Background: An observational pilot study of 41 medical and surgical intensive care patients on infusion insulin at our own institution found that glycemic control rapidly deteriorated within 48 hours of stopping infusion insulin. This prompted the design and testing of a transition protocol.

Methods: The transition protocol identified appropriate patients for subcutaneous (SC) insulin along with the insulin dose and schedule. A pharmacist-hospitalist improvement team offered protocol guidance but adherence was left to the discretion of the provider. The primary endpoints were mean blood glucose the first and second day after stopping the insulin infusion and the number of patients with hypoglycemia (41-70 mg/dL) and severe hypoglycemia (<40 mg/dL) during the 48-hour transition. Secondary endpoints include severe hyperglycemia (>300 mg/dL), length of stay (LOS), re-initiation of the infusion insulin, day-weighted glucose mean 12 days following transition for patients with diabetes, and identification of a new diagnosis of diabetes.

Results: Patients with diabetes transitioned by protocol (n=33) had better glycemic control than those (n=39) transitioned without the protocol (Day 1 population glucose mean of 168 mg/dL vs. 211 mg/dL [P<0.001], Day 2 means of 176 mg/dL vs 218 mg/dL [P<0.001]). Severe hypoglycemia occurred once in each group. There were 14 patients newly diagnosed with diabetes based on an A1c ≥6%. Patients with stress hyperglycemia maintained good glycemic control with correctional insulin only.

Conclusion: Protocol adherence improved glycemic control, reduced unnecessary use of insulin, and identified patients with previously undiagnosed diabetes, without any increase in hypoglycemia. *Journal of Hospital Medicine* 2010;5:446–451.

KEYWORDS: diabetes mellitus, hyperglycemia management/DKA, infusion insulin, transition protocol.

Additional Supporting Information may be found in the online version of this article.
infusion insulin to SC insulin. The successful implementation of this protocol could serve as a blueprint to other institutions without the need for additional technology or personnel.

Methods
Patient Population/Setting
This was a prospective study of patients admitted to either the medical/cardiac intensive care unit (MICU/CCU) or surgical intensive care unit (SICU) at an academic medical facility and placed on infusion insulin for >24 hours. The Institutional Review Board (IRB) approved the study for prospective chart review and anonymous results reporting without individual consent.

Patients in the SICU were initiated on infusion insulin after 2 blood glucose readings were above 150 mg/dL, whereas initiation was left to the discretion of the attending physician in the MICU/CCU. A computerized system created in-house recommends insulin infusion rates based on point-of-care (POC) glucose measurements with a target range of 91 mg/dL to 150 mg/dL.

Inclusion/Exclusion Criteria
All patients on continuous insulin infusion admitted to the SICU or the MICU/CCU between May 2008 and September 2008 were evaluated for the study (Figure 1). Patients were excluded from analysis if they were on the infusion for less than 24 hours, had a liver transplant, were discharged within 48 hours of transition, were made comfort care or transitioned to an insulin pump. All other patients were included in the final analysis.

Transition Protocol
Step 1: Does the Patient Need Basal SC Insulin?
Patients were recommended to receive basal SC insulin if they either: (1) were on medications for diabetes; (2) had an A1c ≥6%; or (3) received the equivalent of ≥60 mg of prednisone; AND had an infusion rate ≥1 unit/hour (Supporting Information Appendix 1). Patients on infusion insulin due to stress hyperglycemia, regardless of the infusion rate, were not placed on basal SC insulin. Patients on high dose steroids due to spinal injuries were excluded because their duration of steroid use was typically less than 48 hours and usually ended prior to the time of transition. The protocol recommends premeal correctional insulin for those not qualifying for basal insulin.

In order to establish patients in need of basal/nutritional insulin we opted to use A1c as well as past medical history to identify patients with diabetes. The American Diabetes Association (ADA) has recently accepted using an A1c ≥6.5% to make a new diagnosis of diabetes.7 In a 2-week trial prior to initiating the protocol we used a cut off A1c of 6.5%. However, we found that patients with an A1c of 6% to 6.5% had poor glucose control post transition; therefore we chose ≥6% as our identifier. In addition, using a cut off A1c of 6% was reported by Rohlffing et al.8 and Greci et al.9 to be more than 97% sensitive at identifying a new diagnosis of diabetes.

