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CASE SERIES

Cognitive effects of a dietary supplement made from extract of Bacopa monnieri, astaxanthin, phosphatidylserine, and vitamin E in subjects with mild cognitive impairment: a noncomparative, exploratory clinical study

Danilo Zanotta Silvana Puricelli Guido Bonoldi

Unità Operativa di Medicina 2. Ospedale di Circolo di Busto Arsizio, Varese, Italy

Correspondence: Danilo Zanotta Unità Operativa di Medicina 2, Ospedale di Circolo di Busto Arsizio, Piazzale G Solaro 3, Busto Arsizio 21052, Varese, Italy Tel +39 0331 699 269 Fax +39 0331 699 370 Email zanottadanilo@gmail.com

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http://dx.doi.org/10.2147/NDT.\$51092

Abstract: A prospective cohort, noncomparative, multicenter trial was conducted to explore the potential of a phytotherapeutic compound, available as a dietary supplement and containing extracts of Bacopa monnieri and Haematococcus pluvialis (astaxanthin) plus phosphatidylserine and vitamin E, in improving cognition in subjects diagnosed with mild cognitive impairment. Enrolled subjects (n=104) were aged 71.2±9.9 years and had a mini-mental state examination score of 26.0 ± 2.0 (mean \pm standard deviation). They underwent the Alzheimer's Disease Assessment Scale-cognitive subscale (ADAS-cog) test and the clock drawing test at baseline and upon completion of a 60-day period of dietary supplementation with one tablet daily of the tested compound. In 102 assessable subjects, total ADAS-cog scores improved from 13.7±5.8 at baseline to 9.7 ± 4.9 on day 60, and the clock drawing test scores improved from 8.5 ± 2.3 to 9.1 \pm 1.9. Both changes were statistically significant (P<0.001). Memory tasks were the individual components of ADAS-cog showing the largest improvements. In a multivariate analysis, larger improvements in total ADAS-cog score were associated with less compromised baseline mini-mental state examination scores. Perceived efficacy was rated as excellent or good by 62% of study subjects. The tested compound was well tolerated; one nonserious adverse event was reported in the overall study population, and perceived tolerability was rated excellent or good by 99% of the subjects. In conclusion, dietary supplementation with the tested compound shows potential for counteracting cognitive impairment in subjects with mild cognitive impairment and warrants further investigation in adequately controlled, longer-term studies.

Keywords: mild cognitive impairment, Bacopa monnieri, astaxanthin, ADAS-cog test, clock drawing test, dietary supplement

Introduction

Most people experience gradual cognitive decline as they age. Some of them are suffering from what is commonly known as mild cognitive impairment (MCI), which is a more serious reduction in mental efficiency than the physiological agerelated decline but does not meet the criteria for a diagnosis of senile dementia or Alzheimer's disease.1-4

The estimated prevalence of MCI is between 10%-20% of people aged over 65,5-10 and the causes of MCI are currently unknown. As with more severe forms of dementia, it is believed that MCI origin is complex, and that it involves factors that are toxic to nerve

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cells, such as β -amyloid protein, Tau protein and Lewy bodies (although these same pathological markers have also been found in elderly asymptomatic individuals), or alterations in cerebral circulation such as localized microstrokes.¹¹

Several longitudinal studies have shown that the majority of people affected by MCI have a substantially increased risk of developing dementia.^{5–7,9} Whereas the incidence of dementia in the general population of the United States is 1%–2% per year, incidence in the MCI population reaches 5%–10% per year¹² and 10%–15% in the subjects who are followed in specialist clinics.¹³

Therefore early recognition of MCI, leading to early intervention, might allow for preventing or delaying the onset of dementia and prolonging the active life of these subjects. However, despite some data suggesting that the rate of reversion of MCI patients to a normal cognitive level could be between 25%–30%, prospective studies have shown lower rates.⁸

The main objective of the present study was to explore the potential of a phytotherapeutic compound, commercially available as dietary supplement and containing extracts of *Bacopa monnieri* and *Haematococcus pluvialis* (astaxanthin) plus phosphatidylserine and vitamin E, in improving cognition in subjects diagnosed with MCI. A secondary objective was to assess the tolerability and safety of use of the compound.

