Review

Carbohydrates and insulin resistance in clinical nutrition: Recommendations from the ESPEN expert group


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Article history:
Received 3 August 2016
Accepted 13 September 2016

Keywords:
Clinical nutrition
Carbohydrates
Insulin resistance

SUMMARY

Growing evidence underscores the important role of glycemic control in health and recovery from illness. Carbohydrate ingestion in the diet or administration in nutritional support is mandatory, but carbohydrate intake can adversely affect major body organs and tissues if resulting plasma glucose becomes too high, too low, or highly variable. Plasma glucose control is especially important for patients with conditions such as diabetes or metabolic stress resulting from critical illness or surgery. These patients are particularly in need of glycemic management to help lessen glycemic variability and its negative health consequences when nutritional support is administered. Here we report on recent findings and emerging trends in the field based on an ESPEN workshop held in Venice, Italy, 8–9 November 2015. Evidence was discussed on pathophysiology, clinical impact, and nutritional recommendations for carbohydrate utilization and management in nutritional support. The main conclusions were: a) excess glucose and fructose availability may exacerbate metabolic complications in skeletal muscle, adipose tissue, and liver and can result in negative clinical impact; b) low-glycemic index and high-fiber diets, including specialty products for nutritional support, may provide metabolic and clinical benefits in individuals with obesity, insulin resistance, and diabetes; c) in acute conditions such as
1. Introduction: carbohydrates, insulin resistance, and clinical nutrition

Carbohydrates in the diet provide an essential metabolic fuel, commonly in the form of glucose. While necessary for life, excess or rapidly changing levels of glucose in the blood can lead to several health problems and contribute to the development of obesity, insulin resistance, and type 2 diabetes mellitus (T2D). Furthermore, poorly controlled glucose levels in critically ill patients or those recovering from surgery can lead to glucose variability with hyper- and hypoglycemia, conditions that can impede recovery and be fatal. In order to summarize recent research findings, share ideas, and discuss how emerging avenues of research may shape clinical nutrition recommendations and guidelines in the future, the authors of this manuscript participated in a workshop hosted by the European Society for Clinical Nutrition and Metabolism (ESPEN) on November 8th and 9th 2015, in Venice, Italy. In this manuscript about glucose and glycemic control in clinical nutrition, we report on key concepts from workshop presentations. This report was prepared from a first draft based on summaries provided by each speaker, professionally edited, and further reviewed and revised in multiple rounds by all authors. In this summary paper, we review how major metabolic organs use glucose and regulate its levels within the body, explain conditions that disrupt glycemic control, and discuss dietary and clinical nutrition guidelines for the treatment of conditions that feature dysglycemia.

Common digestible carbohydrates are classified as monosaccharides (glucose, fructose, and galactose), disaccharides (sucrose, lactose), or polysaccharides (starches, glycogen), based upon chemical structure [1]. Alternatively, carbohydrates are grouped based upon their digestibility and nutritional effect: the alpha bonds between glucose molecules in starch are easily broken down in digestion, whereas beta bonds in fibers are resistant to human digestive enzymes. Digestible carbohydrates break down and provide the body with monosaccharides for energy, while those that resist digestion are non-glycemic, but instead provide energy through fermentation in the colon by the gut microbiota. Carbohydrate quality and digestibility can influence post-prandial plasma glucose concentration and the inflammatory response, which is now known to underlie the development of insulin resistance, metabolic syndrome, and T2D [2]. Foods with high glycemic index (GI) and glycemic load (GL) are associated with increased risk of such diseases [3–5]. Conversely, lowering dietary GI and GL can improve metabolic control [6–11]. Furthermore, increasing the protein-to-carbohydrate ratio can reduce glycemia [12], and inflammation can be tempered through dietary modification [13].

2. Glucose metabolism in the organs

Advances in research have shed light on the ways in which glucose interacts with a number of organ systems. Excess exposure of these organs to glucose as a result of hyperglycemia, as well as uncontrolled spiking of glucose levels after meals, can contribute to the deterioration of an individual’s condition by causing metabolic derangements such as oxidative stress, tissue and systemic inflammation, and insulin resistance. This section summarizes the impact of glucose on major organs involved in substrate metabolism and utilization.

