Cerebral perfusional effects of 1-year rivastigmine treatment in Alzheimer disease: a case report

Effetti perfusionali cerebrali dopo trattamento di 1 anno con rivastigmina nella malattia di Alzheimer: un caso clinico

EVARISTO ETTORRE1, FRANCESCO SAVERIO BERSANI2, FABIOLA COLELLA1, ADRIANA SERVELLO1, AMEDEO MINICHINO2, VALENTINA MEGNA3, MAURO LIBERATORE3, MASSIMO BIONDI2, VINCENZO MARIGLIANO1

E-mail: bersani.fs@gmail.com

1 Department of Cardiovascular, Respiratory, Nefrologic, Anaesthesiologic and Geriatric Sciences, Sapienza University of Rome, Italy
2 Department of Neurology and Psychiatry, Sapienza University of Rome, Italy
3 Department of Radiological, Oncological and Anatomopathological Sciences, Sapienza University of Rome, Italy

SUMMARY. It is described the case of a 74-years-old woman with probable Alzheimer Disease who showed good clinical response to rivastigmine associated with relevant improvement of cerebral perfusion after 1 year of treatment. The single-photon emission computed tomography (SPECT) scan showed a significant improvement in cortical uptake of the tracer in temporo-parietal and frontal regions in comparison to the examination performed before the treatment.

KEY WORDS: rivastigmine, SPECT, Alzheimer’s disease, psychopharmacology, acetylcholinesterase inhibitors.

RIASSUNTO. Viene descritto il caso di una donna di 74 anni con probabile malattia di Alzheimer che presenta una buona risposta clinica alla rivastigmina associata a rilevante miglioramento di perfusione cerebrale dopo 1 anno di trattamento. La tomografia a emissione di fotone singolo (SPECT) mostra un significativo miglioramento nella captazione corticale del tracciante delle regioni temporo-parietali e frontali rispetto all’esame eseguito prima del trattamento.

PAROLE CHIAVE: rivastigmina, SPECT, malattia di Alzheimer’s, psicofarmacologia, inibitori dell’acetilcolinesterasi.

INTRODUCTION

Alzheimer’s disease (AD) is a progressive neurodegenerative condition that predominantly affects the elderly population; it affects about 3 million people in the European Union, and it is predicted to affect 1 in 85 people globally by 2050.1-4

The most strongly supported evidence available today on the pharmacological treatment of the cognitive symptoms of AD concerns the acetylcholinesterase (ChE) inhibitors, especially donepezil and rivastigmine, in addition to cognitive rehabilitation.5-10

Rivastigmine is a cholinesterase inhibitor that inhibits both AChE and butyrylcholinesterase (BuChE). The efficacy of oral rivastigmine has been demonstrated in clinical trials involving more than 3000 AD patients and 500 patients with Parkinson’s dementia.10 Rivastigmine is a small molecule (<400 Da), and is both lipophilic and hydrophilic; it passes easily through the skin into the bloodstream as well as through the blood-brain barrier, making it well-suited to transdermal delivery.17

From a functional point of view, AD is characterized by hypometabolism/hypoperfusion of several different brain areas: bilateral posterior cingulate gyri, bilateral superior and middle frontal gyri, left inferior parietal lobe and medial temporal lobe.10,19 In particular, temporo-parietal hypometabolism on positron emission tomography (PET) is now considered to be one of the three diagnostic criteria for progression to AD in patients with mild cognitive impairment, along with medial temporal lobe atrophy on magnetic resonance imaging (MRI) and tau/amyloid- in cerebrospinal fluid (CSF).20-22

Despite the well-known efficacy of rivastigmine in the treatment of cognitive symptoms of AD, a very small number of studies investigated its capacity of modulating regional cerebral blood flow (rCBF), showing contrasting results.23-29

Later we describe the case of a 74-years-old woman with probable AD who presented good clinical response to rivastigmine associated with relevant improvement of cerebral perfusion after 1 year of treatment.

CASE REPORT

Ms M. is a 74-years-old woman with probable AD (according to NINCDS-ADRDA Criteria).30 She worked as housekeeper; she...
Cerebral perfusional effects of 1-year rivastigmine treatment in Alzheimer disease: a case report

Figura 1. Single photon emission computed tomography (SPECT) of the brain with 99mTc-HMPAO in a patient affected by Alzheimer’s Disease. Sagittal (A, B) and transaxial (C, D) reconstruction before (A, C) and after treatment (B, D). A significant improvement of the cerebral uptake of the radiopharmaceutical after treatment can be seen.

lived with her husband and had three sons. She had clinically controlled hypertension and hypothyroidism (post thyroidectomy).

At the first evaluation (August 2011) Ms. M complained amnesia problems and sleep disturbances gradually occurred in the previous 12 months.

Mini Mental Score Evaluation (MMSE) score was 21; the most affected cognitive areas were recall, attention and calculation and orientation to time.

Ms. M. experienced a conflicting relation with her husband; slight depressive and anxious symptoms were also present (Geri-atric Depression Scale - GDS: 10; Neuropsychiatric Inventory - NPI: 9).

Some basic daily life activities were compromised: she needed help to wash herself, to take medicines and to use money.

Laboratory tests were within the normal ranges with the exception of total cholesterol: 259 mg/dL, low-density lipoprotein (LDL): 157 mg/dL, triglycerides: 205 mg/dL and thyroid-stimulating hormone (TSH): 4.30 mU/L.
DISCUSSION

The present clinical case suggests a relationship between the increase of perfusion patterns in temporoparietal and frontal regions and the improvement of cognitive and behavioural symptoms in a patient with probable AD under rivastigmine treatment.

Previous studies that investigated this issue showed contrasting results. Three studies on AD patients reported linear correlations between the improvement of cognitive and behavioural symptoms and perfusion pattern in right cingulate, frontal, parieto-temporal regions bilaterally after 3-6 months of ChE inhibitors therapy. On the other hand, Nobili et al. and Nakano et al. reported that rCBF in critical areas was maintained or decreased after 6-18 months of treatment with donepezil or rivastigmine. The progressive degenerative course of AD is usually associated with progressive reduction of cerebral perfusion in those brain areas involved in its etiopathogenesis. The different outcomes between above mentioned studies could be partially related to the different intervals between baseline and follow-up neuroimaging assessments. In fact, it is possible that the improvement of cerebral perfusion induced by ChE inhibitors therapy may be affected by the natural progression of the disease, explaining the absence of perfusion improvements in those studies with a longer follow-up period.

To our knowledge, the clinical history of Ms. M. represents the only case of cognitive and behavioural improvements associated with relevant increase of cerebral perfusion after 1 year of rivastigmine treatment. Differently from those studies that already observed a correlation between increased cognitive functions and perfusion patterns, in our case the time interval between the two SPECT evaluations was long enough to potentially observe the progressive deterioration of the disease.

A possible relation between paroxetine and rivastigmine was present: it is known that paroxetine may improve cognitive functions and that rivastigmine may show antidepressant efficacy. In our case the concomitant treatment with paroxetine should not represent a confounding factor in relation to brain perfusion as it has been shown that paroxetine does not significantly affect rCBF.

It is possible that the improvements in cognitive performances after rivastigmine therapy are related to increased perfusion in the temporoparietal and prefrontal regions (i.e. the so-called “cognitive network”), while the behavioural improvements are related to increased perfusion in the dorsolateral frontal and cingulate regions (i.e. the so-called “limbic network”).

REFERENCES

Cerebral perfusional effects of 1-year rivastigmine treatment in Alzheimer disease: a case report


Riv Psichiatri 2015; 50(4): 188-191