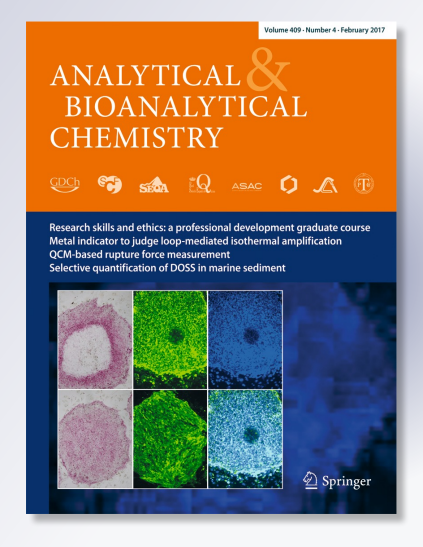
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RESEARCH PAPER



Mediterranean *Azadinium dexteroporum* (Dinophyceae) produces six novel azaspiracids and azaspiracid-35: a structural study by a multi-platform mass spectrometry approach

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Abstract Azadinium dexteroporum is the first species of the genus described from the Mediterranean Sea and it produces different azaspiracids (AZA). The aims of this work were to characterize the toxin profile of the species and gain structural information on azaspiracids produced by the A. dexteroporum strain SZN-B848 isolated from the Gulf of Naples. Liquid chromatography-mass spectrometry (LC-MS) analyses were carried out on three MS systems having different ion source geometries (ESI, TurboIonSpray®, ESI ION MAX) and different MS analyzers operating either at unit resolution or at high resolution, namely a hybrid triple quadrupole-linear ion trap (Q-Trap MS), a time of flight (TOF MS), and a hybrid linear ion trap Orbitrap XL Fourier transform mass spectrometer (LTQ Orbitrap XL FTMS). As a combined result of these different analyses, A. dexteroporum showed to produce AZA-

Rachele Rossi and Carmela Dell'Aversano contributed equally to this work.

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35, previously reported from *Azadinium spinosum*, and six compounds that represent new additions to the AZA-group of toxins, including AZA-54 to AZA-58 and 3-epiAZA-7, a stereoisomer of the shellfish metabolite AZA-7. Based on the interpretation of fragmentation patterns, we propose that all these molecules, except AZA-55, have the same A to I ring system as AZA-1, with structural modifications all located in the carboxylic side chain. Considering that none of the azaspiracids produced by the Mediterranean strain of *A. dexteroporum* is currently regulated by European food safety authorities, monitoring programs of marine biotoxins in the Mediterranean area should take into account the occurrence of the new analogues to avoid an underestimation of the AZA-related risk for seafood consumers.

Keywords Azaspiracids \cdot *Azadinium dexteroporum* \cdot Marine toxins \cdot LC-MS/MS \cdot LC-HRMSⁿ \cdot LC-TOFMS

Introduction

Azaspiracids (AZAs) are a group of polyether toxins first reported to cause seafood poisoning in the Netherlands in 1995, following the ingestion of contaminated shellfish from Killary Harbour, Ireland [1]. Shellfish were contaminated by a unique marine toxin, originally named "Killary-toxin" and shortly after structurally elucidated and renamed azaspiracid-1 (AZA-1) [2–4] (Fig. 1). The name was consistent with its chemical features which includes a six-membered cyclic amine (AZA), a unique tri-spiro assembly (SPIRO), and a carboxylic acid group (ACID). The rest of the molecule consisted of a linear carbon chain cyclized at several points through ether bridges. The biogenetic origin of azaspiracids causing the Killary Harbour shellfish contamination was initially identified in the heterotrophic dinoflagellates



Fig. 1 Planar structure of AZA-1 and characteristic fragmentations occurring in CID mode. The other azaspiracids so far known mainly differ from AZA-1 by substitution at C3 (H or OH), C8 (H or CH₃), C21 (OH or OCH₃), C22 (H, CH₃, or COOH), C23 (H or OH), and/or C39 (H or CH₃)

Protoperidinium crassipes [5], but was later definitely proven to be the phototrophic dinoflagellate Azadinium spinosum [6].

