INTRODUCTION

Delusion is one of the most common behavioral symptoms associated with Alzheimer’s disease (AD), disturbing patient quality of life and cognitive functions. The prevalence of delusion in AD has been reported to be about 36%, ranging from 9.3% to 63%. Among various kinds of delusions, delusion of theft (DT) is the most frequent type in AD. Several prior studies investigated the relationship between cognitive functions and DT in AD, but the results remain inconclusive. Although not specified for DT, some studies reported that delusion correlated significantly with cognitive function test scores such as the Mini Mental State Examination (MMSE), whereas other studies reported that there was no significant correlation.

Although delusion of theft (DT) is the most frequent type of delusion in Alzheimer’s disease (AD), its relationship to cognitive functions remains unclear. The aim of this study was to demonstrate the relationship between DT and cognitive functions in mild AD. Two hundred eighty-nine mild AD patients were enrolled in this study. These subjects were classified into three groups: patients with no delusions (ND, n=82), patients with paranoid delusions (PD, n=114) and patients with DT (n=93). Cognitive functions and their associations with the degree of delusion were compared among the three groups. The results showed that verbal Fluency scores were significantly lower in the PD group than in the DT and ND groups. Word List Recall scores were significantly lower in the DT group than in the PD and ND groups. Interestingly, delusion severity measured with the Neuropsychiatric Inventory delusion subscale correlated negatively with the Word List Recall scores in the DT group. In this study, we demonstrated that episodic memory functions in mild AD patients were associated with DT, but not with PD. Further studies might be needed to clarify the pathophysiology of delusions associated with AD.

METHODS

Two hundred eighty-nine patients who visited St. Vincent’s Hospital were recruited for this study. The inclusion criteria were 1) a diagnosis of probable AD according to the National Institute of Neurological and Communicative Disorders and Stroke (NINCDS) and the Alzheimer’s disease and Related Disorders Association (ADRDA) criteria and 2) a score on the Clinical Dementia Rating Scale (CDR)=1. Subjects who had other neurological or psychiatric conditions were excluded. The Korean version of the Consortium to Establish A Registry for Alzheimer’s Disease (CERAD-K) neuropsychological assessment battery was used to assess cognitive functions, including Verbal Fluency (category switching), Word List Recall (episodic memory), and MMSE (general cognitive function).

Key Words Delusion of theft, Cognitive function, Alzheimer’s disease.

BRIEF REPORT

Relationship between Delusion of Theft and Cognitive Functions in Patients with Mild Alzheimer’s Disease

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Key Words Delusion of theft, Cognitive function, Alzheimer’s disease.
Delusion of Theft and Cognitive Functions

logical battery, including Verbal Fluency (VF), 15-item Boston Naming Test (BNT), MMSE, Word List Memory (WLM), Word List Recall (WLR), Word List Recognition (WLRc), Constructional Praxis (CP), and Constructional Recall (CR), was used to assess cognitive functions. In addition, the Neuropsychiatric Inventory (NPI) was used to assess the severity of behavioral symptoms. This study was approved by the Institutional Review Board of the Catholic University of Korea.

Written informed consent was obtained from all subjects and their guardians (IRB No. VC15EISI0044).

Study participants were classified into three groups according to the presence and type of delusion: AD with no delusions (ND), AD with PD (the false belief that one is being harmed or persecuted by a particular person or group of people) and AD with DT (the false belief that someone has stolen something). Most of paranoid themes were related with ‘my family (someone) is trying to harm or kill me.’ One of the contents was “My daughter-in-law seems to have poisoned my rice.” DT includes 'my daughter stole my jewelry' or 'my son seems to have taken my pocket money.' We excluded subjects who had multiple types of delusions. The severity of delusions was measured with the NPI delusion subscale.

Statistical analyses of demographic data were performed with the Statistical Package for the Social Sciences (SPSS, version 12.0, SPSS Inc., Chicago, IL, USA). The assumption of normality was tested for all continuous variables with the Kolmogorov-Smirnov test. All variables were normally distributed except the NPI delusion subscale scores. The independent t- and χ² tests were used to assess potential differences among the three groups for all demographic variables. In addition, the Mann-Whitney U test was used to compare the group differences in the NPI delusion subscale between the PD and the DT groups.

Analysis of covariance (ANCOVA) and post hoc tests using Tukey’s multiple comparison tests were performed to identify group differences in cognitive functions after controlling for age, education and gender effects. In addition, as the NPI delusion subscale value of the each group was not normally distributed, the non-parametric Spearman correlation analysis was used to identify the association pattern between cognitive functions and the degree of delusion in the PD and DT groups.

For all statistical analyses, a two-tailed p-value less than 0.05 was considered statistically significant.

RESULTS

Demographic data

There were no significant differences in baseline demographic characteristics among the three groups (Table 1). There were no differences in NPI scores between the PD and DT groups.

