

Construction of a Diabetes Prediction Model Based on Machine Learning

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Abstract. This study investigates key predictors of diabetes risk across non-diabetes, prediabetes, and diabetes categories, while developing an optimal prediction model using multiple machine learning algorithms. Biomedical indicators such as HbA1c, urea, and creatinine, along with demographic factors like age and gender, were analyzed to evaluate their predictive value. Among the five algorithms tested, ensemble learning methods (CatBoost and XGBoost) outperformed traditional models, with CatBoost achieving the highest accuracy and demonstrating superior robustness. Feature importance analysis identified HbA1c as the most influential predictor, followed by age and BMI, aligning with established medical knowledge, whereas gender contributed minimally. The findings highlight the potential of advanced machine learning models, particularly CatBoost, in delivering highly accurate and stable diabetes risk prediction. This research provides strong technical support for early screening, targeted intervention, and practical risk assessment in diabetes management.

Keywords: Diabetes Prediction, Machine Learning, CatBoost, HbA1c

1. Introduction

Diabetes is a chronic metabolic disease characterized by persistent hyperglycemia and has become a major global public health challenge. According to statistics from the World Health Organization (WHO), the global prevalence of diabetes among adults aged 20-79 has reached 14%, with more than 800 million confirmed cases, a fourfold increase over 1990 [1]. By 2024, global diabetes-related medical expenditure is expected to exceed US \$1 trillion, accounting for 12% of global health expenditure, which not only increases the economic burden on patients' families but also impacts public health systems [2]. Accurate identification of prediabetes and early prediction are core to prevention and control.

Complex factors influence diabetes and rely on assumptions of linearity and variable independence, struggle to capture synergistic effects between features, and have weak multi-dimensional data fusion capabilities. Its missed diagnosis rate can reach 30%, failing to meet the needs of precise screening [3]. It can efficiently integrate multi-dimensional data, explore non-linear correlations between features, and ensure interpretability through feature importance analysis.

Based on this, this study uses the Diabetes Prediction Dataset from Kaggle to explore core influencing factors of prediabetes via machine learning, construct and optimize risk prediction

models, select the most interpretable optimal model, and provide a practical tool for early prediabetes screening to reduce the risk of progression to type 2 diabetes [4].

2. Literature review

2.1. Review of previous studies on machine learning algorithms for diabetes prediction

In diabetes prediction research, the evolution of machine learning applications progresses from traditional single models to more sophisticated ensemble methods. Conventional algorithms such as Logistic Regression, Decision Trees, and k-Nearest Neighbors are widely adopted in earlier studies owing to their conceptual simplicity and high interpretability. However, subsequent research reveals notable performance limitations of these individual models. A major issue stems from the inherent characteristics of real-world medical data, which often contain high rates of missing values, noise, and inconsistencies. The absence of critical clinical indicators, such as HbA1c and OGTT, can substantially impair model performance [5]. Traditional statistical methods (e.g., mean imputation) fail to handle complex missing data, limiting model performance.

Ensemble learning effectively addresses these limitations. Voting-based ensemble models, for instance, demonstrate improved prediction accuracy through multi-model fusion [6]. CatBoost excels at automatically processing categorical features, maintaining stable performance in samples with discrete variables, and solving the problem of categorical feature encoding in traditional algorithms. XGBoost further optimizes performance through regularization mechanisms, achieving high accuracy with feature importance rankings consistent with clinical consensus. Zhang et al. further optimized XGBoost by addressing its cumbersome parameter tuning and tendency to fall into local optima: they introduced a genetic algorithm for global parameter search (using MSE as the fitness function), and the optimized model outperforms both traditional algorithms and default-parameter XGBoost [7]. However, large prediction deviations for rare high blood glucose values and room for optimization in runtime should be addressed.

Overall, previous studies share common limitations, including insufficient algorithm comparison, rough data preprocessing, and inadequate integration of medical knowledge. By comparing five algorithms (Logistic Regression, Decision Tree, KNN, CatBoost, XGBoost) and combining strict data cleaning with medical cognition verification, this study aims to construct a more accurate and stable prediction model.

2.2. Supplementary applications of machine learning algorithms in medical prediction

Machine learning excels at processing complex data and capturing correlations. It is widely used for risk prediction in fields such as cardiovascular and cerebrovascular diseases, cancer, and infectious diseases, and performs significantly better than traditional methods.

