Machine Learning Models for Inpatient Glucose Prediction

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Abstract
Purpose of Review Glucose management in the hospital is difficult due to non-static factors such as antihyperglycemic and steroid doses, renal function, infection, surgical status, and diet. Given these complex and dynamic factors, machine learning approaches can be leveraged for prediction of glucose trends in the hospital to mitigate and prevent suboptimal hypoglycemic and hyperglycemic outcomes. Our aim was to review the clinical evidence for the role of machine learning–based models in predicting hospitalized patients’ glucose trajectory.

Recent Findings The published literature on machine learning algorithms has varied in terms of population studied, outcomes of interest, and validation methods. There have been tools developed that utilize data from both continuous glucose monitors and large electronic health records (EHRs). With increasing sample sizes, inclusion of a greater number of predictor variables, and use of more advanced machine learning algorithms, there has been a trend in recent years towards increasing predictive accuracy for glycemic outcomes in the hospital setting. While current models predicting glucose trajectory offer promising results, they have not been tested prospectively in the clinical setting.

Summary Accurate machine learning algorithms have been developed and validated for prediction of hypoglycemia and hyperglycemia in the hospital. Further work is needed in implementation/integration of machine learning models into EHR systems, with prospective studies to evaluate effectiveness and safety of such clinical decision support on glycemic and other clinical outcomes.

Keywords Machine learning · Artificial intelligence · Glucose · Insulin · Hospital · Diabetes

Introduction
Diabetes mellitus (DM) is a prevalent condition in the hospital, present in nearly 1 in 5 inpatients [1]. Considering how various dynamic factors (e.g., nutritional status, steroid and antihyperglycemic medications, renal function, and infection) may influence glucose homeostasis over a relatively short time frame, it can be difficult to attain or maintain glycemic targets in hospitalized patients [2, 3]. Attaining and maintaining euglycemia is important in the hospital as both hyperglycemia and hypoglycemia have been associated with increased morbidity, mortality, and healthcare expenditures. [4–7]

To address the challenge of glycemic management throughout a patient’s evolving and often complex hospital course, there has been growing interest in and use of clinical decision support tools that provide insights into glucose trends. Machine learning using large electronic health record (EHR) data has been increasingly utilized to facilitate clinical decision-making for a broad range of clinical conditions [8, 9]. In the ambulatory setting, there is a large body of evidence for machine learning–based glucose and insulin dosing prediction, but there are fewer tested algorithms in hospitalized patients [10]. Tools that have been developed to predict glucose trajectory include continuous glucose monitoring (CGM) technology and machine learning models based on point-of-care and serum glucose measurements or CGM [11]. In recent years, there has been a growing interest in using CGM
in the hospital setting as a decision support tool, akin to telemetry monitoring [12]. However, due to challenges associated with employing CGMs in the hospitalized setting, there has also been interest in employing non-CGM-based glycemic prediction algorithms using large EHR datasets. While glycemic prediction models do not necessarily recommend to a provider the best course of action to maintain euglycemia, they can offer insights into glycemic trends to guide therapeutic adjustments in the antihyperglycemic regimen.

The purpose of this review is to summarize machine learning–based models and clinical decision support tools for glycemic prediction in hospitalized patients. Our review focuses on the methodology used, hospital setting (ICU vs. non-ICU), model performance, and impact on clinical outcomes.

**Machine Learning Overview**

Machine learning is the computational process of taking an input of data (i.e., independent variables) and an output (i.e., dependent variable) and uses different statistical rules based on the parameters of each model to create an algorithm that represents the relationship between the raw data and the output. Machine learning has been used throughout medicine to find patterns in genome sequencing, improve diagnostic pathology, and filter relevant problems during a patient’s admission [13]. EHR data is at the forefront of machine learning techniques due to the amount of raw information it possesses, and has been used to predict various disease phenotypes or diagnoses (e.g., congestive heart failure, chronic obstructive pulmonary disease). [14]

