The Brazilian Amaryllidaceae as a source of acetylcholinesterase inhibitory alkaloids Jean Paulo de Andrade . Raquel B. Giordani . Laura Torras-Claveria .Natalia Bele'n Pigni . Strahil Berkov . Merce` Font-Bardia . Teresa Calvet . Eduardo Konrath . Kelly Bueno . Liana G. Sachett . Julie H. Dutilh . Warley de Souza Borges . Francesc Viladomat . Amelia T. Henriques . Jerald J. Nair . Jose' Angelo S. Zuanazzi . Jaume Bastida J. P. de Andrade · L. Torras-Claveria · N. B. Pigni · S. Berkov · F. Viladomat · J. J. Nair · J. Bastida Departament de Productes Naturals, Biologia Vegetal i Edafologia, Facultat de Farma'cia, Universitat de Barcelona, Av. Diagonal 643, 08028 Barcelona, Spain e-mail: jaumebastida@ub.edu J. P. de Andrade · R. B. Giordani · E. Konrath · K. Bueno · L. G. Sachett A. T. Henriques · J. A. S. Zuanazzi Faculdade de Farma'cia, Universidade Federal do Rio Grande do Sul, 2752 Ipiranga Av., Porto Alegre 90610-000, Brazil J. P. de Andrade · W. de Souza Borges Departamento de Qui'mica, Universidade do Espi'rito Santo, Vitoria, ES 29075-910, Brazil S. Berkov AgroBioInstitute, 8 Dragan Tzankov blvd., 1164 Sofia, Bulgaria M. Font-Bardia Centres Cientifics i Tecnologics, Universitat de Barcelona, Sole' i Sabaris 1-3, 08028 Barcelona, Spain T. Calvet Cristal.lografia, Mineralogia i Dipo'sits Minerals, Facultat de Geologia, Universitat de Barcelona, Marti' i Franque's s/ n, 08028 Barcelona, Spain J. H. Dutilh Departamento de Bota[^]nica, Universidade de Campinas, Cidade Universita'ria, Campinas 13083-970, Brazil

53 54	ABSTRACT:
55	Nine Brazilian Amaryllidaceae species were studied for their alkaloid composition and
56	acetylcholinesterase (AChE) inhibitory activity via GC-MS and a modified Ellman assay, respectively.
57	A total of thirty-six alkaloids were identified in these plants, of which Hippeastrum papilio and H.
58	glaucescens exhibited the highest galanthamine content and the best IC50 values against AChE.
59	Furthermore, Hippeastrum vittatum and Rhodophiala bifida also showed notable AChE inhibitory
60	effects. X-ray crystallographic data for four galanthamine-type compounds revealed significant
61	differences in the orientation of the N-methyl group, which are shown to be related to AChE inhibition.
62	
63	
64	

INTRODUCITON

_	_
o	o

67 68

69

70

71 72

73

74 75

76

77 78

79

80 81

82

83

84

85

86

87 88

89 90

91

65

The Amaryllidaceae alkaloids represent a large group of isoquinoline alkaloids derived from the common biogenetic precursor O-methylnorbelladine through oxidative phenolic coupling, leading to eight distinct structural-types (Bastida et al. 2006). The galanthamine- type skeleton has been the focus of numerous studies since the AChE inhibitor galanthamine was approved by the FDA for the clinical management of mild to moderate Alzheimer's disease (AD) (Maelicke et al. 2001). Although the chemical synthesis of galanthamine has been achieved on several occasions, natural sources still constitute the bulk of its commercial supply chain (Berkov et al. 2011). Apart from this, the other structural representatives of the Amaryllidaceae are known for a diverse array of biological activities including, antitumoral, antiviral, antiparasitic, anti-inflammatory, psychopharmacological and interactions with human cytochrome P450 3A4 (Vrijsen et al. 1986; C. itog lu et al. 1998; da Silva et al. 2006; McNulty et al. 2007, 2009; Zupko' et al. 2009; Giordani et al. 2010). These attributes have showcased the Amaryllidaceae as a promising resource for new and bioactive molecules. The high resolution power of the capillary column technique in gas chromatography (GC) together with the ready availability of libraries of electron impact mass spectrometry (EI-MS) data in the literature facilitate the rapid identification and quantification of known alkaloids. This has been shown to be particularly useful to studies of the Amaryllidaceae, extracts of which contain a large number of alkaloids (Kreh et al. 1995; Wagner et al. 2003). To this extent, several southern Brazilian Amaryllidaceae species have been examined for their alkaloid content and biological activity (da Silva et al. 2006, 2008; Pagliosa et al. 2010; Giordani et al. 2011a, b; de Andrade et al. 2011). In the present study, a GC-MS analysis was undertaken on nine Amaryllidaceae species which allowed for the identification of thirty-six alkaloids belonging to seven skeleton-types. Furthermore, an AChE inhibitory activity assay was carried out with both isolated compounds and alkaloid-rich fractions. In addition, Xray crystallographic analysis was carried out on some galanthamine derivatives, providing insights to the structural features attending AChE activity.

MATERIALS AND METHODS

93

92

94 Chemicals

- 95 Galanthamine (27) and 11b-hydroxygalanthamine (32) used for X-ray crystallography were previously
- obtained from Hippeastrum papilio (de Andrade et al. 2011). Sanguinine (28) and narwedine (31) were
- 97 obtained in previous works from Crinum kirkii Chemicals Galanthamine (27) and 11b-
- 98 hydroxygalanthamine (32) used for X-ray crystallography were previously obtained from Hippeastrum
- 99 papilio (de Andrade et al. 2011). Sanguinine (28) and narwedine (31) were obtained in previous works
- from Crinum kirkii (Machocho et al. 2004) and Leucojum aestivum (Berkov et al. 2008a), respectively.
- MeOH (HPLC grade), CHCl3, Me2CO, H2SO4 and NH4? (analytical grade) were purchased from SDS
- 102 (France). Acetylthiocholine iodide (ATCI), acetylcholinesterase (AChE) from electric eels (type VI-S
- lyophilized powder), and 5,5 V-dithiobis[2-nitrobenzoic acid] (DTNB) were obtained from Sigma-
- Aldrich Chemie (Steinheim, Germany). The n-hydrocarbon mixture (C9–C36, Restek, Cat no. 31614)
- was supplied by Teknokroma (Spain). Galanthamine (purity[99 %) used for the calibration curves was
- previously obtained by the authors, and codeine (purity C 99 %) used as internal standard was purchased
- from Sigma Aldrich (St. Louis, MO, USA).