To ensure an A1c was ordered and available at the time of transition, critical care pharmacists were given Pharmacy and Therapeutics Committee authorization to order an A1c at the start of the infusion. Pharmacists would also guide the primary team through the protocol’s recommendations as well as alert the project team when a patient was expected to transition.

Step 2: Evaluate the Patient’s Nutritional Intake to Calculate the Total Daily Dose (TDD) of Insulin
TDD is the total amount of insulin needed to cover both the nutritional and basal requirements of a patient over the course of 24 hours. TDD was calculated by averaging the hourly drip rate over the prior 6 hours and multiplying by 20 if taking in full nutrition or 40 if taking minimal nutrition while on the drip. A higher multiplier was used for those on minimal nutrition with the expectation that their insulin requirements would double once tolerating a full diet. “Full nutrition” was defined as eating >50% of meals, on goal tube feeds, or receiving total parenteral nutrition (TPN). “Minimal nutrition” was defined as taking nothing by mouth (or NPO), tolerating <50% of meals, or on a clear liquid diet.

Step 3: Divide the TDD Into the Appropriate Components of Insulin Treatment (Basal, Nutritional and Correction), Depending on the Nutritional Status
In Step 3, the TDD was evenly divided into basal and nutritional insulin. A total of 50% of the TDD was given as glargine (Lantus) 2 hours prior to stopping the infusion. The remaining 50% was divided into nutritional components as either Regular insulin every 6 hours for patients on tube feeds or lispro (Humalog) before meals if tolerating an oral diet. For patients on minimal nutrition, the 50% nutritional insulin dose was not initiated until the patient was tolerating full nutrition.

The protocol recommended basal insulin administration 2 hours prior to infusion discontinuation as recommended by the American Association of Clinical Endocrinologists (AACE) and ADA consensus statement on inpatient glycemic control as well as pharmacokinetics.10,11 For these reasons, failure to receive basal insulin prior to transition was viewed as failure to follow the protocol.

Safety features of the protocol included a maximum TDD of 100 units unless the patient was on >100 units/day of insulin prior to admission. A pager was carried by rotating hospitalists or pharmacist study investigators at all hours during the protocol implementation phase to answer any questions regarding a patient’s transition.

Data Collection/Monitoring
A multidisciplinary team consisting of hospitalists, ICU pharmacists, critical care physicians and nursing
representatives was assembled during the study period. This team was responsible for protocol implementation, data collection, and surveillance of patient response to the protocol. Educational sessions with house staff and nurses in each unit were held prior to the beginning of the study as well as continued monthly educational efforts during the study. In addition, biweekly “huddles” to review ongoing patient transitions as well as more formal monthly reviews were held.

The primary objective was to improve glycemic control, defined as the mean daily glucose, during the first 48 hours post transition without a significant increase in the percent of patients with hypoglycemia (41-70 mg/dL) or severe hypoglycemia (<40 mg/dL). Secondary endpoints included the percent of patients with severe hyperglycemia (≥300 mg/dL), length of stay (LOS) calculated from the day of transition, number of restarts back onto infusion insulin within 72 hours of transition, and day-weighted glucose mean up to 12 days following transition for patients with diabetes.

Glucose values were collected and averaged over 6-hour periods for 48 hours post transition. For patients with diabetes, POC glucose values were collected up to 12 days of transition.

