Full Length Research Paper

Isolation and structure elucidation of acetyl cholinesterase inhibitor from *Gyrostoma helianthus* of the Red Sea, Egypt

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Received 16 October, 2012; Accepted 15 March, 2013

The present study was carried out to isolate, purify and elucidate the structure of low molecular weight bioactive compounds from sea anemone, *Gyrostoma helianthus* of the Red Sea environment. The obtained results indicated that the ethanolic crude extract of the sea anemone, *G. helianthus* showed inhibition activity against acetylcholinesterase (AchE). The 0.5 kD fraction inhibited the activity of the AchE that indicated the presence of low M.W active compound(s) of less than 0.5 kD. Two active fractions were obtained after BioGel P2 fractionation of the 0.5 kD of the *G. helianthus*. The first active fraction inhibited the activity of the acetylcholinesterase, while the other fraction did not. High-performance liquid chromatography (HPLC) technique aided by semipreparative columns was used to isolate the target compounds in a pure form for the structure elucidation, one dimensional nuclear magnetic resonance (NMR) analysis (¹H and ¹³C-NMR) and DEPT were carried out to elucidate the structure of the isolated compound which was tentatively identified as N,N’-bis-(1-methyl-pyridin-2-yl)-hydrazine.

Key words: Sea anemone, *Gyrostoma helianthus*, Red Sea, structure elucidation, acetylcholinesterase (AchE) inhibitor.

INTRODUCTION

The study of marine organisms as a source of biologically active compounds is considered a very productive field, having already led to the discovery of various new pharmacological tools and medicines (Bhakuni, 1994; Munro et al., 1999; Faulkner, 2001).

The work of Bergman and Feeney at the beginning of the 1950s initiated the study of marine natural products, and in the last few decades, an appreciable number of new compounds have been isolated from marine organisms (Bhakuni, 1994; Faulkner, 2000a, b; Faulkner, 2001). Many
The search for novel low molecular weight natural products from marine organisms is of crucial importance for different applications as drugs and pesticides. Of special interest is the Red Sea environment, which is unique in its life forms and considered as one of the richest marine areas in biodiversity and richer than similar areas in the eastern Pacific and Atlantic (Gomaa and Aboul-Enein, 2000).

Therefore, the objective of the present study was to isolate, purify and elucidate the structure of low molecular weight bioactive compounds from sea anemone, *Gyrostoma helianthus* of the Red Sea environment using the bioassay guided fractionation technique.

**MATERIALS AND METHODS**

**Organism under study**

Sea anemone, *G. helianthus* was collected from the vicinity of Hurghada, Red Sea, Egypt.

**Sampling locations**

Samples of sea anemone, *G. helianthus* were collected from Hurghada vicinity at latitude from 27°00'N to 27°45'N and longitude from 33°30'E to 34°00'E. The locations of the sampling sites were as follows:

Shaab AbuShaar, Shaab AbuGalawa, Shaab AbuSadaf, ElFanadir Island, Gafton Kabier Island, Gafton Saghier Island, Dishet AbuMinqar, Magawish Kabier Island and Magawish Saghier Island.

**Sample preparation**

Sea anemone, *G. helianthus* samples were collected by trained divers working in the areas known to harbor the specific species under study. The collected samples were kept in seawater during the sampling trip around the vicinity of Hurghada. Upon arrival at the laboratory of Hurghada Marine Station, National Institute of Oceanography and Fisheries, the samples were washed with distilled water followed by gentle centrifugation to remove excess water. Samples were weighed and counted and the wet weight was recorded. Samples were also extracted in the laboratories of the Hurghada Marine Station of the National Institute of Oceanography and Fisheries.

**Extraction of bioactive material form marine organism**

Samples were extracted using a modified technique of Gomaa et al. (2000). Washed intact marine organism samples were weighed and equal amounts of absolute ethanol were added (1:1 w/v) and blended for 3 mins. The extracted samples were centrifuged at 2570 g for 10 min and the supernatant was removed and kept for further extraction. The residues were subjected to a second extraction with absolute ethanol and to a third extraction with 50% aqueous ethanol.

