Plant-derived acetylcholinesterase inhibitory alkaloids for the treatment of Alzheimer’s disease

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Abstract: The inhibition of acetylcholinesterase (AChE) has been one of the most used strategies for the treatment of Alzheimer’s disease (AD). The AChE inhibitors (AChE-I) produce not only short-term symptomatic effects, but can also play a role in other pathological mechanisms of the disease (e.g., formation of amyloid-β plaques), which has renewed interest in the discovery of such inhibitors. Four of the five currently prescribed treatments for AD are AChE-I. Natural alkaloids such as galantamine or alkaloid-related synthetic compounds (such as rivastigmine) are considered beneficial for patients with mild-to-moderate AD. However, there is a need for the discovery of more effective compounds and for this reason, plants can still be a potential source of new AChE-I. Findings and advances in knowledge about natural alkaloids as potential new drugs acting as AChE-I will be summarized in this paper.

Keywords: quinolizidine, steroidal, indole, isoquinoline

Introduction
Alzheimer’s disease (AD), an irreversible neurodegenerative disorder primarily targeting elderly populations, affects approximately 36 million people worldwide according to the 2010 estimations.1–3 This illness is characterized by progressive neurodegenerative disorders, collapse of cognitive functions, and formations of amyloid plaques and neurofibrillary tangles.2,3 AD is the most frequent cause of dementia in mid- to late life, and has a devastating impact on public health and society.

The AD pathogenesis is complex and comprises genetic and environmental factors,4–7 thus AD is considered a multifactorial disease. Different hypotheses regarding the pathological routes of AD have been proposed and the two main hypotheses are related to the cholinergic, amyloid-β (Aβ), and tau proteins. For this reason, drugs acting on acetylcholine (ACh) levels, mainly acetylcholinesterase inhibitors (AChE-I), or that reduce the formation of toxic Aβ peptides, mainly noncompetitive β-secretase (BACE-1) and secretase inhibitors, were studied for the development of anti-AD drugs.2 By its pharmacological nature, cholinesterase inhibitory therapy may be considered as a simple symptomatic intervention. However, therapy with AChE-I can be effective over a longer period,8 and it is believed that AChE also plays an important role in Aβ deposition.8–10 It seems likely that AChE interacts with Aβ and promotes the formation of amyloid fibril through a pool of amino acids located in proximity to the peripheral anionic-binding site of the enzyme.9 Furthermore cholinergic neurons are subject to control via the modulation of nicotinic ACh receptors (nAChRs) and some AChE inhibitors, such as galantamine, bind at allosteric secondary binding sites, and potentiate the receptor response to the available ACh.11 Since AD is a multifactorial
disease, the recent trend in the search for new drugs also focuses on multicated compounds.

Many plant-derived alkaloids that act as AChE-I can be considered as models for the development of AD drugs, and for example galantamine\(^{11–13}\) and rivastigmine\(^{14}\) are available for the symptomatic treatment of patients with mild to moderate AD. Three ChE-Is are the most frequently used drugs to treat patients with mild to moderate AD: donepezil, rivastigmine, and galantamine. Donepezil and galantamine are selective AChE-Is. Rivastigmine inhibits both AChE and butyrylcholinesterase (BuChE) from degrading ACh. The usefulness of donepezil, rivastigmine, and galantamine in the treatment of patients with AD has been recognized in recent health technology assessments.\(^{15}\) However, there is still a need for new AChE-I lead compounds with lower toxicity and higher central nervous system (CNS) penetration. Many plants have been studied by bioassay-guided approaches for the identification of new AChE-Is\(^{16}\) and different classes of plant-derived natural products have been considered as new AChE-Is potentially useful for AD treatment.\(^{17}\) Both nonalkaloids and alkaloid-derivative compounds\(^{16}\) have been studied, but the most potent AChE-I nowadays appear to be natural or semisynthetic alkaloids. This review focuses on plant-derived alkaloids with AChE-I properties and their potential role as models for the development of drugs for AD treatment.

