Wavelet based Machine Learning Approaches towards Precision Medicine in Diabetes Mellitus

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Keywords: Diabetes Mellitus, Discrete M-band Wavelet Transform, Machine Learning, Precision Medicine, Data Visualization, t-SNE, UMAP.

Abstract: It is estimated that 422 million people around the world have diabetes mellitus (DM)—a devastating, complex, and highly heterogeneous disease—requesting better interventions based on disease subtyping. In this research, we utilize the discrete wavelet transform (DWT) to decompose and denoise DM data. Using DWT, we enhance heart rate variability (HRV) based DM diagnosis, data visualization of the disparities in Human Microbiome Project (HMP) data (gut bacteria, metabolomics, proteomics, RNA sequencing, targeted proteomics, and transcriptomics data) using demographic features, and insulin resistance prediction. We also attempt to forecast continuous glucose monitoring (CGM) ahead by 90 minutes because CGM is unable to provide real-time blood glucose measurements. We achieve 91.9% diagnosis accuracy for Type 1 DM using Random Forest on data transformed with DWT, holding the potential for usage in clinics. In addition, our DWT-based t-SNE and UMAP explorative analysis of HMP data support subtypes of prediabetic patients stratified by sex, race, and age. Moreover, DWT-based transformations provide multi-view clustering that any other methods would not provide on metabolomics, proteomics, RNA sequencing, targeted proteomics, and transcriptomics data and outperform those without DWT. Taken together, DWT-based machine learning approaches enable a fine resolution of subtyping DM towards precision medicine.

1 INTRODUCTION

Diabetes Mellitus (DM)—a group of metabolic diseases that manifest themselves with chronic hyperglycemia resulting from issues with insulin absorption or production—is estimated to have a prevalence of 422 million people around the world. DM is a devastating, complex, and highly heterogeneous disease, requesting better interventions based on disease subtyping (Kharroubi & Darwish, 2015).

Hypoglycemia is a condition where blood sugar drops below normal levels. It is a relatively common condition in diabetic patients, although it occurs more frequently in individuals affected by type 1 DM (T1D). While hypoglycemia is usually harmless, prolonged hypoglycemia without action can lead to seizure, brain damage, or even death. Conversely, hyperglycemia is a condition where blood sugar rises above normal levels and this condition is more common with individuals affected by T1D. However, untreated hyperglycemia can result in damage to various tissues, comatose, or even death.

Because of these conditions, monitoring blood glucose levels is vital. While accurate blood glucose can be given in near real-time, predicting blood glucose levels into the near future would be a useful tool in preventing abnormal levels of glucose. By using machine learning methods, we sought to provide accurate predictions for blood glucose levels in individuals with T1D or type 2 DM (T2D).

At the same time, diagnosing DM is essential for the long-term health of patients. Nevertheless, many of these tests are either inaccurate or very inconvenient. For example, the A1C test is affected by many factors, including anemia, smoking, pregnancy, and certain infections (Bonora & Tuomilehto, 2011). Other tests, like the glucose tolerance test, take too much running time for many individuals. By using machine learning methods, we sought to provide accurate predictions for blood glucose levels in individuals with T1D or type 2 DM (T2D).

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sought to classify individuals as healthy or T1D by using heart rate variability (HRV).

In this research, we utilize the discrete wavelet transform (DWT) to decompose and denoise DM data, and to create multi-view clustering viewing window as well.

To address many aspects of the DM illnesses and their treatments, we base our study on CGM forecast, HRV-based diagnosis, and DM subtyping. Across these topics, we identify prediabetes (i.e., subtyping by demographics, prediction of insulin resistance), as well as type 1 (i.e., diagnosis, managing blood glucose) and type 2 DM (i.e., managing blood glucose). Combining all of these topics shines light on DWT-based machine learning approaches towards precision medicine. In addition, our DWT-based t-SNE and UMAP explorative analysis of HMP data support subtypes of prediabetic patients stratified by sex, race, age, insulin resistance (IR)/insulin sensitivity (IS) based on DWT of proteomics, targeted proteomics, and transcriptomics data.