Bacopa monnieri is a medicinal plant species traditionally used as a brain tonic. Studies conducted in animal models suggest that bacoside constituents of *Bacopa monnieri* have an antioxidant effect on the hippocampus, frontal cortex, and the striatum.^{14–17} A systematic review of several randomized, double-blind clinical studies conducted with *Bacopa* has demonstrated efficacy in improving memory and some cognitive functions.¹⁸

Astaxanthin is a carotenoid that is positioned within the cell membranes and the circulating lipoproteins and is not produced by human metabolism. It plays an antiinflammatory and antioxidant role¹⁹⁻²¹ and protects the plasma membrane and the double mitochondrial membrane.²² In a 12-week open study conducted in healthy subjects aged 50 to 69 years, oral administration of astaxanthin led to an improvement in some cognitive functions, such as reaction time, attention, and working memory.²³

Phosphatidylserine is a phospholipid that becomes a key constituent of the cell membrane.²⁴ It is found in high concentrations in the cerebral tissue, where it supports cellular functions such as maintenance of integrity of mitochondrial membranes for the production of energy, electrical depolarization of neuronal membranes, release of neurotransmitters in the synaptic cleft, postsynaptic receptor activity, and activation of protein kinase C.^{25–30} Vitamin E has antioxidant properties: it prevents the spread of oxidation of the polyunsaturated fatty acids by scavenging the lipid peroxyl radicals. The potential therapeutic or preventive role of vitamin E in brain aging and age-related cognitive disorders is still a subject of debate.³¹ While large-scale clinical trials have failed to demonstrate major cognitive benefits, vitamin E deficiency has been consistently found to be associated with Alzheimer's disease and MCI in recent human studies.³¹

Materials and methods Tested compound

The tested compound is available as a dietary supplement in a tablet formulation (Illumina[®]) and was supplied free of charge by Cristalfarma (Milan, Italy) for use in the study. Its composition is shown in Table 1.

Subjects

Subjects aged \geq 50 years referred to outpatient clinics of the participating centers due to forgetfulness, disorientation, difficulty in concentrating or other cognitive difficulties, and diagnosed with MCI were eligible for the study. MCI diagnosis was based on a mini-mental state examination³² (MMSE) score \geq 22 and <28 and on caregiver testimony on cognitive symptoms.

Subjects with known current psychiatric disorders or organic diseases that could interfere with the cognitive status were excluded, as were those undergoing treatment with any psychotropic drug.

All enrolled subjects provided written informed consent for participation in the study.

Study design

This was a prospective cohort, noncomparative, multicenter trial conducted at seven Italian outpatient clinics for agerelated cognitive disorders.

Baseline assessments included medical history, review of current therapies, physical examination, geriatric assessment, and MMSE, followed by the administration of the Alzheimer's

Table 1 Oualitative and duantitative composition of Illumi

Components	Content	
	per tablet	
Microalgae (Haematococcus pluvialis) dry extract 3%	74 mg	
Astaxanthin	2 mg	
Bacopa monnieri dry extract 20%	100 mg	
Bacosides	20 mg	
Phosphatidylserine (extracted from soy)	30 mg	
Vitamin E (vegetable origin)	30 mg	
Note: Illumina [®] (Cristalfarma, Milan, Italy)		

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Disease Assessment Scale-cognitive subscale (ADAS-cog) test³³ and the clock drawing test (CDT).³⁴ Subjects were instructed in the presence of the respective caregivers to take one tablet of Illumina per day immediately before their main meal, for 60 days. Instructions and contact details were provided to subjects and caregivers should the need arise to report to investigators any adverse events occurring over the treatment period.

