Computer-Assisted Cognitive Rehabilitation in Stroke and Alzheimer’s disease

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Received date: Oct 23, 2014, Accepted date: Dec 10, 2014, Published date: Dec 15, 2014

Abstract

Background: The aim of study was to compare the effects of computer-assisted cognitive rehabilitation (CR) in mild cognitive impairment after stroke and in patients with Alzheimer disease.

Methods: The study included 21 patients after ischemic stroke (12 males, 9 females, age median 60.5 age range 38–81 years) and 15 patients with AD (8 males and 7 females 71.5, 50–86 years). We administered WAIS-III, MMSE, and ACE-R to evaluate MCI. NEUROP-4 software was employed for CR. Rehabilitation was carried out for two periods of 1.5 h each week for 3 months.

Results: In the stroke group we recorded significantly higher scores for the following parameters after CR: IQc (median 84 before vs 88 after p = 0.001), IQv (83 vs 92, p = 0.029), IQp (78 vs 86, p = 0.001), VC (91 vs 97, p = 0.017), PO (82 vs 94, p = 0.001), SOP (71 vs 8, p = 0.0003), ACE-R (79 vs 84, p = 0.01) In the AD group only the ACE-R was increased (75 vs 83, p = 0.008).

Conclusions: Our study demonstrates that the beneficial effects of computer-assisted CR in patients with MCI are more significant in stroke patients than in AD patients.

Keywords: Cognitive rehabilitation; Alzheimer disease; Stroke; Cognitive impairment

Abbreviations:

AD: Alzheimer disease; ACE-R: Addenbrooke’s Cognitive Examination, revised; CR: Cognitive rehabilitation; HADS: Hospital Anxiety and Depression Scale; IQc, IQv, IQp: IQ score global; IQc, IQv, IQp: IQ score performance; IQc, IQv, IQp: IQ score verbal; MMSE: Mini-Mental State Examination; WM: Working memory; PO: Perceptual organization; SOP: Speed of processing; VC: Verbal comprehension; WAIS-III Wechsler Adult Intelligence Scale, third revision

Background

Cognitive functions are impaired in brain disorders including neurodegenerative diseases, stroke, and brain trauma. Treatment is typically pharmacological [1,2] and, in some cases, neurosurgical. However, non-pharmacological intervention, especially cognitive rehabilitation, was often neglected in the past but now there have been increasing efforts to develop cognitive interventions to ameliorate cognitive problems experienced by older adults, especially AD, stroke and brain trauma patients [3,4].

Computer-based programs specifically targeted to dementia or mild cognitive impairment have been developed as a support in rehabilitation of cognitive areas and daily living functions [5]. Computer-assisted CR is a cost-effective method for providing individualized treatment, based on each patient’s neuropsychological patterns, in which impaired cognitive areas are repeatedly stimulated [6]. Computer interventions are rapidly becoming popular and cognitive exercise has been successfully implemented because older adults are often the fastest growing group of computer technology users [7].

CR provides a structured practice of complex mental activity designed to enhance cognitive function [3,8]. Neurplasticity is a prerequisite for improvement of cognitive impairment following CR [9,10]. Several studies have been carried out on the effect of cognitive training and rehabilitation in different diseases associated with cognitive impairment. Most studies have reported mild beneficial effects of CR in patients with neurodegenerative disease such as AD [5,9,11-13]. However, studies on CR in stroke patients have been more promising [10,14-17]. Data comparing the effects of CR in patients with acute brain lesions versus chronic progressive disease, notably neurodegenerative conditions such as AD, are needed.

The aim of this pilot study was therefore to compare the effects of computer-assisted CR in patients with mild cognitive impairment following stroke versus patients with mild cognitive impairment attributed to probable AD.

Methods

Cognitive rehabilitation: For computer-assisted rehabilitation we used the software NEUROP-4. This program provides multimodal and
multiple-domain training of cognitive function. In order to train
memory and concentration function, we used non-verbal tasks such as
assembling shapes or figures, getting through a labyrinth, memorizing
cards and shapes. We used tasks focused on planning and strategic
thinking for executive functions training, e.g. London Tower, Hanoi
Tower, etc.

Rehabilitation was carried out for two periods of 1.5 h each week for
3 months. Rehabilitation was supervised by a neuropsychologist. The
battery of the cognitive tests was administered both before the start of
computer-assisted CR and after completion of CR. The effect of CR
was assessed as the difference between baseline scores and post-
training scores within the each group, and also by comparison between the
two groups.

