Are There Any Connections between Language Deficits and Cognitive Slowing in Alzheimer’s Disease?

Michael Schecker, Carsten Kochler, Klaus Schmidtke, Reinhold Rauh

Klinik für Psychiatrie, Psychotherapie und Psychosomatik im Kindes- und Jugendalter, Universität Freiburg, Freiburg, Klinik für Hirnleistungsstörungen Klausenbach, Nordrach, and Neurolinguistisches Labor NLL, Klinik für Psychiatrie, Psychotherapie und Psychosomatik im Kindes- und Jugendalter, Universität Freiburg, Freiburg, Germany

Key Words
Alzheimer’s disease · Brain function in dementia · Cognitive slowing · Deficits in language processing

Abstract
Background: Speech disorders already occur in the early phases of Alzheimer’s disease (AD). As a possible cause, problems of executive processes are discussed. Cognitive slowing is also repeatedly addressed. Aims: Are there any connections between cognitive slowing and speech disorders in AD? And is there a relationship between cognitive slowing and executive processes? Methods: The data of 72 healthy controls and 52 AD patients were examined with regard to their language performance and their response times in a computerized Stroop paradigm. Results: The AD patients showed significantly worse results in all language tests as well as much longer reaction times in all Stroop conditions, especially in the interference condition (Stroop 3). Speech errors and response times correlated with severity (MMSE), and the speech errors correlated with the reaction times in Stroop 3 (interference condition, which reflects the processing time of executive processes). Conclusion: The most interesting question now is: How are language processing and executive processing time (Stroop 3) related?

Introduction
In early stages of Alzheimer’s disease (AD), there are always deficits in language processing. On the pragmatic level, these processing difficulties have been associated with executive processes and the working memory (here the central executive) [1–5] (for the use...
of pronouns, see [6, 7], for the resolution of ambiguity, see [8–10], and for the understanding of idioms, see [11–13]). But there is evidence that, for example, naming disorders or errors on the syntactic level also have something to do with the deterioration of working memory processes [14–17] (for an overview, see [18]).

The fact that the cognitive processing speed in AD patients is significantly reduced is also well known (‘cognitive slowing’, e.g. with regard to the Stroop test [19–22] or to priming experiments [23]). Is there a connection between the disorders of language processing and cognitive slowing in AD? And is it possible to identify a specific link between cognitive slowing and specifically executive processes?

**Methods**

**Subjects**

Our sample consisted of 52 subjects (40 female, 12 male) with mild AD (fulfilling the DSM-IV [24] and NINCDS-ADRDA criteria [25]; see also Petersen [26, 27] on the differences to mild cognitive impairment) and 72 healthy controls (48 female, 24 male). The average age within the experimental group was 73.8 years (SD = 5.23), and they had a mean of 10.8 years (SD = 2.87) of education. The average age within the control group was 72.6 years (SD = 5.43), and they had a mean of 11.9 years (SD = 2.2) of education. Both groups differed in the MMSE score [28] (AD: mean = 22.5, SD = 2.7; controls: mean = 28.4, SD = 1.01). The differences were also statistically significant [ANOVA, F(2, 36) = 65.8, p < 0.001]. All patients were medicated with an acetylcholinesterase inhibitor and were well adjusted.

The participants were recruited through the ZGGF (Center for Geriatrics and Gerontology), University Medical Center Freiburg, and the memory clinic of the University Hospital Heidelberg, both Germany. Additional subjects were recruited through institutions of assisted living and residential care homes for the elderly in Freiburg, as well as through advertisements in the local press.

We used the following exclusion criteria: other psychiatric or neurological disorders; a score on the ‘reduced’ Geriatric Depression Scale [29] of 5–15 points; receiving psychotropic drugs; <6 years of education; mother tongue other than German; impairments of the peripheral vision and hearing (control of the speech-relevant frequencies 500/1,000/2,000/4,000 Hz using an audiometer; excluded if hearing achievements were only about 45 dB), and no right-handedness (Edinburgh Handedness Inventory [30]). All subjects were also tested for their reading comprehension (brief written requests for action).

All participants were considered to have the necessary abilities to give their informed consent for participation. Appropriate care was taken in explaining the study, and they were given sufficient time to reach a decision. It was made clear to all subjects that they would not be treated any differently if they chose not to participate.

