Predicting Glaucomatous Progression with Piecewise Regression Model from Heterogeneous Medical Data

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Abstract: This study aims to accurately predict glaucomatous visual-field loss from patient disease data. In general, medical data show two kinds of heterogeneity: 1) internal heterogeneity, in which the phase of disease progression changes in an individual patient's time series dataset; and 2) external heterogeneity, in which the trends of disease progression differ among patients. Although some previous methods have addressed the external heterogeneity, the internal heterogeneity has never been taken into account in predictions of glaucomatous progression. Here, we developed a novel framework for dealing with the two kinds of heterogeneity to predict glaucomatous progression using a piecewise linear regression (PLR) model. We empirically demonstrate that our method significantly improves the accuracy of predicting visual-field loss compared with existing methods, and can successfully treat the two kinds of heterogeneity often observed in medical data.

1 INTRODUCTION

1.1 Motivation of our Study

The aim of our study is to construct a novel method for the treatment of heterogeneous medical data in the prediction of disease progression. We specifically focus on data from patients with glaucoma to predict visual-field loss progression. Medical datasets are heterogeneous from the following two aspects: external heterogeneity and internal heterogeneity (Fig. 1). External heterogeneity generally refers to the relationship of measured data among patients; e.g., the rates of the disease progression differ from patient to patient. By contrast, internal heterogeneity occurs within an individual patient; e.g., changes in the phase of disease progression over time. Therefore, in order to obtain comprehensive knowledge and trends from medical data, a method for appropriately dealing with the characteristic heterogeneity is required.

It is not straightforward to treat heterogeneous medical data because these two kinds of heterogeneity must be resolved in different ways. The main challenge in this respect is due to the specific structure of large medical datasets. Although the datasets are often composed of data from a large number of patients, data for each patient are usually limited be-

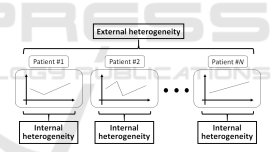


Figure 1: The two kinds of heterogeneity observed in medical datasets. External heterogeneity is caused by differences among patients, whereas internal heterogeneity is caused by changes in each patient's state over time.

cause of the high costs related to diagnosis, both for the patients and clinicians. In particular, a detailed medical examination requires well-trained clinicians, special medical equipment, and a long period of diagnosis. In this paper, we refer to this specific structure of medical data as *medical-data-structure-difficulty*. This difficulty limits the ability to construct a reliable predictive model for each patient using only the patient's information. One way to solve this problem is to take advantage of hidden relationships among patients. However, mining for such hidden relationships is often difficult because of the heterogeneous nature of medical data described above. Therefore, accurate analysis of medical data requires a method for overcoming the two heterogeneity problems and

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the medical-data-structure-difficulty simultaneously.

Various methods have been proposed to overcome the medical-data-structure-difficulty in a medical dataset, which have mostly involved incorporating information from other patients to improve the prediction accuracy (Liang, Z. et al., 2013; Maya, S. et al., 2014; Murata, H. et al., 2014; Morino, K. et al., 2015). Therefore, appropriate information is collected from a patient dataset and a tuned predictor is constructed for the target patient. We here refer to these types of methods as "collective methods." In collective methods, we should carefully analyze the heterogeneity to collect data appropriately. Most of the existing collective methods for dealing with data from glaucoma patients have focused on resolving the external heterogeneity problem. We here propose a new collective method that achieves the better prediction accuracy than existing methods, because our method can cope with both the external and internal heterogeneity in the medical dataset.

Glaucoma data, the focus of our study, critically contain both kinds of heterogeneity as well as the medical-data-structure-difficulty mentioned above. Glaucoma is an eye disease that causes progressive damage to a patient's visual field, which can ultimately lead to blindness, and is the second-leading cause of blindness worldwide (Kingman, S., 2004). Quigley et al. (Quigley, H. A. and Broman, A. T., 2006) estimated that nearly 80 million people will suffer from glaucoma by 2020. The glaucomatous visual-field loss is considered to be irreversible, but glaucomatous progression can be delayed with appropriate treatment. Therefore, suggesting an appropriate treatment plan at an early stage of disease progression is a critical factor for improving patients' quality of life. Accordingly, the development of methods for the early prediction of glaucoma progression is particularly important for effectively treating the disease.

Most of the existing glaucoma prediction methods involve analyses of visual-field data, which are associated with the aforementioned medical-datastructure-difficulty and external heterogeneity problem. However, in reality, the rate of glaucomatous progression changes over time for each eye (internal heterogeneity). Therefore, to improve prediction, a novel collective method is required to deal with the internal heterogeneity of data (within-eye level) in addition to the external heterogeneity (betweeneye level). The internal heterogeneity can be potentially captured with intraocular pressure (IOP) data. This is based on clinical evidence that the progression rate of glaucoma increases with an increase in the IOP value (AGIS Investigators, 2010; Collaborative Normal-Tension Glaucoma Study Group, 1998;

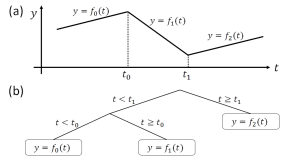


Figure 2: Schematic figure of the piecewise linear regression model. (a) The piecewise linear regression. (b) The tree structure expression corresponding to the piecewise linear regression lines shown in (a).

Satilmis, M. et al., 2003). However, previous collective models for predicting glaucomatous progression (Liang, Z. et al., 2013; Maya, S. et al., 2014; Murata, H. et al., 2014) have been trained only with visual-field data. In this paper, we outline the first application of IOP data to the prediction of glaucoma progression and achieve good prediction accuracy.

1.2 Heterogeneity of Glaucoma Data

Here, we introduce the heterogeneity of glaucoma.