Within the field of diabetes, machine learning has been implemented broadly to improve detection of diabetes diagnosis [15] and complications [16], cardiovascular risk stratification [17], and insulin dosing [18]. An early application of machine learning in diabetes was a neural network model that achieved a sensitivity and specificity of 0.76 in predicting the onset of diabetes [19]. Since then, landmark studies in the field have included a gradient boosting model that predicts glucose response prediction based on personal and microbiome factors [20] and a deep learning model that identifies diabetic retinopathy [21]. Most notably, an initial closed-loop insulin delivery system was based on a rules-based model; [18] since then, bi-hormonal artificial pancreas technology has been associated with improved glycemic control when compared to sensor-augmented insulin pump therapy [22]. Over the past few years, there has been growing scientific interest in the use of machine learning with large EHR datasets to predict glucose trends or required insulin doses in hospitalized patients.

**Phases of Machine Learning Research**

Figure 1 outlines the development phases of machine learning–based research with application to the clinical topic of inpatient glycemic prediction [23]. These steps include problem selection, data collection, machine learning development, model validation, assessment of impact, and deployment and monitoring [24]. It is important to recognize that key differences in each of these phases of machine learning research will influence model results, comparability, and generalizability.

**Problem Selection and Study Population**

The first step in machine learning development is defining the study problem and population of interest. The study problem defines the dependent variable, specifically the outcome that is being predicted (i.e., hypoglycemia over the next 24 h, hypoglycemia during any portion of a patient’s admission, etc.). For inpatient glycemic prediction, potential populations include patients with known diabetes or specific type of diabetes (type 1 vs. type 2), patients with acute/stress hyperglycemia, and exposure to specific anti-hyperglycemic medications; populations may be limited to ICU or general floor patients. Selection of the study population will directly affect the generalizability and clinical application of the model. For example, a machine learning model derived from patients with only diagnosed T2DM at the time of admission could not be universally applied to all patients with diabetes.

The method of evaluating model performance (i.e., how accurately the model predicts the outcome in either past or future patients) will also be driven by the problem of interest. For example, if the problem is predicting the next glucose measurement based on CGM, quantitative values for the error, such as mean average error, can be used to measure how far a prediction deviates from the true value; however, if clinical context is more valued than quantitative accuracy and the goal is to properly treat patients, Clarke error grid proportions can be reported [25]. If the problem is prediction of hypoglycemia or hyperglycemia, these binary outcomes can be reported as sensitivity/specificity, positive and negative predictive values, positive and negative likelihood ratios, or area under the receiver operating curve (AUC) [26]. AUC relates the tradeoff a model makes between sensitivity and specificity—a higher AUC indicates overall higher sensitivity and specificity. Generally, an AUC of 0.5 represents no discriminatory value, 0.7–0.8 is considered acceptable, and greater than 0.9 is considered outstanding. It is worth noting that for rare outcomes such as hypoglycemia, the AUC may not
sufficiently quantify the accuracy of a model because of the imbalance between events (hypoglycemia) and non-events. In such cases, a precision-recall plot may be more informative. [27]

**Data Extraction and Processing**

After defining the study question and population, data related to clinical predictors and the outcome measure must be extracted from the EHR. Challenges at this stage include the availability of predictor variables and the ease of extracting relevant information from within the EHR. A collaborative and iterative approach between clinical content experts and EHR vendor certified analysts is recommended to identify and validate the data elements and sources. Since machine learning entails use of data from thousands of patients, another important consideration at this is ensuring the security of the data and minimizing breach of confidentiality by using secure virtual environments for data analysis and considering use of limited (deidentified) datasets when possible.

After a raw dataset is assembled, it must undergo “data tidying,” which consists of organizing the dataset in such a way that it is usable to answer the question at hand based on the defined exposure variables, index unit of observation (e.g., calendar day, patient day, each glucose reading), and the prediction horizon (e.g., next glucose reading, next 24 h, remainder of admission). For example, if a machine learning algorithm attempts to predict hypoglycemia every calendar day but the dataset was extracted for every glucose observation, the dataset will need to be constructed longitudinally such that each row represents the index unit of observation (e.g., calendar day).

