Abstract

The impressive benefits related to the use of tight glucose control by intensive insulin therapy have not been reproduced until now in multicenter large-scale prospective randomized trials. Although the reasons for these failures are not entirely clear, we suggest the use of a stepwise approach — Safe, Effective Glucose Control — that will essentially target an intermediate blood glucose level. As compared with genuine tight glucose control, Safe, Effective Glucose Control — already used in many intensive care units worldwide — is intended to decrease the rate of hypoglycemia and the workload, while reducing the adverse effects of severe hyperglycemia.

In 2001, following in the path of the glycemic pioneers [1-3], the Leuven 1 investigators published their landmark study of intensive insulin therapy in a population of surgical intensive care unit (ICU) patients, targeting 80 to 110 mg/dl in the interventional arm [4]. This prospective, controlled randomized trial spurred clinicians in ICUs around the world to adopt tight glycemic control (TGC) [5]. Confirmation of the mortality benefit of TGC in a mixed medical–surgical ICU setting was seen in the nonrandomized Stamford study published nearly 3 years later [6,7]. An additional 2 years elapsed until the publication of the Leuven II study, performed in a medical ICU, which demonstrated reduced mortality in the predefined population of patients staying in the ICU for at least 3 days but not in the entire population [8]. Two subsequent multicenter studies — GLUCONTROL and VISEP — were terminated prematurely, mainly because of the occurrence of severe hypoglycemia without concurrent improvement in survival [9,10].

Why were the benefits of TGC apparent in the earlier studies not confirmed in the more recently published work?

TGC demands a complex application of monitoring and dynamic treatment throughout the course of the patient’s ICU stay; deficiencies in any number of institutional factors may doom the intervention to failure. Protocol-driven care is central to TGC, with frequent assessment of glycemic levels and responding adjustments in the administered treatment. The experience and skill of the nursing staff in the use of protocols will materially affect the probability that the treatment goals of the protocol are achieved. Moreover, the structural and organizational characteristics of the ICU may have a strong impact, especially in view of the high work burden imposed by TGC — estimated to consume up to 2 hours out of a 24-hour working day for the ICU nurse [11,12].

Appropriate data outcome tools greatly increase the chance that TGC will be practiced successfully. Glycemic reporting tools allow clinicians to know whether glycemic targets are being reached and, importantly, whether there is a significant rate of treatment-associated hypoglycemia. Ideally an ICU should also have an outcomes reporting tool — the ability to provide information such as severity-adjusted mortality and length of stay, complications and resource utilization. Positive feedback imparts a strong incentive to continue the effort needed to maintain effective implementation of TGC [13,14].

One important difference between the early and later trials of TGC is the rate of treatment-associated severe hypoglycemia, defined as <40 mg/dl, found recently to confer increased risk of mortality in a large cohort of mixed medical–surgical ICU patients [15]. The Leuven I study reported an increase in the number of patients with severe hypoglycemia from 0.8% to 5.1%, with no associated adverse consequences [4]. There was no increase in severe hypoglycemia in the Stamford study [6]. In contrast, the percentages of patients in the corresponding groups of the Leuven II study were 3.1% and 18.7% (25.1% among patients in the ICU for longer than 5 days) [8]. In this study the occurrence of severe hypoglycemia was independently associated with mortality on multivariable analysis and resulted in an attenuation of the survival benefit of TGC [8].

ICU = intensive care unit; SEGC = Safe, Effective Glycemic Control; TGC = tight glycemic control.
Similarly, the percentage of patients sustaining severe hypoglycemia among patients in the interventional arm versus the control arm of the GLUCONTROL trial was 2.7% versus 9.8% [9]; the corresponding rates in the VISEP trial were 4.1% and 17.0% [10]. It is possible that differences in monitoring technology and testing frequency may explain some of the differences in the rates of severe hypoglycemia when comparing the Leuven 1 study, which exclusively used arterial blood from indwelling arterial catheters, with the later studies. A growing literature has described the limitations of capillary glucose measurement in the critically ill patient, especially in the lower ranges targeted by these trials [16,17].

It is likely that an additional factor – glycemic variability – has played a role in explaining the divergent outcomes of these different interventional trials [18,19]. A new evaluation of glycemic variability, defined as the standard deviation of the mean glucose level during the ICU stay, suggests that glycemic variability may be an even more important predictor of mortality in the critically ill patient than is the mean glucose level [20]. It is intriguing to note that while the mean (standard deviation) morning glucose levels of the control and interventional arms of the Leuven I study were 153 (33) mg/dl versus 103 (19) mg/dl [4], the corresponding results for the Leuven II study were 153 (31) mg/dl versus 111 (29) mg/dl [8]. Glycemic control improved in the second study but, perhaps, glycemic variability was unchanged.

We are forced, ultimately, to conclude that the Leuven I study may have set the bar too high: TGC, with a glycemic target of 80 to 110 mg/dl is, simply, too tight to practice safely and effectively. If TGC cannot be implemented safely and effectively in a research setting leading to a published intervention trial, then it probably cannot be implemented safely and effectively by most ICU teams.

Instead of TGC, we propose a stepwise approach defining a new standard – Safe, Effective Glycemic Control (SEGC) [21,22]. SEGC involves, first, adoption of a safe glycemic target appropriate to the skills, experience and available tools of the ICU that does not result in a significant increase in the rate of hypoglycemia. A glycemic target of 80 to 150 mg/dl is not unreasonable for an ICU to choose initially; implementation can subsequently lead to downward revision of the glycemic goal. Effective implementation of TGC involves successful attainment of glycemic goals with minimum variability. The use of appropriate data monitoring tools, for both glycemic results and relevant clinical outcomes, is essential for SEGC. Finally, sensible utilization of existing monitoring technologies is mandatory for SEGC.

Preliminary clinical evaluations of the accuracy of continuous or near-continuous glucose monitors have been published recently [23-25]. These devices offer the promise of a reduction in severe hypoglycemia, glycemic variability and the nursing work burden, and will probably become a cornerstone of SEGC. The goals of the SEGC mandate team collaboration are to create and apply glycemic protocols, and the appropriate use of all of the data and monitoring tools that we currently have in our armamentarium, as well as rapid employment of new tools as they are developed. Our patients deserve no less.

**Competing interests**

The authors declare that they have no competing interests.

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