For defatting and partial depigmentation of sea anemone, *G. helianthus* ethanolic crude extract, as a step of purification, the supernatants were mixed with an equal volume with n-hexane. Defatting with n-hexane was repeated for 3 times. The residues were also blended with equal amounts (w/v) of n-hexane. Both aqueous ethanolic and hexane extracts were evaporated under reduced pressure at 40°C.

**Inhibitory effect on acetylcholinesterase (AchE)**

The *in vitro* AchE inhibition activity of the ethanolic crude extract and the other fractions was determined according to Ellman et al. (1961) using reagent kits purchased from Qunica Clinica Aplicada S.A., Spain.

**Isolation, purification and identification of *G. helianthus* AchE inhibitor**

Fractionation and purification of the *G. helianthus* ethanolic crude extract was carried out using different chromatographic techniques, as described in the following sections, followed the AchE inhibition activity guided fractionation protocol.

**Molecular weight exclusion ultrafiltration**

Crude extract of *G. helianthus* was filtered through membrane filters with cut off 10, 5, 3, 1 and 0.5 k Dalton (76 mm in diameter, Millipore Corporation, Bedford, MA, USA). Ultrafiltration was performed under pressure of nitrogen gas (40 kg.cm\(^{-2}\)).

**BioGel P2 gel filtration**

Gel filtration column chromatography was prepared using a 3.5 x 80 cm Omni LC column, packed with BioGel P2 (BioRad Laboratories, Richmond, CA, USA), to reach a bed height of 75 cm and a bed volume of 728 ml (Shimizu et al., 1975; Buckley et al., 1975; Hall 1982; Gomaa, 1990). 15 ml of Milli Q water was used to redissolve 5 g of the less than 0.5 k Dalton freeze-dried filtrate and then applied on the top of the BioGel P2 column. The column was eluted with 2 bed volumes of milli Q water using a peristaltic pump to provide a flow rate of 48 ml/h, and 5 ml fractions were collected.

Buckley spot plate technique (Buckley et al., 1975) was used to detect fluorescence and quenching activity of the collected fractions.
Table 1. Mobile phase gradient program used for isolation of AchE inhibitor.

| Time (min) | Water (%) | Methanol (%) | Acetonitrile (%) |
|-----------|-----------|--------------|------------------|
| 0         | 5         | 10           | 85               |
| 5         | 15        | 5            | 80               |
| 10        | 30        | 0            | 70               |
| 15        | 5         | 10           | 85               |

along with their reaction with ninhydrin. Aliquots of all the collected fractions, 5 μl each, were spotted on 10x10 cm silica gel TLC plates (precoated, type 60 F254, with fluorescent indicator, aluminum backed, E. Merck, Germany). Fluorescent and quenching activity was observed under long wave (366 nm) and short wave (254 nm) UV. Spot plates were sprayed with 1% ninhydrin in ethanol to detect the presence of the free α-amino groups (purple color) or the substituted α-amino group (yellow color) in the active fractions.

Active fractions, detected by in vitro AchE, were compared with any physical property that may appear by Buckley spot test. Percent bed volume was calculated for each fraction to correlate between the position(s) of the active fractions with the calculated percent bed volume.

**High-performance liquid chromatography (HPLC)**

Different HPLC techniques were tested to compare between the active and nonactive fractions to locate the peak that may account for the detected activity (AchE inhibition). The HPLC system used was Perkin-Elmer series 200 pump system equipped with a Perkin-Elmer series 200 UV absorbance detector set at wavelength of 203, 254 and 365 nm. Different mobile and stationary phases were used to find out the best method for the separation of the toxic compounds. Data were collected and integrated with a Totalchrom Navigator Chromatography Manager.

