Natural sesquiterpen lactones as acetylcholinesterase inhibitors

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ABSTRACT

Background and the purpose of the study: The amount of elder people who suffer from Alzheimer disease is continuously increasing every year. Cholinesterase inhibitors have shown to be effective in alleviating the symptoms of the disease, thus opening a field of research for these treatments. Herbal products, owning a reputation as effective agents in many biological studies are now drawing attention for inhibiting acetylcholinesterase, in other words, Alzheimer disease. In the present study, the ability of three sesquiterpene lactones from Inula oculus-christi and I. aucheriana to inhibit AChE has been evaluated through Ellman assay.

Materials and Methods: Gaillardin and pulchellin C were obtained from I. oculus-christi and britannin from I. aucheriana by chromatographic methods. They were dissolved in methanol in concentration of 3 mg/mL and the AChEI activity of the compounds was determined by Ellman method using Acethylthiocholine iodide as the substrate and 5, 5'-dithiobis-2-nitrobenzoic acid as the reagent, in 96-well plates at 405 nm.

Results: AChEI activity of the examined compounds was obtained as 67.0, 25.2 and 10.9% in concentration of 300 μg/L for gaillardin, britannin and pulchellin C, respectively.

Conclusion: Among the three sesquiterpene lactones, gaillardin with 67% inhibition of AChE could be considered a good candidate for future Alzheimer studies.

Key words: Acetylcholinesterase inhibitor, Alzheimer, britannin, gaillardin, Inula, pulchellin C.
the cholinergic hypothesis, acetylcholinesterase inhibitors (AChEIs) are widely used to alleviate AD. Nowadays investigations to find AChEI materials from natural sources have increased and it has been proven that many compounds of different structures have AChEI property; among which alkaloids are the most ruling compounds. Agents such as the alkaloid galanthamine, are examples possessing the mentioned property (Feitosa et al. 2011). There are not as many reports about AChEI activity of sesquiterpenes as alkaloids, but the existing data are indicative of the mentioned activity in some herbs and their sesquiterpene constituents. The results of isolating sesquiterpene lactones from *Amberboa ramosa*, a plant of Asteraceae, have revealed their excellent AChEI activity (Ibrahim et al. 2013); while the leaves essential oil of *Hedychium gardnerianum* has exhibited AChEI activity (Arruda et al. 2012) recommending the oil to be used in aromatherapy. Since the constituents were mostly found to be sesquiterpenes, the authors had concluded that the AChEI activity was due to the presence of these compounds. The *n*-hexane fractions of the peels and leaves of bitter orange have also demonstrated to be inhibitors of AChE and gas chromatography-mass spectrometry analysis have again deduced the most dominant compounds to be mono- and sesquiterpenes (Loizzo et al. 2012). These results attracted our attention to the possibility of other sesquiterpenes to encompass the ability to inhibit AChE and in the present study, sesquiterpene lactones isolated from two species of *Inula* (Asteraceae) named *I. oculus-christi* L. and *I. aucheriana* DC. have been investigated through Ellman assay.

**MATERIALS AND METHODS**

**CHEMICALS**

Acetylcholinesterase (AChE) was purchased from Sigma, Germany. Acetylthiocholin Iodide (ATCI) was prepared from Fluka, Germany. 5, 5'-dithiobis-2-nitrobenzoic acid (DTNB) and other solvents and chemicals were provided from Merck, Germany.

**PLANT MATERIAL**

In June 2009, *Inula oculus-christi* L. was collected from Golestan province and *Inula aucheriana* DC. was collected from West Azerbaijan province in Iran (July 2010). The species were authenticated by Mrs. Atefeh Pirani (Botanist), Traditional Medicine and Materia Medica Research Center (TMRC), Shahid Beheshti University of Medical Sciences, Tehran, Iran. Voucher specimens of both species have been deposited at TMRC Herbarium for future reference. The aerial parts of the plants were dried in shade and ground.

**EXTRACTION AND ISOLATION**

Gaillardin and pulchellin C were obtained from *I. oculus-christi* (Mosaddegh et al. 2010); briefly, the chloroform extract of 250 g of the aerial parts was further subjected to column chromatography to afford gaillardin and pulchellin C. Britannin was obtained from chloroform fraction of 500 g *I. aucheriana*, through serial extraction with *n*-hexane, chloroform and methanol (Hamzeloo-Moghadam et al. 2012). The chemical structures of the compounds are presented in figure 1.

**ACETYLCOLINESTERASE INHIBITORY ACTIVITY ASSAY**

Ellman assay is usually applied *in vitro* to define the cholinesterase activity of materials. It was established by Ellman et al. (1961) and is based on the reaction of thiocholine (one of the products of enzymatic hydrolysis of acetylthiocholine) with DTNB (Ellman’s reagent), forming a yellow product (5-mercaptop-2-nitrobenzoic acid and its dissociated forms) at pH 8 which can be detected at 405 nm (Komersova et al. 2007). Based on the above mentioned assay, the experiment was conducted according to modified microplate method (Mukherjee et al. 2007). All samples were dissolved in methanol in concentration of 3 mg/mL. In the 96-well plates, 125 μL of 3 mM DTNB, 25 μL of 15 mM ATCI and 50 μL of phosphate buffer (pH 8), 25 μL of sample dissolved in methanol were
added to the wells. The absorbance was read at 405 nm every 13 s for 65 s. 25 μL of 0.22 U/mL of AChE enzyme was then added and the absorbance was again measured every 13 s for 104 s using a TECAN microplate reader at 405 nm. Absorbance was plotted against time and enzyme activity was calculated from the slope of the line and expressed as a percentage compared to an assay using a methanol without any inhibitor. Any increase in absorbance due to the spontaneous hydrolysis of the substrate was corrected by subtracting the rate of the reaction before adding the enzyme, from the rate after adding the enzyme. Inhibition percentage was obtained by comparing the rates for the sample to the blank (methanol). Donepezil was used as the positive control.