1.1 CGM Forecast

The introduction of continuous glucose monitoring (CGM) enables non-invasive and more comprehensive monitoring. CGM sensors can deliver interstitial glucose levels every 1 to 5 minutes, in contrast with previous glucose monitoring methods. This provides a significantly more detailed time series on glucose levels that can be automatically sent to smart devices. However, CGM devices are unable to accurately find blood glucose levels; rather, CGM lags behind the trend of blood glucose levels. As a result, it is necessary to be able to forecast glucose levels by using CGM measurements to provide real-time glucose updates for diabetic patients (Lobo et al., 2021).

Using a modified Artificial Neural Network (ANN), a model that replicates the interconnected neurons of the brain, Bertachi et al. predicted blood glucose 15, 30, 45, and 60 minutes ahead and achieved RMSEs of 6.43, 7.45, 8.13, and 9.03, respectively (Bertachi et al., 2018). In 2017, Fiorini et al. trained 4 different models namely Long Short Term Memory (LSTM), Auto Regressive Integrated Moving Average (ARIMA), Kalman Filter, and Kernel Ridge Regression (KRR) with a dataset of 148 patients (Fiorini et al., 2017). KRR was the most accurate for 30, 60, and 90 minutes, successively. However, each model was trained to fit separate individuals, rather than fitting all of the patients. These methods differed from previous ones with the introduction of methods that are not based on neural networks, such as KRR.

In this research, we propose a method to accurately predict glucose levels using a novel technique developed by Facebook called Prophet. We also use traditional methods ARIMA and KRR as benchmarks (Taylor & Letham, 2017).

1.2 Diabetes Prediction using HRV

Heart rate variability (HRV) refers to the variability in RR intervals, which are the time between consecutive heartbeats. As DM has a harmful impact on the heart, HRV in diabetic patients is reduced (Kardelen et al., 2006). As a result, HRV has been used to detect DM. Seyd et al. used an ANN, to classify HRV signals from 65 healthy people and 65 diabetic patients. Using the ANN, they achieved an accuracy of 93.08%, a precision of 96.67%, and a recall of 89.23% (P.t. et al., 2011). Similar to Seyd et al., Swapna et al. used a Convolutional Neural Network (CNN), but combined with LSTM and support vector machine (SVM) for the classification of echocardiogram (ECG) signals of 20 diabetic patients and 20 healthy individuals. Swapna et al. attained a high accuracy of 95.7% with the combination model (Swapna et al., 2018).

On the other hand, machine learning (ML) algorithms have also been used to classify HRV data with comparably high metrics. Acharya et al., recording ECG signals and obtaining HRV signals from 15 healthy and 15 diabetic patients, achieved an accuracy of 90.0%, precision of 88.9%, and recall of 92.5% with the AdaBoost classifier with the least squares method (Rajendra acharya et al., 2013). Furthermore, in a later work, Acharya et al. carried out the Decision Tree algorithm on the same HRV signals transformed using Wavelet decomposition up to 5 levels, resulting in an accuracy of 92.64%, precision of 92.59%, and a recall of 92.68% (Rajendra acharya et al., 2015).

To predict DM using HRV signals, we examined the SVM, XGBoost, and Random Forest (RF) on HRV signals transformed in a variety of methods via DWT. While RF has been used to diagnose DM (Samant & Agarwal, 2018; Benbekcame, 2019), XGBoost and RF have not been used prior to classify HRV signals transformed using DWT, and our remarkable experimental results strongly support our algorithm.

1.3 Wavelet based t-SNE and UMAP

t-SNE (t-Distributed Stochastic Neighbor Embedding) visualization in biomedical fields has
been recently growing in popularity, particularly for high-dimension single-cell sequencing data. However, its application in the detection of DM still remains rare and limited. UMAP (Uniform Manifold Approximation and Projection) is widely applied in data visualization and dimension reduction (McInnes et al., 2018).

Using t-SNE visualization, Gupta et al. successfully distinguished patients diagnosed with T2DM from the non-diabetic, healthy samples based on a dataset of 9,948 samples (Gupta et al., 2015). However, the visualization was unable to cluster the healthy and diagnosed individuals into individual clusters.