Internal Heterogeneity: The progression rate of glaucoma essentially changes over several time points, even when considering the time series of one eye. Clinical knowledge suggests that the progression rate of glaucoma varies because of high IOP values (AGIS Investigators, 2010; Collaborative Normal-Tension Glaucoma Study Group, 1998; Satilmis, M. et al., 2003). This internal heterogeneity can be difficult to detect because only limited data can be obtained from the target eye at certain time points. Our proposed novel collective method resolves this internal heterogeneity difficulty by using IOP data.

External Heterogeneity: Glaucoma data are highly variable among eyes from the following four perspectives: (1) the disease stage at initial diagnosis; (2) the progression rate of glaucoma; (3) the average and fluctuation levels of IOP; (4) the minimum level of IOP that affects the progress of glaucoma.

1.3 Novelty and Significance

The novelty and significance are summarized below.

1) A Novel Framework for Solving the Heterogeneity Problems of Medical Data with a Collective Method. We here propose a novel collective *piecewise linear regression* (PLR) model to simultaneously deal with the two kinds of heterogeneity of medical data and the medical-data-structuredifficulty. In general, a PLR model is suitable for dealing with the internal heterogeneity problem of medical data; however, it cannot be easily applied to the problem of glaucoma progression prediction. A PLR model (Fig. 2(a)) can be interpreted as a treestructured model (Fig. 2(b)); i.e., the edges carry information about the segmentation, and the nodes carry information about the regression lines. This model is powerful because it can reflect the complex features of medical data by breaking it down into several pieces. However, this benefit comes with a disadvantage in that piecewise regression requires a large dataset for good prediction even if the complexity of the model or the depth of the tree structure is appropriately controlled. Therefore, the medical-datastructure-difficulty is a barrier to effectively analyzing the data with existing PLR methods. Our proposed method overcomes the problems of medical data.

A) **Application of a Collective PLR Model with Medical-data-structure-difficulty.** Our proposed method can be used to train a PLR model with heterogeneous medical data. As described above, only limited data can be obtained for each patient from a large medical dataset owing to the medical-data-structuredifficulty. Although effective training algorithms for a tree-structured model with a very large dataset have been intensively investigated (Natarajan, R. and Pednault, E., 2002; Vogel, D. S. et al., 2007), there have been few studies conducted to develop a training algorithm for this type of "big data." Therefore, our current study sheds new light on this common problem and offers a potential solution.

B) A Useful Framework for the Overall Optimization of a Piecewise Regression Model Considering External Heterogeneity. Our novel method optimizes the piecewise model as well as each regression line for each piece simultaneously. Our model (Fig. 3) consists of two parts: one that controls the model's complexity, and the other that controls the model's prediction accuracy. This clearly divided model structure enables the use of other collective regression algorithms besides those employed in this paper (Liang, Z. et al., 2013). We describe this feature in greater detail in Sec. 3.2. We optimized the whole model, including the segmentation and regression parameters, using data from similar eyes. We did this optimization by applying the statistical model selection criteria. Specifically, we examined a number of existing information criteria to investigate which gave the best prediction accuracy.

C) Good Framework of the Collective Piecewise Regression for Tackling Internal Heterogeneity. For the collective piecewise regression, our proposed method provides a good framework that can effec-

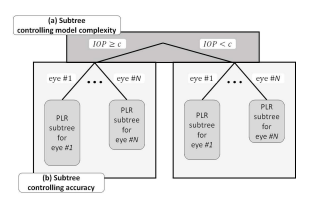


Figure 3: Hierarchical tree structure of our IOP-based piecewise model. The structure of the model can be separated into two parts (a) controlling the model complexity and (b) controlling the model accuracy. The PLR model for each eye also has its tree structure as shown in Fig. 2(b).

tively deal with internal heterogeneity. Our model was carefully designed to collect data of similar eyes while coping with the internal heterogeneity related to disease progression over time. Our method involves segmentation of time-series data, even within one patient's time course, followed by calculation of the regression lines for each piece. We assume that the gradient parameters for each piece in the same state are common and that the intercept parameters for each piece are different, so that the internal heterogeneity can be accurately expressed (see Step 4 in Fig. 5). We note that our model does not have only common gradient parameters but individualized intercept parameters to reflect each patient's characteristics. This individualization cannot be realized by simply sharing the parameters with all the patients. Hence, our model can represent each patient's glaucoma progression efficiently.

2) Big Impact on the Medical Field of Glaucoma

A) First Application of IOP to the Prediction of Glaucoma Progression. To the best of our knowledge, our proposed model represents the first application of IOP to the prediction of glaucoma progression. In existing analyses (Liang, Z. et al., 2013; Maya, S. et al., 2014; Murata, H. et al., 2014; Maya, S. et al., 2015; Holmin, C. and Krakau, C. E. T., 1982; Zhu, H. et al., 2014), only visual-field data have been used for prediction. Although clinical studies suggest the importance of IOP to understand disease progression (AGIS Investigators, 2010; Collaborative Normal-Tension Glaucoma Study Group, 1998; Satilmis, M. et al., 2003), the high external heterogeneity of IOP has limited its application to predictive models (see Sec. 1.2). We believe that this difficulty can be overcome with our novel collective method, which is expected to have a great impact on the medical field of glaucoma.

B) Wide Applicability to other Glaucomatous Prediction Models based on Visual-field Data. Any existing glaucoma prediction method can be converted into an IOP-based piecewise prediction model using our framework, which should increase its impact in the medical field of glaucoma. As mentioned in point 1-B), our model was developed using a general framework. Therefore, our model is not only applicable to various existing glaucoma prediction models but also to models that will be developed in the future. This wide applicability is practically important for the future progress of glaucoma prediction, since any great prediction model will lose its value after an improved method is proposed. Owing to its clearly segmentable structure, our model will be able to embrace other models, indicating that it has a long life span, and is flexible and widely applicable.

