Building a DNA Methylation Aging Clock Model on Less Labelled Data Using Item Response Theory

Keiji Yasuda¹,², Miyuki Nakamura¹, Masatoshi Nagata¹ and Masaru Honjo¹
¹KDDI Research, Inc., 2-1-15 Ohara, Fujimino, Saitama, 356-8502, Japan
²Nara Institutes of Science Technology, 8916-5 Takayama-cho, Ikoma, Nara 630-0192, Japan

Abstract: A method is proposed for DNA methylation analysis using item response theory. The analysis method consists of two steps: a cytosine phosphate guanine (CpG) sites selection step, and a parameters estimation step. Experiments are carried out to compare several CpG site selection conditions and evaluate if item response theory (IRT) can be applied to methylation analysis. According to the results of an experiment on public data measured by infinium HumanMethylation450 BeadChip, even under the condition of less age-labelled epigenetic data, CpG site filtering works well and the following IRT-based epigenetic clock model produced precise performance.

1 INTRODUCTION

The development of DNA sequence analysis technologies has enabled not only DNA genotyping but also measurement of epigenetic information such as DNA methylation level. DNA methylation phenomena are often observed at cytosine phosphate guanine (CpG) sites where a cytosine base is followed by guanine base.

By using methylation analysis, some molecular biology studies have revealed the impacts of DNA methylation within an organism such as repression of gene expression (Jones, 2012). Other studies (Horvath, 2013, Hannum et al., 2013, Liu et al., 2020) in this field have identified the relationship between DNA methylation and aging. This research has resulted in the development of several analysis methods that use DNA methylation information as aging indicators, often referred to as the epigenetic clock.

The main idea of conventional epigenetic clock research is to use methylation levels of CpG sites to predict biological age using a regression model. Since the dimension size of independent variables is very large, regularized regression models are often used to avoid the effect of multicollinearity.

This paper describes a new methylome analysis method for the epigenetic clock. The method incorporates item response theory (IRT) which has been developed and used in the field of psychometric research. This method has two strong advantages. One is robustness against methylation measurement noise such as the batch effect. The other is its applicability to unlabelled (age unknown) epigenome data.

In this paper, section 2 describes related work in the field of epigenetic clock research. Section 3 explains the proposed analysis method including formulation of IRT. Section 4 shows the experimental results. Finally, section 5 concludes the paper.

2 RELATED WORK

It is known that the DNA methylation pattern in the human genome is associated with individual chronological age. Some specific genomic position (CpG site) shows a differential methylation level. To date, several epigenetic clock models have been proposed to predict biological/chronological age using differential methylation patterns (Horvath, 2013, Hannum et al., 2013). These epigenetic clocks consist of different subsets of CpG sites that are selected from 450K or more sites measured by Infinium® HumanMethylation450 BeadChip or a high-throughput sequencer. Notably, the subsets of CpG sites derived from different epigenetic clocks are...
highly divergent (Liu et al., 2020), implying that they reflect distinct aspects of biological age. For age prediction by epigenetic clocks, linear regression models, such as elastic net regression, have been widely used (Horvath 2013, Hannum et al., 2013, Field et al., 2018). In addition to using linear regression models, neural network models have been developed in recent years to predict age (Galkin et al., 2021, Lima Camillo et al., 2022). However, it is known that the DNA methylation level at some CpG sites is susceptible to DNA methylation measurement platforms (Shu et al., 2020). Indeed, the predicted ages obtained using DNA methylation sometimes suffer from technical noise, resulting in relatively large errors of up to 9 years (Higgins-Chen et al., 2022).

The analysis method presented in this paper differs from conventional epigenetic clock research in two crucial aspects. One is the robustness against methylation measurement noise and the other is the ability to use data without an age label.

3 PROPOSED ANALYSIS METHOD

This section explains IRT and the methylome analysis method using IRT. First, formulation of IRT is shown. Then, we explain how IRT can be applied to methylation analysis.

3.1 Item Response Theory

Item response theory (IRT) is a psychometric model which enables the latent traits of human subjects to be estimated from test results. Currently, IRT is being used for crucial tests such as the Test of English for International Communication (TOEIC), Test of English as a Foreign Language (TOEFL) and the Graduate Record Examination (GRE).

IRT research has a long history and several models have been proposed since the advent of modern test theory (Rash, 1960), including 3-parameter logistic models (Barton, 1981), the graded response model (Samejima, 1969), and the continuous response model (Samejima, 1973). In our research, we use a 2-parameter logistic (2PL) model (Birnbaum, 1968), which is the second simplest model among several IRT models. The following parts of this subsection explain the formulation of the 2PL model.