In chronic kidney disease (CKD) prediction, Liu et al. used CatBoost to forecast 2-year end-stage renal disease (ESRD) risk, achieving high accuracy and clinical utility [8]. For myocardial infarction (MI) risk, XGBoost outperformed traditional methods in identifying high-risk groups and explaining prediction logic [9].

Machine learning offers universal advantages in medical prediction, as it can integrate multiple types of data, capture correlations between indicators, and achieve high accuracy with fewer misjudgments including missed judgments, and erroneous judgments compared to traditional methods.

3. Methodology

3.1. Algorithm principles

3.1.1. K-Nearest Neighbors (KNN)

KNN is an inert supervised learning algorithm based on sample similarity to determine category attribution. It lacks an explicit model training process, relying solely on stored training data for prediction. The approach utilizes Euclidean distance as the standard for similarity quantification, which requires determining the hyperparameter K through cross-validation to balance the ability to capture local features with resistance to noise interference. Finally, the K nearest neighbor samples with the smallest distance are selected from the predicted samples, and their categories are deduced based on the majority voting rule. It is a probability voting process based on the distribution of local samples.

3.1.2. CatBoost

CatBoost is an improved ensemble learning algorithm based on Gradient Boosting Decision Trees (GBDT), optimized for categorical feature processing and bias control. It realizes classification by iteratively constructing a weighted ensemble of CART trees. For non-numeric categorical features, it adopts ordered ranking encoding based on the mean value of the target variable, which completes numerical conversion while preserving logical relationships between categories. Through the “ordered boosting” strategy, it calculates gradients using the complete training set to reduce bias caused by sample splitting in traditional GBDT. Additionally, it suppresses overfitting via three regularization mechanisms: tree complexity restriction, learning rate adjustment, and random subspace sampling. It is suitable for modeling scenarios with small samples and multiple categorical features.

3.1.3. Extreme Gradient Boosting (XGBoost)

XGBoost is a theoretically optimized and engineering-enhanced version of GBDT, focusing on accurate loss function optimization and robustness improvement. It iteratively corrects errors of previous models through an additive model. Breaking the limitation of traditional GBDT using only first-order derivatives, XGBoost introduces second-order derivatives (Hessian) and approximates the loss function as a quadratic Taylor polynomial for more precise error optimization. It can automatically learn the optimal splitting path for missing values during training and improve computational efficiency via feature parallelism and sparse-aware optimization. It is suitable for modeling scenarios with multiple features and missing values.

3.2. Data preprocessing

3.2.1. Data cleaning

Data cleaning focused on correcting outliers and unreasonable values in the original dataset to ensure data consistency with reasonable ranges of physiological indicators and improve data accuracy. Specific operations included:

Correcting abnormal physiological indicator values: Adjusting values exceeding normal physiological ranges for indicators such as urea, creatinine (Cr), high-density lipoprotein (HDL),

very low-density lipoprotein (VLDL), and cholesterol (Chol).

Addressing data consistency issues: Calibrating unreasonable values caused by potential input errors (e.g., abnormally high/low values of VLDL and triglycerides [TG]) based on reasonable ranges to avoid interference of extreme values on subsequent analysis.

3.2.2. Data formatting

Data formatting focused on data type conversion, removal of irrelevant variables, and numerical encoding of categorical variables to provide numerical data suitable for subsequent modeling. Specific operations included:

Numerical encoding of categorical variables: Converting character-type categorical variables to numerical variables: Gender (gender): 0 for female (original: F), 1 for male (original: M). CLASS (diabetes status): 0 for Non-diabetic (original: N), 1 for Prediabetic (original: P), 2 for Diabetic (original: Y). This solved the problem that character-type variables cannot be directly used for modeling.

Removing irrelevant identifier variables: Deleting ID and No_Pation columns (both are unique sample identifiers with no correlation to diabetes) to simplify feature dimensions and reduce interference of redundant information on the model.

Ensuring data format consistency: Unifying data types of all features, aligning numerical indicators with encoded categorical variables to lay a foundation for subsequent machine learning model training and statistical analysis.

3.3. Model design

To determine the optimal parameters for the training model, this study uses a grid search method with 5-fold cross-validation [10]. Grid search exhaustively tests all possible hyperparameter combinations by defining the hyperparameters to be adjusted and their possible value ranges, ultimately selecting the combination with the best performance. Accuracy is used as the core evaluation metric for optimal parameter selection during grid search. In the subsequent test phase, this study uses multiple metrics, Accuracy, Recall, Precision, F1, and AUC-ROC for verification. The main adjusted model parameters and their value ranges are shown in Table 1.