1.4 Related Works

Conventionally, linear regression analyses of visualfield time-series data for each eye have been used to predict the glaucomatous progression (Holmin, C. and Krakau, C. E. T., 1982). However, simple linear regression for time series of an individual eye is not effective when the number of data points is small, which is often the case for clinical data. Although various regression models have been applied to the prediction with only data from a target eye, the prediction accuracy was limited due to the shortage of data (Fujino Y. et al., 2015; Taketani Y. et al., 2015). Thus, we must overcome this shortage for better prediction.

To make up for this deficiency in clinical data, some recent studies have proposed collective methods that exploit visual-field data from other eyes to predict the glaucomatous progression in a target eye at an early stage of disease. These studies also proposed some methods for coping with the external heterogeneity and medical-data-structure-difficulty from a data-mining point of view. Liang et al. (Liang, Z. et al., 2013) proposed a spatio-temporal clusteringbased method. They collected similar eyes in terms of their spatial and temporal feature of progression. Then, they used data from the eyes similar to the target eye for prediction. Zhu et al. (Zhu, H. et al., 2014) used Bayesian inference to reflect the spatial correlation of progression. Maya et al. (Maya, S. et al., 2014) used tensor decomposition and a multitask-learning method to extract multiple features from a heterogeneous glaucoma dataset. Murata et al. (Murata, H. et al., 2014) used Bayesian linear regression to utilize the measurements of other eyes when making predictions of a target eye. Maya et al. (Maya, S. et al., Figure 4: Schematic figure of the glaucomatous progression. The total deviation (TD) values on a visual field are schematically shown in shades of gray. Darker mesh shades indicate a more defective visual field. The overall aim was to predict TD values at each mesh at a given target time. 2015) proposed a *hierarchical minimum description length (MDL)based clustering method* for finding progression patterns, and substituted the clusters used in Liang et al. (Liang, Z. et al., 2013) with their newly discovered clusters to more effectively predict glaucomatous progression. However, all of these studies focused on external heterogeneity and did not address the problem of internal heterogeneity.

To overcome the internal heterogeneity problem, we employed a PLR model for predicting glaucomatous progression. As we mentioned above, a PLR model potentially deals with the internal heterogeneity problem. However, it is widely known that sufficient data within an appropriate range are required in order to appropriately estimate each regression line. Therefore, a large dataset should be used in the learning phase, which can sometimes cause problems in applying this model to real situations. Although several studies have focused on training models for a tree structure with a massive dataset (Natarajan, R. and Pednault, E., 2002; Vogel, D. S. et al., 2007), there is barely any information on training a PLR model under a situation of medical-data-structure-difficulty.

1.5 Organization

The rest of this paper is organized as follows. In Section 2, we introduce some prior knowledge for understanding glaucomatous progression. In Section 3, we present our proposed framework for training a collective PLR model for tackling the internal and external heterogeneity. The results of experiments conducted to evaluate our framework with glaucoma dataset are described in Section 4. Finally, we provide an overall conclusion and summary of our method in Section 5.

2 PRELIMINARIES

2.1 Prediction of Glaucoma

Our aim was to predict the visual-field loss precisely at a given target time, as depicted in Fig. 4. In our dataset, the visual-field data were measured on 74 meshes of a visual field to obtain a *total deviation* (TD) value. This value represents the differences in the measured light sensitivity on each mesh compared to age-matched normative data. A negative TD value

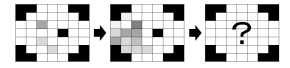


Figure 4: Schematic figure of the glaucomatous progression. The total deviation (TD) values on a visual field are schematically shown in shades of gray. Darker mesh shades indicate a more defective visual field. The overall aim was to predict TD values at each mesh at a given target time.

means that the sensitivity is worse than the normative sensitivity; thus, the TD value decreases as glaucoma progresses. The rates of progression differ among the meshes on a visual field; therefore, the TD value needs to be predicted independently for each mesh.

2.2 Effect of High IOP

Several ophthalmological studies have shown a relationship between IOP and glaucoma progression. The AGIS Investigators (AGIS Investigators, 2010) found that a group of patients with glaucomatous eyes and high IOP lost their visual fields more quickly than patients with lower IOP. The Collaborative Normal-Tension Glaucoma Study Group (Collaborative Normal-Tension Glaucoma Study Group, 1998) observed that the rate of glaucoma progression in the eyes of patients undergoing treatment, which involves reduction in IOP, was significantly lower than that of patients not receiving treatment for IOP reduction. Satilmis et al. (Satilmis, M. et al., 2003) demonstrated that the rate of glaucoma progression was related to the standard deviation in IOP.

Considering these results, the rates of the progression greatly change at some time points that can be detected with IOP values. This means that the internal heterogeneity in the glaucoma dataset can be captured with IOP. However, the relationship between IOP and the degree of glaucoma progression has not yet been fully investigated in ophthalmology. Hence, we do not employ the raw value of IOP as an explanatory variable but rather discretize it into "high" and "low" states to make segmentations of time series according to whether the undelying IOP is high or low. Further, the discretization of IOP into the "two" states makes it effcient to estimate the parameters of the model. This is because the more states we separate IOP into, the smaller the size of each piece becomes.

3 PROPOSED METHOD

3.1 Concept of Proposed Method

Our method makes it possible to decide the appropriate thresholds for separating time series and express the internally heterogeneous progress of glaucoma. We optimize the thresholds using an information criterion and estimate gradient and intercept parameters for each of the separated piece. The outline of our method is described below. Figure 5 helps to understand its procedure. We note that the following procedure was applied to one prediction-target eye.