Over the last years, several analogues of AZA-1 have been discovered either in shellfish [7–11] or in dinoflagellates [6, 12-17] and structurally characterized by tandem mass spectrometry (MS/MS) and partly also by nuclear magnetic resonance (NMR) spectroscopy. Some AZAs (AZA-1 and AZA-2, among others) are actually produced by the dinoflagellates, whereas others (e.g., AZA-3 to AZA-19) seem to derive from biotransformation occurring in shellfish [9]. The formation of extraction artifacts (21-epi-AZA-1, AZA-29, AZA-30, and AZA-32) has also been demonstrated [18], especially when methanol is used as extraction solvent. Consequently, the extraction solvent currently suggested for AZA analysis is acetone or aqueous acetonitrile. Not considering stereoisomers, more than 30 variants of AZAs have been detected so far [19, 20]. Among them, only AZA-1, AZA-2, and AZA-3 are currently regulated in Europe and hence regularly monitored in shellfish to safeguard seafood consumer health [21].

The official technique for the monitoring of regulated azaspiracids in shellfish is LC-MS [22] which provides quantitative data and is also widely used in research settings to gain structural insights into so far unknown molecules. The MS behavior of AZAs is indeed characteristic and it has been studied widely using either tandem MS or high resolution MS analyzers [9, 23–25]. Over the last years, new dinophycean species other than *A. spinosum* have been identified as AZA producers, including *Amphidoma languida* from Bantry Bay and two species in the genus *Azadinium*, i.e., *Azadinium poporum* from the North Sea, Korea [12], and the China Sea [26], and *Azadinium dexteroporum* from the Mediterranean Sea [13]. The latter species, first described based on a strain isolated in the Gulf of Naples [13], was later

retrieved in Arctic waters [27], but the cold water strain slightly differed in molecular sequence data and did not produce any AZA. Preliminary LC-TOF MS analysis of an 80 % aqueous methanol extract from the Mediterranean *A. dexteroporum* strain showed the presence of three potentially new AZA-like compounds with [M+H]⁺ ions at *m/z* 828, *m/z* 858, and *m/z* 830 [13]. These compounds were supposed to be AZA-like compounds, based on their molecular masses and *in-source* fragmentation behavior analyzed in comparison to that of AZA-1, and tentatively assigned to putative AZA-3, AZA-7, and Compound 3 [12], respectively, while several other AZA-like compounds were noticed and left undescribed.

The aim of this work was to further investigate AZAs produced by the Mediterranean *A. dexteroporum* SZN-B848 strain to shed light on their identity, thus providing a better characterization of the whole toxin profile of the strain. The analyses took advantage of several different LC-MS techniques, used in combination in order to obtain structural information on the AZAs produced by this strain.

Materials and methods

Chemicals and standards

Water was deionized and purified through a MilliQ water purification system (Milli-Q, Millipore, Eschborn, Germany) to $18~\mathrm{M}\Omega~\mathrm{cm}^{-1}$ or better quality. The organic solvents acetonitrile and methanol (HPLC grade) and the additives formic acid (90 %, p.a.), acetic acid (p.a.), and ammonium formate (p.a.) were purchased from Sigma-Aldrich (Steinheim, Germany) and Merck (Darmstadt, Germany). Certified reference material of AZA-1 was purchased from NRC, Canada (NRC CRM



AZA-1) and used for setting LC-HRMSⁿ conditions and for quantitative analysis. A mussel tissue extract containing AZA-1, AZA-2, and AZA-3, kindly provided by Dr Philipp Hess (Ifremer, Nantes, France), and reference samples of AZA-7 and AZA-35, kindly provided by Dr Jane Kilkoyne (Marine Institute, Galway, Ireland), were used for toxins identification.

A. dexteroporum batch cultures

The strain SZN-B848 of *A. dexteroporum* was isolated from a water sample collected on May 11, 2010 at the LTER-MC in the Gulf of Naples. The culture was grown in 1 L Erlenmeyer flask in K/10 medium at a salinity of 36, under no axenic conditions. It was maintained at 20 °C under an irradiance of 70–80 μ mol photons m⁻² s⁻¹ in a 12:12 light/dark regime as described in Percopo et al. [13]. Cell density in the culture harvested for chemical analyses during the stationary growth phase was 1.4×10^5 cells mL⁻¹.

Extraction

The 80 % aqueous MeOH extract of A. dexteroporum used in Percopo et al. [13] was re-analyzed by LC-MS 6 months after preparation. Considering that extraction with aqueous methanol may lead to formation of AZAs artifacts, including methyl esters and/or methyl ketals [18], another batch of A. dexteroporum was extracted using the procedure described by Jauffrais et al. [18]. In particular, the pellet $(1.4 \times 10^5 \text{ cells})$ was suspended in 1 mL acetone/H₂O (9:1, v/v), vortexed for 1 min, sonicated for 10 min, and centrifuged at 1000×g for 5 min. The supernatant was transferred into a glass test tube and the pellet was re-extracted twice with 0.5 mL of acetone/ H_2O (9:1, v/v) following the same procedure. The supernatants were combined and evaporated to dryness. The residue was dissolved in 1 mL acetone/ H_2O (9:1, v/v), centrifuged through a 0.22-µm spin-filter (Millipore Ultrafree, Eschborn, Germany) at 3000×g for 5 min, and directly analyzed by LC-MS on system #1 and subsequently on systems #2 and #3.

