Novel Role of Red Wine-Derived Polyphenols in the Prevention of Alzheimer’s Disease Dementia and Brain Pathology: Experimental Approaches and Clinical Implications

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Abstract

Recent studies suggest that by the middle of this century, as many as 14 million Americans will have Alzheimer’s disease, creating an enormous strain on families, the health care system and the federal budget. There are still widespread misconceptions about issues related to the prevention and/or treatment of disease pathogenesis, leaving us unprepared to deal with the disease. To address these challenges, several therapeutic approaches are currently under investigation, mainly in an attempt to delay disease onset and eventually slow down its progression. Recent epidemiological evidence has implicated the protective role of dietary polyphenols from grape products against Alzheimer’s disease. Furthermore, experimental evidence supports the hypothesis that certain bioactive grape-derived polyphenols may protect against Alzheimer’s disease-type cognitive deterioration, in part by interfering with the generation and assembly of β-amyloid peptides into neurotoxic oligomeric aggregated species. Brain-targeting polyphenols have been shown to significantly reduce the generation of β-amyloid peptides in primary cortico-hippocampal neuron cultures, and preliminary results indicate that they may influence neuronal synaptic plasticity. Recent evidence has also implicated the role of certain grape-derived preparations in beneficially modulating tau neuropathology, including reducing tau aggregation. Studies suggest that dietary polyphenolics may benefit Alzheimer’s disease by modulating multiple disease-modifying modalities, both β-amyloid-dependent and independent mechanisms, and provide impetus for the development of polyphenolic compounds for Alzheimer’s disease prevention and/or therapy.

Introduction

Alzheimer’s disease (AD) is the most common type of dementia in the United States. Victims of AD commonly display a loss of memory, inability to learn new things, loss of language function, deranged perception of space, inability to do calculations, depression, delusions, and other cognitive deficits. AD is ultimately fatal within 5 to 10 years of its onset. Approximately 5 million people in the United States currently have AD [1], with an estimated cost to society of more than $100 billion per year. The estimated cost of dealing with AD over the next 40 years is twenty trillion dollars. Up to 14 million people in the United States are projected to be affected by AD by the middle of this century [1]. To date, there is no cure for AD. The few agents that are approved by the FDA for the treatment of AD have only modest efficacy in terms of modifying clinical symptoms, and none appear to affect disease progression or prevention [2]. Scientists are continually exploring novel avenues for preventing or treating this condition.

Alzheimer’s Disease Neuropathological Features: Implications for Therapeutic Developments

While the classification and diagnosis of AD is based on the cognitive behavior of an individual, the roots of the disease lie in the neurological pathology of its victims [3]. The two defining neuropathological features of AD are abnormal aggregation and deposition of certain toxic peptide fragments known as β-amyloid (Aβ) peptides or tau protein in the brain as, respectively, extracellular neuritic plaques (NP) and intracellular neurofibrillary tangles (NFT) [4]. Aβ peptides are derived from the ubiquitous amyloid precursor protein

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(APP) through amyloidogenic processing by β- and γ-secretases, rather than through non-amyloidogenic cleavage by α-secretase. In humans, genetic mutations leading to Aβ neuropathology in at least one of three genes, namely APP, presenilin 1 and presenilin 2, are causally linked to early onset AD cases and are associated with AD dementia [5, 6]. In experimental animal models, these same mutations also accelerate Aβ deposition and cognitive deterioration [7]. Based on this, major efforts are focused on developing pharmacological strategies that delay the initiation and/or slow the progression of Aβ-mediated neuropathological responses. Recent evidence from experimental AD mouse models indicates that the accumulation of soluble high-molecular weight oligomeric Aβ species in the brain, rather than deposition of NP per se, may be specifically related to spatial memory reference deficits [8–12].

Despite strong genetic data arguing that Aβ neuropathology is sufficient to cause AD [13], progressive cognitive decline and neuron and synapse loss in AD are best correlated with tau neuropathology [14]. In the AD brain, tau proteins, particularly hyperphosphorylated tau, are found aggregated into progressively larger polymeric species that are ultimately deposited as insoluble NFTs [15]. NFTs themselves are not necessarily the tau species inducing neurotoxicity [16, 17]. A predominant theory of tau-mediated neurodegeneration is based on a “toxic gain of function” model, in which abnormally phosphorylated tau promotes sequestration of both hyperphosphorylated and normal tau from microtubules, leading to microtubule instability and alterations of microtubule-mediated processes, including abnormalities in axon transport, among others [18].