2.1. Central nervous system

The relationship between glucose and the brain is important for the whole body. Glucose is the major physiological source of energy for the brain, and the brain senses glucose and carbohydrate levels...
throughout the body (Fig. 1A, B). The brain utilizes hormones to signal to other organs (Fig. 1B–E), communicate glucose status, and influence whole-body glucose homeostasis [14–16]. The impaired glucose homeostasis that occurs in T2D may be caused in part by early defects in central nervous system glucose sensing mechanisms [16].

2.2. Skeletal muscle

Skeletal muscle is a major contributor to whole-body glucose utilization, as glucose is a relevant fuel for the maintenance of skeletal muscle energy homeostasis (Fig. 1A, D). However, excess glucose exposure can lead to muscle damage [17], which in turn has health and clinical consequences for the individual. Mechanisms of glucose-induced tissue damage are complex and may vary in acute and chronic conditions. Common fundamental pathways causing muscle damage following exposure to excess glucose however include oxidative stress, inflammation, and insulin resistance, and it may alter tissue cell proliferation and differentiation [18]. Elevated glucose has been shown to cause mitochondrial damage and dysfunction in muscle cell culture experiments [19], thereby potentially leading to impaired tissue energy metabolism and substrate utilization. Through these combined mechanisms, hyperglycemia may enhance muscle protein catabolism leading to reduced lean body mass and strength [20–22]. In agreement with the above observations, people with T2D demonstrated activation of pro-inflammatory signaling pathways [23] and substantially enhanced protein breakdown [24] in skeletal muscle compared to healthy individuals. Muscle alterations are likely to become more clinically relevant when diabetes-induced hyperglycemia is associated with synergistic oxidative, pro-inflammatory, and insulindesensitizing conditions such as aging or chronic and acute disease.

2.3. Adipose tissue

Adipose tissue plays a major role in maintaining whole-body metabolic homeostasis [25], but its accumulation is associated with adverse outcomes such as metabolic syndrome and diabetes, cardiovascular events and several chronic diseases [26]. In recent years, research findings have revealed that qualitative changes in metabolic and endocrine characteristics of adipocytes (adiposopathy) mediate aspects of human disease. Metabolic research breakthroughs have uncovered ways that adipose tissue has substantial impact on energy balance, insulin resistance, inflammation and obesity-associated complications. Recently, differences between white and brown adipocytes have been described. White adipose tissue is the most abundant type of adipose tissue in human adults, and it functions as an energy store as well as a modulator of whole-body substrate utilization and metabolism through its endocrine functions [27]. Brown adipose tissue has an increasingly recognized metabolic importance due to its higher mitochondrial content with high levels of uncoupling. These features lead to generation of heat (thermogenesis) associated with energy dissipation that may favor resistance to obesity and diet-induced weight gain [28]. Lower brown adipose tissue content has been described in people with obesity or T2D than in healthy individuals [29]. Experimental research has indicated that white adipose tissue can be converted into its more beneficial, metabolically active brown counterpart, and this process has become the target of intensive research [27,30–33]. Irisin, an exercise-induced myokine, is thought to underlie the observed browning of adipose tissue in experimental models [30]. Although controversy surrounds the role of irisin in humans [34], this process may further underscore the potential importance of loss of muscle mass and function in the onset of obesity-associated metabolic complications.

Glucose modulates adipose tissue metabolism and mass both directly and indirectly by increasing insulin secretion and plasma concentration. Hyperinsulinemia is a key inducer of lipogenesis and adipose tissue expansion, and selective adipose tissue insulin receptor knockout may protect from fat tissue accumulation [35,36] (Fig. 1E). In agreement with the emerging view of inter-organ cross-talk to regulate metabolism, experimental models have demonstrated that glucose is able to re-direct stem cells derived from non-adipose tissues such as skeletal muscle to differentiate into ectopic adipocytes [37].