Bej et al. analyzed a dataset of 10,125 T2DM patients from the National Family Health Survey-4, involving many features such as medical history, dietary habits, addictions, and socioeconomic status. They found that the conventional application of UMAP was ineffective and uninformative. However, applying a feature type-wise clustering method, Bej et al. enabled visualizing the patients by clusters corresponding to different features. Their findings indicated that age and body mass index (BMI) are the most important factors for T2DM (Bej et al., 2020).

For data visualization, we first denoised the data by applying DWT on the HMP Stanford datasets (iHMP Research Network Consortium, 2014) into multi-view wavelet domain followed by applying t-SNE and UMAP to the transformed data. Our newly derived DWT-based t-SNE and UMAP methods on metabolomics, proteomics, RNA sequencing, targeted proteomics, and transcriptomics data enable better clustering than do those without DWT.

2 RESULTS

2.1 Data Pre-processing

We used CGM and ECG data from the D1NAMO dataset, a collection of data from 20 healthy individuals and 9 patients with type 1 DM. The data contain 4 day and collected ECG, CGM, food, and breathing variables. CGM data were measured in five-minute intervals before each meal and two hours afterwards, for a total of 6 times a day. We transformed the CGM unit from mmol/L to mg/dL. Alongside breathing data and the CGM data, the ECG data were collected at a rate of 250 Hz (Dubosson et al., 2018). The HRV data recorded many RR intervals in succession. We excluded a misclassified healthy control with type 1 DM.

The Human Microbiome Project (HMP) began in 2008 to investigate how microbiomes affect their hosts. Split up into two phases, HMP and Integrative HMP (iHMP or HMP2), HMP has collected over 10000 samples from 300 subjects. HMP took microbiomes from both healthy adults and diseased individuals. iHMP explored how microbiomes interacted with their hosts. Metabolism, immunity, and molecular activity were all investigated, along with how microbiomes might inform us about the onset of type 2 DM. We downloaded the data from the iPOP Project Data Portal from each of the abundance entries (Snyder Lab, n.d.). We describe results from amounts of certain types of bacteria in the gut, amounts of certain products of metabolism, untargeted profile of the amounts of certain proteins, RNA transcripts, targeted profile of the amounts of certain proteins, and also RNA transcripts.

For the glucose data from the DM patients in the D1NAMO dataset, we removed the manual glucose measurements so that the glucose data would solely consist of CGM data. We then used the glucose column, where each index represents five minutes. To preprocess the HRV data, we first had to find intervals of data without any outliers (which we defined as any HRV measurement of under 500 milliseconds or above 2000 milliseconds). We chose to use intervals of length 512 to be able to use DWT to denoise the data and generate TS data. We also decided to treat each interval as an independent sample, which gave us 3003 healthy HRV samples and 769 diabetic HRV samples. To fix this data imbalance, we used SMOTE Tomek resampling, which resulted in about equal numbers of healthy and diabetic HRV samples.

To preprocess the abundance matrices from the HMP dataset, we matched each subject’s VisitID to their IR/IS classification (either IR or IS; we removed subjects with Unknown). Subsequently, we removed subjects for whom there were missing values. In addition, we normalized the data by adding 0.1 and taking the log base 2. We then performed DWT on the normalized data. Figure 1 illustrates the workflow we followed.
2.2 CGM Forecast

In our research we used 3 different methods to predict blood glucose levels. Table 1 bolds the best RMSE and MAE for 30, 60, and 90 minutes, which are shared by KRR and ARIMA. In figure 2, KRR even correctly anticipates a drop in CGM for patient 3. These results support KRR and ARIMA as good forecasters of CGM, whereas Prophet did not yield good forecasts.

Table 1: Each metric is represented in the form mean (standard deviation) based on the results of each patient.