Data analysis: All results are expressed as median and minimum–
maximum values. SPSS v.15 (SPSS Inc., Chicago, USA) statistical
software was used to analyze the data. The Wilcoxon signed-rank test
was applied to compare the paired data, and the Mann–Whitney U test
was used to compare the groups. Spearman’s correlation analysis was
carried out to evaluate the relationship between age, other baseline
variables and differences in cognitive scores in both groups. Analysis
of covariance ANCOVA was used for analysis of differences between
the groups with age as a covariate. Nonparametric tests were used
owing to small sample sizes. P <0.05 was taken to indicate statistical
significance.

Results

We included 21 patients in our prospective study after ischemic
stroke (12 males, 9 females, age median 60.5, age range 38–81 years)
and 15 patients with probable AD (8 males and 7 females, age median
71.5, age range 50–86 years). Patients in both groups were matched
with respect for age and sex. Table 1 presents the results of the
cognitive tests before beginning the treatment in both groups. Table 2
compares the scores on the cognitive test battery at baseline and
following CR, as presents the changes in scores following CR.

Table 1: Baseline characteristics of the study population; a … Fisher’s
exact test, b … Mann-Whitney U test Data in the table are presented
as median (minimum – maximum). *Spearman correlation did not
prove significant relationship between age and other baseline variables
in both groups.

In the stroke group, the Wilcoxon signed-rank test demonstrated
statistically significant increased scores after the therapy for the
majority of parameters, including WAIS-III, ACE-R, and MMSE. IQc (median 84 before vs 88 after, p = 0.001), IQv (median 83 vs 92, p = 0.029), IQp (median 78 vs 86, p = 0.001), VC (median 91 vs 97, p = 0.017), PO (median 82 vs 94, p = 0.001), SOP (median 71 vs 8, p = 0.0003), ACE-R (median 79 vs 84, p = 0.01) Only one parameter, VM, did not change significantly in the stroke group (median 84 vs 85, p = 0.833).

Table 2: Scores on the cognitive test battery at baseline and after
cognitive rehabilitation; p … significance of Wilcoxon signed-rank test
for comparing before and after therapy, score median (min-max)

| Cognitive domains | AD (n=15) | Stroke (n=21) |
|-------------------|----------|--------------|
|                   | Before   | After        | p    |
|                   |          |              |      |
| IQc               | 87 (74-110) | 92 (66-112)  | 0.61 |
| IQv               | 95 (61-119) | 98 (60-119)  | 0.755|
| IQp               | 81 (68-105)| 85 (57-114)  | 0.208|
| VC                | 93 (34-126)| 102 (64-126) | 0.835|
| PO                | 84 (30-109)| 90 (60-107)  | 0.889|
| WM                | 90 (32-113)| 94 (70-110)  | 0.561|
| SOP               | 81 (7-113)| 86 (6-117)   | 0.382|
| ACER              | 79 (54-97)| 84 (58-98)   | 0.01 |
| MMSE              | 27 (24-30)| 29 (24-30)   | 0.094|

Table 3: Differences in cognitive scores between the stroke group and
AD group; Significance of Mann-Whitney U test for comparison of
AD and stroke group median of difference before and after the therapy
(maximum decline and maximum elevation)

| Cognitive domains | AD | Stroke |
|-------------------|----|--------|
|                   | (n=15) | (n=21) |
|                   | Before | After | p    |
| IQc               | 1.5 (-15 ... 10) | 7.0 (-3 ... 16) | 0.001 |
| IQv               | -0.5 (-17 ... 10) | 3.0 (-56 ... 25) | 0.08 |
| IQp               | -4.5 (-14 ... 9)  | 6.0 (-4 ... 20)  | 0.001 |
| VP                | 0 (-12 ... 75)   | 4.0 (-6 ... 31)  | 0.089 |
| PO                | 1.0 (-14 ... 77) | 8.0 (0 ... 28)   | 0.017 |
| WM                | 1.0 (-24 ... 77) | 0 (-12 ... 10)  | 0.66 |
| SOP               | 2.0 (-26 ... 92) | 10.0 (-2 ... 87) | 0.049 |
| ACER              | 4.0 (-7 ... 16)  | 3.0 (-11 ... 15)| 0.904 |
| MMSE              | 1.0 (-2 ... 5)   | 0 (-4 ... 5)    | 0.6  |
| HADS              | 0 (-10 ... 12)   | -2.0 (-9 ... 17)| 0.46 |

By contrast, only ACE-R scores were significantly increased
following rehabilitation in the AD group (median 75 vs 83, p = 0.008);
no significant improvements were recorded in any of the other tests applied.

