**Use of Intensive Insulin Therapy for the Management of Glycemic Control in Hospitalized Patients: A Clinical Practice Guideline From the American College of Physicians**

Amir Qaseem, MD, PhD, MHA; Linda L. Humphrey, MD, MPH; Roger Chou, MD; Vincenza Snow, MD; and Paul Shekelle, MD, PhD, for the Clinical Guidelines Committee of the American College of Physicians*

**Description:** The American College of Physicians (ACP) developed this guideline to present the evidence for the link between the use of intensive insulin therapy to achieve different glycemic targets and health outcomes in hospitalized patients with or without diabetes mellitus.

**Methods:** Published literature on this topic was identified by using MEDLINE and the Cochrane Library. Additional articles were obtained from systematic reviews and the reference lists of pertinent studies, reviews, and editorials, as well as by consulting experts; unpublished studies on ClinicalTrials.gov were also identified. The literature search included studies published from 1950 through March 2009. Searches were limited to English-language publications. The primary outcomes of interest were short-term mortality and hypoglycemia. This guideline grades the evidence and recommendations by using the ACP clinical practice guidelines grading system.

**Recommendation 1:** ACP recommends not using intensive insulin therapy to strictly control blood glucose in non–surgical intensive care unit (SICU)/medical intensive care unit (MICU) patients with or without diabetes mellitus (Grade: strong recommendation, moderate-quality evidence).

**Recommendation 2:** ACP recommends not using intensive insulin therapy to normalize blood glucose in SICU/MICU patients with or without diabetes mellitus (Grade: strong recommendation, high-quality evidence).

**Recommendation 3:** ACP recommends a target blood glucose level of 7.8 to 11.1 mmol/L (140 to 200 mg/dL) if insulin therapy is used in SICU/MICU patients (Grade: weak recommendation, moderate-quality evidence).

Hyperglycemia is a common finding among medical and surgical patients with or without known diabetes during hospital admission (1, 2). Although the prevalence of hyperglycemia in hospitalized patients is not known with certainty, it is estimated to be around 40% (3). Poorly controlled hyperglycemia is associated with increased morbidity, mortality, and costs (4). Hyperglycemia is associated with poor immune response, increased cardiovascular events, thrombosis, inflammatory changes, delayed healing, and other problems (5). Achieving tight glycemic control safely in inpatients is labor intensive and often requires coordination of efforts involving a multidisciplinary team in a hospital setting (4). Most of the evidence on how to best achieve target blood glucose levels centers around the use of intensive insulin protocols.

The purpose of this American College of Physicians (ACP) guideline is to address the management of hyperglycemia and evaluate the benefits and harms associated with the use of intensive insulin therapy (IIT) to achieve tight glycemic control in hospitalized patients with or without diabetes mellitus. We defined “IIT” as use of intravenous insulin to achieve targeted blood glucose level with frequent blood glucose testing and adjustment of insulin doses. In intensive care unit (ICU) settings, the usual target of IIT is normoglycemia (blood glucose level, 4.4 to 6.1 mmol/L [80 to 110 mg/dL]), whereas targets in non-ICU settings have been more variable (ranging from normoglycemia to <11.1 mmol/L [<200 mg/dL]).

The target audience for this guideline includes all clinicians, and the target patient population comprises all adults with hyperglycemia in a hospital setting. These recommendations are based on a systematic evidence review by Kansagara and colleagues (6), from an
METHODS

The objective of this guideline is to present the evidence for the following questions:

1. Does the use of IIT to achieve tight glycemic control compared with less tight glycemic control improve important health outcomes in the following settings or patient populations: surgical intensive care unit (SICU), medical intensive care unit (MICU), general surgical ward, general medicine ward, patients with myocardial infarction or acute stroke, and patients in the perioperative setting?