On day 60±3, subjects were reassessed for intercurrent medical history, including adverse events and therapy changes. ADAS-cog and CDT were readministered by the same investigating physician. Subjects were asked to rate the perceived efficacy and tolerability of the treatment on 5-point scales. Each investigating physician could record free-text comments on individual treatment outcome.

Neuropsychological tests

The MMSE has a total score of 0–30 points. It was given to assess baseline cognitive status and to exclude the presence of any intellectual deficit. The lower limit for inclusion (22 points) was intended to exclude cognitive deterioration due to Alzheimer's disease or other forms of dementia.

ADAS-cog and CDT were given to assess cognitive skills. CDT examines the subject's planning ability and constructive skills, whereas ADAS-cog assesses specific characteristics of cognitive efficiency that are sensitive to deterioration due to primary degenerative dementia. ADAS-cog includes eleven tasks, of which nine are related to functional assessment and two are objective memory tests.

Statistics

Univariate analysis of day 60 versus baseline changes in ADAS-cog and CDT scores was performed using the Student's *t*-test for paired data. Multivariate analysis of variance (MANOVA) for paired data was used to explore relationships of ADAS-cog and CDT score changes to baseline variables such as sex, age, MMSE, family history of cognitive disorders, and concomitant diseases. Data analysis was performed using the Statistical Package for Social Sciences (SPSS) version 20.0 (IBM Corporation, Armonk, NY, USA). The significance threshold was set at P<0.05. Data are reported as means ± standard deviation unless noted otherwise.

Results

A total of 104 subjects were enrolled, 48 men and 56 women with mean age of 71.2 ± 9.9 years. Baseline MMSE score was 26.0 ± 2.0 . A family history of cognitive disorders was present in 14 subjects (13.5%). Among concomitant diseases recorded at baseline, those with a prevalence ≥ 3 subjects included hypertension (n=29), bone and joint disorders (n=9), allergy to drugs or foods or environmental agents (n=8), respiratory disorders (n=4), and diabetes mellitus (n=3). Concomitant use of drugs possibly affecting cognitive status was not reported for any subject at either baseline or over the 60-day observation period.

All 104 subjects were assessed for safety. As two subjects were lost to observation (one of them following a nonserious adverse event, as detailed below), 102 were included in the univariate analysis of efficacy. Of those, 98 had complete information on baseline variables and were thus included in the multivariate analysis.

Changes in total ADAS-cog score and CDT are shown in Table 2. ADAS-cog went from 13.7 \pm 5.8 at baseline to 9.7 \pm 4.9 on day 60, a statistically significant improvement according to univariate analysis (*P*<0.001). A significant improvement was also found in the CDT score, from 8.5 \pm 2.3 to 9.1 \pm 1.9 (*P*<0.001).

Changes in individual components of ADAS-cog are shown in Figure 1. There was a statistically significant improvement in each of the eleven items ($P \le 0.02$). The largest improvements were observed in memory tasks ("word recall" and "word recognition").

Multivariate analysis of total ADAS-cog score showed a statistically significant interaction between treatment effect and baseline MMSE (P=0.009), in the sense that larger ADAS-cog improvements were associated with less compromised MMSE scores. Otherwise, changes in ADAS-cog individual components and in the CDT score did not appear to be significantly influenced by any of the baseline variables included in the multivariate model.

Only one adverse event was recorded, consisting of gastric disturbances in a subject who was taking concomitant oral corticosteroids. This led to discontinuation of Illumina upon the subject's decision.

Assessments of the perceived efficacy and tolerability of treatment were obtained from 93 and 96 subjects, respectively. Efficacy was rated as excellent by 16% of subjects, good by

Table 2 ADAS-cog and clock drawing test scores before and after treatment with Illumina $^{\scriptscriptstyle (\! 8\!)}$

Test administered	Time	N	Mean	Standard deviation	t-test	P-value
ADAS-cog	D0	102	13.65	5.78	11.177	<0.001
	D60	102	9.65	4.89		
Clock drawing	D0	102	8.46	2.29	-4.521	< 0.001
test	D60	102	9.07	1.92		

Note: Illumina® (Cristalfarma, Milan, Italy).