**Acetylcholinesterase inhibitors as anti-Alzheimer drugs**

The AChE present in the CNS catalyzes the hydrolysis of the ACh to choline. ACh is released in the synaptic cleft, where it activates both postsynaptic and presynaptic cholinergic receptors, namely nicotinic and muscarinic, leading to an increase of cholinergic transmission, which results in cognition improvement.\(^{18}\) Its neural-stimulating activity is terminated by the action of the cholinesterase enzymes, namely AChE and BuChE. In AD patients, a deficit of this neurotransmitter was observed and this led to the so-called cholinergic hypothesis.\(^{4}\)

Classified as a serine hydrolase, the enzyme AChE uses three amino acidic residues in the active site: histidine, serine, and glutamic acid. The first two activate the side chain of a serine residue. Once the active site serine forms a tetrahedral intermediate with the ACh carbonyl group, choline is released. Then a water molecule is employed in the active site for releasing acetate from the serine, which returns the enzyme back to its original state. The enzyme active site is located at the bottom of a 20Å gorge acting as a bottleneck in which some aromatic residues line up the entry and contribute to its specificity of action.\(^{2}\) The enzyme AChE is the target of the available AChE-Is for AD treatment and the aim is obviously to restore, at least in part, the levels of the neurotransmitter (Figure 1). This approach can be considered
as a symptomatic short-term intervention, but some data emerging from long-term open-label trials have shown that the effect may be prolonged. For this reason, the observed effect of stabilization of the cognitive status of patients suggests the presence of further mechanisms.\(^\text{16}\) Cholinesterase inhibitors could perhaps help the prevention of the formation of β-amyloid plaques, while the peripheral site of AChE promotes aggregation of β-amyloid peptides.\(^\text{19}\) Other mechanisms are also under investigation and multitargeted drugs will be preferred due to the multifactorial pathogenesis of the disease. The use of AChE-I has been associated with an anti-inflammatory effect mediated by \(\alpha-7\) nAChRs. The inflammatory component in AD is very important and the role of inflammation and AChE-I emphasizes the role of cholinergic balance in this pathology.

**Alkaloids as acetylcholinesterase inhibitors**

On one hand, many natural alkaloids have been studied as AChE-Is, but relatively few compounds have entered therapeutic use. On the other hand, many different classes of compounds have been considered, namely indole derivatives (such as physostigmine and related compounds), isoquinoline and related derivatives (such as galantamine and lycorine-type alkaloids), steroid and terpenoid alkaloids, and many other derivatives that present significant inhibitory effects on AChE.\(^\text{16}\) Plant families that have been considered as potential sources of such alkaloids were Amaryllidaceae, Buxaceae, Apocynaceae, Papaveraceae, Lycopodiaceae, and Leguminosae. The great chemical diversity of the different derivatives studied presents the opportunity to explore various chemical scaffolds for the development of potential new drugs to act on enzymes with different pharmacological profiles. One opportunity is to generate new AChE-Is coupled with properly selected bioactive molecules to obtain homo- and heterodimers endowed with increased potency and supplementary actions.\(^\text{20}\) Furthermore, compounds possessing not only the AChE inhibitory properties, but also other anti-inflammatory or antioxidant qualities, such as protoberberine and related compounds are considered attractive because of the possibility to create multitargeted drugs acting with different mechanisms all related to the disease.

**Indole alkaloids and related compounds**

The indole alkaloid family is probably the most important source of active natural products in compounds that have entered clinical use. Indole alkaloids are found in a large group of plants and also in fungi. Reserpine, yohimbine, vincamine, and vinblastine are only some examples of such compounds with medicinal importance. The majority of these alkaloids are derived from tryptophan and the monoterpeneid sesquiterpan, yielding complex formations of tetracyclic (as agroclavine), pentacyclic (ajmalicine) hexacyclic (apodine), and heptacyclic (brucine) structures.\(^\text{21}\) Physostigmine (Figure 1) is the main alkaloid of the Calabar bean, the seed of *Physostigma venenosum* (Leguminosae), and was one of the first investigated AChE-Is. This compound acts as a parasympathomimetic agent and is able to pass through the blood–brain barrier. It is a potent short-acting inhibitor of the enzyme and has been shown to improve the cognitive function in vivo and both in normal and AD patients.\(^\text{13–16,22,23}\)