### TABLE 1. Patient Demographics

<table>
<thead>
<tr>
<th></th>
<th>Patients With Diabetes</th>
<th>Patients Without Diabetes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Protocol Followed, n = 29 Patients*</td>
<td>Protocol NOT Followed, n = 33 Patients</td>
</tr>
<tr>
<td>Average age, years, mean ± SD</td>
<td>57.7 ± 12.1</td>
<td>57.8 ± 12.3</td>
</tr>
<tr>
<td>Male patients</td>
<td>21 (72%)</td>
<td>21 (63%)</td>
</tr>
<tr>
<td>BMI</td>
<td>30.7 ± 7.2</td>
<td>28.6 ± 6.8</td>
</tr>
</tbody>
</table>
| History of diabetes    | 18 (64%) | 25 (86%) | 0.07 | 0 | 0 | 1
| Mean Hgb A1c (%)       | 6.6 ± 1.2 | 7.3 ± 1.8 | 0.136 | 5.6 ± 0.3 | 5.4 ± 0.4 | 0.995 |
| Full nutrition         | 26 (79%) | 24 (81%) | 0.131 | 23 (76%) | 9 (100%) | 1
| On hemodialysis        | 5 (17%) | 9 (27%) | 0.380 | 3 (10%) | 0 | 1
| On >60 mg prednisone or equivalent per day | 7 (24%) | 10 (30%) | 0.632 | 0 | 0 | 1

**Abbreviations:** BMI, body mass index; SD, standard deviation; TPN, total parenteral nutrition.
*One patient followed the protocol requiring basal insulin solely because they were on high dose steroids. They did not have a medical history of diabetes or an A1c ≥6%.
†Too small of cell size to conduct chi-square analysis.

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FIGURE 1. Patient Flow Chart. *Failure to follow the protocol for diabetic patients was defined as: receiving <80% of the recommended basal insulin or receiving insulin after stopping infusion insulin. **Failure to follow the protocol for non-diabetics was defined as: receiving basal or nutritional insulin when none was recommended.
TABLE 2. Insulin Use and Glycemic Control

<table>
<thead>
<tr>
<th>Value</th>
<th>Patients With Diabetes</th>
<th>Patients Without Diabetes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>Protocol Followed, n = 33 transitions</td>
<td>Protocol NOT followed, n = 39 transitions</td>
</tr>
<tr>
<td>Average infusion rate, hours</td>
<td>3.96 ± 3.15</td>
<td>3.74 ± 3.64</td>
</tr>
<tr>
<td>Average BG on infusion insulin (mg/dL)</td>
<td>122.5 ± 27.5</td>
<td>122.5 ± 21.8</td>
</tr>
<tr>
<td>Average basal dose (units) given</td>
<td>34.5 ± 14.4</td>
<td>14.4 ± 15.3</td>
</tr>
<tr>
<td>Hours before (-) or after (+) infusion stopped basal insulin given</td>
<td>-1.13 ± 0.9</td>
<td>11.6 ± 9.3</td>
</tr>
<tr>
<td>Average BG 6 hours post transition (mg/dL)</td>
<td>143.7 ± 39.4</td>
<td>182 ± 62.5</td>
</tr>
<tr>
<td>Average BG 0 to 24 hours post transition (mg/dL)</td>
<td>167.98 ± 50.24</td>
<td>211.02 ± 81.01</td>
</tr>
<tr>
<td>Total insulin used from 0 to 24 hours (units)</td>
<td>65 ± 32.2</td>
<td>26.7 ± 25.4</td>
</tr>
<tr>
<td>Average BG 25 to 48 hours post transition (mg/dL)</td>
<td>176.1 ± 55.25</td>
<td>218.2 ± 88.54</td>
</tr>
<tr>
<td>Total insulin used from 25 to 48 hours (units)</td>
<td>60.5 ± 35.4</td>
<td>28.1 ± 24.4</td>
</tr>
<tr>
<td># of patients with severe hypoglycemia (&lt;40 mg/dL)</td>
<td>1 (3%)</td>
<td>0</td>
</tr>
<tr>
<td># of patients with hypoglycemia (41-70 mg/dL)</td>
<td>3 (9%)</td>
<td>2 (5.1%)</td>
</tr>
<tr>
<td>% of BG values in goal range (80–180 mg/dL) (# in range/total #)</td>
<td>60.2% (153/254)</td>
<td>38.2% (104/272)</td>
</tr>
<tr>
<td># of patients with severe hyperglycemia (&gt;300 mg/dL)</td>
<td>5 (15.2%)</td>
<td>19 (48.7%)</td>
</tr>
<tr>
<td>LOS from transition (days)</td>
<td>14.6 ± 11.3</td>
<td>14 ± 11.4</td>
</tr>
</tbody>
</table>

Abbreviations: BG, blood glucose; LOS, length of stay; n/a, not applicable.
*Too small of cell size to conduct chi-square analysis.