2. What are the harms of strict glycemic control in the above subpopulations?

The databases used for the literature search were MEDLINE and the Cochrane Database of Systematic Reviews; the search included studies published from database inception through January 2010. The literature search was supplemented by reviews of reference lists, suggestions from consulting experts, and searches on ClinicalTrials.gov for unpublished studies. Each article was reviewed by using the eligibility criteria outlined in the systematic review (6). Eligible articles were published in English and provided primary data relevant to the use of IIT in hospitalized patients. Studies evaluating fixed-dose insulin infusions, including trials of fixed-dose glucose–insulin–potassium infusions, were excluded. The primary outcome of interest was short-term mortality (preferential order: 28-day mortality, hospital mortality, ICU mortality). The major harm of interest was the rate of hypoglycemia, including effects of hypoglycemia on clinical outcomes and length of hospitalization. The quality of each study was rated as good, fair, or poor on the basis of 1) the comparability of treatment groups; 2) the adequacy of randomization; 3) whether treatment allocation was concealed; 4) whether eligibility criteria were specified; 5) whether patients, care providers, and outcome assessors were blinded; 6) whether the analysis was done on an intention-to-treat basis, conducted with postrandomization exclusions, or had extensive or differential loss to follow-up; and 7) whether clearly defined interventions and reliable outcome measurement were used. Given the importance of glucose control and hypoglycemia in assessing the effectiveness and safety of IIT, studies that did not fully report glucose levels achieved or overall hypoglycemia rates were rated as poor quality.

Three investigators reviewed the abstracts of citations identified from literature searches. When reviewers disagreed about the quality rating, consensus was reached through discussion with all authors. Details of the methods for the evidence review are provided in the evidence review (6).

BENEFITS OF IIT FOR IMPROVING HEALTH OUTCOMES

Mortality

A meta-analysis of 21 trials (14 768 participants) found no benefit associated with IIT on short-term mortality (28-day, hospital, or ICU mortality) (relative risk [RR], 1.1 [95% CI, 0.99 to 1.1]) (6). Although the trials differed with regard to target glucose levels for IIT and control groups, achieved glucose levels, IIT protocols, and medical setting (for example, ICU vs. non-ICU), there was no statistical heterogeneity ($I^2 = 0\%$). Results were similar when trials were stratified according to blood glucose level achieved, the percentage of diabetic patients, definition of short-term mortality, or study quality. Meta-analysis also showed no benefit associated with IIT for 90- or 180-day mortality (13 trials; RR, 1.1 [CI 0.99 to 1.1]; $I^2 = 0\%$). Results in specific patient subgroups are described below.

ICUs

MICUs

Evidence from 5 fair-quality trials (7–11) and 1 poor-quality trial (12) of IIT (target glucose levels of normoglycemia [4.4 to 6.1 mmol/L [80 to 110 mg/dL]] compared with control values ranging from 7.8 to 11.1 mmol/L [140 to 200 mg/dL]) consistently found no mortality benefit associated with normoglycemia as a target. The overall quality of evidence that IIT with normoglycemia as a target in patients in the MICU does not improve mortality was rated as high.

SICUs

Three fair-quality trials (7, 9, 13) and 2 poor-quality trials (14, 15) of IIT (target glucose levels of 4.4 to 8.3 mmol/L [80 to 150 mg/dL] vs. control values ranging from...
10.0 to 12.2 mmol/L [180 to 220 mg/dL]) conducted in SICUs showed no benefit of IIT on mortality (7, 9, 13–15). The largest trial (2232 patients in the surgical subgroup) was the recent multicenter NICE-SUGAR (Normoglycemia in Intensive Care Evaluation—Survival Using Glucose Algorithm Regulation) study (9), a fair-quality study with target glucose levels of 4.4 to 6.0 mmol/L (80 to 108 mg/dL) that showed an increase in mortality compared with a target less than 10.0 mmol/L (<180 mg/dL) (RR, 1.31 [CI, 1.07 to 1.61]). However, not all of the trials found no benefit. A large (1600 participants), fair-quality study showed a statistically significant reduction in all-cause ICU mortality in the IIT group (morning blood glucose level, 4.4 to 6.1 mmol/L [80 to 110 mg/dL]) compared with conventional therapy (morning blood glucose level, 10.0 to 11.1 mmol/L [180 to 200 mg/dL]; mortality rates were 4.6% and 8.0%, respectively (RR, 0.58 [CI, 0.38 to 0.78]) (16). The overall quality of evidence that IIT targeted at normoglycemia in patients in the SICU does not improve mortality was rated as moderate.

