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uring the first 24–48 hours of life, as normal neonates transition from intracellular to extracellular life, their plasma glucose (PG) concentrations are typically lower than later in life.1–3 Published guidelines for screening at-risk newborns and managing low PG concentrations in neonates focus on the immediate neonatal period, but do not address the diagnosis and management of disorders causing recurrent and prolonged hypoglycemia.4–6 Distinguishing between transitional neonatal glucose regulation in normal newborns and hypoglycemia that persists or occurs for the first time beyond the first 3 days of life is important for prompt diagnosis and effective treatment to avoid serious consequences, including seizures and permanent brain injury.

Moreover, the evaluation and management of pediatric hypoglycemia differ in several respects from that in adults, for whom guidelines were recently published.7 First, persistent hypoglycemia most often results from a congenital or genetic defect in regulating secretion of insulin, deficiency of cortisol and/or growth hormone, or defects in the metabolism of glucose, glycogen, and fatty acids. Second, it may be difficult to identify and distinguish newborn infants with a persistent hypoglycemia disorder from those with transitional low glucose levels in the initial 48 hours of life, as detailed in the separate document on transitional neonatal hypoglycemia prepared by our committee.8 Third, the first few months of life are the most vulnerable period for developmental disability, which occurs in ~25%–50% of children with congenital hyperinsulinism. Early recognition and treatment are crucial for preventing these sequelae.8–10

To address these deficiencies, the Pediatric Endocrine Society convened an expert panel of pediatric endocrinologists and neonatologists to develop guidelines for managing hypoglycemia in neonates, infants, and children, but excluding children with diabetes. The goals of these guidelines are to help physicians recognize persistent hypoglycemia disorders, guide their expeditious diagnosis and effective treatment, and prevent brain damage in at-risk babies.

Methods

Evidence Retrieval and Rating

The committee searched for existing evidence synthesis reports, systematic reviews, and meta-analyses. The committee also evaluated guidelines published by the Endocrine Society, American Academy of Pediatrics, Canadian Pediatric Society, and others, and reviewed their bibliographies.3–7 Committee members identified additional individual studies.

The committee adopted the framework of the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) Working Group,11 in which guideline developers rate their confidence in the evidence as very low (+000), low (++00), moderate (+++0), or high (++++). Randomized trials start as high, and observational studies start as low.11

Grading the Strength of Recommendations

The guideline developers considered the quality of the evidence. They also considered the balance between benefits and harms, patients’ values and preferences, cost and resource utilization, and other societal and contextual factors, such as availability of technology and health services and implementation barriers. The recommendations according to the GRADE framework are either strong (GRADE 1), stated as “we recommend,” or weak (GRADE 2), stated as “we suggest.”
Section 1: Which Neonates, Infants, and Children to Evaluate for Hypoglycemia

1.1. For children who are able to communicate their symptoms, we recommend evaluation and management only of those in whom Whipple’s triad (see below) is documented. GRADE 1++++.

1.2. For infants and younger children who are unable to reliably communicate symptoms, we suggest evaluation and management only of those whose PG concentrations are documented by laboratory quality assays to be below the normal threshold for neurogenic responses (<60 mg/dL [3.3 mmol/L]). GRADE 2+++.

1.3. For those neonates who are suspected to be at high risk of having a persistent hypoglycemia disorder, we suggest evaluation when the infant is ≥48 hours of age so that the period of transitional glucose regulation has passed and persistent hypoglycemia may be excluded before discharge home. GRADE 2++0.

Clinical Definition of Hypoglycemia

Clinical hypoglycemia is defined as a PG concentration low enough to cause symptoms and/or signs of impaired brain function. Hypoglycemia may be difficult to recognize because the signs and symptoms are nonspecific, and a single low PG concentration may be an artifact. For these reasons, guidelines in adults emphasize the value of Whipple’s triad for confirming hypoglycemia: symptoms and/or signs consistent with hypoglycemia, a documented low PG concentration, and relief of signs/symptoms when PG concentration is restored to normal. Young infants and children often cannot dependably recognize and/or communicate their symptoms; therefore, recognition of hypoglycemia may require confirmation by repeated measurements of PG concentration and formal testing. Nevertheless, suspected hypoglycemia should be treated promptly to avoid potential adverse consequences.