Rivastigmine has a “pseudoirreversible” inhibitor of AChEs and BuChEs with a phenylcarbamate structure, the metabolism of which is almost totally independent of the hepatic cytochrome P450 system. After binding to cholinesterase, the carbamate portion of rivastigmine is slowly hydrolyzed, cleaved, conjugated to a sulfate, and excreted. Rivastigmine has an oral bioavailability of 0.355 and low (40%) binding to plasma proteins. Its elimination half-life is around 2 hours.\(^\text{14}\) Its short half-life, narrow therapeutic index, and secondary effects did not allow its clinical use.\(^\text{16,22,23}\) However, the indole alkaloids family still remain an important source of new AChE-Is.\(^\text{17,23}\)

Therefore, starting with physostigmine, many derivatives were prepared to improve efficacy and the pharmacokinetic profile. Phenserine, tolserine, and several derivatives (Figure 1) present potent AChE-I activity with an half-maximal inhibitory concentration (IC\(_{50}\)) in the nanomolar range.\(^\text{4,17}\) The synthetic modification occurred in the substituent pattern of the phenyl group of the phenylcarbamoyl moiety of the phenserine compound. Simple aliphatic chains (as methyl-, di-, or trimethyl groups) were linked in each position of the aromatic ring leading to a series of derivatives with similar AChE-I properties to the starting compound. As our first example, the phenylcarbamate derivative phenserine was proposed as a new potent inhibitor. The compound was considered less toxic and more active compared with physostigmine, and in vivo studies reported its efficacy in cognition enhancement and β-amyloid peptide formation.\(^\text{17}\) In particular, the compound is able to act as an AChE-I and to inhibit amyloid precursor protein gene.\(^\text{4,24}\) Furthermore the (−) phenserine enhanced cognition in vivo and in AD patients (at a dose of 30 mg/day), but in Phase III clinical studies, there was no difference in effects compared to placebo in AD patients. Also the dextrorotatory enantiomer (+) phenserine
was considered and it was used in a Phase I clinical study, but further clinical studies are not planned.²²,²⁵

Recently, a shrub that was used in folk medicine for the treatment of various diseases, Himatanthus lancifolius, was studied for the possible anti-AChE properties of its alkaloids. The plant contains several indole alkaloids. Rich fractions of lipophilic alkaloids were able to act as AChE-I and uleine (Figure 1) was isolated from them.²²,²⁶ This alkaloid was firstly isolated from the root bark of Aspidosperma ulei MgF and was studied from a chemical and synthetic point of view.²⁷–³⁰ Furthermore, this alkaloid and its related derivatives were also studied for their anticancer properties,³⁰ and, more recently, alkaloids-enriched fractions were also studied for their in vitro anti-inflammatory properties.³¹ It was reported recently that uleine has significant AChE-I activity with an IC₅₀ of 0.45 μM.²⁶ For this reason, uleine is considered as a new possible candidate for the future development of new anti-AD drugs, but no reports about safety, efficacy, and human studies specific to cognition have been reported to date.²²,²⁶

Isoquinoline and Amaryllidaceae alkaloids

Isoquinoline alkaloids are probably the largest single group of plant alkaloids.²¹ Their basic structure is derived from tetrahydroisoquinoline, which can be modified in complex structures that can be all biosynthetically derived from phenylalanine and tyrosine. These alkaloids are aromatic bases; the majority of the natural derivatives present tetracyclic structures, but pentacyclic compounds (such as aporphine and emetines) are also present. The most widely distributed isoquinoline alkaloids are the aporphines and protoberberine types.²¹ The morphine-correlated compounds are the most notable for their medicinal chemistry and clinical use in this important class of compounds. The presence of Amaryllidaceae alkaloids, as their name suggests, is restricted to the Amaryllidaceae family and no other classes of alkaloids were reported from this source. The daffodil family includes narcissus, snowdrops, and amaryllis and is formed by 75 genera and about 1100 species.²¹ All these alkaloids share a C-15 skeleton and their biosynthesis involves the phenolic coupling of two aromatic amino acids with one of the two nitrogen atoms retained in the process.²¹ These two classes of alkaloids are considered very important AChE-Is.