As a precondition, the target test consists of multiple test items \((i \in I)\) which are answered by multiple human subjects \((s \in S)\). As a result of testing, we obtain the response matrix \(X_{i,s} \in \mathbb{R}^{I \times S}\) which has binary elements. Here 0 and 1 stand for the wrong and correct answer, respectively.

2PL expresses each item using 2 parameters \((a_i, b_i)\), which are item discrimination and item difficulty. By using a logistic function, \(p_{i,s}\), which is the probability that human subject \(s\) gives a correct answer to item \(i\), is modeled by the following formula

\[ P_{i,s}(x_{i,s} = 1|a_i, b_i) = \frac{1}{1 + e^{-d_i(a_i x_{i,s} - b_i)}} \] (1)

where \(D(= 1.7)\) is a scaling constant.

Given \(X\), the marginal maximum likelihood method (Hsieh, 2010) estimates item parameters \((a_i, b_i)\). Then, the latent traits of each human subject are estimated on \(X\) and estimated item parameters by the maximum likelihood method. In this study, we use this flow, which is the most basic approach for IRT parameter estimation. However, there are several different algorithms for IRT parameter estimation such as Bayesian estimation or deep learning (Hsieh, 2010, Yeung, 2019).

3.2 Progressive Methylation Status Modelling by IRT

An elastic net model, which is a frequently used regression model in conventional epigenetic clock research, selects useful CpG sites and builds a regression model at the same time. However, a 2PL IRT model does not suitably CpG site selection from all available CpG sites since this model presumes single dimensionality of latent traits, while available CpG sites can contain multiple latent traits. Hence, we apply CpG site selection from all available CpG sites as a first step. As the second step, we apply IRT to selected CpG sites. After item parameter estimation, we exclude CpG sites with too large \((a_i > 10)\) or too small \((a_i < 0.5)\) value discrimination parameters because such CpG sites are regarded as too peaky or less informative to be used for epigenetic clock research.

For CpG site selection, we compare several selection methods as follows

- **Selection with full data:** Selects the CpG sites by the correlation coefficient \((r)\) calculated by the following formula

\[ r_i = \frac{1}{\sqrt{\sum_{s=1}^{S}(z_{i,s}^{\text{aligned}} - \bar{z}_{i,s}^{\text{aligned}})^2 \cdot \sum_{s=1}^{S}(c_s - \bar{c}_s)^2}} \] (2)
Table 1: Data set for the experiments.

<table>
<thead>
<tr>
<th>Data set type</th>
<th>Average age</th>
<th>Minimum age</th>
<th>Maximum age</th>
<th># of subjects (S)</th>
<th># of CpG sites (I)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test set</td>
<td>54.22</td>
<td>20</td>
<td>99</td>
<td>142</td>
<td>13,751</td>
</tr>
<tr>
<td>Development set</td>
<td>56.30</td>
<td>21</td>
<td>90</td>
<td>142</td>
<td>13,751</td>
</tr>
<tr>
<td>Training set</td>
<td>54.91</td>
<td>20</td>
<td>112</td>
<td>1,138</td>
<td>13,751</td>
</tr>
</tbody>
</table>

Table 2: Experimental conditions for CpG site selection.

<table>
<thead>
<tr>
<th>Condition</th>
<th>CpG site selection</th>
<th>IRT (item parameter estimation)</th>
<th>Evaluation (Latent trait estimation)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data set</td>
<td>Feature(s)</td>
<td>Selection criteria</td>
<td>Data set</td>
</tr>
<tr>
<td>Selection with full data</td>
<td>Training set</td>
<td>Age, CpG Methylation rate (raw value)</td>
<td>Eq (2)</td>
</tr>
<tr>
<td>Selection with less data</td>
<td>Development set</td>
<td>Age, CpG Methylation rate (raw value)</td>
<td>Eq (2)</td>
</tr>
<tr>
<td>Selection without age information</td>
<td>Training set</td>
<td>CpG Methylation rate (raw value)</td>
<td>Eq (3)</td>
</tr>
</tbody>
</table>

where $c_s$ is the chronological age of human subjects $s$, and $z_{i,s}^{aligned}$ is preprocessed value of methylation rate at CpG site $i$ of human subject $s$. Details of the preprocessing are explained in 4.1.1. Here, the correlation coefficients are calculated on the training data set.

✓ **Selection with Less Data:** The only difference between this selection and the previously mentioned selection is the data set used for the coefficient calculation. Here, we use a small-sized development data set instead of a training set.