Liquid chromatography-mass spectrometry (LC-MS) analyses

LC-MS analyses were performed according to Quilliam et al. [28] on three different systems, namely a LC-Time of Flight MS (LC-TOFMS) (system #1), a LC-LTQ Orbitrap XL FTMS (LC-HRMSⁿ, n = 1–3) (system #2), and a LC-triple quadrupole MS (LC-MS/MS) (system #3). A Luna 3 μ C(8) 150 × 2.00 mm (system #1) and a Hypersil 3 μ C8 BDS, 50 × 2.00 mm columns (Phenomenex, Torrance, CA, USA) (systems #2 and #3) were kept at room temperature and eluted with water (eluent A) and 95 % acetonitrile/water (eluent B), both containing 50 mM formic acid and 2 mM ammonium formate. Flow rate was 200 μ L min⁻¹ and injection volume

was 5 μ L. MS parameters and gradient conditions on systems #1-3 are detailed below.

System #1 LC-TOF MS analyses were performed at IZSM (Italy) on an Agilent TOF MS model G1969 coupled with an Agilent model 1100 LC (Agilent Palo Alto, CA, USA). The following linear gradient was used: 60 % B for 6 min, 60 to 80 % B in 0.5 min and hold for 2 min, then returning to initial conditions in 6 min. Full scan MS experiments (positive ion mode) were performed in the mass range m/z 100–1000 at a resolving power (RP) = 10,000. The following ESI source settings were used: drying gas (N_2) flow rate = 8 mL min⁻¹, drying gas temperature = 350 °C, nebulizer gas = 45 psi, capillary voltage = 4000 V, fragmentor voltage = 350 V, skimmer voltage = 60 V. A tuning mix (G1969-85001) was used for lock mass calibration. Agilent LC/MS-TOF Software was used for data acquisition and data processing by analyzing all the scans ("walking on the chromatogram") and identifying the characteristic fragment ions in the spectrum preview. Extracted ion chromatograms (EIC) were obtained for all the AZAs by selecting exact masses of their [M+H]⁺ ions at a mass tolerance of 10 ppm.

System #2 LC-HRMSⁿ analyses (n = 1-3) were performed at University of Napoli Federico II (Italy) on a hybrid linear ion trap LTQ Orbitrap XLTM Fourier Transform Mass Spectrometer (FTMS) equipped with an ESI ION MAXTM source (Thermo Fisher, San José, USA) and coupled to an Agilent 1100 LC binary system (Agilent Palo Alto, CA, USA). The following gradient elution was used, 30 to 100 % B in 15 min followed by 100 % B for 9 min, then returning to initial conditions until 27 min, and reequilibrating for 9 min (total run time = 36 min). Full scan HRMS experiments (positive ion mode) were acquired in the m/z 700–900 range at a RP = 60,000 (FWHM at m/z 400). The following source settings were used: spray voltage = 4.8 kV, capillary temperature = 260 °C, capillary voltage = 29 V, sheath gas flow = 36 and auxiliary gas flow = 14 (arbitrary units), tube lens voltage = 125 V. HRMS^n experiments (n = 2, 3) were acquired at RP = 30,000 using either higher energy collisional dissociation (HCD), collision-induced dissociation (CID), or their combinations (CID/CID and CID/HCD). In HR HCD MS² experiments [M+H]⁺ ions of individual AZAs were used as precursors with a collision energy (CE) = 35 %. In HR CID/CID MS³ and HR CID/HCD MS³ experiments, precursors were $[M+H]^+$ ions (CE = 45 %) and [M+H- H_2O]⁺ ions (CE = 35 %). An isolation width (IW) = 2.0 Da, an activation Q = 0.250, and an activation time = 30 ms were used. Calculation of elemental formulae was performed on the mono-isotopic peak of each ion cluster using Xcalibur software v2.0.7 with a mass tolerance of 3 to 5 ppm. The isotopic ion pattern was taken into account in assigning molecular formula. EIC for the detected AZAs were obtained by

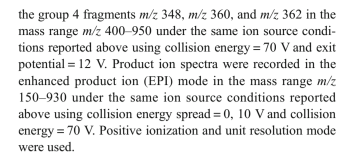


selecting the [M+H]⁺ ion clusters and setting mass tolerance to a value compatible with the error in ion assignment of the AZA standard (1–4 ppm) depending on the daily instrument calibration status.