RESULTS AND DISCUSSION

Acetylcholinesterase inhibitors (AChEIs) as treatments for AD, are the first class of agents approved by the US Food and Drug Administration (FDA). They are thought to bind to acetylcholinesterase in the synaptic cleft, giving more time to acetylcholine which has been released from the pre-synaptic cholinergic terminal; thus increasing the chance of interacting with the postsynaptic cholinergic receptors (Aisen et al. 2012). The administration of AChEIs in some clinical trials have shown advantages in approximately 40–50% of patients, though the effect was temporary. There is also a suggestion of neuroprotective effect of these treatments (Kemp et al. 2003).

Plants have a reputation of being rich sources of effective compounds for drug discovery and might have advantages regarding efficacy (Frydman-Marom et al. 2011). Physostigmine, galanthamine and huperzine A are the alkaloid-type of compounds isolated from the plants which serve as medicines for AD (Orhan et al. 2009). Most researches on acetylcholinesterase inhibitory effect of natural sources have been focused on alkaloids, regarding other groups of natural compounds, as next rates. Some examples of acetylcholinesterase inhibitors include steroidal alkaloids from Sarcoccasa ligna (Atta-ur-Rahman et al. 2004), semi-synthetic analogues piperidine alkaloids of Senna spectabilis (Viegas et al. 2005), triterpenoid alkaloids from Buxus hyrcana (Choudhary et al. 2006), bisbenzylisoquinoline alkaloids from Cocculus pendulus (Atta-ur-Rahman et al. 2009) and alkaloid leptomerine from stems of Esenbeckia leiocarpa (Cardoso-Lopes et al. 2010).

The results of the present investigation proved that sesquiterpene lactone gaillardin inhibited the enzyme by 67% in concentration of 300 μg/mL which showed reasonable inhibition (figure 2). Pulchellin C and britannin showed 10.9% and 25.2% inhibition, respectively (figure 3).

The AChEI property of gaillardin draws attention to these versatile secondary metabolites and the plants containing sesquiterpene lactones, namely the Asteraceae species. The drawback to sesquiterpene lactones might be because of the reports of cytotoxic activity about most of them. Cytotoxicity...
of britannin, gaillardin and pulchellin C have also been reported before (Mosaddegh et al. 2010, Hamzeloo-Moghadam et al. 2012). Their chemical structures make somewhat great differences in their cytotoxic effects which indicate britannin as the most and pulchellin C as the least effective one. Looking back at the results of AChEI activity of the three, demonstrates that gaillardin has presented much more activity than the other two sesquiterpene lactones and pulchellin C again is the least effective. Pulchellin C is an eudesmanolide sesquiterpene lactone, while the other two are both guaianolides. Regarding polarity, pulchellin C is the most polar with the least AChEI activity. It could be concluded that polarity might play a role in AChEI activity. Gaillardin which is the least polar compound, has the most AChEI activity (67%) and britannin and pulchellin C are in the next rates, respectively. In addition to polarity, chemical structure could make a lot of difference in binding to AChE. It is clear that all the three compounds belong to sesquiterpene lactones and they all bear the α-methylene-γ-lactone functional group but they do not simulate the same AChE inhibitory nor cytotoxic activity, suggesting a groundwork where medicinal chemistry rules. Hence derivatization might be a choice to be considered whenever cytotoxic activity is the hindrance, and it could even increase the biological activity, giving a chance to reach to more effective, less cytotoxic derivatives by manipulating the structures. The clinical efficacy and safety of gaillardin and other sesquiterpene lactones are still to be taken into account. There is limited or even no clinical data about these compounds, thus further assessments are crucial.

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RESUMO

Embasamento teórico e proposta de estudo: A quantidade de pessoas idosas que sofrem da doença de Alzheimer está aumentando a cada ano. Inibidores de colinesterase têm mostrado serem eficazes em amenizar os sintomas da doença, abrindo, então, um campo de pesquisa para estes tratamentos. Produtos de origem vegetal, que possuem uma reputação como agentes eficazes em muitos estudos biológicos, ultimamente vem chamando a atenção por inibirem a acetilcolinesterase (AChE), em outras palavras, a doença de Alzheimer. No presente estudo, a capacidade de três lactonas sesquiterpênicas de Inula oculus - christi e I. aucheriana para inibir a AChE foi avaliada pelo ensaio de Ellman.

Materiais e Métodos: Gailardina e pulchelina C foram
obtidas a partir de I. oculus-christi e a britania de I. aucheriana por métodos cromatográficos. Elas foram dissolvidas em metanol na concentração de 3 mg/mL e a atividade AChE dos compostos foi determinada pelo método de Ellman usando iodeto de acetilícololina como substrato e o ácido 5, 5’-ditiobis-2-nitrobenzoico como o reagente, em placas de 96 poços a 405 nm. **Resultados:** A atividade AChEI dos compostos avaliados foi de 67,0, 25,2 e 10,9% na concentração de 300 mg/mL para gailardina, britania e pulchelina C, respectivamente.

**Conclusão:** Entre as três lactonas sesquiterpênicas, gailardina, britanina e pulchelina C, respectivamente.

**Palavras-chave:** inibidor de acetilcolinesterase, Alzheimer, estudos na doença de Alzheimer.

**REFERENCES**


