Incidence Rate Prediction of Diabetes from Medical Checkup Data

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SUMMARY To promote effective guidance by health checkup results, this paper predicts a likelihood of developing lifestyle-related diseases from health checkup data. In this paper, we focus on the fluctuation of hemoglobin A1c (HbA1c) value, which deeply connected with diabetes onset. Here we predict incensement of HbA1c value and examine which kind of health checkup item has important role for HbA1c fluctuation. Our experimental results show that, when we classify the subjects according to their gender and triglyceride (TG) fluctuation value, we will effectively evaluate the risk of diabetes onset for each class.

key words: specific health examination, lifestyle-related disease, machine learning

1. Introduction

Japanese Ministry of Health, Labour and Welfare reported that the percentage of highly suspicious of diabetes is 15.5% for male and 9.8% for female in 2014\textsuperscript{[1]}. To decrease that the percentage of highly suspicious of diabetes is 15.5%, Japanese Ministry of Health, Labour and Welfare reported. A1c (HbA1c) value, which deeply connected with diabetes onset. Here we focus on fluctuations in health checkup results. In this paper, we predict fluctuation of hemoglobin A1c (HbA1c) value, which deeply connected with diabetes onset. Here we focus on fluctuations in consecutive yearly health checkup data to investigate which element is related to HbA1c behavior.

To establish a framework to evaluate medical checkup data consisting on various domains, we first describe a fuzzy calculation method \textsuperscript{[2], [3]}. We also analyze relationships between HbA1c and other items in specific health examination data, as one of the basic researches to establish the above measures, using self-organizing maps (SOMs) \textsuperscript{[4]}. In this paper, to simplify the agenda, we are going to evaluate the risk of developing diabetes from health checkup data by using machine learning techniques; Haffner et al. \textsuperscript{[5]} use multivariate analysis, Wilson et al. \textsuperscript{[6]} use regression model and Yu et al. \textsuperscript{[7]} use support vector machine (SVM) to predict diabetes risk. In \textsuperscript{[8]}, Khalil et al. showed that Randomized Tree (RT) \textsuperscript{[9]} based method outperformed other machine learning meth-

Because this dataset has many missing value, we cannot achieve enough number of successive three-year data. In this dataset, 1569 subjects exceed the HbA1c threshold at second year. So its probability is only 1.75%. Figure 1 shows distribution of HbA1c value at 1st year, solid line represents the number of subjects who exceed the threshold in next year, and dashed line represents who did not exceed the threshold. From the figure, we can see that when a subject has enough small value at 1st-year, most of subjects doesn’t exceed the HbA1c threshold in next year.

So, we narrow the range of 1st year HbA1c value as $5.8 \leq \text{HbA1c} \leq 6.0$. In this condition, 1265 out of 9605 subjects exceeds the threshold in next year and its probability becomes 13.2%. For these dataset, we calculate yearly difference of following eleven health checkup data: body mass index (BMI), triglyceride (TG), high-density lipoprotein cholesterol (HDL), low-density lipoprotein cholesterol (LDL), systolic blood pressure (SBP), gamma-glutamyl trans peptidase (GTP), creatinine (CRE), serum glutamic-oxaloacetic transaminase (GOT), serum glutamate pyruvate transaminase (GPT), uric acid (UA) and waist circumference, to predict the HbA1c value exceeds the threshold in second year.

There are many studies to predict developing diabetes from health checkup data by using machine learning techniques; Haffner et al. \textsuperscript{[5]} use multivariate analysis, Wilson et al. \textsuperscript{[6]} use regression model and Yu et al. \textsuperscript{[7]} use support vector machine (SVM) to predict diabetes risk. In \textsuperscript{[8]}, Khalil et al. showed that Randomized Tree (RT) \textsuperscript{[9]} based method outperformed other machine learning meth-


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ods: SVM, bagging and boosting, in terms of the area under the receiver operating characteristic (ROC) curve (AUC).

In this paper, we also employ RT method to predict developing diabetes. However, our purpose is not further improvement of prediction accuracy but establishing effective report to rise health consciousness of individual subjects. By the analysis of prediction results, we can see that what kind of health checkup item has relation with HbA1c increment and we classify the type of diabetes onset. By evaluating the risk of developing diabetes for each class, we can make personal advice which propose numerical targets of health checkup items and show risk reduction by the achievement.