The analytical column chromatography, Spheri 5 silica (100 x 4.6 mm; 5 μm) (Applied Biosystems Inc. Foster City, A 94404 USA, Brownlee columns) was used to separate the AchE inhibitor. The mobile phase system (H₂O : MeOH : Acetonitrile) gradient program as shown in the Table 1. UV detector wavelength at 254 nm and the flow rate was 1 ml/min. Also, the semi preparative column hypersil HS/Silica (250 x 10 mm; 12 μm) Thermo Hypersil was used to obtain enough materials for the structure elucidation.

**Structure elucidation**

1D nuclear magnetic resonance (NMR) (¹H and ¹³C) analysis and EI-MS was carried out to elucidate the structure of the isolated bioactive compound.

**RESULTS AND DISCUSSION**

**Organism under study**

To identify the collected sea anemone samples from the Red Sea, specimens of the collected sea anemone species were sent to Prof. Dan Hartog of the National Museum of the Netherlands who was assigned by the British Museum of the Natural History as the best expert in this field. According to Prof. Dan Hartog, the scientific name of the species under study was confirmed as *G. helianthus or Entacmaea quadricolor*.

**Crude extract bioactivity**

According to Gomaa et al. (2000), the crude aqueous ethanolic extract of *G. helianthus* when intra peritoneally (i.p.) injected in the 20 g male mice showed symptoms of neurotoxicity. Gomaa et al. (2000) also showed that the part of the neurotoxicity observed in the mouth assay was due to reversible inhibition of the AchE in the brain and blood of the mice, which was confirmed by Gomaa and Aboul-Enien (2000) after using the in vitro AchE assay. The same assay was used in this study to track the bioactive compound during the bioassay-guided fractionation technique. AchE inhibition activity was confirmed in this study in the ethanolic crude extract of sea anemone *G. helianthus*. Different reports showed that sea anemones produce two types of protein toxins: neurotoxins, which act mainly on ion channels (Honma and Shiomi, 2005) and cytolysins (or actinoporins), which exhibit lytic activity on a variety of cells (Anderluh and Shiomi, 2005) and lytic activity on a variety of cells (Anderluh and Shiomi, 2005) and short wave (254 nm) UV. Spot plates were sprayed with 1% ninhydrin in ethanol to detect the presence of the free α-amino groups (purple color) or the substituted α-amino group (yellow color) in the active fractions.

Active fractions, detected by in vitro AchE, were compared with any physical property that may appear by Buckley spot test. Percent bed volume was calculated for each fraction to correlate between the position(s) of the active fractions with the calculated percent bed volume.

**Molecular weight exclusion AchE inhibition activity**

In this study, *G. helianthus* crude extract showed 91.9% inhibition in the AchE activity while the MW exclusion fractions showed lower percent inhibitions (Table 2). This table shows the effect of purification steps of *G. helianthus* on the activity of AchE. The more the purification, from the crude extract until less than 5 kD filtrate, the less the inhibition percent of AchE which means the presence of more than one AchE inhibitors in the crude extract. The only previous report detecting the AchE inhibitor in the extract of *G. helianthus* was that of Gomaa and Aboul-Enien (2000) and Gomaa et al. (2000). However, Mebs et al. (1983) did not detect any AchE inhibitors in *G. helianthus* collected from the Red Sea, the AchE inhibition activity detected in the less than 0.5kD filtrate indicated the presence of low MW active compound (less than 0.5 kD).
Table 2. In vitro AchE inhibition activity of the crude extract and different ultrafiltration fractions of *G. helianthus*.

| Sample            | AchE concentration (U/L) | Inhibition of AchE activity (%) |
|-------------------|--------------------------|---------------------------------|
| Control           | 4201.35                  | ---                             |
| Crude extract     | 340.65                   | 91.9                            |
| 10 kDa fraction   | 2573.8                   | 38.7                            |
| 5 kDa fraction    | 3255.1                   | 22.5                            |
| 3 kDa fraction    | 3180.4                   | 24.3                            |
| 1 kDa fraction    | 3146.8                   | 25.1                            |
| 0.5kDa fraction   | 2384.0                   | 43.3                            |

Biogel P2 fractions bioactivity

Using Biogel P2 fractionation and Buckley spot plate, the two distinct chemically active groups reported by Gomaa and Aboul-Enien (2000) were confirmed. The first group (65 to 72% bed volume) showed quenching under short wave (254 nm) while the second (72 to 77% bed volume) showed fluorescence activity under long wave (366 nm).