Moreover, exposure variables must be temporally related to the outcome variable such that all exposures occurred prior to the outcome (i.e., insulin doses administered in the previous 24 h to predict glucose values in the next 24 h). In some cases, machine learning models have used aggregate data across an entire patient stay to predict glycemic outcomes occurring at any time during admission, which is in fact more of a cross-sectional design (association) rather than longitudinal design (prediction). When insulin or other medications are being used as relevant exposure variables, one should consider the pharmacokinetic profile of the different insulin types to determine insulin dose on board at the time of the prediction. Similarly, glucocorticoids, which can have a significant influence on glucose homeostasis, may need to be normalized to a common equivalent dose (e.g., hydrocortisone or prednisone equivalents) and their

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**Fig. 1** Phases of research in machine learning models for inpatient glucose management. (ICU intensive care unit, CGM continuous glucose monitor, TIR time in range, TAR time above range, RCT randomized controlled trial, ICD International Classification of Diseases)
pharmacokinetic profile would need to be accounted for to determine the active glucocorticoid dose at the time of the index prediction. These data processing steps can be labor intensive, requiring a series of data merges, collapsing of data into summary measures, and computations.

**Machine Learning Development**

**Defining the Outcome**

The next key step is defining the outcome of interest, and variation among different research groups in defining glucose-related outcomes will influence model performance and inter-model comparability. For example, studies exploring hypoglycemia may use different cutoffs (glucose ≤ 70, < 70, < 54, < 50, < 40 mg/dl, etc.), and may apply different criteria to distinguish iatrogenic from spontaneous hypoglycemia. Similarly, various definitions of controlled glucose or hyperglycemia could be considered based on mean glucose values, peak glucose values, or proportion of read-
ings within a defined range (e.g., 70–180 mg/dl). Differences in glycemic outcome definitions will have a direct effect on outcome prevalence and, accordingly, model performance (sensitivity and specificity). [28] For example, choosing more restrictive hypoglycemia cutoffs (< 54 mg/dL vs. < 70 mg/dL) or narrowing the prediction horizon (hypoglycemia with next glucose measurement vs. at any time during admission) will decrease the prevalence of the outcome, thereby affecting the sensitivity and specificity of the model (i.e., it can be harder to accurately predict a more rare outcome). [29]

**Defining the Prediction Horizon**

The prediction horizon (i.e., how far ahead the model attempts to predict the outcome) will have important implications on clinical relevance, usability, and model performance. For example, a shorter prediction window (e.g., next patient day) may be more actionable (i.e., provide a definite time period for clinician intervention), whereas a larger prediction window (e.g., patient admission) increases the prevalence of an outcome and thus model performance, but may have diminished value as a decision support tool throughout the entire hospitalization. Models that attempt to predict risk of a dysglycemic outcome at any time during hospitalization may be useful to flag patients at risk at the time of admission but would have less clinical utility for inpatient clinicians for day-to-day adjustments of the antihyperglycemic regimen.

**Modeling Technique**

Once the outcome and prediction window have been defined, a tidied dataset can be used to create a novel machine learning algorithm. There are many different machine learning techniques—the simplest is linear regression, which infers that there is a linear relationship between exposure variables and the outcome variable and fits slopes for each exposure variable [30]. Although complex modeling techniques can be more robust in creating non-linear models, they are at risk of being computationally inefficient.

Modeling techniques require different amounts of computational storage since they employ unique methodologies to create the algorithm. The simplest of models for quantitative prediction is simple linear regression, which creates a linear model that relates the predictor to the outcome with a slope and y-intercept. For two-class classification (i.e., yes or no), the equivalent would be logistic regression, where each predictor has an associated slope and a y-intercept to output a probability that a prediction is either “yes” or “no.”

The primary limitation to linear and logistic regression is that as relationships between predictors and outcome become less linear, the predictive accuracy of the models worsen [31]. Thus, there has been growing interest in utilizing modeling techniques that do not impose a given relationship on outcome and predictors to leverage information from big datasets. Other modeling techniques used for prediction include k-nearest neighbors, tree-based methods such as random forests, and neural networks.