✓ **Selection without Age Information:** Selects CpG sites based on the Item-Test or Item-Total (I-T) correlation ($r_{IT}$) given by the following formula.

$$r_{IT} = \frac{1}{\sqrt{\sum_{i=1}^{I} (z_{i,s}^{aligned} - \bar{z}_{i,s}^{aligned})(t_s - t_s)}} \frac{1}{\sqrt{\sum_{i=1}^{I} (z_{i,s}^{aligned} - \bar{z}_{i,s}^{aligned})^2} \sqrt{\sum_{i=1}^{I} (t_s - t_s)^2}}$$  \hspace{1cm} (3)

where

$$t_s = \sum_{i=1}^{I} z_{i,s}^{aligned}$$ \hspace{1cm} (4)

This method is often used to filter out unsuitable test items (Henrysson, 1963), when the test is designed.

CpG site selection with full data is the same data usage setting as that employed in conventional research of epigenetic clock that uses age information and methylation rates of CpG sites for all human subjects. Semi-supervised selection and unsupervised settings are new settings for epigenetic clock research. These settings focus on effective utilization of epigenome data, which do not contain age information on human subjects. We refer to a setting of small-size well-regulated data and large-size not well-regulated data availability as “selection with less data setting.” And, “CpG site selection without an age information setting” is a setting where we have large-size data but the data has no age label for each human subject at all.

For the following IRT parts, we simply consider each selected CpG site as a test item, then apply IRT to estimate item (CpG sites) parameters and the DNA methylation aging clock as latent traits ($\theta_s$). Since IRT does not use age information for parameter estimation, it is expected to model progressive methylation status as a latent trait.

Apart from the single-cell experimental setting, methylation measurement results are given as averaged methylation rate on multiple cells. In such a case, we simply binary the methylation rate using a threshold of 0.5 before applying IRT. This binary preprocessing has both an advantage but also a disadvantage. The advantage is gaining robustness against methylation measurement noise such as the batch effect. The disadvantage is that binarization reduces the information. Although binarization preprocessing has this disadvantage, robustness against noise is a very important aspect especially when we use epigenetic data collected by other studies.
4 EXPERIMENTS

4.1 Experimental Settings

For the experiments, we used epigenome open data (Xion et al., 2022) measured by infinium HumanMethylation450 BeadChip (450K). The data consist of epigenetic read data from 21 body tissues. From these 21 tissues, we use whole blood data because the data of the tissue contains the largest number of human subjects.

4.1.1 Data and Pre-Processing

Before we carry out CpG site selection and IRT experiments, we apply the following pre-processing as data cleaning.

- **Step 1:** We discard subjects aged under 20 years. (1,422 subjects remained after this step.)
- **Step 2:** CpG sites containing missing values are discarded. (166,863 CpG sites remained after this step)
- **Step 3:** If the correlation coefficient calculated by Eq. 2 is the negative value for CpG $i$, we covert raw value of the methylated rate $z_{i,s}(\in Z)$ as follows

  $\begin{align*}
  z_{i,s}^{aligned} &= 1 - z_{i,s} \\
  \text{else} & \quad \text{for each } s \in S \\
  z_{i,s}^{aligned} &= z_{i,s}
  \end{align*}$

  The pre-processing in this step is to align methylation direction toward aging.

- **Step 4:** Randomly divide 1,422 subjects into three sets, the test set, development set and training set. (Each of these sets contain 142, 142 and 1,138 subjects, respectively.)
Step 5: Binarize methylation rate of each CpG site applying a threshold of 0.5 as follows

\[ \begin{align*} 
    &\text{If } (z_{ls} \leq 0.5) \\
    &\quad x_{ls} = 0 \\
    &\text{else} \\
    &\quad x_{ls} = 1 
\end{align*} \]

Step 6: Remove CpG sites if 10% or fewer subjects have a value of 0 or 1. (13,751 CpG sites remained after this step.)

Table 1 shows the statistics for the data sets used for CpG site selection and IRT experiments.

4.1.2 Experimental Conditions

As we explained in 3.2, we carried out CpG site selection experiments under three different conditions. Table 2 shows the data usage for CpG selection and the IRT experiments. As shown in the table, item parameter estimation and the evaluation process are performed under the same condition. Since the item parameter estimation process does not require a correct value for latent traits, we only use the binarized CpG methylated rate for the estimation. Each selection condition is evaluated by the results of the following IRT process. For the evaluation metric, we employ the correlation between chronological age and estimated latent traits on the test set. Thus, we use the age information of the test set for the evaluation.