Hypoglycemia cannot be defined as a specific PG concentration, because: (1) thresholds for specific brain responses to hypoglycemia occur across a range of PG concentrations, and these thresholds can be altered by the presence of alternative fuels, such as ketones, and by recent antecedent hypoglycemia; (2) it is not possible to identify a single PG value that causes brain injury, and the extent of injury is influenced by other factors, such as duration and degree of hypoglycemia; and (3) potential artifacts and technical factors that lead to inaccuracies in glucose determination may complicate the interpretation of any single PG value.

Symptoms of Hypoglycemia

The symptoms of hypoglycemia reflect responses of the brain to glucose deprivation and have been well delineated in adults. Neurogenic (autonomic) symptoms result from the perception of physiological changes caused by the sympathetic nervous discharge triggered by hypoglycemia; these include adrenergic responses (eg, palpitations, tremor, anxiety) and cholinergic responses (eg, sweating, hunger, paresthesias). Neuroglycopenic signs and symptoms, including confusion, coma, and seizures, are caused by brain dysfunction resulting from a deficient glucose supply to sustain brain energy metabolism. Awareness of hypoglycemia depends chiefly on perception of the central and peripheral effects of neurogenic (as opposed to neuroglycopenic) responses to hypoglycemia. Brain glucose utilization becomes limited at a PG concentration of approximately 55-65 mg/dL (3.0-3.6 mmol/L). Neurogenic symptoms are perceived at a PG concentration <55 mg/dL (<3.0 mmol/L), which in older children and adults triggers a search for food or assistance, an important defense against hypoglycemia. Cognitive function is impaired (neuroglycopenia) at a PG concentration <50 mg/dL (<2.8 mmol/L).

Glucose Utilization

The adult brain accounts for more than one-half of total glucose consumption. Because of their disproportionately larger brain size relative to body mass, infants and young children have a 2- to 3-fold higher glucose utilization rate (4-6 mg/kg/min) per kilogram of body weight compared with adults. Although the brain has an obligate requirement for glucose, it also can use plasma ketones and lactate as energy sources if the concentrations of these substances are sufficiently elevated. However, in hypoketotic conditions, such as hyperinsulinism or fatty acid oxidation disorders, ketones and lactate are not available in sufficiently high concentrations to substitute for glucose, and the risk of brain energy failure is greater.

Neuroendocrine Defenses against Hypoglycemia

In normal individuals, the maintenance of normal PG concentrations is highly protected. The first defense is suppression of insulin secretion when PG concentration falls below the normal postabsorptive mean of ~85 mg/dL (4.9 mmol/L). Further reduction of PG to 65-70 mg/dL (3.6-3.9 mmol/L) elicits glucagon secretion and activation of the sympathoadrenal system (reflected by increased epinephrine concentration), which increases glucose release from liver glycogen stores to raise the PG concentration. At a PG concentration <65 mg/dL (3.6 mmol/L), levels of plasma cortisol and growth hormone, important for maintenance of glucose during prolonged fasting, increase as well. Because the brain has only a few minutes worth of stored fuel reserves in the form of glycogen, interruption of glucose delivery can have devastating consequences. Whereas recovery from brief periods of hypoglycemia is usually complete, severe and prolonged hypoglycemia can cause permanent brain injury.

Metabolic Defenses against Hypoglycemia

In the postabsorptive phase, the liver supplies the brain and other tissues with glucose by releasing glucose from the breakdown of stored glycogen and by gluconeogenesis, principally from gluconeogenic amino acids, such as...
alanine, and recycled lactate. With longer fasting and further suppression of insulin secretion, glucose utilization is restricted to the brain and a few glycolytic tissues, such as erythrocytes. Adipose tissue lipolysis releases glycerol, a gluconeogenic substrate, and free fatty acids (FFAs) that can replace glucose as an energy substrate in skeletal and heart muscle, but not in brain. FFAs are also converted by the liver to beta-hydroxybutyrate (BOHB) and acetoacetate for use by the brain. BOHB is the predominant ketoadic acid, and its plasma level serves as a measure of ketogenesis. As ketoacid concentrations rise, they can partly support the brain’s energy needs. The changes in fuel metabolism during fasting in normal neonates after 2-3 days of age and in infants and children do not differ substantially from those in adults, except that PG concentrations decrease more rapidly and hyperketonemia develops sooner, because of the energy needs of their relatively larger brains. Measurement of BOHB, FFA, and lactate at the time of hypoglycemia provides important information for diagnosing the cause of hypoglycemia (Figure).