In k-nearest neighbors, observations in the training set are plotted in as many dimensions as there are predictors. Then, a prediction is made based on the average of the k nearest points. Simply, k-nearest neighbors make a prediction based on the outcome of the observations most similar to the point of interest. Tree-based methods create cutoffs of different predictor values that are set as rules, and a prediction is then made based on those rules. To improve model performance, multiple trees can be used with the resulting prediction based on the average prediction from all the trees in the random forest. In neural networks, the predictors are used to build new functions that are then related to the outcome. The term neural network comes from the new intermediate functions built between predictors and outcome, which were considered analogous to neurons either activating or inhibiting a downstream neuron (the prediction).

In addition to logistic regression, most published models in inpatient glycemic prediction use a derivative of tree-based models. The primary derivatives of tree-based models are bagging, random forests, and boosting [30]. In bagging, repeated random samples are taken from the dataset and predicted upon to allow for multiple trees to be created and reduce the variance of the overall model. Random forests go beyond the bagging technique by limiting the predictors that could be chosen from when creating each rule to decorrelate the trees and make them more reliable. The notion behind random forests is that there are a set of variables that are most correlated to the outcome of interest, but by forcing the model to randomly choose from less correlated predictors,
the model will better capture the nuances between predic-
tors. Boosting is unique in that rather than creating multiple
trees attempting to predict the same outcome, each sequen-
tial tree in boosting is built off the residual, or errors, in
the previous tree such that each tree is built sequentially.
Different machine learning methods such as stochastic gra-
dient boosting and XGBoost modify the boosting technique
in an attempt to maximize model performance or improve
processing efficiency. [32]

Model Validation

A developed machine learning model can then be tested on a
set of patients from the same hospital (internal validation) or
different hospitals (external validation) who are naïve to the
model to estimate model performance. Previous research has
highlighted that external validation is crucial to algorithm
development, as internal validation can overestimate model
performance on a future cohort of patients [33]. Differences
in validation methods using retrospective data may affect
how a model performs after prospective evaluation, as ran-
don cross-fold validation may not account for secular trends
in the hospital setting that could affect the prevalence of the
outcome. This technique of blocking observations is based
on the concern that in structured data models may perform
worse if observations are randomly assigned to the training
and test set [34]. For inpatient glucose prediction, data are
inherently structured by time: increasing understanding of
dysglycemia and new protocols that are developed over time
play a role in mitigating hypoglycemia and hyperglycemia,
but the secular trends are poorly captured in machine learn-
ing models. By dividing earlier observations into the training
set and more recent observations into the test set, research-
ers can report model performance statistics that reflect
changes in the healthcare system not directly accounted for
by included predictor variables.

Assessment of Impact and Deployment and Monitoring

After a model has been developed and validated, it can be
integrated into the EHR and evaluated prospectively for
effectiveness in reducing the desired outcome of interest.
This process requires a near real-time data feed from the
EHR into the algorithm, with output returned to the end user
through the EHR. Some EHR vendors, such as Epic Systems
and Cerner, have built-in tools for predictive analytics to
simplify the process of getting data out and back into the
EHR in real time. For example, the Epic Cognitive Compu-
ting platform (EpicCare, Verona), which leverages Microsoft
Azure cloud software, can be used to configure prediction
models with EHR data, create and modify workflows based
on the results of the predictive models, and report outcomes
of the models and expected workflows [35]. Recently, the
Epic Cognitive Computing platform was used to quickly
develop a model for COVID-19-related outcomes [35]. Similarily, Cerner has partnered with Amazon Web Services
to create a cloud-based health platform for predictive ana-
ytics called “Project Apollo.” [35] It is expected that these
embedded tools for predictive analytics will accelerate the
deployment of machine-learning based prediction models by
streamlining the process for clients and by using native func-
tionality of the EHR. Various formats for the output of the
model could be considered, such as a best practice advisory/
alert (either intrusive or non-intrusive) or a report embedded
with the glucose/insulin management section of the EHR.

After deployment of an EHR-based prediction model,
fixed updates may be needed to revise prediction models
as new data become available, while ad hoc updates may
be needed if there are new therapies (e.g., new insulins or
oral antihyperglycemic medications). In addition, feedback
from end users about clinical utility of the EHR-based model
should be considered to adjust the model based on end user
needs or any changes in clinical care practices.