2. Predict HbA1c Increment from Fluctuation of the Other Health Checkup Items

To predict a subject’s HbA1c value exceeds the threshold in 2nd year or not, we use RT method. The RT (sometimes called random forest) is an ensemble learning method for classification, that fits a number of decision tree (DT) classifiers on various sub-samples of the dataset and use averaging to improve the predictive accuracy and control over-fitting. We use the eleven health checkup data as an input vector, and classified whether the HbA1c value exceed the threshold in second year or not. By analyzing the prediction result, we can see what kind of parameter will change with HbA1c.

2.1 Baseline Performance

First of all, by using RT, we test the prediction for 9605 subjects, whose HbA1c value was between 5.8 and 6.0 at 1st year. Figure 2 shows a prediction result as ROC curve. In this figure, horizontal axis represents false positive rate (FPR); the rate of healthy subjects who incorrectly receive a positive prediction result, and vertical axis represents true positive rate (TPR); the rate of unhealthy subjects who receive correct prediction result. The AUC value, which is a common criterion for evaluating ROC curve, becomes 0.67 for RT. This is a baseline performance of this prediction.

From the preliminary experiments, we select RT parameters as follows; max depth is 4 and number of trees is 100.

Here, we also show results of other machine learning methods; DT and logistic regression model (LRM) in Fig. 2. From the results, RT slightly outperforms LRM (AUC = 0.66) and has advantage against to DT (AUC = 0.63).

2.2 Tendency by Gender

Next, we divide the health checkup data by gender, because our earlier study indicates that there exist several routes to diabetes onset, especially, sexual difference has strong influence to HbA1c increment [4]. In the dataset, we have 3636 male subjects and 5761 female subjects who has 1st year HbA1c value between 5.8 and 6.0. For each gender group, we apply RT to predict whether the subject exceeds HbA1c threshold in next year or not.

Figure 3 shows a prediction result by ROC curve. Here, we select RT parameters as follows; number of trees = 100, max depth of tree = 3, and we apply 10-fold cross validation to make the ROC curve by changing weight parameter, which can control class priority. Here, we also show AUC value, which represent the prediction accuracy; 1 represents a perfect prediction and .5 represents a worthless prediction.

From this figure, we can see that the accuracy of male prediction is superior to female’s one. This result implies that the relationship between HbA1c fluctuation and others will have simpler connection than female’s one. To discuss this, we calculate feature importance of each health checkup item for both groups. Here, the importance is the Gini importance, which is computed as the total reduction of the criterion brought by that feature. Figure 4 (a) and 4 (b) shows the feature importance of RT classifier for male and female, respectively. From the result, BMI fluctuation is most important for predicting whether HbA1c excess the threshold, and second and third important item is GPT and GTP which represents liver function for both gender group. On the other hand, waist fluctuation is important only for male group.

2.3 Simple Grouping of Subjects

There will be several causes for increment of HbA1c, so it
will be difficult to predict whole subjects by single predictor. To confirm this assumption, here we test several simple grouping of the subjects. In this section, we pick up three health checkup item, BMI, GPT and TG. As shown in Fig. 4, BMI and GPT have enough importance for predicting HbA1c increment, and our previous study [4] also indicate that TG has some sort of relation with HbA1c increment. Here we simply divide the dataset whether the item value was increased or decreased for each gender set.

Table 1 shows the number of subjects for each group and their prediction accuracy by AUC value. The increment of selected three items has adverse effect for health, so the increased groups have larger HbA1c excess ratio. Contrary to our expression, prediction accuracy drops by grouping at most cases. To separate adverse effect of reducing the number of training samples, we examine the relation between number of training samples and prediction accuracy. Figure 5 shows AUC comparison against to the number of training samples. From these results, we can see that grouping by BMI decreases prediction accuracy significantly. As shown in Fig. 5, BMI fluctuation is a most significant clue for prediction, but the dataset dividing by BMI fluctuation will lose the advantage and complicate the problem.

Figure 6 shows examples of DT for predicting whether HbA1c value exceeds threshold at next year or not. In this figure, TP represents true-positive, which can correctly predict the HbA1c exceedance, and FP, TN, FN are false-negative, true-negative and false-negative, respectively. Figure 6 (a) shows a tree for whole female dataset and Fig. 6 (b) shows a tree for female who increase her TG value in next year. From these figures we can confirm that BMI fluctuation plays important role for decision in each cases, and by grouping dataset, we can simplify the tree and improve its accuracy for HbA1c prediction.