**Mixed MICU and SICU Populations**

Five fair-quality trials that included mixed MICU and SICU populations (glucose target for IIT ranging from 4.0 to 6.1 mmol/L [72 to 110 mg/dL] compared with control ranging from 7.8 to 11.1 mmol/L [140 to 200 mg/dL]) did not demonstrate an overall mortality benefit of IIT (7, 9, 17, 18; Mackenzie I, Blunt M, Ingle S, Palmer C. GLYcaemic Control and Outcome in GENeral Intensive Care. Unpublished report). The largest trial (6104 participants), the NICE-SUGAR study, showed that IIT with target glucose levels of 4.4 to 6.0 mmol/L (80 to 108 mg/dL) was associated with an increase in 90-day mortality compared with a target level less than 10.0 mmol/L (<180 mg/dL) (RR, 1.14 [CI, 1.02 to 1.28]); there was approximately 1 excess death per 39 patients treated with IIT (9). The overall quality of evidence that IIT targeted at normoglycemia in mixed MICU/SICU populations does not improve mortality was rated as high.

**General Medicine Ward**

No studies evaluated IIT in patients on the general medical ward.

**Patients With Myocardial Infarction**

Evidence from 3 fair-quality studies (19–21) and 2 poor-quality studies (22, 23) showed no reduction in mortality among patients with myocardial infarction who received IIT and adjustable-dose glucose–insulin–potassium infusions (target glucose levels ranging from 4.0 to 11.0 mmol/L [72 to 198 mg/dL] vs. unspecified target levels in control groups). One fair-quality trial (620 participants) that compared IIT (target glucose levels ranging from 7.0 to 11.0 mmol/L [126 to 198 mg/dL]) with long-term post-discharge insulin therapy found a mortality reduction at 1 year (18.6% vs. 26.1%, respectively; RR, 0.69 [CI, 0.49 to 0.96]; P = 0.027), but it was not possible to determine whether the results were due to IIT or the use of insulin after discharge (24). In addition, this trial was published about 10 years before the other trials, and given changes over time in management of myocardial infarction, its current applicability may be limited.

Variations in trial design, glucose level achieved, and concomitant therapy for myocardial infarction limit the strength of conclusions that can be drawn from these studies. The overall quality of evidence for IIT to achieve target glucose levels of 4.0 to 11.0 mmol/L (72 to 198 mg/dL) on mortality in patients with myocardial infarction was rated as low.

**Patients With Stroke or Acute Brain Injury**

Two fair-quality trials (25, 26) and 2 poor-quality trials (27–29) that examined the efficacy of IIT (target glucose values ranging from 4.4 to 8.0 mmol/L [80 to 144 mg/dL]) in patients with stroke or brain injury showed no mortality benefit compared with higher targets (ranging from <10.0 to <17.0 mmol/L [<180 to <306 mg/dL]). The overall quality of evidence that IIT targeted to achieve glucose levels of 4.4 to 8.0 mmol/L (80 to 144 mg/dL) in patients with stroke does not improve mortality was rated as low.

**Perioperative Care**

Evidence from 1 fair-quality trial (30) and 2 poor-quality trials (31, 32) that evaluated IIT (glucose targets ranging from 3.9 to 10.0 mmol/L [70 to 179 mg/dL]) versus higher or unspecified target values during the immediate perioperative period (IIT was begun before, during, or immediately after surgery and was continued for less than 24 hours after surgery) did not show a beneficial effect on mortality.

The trials included patients undergoing surgery (mainly cardiac) and had small sample sizes, low event rates, and considerable differences in interventions used and blood glucose targets. The overall quality of evidence that IIT to achieve target glucose levels of 3.9 to 10.0 mmol/L (70 to 179 mg/dL) does not improve health outcomes in patients receiving perioperative care was rated as low.