Machine Learning Models for Inpatient Glucose Prediction

Due to the established adverse effects of poor glycemic con-
trol on patient outcomes, there has been growing research
over the past decade in predicting dysglycemia using
machine learning approaches in the hospital setting [4]. As
studies in the field of inpatient glucose prediction continue
to accrue, there has been a gradual progression from tra-
ditional regression modeling to more advanced machine
learning models, with studies using larger datasets and
much larger numbers of clinical predictors showing con-
tinually improving predictive accuracy. Table 1 summarizes
validated machine learning models for glucose prediction
in the hospital. We review these key studies with respect to
the study populations, outcome definitions, key exposure
variables, prediction horizon, validation method, and model
performance.

Logistic Regression Models for Inpatient Glucose Prediction

Early machine learning methods relied on relatively small
sample sizes and used logistic regression to predict the risk
of hypoglycemia occurring during hospital admission. In
2012, Elliott et al. used a logistic regression model with nine
predictors that included prandial insulin, sulfonylurea, basal
insulin, weight, and renal function [36]. They employed a
training set of 172 patients and presented model performance
on a test set of 3,028 patients; notably, their inclusion crite-
rria for the test set included patients with glucose < 90 mg/
| Author, year | Number of patients | Modeling technique | Clinical predictors | Prediction horizon | Outcome definition | Validation approach | Model performance |
|--------------|-------------------|--------------------|---------------------|-------------------|-------------------|-------------------|-------------------|
| Elliott, 2012 [36] | Training set = 172, test set = 3028 | Logistic regression | 9 total, including prandial insulin, sulfonylurea, basal insulin, weight, and renal function | Hospital stay | Hypoglycemia < 70 mg/dL, < 60 mg/dL, and < 40 mg/dL | Internal validation | 50% sensitivity cutoff for hypoglycemia < 70 mg/dL, < 60 mg/dL, and < 40 mg/dL identified 71%, 70%, and 55% of hypoglycemic events respectively |
| Stuart, 2017 [37] | 9584 admissions | Logistic regression | 13 total including demographics, insulin and sulfonylureas, and electrolytes | Hospital stay | Hypoglycemia < 4.0 mmol/L | Bootstrapping | Model AUC was 0.733 (95% CI: 0.719–0.747) |
| Ena, 2018 [38] | Training set = 839, test set = 561 | Logistic regression | 4 total: GFR < 30 mL/min/1.73 m2, insulin dose > 0.3 units/kg/day, length of stay, previous episode of hypoglycemia during 3 months before admission | Hospital stay | Hypoglycemia < 70 mg/dL | External validation | Model AUC in validation cohort was 0.71 (95% CI: 0.63–0.79) |
| Mathioudakis, 2018 [39] | Training set = 13,360, test set = 5902 | Logistic regression | 44 variables in model predicting ≤ 70 mg/dL and 35 variables in model predicting < 54 mg/dL | 24 h | Hypoglycemia ≤ 70 mg/dL and < 54 mg/dL | Internal validation | C-statistic of 0.77 (95% CI: 0.75–0.78 and 0.80 (95% CI: 0.78–0.82) were achieved for models predicting hypoglycemia ≤ 70 mg/dL and < 54 mg/dL respectively |
| Winterstein, 2018 [40] | 21,840 patients | Logistic regression | 38 variables | 24 h | Hypoglycemia < 50 mg/dL not followed by glucose value > 80 mg/dL within 10 min | Bootstrapping | C-statistic of 0.887 (95% CI: 0.874–0.899) was achieved for predicting hypoglycemia on days 3–5 |
| Shah, 2019 [41] | Training set = 300 patients, test set = 300 patients | Logistic regression | 5 variables: age, ED visit 6 months prior, insulin use, use of oral agents that do not induce hypoglycemia, and severe CKD | Hospital stay | Hypoglycemia ≤ 70 mg/dL | External validation | Validation cohort c-statistic was 0.642 with a sensitivity of 0.77 and specificity of 0.28 when using a risk score cutoff of ≥ 9 |
| Author, year    | Number of patients | Modeling technique       | Clinical predictors | Prediction horizon | Outcome definition | Validation approach                                                                 | Model performance                                                                 |
|---------------|-------------------|--------------------------|---------------------|-------------------|-------------------|-------------------------------------------------------------------------------------|------------------------------------------------------------------------------------|