108

- 109 Plant material
- 110 The species H. papilio (Ravenna) Van Scheepen (bulbs and leaves, UFRGS-ICN 149428), Hippeastrum
- vitattum (L'He'r.) Herb. (bulbs, UFRGS-ICN 8889), Hippeastrum striatum (Lam.) Moore (bulbs,
- 112 UFRGS-ICN 9549), Hippeastrum morelianum Lem. (bulbs, UNICAMP-UCE 14351), Hippeastrum
- santacarina (Traub) Dutilh (bulbs, UFRGS-ICN 149429), Hippeastrum breviflorum Herb. (bulbs,
- UFRGS-ICN 9190), Hippeastrum glaucescens (Mart.) Herbert (bulbs and leaves, UFRGS-ICN 8894),
- Hippeastrum psittacinum Herb. (bulbs and leaves, UNICAMP–UCE 143513) and Rhodophiala bifida
- 116 (Herb.) Traub (bulbs, UNICAMP–UCE 136352) were collected and the extracts obtained according to
- previously described methods (Castilhos et al. 2007; da Silva 2005; da Silva et al. 2008; Pagliosa et al.
- 2010; Giordani et al. 2011a, b; de Andrade et al. 2011; Sebben 2005).

119

- 120 Sample preparation
- The plant material (1 g) was crushed and extracted by stirring at rt with MeOH (3 9 50 ml), the
- 122 combined macerate filtered and evaporated to dryness under reduced pressure. The crude extract was
- acidified to pH 2 with 2 % H2SO4, neutral material removed using Et2O (3 9 25 ml). The aqueous
- phase was then basified up to pH 11 with NH3 (25 %, v/v) and extracted with CHCl3 (3 9 25 ml) to
- afford the chloroform extract.

126

127

- 129 GC–MS and identification of alkaloids
- The chloroform extract (300 ll) was filtered and then used for subsequent GC–MS analysis. EI–MS
- spectra were obtained on an Agilent 6890 N GC 5975 inert MSD operating in EI mode at 70 eV (Agilent
- Technologies, Santa Clara, California, USA) utilizing a DB-5 MS column (30 m 9 0.25 mm 9 0.25 lm,
- 133 Agilent Technologies) with an injector temperature of 280 \(\textstyle \) C. The temperature program was as
- 134 follows: 100–180 \[C at 15 \[Cmin-1, 1 min hold at 180 \[C and 180–300 \[C at 5 \[Cmin-1 and 10 min \]
- hold at 300 \(\bigcap \) C. The flow rate of carrier gas (Helium) was 0.8 ml min-1 and a split ratio of 1:20 was
- followed. The alkaloids were identified by comparing their GC-MS spectra and Kovats retention indices
- 137 (RI) with our in-house library database. This library has been continually updated and reviewed with
- alkaloids isolated by our group and identified using other spectroscopic techniques such as NMR, UV,
- 139 CD and MS. Mass spectra were deconvoluted using AMDIS 2.64 software (NIST). Kovats retention
- indexes (RI) of the compounds were recorded with standard calibration of an n-hydrocarbon mixture
- 141 (C9–C36).
- The proportion of each individual component in the alkaloid fractions analysed by GC–MS (Table 1) is
- expressed as a percentage of the total alkaloids (TIC—total ion current). The area of the GC–MS peak
- depends not only on the concentration of the corresponding compound but also on the intensity of its
- mass spectral fragmentation. Although data given in Table 1 do not express a real quantification, they
- can nevertheless be used for a relative comparison of the alkaloids.
- 147 Quantification of galanthamine in H. papilio The quantification was performed in triplicate using 50 mg
- of dried material (leaves and bulbs, separately) and codeine as i.s. (50 lg) in screw-top 2.0 ml Eppendorf
- tubes. The maceration procedure was carried out with 1 ml of MeOH adjusted to pH 8 with NH3 (25 %,
- 150 v/v). After 2 h of extraction at room temperature assisted by 15 min ultrasonic baths every 30 min, the
- samples were centrifuged at 10,000 rpm for 2 min. An aliquot of 500 ll of methanolic macerate was
- acidified with 500 ll of H2SO4 (2 %, v/v) and neutral material removed with chloroform (2 9 500 ll).
- The aqueous fraction was then basified with 200 ll of NH3 (25 %, v/v) and alkaloids extracted with
- 154 CHCl3 (3 9 500 ll). Finally, the purified alkaloid extract was dried under N2 and redissolved in 100 ll of
- 155 CHCl3 for GC–MS analysis. The GC–MS conditions were the same used for the alkaloid-rich extract
- 156 (Section GC–MS and identification of alkaloids).
- Recovering and repeatibility of the extraction The extraction recovery was performed as described
- above by adding 50, 300 and 500 lg of galanthamine to the dry plant sample (50 mg of powdered bulbs
- and leaves of H. papilio) before the extraction and purification. Intraday (n = 4) and interday (n = 8)
- repeatability was calculated with 50 mg of dried powdered bulbs of H. papilio, extracted, purified and
- analysed via GC–MS on two different days according to Berkov et al. (2008b).
- 163 Samples for X-ray

- Narwedine (31) and 11b-hydroxygalanthamine (32) were dissolved in CHCl3 under a pentane

- MeOH:EtOH mixture (1:1, v/v) under a pentane atmosphere and left in the freezer (less than 5 \(\text{\mathbb{C}} \) for
- two weeks. Galanthamine (27) was dissolved in Me2CO and left in the freezer for a week. Suitable
- 168 crystals for X-ray analysis were preselected under a light microscope. The crystallographic data of 27
- and 31 were in agreement with those previously reported (Carrol et al. 1990; Hemetsberger et al. 2004).
- 170
- 171 X-Ray analysis for sanguinine (28)
- A translucent prism-like specimen of sanguinine with the dimensions 0.192 mm 9 0.278 mm 9 0.457
- mm was used for X-ray crystallographic analysis. First, the X-ray intensity data were determined, with a
- total of 171 frames collected at an exposure time of 1.71 h. The frames were integrated with the Bruker
- 175 SAINT software package using a narrow-frame algorithm. The integration of the data using a
- monoclinic unit cell yielded a total of 19,735 reflections to a maximum h angle of 30.67 $\boxed{0.70 \text{ A}}$ (0.70 A°
- 177 resolution), of which 7366 were independent (average redundancy 2.679, completeness = 94.1 %,Rint =
- 4.80 %, Rsig = 5.54 %) and 6597 (89.56 %) were greater than 2r(F2). The final cell constants of a =
- 9.227(6) A $^{\circ}$, b = 15.095(8) A $^{\circ}$, c = 9.750(5) A $^{\circ}$, b = 102.28(3) , volume = 1327.(2) A $^{\circ}$
- 3, are based upon the refinement of the XYZcentroids of 142 reflections above 20 r(I) with
- 4.944 \2h\49.15 . Data were corrected for absorption effects using the multi-scan method (SADABS).
- The ratio of minimum to maximum apparent transmission was 0.757.
- The structure was solved and refined using the Bruker SHELXTL Software Package, with Z = 2 for the
- formula unit, C16H19NO3. The final anisotropic full-matrix least-squares refinement on F2 with 375
- variables converged at R1 = 4.08 %, for the observed data and wR2 = 10.17 % for all data. The
- goodnessof- fit was 1.047. The largest peak in the final difference electron density synthesis was 0.382
- 187 e-/A $^{\circ}$ 3 and the largest hole was -0.274 e-/A $^{\circ}$ 3 with an RMS deviation of 0.058 e-/A $^{\circ}$ 3. On the basis of
- the final model, the calculated density was 1.367 g/cm3 and F(000), 584 e-.
- 189
- 190 X-Ray analysis for 11b-hydroxygalanthamine (32)
- A prismatic crystal (0.1 9 0.09 9 0.08 mm) was selected and mounted on a MAR345 diffractometer with
- an image plate detector. Unit-cell parameters were determined from 107 reflections (3\h\318) and
- refined by the least-squares method. Intensities were collected with graphite monochromatized Mo Ka
- radiation. 8529 reflections were measured in the range 2.44 B h B 24.10, 2419 of which were non-
- equivalent by symmetry (Rint(on I) = 0.045). 2135 reflections were assumed as observed applying the
- 196 condition I[2r (I). Lorentz-polarization was considered, but no absorption corrections were made. The
- structure was solved by direct methods, using the SHELXS computer program (Sheldrick 2008) and
- refined by the full-matrix least-squares method with the SHELX97 computer program (Sheldrick 2008),
- using 8529 reflections, (very negative intensities were not assumed). The function minimized was R w
- 200 ||Fo|2 |Fc|2|2, where w = [r2(I)? (0.0683P)2]-1, and P = (|Fo|2? 2|Fc|2)/3, f, f' and f'' were taken
- from International Tables of X-Ray Crystallography (1974). All H atoms were computed and refined,
- using a riding model, with an isotropic temperature factor equal to 1.2 times the equivalent temperature