The alkaloid galantamine is one of the most important derivatives with great therapeutic value for AD treatment (Figure 2). This compound was isolated from Galanthus woronowii and G. nivalis and some Narcissus and Leucojum spp. of Amaryllidaceae. This compound is reported to be more selective to AChE than BuChE and provides good oral bioavailability (between 80% to 100%) with linear pharmacokinetics.³²–³³ Galantamine inhibits the breakdown of ACh by binding competitively and reversibly to the active site on AChE. Of particular importance are the inhibitory effects of galantamine on AChE in the frontal cortex and hippocampal regions of the brain, the two areas in which cholinergic neurotransmission is most affected in patients with AD.¹¹ In addition galantamine has a peculiar mechanism of action because it both acts as competitive inhibitor of the enzyme and as a positive allosteric modulator of nAChRs.¹¹,²² The inhibition of AChE is potent and selective (53-fold greater compared to BuChE)¹¹ and the effect is observed in the frontal cortex and hippocampal region of the brain, which are areas that present poor cholinergic neurotransmission in the AD patients. The relatively low affinity for BuChE, mainly diffused in plasma, is considered an advantage in terms of the poor peripheral effect of the drug.¹¹ The second mechanism of action of galantamine is the allosteric modulation of nAChRs. Galantamine binds to a secondary (allosteric) site in the nAChR, and this makes the receptor more sensitive to the physiologic ligand (ACh), which enhances the response to the available ACh. Furthermore, in vivo studies in animal models suggest that the modulatory activity of galantamine in nAChRs increases the number of such receptors in long-term treatments.¹¹,³² Galantamine is actually used in therapy (as Reminyl®(Janssen-Cilag SpA,)) and its use is associated
with a preservation of cognitive function in patients with AD. It also stabilizes patients and postpones the emergence of behavioral symptoms, which improves aberrant motor behavior, anxiety, disinhibition, and hallucinations.\textsuperscript{1,11} Of the various natural products that have been studied, galantamine is considered a well-tolerated treatment for the control of AD progression.\textsuperscript{11}

Several derivatives were prepared from the galantamine scaffold including C and D ring modifications, quaternary ammonium salts, carbamate and esters, and bis-interacting ligands, but pharmacology studies on the possible efficacy of these compounds are still lacking.\textsuperscript{17}

Ungeremine and lycorine (Figure 2) were isolated from \textit{Nerine bowdenii} W Watson (Amaryllidaceae) and from \textit{Narcissus pseudonarcissus} (Amaryllidaceae). These compounds are moderate AChE-Is, and have been used as models for the preparation of several synthetic derivatives with the aim of improving the AChE-I activity of the starting compounds.\textsuperscript{17} Considering the different Amaryllidaceae alkaloids, the lycorine type derivatives exhibited the higher activity, namely O-acetyl lycorine was twofold more potent than galantamine.\textsuperscript{17}

New therapeutic treatments should target multiple AD pathways to prevent the rapid progression of AD as opposed to the traditional “one drug, one target” approach, and for this reason some isoquinoline and benzylisoquinoline derivatives were considered potentially useful compounds. Alkaloids from \textit{Colchicum speciosum} Steven (Colchicaceae), \textit{Coptis} spp. (Ranunculaceae) and \textit{Corydalis} spp. (Papaveraceae) act as AChE-Is.\textsuperscript{12} Epiberberine, pseudoberberine, and pseudocoptisine are examples of such compounds. Berberin was recently considered as an attractive compound for AD treatment not only due to its AChE-I, but also for BuChE, antioxidant, monoamine oxidase inhibitory, amyloid-\(\beta\) peptide level-reducing and cholesterol-lowering activities which suggest a mult-targeted approach for the disease’s treatment.\textsuperscript{33} Berberin-based derivatives were developed to prepare more potent AChE-Is.\textsuperscript{34} Pseudoberberine and pseudocoptisine are also able to alleviate scopolamine-induced memory impairment in vivo models.\textsuperscript{35,36}

Structures of some derivatives that have been studied for their AChE-I properties are reported in Figure 3.

