Noninvasive prediction of shunt operation outcome in idiopathic normal pressure hydrocephalus

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Idiopathic normal pressure hydrocephalus (iNPH) is a neuropsychiatric disorder characterized by gait disturbance, cognitive deterioration and urinary incontinence in elderly individuals. These symptoms can be improved by shunt operation in some but not all patients. Therefore, discovering predictive factors for the surgical outcome is of great clinical importance. We used normalized power variance (NPV) of electroencephalography (EEG) waves, a sensitive measure of the instability of cortical electrical activity, and found significantly higher NPV in beta frequency band at the right fronto-temporo-occipital electrodes (Fp2, T4 and O2) in shunt responders compared to non-responders. By utilizing these differences, we were able to correctly identify responders and non-responders to shunt operation with a positive predictive value of 80% and a negative predictive value of 88%. Our findings indicate that NPV can be useful in noninvasively predicting the clinical outcome of shunt operation in patients with iNPH.
iNPH patients. In support to this notion, Miyoshi et al. revealed that frontal lobe dysfunction has significant correlation with gait disturbance in iNPH patients. In a previous study, we also suggested a way to distinguish shunt responders from non-responders using a combination of NPV analysis and CSF tapping. However, CSF tapping is an invasive procedure and has risks of infection, post-procedural headache and bleeding. To make things worse, iNPH patients often have suffered myocardial or cerebral infarction in their past and take anti-platelet drugs, which increases the risk of bleeding by CSF tapping. In addition, the CSF tap test, a diagnostic procedure for predicting the shunt operation outcome using CSF tapping and clinical assessments, has a low negative predictive value.

EEG has become a useful tool in neuroscience to investigate cortical functions, and also in clinical practice for objective clinical assessment of neurological and neuropsychiatric disorders, such as epilepsy and Alzheimer's disease. This usefulness mainly arises from the fact that EEG can directly measure brain electrical activity. In specific, EEG time-series signals relate to dynamic postsynaptic potentials of the cerebral cortex, with millisecond temporal resolution.

In the present study, we aimed to distinguish shunt responders from non-responders using only pre-CSF tapping EEG data, that is to say without performing CSF tapping, something that would significantly benefit iNPH patients as an alternative to tapping procedures.

Results

Demographic and clinical results. We classified responders and non-responders according to shunt operation outcome as mentioned in the Materials and Methods section. One responder showed improvement in gait and cognitive domains and the other responders showed improvement only in the gait domain. Demographic and clinical data are shown in Table 1. There were no differences in age, gender, and clinical scores between groups. The rate of responders was relatively low because we adopted stricter classification criteria of shunt response than the Japanese Clinical Guidelines in order to select patients who showed significant improvements in iNPH symptoms as responders.

NPV analysis results. Statistically significant NPV differences between responders and non-responders in each frequency band are listed in Table 2. Responders had higher beta NPV values at the right anterior prefrontal electrode (Fp2), the right temporal electrode (T4) and the right occipital electrode (O2) compared to non-responders. Maps of beta NPV results of responders and non-responders and the t-values between them are shown in Fig. 1 (left, center and right). Using beta NPVs at Fp2, T4 and O2 electrodes simultaneously as discriminating variables, linear discriminant analysis yielded a discriminant score that most accurately distinguishes shunt responders from non-responders. The equation was: Discriminant score = 0.30 × betaFp2 NPV + 0.60 × betaO2 NPV + 0.57 × betaT4 NPV − 0.24. The cut-off value of the score is zero, and positive or negative scores indicate responders or non-responders, respectively. This score could correctly identify responders and non-responders with a positive predictive value of 80% (8/10) and a negative predictive value of 88% (7/8) (Fig. 2).

Power spectral analysis results. There were no significant power differences between responders and non-responders.

Discussion

In the present study, we employed NPV analysis, which sensitively detects the instability of cortical electrical activity, to find objective predictive EEG markers of shunt operation outcome in patients with iNPH. Our main findings were that: (1) responders had significantly higher values of beta NPV at the right fronto-temporo-occipital cortices [the right anterior prefrontal electrode (Fp2), the right temporal electrode (T4) and the right occipital electrode (O2)] compared to non-responders as shown in Figure 1, and (2) using these beta NPVs, we could correctly identify responders and non-responders with a positive predictive value of 80% (8/10) and a negative predictive value of 88% (7/8), as shown in Figure 2.