System #3 LC-MS/MS analyses were performed at AWI (Germany) on an AB-SCIEX-4000 Q-Trap MS equipped with a TurboSpray® interface coupled to an Agilent model 1100 LC. The LC equipment included a solvent reservoir, in-line degasser (G1379A), binary pump (G1311A), refrigerated autosampler (G1329A/G1330B), and temperature-controlled column oven (G1316A). The following gradient elution was used, 30 to 100 % B in 8 min followed by 100 % B for 10 min, then returning to initial conditions until 21 min and reequilibrating for 8 min (total run time = 29 min). In addition, a guard column (Securiguard, Phenomenex, Aschaffenburg, Germany) was used. AZA profiles were determined in one period (0-18) min with curtain gas = 10 psi, CAD = medium, ion spray voltage = 5500 V, temperature = ambient, nebulizer gas = 10 psi, auxiliary gas = off, interface heater = on, declustering potential = 100 V, entrance potential = 10 V, exit potential = 30 V. Selected reaction monitoring (SRM) experiments were carried out in positive ion mode by selecting the transitions (precursor ion > fragment ion) shown in Table 1. AZAs were calibrated against AZA-1 CRM and expressed as AZA-1 equivalents. Precursor ion scanning (PI) experiments were carried in the positive ion mode scanning precursors of

Table 1 Mass transitions m/z (Q1>Q3 mass) and collision energy (CE) used to monitor azaspiracids in SRM experiments

Azaspiracid (AZA)	Mass transition	CE (V)
AZA-33	716>698	40
AZA-39	816>798	40
AZA-39	816>348	70
AZA-3	828>658	70
AZA-3, AZA-58	828>810	40
AZA-35, AZA-38	830>812	40
AZA-38	830>348	70
AZA-1	842>672	70
AZA-1, AZA-6, AZA-41	842>824	40
AZA-4, AZA-5, AZA-56	844>826	40
AZA-37	846>828	40
AZA-2, Me-AZA-1	856>672	70
AZA-2, Me-AZA-1	856>838	40
AZA-7, AZA-8, AZA-9, AZA-10, AZA-36	858>840	40
AZA-36	858>348	70
AZA-55	868>362	70
AZA-54, Me-AZA-2	870>852	40
AZA-11, AZA-12	872>854	40
AZA-57	884>866	40



Quantitation

Quantitation was performed on system #3 by external calibration in single measurements against an AZA-1 CRM solution and all values were expressed as AZA-1 equivalents. Relative percentages of AZAs were measured based on peak areas of individual AZAs obtained in SRM experiments and cross-checked with EIC of individual AZAs obtained on system #2. Cell quota was extrapolated from the data, assuming that the detected AZAs had the same molar response as AZA-1.

Derivatization with diazomethane

A 50- μ L aliquot of the algal extract was mixed with an equal volume of a 2 M trimethylsilyl diazomethane (TMSDM) solution (product no: 527524, Sigma-Aldrich, Steinheim, Germany). The mixture was vortexed six times every 5 min for complete mixing. After 30 min, excess TMSDM was destroyed by the addition of 5 μ L glacial acetic acid and vortexing. The derivatized sample was taken to dryness in a gentle nitrogen stream and the residue taken up in 50 μ L methanol.

Results and discussion

The aqueous methanol extract studied by Percopo et al. [13] and an acetone extract of the same A. dexteroporum SZN-B848 strain were analyzed by LC-MS on three MS systems having different ion source geometries (ESI, TurboIonSpray®, ESI ION MAX) and different MS analyzers operating either at unit resolution or at high resolution (Q-Trap MS, TOF MS, and LTQ Orbitrap XL FTMS). Both extracts gave the same results confirming that none of the AZAs detected in the methanolic extract was an extraction artifact. Seven azaspiracids were detected (Fig. 2): AZA-35, which had been reported previously from A. spinosum [29], and six novel compounds that represent new addition to the AZAgroup of toxins, including AZA-54 to AZA-58 and 3-epi-AZA-7, a stereoisomer of the shellfish metabolite AZA-7 [5]. A total AZA cell quota of 24.5 fg/cell was estimated from a cross check of the Q-Trap MS data with relative abundances and concentrations measured on the LTQ Orbitrap XL FTMS



