CLASSIFICATION OF BIPOLAR DISORDER, MAJOR DEPRESSIVE DISORDER, AND HEALTHY STATE USING VOICE

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ABSTRACT

Objective: In this study, we propose a voice index to identify healthy individuals, patients with bipolar disorder, and patients with major depressive disorder using polytomous logistic regression analysis.

Methods: Voice features were extracted from voices of healthy individuals and patients with mental disease. Polytomous logistic regression analysis was performed for some voice features.

Results: With the prediction model obtained using the analysis, we identified subject groups and were able to classify subjects into three groups with 90.79% accuracy.

Conclusion: These results show that the proposed index may be used as a new evaluation index to identify depression.

Keywords: Voice, Bipolar disorder, Major depressive disorder, Polytomous logistic regression analysis.

INTRODUCTION

Due to the stress in modern society, mental health care has become an important issue. In recent years, mental disorders resulting from stress were shown to cause major social loss by declining labor productivity[1]. Therefore, it is important to create a system where emotional problems can be discovered in the early stages and to develop a technology that can easily assess depression and stress.

The current screening methods for mental disorders use biomarkers such as saliva[2], blood[3], electrocardiogram[4], and electroencephalogram[5], but these are invasive and high cost because special equipment and medicines are needed. Some non-invasive methods include self-administered psychological tests such as the General Health Questionnaire[6] and the Beck Depression Inventory[7]. Self-administered psychological tests are relatively easy, but reporting bias cannot be eliminated. Reporting bias is defined as selective under- or overestimation of certain information influenced by the responder’s consciousness/unconsciousness[8]. Bipolar disorder and major depressive disorder are two types of depression and divided based on their respective symptoms. Bipolar disorder is a mental disease with alternating manic and depressive states, with a difficult differential diagnosis from unipolar depressive disorder during a depressive episode[9]. It is particularly difficult to diagnose using a single self-administered psychological test, and the differences between these two diseases can be difficult to identify.

On the contrary, it is empirically known that changes in mood are expressed in a person’s voice, and in a previous study, the authors developed a method to estimate mental health states, such as depression and stress, using a person’s voice[10-12]. Voice analysis is non-invasive, does not require a specialized device, and can be performed easily and remotely. Furthermore, it may solve the reporting bias associated with self-administered psychological tests and other various problems encountered while detecting mental diseases. Thus, the stress evaluation method using voice analysis has recently been garnering attention.

The authors had conducted a study on voice index that could detect bipolar disorder and major depressive disorder[13]. In this previous study, we identified healthy individuals and patients with mental disease based on a single classic voice index and showed that, with another single classic voice index, bipolar disorder and major depressive disorder could be identified. However, the detection precision was not sufficient.

Therefore, in this study, we targeted groups of healthy individuals, patients with bipolar disorder, and patients with the major depressive disorder and proposed a new voice evaluation index to identify these three groups with improved precision using polytomous logistic regression analysis[14,15].

METHODS

Subjects

Our subjects were outpatients at the National Defense Medical College Hospital who were undergoing treatment for bipolar disorder and major depressive disorder. They were diagnosed by psychiatrists using the Mini-International Neuropsychiatric Interview[16]. Subjects who had no issues with their daily living were enrolled as healthy volunteers. There were eight patients with bipolar disorder and 14 patients with major depressive disorder. There were nine and 23 healthy people at the National Defense Medical College Hospital and Tokyo Medical...
University Hospital, respectively, with a total of 32 healthy individuals. Depending on their situations, patients received multiple examinations, and their voices were recorded at each examination. In total, there were 14 data of voices with bipolar disorder, 30 data of voices with major depressive disorder, and 32 healthy individuals.

Table 1 shows the details of the data, where values outside the brackets represent the number of patients, and values inside the brackets represent the number of voice data.

The mean age of the healthy group was 50.48±13.45 years (age could not be confirmed for three patients). The mean age of the patients with bipolar disorder was 46.50±13.06 years. The mean age of the patients with major depressive disorder was 43.71±11.57 years.

As other tests, all patients recorded scores of Hamilton Depression Rating Scale [17] and Young Mania Rating Scale [18] at each examination.