<table>
<thead>
<tr>
<th></th>
<th>30 min</th>
<th>60 min</th>
<th>90 min</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RMSE</td>
<td>MAE</td>
<td>RMSE</td>
</tr>
<tr>
<td>KRR</td>
<td>24.27</td>
<td>26.99</td>
<td>43.05</td>
</tr>
<tr>
<td></td>
<td>(25.37)</td>
<td>(29.09)</td>
<td>(44.74)</td>
</tr>
<tr>
<td>ARIMA</td>
<td>25.92</td>
<td>17.50</td>
<td>54.41</td>
</tr>
<tr>
<td></td>
<td>(7.22)</td>
<td>(4.23)</td>
<td>(16.95)</td>
</tr>
<tr>
<td>Prophet</td>
<td>60.43</td>
<td>45.79</td>
<td>67.17</td>
</tr>
<tr>
<td></td>
<td>(26.27)</td>
<td>(21.69)</td>
<td>(28.17)</td>
</tr>
</tbody>
</table>

The best metrics for each category are bolded.

Of the four, KRR had the lowest RMSE for each prediction horizon while ARIMA had a lower MAE. Because of this, KRR may not be a reliable model to use when forecasting blood glucose. KRR had difficulty predicting when blood glucose would suddenly rise or drop. ARIMA had the lowest standard deviations of any group. This is represented in its MAE, where it consistently has the lowest value. Prophet performed poorly. Its RMSE and STD were both relatively high.

A limitation in our work was the limited datasets that were available to us. The small training size that we had, nine diabetic patients, causes our models difficult to generalize. The accuracy of our models was most likely negatively affected by the shorter time series data. For future research, a larger dataset of patients would be desirable.

2.3 Diabetes Prediction using HRV

In this section, we assess machine learning algorithms on data transformed in a variety of ways using DWT (Table 2). RF algorithm on HRV signals transformed to the Wavelet Domain (TS) and resampled using the SMOTE Tomek method; the algorithm yielded an accuracy of 91.9%, precision of 95.5%, and recall of 87.9%. These results, while the accuracy and recall are lower and the precision is higher, are comparable to those of Acharya. Moreover, Acharya used 1000 HRV samples at a time while we only used 512; as a consequence, our model classifies HRV signals with nearly as much accuracy while taking about half as much time to record a patient’s HRV.

Table 2: Best overall results out of all data transformations tested.

<table>
<thead>
<tr>
<th></th>
<th>Accuracy</th>
<th>Balanced Accuracy</th>
<th>Precision</th>
<th>Recall</th>
<th>ROC AUC</th>
<th>F1</th>
<th>AUC Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF</td>
<td>0.919</td>
<td>0.919</td>
<td>0.955</td>
<td>0.880</td>
<td>0.968</td>
<td>0.916</td>
<td></td>
</tr>
<tr>
<td>XGBoost</td>
<td>0.788</td>
<td>0.788</td>
<td>0.788</td>
<td>0.789</td>
<td>0.871</td>
<td>0.788</td>
<td></td>
</tr>
<tr>
<td>SVM</td>
<td>0.775</td>
<td>0.775</td>
<td>0.771</td>
<td>0.783</td>
<td>0.853</td>
<td>0.777</td>
<td></td>
</tr>
</tbody>
</table>

RF: Random Forest. The best metrics for each category are bolded.

As shown in Figure 3, it appears that the best recall of 88.4% occurred with RF for classifying HRV signals. For RF, all data were classified with comparable ROC AUC and average precision (Figure 4 & 5). Nevertheless, (TS)* data performed best in diagnosing diabetes.
Figure 3: The four best confusion matrices for Random Forest are (TS)*, a1, TS, and No DWT, in that order. Each entry is of the form mean ± standard deviation.

Figure 4: The four best ROC curves for Random Forest are (TS)*, a1, TS, and No DWT with mean AUC of 0.97, 0.96, 0.96, and 0.96 respectively. Each ROC curve displays 10 folds as well as a mean curve (mean ± standard deviation).

Previously, Acharya et al., using the Decision Tree algorithm on the same HRV signals transformed using DWT decomposition up to 5 levels, obtained an accuracy of 92.64%, precision of 92.59%, and recall of 92.68% (Acharya et al., 2015).

However, compared to the neural networks of other papers, RF on HRV data transformed to the Wavelet Domain performed poor. Seyd P.T.’s ANN

Figure 5: The four best Precision v. recall curves for Random Forest are (TS)*, a1, TS, and No DWT with mean average precision of 0.97, 0.96, 0.97, and 0.97 respectively. Displayed are the results from each of the 10 folds.