Table 3 presents the differences in cognitive scores between the stroke group and the AD group. The Mann–Whitney U test demonstrated a statistically significant difference between the stroke and AD groups in changes in the parameters IQc (p = 0.001), IQv (p = 0.001), PO (p = 0.017), and SOP (p = 0.049).

Table 4 shows ANCOVA analysis of differences between the groups with age as a covariate. ANCOVA proves the significant difference between the groups only in one outcome parameter IQp (p = 0.015).

| Adjusted mean difference before-after | AD (n=15) | Stroke (n=21) | p   |
|--------------------------------------|----------|--------------|-----|
| IQc                                  | 0.002    | 5.311        | 0.052|
| IQv                                  | 1.39     | 2.221        | 0.747|
| IQp                                  | -1.294   | 6.757        | 0.015|
| VP                                   | 5.359    | 3.665        | 0.803|
| PO                                   | 5.688    | 9.091        | 0.636|
| PP                                   | 5.75     | -1.5         | 0.316|
| RZ                                   | 9.242    | 10.04        | 0.925|
| ACER                                 | 4.242    | 3.974        | 0.909|
| MMSE                                 | 0.889    | 0.504        | 0.678|
| HADS                                 | -0.046   | 0.601        | 0.904|

Table 4: ANCOVA analysis of differences between the groups with age as a covariate.

Spearman’s correlation analysis showed the significant relationship between age and differences in some cognitive scores only in the stroke group (IQc ρ = 0.601; IQv ρ = 0.565, PO ρ = 0.535). In the AD group, Spearman’s correlation analysis does not prove any relationship between age and any outcome variables.

We conclude that cognitive parameters were significantly improved by CR in the stroke group, whereas there was little significant change in the AD group.

Discussion

This study demonstrates that the beneficial effects of computer-assisted CR in patients with mild cognitive impairment are much more significant in stroke patients than in patients with similar mild cognitive impairment attributable to probable Alzheimer disease.

This result is supported by other studies. Systemic review and meta-analysis by Cha and Kim [16] indicated that the overall effect size of computer-assisted CR in patients with stroke is 0.54 (95% CI; 0.33–0.74). That can be interpreted as a medium effect size. By contrast, the beneficial effects of CR in AD patients are at best ‘mild’, [5,13] although no direct comparison between these groups has yet been performed [18]. In addition, previous studies found no significantly different effects of CR in the patients in the acute versus chronic stroke phases [14]. In our study we therefore enrolled stroke patients from a single period (3 months after stroke).

Our results confirm the hypothesis that computer-assisted CR is more effective in patients with acute structural brain lesions (stroke) than in neurodegenerative diffuse brain disease (AD). Because brain plasticity is probably crucial for successful CR, one might anticipate a greater effect of CR in patients with acute demarcated brain lesions (as in stroke) than in patients with a diffuse neurodegenerative disorder.

We also report some improvement of cognitive function in the AD group, but the improvement was only seen in one test (ACE-R) and the extent of improvement was limited. Generally, the beneficial effects of CR were not observed using MMSE; we surmise that MMSE is unsuitable for screening patients before CR, and ACE-R is likely to be a more appropriate screening test in this context.

Study limitations: First, heterogeneity, in our study we included patients after stroke whose lesions were located variably (11 patients with lesions in the left hemisphere, 7 patients with lesions in the both hemispheres, and 3 with lesions in the right hemisphere).

Second, owing to the small number of patients in our study it was not possible to analyze cognitive subgroups such as memory, thinking operations, executive functions, and orientation. Third, there was a significant difference in age between the groups, but we excluded the relationship between age and other baseline variables in both groups.

ANCOVA, age as a covariate, showed the significant difference between the groups only in one outcome variable (IQ score global).

Despite the limitations of our study, statistical analysis provides preliminary evidence for the differential effects of CR in patients with mild cognitive impairment following stroke versus similar impairment attributed to probable AD. The effectiveness of CR in AD is unsatisfactory and there is a question of future aiming of the computer-assisted CR in AD patients. There is also no clear long-time effect of CR after stroke, and therefore we plan to retest our patients after one year.

Conclusion

We report that computer-assisted CR is likely to provide greater cognitive benefits in patients with mild cognitive impairment after stroke than in patients with similar impairment attributable to AD. For screening of patients before CR, ACE-R testing may be superior to MMSE. Further studies are necessary to evaluate the generalizability of our results.

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