Altered Hypoglycemia Awareness
Previous exposure to an episode of hypoglycemia can blunt, and repeated episodes can eliminate, neurogenic responses to subsequent hypoglycemic episodes. This leads to reduced or absent awareness of hypoglycemia and impairs hepatic glucose release, perpetuating hypoglycemia. This combination of events has been termed hypoglycemia-associated autonomic failure (HAAF). HAAF can persist for >24 hours after a single episode of hypoglycemia or even longer after repeated episodes of hypoglycemia. A similar impairment in neuroendocrine responses to hypoglycemia also occurs during sleep and exercise. Thus, exposure to recurrent hypoglycemia can shift the usual glucose threshold for recognition of neurogenic symptoms of 55 mg/dL (3.0 mmol/L) to a lower level. Although previous exposure to hypoglycemia lowers the glucose threshold for neurogenic responses, the threshold for neuroglycopenic symptoms is not altered acutely; whether adaptation occurs with repeated exposure to hypoglycemia is unknown. Features of HAAF have been demonstrated in infants as young as age 10-13 weeks.

Potential Artifacts in Measurements of PG Concentration
To diagnose hypoglycemia, PG concentration should be measured using a clinical laboratory method. Important considerations are that whole blood glucose values are 15% lower than PG concentrations, and that because of red cell glycolysis, delays in processing and assaying glucose can reduce the glucose concentration by up to 6 mg/dL/hour (0.3 mmol/L/hour). Point-of-care meters provide a convenient screening method for detecting hypoglycemia,

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**Figure.** Algorithm showing how the major categories of hypoglycemia can be determined with information from the critical sample. GH, growth hormone.
but their accuracy is limited to approximately ±10-15 mg/dL (0.6-0.8 mmol/L) in the range of hypoglycemia. Therefore, before establishing a diagnosis of hypoglycemia in neonates, infants, and children, it is essential to confirm low PG concentration using a clinical laboratory method.

**Normal PG Concentrations in Neonates Aged >48 Hours, Infants, and Children**

After the first 48 hours of life, PG concentration and the physiology of glucose homeostasis do not differ to any great extent with age. Mean PG concentration in the postabsorptive state in normal neonates after ~2 days of age and in infants and children does not differ from that in adults (70-100 mg/dL [3.9-5.5 mmol/L])

**PG Concentrations in Neonates Aged <48 Hours**

In normal newborn infants, PG concentration commonly decreases immediately after birth to levels below those in older infants and children. The interpretation and response to PG concentration during the first 2 days of life have been controversial. Whether the brain of newborn infants has greater or lesser susceptibility to hypoglycemic injury is controversial as well. As discussed previously, the committee’s review of available data on transitional neonatal hypoglycemia in normal newborns (ie, hypoketonemic hypoglycemia with inappropriate large glycemic responses to glucagon or epinephrine stimulation) suggests that it is a mild and transient form of hyperinsulinism in which the mean PG threshold for suppression of insulin secretion is ~55-65 mg/dL (3.0-3.6 mmol/L) shortly after birth, compared with ~80-85 mg/dL (4.4-4.7 mmol/L) in older infants, children, and adults. As the glucose stimulated-insulin secretion mechanism matures, mean PG concentration in normal newborns increases and by 72 hours of age is similar to those in older infants and children; therefore, the standards for normal neonates must never be extrapolated beyond 2-3 days after birth. Because of the difficulty in distinguishing a suspected persistent hypoglycemia disorder from transitional neonatal glucose concentrations during the first 48 hours of life, we suggest delaying diagnostic evaluations until 2-3 days after birth.