2.4. Liver

Interactions between carbohydrate and fat substrate availability may affect non-adipose tissues in the body by favoring ectopic lipid deposition [38]; non-alcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH) represent a relevant example of this negative interplay. NAFLD is an early indicator of insulin resistance and metabolic syndrome in people with obesity [39], triggered by inflammatory processes induced by overfeeding. Carbohydrates play a key role in the onset of NAFLD, with a negative impact for both excess glucose and fructose, whereas complex and non-digestible carbohydrates may be protective [40,41]. High glucose contributes to the onset of NAFLD also by enhancing circulating insulin, which in turn contributes to hepatic lipogenesis. Inflammation and pro-inflammatory signals are generated by excess fat accumulation [26,42,43] and are also directly triggered by endotoxin translocation from the gut to the liver, with fructose being a major regulator of this process [44]. Underlying molecular mechanisms of liver damage are not fully understood but may involve the immune response and activation of immune signaling pathways that can cause liver damage and fibrosis [45–48]. Steatosis is reported to be ameliorated by intake of omega-3 polyunsaturated fatty acids (PUFAs) [49] and complex carbohydrates [50,51]; elements of the Mediterranean diet [52,53] and the Asian diet [54,55] may therefore prevent metabolic liver disease. NAFLD can also be combated by the fostering of a healthy and diverse population of gut microbiota, as discussed below [44,56–59].
Ingestion of a fructose-free diabetes-specific disease (NAFLD), involving stimulation by fructose ingestion of non-alcoholic fatty liver disease mainly by triggering pro-inflammatory responses, and by favoring efficient nutrient absorption [60,61]. On the other hand, beneficial bacterial strains may result in protection from metabolic disease, and interaction with non-digestible dietary carbohydrates contributes to this effect. In particular, dietary fibers interact with the gut microbiota and may reduce inflammation and unfavorable metabolic responses, thereby also reducing hepatic steatosis [41,62]. Gut microbiota-driven fermentation of non-digestible carbohydrates or prebiotics can decrease carbohydrate-induced blood glucose spikes that occur after a meal [63]. Probiotics may further modulate release of gut peptides including glucagon-like peptide 1 (GLP-1), also potentially contributing to limit obesity and its metabolic complications (45, 82). Fermentation of non-digestible carbohydrates also results in production of short chain fatty acids (SCFAs) that may play protective roles and reduce the risk for systemic and local disease including cancer. Obese individuals are reported to display metabolically unfavorable populations of gut microbes, and weight loss after gastric bypass surgery may shift this pattern towards one resembling normal weight individuals [64,65]. The possibility of harnessing microbiota to treat obesity and metabolic disease is under intensive investigation. Small-scale clinical studies of probiotic supplementation have found favorable changes to glucose and fat metabolism [61,66–68]. Research has identified metabolically beneficial bacterial strains in the gut microbiota, like Lactobacillus, and Bifidobacterium, or Akkermansia, though their role as modulators of the host metabolism is still debated [69,70]. Larger and longer-term human trials are still necessary before tailored probiotic use can be incorporated into official guidelines for the treatment of obesity and metabolic syndrome [61,71].

### 2.5. Gut and gut microbiota

The gut plays central roles in the processing of carbohydrates and thereby influences glucose balance in the body (Fig. 1A). Gut endocrine functions and the gut bacterial population (microbiota) are emerging key players in the regulation of intermediary metabolism. Unfavorable microbiota may contribute to the onset of obesity and metabolic syndrome mainly by triggering pro-inflammatory responses, and by favoring efficient nutrient absorption [60,61]. On the other hand, beneficial bacterial strains may result in protection from metabolic disease, and interaction with non-digestible dietary carbohydrates contributes to this effect. In particular, dietary fibers interact with the gut microbiota and may reduce inflammation and unfavorable metabolic responses, thereby also reducing hepatic steatosis [41,62]. Gut microbiota-driven fermentation of non-digestible carbohydrates or prebiotics can decrease carbohydrate-induced blood glucose spikes that occur after a meal [63]. Probiotics may further modulate release of gut peptides including glucagon-like peptide 1 (GLP-1), also potentially contributing to limit obesity and its metabolic complications (45, 82). Fermentation of non-digestible carbohydrates also results in production of short chain fatty acids (SCFAs) that may play protective roles and reduce the risk for systemic and local disease including cancer. Obese individuals are reported to display metabolically unfavorable populations of gut microbes, and weight loss after gastric bypass surgery may shift this pattern towards one resembling normal weight individuals [64,65]. The possibility of harnessing microbiota to treat obesity and metabolic disease is under intensive investigation. Small-scale clinical studies of probiotic supplementation have found favorable changes to glucose and fat metabolism [61,66–68]. Research has identified metabolically beneficial bacterial strains in the gut microbiota, like Lactobacillus, and Bifidobacterium, or Akkermansia, though their role as modulators of the host metabolism is still debated [69,70]. Larger and longer-term human trials are still necessary before tailored probiotic use can be incorporated into official guidelines for the treatment of obesity and metabolic syndrome [61,71].