The regions indicated by our main findings, that is the right fronto-temporo-occipital cortices, may correspond to the area of the right ventral attention network (VAN). Right VAN was originally discovered as a visual recognition network, where visual information that has flowed from the occipital lobe is compared to visual/spatial memory in the right temporal cortex and then identified in right temporal or frontal cortex. In support to this notion, Miyoshi et al., using the walking test in dual task conditions, have reported that after CSF tapping iNPH patients significantly improved allocation of attention to gait during dual task, which is a function of right VAN, compared to iNPH-like patients. Also, in a previous study, we found that alpha and beta NPV changes at the right anterior prefrontal electrode (Fp2) correlated with changes in walking time, which suggests a recruitment of the right VAN in gait control in iNPH patients. This recruitment of VAN during gait has also been seen in patients with Parkinson’s disease, and these authors discussed that an associated burn-out of VAN may lead to the freezing of gait. Recruitment of the right VAN has also been found in healthy subjects during line tracing by hand. These authors suggested that the role of the right VAN during motion is to modulate attention and adapt the movement using spatial information. Taking into account all of this information, our main finding (1) indicates that the area which showed a differ-

| Table 1 | Cognitive and gait function test scores of responders and non-responders |
| --- | --- |
| Test | Responders | Non-responders |
| Age | 76 ± 5.2 | 74 ± 7.0 |
| Gender (F/M) | 4/9 | 3/6 |
| WTT | 24.6 ± 6.8 | 19.7 ± 5.9 |
| TUG | 15.6 ± 5.3 | 12.0 ± 2.9 |
| GSS | 6.3 ± 3.5 | 5.0 ± 1.8 |
| MMSE | 22.7 ± 3.6 | 23.8 ± 3.6 |
| FAB | 11.2 ± 3.3 | 11.5 ± 2.8 |
| TMT-A | 138 ± 95 | 97 ± 48 |
| WMS-R_Attention/Concentration index | 77.4 ± 15.8 | 80.6 ± 13.6 |
| WAIS-III-Digit SymbolCoding | 5.6 ± 2.6 | 6.7 ± 1.9 |
| WAIS-III-Block Design | 6.5 ± 3.6 | 7.1 ± 2.1 |

| Table 2 | NPV differences between responders and non-responders |
| --- | --- |
| NPV at band-electrode | t-value | p-value |
| Beta-Fp2 | 2.1 | 0.047 |
| Beta-O2 | 2.1 | 0.047 |
| Beta-T4 | 2.6 | 0.020 |
ence in cortical electrical activity corresponds to the right VAN, which supports gait function in iNPH patients. This is also consistent with our results of gait and cognitive assessment. Functional recovery induced by shunt operation occurs mainly in the gait domain but not in the cognitive domain25.

Since responders and non-responders exhibit different cortical electrical activities in the beta band, it is worthwhile to comment on the role and spatial distribution of the beta frequency band in cortical electrical activity. Beta oscillation is activated during isometric contraction, suppressed by voluntary movements, and greatest after movement termination, something that is described as post movement beta rebound (PMBR). This characteristic modification of beta is observed in motor related areas (sensorimotor and premotor areas, the basal ganglia and the cerebellum). Beta oscillation in motor related areas is thought to maintain the ongoing motor set and disorders of beta oscillation are closely related to motor impairment diseases26. For example, motor impaired patients with Parkinson’s disease show abnormal enhancement of contralateral beta power in primary sensorimotor areas during isometric contraction, related to the severity of motor impairment27. These patients also have disturbance of the PMBR28. In addition, in a previous study, we found that beta oscillation in right anterior PFC, which may be an anterior part of motor supporting areas (right VAN) was implicated in gait control in iNPH patients8. However, iNPH “responder” patients showed significant difference of beta activity in motor supporting areas (right VAN) and not in motor and premotor areas, compared to non-responders. This may be explained by the fact that almost all non-responders improved their gait status, without accompanying improvements of the other walking tests. Gait status is closely associated with the left dorsal PMA8,24 and the higher-order motor areas including PMA control the cortical electrical activity of the primary motor cortex29. Therefore, we can presume that non-responders also recovered activity in motor and premotor areas to some extent, thus showing no difference compared to responders.

In a previous study, we found a correlation of NPV with brain function: an NPV increase, which describes the destabilization of cortical electrical activity, reflects functional worsening, while an NPV decrease, describing the stabilization of cortical electrical activity, reflects functional improvement8. Thus, our present study result that responders had higher beta NPV values in the right VAN is consistent with our results of gait and cognitive assessment.

Figure 1 | Beta band NPV in Responders and Non-responders and t-value between them. Beta band NPV in responders (left) and non-responders (center) and t-value between them (right). Responders had higher beta NPV values at the right anterior prefrontal electrode (Fp2), the right temporal electrode (T4) and the right occipital electrode (O2).