| Kim, 2020 [42]| 20 patients       | Recurrent neural network | CGM data            | 30 min            | Quantitative glucose value           | Internal validation                                                                 | Average root mean squared error of prediction was 21.5 mg/dL and mean absolute percentage error was 11.1% |
| Kyi, 2020 [43]| 594 patients      | Logistic regression      | 10 variables        | Hospital stay     | At least 2 days with capillary glucose < 72 or > 270 mg/dL | Internal validation                                                                 | Early identification of persistent adverse glycemia had ROC of 0.806, with sensitivity, specificity and PPV of 0.84, 0.66, and 0.53 respectively |
| Ruan, 2020 [8]| 17,658 patients   | XGBoost                  | 42 variables        | Hospital stay     | Hypoglycemia < 4.0 mmol/L and < 3.0 mmol/L | Internal validation                                                                 | AUROC was achieved of 0.96 for the XGBoost model in predicting hypoglycemia < 4.0 mmol/L and < 3.0 mmol/L |
| Elbaz, 2021 [44]| 3,605 patients in training set, 2,425 patients in validation set I and 3,635 patients in validation set II | Logistic regression | 10 variables | 1st week of admission | Hypoglycemia ≤ 70 mg/dL | Internal validation and external validation | AUC in the two validation sets was 0.72 and 0.71 |
| Fitzgerald, 2021 [45]| 10,938 patients in training set and 5,172 patients in test set | Boosted tree | Time series data including glucose, nutrition and insulin dosing | 2 h | Quantitative glucose value | Internal validation | Mean absolute percentage error estimated 16.5–16.8% with 97% of predictions clinically acceptable |
| Mathioudakis, 2021 [9]| 35,147 patients | Stochastic gradient boosting | 43 variables | 24 h after each glucose measurement | Hypoglycemia ≤ 70 mg/dL | Internal validation and external validation | Internal validation C-statistic was 0.90 (95% CI: 0.89–0.90) and external validation C-statistic ranged from 0.86 to 0.88 |
| van den Boorn, 2021 [46]| 94 patients      | First and second derivatives | CGM data            | 30 min            | Quantitative glucose value           | Internal validation                                                                 | Mean squared difference was 7.39 mg/dL and AUC on external validation was 0.79 |
| Horton, 2022 [47]| 11,847 patients   | Logistic regression      | 41 variables        | Impending during ICU stay | Hypoglycemia < 70 mg/dL requiring 50% dextrose within 1 h | External validation                                                                 | AUC on external validation was 0.79 |
as they were predicting hypoglycemic events based on antecedent mild hypoglycemia. Their model, which included predictor variables of insulin doses, weight, creatinine clearance, and sulfonylurea use, achieved a sensitivity of 71% for a hypoglycemic event < 70 mg/dL. This predictive model was coupled with therapeutic recommendations and, to our knowledge, is the only published model that has been deployed in the EHR as an informatics alert and evaluated prospectively. [48]

Ena et al. also used a logistic regression model that included four predictors: GFR < 30 mL/min/1.73 m², insulin dose > 0.3 units/kg/day, length of stay, and previous episode of hypoglycemia during the 3 months prior to admission [38]. Their training and test sets were 839 patients and 561 patients, respectively, from two different nationwide retrospective cohort studies. The authors achieved a validation set AUC of 0.71 for prediction of a hypoglycemic event < 70 mg/dL during a hospital admission. Stuart et al. utilized logistic regression that included thirteen predictors to predict a hypoglycemic event < 4.0 mmol/L (< 72 mg/dL) during hospital admission. [37] Their model achieved an AUC of 0.733 using a cohort of 9,584 admissions. Shah et al. developed a risk prediction tool based on five variables (age, ED visit 6 months prior to admission, insulin use, use of oral agent that does not induce hypoglycemia, and severe CKD) from 300 patients using logistic regression. Using a risk score cutoff of at least 9, their model achieved a sensitivity of 0.77 and specificity of 0.28.