Table 1. Adjusted parameters and value ranges of each model

Model	Parameters	Value Range
Logistic Regression	C (regularization strength)	[0.01,0.1,1,10]
	Penalty (regularization method)	['l1', 'l2']
Decision Tree	max_depth (maximum tree depth)	[3,5,7, None]
	min_samples_leaf(min samples per leaf node)	[1,5,10]
	criterion (splitting criterion)	[gini (Gini coefficient),entropy (information, entropy)]
KNN	n_neighbors	[1,3,5,7,9]
	weights	['uniform' (equal weight), 'distance'(distance-weighted)]
	p (distance metric)	[1 (Manhattan distance),2 (Euclidean distance)]
CatBoost	iterations	[100,200,1000]
	learning_rate	[0.01,0.1,0.3]
	depth (tree depth)	[3,5]
XGBoost	n_estimators (number of decision trees)	[100,200,1000]
	learning_rate	[0.01,0.1,0.3]
	max_depth	[3,5,7]

4. Experiment

4.1. Experimental design

First, the dataset was split using Python's `train_test_split` function, with 80% allocated to the training set and 20% to the test set. `random_state=42` was set to ensure experimental reproducibility.

Logistic Regression and Decision Tree served as basic prediction methods for result comparison. KNN, CatBoost, and XGBoost were the core research objects, with hyperparameters optimized via grid search (`GridSearchCV`), with five-fold cross-validation (target: accuracy). Subsequently, the optimal model of each algorithm was evaluated on the test set, and accuracy, macro-averaged recall, macro-averaged precision, macro-averaged F1-score, and multi-class AUC-ROC were calculated. Feature importance was extracted and visualized for four models (Decision Tree, CatBoost, XGBoost, and Logistic Regression). Finally, the optimal prediabetes prediction model was selected based on comprehensive evaluation metrics.

4.2. Analysis of feature importance

As shown in Figure 1, feature importance rankings across the four models (Logistic Regression, Decision Tree, CatBoost, and XGBoost) exhibit both consistencies and reasonable differences, all of which align closely with clinical knowledge and validate key diabetes risk factors.

HbA1c is consistently identified as the most important predictive feature in all models, reflecting its status as an internationally recognized gold standard for diabetes diagnosis. As it indicates average blood glucose levels over the past 2–3 months, thereby validating the feature selection approach in this study [11].

Beyond HbA1c, the hierarchy of influencing factors reflects clinically relevant patterns. Ensemble models emphasize the significance of age, corroborating the notion that “individuals over 40 belong to the high-risk group for diabetes.” In the Decision Tree model, BMI ranks second in importance, supporting the clinical understanding that obesity-induced insulin resistance is a core

mechanism in type 2 diabetes pathogenesis [12]. The importance of blood lipid and renal function indicators varies complementarily across models, illustrating the systemic nature of metabolic dysregulation and the ability of different algorithms to capture distinct correlation patterns.

In contrast, gender is the least important feature in all models, indicating that its influence on diabetes onset is negligible compared to core physiological, metabolic, and demographic factors (age, BMI). This observation is consistent with previous studies that found no significant impact of gender on incidence rates or mortality [13]. Therefore, gender need not be considered a core indicator in diabetes screening. Differences in secondary feature weights, such as linear models focusing on linear correlations of blood lipid indicators and ensemble models emphasizing interactions between age and renal function indicators are inherent to algorithmic principles and do not alter the identification of central risk factors.

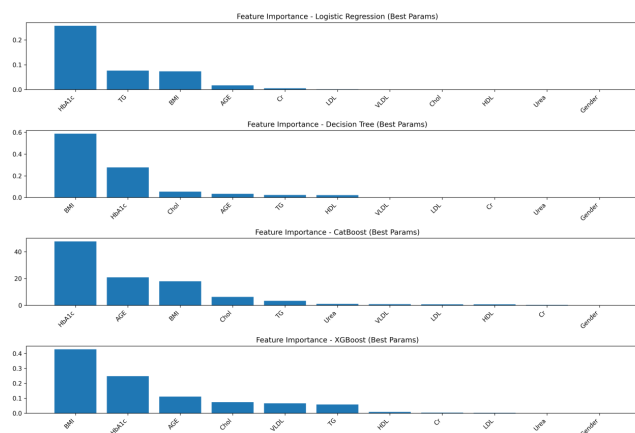


Figure 1. Model feature importance under optimal parameters

4.3. Model performance comparison

Table 2 presents the test set performance metrics of each model under optimal parameter configuration. The models exhibited a clear hierarchy: ensemble learning models (CatBoost,XGBoost) > single-tree model (Decision Tree) > lazy learning model (KNN) > linear model (Logistic Regression). Ensemble learning models performed particularly well in metrics critical to medical scenarios.