Step 1: Normalization for the External Heterogeneity of IOP. The mean and standard deviation of the IOP distribution are highly variable among eyes. Therefore, the IOP distribution of each eye was normalized to effectively analyze the external heterogeneity of IOP.

Step 2: Selection of an IOP Threshold Value c. An IOP threshold value c is selected to construct the PLR model from a possible list of IOP threshold values. We note that a threshold value that has in the previous iterations is not selected again for model construction. Step 3: Division of the Visual-field Time Series into Pieces. The state of an eye (high- or low-IOP state) is decided using the given IOP threshold c. At each time point when visual-field data are recorded, if the standardized IOP score is larger than the given c, the IOP state at that time point is determined to be in a high-IOP state (see Sec. 3.4). The time series is then divided into pieces when the state changes. This procedure reflects actual clinical knowledge based on the internal heterogeneity of glaucoma progression.

Step 4: Construction of PLR Models using Our Collective Method. A PLR model was constructed using our proposed collective method with the divided pieces. First, a set of similar eyes E was selected to estimate the parameters of the PLR models. In this case, the training dataset consisted of |E| eyes that showed similar behavior to the target eye. In this study, we employed an existing clustering method (Liang, Z. et al., 2013) to collect data from similar eyes. However, another collective method could be used for the same process. Next, we trained the PLR model only using the eye set E. For each piece, we fit a linear function of time; i.e., we estimate the gradient and intercept parameters. We applied the same gradient parameters to pieces in the same IOP state. On the other hand, the intercept parameters were individually determined for each piece. This proper estimation of the gradient and intercept parameters is the key for effectively constructing a PLR model (see Sec. 3.5).

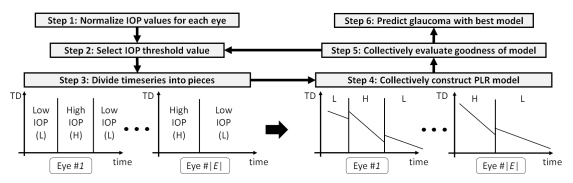


Figure 5: Flow of our algorithm. First, we calculate the standard scores of IOP in Step 1. Then, we iteratively obtain one PLR model by changing IOP threshold values c in Steps 2-5. Finally, we determine the best PLR model based on an information criterion and predict the glaucoma progression in Step 6. We note that the gradients corresponding to the same IOP state are the same for all the eyes belonging to the eye set E composed of eyes similar to the target eye.

Step 5: Evaluation of the Generated PLR Models. Different PLR models were generated when a different threshold c was chosen. Therefore, the best c is required to obtain the best PLR model. We evaluated the quality of the generated PLR models on the basis of statistical model selection criteria. If all possible thresholds were already selected, the best PLR model was chosen as the predictor of the target eye, which was then applied to Step 6. Otherwise, Steps 2-5 were repeated to construct a new PLR model.

Step 6: Prediction of progression with the best PLR model. The visual-field loss of the target eye was predicted with the best PLR model incorporating both the external and internal heterogeneity of the glaucoma dataset.

3.2 Shared and Unshared Parts of Our Collective Method

Our collective method is different from those that simply use data from other eyes for prediction of a target eye. The important difference is that we carefully estimated the model parameters shared among other eyes.

In the Model Selection Phase (Fig. 3 (a)): The fitness and simplicity of the model are evaluated from the overall model. The threshold value for the standardized IOP is common to all similar eyes. This value is decided by considering the whole model and each of the predictors for each eye.

In the Estimation Phase (Fig. 3 (b)): The gradient parameter is shared among each piece in the same IOP state; therefore, this parameter is estimated using the data from all similar eyes. Meanwhile, the intercept parameters differ among each piece; therefore, this parameter is estimated for each eye individually. This modeling procedure allows for the internal heterogeneity of progression to be expressed precisely.

3.3 Problem Settings

Let *N* denote the number of observed eyes, $t_{i,j}$ the time of the *j*th measurement of the *i*th eye, and n_i the number of measurements for the *i*th eye. We represent the vector of the TD values measured at $t_{i,j}$ as $\mathbf{y}_{i,j} = (\mathbf{y}_{i,j}^{(1)}, \dots, \mathbf{y}_{i,j}^{(K)}) \in \mathbb{R}^K$, where *K* is the number of meshes on a visual field, and $p_{i,j} \in \mathbb{R}$ is the IOP value at $t_{i,j}$. We set $T_i := (t_{i,1}, \dots, t_{i,n_i}), Y_i := (\mathbf{y}_{i,1}, \dots, \mathbf{y}_{i,n_i})$, and $P_i := (p_{i,1}, \dots, p_{i,n_i})$. The whole measured dataset is represented as $D := \{(T_1, Y_1, P_1), \dots, (T_N, Y_N, P_N)\}$. We predict the TD value at an arbitrary time point after the last measurement, given a dataset of *N* eyes that includes measurement time, TD and IOP values.

3.4 Judging the IOP State

As mentioned in Sec. 2.2, the higher the IOP, the more rapidly glaucoma progresses; therefore, the progression rate changes at certain time points depending on fluctuations in IOP. To model this internal heterogeneity, we propose an IOP-based PLR model. The two IOP states are defined as the high-IOP state (denoted as H) and the low-IOP state (denoted as L).