**Materials and Procedure**

We used 2 types of tests. First, we tested the cognitive processing speed with the Stroop Color-Word Test [19, 21, 22, 31]. It is generally accepted that the third condition (interference condition) measures higher cognitive abilities or, more precisely, executive performance. We used a modified computerized Stroop paradigm [32]. When measuring time, we considered only correct responses.

Second, we tested the ability to represent context and to integrate contextual cues in information processing using the PMA® (Predictive Monitoring in AD, a test package that we developed at the Neurolinguistic Laboratory Freiburg [10]). Here we used the 3 subtests shown in figures 1–3 (number of test items per test = 10, no time limit). The statement in
Fig. 1. Comprehension of homonyms.

‘Er bindet sich die Fliege um’ / ‘He puts on the bow tie’
Which picture agrees best with the statement?
(Here, the correct image is highlighted by a red arrow.)

Fig. 2. Comprehension of idioms.

‘What does this usually mean?’
(Here, the correct answer is marked by a red arrow.)

Fig. 3. Processing of pronouns.

‘Please insert the appropriate expressions in the gaps of the text.’
(Here, the correct expressions are framed in red.)
Table 1. Stroop performance A

| Condition               | Response time (average), ms | Group differences (one-way ANOVA with Brown-Forsythe correction) |
|-------------------------|-----------------------------|---------------------------------------------------------------|
|                         | controls (n = 72)           | AD (n = 52)                                                  | F(1, 13.09) = 31.00, p < 0.0005 |
| Stroop 1                | 821 (132)                   | 1,468 (465)                                                 | F(1, 13.28) = 28.19, p < 0.0005 |
| Stroop 2                | 856 (109)                   | 1,523 (405)                                                 | F(1, 16.24) = 50.00, p < 0.0005 |
| Stroop 3 (with interference) | 941 (152)               | 2,372 (726)                                                 |                                             |

Stroop 1: color > color (e.g. ◼ > ◼). Stroop 2: meaning > color (e.g. ‘green’ > ■). Stroop 3: color of the letters > color (e.g. ‘green’ > ●). In the evaluation of the processing times only the correct trials were included.

Table 1 shows the processing times (average) of the 2 simple conditions and the interference condition. The repeated-measures ANOVA shows an interaction of group and condition [F(2, 28) = 9.80, p < 0.001] and main effects for both, condition [F(2, 28) = 37.10, p < 0.0005] and group [F(1, 29) = 65.09, p < 0.0005]. The multiple comparisons with Bonferroni adjustment between the different Stroop conditions in the total sample found no significant differences between the 2 simple conditions Stroop 1 and Stroop 2 [F(1, 29) <1, not significant]. However, the simple conditions differ from the complex condition [Stroop 1 to Stroop 3: F(1, 29) = 62.27, p < 0.0005; Stroop 2 to Stroop 3: F(1, 29) = 67.76, p < 0.0005]. The same pattern is also found in the AD group [Stroop 1 to Stroop 2: F(1, 12) <1, not significant; Stroop 1 to Stroop 3: F(1, 12) = 36.55, p < 0.0005; Stroop 2 to Stroop 3: F(1, 12) = 38.25, p < 0.0005]. Table 1 shows the differences between AD and control subjects.

AD patients (early stages) need significantly more time than the controls, and this slowdown is already present in the simple conditions. However, it is particularly interesting that the processing times of the 2 simple conditions and the interference condition do not rise in parallel. The patients are 1.78 and 1.77 times slower than the healthy controls in Stroop 1 and Stroop 2, respectively, and 2.52 times slower in Stroop 3. Moreover, in the AD group, the
increase in processing time of Stroop 2 to Stroop 3 is 10 times the corresponding increase in the controls (fig. 4).

Are there any connections between the cognitive slowing and the severity of the disease (MMSE)? Yes, but only in the AD group and only for Stroop 3 (table 2).

Table 3 shows the error rates of the PMA subtests. Because the assumption of a normal distribution for the PMA must be rejected (significant Kolmogorov-Smirnov test for all variables), we used non-parametric tests for 2 independent samples (Wilcoxon-Mann-Whitney test) to test for group differences.

In the naming task (naming of real objects), the AD patients did not differ from the controls.