factor of the atom which are linked. The final R(on F) factor was 0.047, wR(on |F|2) = 0.117 and 203 goodness of fit = 1.069 for all observed reflections. The number of refined parameters was 200. Max. 204 shift/esd = 0.00, mean shift/esd = 0.00. Max. and min. peaks in final difference synthesis were 0.395 and 205 $-0.169 \text{ e.A}^{\circ}$ -3, respectively. 206 207 208 AChE inhibitory activity 209 The assay for measuring AChE inhibitory activity was performed as described by Lo'pez et al. (2002). 210 Galanthamine hydrobromide was used as a positive control. A solution of the initial alkaloid-rich extract (chloroform fraction) at 1 mg/ml was taken up in MeOH and diluted further with phosphate buffer to 211 212 give 100, 10, 1, 0.1, 0.01, 0.001 lg/ml solutions. Only IC50 values less than 100 lg/ml were considered. Compounds 27, 28, 31, and 32 were used in dilutions at the range of 10-8 to 10-3 M. Dilutions at 10-4 213 M were prepared in MeOH and further dilutions were carried out using phosphate buffer. IC50 of all 214 extracts/compounds were measured in triplicate and the results are presented as a mean \pm standard 215 deviation using the software package Prism (Graph Pad Inc., San Diego, USA). 216 217

219 Results and discussion 220 221 GC-MS results 222 GC-MS analysis has here proved to be a robust and efficient technique for the rapid identification and 223 quantification of a large number of alkaloids from Amaryllidaceae plant extracts. In this study, nine 224 Brazilian species were analysed and thirty-six compounds belonging to seven skeleton-types were 225 identified (see Fig. 1; Table 1). Lycorine- and homolycorine-type: an 'ortho-para' phenolic coupling 226 As the lycorine skeletal-type is widely distributed in the Amaryllidaceae, it was surprising to find few 227 representatives of this group in the Hippeastrum species and Rodophiala bifida surveyed. The alkaloid 228 229 lycorine (3) is known to be poorly soluble in both CHCl3 and MeOH, which impedes its correct 230 quantification by GC-MS (de Andrade et al. 2012). This might explain the low relative percentage observed for H. santacatarina (19.18 %, see Table 1), in contrast with a recent study of the same species, 231 232 in which it was isolated as the main compound (Giordani et al. 2011b). Overall, homolycorine-type alkaloids were observed in higher variety and quantity, indicating that conversion of lycorine-to 233 234 homolycorine-type alkaloids is an active chemical transformation in these species. 235 236 Crinine-, haemanthamine-, tazettine-, narciclasine- and montanine-type alkaloids: a 'para-para' phenolic 237 coupling 238 A major mechanistic consideration in the biosynthesis of Amaryllidaceae alkaloids is 'para-para' 239 coupling, since it gives rise to five distinct skeleton-types. The crinine-type skeleton is uncommon in the 240 genus Hippeastrum and the absolute configuration of its 5,10b-ethano bridge is ratified only by CD 241 spectra or X-ray crystallography (Wagner et al. 1996). As shown in Table 1 and Fig. 1, the 5,10b-242 ethanophenanthridinealkaloids described in this study possess the haemanthamine-type skeleton as previously confirmed (da Silva et al. 2008; de Andrade et al. 2011; Giordani et al. 2011a). 243 244 With respect to the tazettine skeleton, there are important features concerning epimerisation at C-3. 245 Duffield et al. (1965) showed that the stereochemistry of the substituent at C-3 effects marked variations in the relative abundance of ions in EI-MS spectra. The b-configuration of the methoxyl group at C-3 246 247 facilitates a Retro-Diels-Alder (RDA) process in ring-B and loss of the neutral fragment [C5H8O], yielding diagnostic ion peaks at 247 and m/z 231 (M-84) for tazettine (19) and deoxytazettine (17), 248 249 respectively. The fragment ion at m/z 70 is a small peak for both epimers (Duffield et al. 1965). As such, 250 the ion peak at m/z 70 for criwelline and 16 is much more pronounced than those observed in 17 and 19, 251 indicating that compound 16 is the 3-epideoxytazettine variant. The a-configuration of the 3-OMe 252 substituent also induces a RDA fragmentation process, but in this case with the loss of the [C4H8N]?

fragment, while the m/z 70 ion peak abundance establishes the C-3 configuration in tazettine derivatives

253

254

(Duffield et al. 1965).