The next phase in machine learning for prediction of hypoglycemia in hospitalized patients used logistic regression to make predictions using a shorter prediction horizon. In 2018, Mathioudakis et al. predicted hypoglycemia occurring on the next calendar day, and achieved a C-statistic of 0.77 when predicting a hypoglycemic event ≤ 70 mg/dL using a logistic regression model based on 44 predictors from 13,360 hospitalized patients in the training set and 5,902 patients in the test set [39]. Similarly, in 2018, Winterstein et al. published a logistic regression model that predicted a hypoglycemic episode defined as a glucose < 50 mg/dL not followed by a glucose value > 80 mg/dL within 10 min in a prediction horizon of 24 h; their model, which used a large cohort of 21,840 patients and 38 predictor variables, achieved a C-statistic of 0.887 when limited to admission days 3–5. [40]

Since the publication of those two studies in 2018, researchers have employed logistic regression in large datasets to answer different questions about predicting inpatient hypoglycemia. Elbaz et al. asked if they could predict glucose ≤ 70 mg/dL in the first week of a patient’s admission [44]. Their dataset included a training set and two validation sets of 3,605, 2,425, and 3,635 patients, respectively. Their externally validated logistic regression achieved AUCs of 0.72 and 0.71 when predicting hypoglycemia during the
first week of a patient’s admission. Horton et al. published a logistic regression model that included 41 predictors for impending hypoglycemia during a patient’s ICU stay [47]. They trained their model using physiologic data up to 12 h prior to a hypoglycemic episode that required treatment with 50% dextrose and excluded any subsequent episodes of hypoglycemia in that patient’s admission. Their training set included 11,847 patients. Upon externally validating their model on the MIMIC-III data, they achieved an AUC of 0.79. Another approach to incorporating both hypoglycemia and hyperglycemia into glucose prediction was presented by Kyi et al. [43] Their model sought to predict persistent dysglycemia (glucose ≤ 72 mg/dL or ≥ 270 mg/dL on two admission days) at the time of admission. Their dataset included 594 patients and ten predictors such as admission dysglycemia, HbA1c ≥ 8.1%, sulfonylurea or insulin use, glucocorticoid use, Charlson Comorbidity Index score, and admission days. Their logistic regression model achieved an ROC of 0.806.

### Tree-Based Models for Inpatient Glucose Prediction

Model performance vastly improved when more advanced machine learning techniques were employed, since a patient’s glucose is not simply linearly related to factors such as antihyperglycemic medication dosing, kidney function, and carbohydrate intake. In 2020, Ruan et al. published an XGBoost model that predicted whether a patient would be hypoglycemic at any time during their admission [8]. Their model, which included 42 clinical covariates and a total sample size of 17,658 patients, achieved an AUROC of 0.96, outperforming a multivariate logistic regression AUROC of 0.75. Seeking to decrease the prediction interval for potentially increased clinical utility, Mathioudakis et al. published in 2021 a stochastic gradient boosting model using 43 static and time-varying clinical covariates that predicted whether or not a patient would have at least one hypoglycemic reading in a rolling window of 24 h from each glucose measurement, which could act as a warning for a provider to adjust a patient’s insulin regimen [9]. The total sample size across internal and external validation hospitals was 35,147 patients, and the stochastic gradient boosting model achieved a C-statistic of 0.90 upon internal validation and ranged from 0.86 to 0.88 upon external validation in four different hospitals.