However, IOP also shows external heterogeneity. To overcome this problem, we focused on the temporal differences within the IOP time series for each eye. For the *i*th eye, we used a standardized score $\tilde{p}_{i,j} = (p_{i,j} - \bar{p}_i)/\sigma_i$ at $t_{i,j}$ calculated with the mean \bar{p}_i and standard deviation σ_i of IOP. Through this normalization of the external heterogeneity of IOP, the IOP data from other eyes can be treated in the same manner. Let $s_{i,j}$ denote the IOP state of the *i*th eye at time $t_{i,j}$, then $s_{i,j}$ is defined as

$$s_{i,j} = \begin{cases} H, & \text{(if } \tilde{p}_{i,j} \ge c), \\ L, & \text{(if } \tilde{p}_{i,j} < c), \end{cases}$$

where c is the threshold constant. Figure 6(a) displays

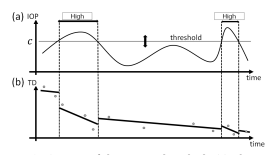


Figure 6: Concept of the proposed method. (a) The timeseries data were divided into intervals with high- or low-IOP states using the threshold c. (b) The PLR model was trained based on these intervals. The gradients of the lines within the same IOP states were the same, and these regression lines were allowed to be disconnected from the subsequent lines. The dots on the graph represent the data points.

the classification protocol. Here, we represent $I_{i,k}$ ($k = 1, ..., L_i$) as the set of indices in the interval where the IOP state remains the same among the series of IOP states $s_i = \{s_{i,j}\}_{j=1}^{n_i}$, calculated as shown above. We denote L_i as the number of intervals for the *i*th eye. We further assume that this threshold constant is common between the target eye and similar eyes.

It is worth noting that the training data should not be partitioned from the target eye data for the following two reasons: first, it is difficult to estimate the correct IOP state, because of the very limited data from the target eye with a small number of diagnoses; second, if the data from the target eye are partitioned, the number of visual-field data points for the target eye will be too small to construct the PLR model, because segmenting the data would further decrease the amount of training data. Thus, we did not partition the time series of the target eye, and assumed that it was in a low-IOP state. This assumption is valid and realistic, because the interval for the high-IOP state is usually very short, and therefore the progression after the final measurement should be in the low-IOP state.

3.5 IOP-based Collective PLR Model

Once the visual-field data for each eye are divided using the standardized IOP scores, a segmented timeseries dataset is obtained for each eye, or a tree structure, as shown in Fig. 3. For each high- and low-IOP state, the proposed model represents a different rate of progression, or the internal heterogeneity of progression. Fig. 6 (b) shows our PLR model as applied to Liang et al.'s method (Liang, Z. et al., 2013).

Liang et al. (Liang, Z. et al., 2013) proposed a spatio-temporal clustering-based linear-regression model using data from other eyes. They assumed the same glaucomatous progression within the same cluster. Here, we show an application of our

developed PLR model to the temporal-shift linear regression method (TSLR) using k-NN as a clustering method (Liang, Z. et al., 2013). They extracted the feature vector of *i*th eye via singular value decomposition of the matrix Y_i to collect data from the eyes similar to the target eye. They recognized the first left-singular vector of Y_i as the spatial feature vector of *i*th eye, and the *k*-nearest eyes in the spatial feature vector space are collected as the similar-eyes cluster. Let $E \subset \{1, \ldots, N\}$ denote the set of indices within a cluster, and let $w_{i,k}$ and $b_{i,k}$ denote the gradient and intercept of the kth piece of the regression line for the *i*th eye, respectively. Then, the assumption above can be formulated as $\forall i, j \in E, \forall k, h \text{ s.t. } s_{i,k} = s_{j,h} =$ S, $\exists w_S, w_{i,k}^{(l)} = w_{j,h}^{(l)} = w_S^{(l)}$, where $S \in \{H, L\}$. As stated above, the intercept parameters are not shared among pieces in the same IOP state within all the eyes in E, while the gradient parameter is shared. This gap enables specificity for each piece and can reflect the internal heterogeneity. Then, the optimal param-eters $\hat{\Phi}_{S}^{(l)} = (w_{S}^{(l)}, \sigma_{S}^{(l)}, \{b_{i,S,k}^{(l)}\})$ are calculated as $\hat{\Phi}_{S}^{(l)} =$ $\arg\min_{\Phi} \sum_{i \in E} \sum_{k=1}^{L_{i}} \sum_{j \in I_{i,k}} \left\{ y_{i,j}^{(l)} - \left(w_{S}^{(l)} t_{i,j} + b_{i,S,k}^{(l)} \right) \right\}^{2}$, where $\sigma_{S}^{(l)}$ is the standard deviation of the above errors. Hereafter, we omit the mesh index l for simplicity since each mesh is processed independently.

3.6 Evaluating the Generated Models

As shown above, several tree-structured PLR models could be obtained with respect to each threshold value *c*. The easiest way to evaluate the models is to select the one with the smallest residual sum of squares (*ERROR*). However, this can result in overfitting, because the ERROR becomes smaller as the tree structure of the PLR models deepens and more detailed data are trained. Therefore, we chose the best model based on information criteria, considering a trade-off between simplicity and fitness of a model. Toward this end, the *Akaike Information Criterion* (AIC) (Akaike, H., 1973), *Bayesian Information Criterion* (BIC) (Schwarz, G., 1978), and *Minimum Description Length Criterion* (MDL) (Rissanen, J., 1986) are well-known information criteria.