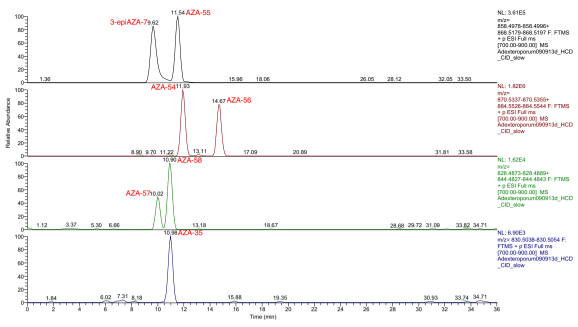


Fig. 2 LC-HRMS analysis (system #2) of the Mediterranean A. dexteroporum strain. Extracted ion chromatograms (1 ppm mass tolerance, normalized peak height) of the seven azaspiracids contained in the acetone extract

using AZA-1 CRM as standard. This cell quota is comparable to that of other AZA-producing species such as *A. spinosum* (5–50 fg/cell) [6] and *A. poporum* (0–10 fg/cell) [14, 16] and is quite different from that reported by Percopo et al. (2–3 fg/cell) for the same strain, based on TOF MS analyses of only three minor components of the *A. dexteroporum* profile [13].

The available material was not sufficient for even attempting isolation and subsequent full structural elucidation based on NMR spectroscopy. Thus, MS fragmentation analysis was the method of choice because of its sensitivity and ability to provide structural information on compounds contained at trace levels in crude extracts. AZA-1, which was not found in the A. dexteroporum extract, was used as reference for the comparative interpretation of the fragmentation patterns. It is worth mentioning that fragmentation of AZAs occurs by elimination of molecule portions of variable size from their charged end—the amine nitrogen in ring I producing group 1 fragments $[M+H-nH_2O]^+$ with n=1-6 and several structurally informative fragment ions due to cleavages at specific sites of the polyketide chain, each generally undergoing multiple water losses (group 2-6 fragments) (Fig. 1). The main cleavages include the following:

- (a) A pseudo retro Diels-Alder (RDA) reaction of ring A [9] leading to loss of the C1-C9 part structure with its substituents and formation of a fragment which, along with its water losses, results in the group 2 fragment cluster (starting at *m*/*z* 672 in AZA-1).
- (b) A cleavage of ring C leading to elimination of the C1-C15 part structure and formation of a fragment which,

- together with its water losses, gives the group 3 fragments (starting at m/z 462 in AZA-1).
- (c) A pseudo RDA of ring E where C1–C23 are eliminated resulting in the formation of fragment ions that barely undergo water losses, named group 4 fragments (starting at m/z 362 for AZA-1).
- (d) A cleavage between C27 and C28 where C1–C27 are eliminated resulting in formation of group 5 fragments (at *m/z* 262 in AZA-1)
- (e) A pseudo RDA of ring G resulting in the formation of group 6 fragments (at *m*/*z* 168 in AZA-1).

Apart from these general fragmentation patterns, 3-hydroxy-AZAs, such as AZA-36 and AZA-37—compounds 1 and 2 in [12]—present a typical fragmentation pathway in their CID spectra: in addition to group 1 fragments (common to all AZAs), their pseudomolecular ion also eliminates CO_2 (44 Da) and several water molecules. The $[M-CO_2-nH_2O]^+$ (n=1-3) fragments usually represent up to 20 % of the intensity of the highest peak but are clearly visible and represent a key feature of all the 3-hydroxy-AZAs analyzed so far [19].

The AZAs produced by the *A. dexteroporum* SZN-B848 strain were first revealed by the *in-source* fragmentation spectra obtained on the TOF MS, which were dominated by the [M+H]⁺ ions of each analogue and relevant group 1 fragments. The structurally informative group 2–4 fragments presented very low intensity on this system and in some cases could hardly be distinguished from the background noise, while the group 5–6 fragments were not detected at all. Therefore, further analyses of the fragmentation patterns contained in