Figure 2 | Distributions of the discriminant score in responders and non-responders. Using beta NPVs at Fp2, T4 and O2 electrodes as discriminating variables, linear discriminant analysis yielded a discriminant score that separates responders and non-responders. With zero as the cut-off value, a discriminant score > 0 indicates shunt responders and a discriminant score < 0 indicates non-responders. This score could correctly identify responders and shunt non-responders with a positive predictive value of 80% (8/10) and a negative predictive value of 88% (7/8).
consistent with the symptom of gait disturbance in iNPH patients. However, non-responders had lower beta NPV values in the right VAN, something that contradicts gait disturbance status. This paradoxical stabilization of beta cortical electrical activity might indicate irreversible cortical impairment.

Altogether, NPV analysis can sensitively detect the difference of right VAN activity between responders and non-responders even in pre-CSF tapping EEG data, unlike other neuroimaging methods, such as fMRI, SPECT and EEG power analysis. This may rise from the fact that EEG directly detects cortical electrical activity with high temporal resolution and that NPV analysis is highly sensitive to instability of cortical electrical activity. Our findings indicate that NPV in the beta frequency band can be an electrophysiological hallmark for the prediction of shunt operation outcome without the need of performing CSF tapping. An important therapeutic implication of our results is that NPV analysis could be a useful non-invasive method to discriminate shunt responders from non-responders among the many untreated iNPH patients of the general population.

Our results should be interpreted with caution because of the following limitations: First, the sample size of iNPH patients was small and our results were not corrected for multiple comparisons. Therefore, our findings were preliminary and their validity should be further tested in notably larger size of patients. However, our main findings of NPV differences between responders and non-responders are consistent with the neuroimaging findings of iNPH and the role and spatial distribution of relative cortical functional networks. Therefore, we can assume that NPV is a reliable measure of cortical electrical activity. Second, we did not directly detect an involvement of the right VAN in gait recovery before and after shunt operation. Therefore, further studies are needed to ascertain the correlation between the right VAN activity and gait recovery.

In conclusion, our research showed that NPV reliably detects the differences of cortical electrical activity between shunt responders and non-responders in the beta frequency band, at right VAN areas of patients with iNPH. This could be done without the need of performing CSF drainage, when no other neuroimaging method has so far been able to do so. Based on our results, we suggest that NPV in the beta frequency band at right VAN areas can correctly distinguish responders from non-responders to shunt operation before the operation itself. Therefore, NPV in beta frequency band could be a powerful non-invasive tool in predicting shunt operation outcome in patients suffering from iNPH, a conclusion of important therapeutic implications.

Methods

Subjects. iNPH patients who underwent shunt operation between April 2010 and December 2013 were consecutively recruited from the Neuropsychological Clinic of the Department of Neuropsychiatry of Osaka University Hospital. The inclusion criteria were: (1) age > 60 years; (2) at least one of the following three symptoms: gait disturbance, cognitive impairment and urinary incontinence; (3) dilated cerebral ventricles and narrowed subarachnoid space at the high convexity without severe cortical atrophy on MRI; (4) absence of diseases or conditions that might cause relative clinical symptoms or imaging findings; (5) no history of severe head trauma, subarachnoid hemorrhage, meningitis, tumor and aqueduct stenosis; (6) normal CSF pressure (<200 mm H2O) and contents at lumbar puncture, (7) right handedness and (8) a positive response to CSF tap test based on the Japanese Clinical Guidelines for iNPH16 and undergoing a shunt operation. Exclusion criteria were: (1) comorbidities of motor or psychiatric disorders, such as Parkinson’s disease, Alzheimer’s disease or schizophrenia (Alzheimer’s disease was defined by severe episodic memory impairment14 and a higher Alzheimer index (>3438), evaluated using the measures of amyloid beta and tau protein in CSF12) and (2) a lack of 500 seconds (500-s) artifact free epochs in the EEG recordings. A total of 23 patients met the inclusion criteria, of which one patient was excluded due to comorbidities and four patients due to lack of artifact free epochs. In the end, 18 patients were included in this study. 13 patients were followed up for one-year and five patients for six months after shunt operation.

This study was approved by the Ethics Committee of Osaka University Hospital and written informed consent was obtained from the participants or their families. CSF tap test and shunt operation were carried out in accordance with the Japanese Clinical Guidelines for iNPH.

Gait assessment. Assessments of gait and cognition have already been described previously; for further details, please refer to our previous study4. Gait disturbance was assessed by the 10-meter reciprocating walking test (WT), the 3 m Timed Up and Go (TUG) test17 and the Gait Status Scale (GSS)18. The improvement thresholds were set at 10% improvement in the WT and TUG, and 1 point improvement in the GSS, based on standard deviations of gait function tests in iNPH patients in previous studies13,19–20. Improvement in WT, TUG and GSS was identified as clinical improvement in gait domain.