4.2 Experimental Results

First, we show the experimental results of CpG site selection with full data. Here, we select the top 100 CpG sites that were highly correlated with subjects’ age on the training set. Figure 1 shows item characteristic curves which are the results of item parameter estimation on the 100 CpG sites. In this figure, the horizontal axis indicates latent trait (θ) and vertical axis indicates methylation probability, which is \( P(x_i = 1 | \theta) \). As shown in the figure, many of the 100 CpG sites tend to be methylated in high latent trait, which indicates an older person. However, there are still some CpG sites which are methylated in lower latent trait. By using the estimated item parameters, we estimate the latent trait of human subjects on the test set. Figure 2 shows the relationship between estimated latent trait and the chronological age of human subjects. As shown in the figure, there is high correlation between the two sets of values (\( r = 0.850 \)). These results demonstrate the efficiency of 2PL IRT for methylation analysis in epigenetic clock research.

Second, we show the results of CpG site selection with less data. Figure 3 shows the relationship between estimated latent traits and the chronological age of human subjects. Also 100 CpG sites are selected by the correlation with age on the development set. The only condition that differs between the settings of Fig. 2 and 3 is the data used for CpG site selection. As shown in the figure, a high correlation is observed between the two sets of values (\( r = 0.836 \)). These results show the possibility of using age label-less epigenetic data, provided smaller age labelled data are available.
Figure 6: Test information function of CpG site set selected without age information.

Third, we show the results of CpG site selection without age information. Figure 4 shows the relationship between estimated latent traits and the chronological age of human subjects in the case where age label-less selection is applied. Here, CpG sites are selected by IT correlation calculated by Eq. 3 on the training set. The same number of CpG sites as the previous experiments are used for item parameter estimation. Comparing the results in Fig 4 to Fig. 2, the correlation between the two sets of values is very low ($r = 0.528$). Figure 5 shows the item characteristic curves of the selected 100 CpG sites. Looking at the figure, variation in the $b_i$ parameter is very small. Most of them have a value greater than 0. These findings indicate that these CpG sites are not useful for estimating latent traits less than 0. Figure 6 shows fisher information ($I_{testInfo}(\theta)$) given by Eq. 5.

$$I_{testInfo}(\theta)=\sum_{i=1}^{l}D^2a_i^2p_i(\theta)(1-p_i(\theta)) \quad (5)$$

$I_{testInfo}(\theta)$ is also called as test information function which is used to calculate variance of estimated $\theta$ ($V[\hat{\theta}|\theta]$) by the following formula.

$$V[\hat{\theta}|\theta] = 1/I_{testInfo}(\theta) \quad (6)$$

In the figure, the vertical and horizontal axes indicate $I_{testInfo}$ and $\theta$, respectively. According to the figure, the set of CpG sites has a very small amount of information to estimate $\theta$ where it is less than 0. By incorporating viewpoint of test information in the future work, we may improve the deviation in low value range of $\theta$ ($\theta < 0$ in Fig 4).

$$b_i = \theta, \text{ when } p_i(x_i = 1) = 0.5$$

4.3 Discussion on Data Size

Here, we discuss data size for selection with less data condition. The previous subsection showed the results of the selection using the development set which consists of methylation data of 142 age-known human subjects. In this subsection, we change development set size from 10 to 142 subjects by random sampling from the original development set. Then, apply selection and IRT experiments in the same manner as the previous experiments.

Figure 7 shows the results of the experiments. In the figure, the horizontal axis indicates data size of the development set. And the vertical axis indicates correlation between the chronological age and latent trait on the test set. For each condition, top 100 CpG sites are selected based on correlation calculated by Eq. 2. Then, apply IRT parameters estimation for each selected CpG sites.

As shown the figure, estimated latent trait correlates well with chronological age, even if only 20 age-known subjects are available for the CpG site selection.

5 CONCLUSIONS AND FUTURE WORK

We proposed an IRT-based method for methylation analysis. First, the method selects CpG sites using a correlation-based metric, then applies IRT to selected data. In the experiments, we compared three selection conditions using open epigenome data measured on whole blood by infinium Human Methylation 450 BeadChip.
According to the experiments, CpG site selection with partially or fully age-labeled data works well with following IRT parameter estimations. Since the IRT parameter estimation part does not use any age information, the model is thought to estimate progressive methylation status as a latent trait. Meanwhile, CpG site selection with no age information does not work well. Further analysis shows that the selected CpG sites have small variation in the difficulty parameter ($b_i$), and this causes deviation in low value range of $\theta$. In future research, we will improve age label-less selection by considering not only IT-correlation but also the test information of a set of CpG sites. Another unfinished work is an evaluation of each item (CpG site) based on item fit statistics. By removing the CpG site whose response does not fit to logistic function, we may improve the analysis results.

Differing from the conventional regression method, IRT-based analysis gives rich information such as discrimination and difficulty parameters for each CpG site. These parameters are thought to indicate “how fast” and “when” CpG sites will be methylated. We will also advance our research in relation to these points of view.

REFERENCES