In general, montanine-type alkaloids are sparsely encountered and are thus poorly represented in the 255 256 Amaryllidaceae. However, montanine (25) was here found as the main constituent in H. vittatum and R. 257 bifida, while trisphaeridine (23) was the only representative of the narciclasine-type skeleton detectable as a minor compound or in trace amounts in most species (Table 1). Trisphaeridine has been considered 258 259 a catabolic product (Bastida et al. 2006) and this hypothesis is supported by its presence in many species 260 but hardly ever as the main alkaloid. Galanthamine-type alkaloids: a 'para-ortho' phenolic coupling 261 Galanthamine-type compounds were found mainly in H. papilio and H. glaucescens, with galanthamine 262 (27) being the main constituent in both cases (Table 1). Galanthamine was previously detected in H. 263 papilio (de Andrade et al. 2011), but it is here reported for the first time in H. glaucescens. The 264 remaining galanthamine-type representatives were detected in both species, but to a lesser extent. Miscellaneous alkaloids Ismine (34) and galanthindole (35) were identified in H. breviflorum, H. 265 morelianum, H. psittacinum and H. glaucescens. Alkaloid 34, like 23, is also considered a catabolic 266 product arising from the haemanthaminetype skeleton (Bastida et al. 2006). Galanthindole (35) and 267 lycosinine B (36) have been considered representatives of a new skeleton containing a non-fused indole 268 ring (U" nver 2007), although the possibility that they are artifacts of homolycorine- or tazettine-type 269 270 derivatives cannot be overlooked. 271

272

- Galanthamine quantification
- 273 H. papilio and H. glaucescens showed highest levels of galanthamine by GC-MS (Table 1) and the
- 274 availability of H. papilio allowed the accurate quantification of galanthamine content from dried plant
- 275 material. Bulbs and leaves exhibited values of 0.51 % (± 0.012) and 0.33 % (± 0.007), respectively (mg
- 276 GAL/100 mg DW). These values are larger than those observed for Galanthus and Leucojum species
- 277 used commercially by pharmaceutical companies for extraction of galanthamine (Cherkasov and
- 278 Tolkachev 2002; Berkov et al. 2008b, 2009). The extraction recovery was 95 % (RSD 1.73 %), 93 %
- (RSD 2.20 %) and 91 % (RSD 0.81 %) for 50, 300 and 500 lg of added galanthamine, respectively. 279
- 280 Intra-day repeatability (n = 4) expressed as RSD was determined as 1.60 for the first day and 2.21 for
- 281 the second, while inter-day repeatability (n = 8) was 2.94 with adequate values of precision (RSD\5 %).

- AChE inhibitory assay for alkaloid-rich extracts 283
- The results from the microplate AChE inhibition assay of plant extracts are shown in Table 2. H. papilio 284
- 285 and H. glaucescens presented the lowest IC50 values as determined via the Ellman method (Section ChE
- inhibitory activity). These are stronger activities than those observed for Galanthus elwesii and G. 286
- 287 nivalis (at 0.1 and 10 lg/ml) as well as Leucojum aestivum (at 10 lg/ml) (Berkov et al. 2008c). The
- possibility of false-positive results in the AChE inhibitory activity values due to chemical inhibition 288
- 289 (Rhee et al. 2003) should not be ruled out.
- H. vittatum and R. bifida, in which elevated levels of montanine were detected, also exhibited notable 290
- 291 AChE inhibitory effects (Table 2). Montanine (25) has previously demonstrated remarkable activity

against AChE obtained from rat brain, with more than 50 % inhibition at 1 mM (Pagliosa et al. 2010). 292 293 These results, together with psychobiological activities reported earlier for montanine (da Silva et al. 2006), reinforce the potential of montanine-type derivatives as therapeutic candidates for AChE 294 inhibition or other functions related to the central nervous system (da Silva et al. 2006; Pagliosa et al. 295 296 2010). 297 X-ray crystallography and AChE assay for galanthamine-derivatives 298 299 In agreement with previous reports (Lo'pez et al. 2002; Berkov et al. 2008c), galanthamine (27) and sanguinine (28) (Fig. 4) were the most active AChE inhibitory alkaloids (IC50s 0.35 and 0.06 lM, 300 301 respectively). Narwedine (31) and 11b-hydroxygalanthamine (32) showed IC50 values of 9.38 and 3.49 IM, respectively. Some studies have been carried out to understand the binding of galanthamine and 302 galanthamine-type alkaloids at the AChE active site (Bartolucci et al. 2001; Greenblatt et al. 1999). 303 Although these have provided useful insights to the binding of the aromatic methoxyl group, the furan 304 and cyclohexene rings as well as the 3-hydroxyl substituent, the effects of the N-methyl group remain 305 largely unresolved. However, it is noteworthy that galanthamine adopted the same conformation at the 306 active site gorge as that determined by X-ray crystallographic analysis (Bartolucci et al. 2001; Carrol et 307 al. 1990). 308 The X-ray data obtained for galanthamine (27) and narwedine (31) are in agreement with previously 309 310 published work (Carrol et al. 1990; Hemetsberger et al. 2004). The X-ray data for sanguinine (28)1 and 311 11b-hydroxygalanthamine (32)2 are reported here for the first time. Interestingly, narwedine (31) and 312 11bhydroxygalanthamine (32) (Fig. 2) showed an axial orientation for the NMe group, opposite to that 313 seen for galanthamine. Sanguinine (28) (Fig. 3), the most potent AChE inhibitor known from the Amaryllidaceae, exhibited both orientations for the NMe group with 50 % of the molecules having the 314 315 NMe group in the axial orientation and the other 50 % with the equatorial orientation. AChE inhibition curves together with the X-ray structures of all tested galanthamine alkaloids are shown in Fig. 4. 316 317

CONCLUSIONS

320

321

322

323324

325

326 327

328

329

330

331

332333

334

335

336337

319

Some indigenous Brazilian species are shown to produce high quantities of the AChE inhibitors galanthamine and montanine. Following the approval of galanthamine by the FDA for clinical management of AD, galanthamine-type alkaloids have been the most commonly studied constituents of the Amaryllidaceae. Herein is reported for the first time the high levels of galanthamine detected via GC-MS in H. glaucescens. Galanthamine levels in leaves and bulbs of H. papilio were higher than those found in Leucojum, Galanthus and Narcissus, species traditionally used for commercial exploitation (Berkov et al. 2009). In addition, H. papilio and H. glaucescens extracts showed the lowest IC50 AChE inhibition values. Since evidence from docking studies of galanthamine analogs are inconclusive, further investigation is required to clarify the role of N-methyl orientation at the AChE active site gorge (Bartolucci et al. 2001). Galanthamine has the N-methyl group in an equatorial disposition and showed better AChE inhibitory activity than narwedine and 11b-hydroxygalanthamine, wherein the N-methyl group is axially-orientated. Chlidanthine also displays an axial orientation for the N-methyl group and exhibits noticeably lower AChE inhibition (IC50 24.1 lM) (Reyes-Chilpa et al. 2011). However, sanguinine exhibits the best IC50 inhibition values and has the N-methyl group in both axial and equatorial orientations. It is known that N-methyl conformers interchange rapidly in the naturally bound ligand, thereby restricting N-methyl orientation to a secondary role in new drug design. Nevertheless, further protein-ligand crystallography and protein-ligand docking studies should clarify the exact role of N-methyl orientation in galanthamine-type alkaloids.