Analysis
Subjects were divided by the presence or absence of diabetes. Those with diabetes were recommended to receive basal SC insulin during the transition period. Within each group, subjects were further divided by adherence to the protocol. Failure to transition per protocol was defined as: not receiving at least 80% of the recommended basal insulin dose, receiving the initial dose of insulin after the drip was discontinued, or receiving basal insulin when none was recommended.

Descriptive statistics within subgroups comparing age, gender, LOS by analysis of variance for continuous data and by chi-square for nominal data, were compared. Twenty-four and 48-hour post transition mean glucose values and the 12 day weighted-mean glucose were compared using analysis of variance (Stata ver. 10). All data are expressed as mean ± standard deviation with a significance value established at P < 0.05.

Results
A total of 210 episodes of infusion insulin in ICU patients were evaluated for the study from May of 2008 to September 2008 (Figure 1). Ninety-six of these episodes were excluded, most commonly due to time on infusion insulin <24 hours or transition to comfort care. The remaining 114 infusions were eligible to use the protocol. Because the protocol recommends insulin therapy based on a diagnosis of diabetes, patients were further divided into these subcategories. Of these 114 transitions, the protocol was followed 66 times (58%).

![FIGURE 2. Transition Graph. Glucose averages before and after transition from infusion insulin.](https://example.com/transition-graph)
both 24 hours and 48 hours after transition than those patients transitioned without the protocol. Day 1 mean blood glucose was 168 mg/dL vs. 211 mg/dL ($P < 0.001$) and day 2 mean blood glucose was 176 mg/dL vs. 218 mg/dL ($P < 0.001$) in protocol vs. nonprotocol patients with diabetes respectively (Figure 2).

There was a severe hypoglycemic event ($\leq$40 mg/dL) in 1 patient with diabetes following the protocol and 1 patient not following the protocol within 48 hours of transition. Both events were secondary to nutritional-insulin mismatch with emesis after insulin in one case and tube feeds being held in the second case. These findings were consistent with our prior examination of hypoglycemia cases.13 Severe hyperglycemia (glucose $\geq$300mg/dL) occurred in 5 (15 %) patients following the protocol vs. 19 (49%) patients not following protocol ($P = 0.002$). Patients with diabetes following the protocol received significantly more insulin in the first 24 hours (mean of 65 units vs. 27 units, $P \leq 0.001$) and 24 to 48 hours after transition (mean of 61 units vs. 28 units, $P \leq 0.001$) than those not following protocol.

An alternate method used at our institution and others14,15 to calculate TDD is based on the patient’s weight and body habitus. When we compared the projected TDD based on weight with the TDD using the transition protocol, we found that the weight based method was much less aggressive. For patients following the protocol, the weight based method projected a mean TDD of 46.3 $\pm$ 16.9 units whereas the protocol projected a mean TDD of 65 $\pm$ 33.2 units ($P = 0.001$).

Patients with diabetes following protocol received basal insulin an average of 1.13 hours prior to discontinuing the insulin infusion versus 11.6 hours after for those not following protocol.

Three patients with diabetes following the protocol and 3 patients with diabetes not following the protocol were restarted on infusion insulin within 72 hours of transition.

LOS from final transition to discharge was similar between protocol vs. nonprotocol patients (14.6 vs. 14 days, $P = 0.836$).

Figure 3 demonstrates that when used correctly, the protocol provides an extended period of glycemic control up to 12 days post transition. Patients transitioned per protocol had a day-weighted mean glucose of 155 mg/dL vs. 184 mg/dL ($P = 0.043$) in patients not following protocol. There was only 1 glucose value less than 40 mg/dL between days 2 to 12 in the protocol group.