**Voice recording**

Voice recording was performed in examination rooms at each hospital, and patients were asked to read a fixed sentence consisting of 17 phrases. For healthy individuals, voice recording was conducted for the nine individuals at the National Defense Medical College Hospital and for the 23 individuals at the Tokyo Medical University Hospital (six individuals overlapped) in the same environments as that of the patient voice recordings. Voices were recorded with a ME52W microphone (OLYMPUS, Tokyo, Japan) attached on the chest, about 100 mm below the subject’s mouth. The recording device used was the Portable Recorder R-26 (Roland, Shizuoka, Japan). The sampling rate of recording was 96 kHz, and the data resolution was 24 bit.

**Voice analysis**

After this, healthy individuals are referred to as “HE,” patients with bipolar disorder as “BP,” and patients with major depressive disorder as “MD.”

To extract features from the voices, we used the free software openSMILE (v. 2.3) [19].

The openSMILE uses a script that automatically extracts a set of various features from the voice, and in this study, we extracted feature sets used for emotion recognition (The large openSMILE emotion feature set) from each voice. In this manner, we extracted 6.552 voice features from each voice. From these features, we extracted features that suited the classifier of healthy and patient groups. The procedure was as follows:

1. Eliminating the difference in recording environment at the National Defense Medical College Hospital and the Tokyo Medical University Hospital. We separated healthy individuals into two groups: Those recorded at the National Defense Medical College Hospital (HE) and those recorded at the Tokyo Medical University Hospital (HE). and for each extracted feature, we calculated the effect size between the two groups (HE and HE). Effect size is an index that evaluates the difference in mean values between two groups, and this difference is defined based on the equations given below, as a value standardized with standard deviation:

\[
ES = \frac{\mu_A - \mu_B}{\sigma} = \sqrt{\frac{(n_A - 1)\sigma_A^2 + (n_B - 1)\sigma_B^2}{n_A + n_B - 2}}
\]

(1)

In these equations, \(\mu_A\) and \(\mu_B\) are means, \(\sigma_A\) and \(\sigma_B\) are standard deviations, and \(n_A\) and \(n_B\) are the number of samples for Groups A and B, respectively. If the effect size is small, the difference is lower between the two groups. In this study, we selected features for which the effect size was lower than 0.5.

Furthermore, to eliminate data dependence on effect size, we tested the difference in mean values for each feature between two of the groups. For this, we used a t-test if feature distribution of HE, and HE satisfied normality and the Mann–Whitney U-test if it did not. We also selected features in which the p-value obtained in the test was larger than 0.05.

Using these processes, we selected 1.553 features from 6.552.

2. We selected features that were effective for identifying HE, BP, and MD. For each one of the 1.553 features, we calculated the effect size between two of the groups - HE-BP, HE-MD, and BP-MD - using the equations from the first step of this procedure. In addition, we performed a multiple comparison test to compare the three groups (HE, BP, and MD). We used the Steel-Dwass test, which does not restrict the distribution shape of the three groups [20].

In this study, we selected features in which the effect size between all combinations of two groups among the three groups (HE, BP, and MD) was >0.5, and p values of paired comparison tests were all smaller than 0.1. Results led to a selection of nine features from 1.553 features.

**Polytomous logistic regression analysis**

A polytomous logistic regression analysis is a multivariate analysis that classifies data into three or more groups based on predicted values. It is an expansion of a normal logistic regression analysis that classifies data into two groups. Model equations for three groups (A, B, and C) are shown below:

\[
\log \left( \frac{P_A}{P_B} \right) = \alpha_A + \beta_1 x_1 + \beta_2 x_2 + \ldots + \beta_n x_n
\]

(2)

\[
\log \left( \frac{P_A}{P_C} \right) = \alpha_A + \beta_1 x_1 + \beta_2 x_2 + \ldots + \beta_n x_n
\]

(3)