### 2.6. Fructose

Glucose is the body’s key form of energy and the most clinically relevant carbohydrate employed in patient nutritional support. For these reasons, glucose is the main focus of the current review. However, glucose is not the only simple sugar available through the diet. Fructose (as a monosaccharide or in the disaccharide sucrose) is also found in a variety of foods, but is processed differently by the body. Fructose has also been a focus of research, as it not only enters the diet through fruits but also is added to juices and other food products as a sweetener, and therefore is widely consumed. After absorption, fructose is metabolized by the liver and can be converted into glucose, lactate, and fatty acids. Fructose-induced hepatic lactate release is a unique feature and opposite to extrahepatic lactate flux to the liver for de novo glucose production. High-fructose diets have been reported to decrease insulin-mediated suppression of glucose production and to increase hepatic lipogenesis and plasma triglyceride concentrations [72], although recent meta-analyses have failed to confirm associations between fructose intake and several metabolic alterations potentially due to additional adaptive changes [73]. As introduced above, a stronger link has been established between fructose and non-alcoholic fatty liver disease (NAFLD), involving stimulation by fructose ingestion of pro-inflammatory signals reaching the liver from the gut [44,46,74]. Ingestion of a fructose-free diabetes-specific nutrition supplement formula (DSF) was shown to cause lower blood glucose concentrations in patients with diabetes than formulas with fructose [75], and physical activity has been shown to attenuate its deleterious effects on glycemic control [76]. However, as these effects of fructose are still debated [73], additional trials to determine whether fructose in particular should be avoided in the diet are necessary.

### 3. Recommendations for glycemic management and nutritional support

#### 3.1. Obesity, metabolic syndrome, and diabetes

**3.1.1. Diet and lifestyle**

Obesity and excess adiposity can lead to the development of glucose insensitivity, impaired insulin action, and inability to properly regulate glycemic variations. Although dietary recommendations aimed at weight loss have recently emphasized the importance of inducing energy deficits, at least in part independently of diet composition, high GI and GL foods are associated with metabolic disease risk and health complications [3–5]. Lowering dietary GI and GL may conversely improve these outcomes and benefit patients with obesity and diabetes [6–11]. Non-digestible carbohydrates may also provide beneficial metabolic effects. Soluble fiber is reported to decrease postprandial plasma glucose concentration and it may additionally decrease blood LDL-cholesterol concentration [7,77]. Insoluble fiber, especially cereal fiber, decreases the risk of T2D and cardiovascular disease [78]. High fiber intake is therefore recommended for people with diabetes or at risk of developing diabetes, including people with obesity and metabolic syndrome (i.e. the cluster of cardiometabolic risk factors including high waist circumference, high blood pressure, elevated blood glucose and dyslipidemia with high triglycerides and low HDL-cholesterol). Such nutritional recommendations (Tables 1 and 2) have been increasingly introduced by several health care organizations and are currently included in guidelines for patients with or at risk of developing T2D, and they are also appropriate for the management of plasma glucose concentration in type 1 diabetes (T1D) [79–81].