Patients Without Diabetes

Of the 39 individual patients without diabetes there were 42 transition events, 33 transitions (78.6%) were per protocol and placed on correctional insulin only. The remaining 9 transitions failed to follow protocol in that basal insulin was prescribed, but these patients maintained comparable glycemic control without an increase in hypoglycemic events. Following transition, patients without diabetes on protocol maintained a mean glucose of 150 mg/dL in the first 24 hours and 153 mg/dL in 24 to 48 hours post transition. They required a mean daily correctional insulin dose of 3.2 units on Day 1 and 2.8 units on Day 2 despite having an average drip rate of 2.3 units/hour at the time of transition (Table 2). There were no severe hypoglycemic events and 80% of blood sugars were within the goal range of 80 mg/dL to 180 mg/dL. Only 1 patient had a single blood glucose of $>300$mg/dL. No patient was restarted on infusion insulin once transitioned.

Patients without diabetes had a longer LOS after transition off of infusion insulin when compared to their diabetic counterparts (22 vs. 14 days).

Discussion

This study demonstrates the utility of hospitalist-pharmacist collaboration in the creation and implementation of a safe and effective transition protocol for patients on infusion insulin. The protocol identifies patients appropriate for transition to a basal/nutritional insulin regimen versus those who will do well with premeal correctional insulin alone. Daily mean glucose was improved post transition for diabetic patients following the protocol compared to those not following the protocol without an increase in hypoglycemic events.

We found an equal number of insulin infusion restarts within 72 hours of transition and a similar LOS in protocol vs. nonprotocol patients with diabetes. The LOS was increased for patients without diabetes. This may be due to worse outcomes noted in patients with stress hyperglycemia in other studies.1,16

The use of the higher multiplier for patients on minimal nutrition led to confusion among many protocol users. The protocol has since been modified to start by averaging the infusion rate over the prior 6 hours and then multiplying by 20 for all patients. This essentially calculates 80% of
Projected insulin requirements for the next 24 hours based on the patient's current needs. This calculation is then given as 50% basal and 50% nutritional for those on full nutrition vs. 100% basal for those on minimal nutrition. This protocol change has no impact on the amount of insulin received by the patient, but is more intuitive to providers. Instead of calculating the TDD as the projected requirement when full nutrition is obtained, the TDD is now calculated based on current insulin needs, and then doubled when patients who are receiving minimal nutrition advance to full nutrition.

Our study is limited by the lack of a true randomized control group. In lieu of this, we used our patients who did not follow protocol as our “control.” While not truly randomized, this group is comparable based on their age, gender mix, infusion rate, mean A1c, and projected TDD. This group was also similar to our preprotocol group mentioned in the Introduction.

Additional study limitations include the small number of nondiabetic patients not following the protocol (n = 9). We noted higher infusion rates in nondiabetics not following protocol versus those following protocol, which may have driven the primary team to give basal insulin. It is possible that these 9 patients were not yet ready to transition from infusion insulin or had other stressors not measured in our study. Unfortunately their small population size limits more extensive analysis.

The protocol was followed only 50% of the time for a variety of reasons. Patients who transitioned at night or on weekends were monitored by covering pharmacists and physicians who may not have been familiar with the protocol. Many physicians and nurses remain fearful of hypoglycemia and the outcomes of our study were not yet available for education. Some reported difficulty fully understanding how to use the protocol and why a higher multiplier was used for patients who were on minimal nutrition.

Efforts to improve adherence to the protocol are ongoing with some success, aided by the data demonstrating the safety and efficacy of the transition protocol.

Conclusion
By collaborating with ICU pharmacists we were able to design and implement a protocol that successfully and safely transitioned patients from infusion insulin to subcutaneous insulin. Patients following the protocol had a higher percentage of glucose values within the goal glucose range of 80 mg/dL to 180 mg/dL. In the future, we hope to automate the calculation of TDD and directly recommend a basal/bolus regimen for the clinical provider.

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