These considerations strongly suggest that reducing the accumulation of soluble oligomeric Aβ peptides and tau species in the brain, as opposed to dissociating or preventing NP and/or NFT formation or their depositions, may be a more productive approach to AD therapy. As discussed in more detail below, we recently demonstrated for the first time that dietary supplementation with red wines equivalent to moderate wine consumption in humans effectively attenuates the development of Aβ-mediated neuropathology and cognitive dysfunction in a mouse model of AD. Moreover, our evidence demonstrates that the grape-derived polyphenolics commonly found in red wines may also modulate tau-mediated neuropathology responses (Fig. 1).
Red Wine-Derived Polyphenols and Alzheimer’s Disease

While genetic factors are highly relevant in early-onset AD cases, their significance diminishes in late-onset sporadic AD cases, which is the most common form of AD [2]. Nongenetic factors, including modifiable lifestyle and dietary regimens such as moderate consumption of alcoholic beverages, are receiving increasing attention in AD research, especially in light of the recent epidemiological studies indicating that moderate wine consumption may influence the relative risk for AD clinical dementia [19]. Little is known about the beneficial role of red wine in AD dementia onset. The neuroprotective efficacy of red wine is typically attributed to the antioxidant activities of polyphenols in the wine. To explore how red wines might benefit AD, we tested whether dietary supplementation with red wines may beneficially modulate AD phenotypes in the Tg2576 AD mouse model [20]. Recapitulating select features of AD, Tg2576 mice are characterized by progressive development of Aβ neuropathology and cognitive decline with increasing age.

Potential Benefits of Moderate Consumption of Red Wine and Other Dietary Grape Polyphenol Products in Alzheimer’s Disease

Polyphenols are members of a very large family of plant-derived compounds containing one or more phenolic group. Thousands of polyphenols have been identified to date, including bioflavono-
ploring the potential beneficial role of grape-derived polyphenols in AD should be conducted using specific metabolites that are accumulated in the brain. For example, we have recently identified selected epicatechin glucuronide derivatives from a grape seed polyphenolic extract that are capable of penetrating the brain [34] and potentially attenuating AD, in part by promoting neuroplasticity processes [40]. The structure of this bioactive epicatechin glucuronide as well as the structure of epicatechin, its parent compound, is illustrated in Fig. 3.

Consistent with epidemiological evidence implicating the protective role of dietary polyphenols from grape products against AD [19], our preclinical studies have demonstrated that dietary supplementation with a grape seed polyphenol extract [41], or moderate consumption of red wines [42, 43] containing high contents of grape polyphenols, is effective in attenuating the onset and progression of Aβ-mediated AD-type neuropathology and cognitive deterioration in transgenic mouse models of AD. As schematically shown in Fig. 1A, Aβ peptides are derived from the ubiquitous protein APP through amyloidogenic processing by β- and γ-secretases rather than through non-amyloidogenic cleavage by α-secretase. Monomeric Aβ peptides (e.g., Aβ1–40 or Aβ1–42 peptides) tend to assemble into soluble, high-molecular weight neurotoxic Aβ aggregates that are key contributory factors to AD dementia. Our evidence has demonstrated that polyphenolic components from grape products may protect against AD dementia, in part by reducing Aβ-mediated neurotoxic mechanisms. Mechanistically, we found that grape-derived polyphenols may modulate Aβ toxicity by either reducing the generation of Aβ peptides from the amyloid precursor protein and/or by interfering with the assembly of Aβ peptides into high-molecular weight neurotoxic Aβ aggregated species (Fig. 1A). For example, using a PICUP assay [44] we demonstrated that polyphenolic components from a red Muscadine wine, made from Vitis rotundifolia (Vitaceae), significantly interfere with Aβ protein-to-protein interactions critical for the initial assembly of monomeric Aβ peptides into increasingly large aggregated species (Fig. 1B). In another example, we showed that treatment with a polyphenolic extract from another red wine (Cabernet Sauvignon) significantly reduced the generation of Aβ1–40 (Fig. 1C, left panel) and Aβ1–42 (Fig. 1C, right panel) in a dose-dependent manner in primary cortico-hippocampal neuron cultures generated from the Tg2576 transgenic AD mouse model. As an example illustrating the efficacy of dietary supplementation with grape products in protecting against cognitive deterioration, Fig. 1D shows that dietary supplementation with the Muscadine wine, equivalent to moderate consumption in humans, significantly attenuates the development of cognitive deterioration in the Tg2576 AD mouse model.