The log-likelihood function for the TSLR is calculated for each IOP state *S* as $\log L(\phi_S \mid D) = -\frac{1}{2\sigma_S^2} \sum_{i \in E} \sum_{k=1}^{L_i} \sum_{j \in I_{i,k}} \{y_{i,j} - (w_S t_{i,j} + b_{i,S,k})\}^2 - n_S \log \sigma_S - \frac{n_S}{2} \log(2\pi)$, where $\phi_S := (w_S, \sigma_S, \{b_{i,S,k}\})$ and n_S denote the parameters and the number of data points for the IOP state *S*, respectively. Therefore, AIC, BIC, and MDL are calculated as follows:

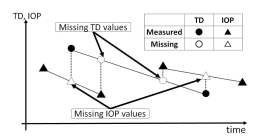


Figure 7: Interpolation of missing TD and IOP values.

$$\begin{aligned} \text{AIC} &= -2\log L(\phi_S \mid D) + 2\left(\sum_{i \in T} L_i + 2\right), \\ \text{BIC} &= -2\log L(\phi_S \mid D) + \left(\sum_{i \in T} L_i + 2\right)\log\sum_{i \in T} n_i, \\ \text{MDL} &= -\log L(\phi_S \mid D) + \left(\sum_{i \in T} L_i + 2\right)\log\sum_{i \in T} n_i \\ &+ \frac{C}{2V}\left[\arcsin x + x\sqrt{1 - x^2}\right]_{b_W V/U}^{a_W V/U}, \end{aligned}$$

where $U = \sqrt{\left\{p - \sum_{i=1}^{P_S} (q_i^2/n_i)\right\}/2}$, *V* is the variance of all time stamps in IOP state *S*, and $C = 2\sqrt{\prod_{i=1}^{P_S} n_i} \left(1/(b_{\sigma}^{P_S+1}) - 1/(a_{\sigma}^{P_S+1})\right) \prod_{i=1}^{P_S} (a_i - b_i)/(P_S+1)$, where (a_w, b_w) , (a_i, b_i) , and (a_{σ}, b_{σ}) represent the upper and lower bounds of the gradient w_S , intercept b_i , and standard deviation σ_S , respectively. We designate *p* as the squared sum of time stamps, q_i as the sum of time stamps in the *i*th interval, P_S as the number of pieces, and n_i as the quantity of data in *i*th interval. The best model and the best threshold are determined by minimizing the sum of the criteria for the high- and low-IOP states.

4 EXPERIMENTS

4.1 Data and Parameters

The dataset used in this paper was provided by the Department of Ophthalmology, The University of Tokyo. This dataset includes visual-field data and IOP data obtained from N = 939 glaucomatous eyes. The visual-field data (TD values) were measured with the Humphrey Field Analyzer (Carl Zeiss Meditec AG, Dublin, CA, USA), using the SITA-standard 30-2 method (K = 74), controlling for the effects of increasing age on the degree of visual-field loss. Therefore, the TD value should decrease if glaucoma has progressed. The TD values were within the range

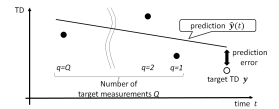


Figure 8: Predicting the last measurement of a target eye. We use Q data points from the target eye for training in addition to data from other eyes.

[-37.0, 4.0] (median: -4, mean: -8.662, standard deviation: 10.56). The mean number of measurements of TD and IOP for each patient was around 11 and 21, respectively. The standardized-score threshold was set within [0.1, 3.0] (step: 0.1).

4.2 Data Interpolation

Our method requires both visual-field data and IOP data for each data point. However, most data points only contain one or the other measure. To facilitate the use of the dataset, we employed linear interpolation to fill in the missing data using the two neighboring measurements for the missing measurement, as shown in Fig. 7. We used these preprocessed data in all of the experiments, even when IOP values were not required. The gap between the time stamp from the last measurement in the training dataset and the target dataset was within [28.9,889] (median: 227, mean: 258, standard deviation: 117) days.

4.3 Evaluation of Prediction Accuracy

We predict the TD value at the last measurement for the target eye using the previous Q data points with data from another N-1 eyes, as depicted in Fig. 8. We set Q = 1,...,6. The prediction accuracy was evaluated with the *Root Mean Square Error* (RMSE):

RMSE_i =
$$\sqrt{\sum_{d=1}^{K} (y_i^{(d)} - \hat{y}_i^{(d)})^2 / K}$$
,

where $y_i^{(d)}$ and $\hat{y}_i^{(d)}$ denote the measured and predicted TD value for the *d*th mesh of the *i*th eye, respectively. Smaller RMSE means better prediction. We also evaluated the accuracy of prediction gained in applying our meta-algorithm to existing methods based on the *Improvement Rate* (IR):

$$IR = \frac{100}{N} \sum_{i=1}^{N} \left[1 - \frac{(RMSE_i \text{ of applied})}{(RMSE_i \text{ of original})} \right]$$

Greater IR indicates larger enhancement of prediction accuracy among N eyes by applying our metaalgorithm. In the following sections, we evaluate the efficacy of our proposed method applied to Liang et al.'s method(Liang, Z. et al., 2013) in terms of RMSE and IR. We define the optimal selection procedure as *BEST* that achieves the smallest RMSE by choosing the optimal model for each eye; i.e., BEST selects the best model from all the generated models on the basis of the TD value to be predicted. We employed leaveone-out cross-validation in the following analyses.

4.4 Experiment 1: The Best Use of IOP

Experimental Settings: There are several options to employ IOP information in our method, including the use of raw values, raw deviations, and standardized scores. Here, we show the superiority of using the standardized scores to the other two options in terms of improvement in prediction accuracy for the BEST procedure. The thresholds for the raw value and raw deviation were set within [10,30] (step: 1) and [0.1,3] (step: 0.1), respectively. Both units are mmHg.

Results: Table 1 displays the median, mean, and 10%-trimmed mean of the RMSEs using our method with the three analytical approaches of IOP. As for the mean and 10%-trimmed mean, the standardized score significantly improved prediction accuracy for all Q according to a one-sided Student's *t*-test. There was no significant difference in terms of the median. Table 2 shows the proportion of cases in which our method using the standardized IOP scores performed better in terms of prediction compared to that using the other two scores. The use of our standardized scores significantly outperformed the use of the other two according to a one-sided binomial test.

4.5 Experiment 2: Best Information Criterion

Experimental Settings: We compared the ERROR, AIC, BIC, and MDL to determine which criterion is most suitable for prediction with our method. We also analyzed the BEST case as reference.