**3.1.2. Disease-specific nutritional supplement formulas for diabetes**

Nutritional support can cause or exacerbate hyperglycemia, especially in obese and diabetic patients, and hyperglycemia is associated with higher morbidity and mortality [91,92]. In the clinical nutrition setting, a burgeoning field of research is dedicated to designing nutritional support products for people with diabetes. Such products aim to limit glycemic variation after administration [93]. Diabetes-specific formulas (DSF) have many of the following ingredients in common: a) lower carbohydrate content than standard formulas (SFS); b) higher proportion of complex carbohydrates that are slowly digestible to reduce blood glucose spiking; c) modified maltodextrin, starch, fructose, isomaltulose, and sucromalt, rather than the maltodextrin, starch, and sucrose found in SFS [94]; d) fat content enriched in unsaturated fatty acids, especially monounsaturated fatty acids, in higher proportion than in SFS [87]; e) fiber content higher than in SFS [95].

Based on this available evidence, the ESPEN expert group endorses the utilization of DSFs for nutritional support of people with obesity and diabetes. When parenteral nutrition must be used, the risk of hyperglycemia in obese and diabetic patients can be reduced if the initial amounts of glucose provided in the TPN bag are limited to less than 2 g/kg/day until proper glycemic control is observed [96]. With the use of enteral nutrition, the risk of hyperglycemia can be decreased by modification of the total amount and of the quality of carbohydrates used. Numerous short- and mid-term studies prove that enteral DSFs are associated with reduced postprandial blood glucose, postprandial blood insulin, mean blood glucose values, glycemic variability, short-acting insulin
requirements, and changes in HbA1c [75,97–102]. As with any other treatment decision, the individual patient features, different clinical settings, and various combinations of insulin therapy may influence the choice between an enteral SF or DSF for patients with obesity or diabetes. Additional randomized controlled studies are desirable to identify optimal formula composition for different clinical conditions.

3.1.3. Recovery from surgery and critical illness

Acute states of metabolic stress often occur in the presence of disease. Such alterations may occur in individuals with otherwise normal weight and glucose metabolism, or in patients with obesity, metabolic syndrome, or diabetes for reasons related or unrelated to the illness causing the metabolic stress. Critical illness and recovery from surgery are common clinical conditions requiring specific consideration.

3.1.4. Nutrition for enhanced recovery in surgical patients

Conventional thinking about the nutritional support of surgical patients has been challenged in recent years by a body of evidence demonstrating the relevant negative impact of metabolic complications on outcome, as well as the importance of nutrition to limit acute metabolic derangements. In particular, it has been clearly established that insulin resistance is a key mechanism behind developments of complications and delayed recovery in surgical patients [103]. Enhanced Recovery After Surgery (ERAS®) is a multimodal perioperative care pathway shown to lead to major improvements in outcomes in patients undergoing abdominal surgery; many ERAS elements reduce insulin resistance, as summarized in recent guidelines [104]. Nutritional intervention may focus on overcoming the traditional concept of fasting as well as on the general indication for immune-nutrition to reduce morbidity. Traditional surgical practices have emphasized the importance of fasting overnight before the procedure, but new research has exposed this protocol as harmful to recovery [105]. Studies had originally indicated that a fixed amount of mixed complex carbohydrates can be administered orally as a drink on the evening before surgery and in the morning up to two hours before anesthesia, resulting in lower insulin resistance following surgical stress with a positive impact on recovery and length of hospital stay [106]. In

### Table 1

Nutrition support guidelines and expert opinions for glycemic management in patients with diabetes mellitus types 1 and 2, T1D, type 1 diabetes; T2D, type 2 diabetes; IDF, International Diabetes Foundation; ADA, American Diabetes Association; NICE, National Institute for Health and Care Excellence; EASD, European Association for the Study of Diabetes; ESE, European Society of Endocrinology; SFAR, French Society of Anesthesia and Intensive Care; SRLF, Intensive Care Society (French language); SCCM, Society of Critical Care Medicine; ASPEN, American Society for Parenteral and Enteral Nutrition.