A qualitative analysis of the errors in the pronoun test showed that (a) in 56.2% of the cases, a definite noun phrase was chosen (e.g. ‘die Frau’/’the woman’) instead of the correct definite pronoun; (b) in less than 8% of the cases, the morphological genus was wrong, obviously referenced incorrectly, and (c) in more than 89% of the cases, the errors reflected no wrong reference; the patients had very well understood to which reference the objects were addressed in the text.

Are there any connections between the PMA errors and the severity of the disease (MMSE)? Yes, but only in the AD group (table 4). In the AD group (and only there), we have also found correlations between the PMA errors and the Stroop response time in the interference condition (table 5).

Fig. 4. Stroop performance B. RT = Response time.

### Table 2. Correlations between response time and MMSE in the AD group

| Condition       | Spearman correlations (one-sided) |
|-----------------|-----------------------------------|
| Stroop 1 – MMSE | r = -0.486, p = 0.06, n.s.        |
| Stroop 2 – MMSE | r = -0.344, p = 0.06, n.s.        |
| Stroop 3 – MMSE | r = -0.655, p = 0.002             |

The correlation coefficient r > 0.5 indicates a large effect [36]. n.s. = Not significant.
There seems to be no general relationship between time consumption in Stroop and errors in error scoring tests (PMA errors). But there is a connection between the time consumption in Stroop 3 (regarding executive processes) and the PMA errors. How can we explain this?

### Discussion

The severe deficits in language tests and the positive correlation between PMA errors and the MMSE in the AD group were to be expected [18]. Also expected was the significant slowing of cognitive processing [19–23] and their relationship to the severity of the disease (MMSE). But why is the time consumption increased so much just with executive processes?

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**Table 3. PMA performance**

| PMA subtests | PMA errors controls | PMA errors AD patients | Group differences (Wilcoxon-Mann-Whitney test) |
|--------------|---------------------|------------------------|-----------------------------------------------|
| Homonyms     | 9.9 (0.3)           | 9.0 (1.3)              | Z = –2.66, p = 0.009                          |
| Idioms       | 9.4 (0.9)           | 8.0 (2.9)              | Z = –2.48, p = 0.014                          |
| Pronouns     | 5.4 (0.6)           | 3.7 (1.6)              | Z = –2.88, p = 0.005                          |

The mean number of correct responses (SD) on the language tasks for each group and results of group differences on the language tasks (computed value of Wilcoxon-Mann-Whitney test).

**Table 4. Correlations between PMA errors and MMSE in the AD group**

| Subtests              | Spearman correlations (one-sided) |
|-----------------------|-----------------------------------|
| Homonyms – MMSE       | r = –0.55; p = 0.004              |
| Idioms – MMSE         | r = –0.59; p = 0.018              |
| Pronouns – MMSE       | r = –0.69; p = 0.002              |

The correlation coefficient r > 0.5 indicates a large effect [36].

**Table 5. Spearman correlation (one-tailed): PMA errors with Stroop response time**

| PMA subtests and Spearman correlation | Stroop conditions  |
|--------------------------------------|-------------------|
|                                      | 1                |
|                                      | 2                |
|                                      | 3 (interference)  |
| Homonyms                             | +0.23            |
| Correlation                           | +0.26            |
| Significance (one-tailed)            | +0.53            |
| Idioms                               | 0.11             |
| Correlation                           | 0.08             |
| Significance (one-tailed)            | 0.001            |
| Pronouns                             | +0.06            |
| Correlation                           | +0.16            |
| Significance (one-tailed)            | +0.55            |
|                                      | 0.37             |
|                                      | 0.19             |
|                                      | 0.001            |
| Pronouns                             | +0.18            |
| Correlation                           | +0.29            |
| Significance (one-tailed)            | +0.60            |
|                                      | 0.172            |
|                                      | 0.06             |
|                                      | 0.001            |

The results show a correlation only between the cognitive slowing of executive processes (Stroop 3) and PMA errors.
We suspect that the executive subprocesses essentially run in a linear sequence, so that the slowdown of the individual processing steps added up.

How can we explain the positive correlation of PMA errors and Stroop 3 response time (interference condition)? There are 2 complementary explanations: (a) working memory is limited in time, and the strong slowdown of executive subprocesses overwhelmed working memory, and (b) the very strong slowdown of executive subprocesses leads to desynchronization phenomena, to a disintegration of basal and higher cognitive processing steps.

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