Only one of these two active groups showed *in vitro* AchE inhibition activity which confirmed the reported results obtained by Gomaa and Aboul-Enien (2000), while the second group showed no inhibition activity toward AchE. The AchE inhibition activity was detected along the purification steps from the crude to the 0.5 kDa filtrate until the first active group of the Biogel P2 fractions. While AchE inhibition activity was not detected in the second active group.

HPLC fractions activity

Different HPLC methods were tested to compare between the toxic (AchE inhibitor) and nontoxic fractions to locate the peak that may account for the detected AchE inhibition activity. The best result was obtained when the 254 nm wavelength was used. The first active group, after Biogel P2 fractionation, that showed the inhibition of AchE was tested by the HPLC technique to define the peak responsible for such activity. AchE inhibition and non-inhibition fractions were injected in the HPLC using a spheri 5 silica column (100 x 4.6 mm; 5 µm). Different mobile phase systems were used and the best resolution was achieved using a mixture of H₂O : methanol : acetonitrile in a gradient program. The peak responsible for AchE inhibition was observed at 8.5 min retention time. The active fractions of the Biogel P2 have a higher peak area at this retention time while the non-active fractions before and after these active ones showed no peaks or a very small one. Also, the peak areas were proportionally related to the activity level.

HPLC purification of AchE inhibitor

The HPLC technique described in the current study aided by semipreparative columns was used to isolate the target compounds in a pure form for the structure elucidation. Thus, the present study may be a step towards the discovery of novel compounds from the sea anemone *G. helianthus*.

Figures 1 to 4 illustrate the HPLC chromatograms of the different purification steps of *G. helianthus* ethanolic crude extract to isolate the AchE inhibitor compound through ultrafiltration steps reaching less than 0.5 kDa filtrate to the Biogel filtration and finally to obtain the pure bioactive compound from the HPLC aided by semi-preparative column. The percent of the peak area of the desired AchE inhibitor bioactive compound in the HPLC chromatograms were 29.9, 51.6, 75.8 and 98% in the crude extract, 0.5 kDa filtrate, AchE inhibition fraction after Biogel P2 and the isolated pure toxin after HPLC, respectively.

Structure elucidation of AchE inhibitor

Further purification on a semi-preparative HPLC led to the isolation of one major constituent in a pure form from the first active fraction after Biogel P2. The purity of the isolated bioactive compound was examined by TLC and HPLC where it gave only one spot on TLC and one peak in the HPLC chromatogram (Figure 4). One dimensional NMR analysis ¹H, ¹³C-NMR and DEPT were carried out to elucidate the structure of the AchE inhibitor isolated from the sea anemone, *G. helianthus* (Figure 5 to 7).

¹H NMR (600MHz, MeOH-d₄/Acetone-d₆) (Figure 5): δ ppm 8.74 (2H, br., J = 6.2 Hz, H-6/6′), 8.51 (2H, ddd=td, J = 7.8, 1.0Hz, H-4/4′), 8.07 (2H, br., J = 7.8 Hz, H-3/3′), 7.92 (2H, ddd=td, J = 6.2, 1.3 Hz, H-5/5′), 4.41 (6H, s, 2xN-Me)

¹³C NMR (150 MHz, MeOH-d₄/Acetone-d₆) (Figure 6): δ ppm 165.16 (C-2/2′), 146.98 (C-4/4′), 146.43 (C-6/6′), 127.57 (C-5/5′), 127.46 (C-3/3′), 47.41 (2xN-CH₃).