Performing with an accuracy of 93.08%, precision of 96.67%, and recall of 89.23%. Of note, RF is a simpler model than a neural network, so our model is promising for relatively fast and accurate DM diagnosis for use by clinicians.

However, a limitation of our classification work with HRV is that we resample the HRV signals to correct the data imbalance between healthy and DM. This means that some of the samples may not be accurate representations of HRV. Also, given that we do not know how long the DM patients in the D1NAMO dataset have had DM, and as DM worsens cardiac health over time, it is most likely that people with DM who have not had DM for very long may be classified with lower accuracy. Thus, for future research, we recommend using RF and transforming the HRV signals to the DWT domain using DWT on HRV signals from DM patients who have not had the condition for more than a few years.

2.4 t-SNE and UMAP

Given the high complexity of the HMP Stanford datasets, t-SNE and UMAP figures reflect the heterogeneity and homogeneity in different datasets. Both t-SNE and UMAP enable successfully distinguishing several qualities within prediabetes patients. We first applied 6 different treatments for the dataset, namely: Preprocessed, Preprocessed with Wavelet Denoise, Preprocessed in Wavelet Domain,
Normalized, Normalized with Wavelet Denoise, and Normalized in Wavelet Domain. In addition, t-SNE and UMAP were each used to test the separability of 5 different bases for separation, which include: Gender, Race, Insulin Resistance (IR)/Insulin Sensitivity (IS) Classification, Age Group, and BMI Group.

We were able to utilize UMAP to obtain well-defined clusters, particularly in gender classification (Figure 6) to the Targeted Proteomics dataset.

![Figure 6: UMAP on the Targeted Proteomics dataset.](image)

The ‘Preprocessed’ data presents two clusters, largely separating males and females. On the other hand, the ‘Normalized’ data presents a scatter of points, with the males and females largely mixed together. The ‘Preprocessed WT Domain’ figure displays two compact clusters, comparatively cleaner than both the ‘Wavelet Denoised’ and preprocessed figures.

Of the Lipidomics dataset figures, the t-SNE results contain noticeable clusters for Black and Hispanic, particularly in the ‘Preprocessed’, “Denoised”, and ‘WT Domain’ with figure presenting a more compact result (Figure 7).

![Figure 7: t-SNE on the Lipidomics dataset.](image)

We analyzed the Cytokine dataset, resulting in distinct BMI-based clusters observed with both the t-SNE and UMAP. In t-SNE, the Figure 8 shows that ‘Denoised’ and ‘Normalized Denoised’ pretreated datasets present more compact clusters and points. The other classification subjects also aggregate into tight, compact clusters. It appears that certain prediabetic subjects share different characteristics, possibly presenting a stratification of these patients in each cluster (Figure 8).

![Figure 8: t-SNE on Cytokine abundance dataset.](image)

Of the Transcriptomics dataset figures, the UMAP results contain well-defined clusters, particularly in gender classification (Figure 9). In all gender classification figures for Transcriptomics, we observed a gender-based cluster pattern. Thus, it has implications in gender-based stratification of patients in precision medicine for prediabetic patients.

![Figure 9: UMAP on the Transcriptomics dataset.](image)

The application of UMAP on the Gut 16 Microbiome dataset resulted in clear separation based on IR/IS classification (Figure 10). These results allow us to observe the patterns in clusters in the set of prediabetic subjects, based on gut microbiomes.

![Figure 10: UMAP on the Gut 16 Microbiome dataset.](image)
2.5 Prediabetes Insulin Resistance Prediction

For SVM on the metabolomics data, both the normalized and (TS)* normalized data obtained the best results, with 77.7% accuracy, 75.0% balanced accuracy, 100% precision, 50% recall, AUC of 1.000, and F1 of 0.661. Similarly, the best results for SVM on the proteomics data were from the normalized and TS normalized data, bolded in Table 3, with 91.6% accuracy, 90.5% balanced accuracy, 100% precision, 81.1% recall, AUC of 1.000, and F1 of 0.892. We observed a similar propensity for SVM on the Targ.proteomics data and the Transcriptomics data. The normalized and TS normalized data enable perfect accuracy, balanced accuracy, precision, recall, AUC, and F1. For SVM on the Transcriptomics data, denoised normalized data also got perfect metrics.