With an interesting mechanism of action, some \textit{C. chinensis} alkaloids with AChE-I properties also

\begin{figure}[h]
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\includegraphics[width=\textwidth]{structures.png}
\caption{Structures of protoberberine-related alkaloids.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{carinatumin.png}
\caption{Carinatumin and annotine alkaloids.}
\end{figure}
displayed BACE-1 inhibitory activity. These compounds are groenlandicine and jatrorrhizine and their derivatives. These alkaloids also present antioxidant activity, therefore they possess multiple activities relevant to AD.\textsuperscript{37}

Alkaloids from \textit{Stephania venosa} Spreng. (Menispermaceae) such as stepharanine, cyclanoline, and N-methyl stepholidine are potent AChE-I\textsuperscript{3}. The structural activity relationships of this protoberberine derivative revealed that a positive charge at the nitrogen of the tetrahydroisoquinoline portion, a steric substitution at the same nitrogen, and a planarity of the compound are important features for AChE-I. Pattern substitution at positions 2, 3, 9 and 10 can also modulate activity and the \textit{S. venosa} derivatives stepharanine and cyclanoline were more potent than cordylamine.\textsuperscript{38}

\textbf{Quinolizidine and Lycopodium alkaloids}

The quinolizidine alkaloids possess a structural unit in which a nitrogen atom occupies a central position in two six-member fused rings. In general, such alkaloids are found in Leguminosae and mainly in plants of the genus \textit{Lupinus}. A typical bicyclic structure is found in lupin and in other more complex structures such as anagyrine, which possess more cycles. These compounds are quite toxic and their occurrence in trees and shrubs is the cause of accidental poisoning in sheep and in some cases also in humans. Some of these alkaloids have been studied for their potential medicinal use. Lycopodium alkaloids are related to quinolizidine, but possess an unusual tetracyclic ring system. Several derivatives have been reported, which differ in their ring closures and rearrangements as well as for methyldihydroxy or other substituent positions. As a class, the Lycopodium alkaloid compounds are moderately toxic.\textsuperscript{21}

The carinatumins A and B (from \textit{Lycopodium carinatum} Desv. ex Poir.), and other alkaloids isolated from \textit{Lycopodium} spp. present interesting AChE-I activity. Their structures are reported in Figure 4. For example, carinatumins A and B present an \textit{IC}_{50} of 4.6 and 7.0 \textmu M, respectively.\textsuperscript{39} Carinatumins A and B, lycoparin C, and other derivatives present significant AChE-I activity, but none of these alkaloids are considered of therapeutic relevance to date. Lycojapodine A is an unusual C16 N type alkaloid isolated from \textit{L. japonicum} that possesses a 6/6/6/7 tetracyclic ring system.\textsuperscript{4} Its AChE-I activity is similar to galantamine and it could be a future potential new candidate for the development of AD drugs. Annotine from \textit{L. annotinum} and other lycoparin C derivatives from \textit{L. casuarinoides} (Spring) were studied for their AChE-I properties, but data about efficacy or safety for possible clinical use are lacking.\textsuperscript{40,41}