| Table 1. Demographic and clinical characteristics of the study participants |
|------------------------|------------------------|------------------------|------------------------|
|                        | ND (N=82)              | PD (N=114)             | DT (N=93)              |
| Age (years±SD)         | 76.9±6.5               | 78.4±6.0               | 77.5±6.7               |
| Education (years±SD)   | 7.8±2.9                | 7.9±2.9                | 7.6±3.2                |
| Sex (M:F)              | 28:54                  | 27:87                  | 29:64                  |
| CDR                    | 1                      | 1                      | 1                      |
| CDR-SB                 | 4.0±3.7                | 4.1±2.9                | 4.1±3.1                |
| S-IADL                 | 19.8±5.5               | 19.5±6.4               | 19.2±6.8               |
| S-ADL                  | 0                      | 0                      | 0                      |
| CERAD-K battery (SD)   |                        |                        |                        |
| VF                     | 6.6±3.2                | 5.4±2.6                | 7.4±3.6                |
| BNT                    | 5.6±2.7                | 5.2±2.8                | 6.0±2.9                |
| MMSE                   | 20.8±3.6               | 20.0±3.6               | 20.5±2.5               |
| WLM                    | 7.2±3.6                | 6.8±3.6                | 6.5±2.1                |
| CP                     | 7.2±2.2                | 7.1±2.2                | 7.5±2.3                |
| WLR                    | 1.7±1.1                | 1.8±1.6                | 0.9±1.4                |
| WLRc                   | 3.2±2.6                | 3.7±3.1                | 3.5±2.9                |
| CR                     | 1.0±1.4                | 1.0±1.7                | 0.9±1.6                |

ND: AD without any delusions, PD: AD with paranoid delusion, DT: AD with delusion of theft, SD: standard deviation, NS: not significant, CDR: clinical dementia rating scale, CDR-SB: sum of box of the clinical dementia rating scale, S-IADL: Seoul instrumental activities of daily living, S-ADL: Seoul basic activities of daily living, CERAD-K: Korean version of Consortium to Establish A Registry for Alzheimer’s disease, VF: verbal fluency, BNT: 15-item Boston Naming Test, MMSE: Mini Mental State Examination, WLM: word list memory, CP: constructional praxis, WLR: word list recall, WLRc: word list recognition, CR: constructional recall
Cognitive functions and delusions

Figure 1 and Table 1 present group differences in cognitive function scores. The VF (p<0.001) and WLR (p<0.001) scores differed significantly among the three groups. Post hoc analysis revealed that the VF scores of the PD group were significantly lower than those of the DT (p<0.001) and ND (p=0.02) groups. The WLR scores of the DT group were significantly lower than those of the PD (p<0.001) and ND (p=0.004) groups. On the other hand, WLR scores did not differ significantly between the ND and PD groups. Interestingly, WLR (Spearman's rho=-0.23, p=0.026) and CR (Spearman's rho=-0.21, p=0.045) scores correlated negatively with delusion severity measured by the NPI delusion subscale in the DT group. However, there were no significant relationships between cognitive functions and the degree of delusion in the PD group.

DISCUSSION

To the best of our knowledge, this study is the first to demonstrate distinctive association patterns between DT and episodic memory disturbances in mild AD patients. The strength of our study resulted from our recruitment of a relatively large sample and our exclusion of subjects with co-existing multiple delusions.

In this study, we found that WLR scores were significantly lower in the DT group than in the other two groups. In addition, delusion severity correlated negatively with WLR/CR scores in the DT group. These relationships might be attributable to the memory-associated nature of DT itself (i.e., the belief that someone has stolen something). In other words, DT may be the most common delusion associated with AD because the primary symptom of AD is memory impairment. On the other hand, PD group who had preserved memory functions compared to the theft delusion group, showed decreased verbal fluency. In general, verbal fluency is known to reflect executive function of frontal lobe. It means PD might be related with executive functions perceive or integrate situations rather than the memory impairments. We assume that this difference between two groups might originate from distinct neural substrates. To date, several neuroimaging studies have found that DT is associated with memory-related neuronal substrates in AD patients. A structural neuroimaging study using computed tomography revealed that right-left medial temporal lobe asymmetry was associated with DT in AD. The present study suggests that this asymmetrical pattern might be related to the memory disturbances in AD patients. Another functional neuroimaging study using SPECT investigated the neuronal substrate for DT in AD. Subjects with DT were found to have significantly lower perfusion in the right medial posterior parietal region (which is known for episodic memory processing) than patients without delusions. Despite the evidence from neuroimaging studies suggesting a distinctive association between episodic memory impairments and DT, few clinical studies have found this association. A previous study identified no significant differences in the scores on Hasegawa’s dementia scale (HDS-R), the MMSE or the Neurobehavioral Cognitive Status Examination (COG-NISTAT) sub-scales between the DT (n=14) and ND (n=42) groups. In addition, another study indicated that delayed recall scores did not correlate significantly with the severity
of DT in 22 AD subjects. These findings might be attributable to the small sample sizes and the non-exclusion of patients with other types of delusions, such as persecutory, phantom and misidentification delusions.

In this study, we found that the VF scores of the PD group were significantly lower than those of the ND and DT groups. These findings are consistent with previous reports of deficient VF and conceptualization in AD patients and of VF and executive functioning deficits in non-demented elderly people with PD.

Several limitations of our study must be taken into consideration. First, to clearly distinguish DT from PD, we could not include AD patients with mixed delusions and other types of delusions, such as misidentification and phantom delusions. In real clinical settings, mixed delusional AD patients are more common. Second, as we included subjects with mild-stage AD, our results cannot be generalized to subjects with more severe AD. Larger studies including a broader spectrum of AD subjects would be needed. Third, despite the decent sample size, the NPI delusion subscale value of the each group was not normally distributed. These skewed distributions of the NPI delusion subscale values may affect the association pattern between the NPI delusion subscale and the VF values which were significantly lower than any other group in the PD group. Therefore, regression analysis using normally distributed between the NPI delusion subscales and the cognitive function measures may helpful to predict impact of cognitive dysfunction on delusion severity precisely.

In conclusion, we demonstrated that episodic memory functions in mild AD patients were associated with DT, but not with PD. Furthermore, mild AD with delusions, regardless of the type, might have a distinct pathophysiology from mild AD without delusions. Further studies are needed to clarify the pathophysiology of delusions associated with AD.

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