**Infection**

Nine fair-quality trials (7, 9, 13, 17, 30, 33–36) and 7 poor-quality trials (12, 14, 15, 29, 32, 37, 38) evaluated the effect of IIT on the incidence of infection in various patient populations. For sepsis, evidence from 9 trials (7, 9, 12–14, 16, 17, 29, 36) showed a marginally significant reduction in the risk for sepsis (RR, 0.79 [CI, 0.62 to 1.00]). Seven other studies (15, 30, 32–34, 38) reported the occurrence of wound infections, urinary tract infections, pneumonia, or a combination of these infections. A pooled analysis of these outcomes showed a non–statistically significant reduction in infection (RR, 0.68 [CI, 0.36 to 1.30]).
Length of Stay

Eight fair-quality trials (7, 9, 18, 24, 30, 33, 34; Mackenzie I, Blunt M, Ingle S, Palmer C. GLYcaemic Control and Outcome in GENeral Intensive Care. Unpublished report) and 5 poor-quality trials (15, 31, 32, 37) reported the effects of IIT on hospital length of stay. Four trials (7, 9, 18; Mackenzie I, Blunt M, Ingle S, Palmer C. GLYcaemic Control and Outcome in GENeral Intensive Care. Unpublished report) in the mixed MICU/SICU environment found a neutral effect of IIT on overall hospital length of stay (0.008 day [CI, −0.84 to 0.85 day] or ICU length of stay (−0.04 day [−0.34 to 0.26 day]; I² = 0%)]. In contrast, 4 SICU studies (13–15, 39) found that IIT was associated with a reduction in ICU length of stay (−1.5 days [CI, −2.2 to −0.73 day]; I² = 50%; P = 0.11). The overall quality of evidence on effects of IIT on length of hospital or ICU stay in patients in the ICU was rated as moderate.

Two fair-quality perioperative trials (33, 40) showed neutral results for hospital length of stay, and 1 fair-quality trial (34) showed on average a 1-day reduction in length of stay. The overall quality of evidence was rated as low.

Harms of Intensive Insulin Therapy: Hypoglycemia

The use of IIT was associated with an excess risk for hypoglycemia in almost all trials; critically ill patients receiving IIT aimed at achieving normoglycemia had the highest occurrence of hypoglycemia (RR, 5.32 [CI, 4.21 to 6.73]) (41–44). The overall quality of evidence that IIT is associated with hypoglycemia in all subgroups except the general medical unit (for which there were no studies) was rated as high.

The consequences of hypoglycemia in hospitalized patients are unclear because few of the studies reviewed reported adverse effects and few studies have examined the long-term consequences of hypoglycemia. There is some evidence for excess mortality or extended length of stay among patients in the MICU experiencing 1 or more episodes of severe hypoglycemia related to IIT (7, 8, 45, 46). However, it is unclear whether hypoglycemia was a causative factor or whether it was a marker for more severe disease. Some studies have suggested that hypoglycemia is associated with an increased risk for dementia in patients with type 2 diabetes (47) and a 2-fold increase in risk for mortality (48) and that it may induce transient ischemia and catecholamine surges (49–51).

Implementation of Effective and Safe Intensive Insulin Therapy

Fair-quality evidence (52–54) on the effects of different insulin infusion protocols on glycemic control differed in terms of patient characteristics, target glucose ranges, the time required to achieve the target glucose, the incidence and definition of hypoglycemia, the rationale or algorithm used for adjusting the insulin infusion rates, the methods used to assess effectiveness, and the methods of glucose monitoring. Two reviews suggested that in light of variability among protocols, each institution should individualize protocol implementation on the basis of its patient population, institutional resources, and provider resources (52, 53). A third review concluded that protocols that incorporate such factors as the rate of change in glucose level, current blood glucose level, and insulin infusion rate may be more effective than simple sliding-scale infusion protocols in decreasing blood glucose levels while maintaining relatively low rates of hypoglycemia (54). However, this conclusion is not based on direct comparisons of protocols.

Evidence evaluating subcutaneous sliding-scale insulin regimens suggests that this regimen may be relatively ineffective in achieving lower target blood glucose values (38, 55–57).