Abbreviations: ADAS-cog, Alzheimer's Disease Assessment Scale-cognitive subscale; D0, day 0; D60, day 60.



Figure I Mean ± standard deviation scores of individual ADAS-cog components before and after treatment with Illumina[®]. Note: Illumina[®] (Cristalfarma, Milan, Italy). Abbreviations: ADAS-cog, Alzheimer's Disease Assessment Scale-cognitive subscale; D0, day 0; D60, day 60.

46%, fair by 28%, and poor or of no use by the remaining 10%. Tolerability was rated as excellent by 67% of the subjects, good by 32%, and poor by 1%.

Specific comments on treatment outcome were recorded by the investigating physicians for 17 subjects. All of the comments highlighted improvements in cognitive performance and/or, interestingly, in the quality or duration of night-time sleep.

Discussion

In this prospective cohort, noncomparative multicenter trial, short-term (60-day) dietary supplementation with Illumina, a phytotherapeutic compound containing extracts of *Bacopa monnieri* and *Haematococcus pluvialis* (astaxanthin), soy-extracted phosphatidylserine, and vitamin E of vegetable origin, was associated with an improvement in cognitive and mnemonic skills in subjects diagnosed with MCI. Cognitive and mnemonic performance was assessed with validated instruments (ADAS-cog and CDT) having predictive value for the risk of MCI progression to Alzheimer's disease.³⁵ The observed 60-day improvements in ADAS-cog and CDT were statistically significant as compared with baseline values and were borne out by subject perception of treatment efficacy, which was rated as "excellent" or "good" in 62% of the cases.

The consumption of Illumina over 60 days was extremely well tolerated, as highlighted by rarity of the reported adverse events (one case out of 104 subjects) and by subject perception of compound tolerability ("excellent" or "good" in 99% of the cases).

These promising findings are of particular interest and relevance when considered in the context of MCI, a condition still posing a largely unmet medical need due to its prevalence in the elderly population, high associated risk for progression to Alzheimer's disease, and current lack of effective therapeutic or preventive measures.³⁶

A basic limitation of the present study was the lack of a control treatment arm with consequent openness to selection bias and placebo effect. Thus the cognitive benefits observed with Illumina, while looking consistent with earlier observations made with other *Bacopa monnieri* or astaxanthin compounds,^{14–23} should be regarded as exploratory (hypothesis-raising). On the other hand, efficacy data were likely exempt from attrition bias, as the main analyses of ADAS-cog and CDT accounted for 98% of the enrolled subjects (102 out of 104). Other study limitations include a period of observation which may have been too short to adequately reflect the natural course of MCI and the lack of assessment of compliance to the tested dietary supplement.

Conclusion

The results of this study show that dietary supplementation with Illumina has potential for counteracting cognitive impairment in subjects with MCI. Further investigation of this compound in adequately controlled, longer-term studies is warranted to formally assess its cognitive effects in mild or moderate mnemonic-cognitive disorders.

Acknowledgments

In addition to the authors, the following clinical investigators, all in Italy, collected data for this study: Orazio D'Alessio, consulting geriatrician, Hospital in Vasto; M Letizia Del Re, Psychogeriatrics Clinic and Alzheimer's Centre, Hospital in Pescara; Alberto Edalli, Raimondi Foundation, Gorla Minore; Paolo Genoni, "Scuola di Medicina Carnaghi e Brusatori" Foundation, Busto Arsizio; Walter Polinelli, DEWA Srl Milan; Fabio Matascioli, psychologist and psychotherapist, San Giorgio a Cremano. Statistical analysis was performed by Stella Forti, freelance consultant in medical statistics, Milan, Italy. The study was supported in part by an unrestricted grant from Cristalfarma, Milan, Italy.

Disclosure

The authors report no conflicts of interest in this study.

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