**Neonates Aged >48 Hours Who May Be at Risk for Persistent Hypoglycemia Disorders**

Some neonates can be identified by various clinical features as being at high risk for severe hypoglycemia during the first 48 hours after delivery, and a subset of those neonates are also at increased risk for persistent hypoglycemia beyond 48 hours of life (Table). These include not only the rare infants with genetic hypoglycemia disorders, such as congenital hyperinsulinism or hypopituitarism, but also those with relatively more common prolonged neonatal hyperinsulinism (also referred to as perinatal stress hyperinsulinism) associated with birth asphyxia, intrauterine growth restriction, or toxemia.

To provide a practical guide and in the light of scarce evidence in this area, the committee used a consensus process to make the following technical suggestions:

For neonates with a known risk of a genetic or other persistent form of hypoglycemia (eg, congenital hyperinsulinism, glucose-6-phosphatase deficiency, fatty acid oxidation disorder), consultation with a specialist should be considered before planning discharge from the nursery. Diagnostic tests may be available to exclude the possibility of a specific disorder within a few weeks (eg, genetic mutation analysis, plasma acyl-carnitine profile). In these cases, a “safety” fasting test

<table>
<thead>
<tr>
<th>Table. Recognizing and managing neonates at increased risk for a persistent hypoglycemia disorder</th>
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<tbody>
<tr>
<td><strong>Neonates at increased risk of hypoglycemia and require glucose screening:</strong></td>
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<tr>
<td>1. Symptoms of hypoglycemia</td>
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<tr>
<td>2. Large for gestational age (even without maternal diabetes)</td>
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<tr>
<td>3. Perinatal stress</td>
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<tr>
<td>a. Birth asphyxia/ischemia; cesarean delivery for fetal distress</td>
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<tr>
<td>b. Maternal preeclampsia/eclampsia or hypertension</td>
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<tr>
<td>c. Intrauterine growth restriction (small for gestational age)</td>
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<tr>
<td>d. Meconium aspiration syndrome, erythroblastosis fetalis, polycythemia, hyperthermia</td>
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<tr>
<td>4. Premature/postmature delivery</td>
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<tr>
<td>5. Infant of diabetic mother</td>
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<tr>
<td>6. Family history of a genetic form of hypoglycemia</td>
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<tr>
<td>7. Congenital syndromes (eg, Beckwith-Wiedemann), abnormal physical features (eg, midline facial malformations, microphthalmus)</td>
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<tr>
<td><strong>Neonates in whom to exclude persistent hypoglycemia before discharge:</strong></td>
</tr>
<tr>
<td>1. Severe hypoglycemia (eg, episode of symptomatic hypoglycemia or need for IV dextrose to treat hypoglycemia)</td>
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<tr>
<td>2. Inability to consistently maintain preprandial PG concentration &gt;50 mg/dL up to 48 hours of age and &gt;60 mg/dL after 48 hours of age</td>
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<tr>
<td>3. Family history of a genetic form of hypoglycemia</td>
</tr>
<tr>
<td>4. Congenital syndromes (eg, Beckwith-Wiedemann), abnormal physical features (eg, midline facial malformations, microphthalmus)</td>
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</table>
should be done before discharge from the nursery to ensure that PG concentration can be maintained above 70 mg/dL if a feeding is missed (ie, a minimum of 6-8 hours). If no definitive test is available (which is often the case with hyperinsulinism), then the fasting test should be extended to include diagnostic features of the suspected disorder following the paradigm shown in the Figure.

For at-risk neonates without a suspected persistent hypoglycemia disorder and in whom hypoglycemia is considered likely to resolve within a short time, a “safety” fast of 6-8 hours should be considered before discharge, to determine whether a PG concentration >60 mg/dL can be maintained, or whether additional management or investigations may be required.