338339

341	References	
342		
343	Bartolucci C,	Perola M, Christian P et al (2001) Three-dimensional structure of a complex of
344	galantha	mine (Nivalin) with acetylcholinesterase from Torpedo californica: implications for the
345	drug des	sign of new anti-Alzheimer drugs. Proteins 42:182-191
346	Bastida J,	Lavilla R, Viladomat F (2006) Chemical and biological aspects of Narcissus alkaloids.
347	In: Cord	lell GA (ed) The alkaloids, vol 63. Elsevier Inc, Amsterdam, pp 87–179
348	Berkov S,	Codina C, Viladomat F et al (2008a) N-Alkylated galanthamine derivatives: potent
349	acetylch	olinesterase inhibitors from Leucojum aestivum. Bioorg Med Chem Lett 18:2263–2266
350	Berkov S,	Bastida J, Nikolova M et al (2008b) Analysis of galanthamine-type alkaloids by
351	capillary	gas chromatography- mass spectrometry in plants. Phytochem Anal 19:285-293
352	Berkov S,	Bastida J, Nikolova M et al (2008c) Rapid TLC/GCMS identification of
353	acetylch	olinesterase inhibitors in alkaloids extracts. Phytochem Anal 19:411-419
354	Berkov S,	Georgieva L, Kondakova V et al (2009) Plant source of galanthamine: phytochemical
355	and biot	echnological aspects. Biotechnol Biotechnol Equip 23:1170-1176
356	Berkov S,	Bastida J, Viladomat F et al (2011) Development and validation of a GC-MS method for
357	a rapid o	determination of galanthamine in Leucojum aestivum and Narcissus ssp.: A metabolomic
358	approac	h. Talanta 83:1455–1465
359	Carrol P, F	urst GT, Han SY et al (1990) Spectroscopic studies of galanthamine and galanthamine
360	methiod	ide. Bull Soc Chim Fr 127:769–780
361	Castilhos TS,	Giordani RB, Henriques AT et al (2007) Avaliac ,a o in vitro das atividades
362	antiinfla	mato'ria, antioxidante e antimicrobiana do alcalo'ide montanina. Rev Bras Farmacogn
363	17:209–	214
364	Cherkasov OA	, Tolkachev ON (2002) Narcissus and other Amaryllidaceae as sources of galanthamine.
365	In: Hank	ss G (ed) Medicinal and aromatic plants-industrial profiles: Narcissus and Daffodil, the
366	genus N	arcissus. Taylor and Francis, London and New York, pp 242–255
367	C, itogʻlu G, T	anker M, Gu"mu"s,el B (1998) Antiinflamatory effects of lycorine and haemanthidine.
368	Phytothe	er Res 12:205–206
369	da Silva AFS	(2005) Hippeastrum vittatum (L'He'r) Herbert e Hippeastrum striatum (Lam.) Moore:
370	Ana'lise	quı'mica e avaliac,a~o biolo'gica dos alcaloides isolados. Dissertation, Universidade
371	Federal	do Rio Grande do Sul
372	da Silva AFS,	de Andrade JP, Bevilaqua LR et al (2006) Anxiolytic-, antidepressant- and
373	anticonv	rulsivant-like effects of the alkaloid montanine isolated from Hippeastrum vittatum.
374	Pharmac	col. Biochem Behav 85:148–154

e Andrade JP, Machado KRB et al (2008) Screening for cytotoxic activity of

extracts and isolated alkaloids from bulbs of Hippeastrum vittatum. Phytomedicine 15:882-885

da Silva AFS, d

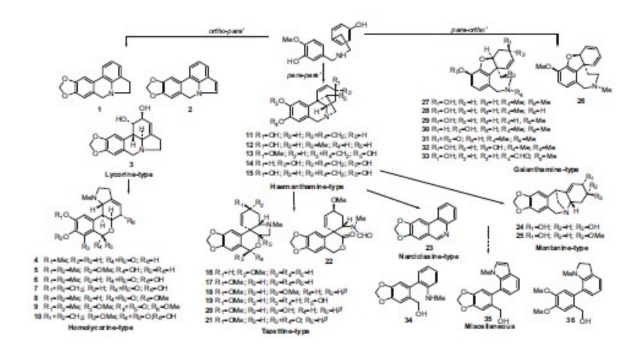
375

Berkov S, Viladomat F et al (2011) Alkaloids from Hippeastrum papilio. 377 de Andrade JP, 378 Molecules 16:7097-7104 de Andrade JP, Pigni NB, Torras-Claveria L et al (2012) Bioactive alkaloids from Narcissus 379 broussonetii: mass spectral studies. J Pharm Biomed Anal 70:13-25 380 381 Duffield AM, Aplin RT, Budzikiewicz H et al (1965) Mass spectrometry in structural and stereochemical problems. LXXXII. A study of the fragmentation of some Amaryllidaceae 382 alkaloids. J Am Chem Soc 87:4902-4912 383 384 Giordani RB, Vieira PB, Weizenmann M et al (2010) Candimine-induced cell death of the 385 amitochondriate parasite Trychomonas vaginalis. J Nat Prod 73:2019–2023 386 de Andrade JP, Verli H et al (2011a) Alkaloids from Hippeastrum morelianum Lem. 387 (Amaryllidaceae). Magn Reson Chem 49:668–672 388 Giordani RB, Vieira PB, WeizenmannMet al (2011b) Lycorine induces cell death in the amitochondriate parasite, Trichomonas vaginalis, via an alternative non-apoptotic death pathway. 389 Phytochemistry 72:645–650 390 391 Greenblatt HM, Kryger G, Lewis T et al (1999) Structure of acetylcholinesterase complexed with (-)-galanthamine at 2.3 A ° resolution. FEBS Lett 463:321–326 392 Treu M, Jordis U et al (2004) 1-methylgalanthamine derivatives. Monatsh Chem 393 Hemetsberger M, 135:1275-1287 394 395 International Tables of X-Ray Crystallography (1974) Kynoch Press. Birmingham Matusch R, Witte L (1995) Capillary gas chromatography-mass spectrometry of 396 Kreh M, 397 Amaryllidaceae alkaloids. Phytochemistry 38:773–776 Lo'pez S, Bastida J, Viladomat F et al (2002) Acetylcholinesterase inhibitory activity of some 398 399 Amaryllidaceae alkaloids and Narcissus extracts. Life Sci 71:2521-2529 Machocho AK, Bastida 400 J, Codina C et al (2004) Augustamine type alkaloids from Crinum kirkii. Phytochemistry 401 65:3143-3149 402 Maelicke A, Samochocki M, Jostock R et al (2001) Allosteric sensitization of nicotinic receptors by 403 galantamine, a new treatment strategy for Alzheimer's disease. Biol Psychiatry 49:279-288 McNulty J, Nair JJ, Codina C et al (2007) Selective apoptosis inducing activity of crinum-type 404 Amaryllidaceae alkaloids. Phytochemistry 68:1068–1074 405 406 McNulty J, Nair JJ, Singh M et al (2009) Selective cytochrome P450 3A4 inhibitory activity of Amaryllidaceae alkaloids. Bioorg Med Chem Lett 19:3233-3237 407 408 Monteiro SC, Silva KB et al (2010) Effect of isoquinoline alkaloids from two Hippeastrum species on in vitro acetylcholinesterase activity. Phytomedicine 17:698-701 409 410 Reyes-Chilpa R, Berkov S, Herna'ndez-Ortega S et al (2011) Acetyl-cholinesterase inhibiting alkaloids from Zephyranthes concolor. Molecules 16:9520-9533 411 412 Rhee IK, van Rijn RM, Verpoorte R (2003) Qualitative determination of false-positive effects in