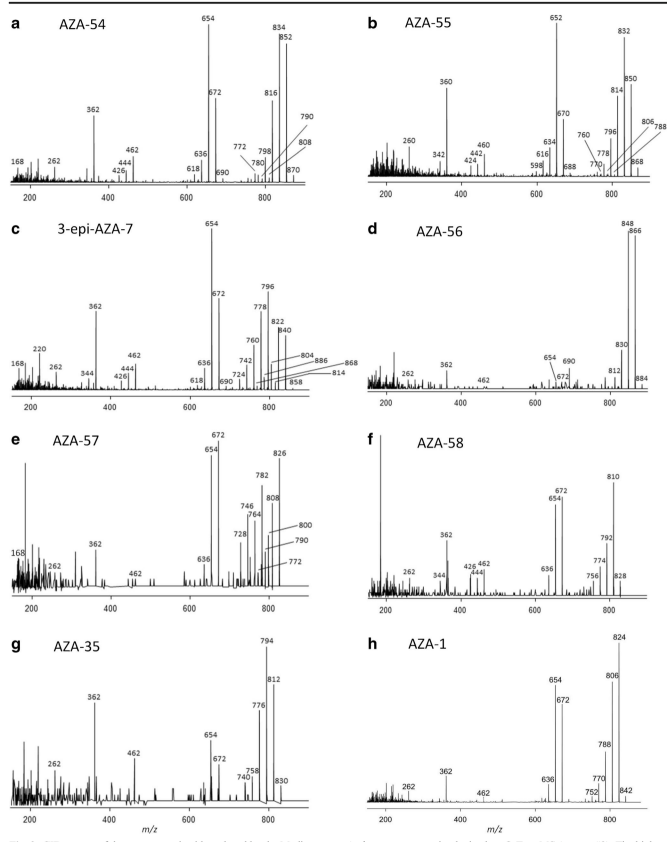


Fig. 3 CID spectra of the seven azaspiracid produced by the Mediterranean *A. dexteroporum* strain obtained on Q-Trap MS (system #3). The highest labeled m/z value in each CID spectrum refers to the [M+H]⁺ peak



Table 2 Exact masses of the seven azaspiracids produced by the Mediterranean *A. dexteroporum* strain and of AZA-1 standard measured on the LTQ-Orbitrap MS. Molecular formula (MF), ring double bond (RDB) equivalents, and errors (Δ, ppm) are also reported together with relative retention times (RRt) to AZA-1 obtained under the different chromatographic conditions (system #1–3)^a

Name	$[M+H]^+$, m/z	Formula	RDB	Δ , ppm	RRt #1	RRt #2	RRt #3
AZA-54	870.5346	C ₄₉ H ₇₆ O ₁₂ N	12.5	-1.553	-3.93	-0.71	-0.61
AZA-55	868.5188	$C_{49}H_{74}O_{12}N$	13.5	-1.773	-2.02	-0.32	-0.30
3-epiAZA-7	858.4987	$C_{47}H_{72}O_{13}N$	12.5	-1.068	2.83	1.60	0.94
AZA-56	884.5535	$C_{50}H_{78}O_{12}N$	12.5	1.597	-5.59	-3.45	-2.82
AZA-57	844.4835	$C_{46}H_{70}O_{13}N$	12.5	-0.688	1.96	1.2	0.65
AZA-58	828.4881	$C_{46}H_{70}O_{12}N$	12.5	-1.183	-0.02	0.32	0.09
AZA35	830.5046	$C_{46}H_{72}O_{12}N$	11.5	-0.283	-0.32	0.24	0
AZA-1	842.50548	$C_{47}H_{72}O_{12}N \\$	12.5	0.685	7.24	11.22	12.09

^a Columns and gradients used for the chromatographic separation on systems #1 (TOF MS), #2 (LTQ Orbitrap XL FTMS), and #3 (O-Trap MS) are reported in the experimental section

CID MS spectra obtained on the Q-Trap MS (Fig. 3) and HRMSⁿ (n = 1, 2, and 3) spectra obtained on the LTQ-Orbitrap FTMS (Tables 2, 3, and 4) were needed to gain confirmation of the structural features of the analogues.

LC-HRMS experiments on the LTQ Orbitrap XL allowed to assign molecular formula and ring double bond (RDB) equivalents to each of the seven AZAs contained in the *A. dexteroporum* extract (Table 2). The fragmentation pattern

of each compound was studied through the analysis of LC-HRMS² spectra in HCD mode and confirmed through LC-HRMS³ spectra in CID/CID or CID/HCD modes (Tables 3 and 4). HCD fragmentation mode provided spectra containing all the expected group 1–6 fragments, with the fragments in the low mass range being the most intense ones (see Electronic Supplementary Material (ESM), Fig. S1). However, for structural characterization of the minor

Table 3 Fragmentation pattern of 3-epiAZA-7 and AZA-35 produced by the Mediterranean *A. dexteroporum* strain versus that of AZA-1 standard measured on the LTQ-Orbitrap FTMS in various HRMSⁿ modes (n = 2, 3 either HCD, CID/HCD, or CID/CID). Errors in ion assignments were in the range 0.5–4 ppm