A recent trend in the field of inpatient glucose prediction has been to shorten the prediction horizon while incorporating as much data present in the EHR as possible. One of the shortcomings of predicting hypoglycemia at the admission level is that the only data that can be included in the model are those that are known at the time of admission, which prevents changes in labs, vital signs, and medications from being incorporated. Risk of hypoglycemia is inherently dynamic within a patient’s hospital course and being able to incorporate new data with a shorter prediction horizon allows for more clinically meaningful predictions. Zale et al. published this year a random forest model employing 59 predictors that classified a patient as hypoglycemic, controlled, and hyperglycemic using a large dataset with over 100,000 patients from five hospitals [29]. Unlike earlier glycemic prediction models, this model predicted the glycemic outcome for the next glucose observation at the time of each index glucose measurement, with the interquartile range for time to next glucose measurement of 1.63 to 4.37 h. Four of those hospitals were used for external validation and achieved sensitivities for controlled, hyperglycemic, and hypoglycemic of 0.64–0.70, 0.75–0.80, and 0.76–0.78, respectively; specificities for controlled, hyperglycemic, and hypoglycemic ranged from 0.80 to 0.87, 0.82 to 0.84, and 0.87 to 0.90, respectively, across the external validation hospitals.

### Predicting Glucose as a Quantitative Variable

With the increase in dataset sizes, several studies have been published in the last 2 years using CGM data and time series analyses to predict glucose over a shorter prediction horizon and to treat glucose as a quantitative (discrete value) rather than a binary outcome. Kim et al. published a recurrent neural network model to predict a patient’s glucose with a prediction horizon of 30 min and achieved an average root mean squared error of 21.5 mg/dL [42]. Their sample size was 20 patients and included 99.4 patient-days of CGM data. Van den Boorn et al. utilized a CGM dataset with a larger sample of ICU patients (N=94) that totaled 134,673 glucose measurements. Their model attempted to predict the glucose reading with a prediction horizon of 30 min based on the first and second derivatives of the CGM data, and achieved a mean squared difference of 7.39 mg/dL [46]. The time series approach to predicting glucose in the short-term horizon has not been limited to CGM data, but has also been applied to the MIMIC-III dataset from critical care unit patients. Fitzgerald et al. applied a boosted tree model to this dataset of ICU patients, dividing the patients into a training set of 10,938 patients and a test set of 5,172 patients. The authors achieved 97% clinically acceptable predictions with mean absolute percentage error estimated between 16.5 and 16.8% with a 2-h prediction horizon. [45]

### Building upon CGM Prediction

Commercially available CGM devices offer real-time alerting for both hypoglycemia and hyperglycemia and could therefore be considered an alternative strategy to EHR-derived machine learning prediction models in the hospital. Although a CGM-based glucose telemetry system...
Future Directions

Despite the growing body of research in the past decade on machine learning–based tools for glucose prediction in the hospital setting, this field is still in a relatively nascent state, with most prediction models having been validated using retrospective data, but only one published model [51] (developed using a relatively small sample) having been integrated into EHR systems and tested prospectively for clinical effectiveness. Although machine learning holds potential for glycemic prediction in the hospital, it is unknown whether providing hospital-based clinicians more accurate predictions in real time will modify prescribing practices and improve glycemic outcomes. We previously found that there is a high degree of clinical inertia in the hospital even after an overt hypoglycemic episode [52], so the clinical value of machine learning–based glycemic prediction remains to be demonstrated. In the meantime, additional published studies from different healthcare systems using EHR data are needed to refine our understanding of parameters that affect insulin dosing in the hospital so that glycemic predictions can ultimately be coupled with actionable insulin dosing recommendations [53]. Considering that many of the insulin dosing algorithms are proprietary and unpublished, publication of methods and results will allow the scientific community to advance our understanding of factors involved in glucose homeostasis in the hospital and to continually improve the predictive accuracy of insulin dosing algorithms.

Conclusion

Advanced machine learning models using large EHR datasets with large numbers of clinical predictors achieve greater predictive accuracy for glucose than more traditional regression modeling techniques in hospitalized patients. Inferences and comparisons of different machine learning algorithms for inpatient glucose prediction require careful consideration of the study population, outcome and exposure variables, and prediction horizon used to derive the model, as well as the validation approach used to describe model performance. Further prospective studies would be needed to demonstrate the added value of deploying machine learning derived predictive analytic support on improving glycemic management and other outcomes in hospitalized patients.

Declarations

Conflict of Interest The authors declare no competing interests.

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