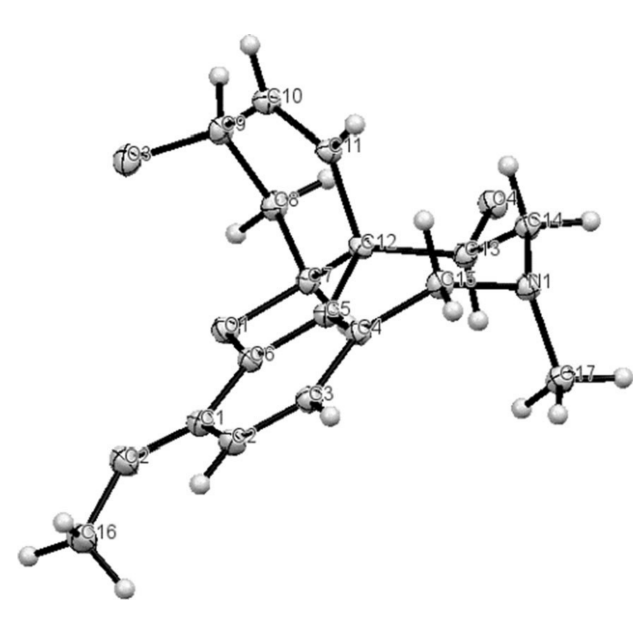
the acetylcholinesterase assays using thin layer chromatography. Phytochem Anal 14:127–131

414	Sebben C	(2005) Investigac, ao qui mica e biolo gica em Hippeastrum breviflorum Herb.
415	(Amaryl	lidaceae). Dissertation, Universidade Federal do Rio Grande do Sul
416	Sheldrick GM	(2008) A program for automatic solution of crystal structure refinement. Acta
417	Crystallo	ogr A 64:112–221
418	U"nver N	(2007) New skeletons and new concepts in Amaryllidaceae alkaloids. Phytochem Rev
419	6:125–13	35
420	Vrijsen R,	Berghe DAV, Vlietinck AJ et al (1986) Lycorine: an eukaryotic terminator inhibitor? J
421	Biol Che	em 261:505–507
422	Wagner J,	Pham HL, Do"pke W (1996) Alkaloids from Hippeastrum equestre Herb5. Circular
423	dichroisi	n studies. Tetrahedron 52:6591–6600
424	Wagner C,	Sefkow M, Kopka J (2003) Construction and application of a mass spectral and
425	retention	time index database generated from plant GC/EI-TOF-MS metabolite profiles.
426	Phytoche	emistry 62:887–900
427	Zupko' I,	Re'thy B, Hohmann J et al (2009) Antitumor activity of alkaloids derived from
428	Amaryll	idaceae species. In Vivo 23:41–48

430	Legends to figures
431	
432	Fig. 1 Alkaloids found in the Brazilian species
433	
434	Fig. 2 ORTEP projection of 11b-hydroxygalanthamine (32)
435	
436	Fig. 3 Top, a view of the molecular structure of compound 3. Bottom, the labeled core of the cubane.
437	Bonds depicted in orange correspond to the short Cu–O distances inside the {Cu4O4} cage and the
438	dashed red bonds show the H-bonds involving the coordinated water molecule
439	
440	Scheme 3 ORTEP projection of sanguinine (28)
441	
442	Fig. 4 Acetylcholinesterase inhibition curve and X-ray structures of sanguinine, galanthamine, 11b-
443	hydroxygalanthamine and narwedine showing the N-methyl orientation
444	
445	
446	



452 FIGURE 2



458 FIGURE 3

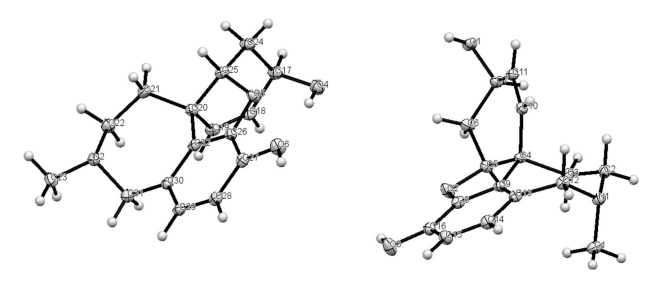
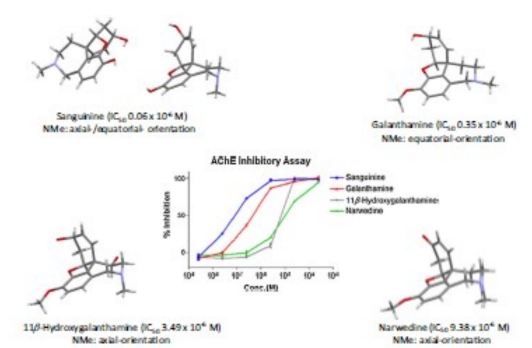