In these equations, \(P_A\), \(P_B\), and \(P_C\) indicate occurrence probabilities for each group. \((x_1, x_2, \ldots, x_n)\) indicates one data set, and \(\alpha_A, \beta_1, \ldots, \beta_n\) indicate model coefficients. To perform polytomous logistic regression analysis, a standard group must be set first; (2) and (3) are model equations that use the Group C as the standard group. Groups other than the standard group are target groups. Logit of the model equation expresses the likelihood of a target group relative to that of the standard group; if the logit is smaller, the standard group is more likely, and if the logit is larger, the target group is more likely. If the coefficient of the model equation is not statistically 0, the variable for such coefficient has an impact on the logit. The occurrence probability for each group based on (2) and (3) can be calculated using the following equations:

\[
P_A = \frac{\exp \left( \log \left( \frac{P_A}{P_B} \right) \right) \cdot \exp \left( \log \left( \frac{P_A}{P_C} \right) \right)}{1 + \exp \left( \log \left( \frac{P_A}{P_B} \right) \right) + \exp \left( \log \left( \frac{P_A}{P_C} \right) \right)}
\]

(4)

\[
P_B = \frac{\exp \left( \log \left( \frac{P_B}{P_C} \right) \right)}{1 + \exp \left( \log \left( \frac{P_A}{P_B} \right) \right) + \exp \left( \log \left( \frac{P_B}{P_C} \right) \right)}
\]

(5)

\[P_C = 1 - P_A - P_B\]

(6)

For each data set, \(P_A\), \(P_B\), and \(P_C\) are estimated from model equations, and the data are classified as the most probable group.

In this study, we used categorical information of each group (HE, BP, and MD) as dependent variables and the nine selected voice features as
RESULTS

We recorded the effectiveness of the nine voice features that were selected through voice analysis as follows:

1. **mfcc_sma1_quartile2**: The median of the three-point moving average in mel-frequency sub-band (second).
2. **mfcc_sma2_linefregc**: The intercept of a regression line of the three-point moving average in mel-frequency sub-band (second).
3. **mfcc_sma2_gregc3**: The secondary regression curve coefficient (constant term) of the three-point moving average in mel-frequency sub-band (third).
4. **mfcc_sma2_quartile1**: The first quartile of the three-point moving average in mel-frequency sub-band (third).
5. **mfcc_sma2_amean**: The arithmetic mean of the three-point moving average in mel-frequency sub-band (third).
6. **mfcc_sma2_linregc2**: The intercept of a regression line of the three-point moving average in mel-frequency sub-band (fourth).
7. **mfcc_sma10_quartile1**: The first quartile of the three-point moving average in mel-frequency sub-band (fourth).
8. **F0env_sma_qregerrQ**: The square error of the secondary regression curve and the original data of the three-point moving average for the F0 envelope (smoothed by the attenuating exponential function).
9. **pcm_fftMag_spectralCentroid_sma_linregerrA**: The primary error of the regression line and the original data of the three-point moving average for gravity center frequency of spectral power.

Table 2 shows the results of the polytomous logistic regression analysis, which also used stepwise regression where the categorical information of subject groups (HE, BP, and MD) was used as dependent variables, the abovementioned nine voice features were dependent variables, and the HE group was the standard group.

Coefficients in the table represent coefficients to the independent variables from the prediction model. The features ultimately selected by the stepwise regression were the following seven:

- **mfcc_sma1_quartile2**
- **mfcc_sma2_linefregc**
- **mfcc_sma2_amean**
- **mfcc_sma2_linregerrA**
- **mfcc_sma3_linefregc2**
- **mfcc_sma2_qregc3**
- **pcm_fftMag_spectralCentroid_sma_linregerrA**

The features **mfcc_sma2_qregc3** and **mfcc_sma2_quartile1** were excluded because they were determined to be features that did not contribute to the dependent variables. Therefore, the coefficient for these two features is 0, and the prediction equation is expressed as follows:

$$
\begin{align*}
\chi_1 &= \text{mfcc_sma1_quartile2}, \\
\chi_2 &= \text{mfcc_sma2_linefregc}, \\
\chi_3 &= \text{mfcc_sma2_amean}, \\
\chi_4 &= \text{mfcc_sma3_linefregerrA}, \\
\chi_5 &= \text{mfcc_sma10_quartile1}, \\
\chi_6 &= \text{F0env_sma_qregerrQ}, \\
\chi_7 &= \text{pcm_fftMag_spectralCentroid_sma_linregerrA},
\end{align*}
$$

In these equations, \( P_{HE}, P_{BP}, \) and \( P_{MD} \) express the probability of the data being classified as the HE group, BP group, and MD group, respectively.