Collectively, we have studied two unrelated red wines, a Cabernet Sauvignon and a Muscadine wine, for their efficacy in modulating AD phenotypes in the Tg2576 transgenic AD mouse model [42, 43] and found that both wines are effective in attenuating Aβ-related mechanisms. Moreover, our evidence suggests that polyphenols from the two wines benefit Aβ-mediated neuropathology and cognitive dysfunction. Therefore, our evidence suggests that polyphenols from Cabernet Sauvignon wine are bioactive in terms of inhibiting Aβ generation. We subfractionated total polyphenolics from the red Cabernet Sauvignon wine into multiple preparations with different polyphenolic composition profiles. Shown are HPLC analysis of total polyphenolics from Cabernet Sauvignon (top panel) and a subfraction comprised primarily of anthocyanin components (bottom panel). Abbreviation: AC, anthocyanin. B Generation of Aβ1–40 peptide by primary cortico-hippocampal neuron cultures from Tg2576 mice in the presence of vehicle (control) or presence of total polyphenols from the wine or in the presence of the anthocyanin polyphenol subfraction.
identification of specific wines and other dietary products that might prove to be effective in AD prevention and therapy. Accumulating evidence suggests that resveratrol, a naturally occurring polyphenolic compound that is associated with beneficial effects on aging, metabolic disorders, inflammation and cancer in animal models [45] and that is found in varying concentrations in red wine and many food products, may enhance Aβ clearance by promoting intracellular proteosome activity in vitro [46]. However, the role of resveratrol in our study on Cabernet Sauvignon treatment in Tg2576 mice [42] is not clear since the Cabernet Sauvignon used had only 0.2 mg/L resveratrol, a concentration 10-fold lower than the minimal effective concentration shown to promote Aβ clearance in vitro [46]. To gather insights into the specific dietary grape-derived polyphenol(s) that might be relevant to AD, we subfractionated polyphenols from bioactive grape products (e.g., red wine and grape seed extract) containing increasingly less complicated polyphenol compositions and conducted in vitro and in vivo studies for exploiting potential beneficial AD-modifying activities. We fractionated total polyphenols from the red Cabernet Sauvignon wine and found that the Aβ-lowering activity of Cabernet Sauvignon can be attributed to its anthocyanin polyphenolic components (Fig. 4).

**Grape Polyphenols Beneficially Modulate Tau-Mediated Neuropathological Responses**

In recent studies, we found that grape seed polyphenolic extracts (GSPE) are capable of interfering with tau-mediated toxicity by interfering with the abnormal aggregation of tau [47–49]. We used both the TMHT [48] and JNPL3 [49] mutant tau mouse models of AD, which overexpress the human TAU441 gene bearing missense mutations V337M and R406W and express human tau protein containing the P301L mutation, to test the efficacy of GSPE in interfering with dementia resulting from abnormal tau functions. We found that dietary supplementation with GSPE in these tau mouse models effectively reduced the severity of abnormal tau aggregation and neuropathology in the brain [47–49]. While ongoing studies are evaluating the efficacy of grape-derived preparations in preserving cognitive function in these tau mouse models, our data suggest that GSPE might also protect against AD and other dementias in which tau neuropathology is a major contributory factor in the development of cognitive impairment.

**Dietary Grape-Derived Bioactive Polyphenolic Components in Alzheimer’s Disease Dementia**

Evidence from our studies strongly supports the hypothesis that moderate red wine consumption might provide preventive and/or therapeutic value in AD. Our experimental evidence suggests that in addition to providing antioxidant activities, polyphenols from red wines and other grape products may also benefit AD by directly modulating Aβ- as well as tau-related pathological mechanisms in the brain (Fig. 5). Based on our observation that multiple dietary grape products with distinct polyphenolic component compositions effectively protect against the onset and progression of AD phenotypes, we hypothesize that additional dietary products containing similar polyphenol forms as grapes, including other red wines, cocoa, tea, apple and berries, might also provide beneficial disease-modifying activities in AD.

There is an urgent need for additional studies in order to identify specific bioactive polyphenolics and corresponding polyphenolic metabolic derivatives from red wines or other grape-derived dietary products that are physiologically bioavailable in target tissues and in order to characterize the mechanisms of action of these bioactive polyphenols. Such information will provide the rational basis for developing selective bioactive dietary polyphenol(s) as lead compounds for clinical testing in AD. Moreover, this information will facilitate the selection of food sources enriched in targeted bioactive polyphenols that ultimately could be incorporated as key components in the development of dietary guidelines for AD prevention and/or management.

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**Conflict of Interest**

There are no relevant conflicts of interest to be reported.

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