		AZA-1			3-epiAZA-	.7		AZA-35		
	Loss	m/z	Formula	RDB	m/z	Formula	RDB	m/z	Formula	RDB
Precursor		842.5	C47 H72 O12 N	12.5	858.5	C47 H72 O13 N		830.5	C46 H72 O12 N	
Group 1	-1H2O	824.4917	C47 H70 O11 N	13.5	840.4897	C47 H70 O12 N	13.5			
	-2H2O	806.4813	C47 H68 O10 N	14.5	822.4763	C47 H68 O11 N	14.5	794.4797	C46 H68 O10 N	13.5
	-3H2O	788.4713	C47 H66 O9 N	15.5	804.4660	C47 H66 O10 N	15.5	776.4702	C46 H66 O9 N	14.5
	-4H2O	770.4616	C47 H64 O8 N	16.5				758.4598	C46 H64 O8 N	15.5
	-5H2O	752.4512	C47 H62 O7 N	17.5						
	-6H2O	734.4421	C47 H60 O6 N	18.5						
	-CO2-H2O				796.4974	C46 H70 O10 N	12.5			
	-CO2-2H2O				778.4866	C46 H68 O9 N	13.5			
	-CO2-3H2O				760.4766	C46 H66 O8 N	14.5			
	-CO2-4H2O				742.4660	C46 H64 O7 N	15.5			
	-CO2-5H2O				724.5551	C46 H62 O6 N	16.5			
Group 2	-H2O	672.4085	C38 H58 O9 N	10.5	672.4094	C38 H58 O9 N	10.5	672.4088	C38 H58 O9 N	10.5
		654.3979	C38 H56 O8 N	11.5	654.3984	C38 H56 O8 N	11.5			
Group 3	-H2O	462.3198	C27 H44 O5 N	6.5	462.3201	C27 H44 O5 N	6.5	462.3196	C27 H44 O5 N	6.5
		444.3096	C27 H42 O4 N	7.5	444.3102	C27 H42 O4 N	7.5			
Group 4	-H2O	362.2679	C22 H36 O3 N	5.5	362.2681	C22 H36 O3 N	5.5	362.2681	C22 H36 O3 N	5.5
		344.2574	C22 H34 O2 N	6.5	344.2571	C22 H34 O2 N	6.5			
Group 5		262.1795	C16 H24 O2 N	5.5	262.1794	C16 H24 O2 N	5.5			
Group 6		168.1377	C10 H18 O N	2.5	168.1378	C10 H18 O N	2.5			



Table 4 Fragmentation pattern of the AZA-54 to AZA-58 produced by the Mediterranean A. dexteroporum strain measured on the LTQ-Orbitrap FTMS in various HRMS" modes (n = 2, 3 either HCD, CID/HCD, or CID/CID). Errors in ion assignments were in the range 0.5-4 ppm