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<th>Region</th>
<th>Source</th>
<th>Title</th>
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<td>T1D</td>
<td>Europe, Worldwide</td>
<td>IDF</td>
<td>IDF 2011 Postmeal glucose guidelines [82]</td>
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<td></td>
<td>US</td>
<td>ADA</td>
<td>Standards of Medical Care in Diabetes-2016 [79]</td>
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<td></td>
<td>UK</td>
<td>NICE</td>
<td>NICE NG17. Type 1 diabetes in adults: diagnosis and management 2015 [83]</td>
</tr>
<tr>
<td>T2D</td>
<td>Europe and US</td>
<td>ADA</td>
<td>Management of hyperglycaemia in type 2 diabetes: a patient-centered approach 2012 [84]; Updated in 2015 [85]</td>
</tr>
<tr>
<td></td>
<td>US</td>
<td>EASD</td>
<td>Management of hyperglycaemia in type 2 diabetes: a patient-centered approach 2012 [84]; Updated in 2015 [85]</td>
</tr>
<tr>
<td></td>
<td>Europe and US</td>
<td>NICE</td>
<td>NICE NG28. Type 2 diabetes in adults: management 2015 [86]</td>
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### Table 2

Nutrition support guidelines and expert opinions for glycemic management in patients with stress metabolism or metabolic syndrome/obesity, ADA, American Diabetes Association; EASD, European Association for the Study of Diabetes; ESE, European Society of Endocrinology; SFAR, French Society of Anesthesia and Intensive Care; SRLF, Intensive Care Society (French language); SCCM, Society of Critical Care Medicine; ASPEN, American Society for Parenteral and Enteral Nutrition.

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<th>Patient population</th>
<th>Region</th>
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<td>SFAR, SRLF</td>
<td>International recommendations for glucose control in adult non-diabetic critically ill patients; Icha 2010 [88]</td>
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<td>ASPEN, SCCM</td>
<td>Guidelines for the provision and assessment of nutrition support therapy in the adult critically ill patient; McClave 2016 [89]</td>
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<td>US</td>
<td>Expert opinion</td>
<td>Management of hyperglycaemia; Mannon 2013 [90]</td>
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<td>Canada</td>
<td>Critical Care Nutrition</td>
<td>2015 Clinical Practice Guidelines [79]</td>
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<td>Europe</td>
<td>EASD</td>
<td>Evidence-based nutritional approaches to the treatment and prevention of diabetes mellitus; Mann 2004 [81]</td>
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general, as reflected in fasting guidelines for the past 20 years, evidence shows that clear fluids can be taken up to two hours before and that solids can be ingested up to six hours before surgery [107]. Efforts should be made to perform surgical procedures under the best attainable nutritional conditions, which may include nutritional support in combination with exercise before intervention [108]. Finally, the health care provider can prescribe pharmaco-nutrient support, including arginine and omega-3 fatty acids, to positively modulate immune response and limit inflammation to reduce morbidity, with particular regard to infectious complications. These can attenuate the inflammation and improve immune responses that may be impaired by surgery [105,107], thereby lessening the risk for infection as well as insulin resistance and hyperglycemia [109].

3.1.5. Glucose and nutritional support in critically ill patients

Glucose is the preferential physiological substrate for the production of energy in emergency conditions, including the acute phase of critical illness. However, in the intensive care unit (ICU), acute metabolic stress commonly leads to insulin resistance and hyperglycemia. Avoiding high blood glucose concentrations with insulin infusion improves the outcomes (mortality and morbidity) of ICU patients in some studies, but not in others [110–112]. The optimal glycemic target is hence undefined and could differ between patients, time from injury, and setting. A strong association has also been reported between high glucose variability as well as hypoglycemia and poor outcomes in the critically ill [113–116]. There is, however, consensus on the importance of effectively and closely monitoring plasma glucose during critical illness to reduce variability. To this aim, automated systems for glucose control and near-continuous glucose monitoring may provide more reliable tools to stabilize glycemia, and their implementation is therefore recommended.