FIGURE 4 462

463

464



NMe: axial-orientation

Compound	Z	M+	Rel. int. (%)	H. striatum Bulbs	H. vinatum Bulbs	H. brevifionem Bulbs	H. strianım H. vinatum H. broviforum H. morelianum H. papilio H. papilio Bults Bults Bults Bults Leaves	H. papilio Bulbs	H. popilio Leaves
Anhydrolycorine (1)	2501	251 (43)	250 (100), 192 (13), 191 (11), 165 (4), 164 (3), 139 (2), 124 (7)	1	tt.	1	1	1	1
11,12-Dehydroath ydrolycorine (2)	3606	249 (60)	248 (100), 191 (10), 190 (24), 189 (7), 163 (7), 95 (17)	E		1.17	c	1	į.
Lycorine (3)	2746	287 (31)	286 (19), 268 (24), 250 (15), 227 (79), 226 (100), 211 (7), 147 (15)	ь	09'0		1	1	1
8-O-Demethylhomolycotine (4)	284	301(-)	192 (0.5), 164 (2), 110 (8), 109 (100), 108 (23), 94 (3), 82 (3)	1.	1		1	1	1
Norinine (5)	2476	347 (-)	330 (7), 329 (3), 236 (1), 221 (9), 191 (2), 109 (100), 94 (2)	1	1	9	1.86	1	1
2x-Hydroxytromolycorine (6)	2970	331(-)	178 (3), 126 (8), 125 (100), 124 (7), 96 (31), 94 (4)	1	1	ī	ь	1	1
Hippeastrine (7)	2917	315(-)	190 (1), 162 (4), 134 (2), 125 (100), 96 (40), 82 (3)	1	1	i	1	1	1
2a-Methoxyhomoly corine (8)	2870	345(-)	178 (5), 140 (11), 139 (100), 124 (67), 94 (7), 77 (5)	i i	1	1	ь	1	1
2a,7-Dimethoxyhomolycorine (9)	2962	375(-)	221 (2), 140 (9), 139 (100), 125 (6), 124 (55), 94 (4)	E	i	r.	b	į.	i
Candimine (10)	3000	345(-)	192 (1), 177 (2), 163 (1), 147 (1), 125 (100), 96 (90), 82 (2)				b		
Vitatine (11)	2472	271 (100)	272 (20), 252 (35), 199 (70), 187 (61), 173 (22), 115 (28)	1	1.23	1	1	1	b
8-O-Demethylmarifidine (12)	2510	273 (100)	274 (17, 230 (24), 201 (83), 189 (52), 175 (20), 115 (18)	1	1.62	,	1	5	1
Haemanthamine (13)	2641	301 (13)	272 (100), 240 (16), 211 (13), 199 (7), 181 (21), 153 (8)	ı.	1		1	91'91	21.60
Hamayne (14)	5696	287 (5)	259 (18), 258 (100), 214 (10), 186 (14), 181 (14), 115 (13)	E	i	r	b	į.	i
11-Hydroxyvictatine (15)	2728	287 (6)	259 (18), 258 (100), 242 (10), 211 (15), 181 (20), 128 (13)	1	1	,	1	5	1
3-Epideoxytazettine (16)	2341	315(21)	300 (41), 232 (14), 231 (100), 185 (12), 115 (15), 70 (65)	1	1	3.12	7.55	1	1
Deoxytazenine (17)	2486	315 (21)	300 (15), 260 (5), 231 (100), 227 (10), 211 (15), 197 (10), 115 (9)	1	1	4.10	3.12	1	1

Compound	RI	tw M	Rel int (%)	H. striatum Bulbs	H. virtatum Bulbs	H. breviforum Bulbs	H. morelianum Bulbs	H. papilio Bulbs	H. popilio Leaves
6-Methoxypretazenine (18)	2610	345 (26)	330 (21), 262 (21), 261 (100), 239 (40), 228 (30), 201 (28)	10	1	b	ı	1	1
Taxettine (19)/Protazettine (20)*	2663	331 (31)	316 (15), 298 (23), 247 (100), 230 (12), 201 (15), 181 (11), 152 (7)	1	1	26.50	58.83	1	1
3-Epimacronine (21)	28	329 (27)	314 (23), 245 (100), 225 (14), 201 (83), 139 (16), 70 (18)	ī	i	69'0	3.18	1	1
Tazetami de (22)	2914	313 (30)	260 (100), 229 (20), 201 (49), 171 (12), 143 (9), 115 (26)	î	1	1	מ	1	1
Trisphaendine (23)	2282	223 (100)	222 (38), 167 (8), 165 (9), 164 (14), 138 (20), 137 (9), 111 (13)	b	1	0.75	1.5	1	1
Pancracine (24)	2718	287 (100)	270 (22), 243 (22), 223 (25), 199 (29), 185 (34), 115 (18)	í.	ti di	i.	i	1	1
Montanine (25)	2611	301 (100)	270 (90), 257 (39), 252 (26), 223 (33), 185 (37), 115 (30)	1	86.62	1	1	1	1
Anhydrogalanthamine (26)	1766	269 (100)	268 (38), 211 (43), 195 (22), 193 (31), 165 (61), 115 (26)	1	1	1	1	132	1
Galanthamine (27)	2395	287 (83)	288 (14), 286 (100), 270 (13), 244 (26), 216 (37), 174 (34)	b	1	1	t	63.24	58.97
Sanguinine (28)	2422	273 (100)	272 (79), 256 (18), 216 (18), 202 (37), 160 (44), 115 (25)	i i	1	ı	ı	1	b
A:Demethylg danthamine (29)	2442	273 (98)	272 (100), 230 (44), 202 (34), 201 (12), 174 (13)	1	1	1	1	1	ı
3-Epigalandramine (30)	2443	287 (77)	286 (100), 270 (15), 244 (16), 216 (70), 211 (14), 174 (26)	ı	1	1	1	1	1
Narwedine (31)	2483	285 (95)	284 (100), 242 (30), 228 (25), 216 (40), 199 (35), 174 (40)	1	1	1	1	1.62	285
11 p-Hydroxyg alanthanine (32)	2597	303 (24)	231 (21), 230 (100), 213 (27), 181 (13), 174 (13), 115 (15)	ī	ı	ī	1	380	996
A-Formylhorgalanthamine (33)	2816	301 (100)	230 (9), 225 (16), 211 (18), 165 (9), 128 (10), 115 (13)	í	1	1	1	1	1
Smitne (34)	2280	257 (35)	238 (100), 211 (6), 196 (8), 168 (6), 154 (3), 106 (4), 77 (3)	1	1	1.41	0.75	1	1
Gallanthindole (35)	2487	281 (100)	280 (7), 264 (13), 263 (17), 262 (20), 252 (15), 191 (14)	i i	ı	1.70	1.49	1	1
Lycosinine B (36)	2520	297 (100)	298 (19), 269 (72), 268 (56), 254	1	1	5.12	1	1	1

