Healthy Nutrition, Phytonutrients and Alzheimer’s Disease

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Introduction

In most areas of the world malnutrition in early life is directly or indirectly responsible of more deaths in children than other causes. Malnutrition is also a main concern in aged population. Nutritional deficiencies affect physical growth but also may produce irreversible mental and emotional changes. Many of these undesirable effects have been studied in animals [1]. Nutrition is of growing importance in degenerative diseases. Nilsson and coworkers have published that episodic memory performance i.e. the ability to recall events in time and place declines linearly with age. This decline in episodic memory began as early as at 20 years of age [2]. Thus, studies about nutrition influences in cognition should begin at early ages (20 years or even less). Recent investigations in animals, on the physical and biochemical aging of the central nervous system, have provided helpful information to understand the effects of malnutrition. During maturation and growth, the amount of water in the brain gradually decreases, whereas the amount of cholesterol steadily increases [3, 4]. Maturation consists of a rapid increase in cells, as indicated by the increased concentrations phosphorus of DNA. Whereas growth parallels myelination and is represented by increased amounts of cholesterol [4,5]. Inadequate nutrition in calories and proteins, coinciding with the period in life in which the brain is growing most rapidly, may result in a smaller (than in controls) brain at maturity. Also, in a brain which matures biochemically and functionally at a slower rate [5]. In the adult rat starvation does not result in significant changes in brain weight or in damages to neural tissue. However, malnutrition associated -oxidative stress and -inflammation occur in neurons and astrocytes in elderly brains which may damage differently neural tissues in a differential fashion [6]. A biological mechanism which can be linked to nutrition and have been associated with aging includes oxidative stress [7], inflammation [8], homocysteine [9], advanced glycation end products [10] and the provision of fatty acids [11]. These mechanisms have in common that they reflect the nature of the existing diet or eventually respond to changes in what we eat. Changes from normal cognition to impaired, glucose tolerance, or related patterns may prove to be useful endpoints [12].

Dementia is not a disease but a group of symptoms which are common to different diseases, such as problems affecting, memory, and reasoning. Alzheimer’s disease (AD) accounts for 50% to 70% of all dementia cases, and vascular dementia accounts for 10-15%, whereas the rest includes about 50 other etiologies, so nutrition should represent in common to different diseases, such as problems affecting, memory, and reasoning. Alzheimer’s disease (AD) accounts for 50% to 70% of all dementia cases, and vascular dementia accounts for 10-15%, whereas the rest includes about 50 other etiologies, so nutrition should be expected to have an impact in different ways [13,14]. The role of nutrition in AD’s disease has not been studied and, phytonutrients in particular appear a very interesting subject for their health benefits [15]. Natural Polyphenols are present in many fruits and vegetables in particular appear a very interesting subject for their health benefits [15]. Among polyphenols, flavonoids represent the largest group. For instance the major phytoestrogen component of soy is genistein. We have found that which appears effective in preventing neuronal death caused by Aβ peptide in neurons in primary culture [17]. In fact the beneficial effects of phytoestrogens on central nervous system functions have received attention in the last decade, and estrogen therapy is one of the potential strategies for the prevention of dementia [18]. Genistein has immunosuppressive and anti-inflammatory properties [19,20]. Data from our laboratory have shown that estradiol or genistein abolishes neuronal reactive oxygen species and attenuates Aβ-induced cell death in cortical neurons by preventing p38 activation and mitochondrial aggregation [17]. Genistein, at nutritionally relevant concentrations, can reproduce the protective effects of estradiol in neurons. In astrocytes, genistein is able to prevent AD-associated inflammation by increasing PPARy expression, thus suggesting potential anti-inflammatory properties against AD [21].

Other studies have shown that excessive alcohol uptake in rats can result in mitochondrial dysfunction in neurons and astrocytes leading to neurodegeneration [22, 23]. Paradoxically, it is reported that moderate alcohol uptake associates with a lower risk of clinical stroke [24].

Researchers have shown a relationship between alcoholism and AD. Moderate alcohol intake produced benefits to AD patients compared to those who did not drink alcohol [25]. Grapes contain flavonoids such as resveratrol or its dimethylated analog pterostilbene, used as antifungal agents, anti-inflammatory drugs, antioxidants, and anti-inflammatory agents [26]. Thus the beneficial effects of wine consumption on neurodegeneration are attributed to this type of molecules [27,28]. Resveratrol was reported to reduce Aβ production in the HEK293 cell line expressing wild type or Swedish-mutant APP695 and Aβ damage mechanisms and the role of the proteosome in the clearance of Aβ have been determined [29]. Therefore, a possible proteasome-dependent anti-amyloidogenic activity of natural polyphenols, such as resveratrol or pterostilbene appears an interesting (nutrition-related) next step in AD research.

Interestingly the prevalence of AD in Indian people 70 to 79 years old is 4.4 fold less than in a similar population of the United States [30]. The Indian diet is rich in spices such as chili, curry or turmeric. Curcumin in particular is a potent free radical scavenger, better than vitamin E, which can provide effective protection against lipid peroxidation. Curcumin is also effective against the amyloid plaques in animal models. Thus, it appears be reasonable to speculate that curcumin could possibly prevent the onset of AD associated with oxidative stress [31].

In the United States, South American Indians knew the Stevia plant as “medicine for women” and use it in menopause an also to calm pain during menstruation. Probably its magnesium combats the cramps and its seven flavonoids (with estrogen-like effects) can ameliorate menopause symptoms [32]. Stevia has an additional property useful in alimentation, since it does not alter glucose blood levels but simultaneously satisfies appetite for something sweet. In animal experiments, stevia prevents atherosclerosis because increases superoxide dismutase enzyme
activists, thus removing oxidized LDL cholesterol and other harmful lipids from blood vessels wall [33]. The preventive effects of steviosides on the development of AD and its relationship with soy-based diet surely deserve further studies.

References