Summary

Poorly controlled hyperglycemia is associated with increased morbidity, mortality, and worsening health outcomes in hospitalized patients. Most clinicians make efforts to prevent and control hyperglycemia in inpatient settings. However, the optimal blood glucose range to target in hospitalized patients is uncertain. Many trials have shown no effect of IIT targeted to different blood glucose levels on mortality, and pooling of trials does not suggest any trend toward benefit. Among patients in ICUs, in whom there are theoretical reasons to target normoglycemia or near-normoglycemia, the evidence also shows no mortality benefit associated with IIT for targeted glucose levels of 4.4 to 6.1 mmol/L (80 to 110 mg/dL). Although an SICU study (16) showed evidence supporting the link between reduced mortality and the use of IIT to targets of 4.4 to 6.1 mmol/L (80 to 110 mg/dL), the study used aggressive parenteral nutrition, which is not standard practice in many hospitals (16, 36, 58). Parenteral nutrition has been associated with hypertriglyceridemia, insulin resistance, increased infection rates, and increased mortality. Furthermore, this study had a relatively low event rate and was stopped early because of benefit, raising concerns that the reported treatment effect was larger than the “true” treatment effect (59). Finally, the results of this study have not been replicated in other settings. In contrast, several other trials were stopped early owing to an excess risk for hypoglycemia in the intervention groups. This raises the possibility that the lack of observed benefit may reflect inadequate power to detect health benefit (7, 8).

The consequences of severe hypoglycemia in hospitalized patients have not been well studied. There is some evidence for excess mortality or extended length of stay among patients experiencing 1 or more episodes of hypoglycemia. However, from the currently available evidence, it cannot be established whether hypoglycemia was the causative factor.

The evidence evaluating insulin protocols that have achieved normoglycemic targets with low rates of hypoglycemia is sparse. The ability to achieve glucose targets safely
probably depends on multiple factors, including the titration characteristics of the protocol, patient characteristics, staffing ratios, and provider acceptance. Characteristics of these studies suggest that several variables may be responsible for the lower rates of hypoglycemia: modest glucose targets (approximately 5.6 to 8.3 mmol/L [100 to 150 mg/dL]); an iterative, institution-based protocol development and deployment process; and innovations in insulin titration protocols.

**RECOMMENDATIONS**

**Recommendation 1:** ACP recommends not using intensive insulin therapy to strictly control blood glucose in non-SICU/MICU patients with or without diabetes mellitus (Grade: strong recommendation, moderate-quality evidence).

**Recommendation 2:** ACP recommends not using intensive insulin therapy to normalize blood glucose in SICU/MICU patients with or without diabetes mellitus (Grade: strong recommendation, high-quality evidence).

**Recommendation 3:** ACP recommends a target blood glucose level of 7.8 to 11.1 mmol/L (140 to 200 mg/dL) if insulin therapy is used in SICU/MICU patients (Grade: weak recommendation, moderate-quality evidence).

Although the target blood glucose levels in the current trials ranged widely, avoiding targets less than 7.8 mmol/L (<140 mg/dL) should be a priority because harms are likely to increase at lower blood glucose targets. Although the consequences of hypoglycemia in hospitalized patients are unclear, there is some evidence for increased mortality or extended length of stay among patients experiencing 1 or more episodes of hypoglycemia. However, optimal targets in patients not receiving care in the SICU or MICU cannot be precisely defined, because IIT was associated with an excess risk for hypoglycemia in almost all trials and no clear differences in mortality were observed at any target level.

**Figure.** The American College of Physicians guideline on the use of intensive insulin therapy for the management of glycemic control in hospitalized patients.
Data on the effects of IIT targeted to normoglycemia on reduction in length of ICU stay are mixed.

**Recommendation 3: ACP recommends a target blood glucose level of 7.8 to 11.1 mmol/L (140 to 200 mg/dL) if insulin therapy is used in SICU/MICU patients (Grade: weak recommendation, moderate-quality evidence).**

Although IIT to achieve normoglycemia is not associated with improved health outcomes and increases the risk for hypoglycemia, poorly controlled hyperglycemia is associated with increased morbidity, mortality, and worsened health outcomes in patients in the ICU. While the evidence is not sufficient to give a precise range for blood glucose levels, target values of 7.8 to 11.1 mmol/L (140 to 200 mg/dL) is a reasonable option in patients in the ICU, because insulin therapy targeted at blood glucose levels of 7.8 to 11.1 mmol/L (140 to 200 mg/dL) is associated with similar mortality outcomes as IIT targeted at blood glucose levels of 4.4 to 6.1 mmol/L (80 to 110 mg/dL) and is associated with a lower risk for hypoglycemia. Current studies do not provide enough information to determine whether allowing blood glucose levels to increase above 10.0 to 11.1 mmol/L (180 to 200 mg/dL) is associated with similar outcomes to those seen at lower target levels.