**Values**

Hypoglycemia disorders in older infants and children are uncommon. Thus, the recommendation to require the presence of Whipple’s triad before undertaking an evaluation avoids exposing patients without a specific disorder to unnecessary procedures, potential risks, and expense without any likely benefit. For neonates, infants, and younger children in whom Whipple’s triad cannot be applied, unnecessary investigation can be avoided by confirming a PG concentration at or below the range at which neurogenic symptoms usually occur. The recommendations regarding which neonates to evaluate for hypoglycemia focus on those high-risk neonates who require close monitoring and/or intervention, such as intravenous (IV) dextrose, for treatment of hypoglycemia (Table). They take into account that transitional hypoglycemia in normal newborns should be completely resolved before the usual time for discharge from the nursery at 2-3 days. The committee emphasizes the necessity of carefully excluding persistent hypoglycemia disorders before discharge to home in the small subset of high-risk neonates (Table) to ensure recognition and facilitate treatment.

### Section 2. Workup/Investigation of Persistent Hypoglycemia

**2.1.** We recommend that investigations be carried out to diagnose the underlying mechanism of persistent hypoglycemia disorders to provide specific management. GRADE 1++++.

In patients with hypoglycemia, a thorough history should include the episode’s timing and its relationship to food, birth weight, gestational age, and family history. Physical examination should include looking for evidence of hypopituitarism (eg, micropenis or cleft lip or palate, short stature), glycochenosis (eg, hepatomegaly), adrenal insufficiency (eg, recurrent abdominal pain, hyperpigmentation, anorexia, weight loss), or Beckwith-Wiedemann syndrome (eg, omphalocele, hemihypertrophy, macroGLOSSIA).

Whenever possible, specimens (a “critical sample”) for identifying the etiology of hypoglycemia should be obtained at the time of spontaneous presentation and before treatment that may rapidly alter the levels of BOHB, FFA, and insulin. Assays for PG, bicarbonate, BOHB, and lactate are readily available and useful for distinguishing categories of hypoglycemia disorders (Figure). Extra plasma can be held in reserve for specific tests (eg, plasma insulin, FFA, and C-peptide for suspected hyperinsulinism; plasma total and free carnitine and acyl-carnitine profile for a suspected disorder of fatty acid oxidation). Laboratory reference ranges for some of these tests (eg, insulin, BOHB) may be inappropriate for interpreting values obtained during childhood hypoglycemia.

In the absence of a “critical sample,” a provocative fasting test is the most informative method for identifying the etiology of hypoglycemia disorders.20,32 The duration of fasting and the specimens obtained should be tailored to the suspected diagnosis. With few exceptions, blood tests performed while glucose concentrations are normal are not informative. For safety, a fasting test should be done with frequent monitoring of vital signs, PG, and BOHB concentrations. A PG concentration of 50 mg/dL (2.8 mmol/L) is sufficiently low to elicit the metabolic and neuroendocrine responses required for diagnosis. After appropriate specimens have been obtained, the fast may be terminated. The fast also can be stopped earlier for signs or symptoms of distress or a plasma BOHB measurement of >2.5 mmol/L.

For suspected hyperinsulinism, the fasting test can be terminated when the PG concentration is <50 mg/dL (<2.8 mmol/L) with administration of glucagon (1 mg IV, intramuscularly, or subcutaneously) to evaluate the glycemic response. An exaggerated glycemic response (>30 mg/dL [>1.7 mmol/L]) is nearly pathognomonic of hyperinsulinism.33 Because plasma insulin concentration is sometimes not above the lower limit of detection,34 it is important to include the following tests when assessing the possibility of hypoglycemia due to hyperinsulinism: plasma BOHB and FFA (both inappropriately low; BOHB <1.5 mmol/L, [<15 mg/dL] and FFA <1.0-1.5 mmol/L [<28-42 mg/dL]), and an increased glycemic response to glucagon. Based on the suspected diagnosis, additional tests performed on specimens obtained at the time that fasting is terminated should be considered, such as growth hormone, cortisol, total and free carnitine, acyl-carnitine profile, C-peptide, proinsulin, and drug screening.

Adrenal insufficiency, congenital or acquired, may manifest with hypoglycemia. Simultaneous measurement of plasma adrenocorticotropic hormone and cortisol is recommended if primary adrenal insufficiency is suspected. An adrenocorticotropic hormone stimulation test may be needed to confirm suspected secondary or tertiary adrenal insufficiency.