A new Lycopodium alkaloid with a peculiar structure was isolated from the traditional Chinese medicine qing ceng ta, namely the \textit{Huperzia serrata}, Huperzine A (Hup A; Figure 5). This compound is considered a new potential drug for the treatment of AD patients. This alkaloid is a potent, reversible AChE-I, whose activity is comparable to the other well-known alkaloids, physostigmine, galantamine, or tacrine. In the 1990s, this alkaloid was approved in China as an anti-AD drug and is currently marketed and patented in the USA as a dietary supplement.\textsuperscript{42–44} Compared to physostigmine or galantamine, Hup A presents higher selectivity to AChE versus BuChE. After oral or intraperitoneal administration, Hup A produced significant inhibition of AChE in the cortex, hippocampus, corpus striatus, medial septum, medulla oblongata, cerebellum, and hypothalamus in rats. Furthermore, studies in animal models revealed significant neuroprotective effects against various neurodegenerative conditions, including AD.\textsuperscript{45–50}
that, compared to donepezil and tacrine, Hup A has higher bioavailability and more easily penetrates the blood–brain barrier.\textsuperscript{42–44} Hup A acts as a mixed competitive inhibitor and some findings can elucidate in part how this alkaloid can bind to the enzyme. The principal interactions include a direct hydrogen bond between the carbonyl group and the hydroxyl oxygen of Tyr 130 at a peripheral site of the enzyme. A further interaction is between the ethylidene methyl group and the main chain oxygen of His 440. Another interaction is the indirect hydrogen bond mediated by water molecules with the aminoacidic residues that constitute the active centre of the enzymes, and the cation interactions induced upon inhibitor binding, between the aminogroup of the alkaloid and the aromatig of Trp 84 and Phe 330 at the choline site. This binding site is shared with other AChE-Is such as tacrine and edrophonium.\textsuperscript{45} The high selectivity of this alkaloid was also studied with the aid of computer-assisted docking studies and with the crystal structure of the AChE–Hup A complex. For example, Hup A is able to form an hydrogen bond with the residue of Tyr 337 within a choline site that exists only in the mammalian homolog of the enzyme, but is not present in the torpedo enzyme and also in BuChE.\textsuperscript{44,46} The alkaloid Hup A is considered a promising compound due to other mechanisms of action that have been studied recently, namely the reduction of $\beta$-amyloid precursor protein metabolism, and the decrease of Aβ-associated neurotoxicity.\textsuperscript{47,48}

Some clinical studies have reported significant improvements in memory deficiencies in aged patients and AD patients. A recent meta-analysis of Hup A’s efficacy and safety showed it is well tolerated and significantly improves cognitive performance and activities of patients with AD. The closely related Hup B is a less potent AChE-I, and probably for this reason it has not been extensively investigated for its possible clinical uses.

**Triterpenoid steroidal alkaloids**

The triterpenoid skeleton appears to be a potential scaffold for the development of AChE-I. Some reports describe the AChE-I of nonalkaloidal triterpenoids.\textsuperscript{16} However, nitrogen-containing compounds such as hyrcanone, hyrcamine, or buxidine (isolated from *Buxus* spp.) were described as moderate inhibitors of the enzyme AChE. A number of steroidal alkaloids were isolated from *Buxus*, *Sarcococca*, and *Fritillaria* spp. and, for example, (+)-O$^6$-buxafurandiene

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**Table 1** Alkaloids with AChE-I activity with a potential use in AD disease

<table>
<thead>
<tr>
<th>Alkaloid class</th>
<th>Structure</th>
<th>Main compound</th>
<th>Plant family</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indole</td>
<td><img src="image1" alt="Indole Structure" /></td>
<td>Physostigmine</td>
<td>Leguminosae</td>
<td>Physostigmine is not clinically used anymore.</td>
</tr>
<tr>
<td>Isoquinoline</td>
<td><img src="image2" alt="Isoquinoline Structure" /></td>
<td>Galantamine</td>
<td>Amaryllidaceae</td>
<td>Galantamine acts as both AChE-I and nAChR and is currently in clinical use.</td>
</tr>
<tr>
<td>Lycopodium type</td>
<td><img src="image3" alt="Lycopodium Structure" /></td>
<td>Huperzine A</td>
<td>Huperziaceae</td>
<td>AChE-I activity, reduction of Aβ deposition, decreases Aβ related neurotoxicity.</td>
</tr>
</tbody>
</table>

**Abbreviations:** AChE-I, acetylcholinesterase inhibitors; AD, Alzheimer’s disease; nAChR, nicotinic acetylcholine receptors.
and (+)-7-deoxy-Oβ-buxafurandiene (Figure 6) presented significant AChE-I activity. Several different papers have been published about this compound and the structural activity relationships are described at least in part.12,49–51 In many derivatives that have been isolated, the activity may be associated with the presence of an α,β-unsaturated amide at C-3 of the ring A, and with the presence of a α,β-unsaturated carbonyl moiety.4,12,49–51 Furthermore, the presence of the dimethylamino moiety at C-20 and the basic nitrogen atom can enhance the structural activity and also improve the active transport of the compounds in the cell. Such compounds are studied as new candidates for the development of new AD drugs, but reports about their in vivo or clinical studies are still lacking.4