Glucose control may become more problematic while implementing effective nutritional treatment in acute critical illness. Enteral nutrition (EN) support has been shown to increase hyperglycemia risk in hospitalized patients. However, this increase is less important than for parenteral nutrition, as enteral feeding triggers an elevation of insulin known as the incretin effect [117–119]. When EN cannot be tolerated and parenteral nutrition (PN) is necessary, the high dextrose delivered by standard PN formulas can further exacerbate the stress-related hyperglycemia, even in non-diabetic critically ill patients [91,92,96,115,120]. Thus both calorie and glucose administration, particularly in early phases of critical illness, also commonly lead to higher insulin requirements to control glycemia, with higher risk for glycemic variability and potential stimulation of lipogenesis. Additional care should be taken to minimize these risks.

Furthermore, it is difficult to determine the optimal carbohydrate amount to administer to critically ill patients for several reasons. These include difficulty in assessing energy requirements, altered enteral absorption, and impaired suppression of endogenous glucose production. One study compared glucose-based energy to lipid-based energy provision in ICU patients and found that glucose was associated with trends for hyperglycemia, higher insulin requirements, enhanced lipogenesis, and no improvement in protein sparing [121]. DSFs containing higher proportions of fat and modified carbohydrates have not been extensively assessed in ICU patients, but recent data suggest that the use of these formulas improves glycemic control, and, in at least one study, this was shown to provide clinical benefit [100]. Further studies should address interactions between glucose, lipid, and protein substrates, as well as the potential metabolic impact of higher utilization of lipid substrates for energy provision.

Guidelines for nutritional strategy and composition of nutritional supplements have been published for practical indications to achieve glucose control in critically ill patients [79,87–90] (Tables 1 and 2). Suggestions from these published studies include: 1) intervene with EN support as soon as possible to limit caloric debt [122]; 2) minimize glycemic variability in patients who must take PN, with a target blood glucose of 90–150 mg/dl (5–8 mM) [123]; 3) avoid hypoglycemia as a result of these approaches. For the avoidance of hyperglycemia, predisposing factors should be identified [124–126], and administration of intravenous insulin to critically ill patients should be restricted when appropriate.

Based on the above considerations and the impact of calories on glucose metabolism and plasma concentrations, the issue of limiting caloric administration to critically ill patients, particularly those with obesity, has been considered [89]. Moderately hypocaloric feeding with high-protein content aimed at counteracting protein catabolism and muscle loss has been suggested in recent guidelines for critically ill obese patients (22–25 kcal/kg of ideal body weight per day, 2 g/kg protein of body weight if BMI is less than 40 kg/m², or 2.5 if BMI is greater than 40 kg/m²) [89]. It should be pointed out that such recommendations are mainly aimed at minimizing metabolic abnormalities such as glucose variability and potential hyperlipidemia, rather than directly inducing weight loss. Additional research is desirable on optimal calorie provision for obese hospitalized patients with acute disease conditions requiring nutritional support.

4. Summary and conclusions

While carbohydrates, which provide glucose to the body to support metabolism, are crucial to the diet, inappropriate intake can lead to hyperglycemia, hypoglycemia, and glycemic fluctuations that are harmful to health outcomes (Fig. 2).

![Fig. 2. Consequences of glucose imbalance.](image-url)
Excess glucose ingestion interacts with the gut and its micro-biome and ultimately affects a number of organs including skeletal muscle, adipose tissue, and the liver. Excess glucose availability may induce expansion of adipose tissue and may favor ectopic fat deposition into liver and muscle tissues, which further exacerbates insulin resistance and glycemic imbalances. Insulin resistance is associated with, and can promote progression of, metabolic syndrome and eventually T2D, and it represents a factor contributing to hyperglycemia, glucose variability, and poor outcomes in the critically ill or those recovering from surgery.

Optimal nutritional support for patients with obesity and T2D should limit glucose provision, and plasma glucose should be critically low. In the surgical situation, preoperative fasting should be avoided as it should be assessed. It should be

The authors thank Dr. Cecilia Hofmann (C. Hofmann & Associates, Western Springs, IL, USA) for her capable assistance with writing, reference management, and editing the manuscript.

Conflict of interest
None.

Acknowledgment

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