### Table 1  HbA1c prediction accuracy by grouping.

<table>
<thead>
<tr>
<th>Gender</th>
<th>Group</th>
<th>Subjects</th>
<th>HbA1c excess number ratio(%)</th>
<th>AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>3636</td>
<td>552</td>
<td>15.2</td>
<td>0.698</td>
</tr>
<tr>
<td>Δ BMI&gt;0</td>
<td>1637</td>
<td>361</td>
<td>22.1</td>
<td>0.652</td>
</tr>
<tr>
<td>Δ BMI&lt;0</td>
<td>1999</td>
<td>191</td>
<td>9.6</td>
<td>0.618</td>
</tr>
<tr>
<td>Male</td>
<td>Δ GPT&gt;0</td>
<td>1796</td>
<td>20.0</td>
<td>0.668</td>
</tr>
<tr>
<td></td>
<td>Δ GPT&lt;0</td>
<td>1840</td>
<td>10.5</td>
<td>0.661</td>
</tr>
<tr>
<td></td>
<td>Δ TG&gt;0</td>
<td>1906</td>
<td>17.3</td>
<td>0.689</td>
</tr>
<tr>
<td></td>
<td>Δ TG&lt;0</td>
<td>1730</td>
<td>12.8</td>
<td>0.683</td>
</tr>
<tr>
<td>All</td>
<td>5671</td>
<td>664</td>
<td>11.7</td>
<td>0.644</td>
</tr>
<tr>
<td>Δ BMI&gt;0</td>
<td>2584</td>
<td>428</td>
<td>16.6</td>
<td>0.585</td>
</tr>
<tr>
<td>Δ BMI&lt;0</td>
<td>3087</td>
<td>236</td>
<td>7.6</td>
<td>0.558</td>
</tr>
<tr>
<td>Female</td>
<td>Δ GPT&gt;0</td>
<td>2720</td>
<td>13.8</td>
<td>0.638</td>
</tr>
<tr>
<td></td>
<td>Δ GPT&lt;0</td>
<td>2951</td>
<td>9.8</td>
<td>0.606</td>
</tr>
<tr>
<td></td>
<td>Δ TG&gt;0</td>
<td>2799</td>
<td>12.6</td>
<td>0.647</td>
</tr>
<tr>
<td></td>
<td>Δ TG&lt;0</td>
<td>2872</td>
<td>10.8</td>
<td>0.641</td>
</tr>
</tbody>
</table>

![Figure 4](image1.png)  
**Fig. 4** Feature importance of each health checkup item.

![Figure 5](image2.png)  
**Fig. 5** Average AUC of grouping dataset according to the number of samples.

![Figure 6](image3.png)  
**Fig. 6** Examples of DT for predicting whether HbA1c value exceeds threshold at next year or not.
3. Incidence Rate Prediction by Using Decision Tree

In the previous section, we are going to predict whether the HbA1c value exceeds a threshold or not, by using fluctuation of other 11 health checkup items, but its prediction accuracy is not so high. This is because that there are many causes to onset diabetes and subjects have wide individual differences. However, from the generated decision tree, we can estimate incidence rate of HbA1c exceedance in detail. It will be useful to rise health consciousness by showing specific guidance.

Figure 7 shows a decision tree to predict the HbA1c exceedance for female dataset whose HbA1c value of the first year was 5.8. The total exceedance risk of female, whose HbA1c value was 5.8, is 4.5%, but if the person increase BMI more than 0.95 and GTP more than 6.5 until next year, her exceedance risk becomes 27.9%. On the other hand, if she can keep BMI increment under 0.95, her exceedance risk falls below 3.5%. A specific guidance showing numerical target will enhance the motivation to avoid lifestyle-related diseases.

4. Conclusion

In this paper, we have tried to predict the HbA1c increment by a Machine Learning method from 11 health checkup data fluctuations. Some experimental result show that by grouping subject according to their TG fluctuation, we can slightly improve the prediction accuracy. We also show that by showing tree structure, we will make an effective health guidance by showing specific numerical target. In future study, we are going to improve prediction accuracy of diabetes incidence, by establishing effective classification of subjects.

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References


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