Table 3: Results of classifying SVM model data as IR or IS on Proteomics.

<table>
<thead>
<tr>
<th>Data Type</th>
<th>Accuracy</th>
<th>Balanced Accuracy</th>
<th>Precision</th>
<th>Recall</th>
<th>ROC AUC</th>
<th>F1 Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>No DWT</td>
<td>0.558</td>
<td>0.500</td>
<td>0.000</td>
<td>0.000</td>
<td>0.500</td>
<td>0.000</td>
</tr>
<tr>
<td>(TS)*</td>
<td>0.558</td>
<td>0.500</td>
<td>0.000</td>
<td>0.000</td>
<td>0.500</td>
<td>0.000</td>
</tr>
<tr>
<td>Denoised</td>
<td>0.558</td>
<td>0.500</td>
<td>0.000</td>
<td>0.000</td>
<td>0.500</td>
<td>0.000</td>
</tr>
<tr>
<td>N</td>
<td>0.916</td>
<td>0.905</td>
<td>1.000</td>
<td>0.811</td>
<td>1.000</td>
<td>0.892</td>
</tr>
<tr>
<td>(TS)* N</td>
<td>0.916</td>
<td>0.905</td>
<td>1.000</td>
<td>0.811</td>
<td>1.000</td>
<td>0.892</td>
</tr>
<tr>
<td>Denoised N</td>
<td>0.558</td>
<td>0.500</td>
<td>0.000</td>
<td>0.000</td>
<td>0.500</td>
<td>0.000</td>
</tr>
</tbody>
</table>

N means Normalized. The best metrics for each model are bolded.

On the other hand, the TS data yielded the best overall results for Random Forest on the RNAseq abundance data (67.2% accuracy, 51.0% balanced accuracy, 30.0% precision, 2.0% recall, F1 of 0.037; denoised got an AUC of 0.988). The TS data also did best overall for Random Forest on the Transcriptomics data, with 92.2% accuracy, 91.2% balanced accuracy, 84.5% recall, AUC of 0.990, and F1 of 0.890, though the results for the denoised data are comparable (had highest precision of 98.0%). Thus, across all HMP data, DWT transformed data yielded better performance in predicting IR or IS in prediabetes.

Our application of the DWT based two experimental visualization techniques to the HMP Stanford dataset hold promising potential in allowing biomedical specialists to interpret the data by studying different visualizations of the dataset. We envision that our approach has the potential to uncover correlations between certain microbiomes and attributes in prediabetes patients, holding a promise for earlier detection and investigation of diabetic behavior in patients.

3 CONCLUSIONS

With our decomposed and denoised DM data, we enhance HRV based DM diagnosis, data visualization of the disparities in HMP data demographic features, and insulin resistance prediction. We forecast continuous glucose monitoring (CGM) 90 minutes in the future because CGM is unable to provide real-time blood glucose measurements. Moreover, we achieved 91.9% T1D diagnosis accuracy using Random Forest on data transformed with DWT, holding the potential for usage in clinics. Furthermore, our DWT-based t-SNE and UMAP explorative analysis of HMP data supports subtypes of prediabetic patients stratified by sex, race, and age. Thus, DWT-based transformations on metabolomics, proteomics, RNA sequencing, targeted proteomics, and Transcriptomics data yield better separation and clearer clusters than those without DWT.

Our results have implications in precision medicine. Precision medicine is based on stratification of patients, taking into account personal lifestyle, genetic information, and biomarkers. Our research involves two datasets, the HMP Stanford dataset along with the D1NAMO dataset, both of which contain extremely detailed information upon microbiomes, protein structure, CGM, ECG, food consumption, and other vital data, suitable for exemplification of machine learning modeling in precision medicine. Taken together, DWT-based...
machine learning approaches enable a fine resolution of subtyping DM towards precision medicine.

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