Compound	B	₩ ₊	Rel. int (%)	H. psirkacinum Bulbs	H. psütacinam Leaves	H. sawacatarina Leaves	H. glaucescens Bults	H. glaucescens Leaves	R. hifsån Bulbs
Anhydrolycorine (1)	2501	251 (43)	250 (100), 192 (13), 191 (11), 165 (4), 164 (3), 139 (2), 124 (7)	1	1	ь	1	1	1
11,12. Dehydroanhydrolycorine (2)	3606	249 (60)	248 (100), 191 (10), 190 (24), 189 (7), 163 (7), 95 (17)	1	1	14.64	1	1	1
Lycorine (3)	2746	287 (31)	286 (19), 268 (24), 250 (15), 227 (79), 226 (100), 211 (7), 147 (15)	1	1	19.18	i	1	1
8-0- Demethylhomolycorine (4)	284	301 (-)	192 (0.5), 164 (2), 110 (8), 109 (100), 108 (23), 94 (3), 82 (3)	1	1970	1	1	1	1
Northine (5)	2476	347 (-)	330 (7), 329 (3), 236 (1), 221 (9), 191 (2), 109 (100), 94 (2)	i	1	1	ī	ī	i
2x-Hydroxyhomolyconne (6)	2970	331 (-)	178 (3), 126 (8), 125 (100), 124 (7), 96 (31), 94 (4)	t	1	1	1	1	1
Hippeastrine (7)	2917	315 (-)	190 (1), 162 (4), 134 (2), 125 (100), 96 (40), 82 (3)	8.82	23.90	1	í	b	ı
2a-Methoxyhomoly corine (8)	2870	345 (-)	178 (5), 140 (11), 139 (100), 124 (67), 94 (7), 77 (5)	1	1	1	ı	ı	100
2a,7- Dimethoxyhomolycorine (9)	2962	375(-)	221(2), 140(9), 139(100), 125(6), 124(55), 94(4)	1	1	1	1	1	1
Candimine (10)	3000	345 (-)	192 (1), 177 (2), 163 (1), 147 (1), 125 (100), 96 (30), 82 (2)		1	1	1	,	31
Vitratine (11)	2472	271 (100)	272 (20), 252 (85), 199 (70), 187 (61), 173 (22), 115 (28)	1	1	13	1	1	ь
8-O-Demethylmarifidine (12)	2510	273 (100)	274 (17), 230 (24), 201 (83), 189 (52), 175 (20), 115 (18)	1	1	1	1	1	1
Haemanthamine (13)	2641	301 (13)	272 (100), 240 (16), 211 (13), 199 (7), 181 (21), 153 (8)	ı	1	3.61	ī	ī	1
Hamayne (14)	3699	287 (5)	259 (18), 258 (100), 214 (10), 186 (14), 181 (14), 115 (13)	1	i	i.	i.	r.	C
11-Hydroxyvitutine (15)	2728	287(6)	259(18), 258(100), 242(10), 211(15), 181(20), 128(13)	1	1	8.51	1	1	SI.
3-Epideoxytazettine (16)	2241	315 (21)	300 (41), 232 (14), 231 (100), 185 (12), 115 (15), 70 (65)	1	1	1	1.26	1	1
Deoxytaxetine (17)	2486	315 (21)	300 (15), 260 (5), 231 (100), 227 (10), 211 (15), 197 (10), 115 (9)	b	0.82	5	030	ь	b

Compound	2	TW.	Rel. int. (%)	H. psiracinum Bulbs	H. psittacinum Leaves	H. santicatarina Leaves	H. glawescens Bulbs	H. glaucescens Leaves	R. byfala Bulbs
6-Methoxypretazettine (18)	2610	345 (26)	330 (21), 262 (21), 261 (100), 239 (40), 228 (30), 201(28)	1	1	t	1	Е.	ı
Taxettine (19)/Protazettine (20)*	2663	331 (31)	316 (15), 298 (23), 247 (100), 230 (12), 201 (15), 181 (11), 152 (7)	36.84	14.83	b	7.62	14.89	ti.
3-Epimacronine (21)	281	329 (27)	314 (23), 245 (100), 225 (14), 201 (83), 139 (16), 70 (18)	5.78	7.31	b	160	3.64	tt.
Taxetamide (22)	2914	313 (30)	260 (100), 229 (20), 201 (49), 171 (12), 143 (9), 115 (26)	1.84	127	1	1	1	ı
Trisphaeridine (23)	2282	223 (100)	222 (38), 167 (8), 165 (9), 164 (14), 138 (20), 137 (9), 111 (13)	91.1	5	19.34	5900	b	į.
Pancracine (24)	2718	287 (100)	270 (22), 243 (22), 223 (25), 199 (29), 185 (34), 115 (18)	1	1	1	1	91	1
Montanine (25)	3611	301 (100)	270 (90), 257 (39), 252 (26), 223 (33), 185 (37), 115 (30)	1	1	1	1	1	91.94
Anhydrogalandiamine (26)	1766	269 (100)	268 (38), 211 (43), 195 (22), 193 (31), 165 (61), 115 (26)	1	1	1	16.38	ь	ı
Galanthamine (27)	2395	287 (83)	288 (14), 286 (100), 270 (13), 244 (26), 216 (37), 174 (34)	b	1	ь	55.30	66.15	1
Sangui nine (28)	2422	273 (100)	272 (79), 256 (18), 216 (18), 202 (37), 160 (44), 115 (25)	1	1	1	1.38	b	ı
A-Demethylg alanthamine (29)	2442	273 (98)	272 (100), 230 (44), 202 (34), 201 (12), 174 (13)	1.	1	1	ь	1	1
3-Epigalandamine (30)	2443	287 (77)	286 (100), 270 (15), 244 (16), 216 (70), 211 (14), 174 (26)	1	r i	1	223	1	1
Narwedine (31)	2483	285 (95)	284 (100), 242 (30), 228 (25), 216 (40), 199 (35), 174 (40)	1	1	1	5.51	2.42	1
11 ft-Hydroxyg alanthamine (32)	2597	303 (24)	231 (21), 230 (100), 213 (27), 181 (13), 174 (13), 115 (15)	1	i	1	ı	t	1
A-Formylhorgalanthamine (33)	2816	301 (100)	230 (9), 225 (16), 211 (18), 165 (9), 128 (10), 115 (13)	1	t	1	ı	b	t
Samine (34)	2280	257 (35)	238 (100), 211 (6), 196 (8), 168 (6), 154 (3), 106 (4), 77 (3)	13.9	10.46	1	1	2.10	l.
Galanthindole (35)	2487	281 (100)	280 (7), 264 (13), 263 (17), 262 (20), 252 (15), 191 (14)	7.40	12.73	1	1	5.41	1
Lycosinine B (36)	2520	297 (100)	298 (19), 269 (72), 268 (56), 254 (32), 237 (19), 222 (16)	ı	1	ı	1	1	1

Plant species	IC ₅₀ (µg/ml)	AChE inhibiti	on %
		10 μg/ml	0.1 μg/ml
Hippeastrum striatum bulbs	nd	-	-
Hippeastrum vittatum bulbs	4.67	31.0	2.0
Hippeastrum breviflorum bulbs	nd	_	-
Hippeastrum morelianum bulbs	nd	-	-
Hippeastrum papilio bulbs	0.45	93.0	23.0
Hippeastrum papilio leaves	0.41	96.0	24.0
Hippeastrum psitaccinum bulbs	nd	-	-
Hippeastrum psitaccinum leaves	nd	_	_
Hippeastrum santacatarina bulbs	nd	-	-
Hippeastrum glaucescens bulbs	0.33	93.0	26.0
Hippeastrum glaucescens leaves	0.49	94.0	20.0
Hippeastrum aulicum leaves	nd	-	_
Rhodophiala bifida bulbs	8.45	28.0	3.0