The possibility of drug administration (accidental or surreptitious) or a toxin may need to be considered. Specimens of plasma and urine obtained at the time of an acute episode may be informative. High plasma insulin level but low C-peptide level during hypoglycemia suggest exogenous insulin administration. The drugs most commonly associated with hypoglycemia include insulin, sulphonylureas, alcohol,
beta-blockers, and salicylates. Not all commercially available insulin assays measure the insulin analogues; thus, the finding of low insulin and C-peptide levels at time of hypoglycemia might not rule out surreptitious insulin administration. (Consultation with the testing laboratory should be considered if clinical suspicion is high.)

Postprandial hypoglycemia in infants and children is usually a complication of fundoplication for gastroesophageal reflux (incidence 25%-30%) or gastric bypass bariatric surgery for morbid obesity. In these patients, a meal may trigger hypoglycemia 1-3 hours later. Postprandial hypoglycemia is often preceded by exaggerated glucose, glucagon-like peptide-1, and insulin responses, which can be detected on an oral glucose or mixed-meal tolerance test.

### Values
Failure to investigate a neonate, infant, or child with suspected hypoglycemia increases the risk of delaying a definitive diagnosis and instituting effective treatment. Expediously identifying the specific cause of hypoglycemia, as outlined above, will enable prompt institution of appropriate treatment and decrease the risk of permanent brain injury from persistent and recurrent severe hypoglycemia. The decision to subject an infant or child to a diagnostic investigation for suspected hypoglycemia, which includes a monitored assessment of fasting adaptation that may expose the patient to the risk of another episode of hypoglycemia, places a higher value on achieving diagnostic certainty and a lower value on avoiding the discomfort, inconvenience, and cost of such procedures.

### Section 3. Management of Neonates, Infants, and Children with a Persistent Hypoglycemia Disorder

3.1. For neonates with a suspected congenital hypoglycemia disorder and older infants and children with a confirmed hypoglycemia disorder, we recommend that the goal of treatment be to maintain a PG concentration >70 mg/dL (3.9 mmol/L). GRADE 1++.00

3.2. For high-risk neonates without a suspected congenital hypoglycemia disorder, we suggest the goal of treatment be to maintain a PG concentration >50 mg/dL (>2.8 mmol/L) for those aged <48 hours and >60 mg/dL (>3.3 mmol/L) for those aged >48 hours. GRADE 2+00.

3.3. We recommend an individualized approach to management with treatment tailored to the specific disorder, taking into account patient safety and family preferences. Ungraded best practice statement.

### Neonates, Infants, and Children with Hypoglycemia Disorders

Neurogenic and neuroglycopenic symptoms usually occur when the PG concentration decreases to 50-70 mg/dL (2.8-3.9 mmol/L). Recurrent PG levels in this range may result in the development of HAAF, which increases the risk of subsequent hypoglycemia and attenuates its recognition. Therefore, treatment targets are generally aimed at avoiding activation of neuroendocrine responses and HAAF by maintaining PG concentration within the normal range of 70-100 mg/dL (3.9-5.6 mmol/L). For this reason, we recommend that the same goal for treatment of adults and children with diabetes (ie, PG >70 mg/dL [>3.9 mmol/L]) be used for neonates with a confirmed hypoglycemia disorder and for infants and children with a persistent hypoglycemia disorder.

For disorders such as hyperinsulinism, the aim is to prevent recurrent hypoglycemia that increases the risk of subsequent, possibly unrecognized, hypoglycemic episodes. For disorders such as defects in glycogen metabolism and gluconeogenesis, maintenance of PG concentration in the normal range prevents metabolic acidosis and growth failure, and possibly the development of long-term complications.

Any episode of severe symptomatic hypoglycemia should be rapidly corrected with IV dextrose infusion. The initial dose is 200 mg/kg, followed by infusion of 10% dextrose at a maintenance rate for age. In cases of hyperinsulinism, glucagon can be expected to raise PG concentration to normal or above within 10-15 minutes and to maintain that concentration for at least 1 hour. Doses of 0.5-1.0 mg (independent of weight), given IV, intramuscularly, or subcutaneously, are usually effective; lower doses (0.03 mg/kg) may carry less risk of transient nausea and vomiting, but may be ineffective unless given IV.