EI. MS, m/z (%) (Figure 8): 186.0 [M-C₇H₄]⁺, (40%); 171.0 [M-C₅H₃]⁺, (90%); 94.0 [M-C₇H₁₀N₂]⁺, (100%); 79.0 [M-C₇H₁₁N₂]⁺, (23%); 52.0 [M-C₇H₁₃N₃]⁺, (23%) As it was interpreted above, the isolated compound was expected to be 2-substituted pyridine-like structure on...
the basis of its splitting pattern, $\delta$- and $J$-values in the $^1$H NMR data (Figure 5). The presence of $N$-CH$_3$ functionality was deduced from the singlet at 4.41 ppm. Depending on its M.S fragments (Figure 8), the isolated bioactive compound was tentatively identified as a symmetric dimer of 2-amino-N-methylpyridine. The base peak (100%) at $m/z$ 94.0 mu was confirmative evidence for the homolytic cleavage of N-N bond. The fragment was diagnostic for the aminopyridine structure, which was followed by loss of 15 mu of NH to give a relatively weak abundant ion at $m/z$ 79.0 (23%). In its $^{13}$C-NMR spectrum (Figure 6), five characteristic carbon resonances were assigned to two pyridine moieties together with an aliphatic C-resonance at 47.41 of two N-CH$_3$ groups. Large down field shift (+16 ppm) of the C-2 to 165.16 ($\sim$149 in pyridine) was indicative of the attachment of amino group to C-2 and accumulation of positive charge on the ring nitrogen.

All $^{13}$C-resonances were assigned by aid of increment of subsistent additive rule and comparison with structure related compounds (Eberhard and Wolfgang, 1987). The appearance of only four $^1$H-resonances in the aromatic

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**Figure 1.** HPLC chromatogram of the sea anemone, *G. helianthus* ethanolic crude extract.
Figure 2. HPLC chromatogram of the less than 0.5 kD ultrafiltration fraction of sea anemone, *G. helianthus* ethanolic crude extract.

Figure 3. HPLC chromatogram of toxic fraction after BioGel P2 of the less than 0.5 kD ultrafiltration fraction of sea anemone, *G. helianthus*.
Figure 4. HPLC chromatogram of the pure isolated bioactive compound from sea anemone, *G. helianthus*.

Figure 5. $^1$H-NMR spectrum of the pure isolated sea anemone, *G. helianthus*, AchE inhibitor.
together with the aliphatic (CH$_3$) regions, respectively and six $^{13}$C-resonances (each of two equivalent carbons), was indicative of an asymmetric bis-(2-amino-N-methylpyridine) structure. Finally, DEPT spectrum (Figure 7) confirmed the suggested structure of the AchE inhibitor toxin as N,N'-bis-(1-methyl-pyridin-2-yl)-hydrazine (Figure 9) through the differentiation among C, CH and CH$_3$ resonances.
Figure 8. Mass spectrum of the pure isolated sea anemone, *G. helianthus*, AchE inhibitor.

Figure 9. Chemical structure of the AchE inhibitor compound isolated from sea anemone, *G. helianthus*.

**Conclusion**

The study of marine organisms as a source of biologically active compounds is considered a very productive field, having already led to the discovery of various new pharmacological tools and medicines. The inhibition activity towards AchE that was detected in crude extract of the sea anemone, *G. helianthus* and in the 0.5 kD fraction indicates a presence of low molecular weight active compound of less than 0.5 kD, this hypothesis coincided with the calculated molecular weight (216 D) after the structure elucidation of the isolated compound using mass spectrometer, NMR analysis (\(^{1}\)H and \(^{13}\)C-NMR) and DEPT. The isolation of a low molecular weight AchE inhibitor compound in a pure form with known chemical structure will be a step forward to discover a new drug.

**Conflict of Interests**

The authors have not declared any conflict of interests.

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