Results: Table 3 presents the median and mean RMSEs of the predictions obtained with the model for the ERROR, AIC, BIC, MDL, and BEST. The MDL performed significantly better than the others in terms of the mean RMSEs at the 1% significance level with a one-sided Student's *t*-test in most cases, while it performed better at the 5% significance level when Q = 3. As for the case of the median RMSEs, the MDL performed well in most cases but there was no significant difference. Table 4 shows the proportion of cases in which our method using the four criteria most accurately predicted the value. A one-sided binomial

test verified that the MDL significantly outperformed the other criteria at the 0.1% level in all the cases.

4.6 Experiment 3: Effectiveness of IOP

Experimental Settings: We compared our method as applied to the original Liang et al.'s method (Liang, Z. et al., 2013) with the original to investigate the improvement in predictive ability. Based on the results of Experiment 2, we used the MDL. In addition, we compared our method with the BEST to evaluate the predictive potential obtained by introducing the IOP.

Results: Table 5 shows the median, mean, and 10%-trimmed mean of the RMSEs for predictions obtained with our proposed method compared to the original method. As for the mean, the Student's *t*-test verified that our model significantly outperformed the original at the 0.1% level when the null hypothesis IR = 0 was set for both trimmed and non-trimmed cases. For the median, a Mann-Whitney-Wilcoxon test verified that our model exceeded the predictive power of the original at the 5% level except when we set Q = 1. Table 6 shows the proportion of cases in which our method predicted the value most accurately. A binomial test determined that our method was significantly better than the original. The results shown in Tables 5 and 6 revealed that our method applied with the BEST procedure is highly significantly superior to the original for prediction accuracy.

4.7 Discussion

Experiment 1: The use of a standardized IOP score was more effective compared to the use of raw IOP values or deviations as an indicator of the IOP state. We believe that this is because the usual state of the IOP differs from eye to eye; thus a standardized score can normalize IOP values across eyes, whereas the raw value and deviation cannot. With the three indicators of IOP, we segmented the IOP states into high and low states. The fact that the standardized score was the best indicator of the IOP states indicates a method for controlling for the heterogeneous differences among individuals; i.e., solving the external heterogeneity of IOP. Thus, the scores could correctly represent the fundamental states of most eyes.

Experiment 2: The results suggested that MDL was a better choice for improving predictions compared to others. This result could be due to the following factors. The framework of MDL, which can take the structure of a model into account by calculating the optimal code length, fits well with the property of our PLR model owing to its clear tree structure (see Fig. 3). Indeed, several studies have shown that the

Table 1: Median, mean, and 10%-trimmed mean of RMSEs using the raw values, raw deviations, and standardized scores of the IOP for each eye. The BEST optimal selection procedure was used for prediction. The symbol \sharp indicates statistical significance at the 0.1 % level.

			Q	1	2	3	4	5	6			
					(a) Me	dian RMS	SEs					
		_	Std. score Raw val. Raw dev.	4.026 4.112 4.139	3.834 3.879 3.861	3.726 3.828 3.770	3.673 3.759 3.739	3.652 3.756 3.724	3.621 3.716 3.666			
Q	1	2	3	4	5	6	1	2	3	4	5	6
		(b) N	Iean RMS	Es				(c) 10	%-trimme	ed Mean R	RMSEs	
Std. score	4.314	4.081	3.991	3.969	3.933	3.875	4.161	3.918	3.809	3.775	3.750	3.708
Raw val.	4.376	4.144	4.054	4.029	3.998	3.939	4.226	3.984	3.872	3.838	3.817	3.771
IR	1.463#	1.563#	1.576#	1.606#	1.707‡	1.834‡	1.262#	1.402#	1.376‡	1.391‡	1.472	1.834‡
Raw dev.	4.456	4.184	4.091	4.100	4.023	3.946	4.241	3.977	3.879	3.851	3.811	3.756
IR	1.550 #	1.252‡	1.300 #	1.501‡	1.139#	1.214#	0.880#	0.803‡	0.862#	0.830#	0.777 \sharp	0.844

Table 2: Proportion (%) of cases for which our proposed standardized score of IOP yielded better prediction performance. Cases for the BEST procedure are shown. The symbol \sharp indicates statistical significance at the 0.1 % level.

Q	1	2	3	4	5	6
v.s. Raw val. v.s. Raw dev.	11	11	T I	11	41	11

Table 3: Median and mean of RMSEs for predictions with the models selected according to the ERROR/AIC/BIC/MDL, and the possible best model (BEST). The symbols * and \dagger indicate statistical significance at 5% and 1%, respectively.

Q	1	2	3	4	5	6	1	2	3	4	5	6
5016		(a) Me	edian RM	SEs				GY	(b) Mear	n RMSEs	:ATI	ON
ERROR	4.461	4.179	4.097	4.083	4.104	4.085	4.738	4.509	4.425	4.425	4.391	4.322
AIC	4.461	4.179	4.096	4.088	4.104	4.087	4.737	4.508	4.425	4.425	4.391	4.323
BIC	4.456	4.173	4.101	4.083	4.099	4.092	4.731	4.505	4.424	4.423	4.387	4.320
MDL	4.422	4.163	4.113	4.087	4.064	4.076	4.682†	4.471 †	4.394*	4.388 †	4.351†	4.291 †
BEST	4.026	3.834	3.726	3.673	3.652	3.621	4.314	4.081	3.991	3.969	3.934	3.875

Table 4: Proportion (%) of cases for which each criterion gave the best performance for prediction. The symbol \sharp indicates statistical significance at the 0.1 % level.