Miscellaneous alkaloids

Ribalone and methyl isoplatydesmine were isolated from Skimmia laureola (Rutaceae) and present significant AChE-I activity. Quinoline (nigellastrine), and β-carboline (harmane and related products), tetrahydroisoquinoline (salsoline and derivatives) from Salsola spp. (Chenopodiaceae) present significant AChE-I activity.4,52 Nevertheless, pharmacological studies about efficacy and safety are still lacking. Compared to other alkaloid classes, the piperidines are less studied as AChE-Is. (−)-spectaline and derivatives isolated from Cassia spectabilis are an example of this and the compounds were used to develop the synthetic derivative. Other alkaloids that are considered of therapeutic interest are sinapine that is an ester of sinapinic acid and choline that is present is Raphanus sativus, which acts as a powerful AChE-I in vitro, and tapsin isolated from Magnolia x solangiana Soul.-Bod (Magnoliaceae), which presents an AChE-I activity greater than galantamine (IC₅₀ = 0.3 μM vs 3.2 μM, respectively).4,51

As observed previously, many natural derived alkaloids have been studied as an AChE-I with potential use in AD, but they also have been used as models to synthesize new compounds. One example is rivastigmine (Figure 7), which is a simple synthetic derivative approved as treatment for moderate-to-mild AD in the USA in 2000, with doses of 6–12 mg/day resulting in improvement of cognition in patients.22,54 Recently the drug was also proposed in the form of a transdermal patch to reduce the side effects that are mainly nausea, vomiting, anorexia, and diarrhea.22 The derivative, xan-tostigmine (Figure 7) is able to inhibit AChE-induced β-amyloid aggregation.55

Conclusion

AD is the most frequent form of dementia and is characterized by memory loss, mental and physical behavioral changes, and reduced quality of life for patients, with an important impact on public health. The pathology is progressive and causes β-amyloid plaque formation and deficiency in the neurotransmitter acetylcholine in the brain. Nowadays the available drugs have limited effect on this pathology, and in fact are only able to control symptoms to preserve cognitive function and slow down the progression of the illness. AD is a multifactorial disease and several pathogenic events are involved in this disease including primary events such as genetics, neuronal death, brain dysfunction. Furthermore, secondary (β-amyloid deposition, tau protein hyperphosphorylation), tertiary (neurotransmission deficiency, neuroinflammation), and quaternary (neuronal death, free radical formation, cerebrovascular dysfunction) events also play a complex role in this pathology. Despite the new findings, AChE-I still represents an important symptomatic therapy and research of new drugs acting on this enzyme is essential. The plant kingdom, an important source of several drugs or “lead compounds” for medicinal chemistry, is still largely unexplored despite the relatively large amount of tested plant extracts for anti-AChE-I. In recent years, many botanical species have been studied as potential sources of AChE-I alkaloids, but relatively few isolated compounds were considered as potential new “leads” for the development of drugs for AD treatment (Table 1). Different compounds belonging to various alkaloids classes were identified as enzyme inhibitors, in some cases not only from plants, but also from other natural organisms.12,16,56 These findings produce an opportunity to modify and prepare several useful derivatives by synthesis or semisynthesis to understand their structural activity relationships. Many of the studied alkaloids have been studied by in vitro or in some cases in vivo models, but future studies are needed to demonstrate their therapeutic role. No human or animal studies have been performed in many compounds, therefore studies of their safety, efficacy, and toxicity are needed. Many approved drugs in the market for AD treatment are natural alkaloids and galantamine is an example. Furthermore, many AChE-I compounds studied as potentially relevant agents in AD treatment are primarily from plants. The evidence that AD is a multifactorial disease is also guiding the research in the direction of multiple targeted compounds. In this case, many natural alkaloids (such as protoberberine and related compounds) have been studied not only for their AChE-I properties, but also for their
anti-inflammatory and antioxidant activities that can play an important role in the development of new anti-AD drugs.

Disclosure

The author reports no conflicts of interest in this work.

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