diseases, such as obesity and type 2 diabetes, which are a major health concern in the adult population (27). Furthermore, quality-of-life scores tend to increase rather than decrease with diet therapy (28).

FIGURE. Comparison of classic (4:1 ratio) ketogenic diet and ketogenic diet variant composition used for adults. GI, glycemic index.
use of these drugs. Moreover, there may be an additional neuroprotective benefit related to improved mitochondrial function due to increased energy reserves combined with decreased production of reactive oxygen species (38). For example, the ketogenic diet has been shown to stimulate mitochondrial biogenesis, increase cerebral ATP concentrations, and result in lower reactive oxygen species production in animal models (39, 40). There is additional growing evidence that ketone bodies exhibit protective anti-inflammatory effects (41). In preclinical studies, KD treatment reduces microglial activation, expression of proinflammatory cytokines, and pain and inflammation after thermal nociception (42–44); it also improves outcomes in models of Parkinson disease and multiple sclerosis through proposed mechanisms that include reduced brain inflammation and neurodegeneration (43, 45). These anti-inflammatory properties may explain the observed benefit of KD in treating patients with SRSE secondary to autoimmune and presumed autoimmune encephalitis, such as those with new onset refractory status epilepticus and febrile infection-related epilepsy syndrome.

Management of Adverse Effects in Adults

The most commonly reported adverse effects associated with KD use in adults are gastrointestinal effects, weight loss, and a transient increase in lipids. The gastrointestinal side effects, which include constipation, diarrhea, and occasional nausea and vomiting, are typically mild, improve with time, can often be managed with diet adjustments under the guidance of a dietitian or nutritionist, and infrequently require medical intervention. Smaller meals, increased fiber intake, exercise, and increased sodium and fluid intake can often prevent or alleviate these complaints. Weight loss may be an intended positive effect in patients who are overweight, but for those who want to maintain or gain weight, adjustments in caloric intake can be recommended. Increases in serum lipids have been shown to normalize with continued diet therapy (after 1 year) or return to normal after cessation of diet therapy (46–48). In addition, very-low-carbohydrate diets that induce ketosis have been shown to lead to reductions in serum triglycerides, low-density lipoprotein, and total cholesterol and increased levels of high-density lipoprotein cholesterol in adults (27). Whether ketogenic diets influence extended serum and vascular markers of atherogenesis in adults with epilepsy beyond standard lipid profiles warrants further investigation. Adults with persistent hyperlipidemia who are responding well to diet therapy may consider increasing fat sources of omega 3 or adhering to published lipid-lowering recommendations for the MAD, although further research is needed to demonstrate that these interventions provide benefit (48). Other potential side effects can result from vitamin and mineral deficiencies secondary to restricting carbohydrates and prolonged ketonemia, including osteopenia and osteoporosis (12, 49, 50), although the precise mechanism remains unclear. The standard practice of supplementing a recommended daily allowance of multivitamin and mineral supplements can reduce the risk of such deficiencies.

Conclusions

Clinical evidence now demonstrates the effectiveness of ketogenic diets in treating drug-resistant epilepsy in adults. Potential side effects of diet therapy are often preventable and manageable; however, additional strategies are needed to improve long-term adherence.

Highlights

1. Ketogenic diets are an important adjunct to AEDs for management of chronic epilepsy and RSE in adults.

2. Studies support the feasibility, tolerability, and efficacy of a KD in adults.

3. Ketogenic diets may have anti-inflammatory properties that could benefit patients with epilepsy caused by an autoimmune process.

4. Compliance and long-term side effects remain major considerations.

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Conflict of Interest

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References