Q	1	2	3	4	5	6
ERROR	17.26	16.77	17.76	15.80	17.55	15.97
AIC	14.98	13.53	14.75	15.80	13.40	15.26
BIC	19.33	19.81	21.74	19.19	19.63	20.96
MDL	48.43 ‡	49.89 ‡	45.75 ‡	49.20 ‡	49.43 ‡	47.81 ‡

MDL works well for tree-structured models (Mehta, M. et al., 1995; Robnik-Šikonja, M. and Kononenko, I., 1998). In addition, extremely ineffective models were not selected when using the MDL, as implied by the fact that there were significant differences in the mean, but almost no differences in the median.

Meanwhile, because the MDL is difficult to calculate analytically, it is difficult to apply the MDL to our proposed method for complex models such as Murata et al.'s model (Murata, H. et al., 2014). However, sufficiently accurate results were gained with our method just using ERROR This might be because we assumed that the gradients of each piece were the same in the same IOP state, which worked as a kind of regularizer of the parameters. This result suggests that using ER-ROR is a feasible solution for more complex models.

			Q	1	2	3	4		5	6				
					(a) Me	edian RM	ISEs							
			Original Proposed (MDL)				4.386 4.24 4.163* 4.113					.205 . 076 *		
		Propos	ed (BEST)	4.026‡	3.834‡	3.726	5# 3.67	73‡ 3.6	52‡ 3	.621‡				
Q	1	2	3	4	5	6	1	2	3	4	5	6		
		(b)]	Mean RM	SEs				(c) 10	%-trimm	ed Mean F	RMSEs			
Original	369.8	566.6	300.0	262.1	391.4	235.4	4.681	4.487	4.394	4.357	4.317	4.266		
Proposed (MDL)	4.682	4.471	4.394	4.388	4.351	4.291	4.525	4.299	4.208	4.181	4.155	4.116		
IR	4.541‡	5.177‡	4.985‡	4.853#	4.205\$	4.182\$	2.670#	2.944‡	2.932‡	3.108‡	2.811#	2.833#		
Proposed (BEST)	4.314	4.081	3.991	3.969	3.934	3.875	4.161	3.918	3.809	3.775	3.750	3.708		
IR	12.78‡	14.19‡	14.54‡	14.83\$	14.32#	14.50\$	10.92\$	11.95‡	12.49\$	12.76‡	12.54#	12.89#		

Table 5: Median, mean, and 10%-trimmed mean of RMSEs. Liang et al.'s method (Liang, Z. et al., 2013) is referred to as the "original", and our method as applied to the original method is referred to as the "proposed". The symbols * and \sharp indicate statistical significance at the 5% and 0.1 % level, respectively.

Table 6: Proportion (%) of cases in which our method applied to Liang et al.'s method is superior to the original method based on MDL and BEST. The \sharp indicates that the values are statistically significant at the 0.1 % level.

Q	1	2	3	4	5	6
Proposed (MDL)	71.41#	71.38‡	71.57‡	71.74 <u></u> ‡	72.74 ‡	67.63
Proposed (BEST)	99.24#	99.35‡	99.78	99.45#	99.23	99.11

Experiment 3: Table 5 demonstrates that the application of our proposed method to the original Liang et al.'s method showed much better performance than the original method. Moreover, our method achieved better prediction accuracy with small *Q*. It is well-known that high IOP exacerbates the progression of glaucoma (see Sec. 2.2). Therefore, data incorporating eyes at different IOP states can be anomalous for long-term prediction of the original method. In our method, such noise is excluded by segmenting the data with IOP values, which cannot be realized in the original method. Therefore, we suppose that this separation of the data enables our method to produce accurate predictions with less data owing to its stronger power of expression and purity of the training data.

Table 6 also indicates that our method with MDL represents a significant improvement over the original, and that it can help to improve the outcome for a large number of eyes. However, about 30 % of patients would not benefit from our method judging from the results. This demonstrates that the best model is not always selected with MDL, and that there is still room for improvement in considering information criteria. Nonetheless, Table 6 shows that insofar as our framework exploits the IOP and copes with external and internal heterogeneity, it offers more accurate predictions compared to existing methods.

Overall Discussion: We have shown the efficacy of our collective PLR model in predicting the progression of glaucoma. Since our method can be applied to other existing methods, it is expected to serve as an improvement of current methods by exploiting supplemental data. However, there is still room for improvement in the model-selection phase, because the accuracy of the prediction with the BEST procedure was much better than that when using other information criteria. One possible explanation for this result is the small number of data entries for the high-IOP state, making it difficult to comprehensively evaluate the model for this state. Thus, our future work will focus on such situations to improve the proposed model.

5 CONCLUSION

We have proposed a novel collective PLR method that copes with external and internal heterogeneity as well as the medical-data-structure-difficulty of medical datasets. Existing methods cannot cope with the internal heterogeneity, i.e., a situation where the rate of progression for individual eyes changes over time. We have dealt with this internal heterogeneity using a PLR model based on clinical knowledge regarding the relationship between IOP and the rate of glaucomatous progression (AGIS Investigators, 2010; Collaborative Normal-Tension Glaucoma Study Group, 1998; Satilmis, M. et al., 2003). Our method can also deal with the external heterogeneity and medicaldata-structure-difficulty by incorporating a collective method. Therefore, our method is a novel extension of previous collective methods from both theoretical and practical aspects, which increases prediction accuracy. Similarly, other methods (Maya, S. et al., 2014; Murata, H. et al., 2014) are expected to be improved by incorporation of our method.

Medical datasets are commonly plagued by high levels of heterogeneity, and we have here proposed a new method that shows good performance in overcoming this heterogeneity in a glaucoma dataset for effective predictions of disease progression. We believe that our method can be extended to tackle similar difficulties in other medical datasets and we have provided standardized directions for such analyses.

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