Table 2. Test set performance metrics of each model

Model	Accuracy	Recall	Precision	F1	AUC-ROC
Logistic Regression	0.900	0.598	0.567	0.582	0.941
Decision Tree	0.975	0.967	0.967	0.967	0.970
KNN	0.950	0.934	0.832	0.869	0.995
CatBoost	0.993	0.997	0.986	0.991	0.999
XGBoost	0.987	0.994	0.974	0.984	0.999

Among all models, CatBoost stood out: it achieved 0.993 accuracy, 0.997 recall (nearly zero missed diagnosis), 0.986 precision (high prediction credibility), plus the highest macro-averaged F1-score 0.991 and multi-class AUC-ROC 0.999. It fully meets diabetes prediction's core needs: low missed diagnosis rate, low misdiagnosis rate, and stable classification.

For other models: XGBoost matched CatBoost in AUC-ROC but performed slight worse in core accuracy and recall. Decision Tree, KNN, and Logistic Regression were inferior overall. In particular, Logistic Regression failed to capture non-linear feature correlations, leading to poor performance (accuracy 0.900, recall 0.598). Notably, AUC-ROC in medical scenarios must be comprehensively judged with specific classification indicators and cannot be used as the sole evaluation criterion. Thus, the CatBoost model is determined to be highly accurate and stable, making it the optimal high-performance prediction model.

5. Discussion

5.1. Limitations

Although this study constructed and selected an optimal diabetes prediction model using multiple algorithms, it has notable limitations in model design and applicability, mainly in two aspects.

Insufficient data support and limited generalization ability. The model relies on single-source data (from Kaggle), which has significant flaws: Lack of labels for region, ethnicity, economic status, and underlying diseases, failing to cover population heterogeneity in diabetes onset. This leads to reduced generalization performance of the model in specific populations.

No longitudinal time-series data in the dataset. Feature processing is confined to basic operations, such as variable encoding and redundancy removal, with no construction of clinically meaningful composite features. Moreover, feature selection relies entirely on the built-in mechanisms of algorithms, without prior screening based on clinical guidelines. This results in insufficient clinical integrity of the feature system and inaccurate characterization of diabetes pathogenesis.

5.2. Future directions

Future research can advance in four key directions. First, expand multi-source longitudinal data by incorporating population heterogeneity features and dynamic monitoring indicators [14]. Second, construct composite indicators based on clinical needs, optimize feature dimensions, develop population-specific sub-models and improve interpretability. Third, adopts methods such as knowledge-guided graph attention networks to enhance disease prediction and incorporate medical knowledge graphs to help infer missing values through graph-based representation learning [15]. Ultimately, promote clinical translation by integrating models into medical systems, developing lightweight tools, and conducting prospective verification to support diabetes prevention and control, ultimately reducing disease burden.

6. Conclusion

Using the Diabetes Prediction Dataset from Kaggle, this study first corrected outliers in physiological indicators, encoded categorical variables, and removed irrelevant variables. Then, five algorithms (Logistic Regression, Decision Tree, KNN, CatBoost, XGBoost) were used for modeling, with hyperparameters optimized via five-fold cross-validation combined with grid search. The model's performance was evaluated using multiple metrics. Results showed that the ensemble learning model, CatBoost, performed best, with an accuracy of 0.993 and AUC-ROC of 0.999 on the test set—outperforming other model types. HbA1c was the primary predictor of diabetes risk, while the gender had the least impact in the model. This study overcomes the limitations of traditional linear statistics, provide an efficient technical tool for early diabetes screening and risk assessment, and clarified the predictive priority of clinical indicators.

This research also has limitations in terms of data support, feature engineering, and model interpretability. In the future, improvements can be implemented in the following aspects. First, collecting multicenter longitudinal data and balancing sample distribution to enhance the model's generalization ability. Second, integrating clinical knowledge to construct composite features, screen indicators based on guidelines, and refine the feature system. Third, leveraging tools such as SHAP values to quantify the mechanism underlying feature action and improve the model's interpretability.

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