Cognitive assessment. Cognitive impairment was assessed by the Frontal Assessment Battery (FAB), Mini-Mental State Examination (MMSE), Wechsler Memory Scale-Revised (WMS-R) Attention-Concentration Index, Wechsler Adult Intelligence Scale-III (WAIS-III)-Block Design/Digit Symbol Coding, and Trail Making Test Part A (TMT-A). The improvement thresholds of the cognitive tests were set at 2, 4, 15, 3, 3 points and 30% respectively, based on previous studies13,19–20. Improvement in more than half of these cognitive tests was identified as clinical improvement in cognitive domain.

Assessment of shunt operation outcome. Before shunt operation, WT and TUG were performed in the morning and afternoon for five consecutive days, in order to reach a plateau of improvement by practice, and then the fastest time was adopted as the time of WT and TUG before CSF tapping. Gait disturbance and cognitive impairment were assessed by the aforementioned tests. Urinary incontinence was excluded from assessment in our study because of low reliability, as the frequency of micturition was sometimes only self-reported. After shunt operation, gait disturbance and cognitive impairment were evaluated at post-operative visits of 1, 3, 6 months, and 1 year. Shunt operation was considered positive if at least one symptom showed substantial improvement at any time of assessment after operation; otherwise it was considered negative. Consequently, the patients were classified as responders and non-responders to shunt operation. This classification criteria is stricter than the Japanese Clinical Guidelines for iNPH15, because we selected patients who showed significant improvements in iNPH symptoms as responders.

EEG recording. EEG data acquisition and the procedure of power spectral and NPV analysis have already been described in detail elsewhere21. EEG data during awake, eyes-closed resting condition were recorded for about 20 minutes using a 19-electrode EEG system (EEG-1000/EEG-1200; Nihon Kohden Inc., Tokyo, Japan), and filtered through a band-pass filter of 0.53 to 120 Hz with a sampling rate of 500 Hz. Subjects were instructed to close their eyes, relax, but stay awake. During the EEG sessions, drowsiness was avoided by providing instructions once again. For each recording, 500-s artifact free epochs were selected off-line, and imported into NPV and Power analyses.

NPV analysis. In NPV analysis program, all electrodes were re-referenced to the average reference (i.e. mean electrical potential of the 19 electrodes). NPV was calculated for every 2.56-s EEG epoch, where NPV was defined as the variance of power divided by the square of mean power, in order to obtain relative values correctly among different subjects. The output of the NPV analysis program was a z-score spatial map, which displays how many units of the standard deviation the observed NPV values at each electrode site were above or below the mean NPV values of healthy controls22. The healthy controls used in the program (27 men and 25 women, age 71.5 ± 8.4 years) had normal results in cognitive tests and magnetic resonance imaging (MRI), without any history of psychiatric or neurological disorders.

In our study, NPVs of 2.56-s EEG epochs were collected at 0.64-s steps for the whole EEG epoch (moving average filter method). Then, collected NPVs were averaged and finally the stationary mean NPV value was obtained for each subject21. These analyses were executed for five frequency bands: delta (2–4 Hz), theta (4–8 Hz), alpha (8–13 Hz), beta (13–30 Hz), and gamma (30–40 Hz).

Power analysis. Power was defined as the square of the amplitude of the EEG signal at each frequency band. Power spectral analysis was done by QP-229A Neurormap software (Nihon Kohden Inc., Tokyo, Japan) in which all electrodes were re-referenced to the average reference. In this study, the output of the power spectral analysis was also displayed as a z-score spatial maps, using the same healthy controls as in NPV analysis.

Statistical group analysis and linear discriminant analysis. The differences in NPV or EEG power in each frequency band at each electrode site between responders and non-responders were assessed by the independent Student’s t-test. The level of significance for these tests was set at p < 0.05. These results were not corrected for multiple comparisons, and the validity of each significant difference in NPV is discussed in the discussion section. In order to find an index that most accurately distinguished responders from non-responders, we selected NPVs in beta frequency band that had significant differences between responders and non-responders as discriminating variables and performed linear discriminant analysis using the SPSS software version 12.0 (SPSS Japan Inc., and IBM Company Tokyo, Japan).

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Author contributions

Y.A., Hiroaki K. and R.I. designed the study. Y.A. and R.I. analyzed the EEG data and wrote the manuscript. T.T. provided data of Alzheimer index. R.I., L.C., T.W., S.I., M.H., T.K., H.K., T.Y., K.N., K.Y., M.I. and M.T.) reviewed the acknowledgment programs of NPV and power analyses. H.K., T.W., H.K., T.Y., K.N. and K.Y. recruited the patients. T.T. provided data of Alzheimer index. T.M., H.M. and K.I. provided additional information.

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