Although the risk for hypoglycemia was higher in studies with lower target glucose values, hypoglycemia was also observed among patients who received insulin therapy with target blood glucose levels ranging from 7.8 to 11.1 mmol/L (140 to 200 mg/dL). Therefore, minimizing hyperglycemia associated with IIT is critical in institutions that choose to implement insulin therapy in patients in the ICU. Factors that may be associated with achievement of glucose targets with low rates of hypoglycemia include titration characteristics of the protocol, patient characteristics, staffing ratios, and clinician acceptance. Institutions that implement insulin therapy in patients in the ICU should incorporate quality improvement and training initiatives in order to achieve target glucose levels while minimizing rates of hypoglycemia.

**SUMMARY OF RECOMMENDATIONS AND EVIDENCE**

See the Figure for a summary of the recommendations and clinical considerations. The Table describes the ACP’s guideline grading system.

From the American College of Physicians, Philadelphia, Pennsylvania; Pfizer, Collegeville, Pennsylvania; Oregon Health & Science University and Portland Veterans Affairs Medical Center, Portland, Oregon; and West Los Angeles Veterans Affairs Medical Center, Los Angeles, California.

**Note:** Clinical practice guidelines are “guides” only and may not apply to all patients and all clinical situations. Thus, they are not intended to override clinicians’ judgment. All ACP clinical practice guidelines are considered automatically withdrawn or invalid 5 years after publication, or once an update has been issued.

**Disclaimer:** The authors of this article are responsible for its contents, including any clinical or treatment recommendations. No statement in this article should be construed as an official position of the U.S Department of Veterans Affairs.

**Acknowledgment:** The authors thank Melissa Starkey, PhD, for her help in putting together this document.

**Financial Support:** Financial support for the development of this guideline comes exclusively from the ACP operating budget.

**Potential Conflicts of Interest:** Any financial and nonfinancial conflicts of interest of the group members were declared, discussed, and resolved. Dr. Vincenza Snow was an employee of the American College of Physicians at the time of the writing of this guideline. Dr. Snow: Employment: American College of Physicians, Pfizer. Dr. Shelleke: Grants/Grants pending (money to institution): Agency for Healthcare Research and Quality; Royalties: UpToDate. Disclosures can also be viewed at www.acponline.org/authors/icmje/ConflictOfInterestForms.do?msNum=M10-2725.

**Requests for Single Reprints:** Amir Qaseem, MD, PhD, MHA, American College of Physicians, 190 N. Independence Mall West, Philadelphia, PA 19106; email, aqaseem@acponline.org.

Current author addresses and author contributions are available at www.annals.org.

**References**


Current Author Addresses: Dr. Qaseem: American College of Physicians, 190 N. Independence Mall West, Philadelphia, PA 19106.
Dr. Humphrey: Portland Veterans Affairs Medical Center, 3710 SW US Veterans Hospital Road, Portland, OR 97201.
Dr. Chou: Oregon Health & Science University, 3181 SW Sam Jackson Park Road, Portland, OR 97239.
Dr. Snow: Pfizer, 500 Arcola Road, Collegeville, PA 19426.
Dr. Shekelle: West Los Angeles Veterans Affairs Medical Center, 11301 Wilshire Boulevard, Los Angeles, CA 90073.

Author Contributions: Conception and design: A. Qaseem, L.L. Humphrey, V. Snow, P. Shekelle.
Analysis and interpretation of the data: A. Qaseem, L.L. Humphrey, R. Chou.
Drafting of the article: A. Qaseem, V. Snow.
Critical revision of the article for important intellectual content: A. Qaseem, L.L. Humphrey, R. Chou, V. Snow, P. Shekelle.
Final approval of the article: A. Qaseem, L.L. Humphrey, R. Chou, V. Snow, P. Shekelle.
Administrative, technical, or logistic support: A. Qaseem.