To determine if the prediction model was significant, we performed a likelihood ratio test with a model using only the intercept. The results had a p value of \( p=1.22E^{-01} \), which confirmed the significance of the prediction model.

To eliminate sex dependence in features useful to the model, we tested the difference between two groups based on the sex of healthy individuals. If the two groups satisfied normalcy, we performed a \( t \)-test, and if they did not, we performed a Mann–Whitney \( U \)-test. Results are shown in Table 3.

We calculated the probabilities of being classified into each of the three groups for each data used in the analysis using the prediction model equations \( (P_{HE}, P_{BP}, \) and \( P_{MD} \)), and by classifying the data in the most probable group, we organized the data. Results are shown in Table 4.
The HE group was identified with 90.63% accuracy. The BP group was identified with 85.71% accuracy. The MD group was identified with 93.33% accuracy. Overall accuracy was 90.79%.

Fig. 1 shows the probability distribution of each subject being identified in each group.

Fig. 1 shows that the probability of subjects in the HE group being classified as the HE group was much higher than the probability of being classified into either of the other two groups. Although the probability of subjects in the BP group being classified as the BP group has a wider range, it was still higher than the probability of being classified into either of the other two groups. The probability of subjects in the MD group being classified as the MD group was much higher than the probability of being classified into either of the other two groups.

**DISCUSSION**

Within the present analytical data, there were data recorded at two different locations for the healthy group. The analysis using only voice data from the National Defense Medical College Hospital led to a bias in the number of male and female subjects in the healthy group, and as the number of samples was small, it was difficult to obtain a model with sufficient precision. Therefore, to eliminate the sex bias and acquire a sufficient number of samples, we added voice data from the Tokyo Medical University Hospital. This addition required that we eliminate the impact of different recording locations on the model, but in the first step of feature selection, such impact was mostly eliminated.

The predicted model equation had a significant coefficient that was 43% of the whole (excluding the intercept), which is a relatively low percentage. However, the overall model had statistical significance, and the data fit was good. In the present analysis, the feature selection condition was somewhat relaxed, which might have been the reason for the coefficient not being significant. In the future, we will apply the prediction model equation obtained from the analysis to another data set and verify the impact of the coefficient on classification precision.

For the seven features useful for the model, the difference between sexes could not be detected for healthy individuals in the present number of samples. In other words, the way in which patients are classified into two groups based on the seven features is not influenced by differences in sexes.

Using the prediction model equation based on the BP group data, one datum was classified as belonging to the major depressive disorder group. This datum was recorded during the second examination of the patient, and this patient’s Hamilton Depression Scale score at that time was higher than it was at the first examination. The same patient’s Young Mania Rating Scale score was lower than it was at the first examination. Thus, as the patient’s symptoms of major depressive disorder became more prominent than those of bipolar disorder, the patient was classified as belonging to the major depressive disorder group. There was another datum in the MD group that was classified as healthy using the prediction model equation. This datum was recorded during the second examination of the patient, and it is possible that, after outpatient treatment, the patient’s depressive symptoms improved, and the patient could be classified as healthy. It is our future challenge to verify how patients are classified when symptoms change with time.

In this study, we did not discuss the impact of the voice features used in the prediction model on the model itself. It is another future challenge for us to examine the features that can most effectively identify the diseases and the characteristics of patients’ voices that each feature captures.

**CONCLUSION**

In this study, we targeted groups of healthy individuals, patients with bipolar disorder, and patients with major depressive disorder and, based on their voices, extracted a large-scale voice feature set using voice feature extraction software. We selected voice features useful as the classifier. Based on the selected features, we performed the polytomous logistic regression analysis and proposed a voice index that identifies healthy individuals, patients with bipolar disorder, and patients with major depressive disorder. Using the prediction model obtained from the analysis, subjects could be classified into the three groups with 90.79% accuracy. The above results indicated that the proposed index could be useful as a new evaluation index to identify depression.

**CONFLICTS OF INTEREST**

All authors have none to declare.

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