Long-term therapy for hypoglycemia disorders should be based on the specific etiology of the disorder, in consultation with a physician experienced in the diagnosis and management of hypoglycemia in infants and children and with careful consideration of patient and family preferences. Medications are available for some disorders, such as hyperinsulinism, and for cortisol and growth hormone deficiency. Surgery may be necessary in some children with hyperinsulinism who are unable to maintain PG concentration in a safe range through medical therapy. Nutritional therapy is the cornerstone of treatment for some disorders, such as disorders of glycogen metabolism or hereditary fructose intolerance. Some milder disorders may be treated adequately by avoidance of prolonged fasting. Patients diagnosed with ketotic hypoglycemia who have recurrent episodes should be reevaluated for other specific disorders, such as glycogen synthase deficiency (GSD 0), GSD type VI, GSD type IX (especially in males), and MCT1 gene deficiency. Nonspecific treatment with glucocorticoids for neonates with hypoglycemia is discouraged.

### High-Risk Neonates without a Suspected Congenital Hypoglycemia Disorder

The mean PG concentration in normal newborns during the first hours after birth (~55-60 mg/dL (~3 mmol/L) is similar to the threshold for neurogenic symptoms of hypoglycemia in older children and adults, but above the threshold for neuroglycopenic symptoms. This appears to
be tolerated during the first few hours after birth. In normal newborns, the mean PG concentration rises to >70 mg/dL (3.9 mmol/L) after 48 hours of age. This is similar to the mean glucose concentration in older children and adults. There is no evidence that the concentration at which PG becomes limiting for brain function is different in neonates than in older children and adults. Of concern, as noted previously, plasma ketone levels are suppressed during hypoglycemia in high-risk neonates, making them particularly vulnerable to hypoglycemia-induced brain damage, and there is no assurance the plasma lactate level will be sufficiently elevated to compensate for low glucose.27,28,30,39

Given the absence of evidence on the short-term or long-term consequences of different treatment targets, the committee focused on evidence related to physiology, etiology, and mechanism, and balanced the risks and benefits of interventions. The committee also focused on postnatal age, because ongoing hypoglycemia beyond the first 48 hours of life (and particularly beyond the first week) increases the concern for an underlying hypoglycemia disorder, such as prolonged neonatal hyperinsulinism, or a genetic disorder. The committee’s consensus was that during the first 48 hours of life, for a high-risk neonate without a suspected congenital hypoglycemia disorder and on normal feedings, a safe target PG concentration should be close to the mean for healthy newborns on the first day of life and above the threshold for neuroglycopenic symptoms (>50 mg/dL [2.8 mmol/L]). The committee recommended raising the glucose target after 48 hours of age (>60 mg/dL [3.3 mmol/L]) to above the threshold for neurogenic symptoms and close to the target for older infants and children, because of the increased concern for an underlying hypoglycemia disorder. The suggested target glucose concentrations were considered adequate while assessing whether hypoglycemia will resolve over time. Because at-risk neonates who require interventions beyond normal feedings, such as IV dextrose, to treat hypoglycemia in the first 48 hours of life are likely to have more severe and prolonged hypoglycemia, the initial treatment target should be above the thresholds for both neurogenic and neuroglycopenic symptoms (>60 mg/dL [3.3 mmol/L]). For neonates who require dextrose infusion, transitioning to normal feedings alone can be attempted once PG concentration is stabilized at >60 mg/dL (>3.3 mmol/L).

The committee’s consensus to accept a lower PG target in high-risk neonates without a suspected congenital hypoglycemia disorder (Section 3.2) than in those with a suspected congenital hypoglycemia disorder (Section 3.1) was based on balancing the risks of intervention compared with a brief period of undertreatment in this group. Higher treatment targets (Section 3.1) should be considered for those with a suspected genetic hypoglycemia disorder and for symptomatic neonates, because the risks of undertreatment outweigh those of overtreatment in these patients.

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