		AZA-54			AZA-55			AZA-56			AZA-57			AZA-58		
	Loss	z/m	Formula	RDB m/z		Formula]	RDB		Formula	RDB	z/m	Formula	RDB	z/w	Formula	RDB
Precursor		870.5	C49 H76 O12 N 12.5	12.5	868.5	C49 H74 O12 N 13.5		884.5	C50H78O12N 12.5		844.5	C46 H70 O13 N 12.5 828.5	12.5	828.5	C46 H70 O12N 12.5	12.5
Group 1 -1H2O	-1H2O	852.5241	852.5241 C49 H74 O11 N 13.5 850.5081 C49 H72 O11 N 14.5	13.5	850.5081	C49 H72 O11 N	14.5									
	-2H2O	834.5135	834.5135 C49 H72 O10 N 14.5	14.5	832.4977	832.4977 C49 H70 O10 N 15.5		848.5266 (848.5266 C50H74O9N	14.5	808.4601	808.4601 C46 H66 O11 N 14.5 792.4643 C46 H66 O10 N 14.5	14.5	792.4643	C46 H66 O10 N	14.5
	-3H2O	816.5015	816.5015 C49 H70 O9 N	15.5	814.4861	814.4861 C49 H68 O9 N	16.5	830.5161	830.5161 C50H72O7N	15.5	790.4488	790.4488 C46 H64 O10 N 15.5	15.5	774.4539	774.4539 C46 H64 O9 N	15.5
	-4H20	798.4908	798.4908 C49 H68 O8 N	16.5	796.4764	796.4764 C49 H66 O8 N	17.5	812.5051	812.5051 C50H70O6N	16.5	772.4407	772.4407 C46 H62 O9 N 16.5		756.4492	756.4492 C46 H62 O8 N	16.5
	-5H2O	780.4820	780.4820 C49 H66 O7 N	17.5	778.4659	778.4659 C49 H64 O7 N	18.5									
	-6H2O				760.4576	760.4576 C49 H62 O6 N	19.5									
	-НСООН										798.4769	798.4769 C45 H68 O11 N 12.5	12.5			
	-CO2-H2O	808.5342	-CO2-H2O 808.5342 C48 H74 O9 N	12.5							782.4808	782.4808 C45 H68 O10 N 12.5	12.5			
	-CO2-2H2O	790.5234	-CO2-2H2O 790.5234 C48 H72 O8 N	13.5			15.5				764.4705	764.4705 C45 H66 O9 N	13.5			
	-CO2-3H2O	772.5134	-CO2-3H2O 772.5134 C48 H70 O7 N	14.5	770.4969	14.5 770.4969 C48 H68 O7 N	16.5									
	-CO2-4H2O				752.4868	752.4868 C48 H66 O6 N										
	-CO2-5H2O															
Group 2	-H20	672.4090	672.4090 C38 H58 O9 N	10.5	670.3928	10.5 670.3928 C38 H56 O9 N	11.5	672.4069 (11.5 672.4069 C38 H58 O9 N		672.4087	672.4087 C38 H58 O9 N	10.5	672.4074	10.5 672.4074 C38 H58 O9 N	10.5
		654.3981	654.3981 C38 H56 O8 N	11.5	652.3829	652.3829 C38 H54 O8 N	12.5							654.3984	654.3984 C38 H56 O8 N	11.5
Group 3 -H2O	-H20	462.3201	462.3201 C27 H44 O5 N	6.5	460.3043	460.3043 C27 H42 O5 N	7.5	462.3203 (462.3203 C27 H44 O5 N		462.3215	462.3215 C27 H44 O5 N	6.5	462.3198	462.3198 C27 H44 O5 N	6.5
		444.3103	444.3103 C27 H42 O4 N	7.5	442.2941	442.2941 C27 H40 O4 N	8.5									
Group 4 -H2O	-H20	362.2682	362.2682 C22 H36 O3 N	5.5	360.2524	360.2524 C22 H34 O3 N	6.5	362.2682	362.2682 C22 H36 O3 N					362.2684	362.2684 C22 H36 O3 N	5.5
		344.2581	344.2581 C22 H34 O2 N	6.5	342.2421	342.2421 C22 H32 O2 N	7.5									
Group 5		262.1797	262.1797 C16 H24 O2 N	5.5	260.1640	260.1640 C16 H22 O2 N	6.5									
Group 6		168 1379	168 1379 C10 H18 O N	2.5	166 1222	166.1222 C10 H16 O N	3.5									



components of the toxin profile (such as AZA-35 and AZA-57), the use of CID fragmentation mode was necessary because of its enhanced sensitivity. CID spectra were dominated mainly by group 1 fragments $[M+H-nH_2O]^+$ (n=1-6) and in some cases by the $[M+H-CO_2-nH_2O]^+$ ions, with the group 2–5 fragments presenting very low intensity (<10 % of the base peak) and the group 6 fragments falling out of the ion stability range (see ESM, Figs. S2 and S3).

SRM experiments on the hybrid triple quadrupole-linear ion trap mass spectrometer excluded the presence of most of the known AZAs (Table 1) in the *A. dexteroporum* extract. However, product ion scans revealed that six of the seven molecules detected by TOF MS were actually AZA-like compounds sharing the typical AZA group 3–5 fragments at m/z 262, 362, and 462, whereas one compound showed the same pattern, but shifted to 2 Da lower masses. CID spectra of all the seven compounds (Fig. 3) provided some structural information of each individual compound.

Following derivatization with TMSDM, all compounds contained in the *A. dexteroporum* acetone extract formed the respective methyl esters, with significantly longer retention times (Table 5). This demonstrated the presence of a carboxylic acid functionality in the molecules. The only exception was presented by AZA-56 and AZA-58, which, however, were among the least abundant compounds and possibly just fell below the LOD, as the respective original peaks (*m*/*z* 884 and 828, respectively) disappeared after TMSDM treatment. These results suggested that none of the azaspiracids contained in the *A. dexteroporum* extract were methyl esters.

The azaspiracids detected in the *A. dexteroporum* strain SZN-B848 and their fragmentation patterns are discussed below listing them in decreasing order of abundance in the extract and molecular weight (MW). Structural hypotheses are summarized in Fig. 4.

Novel AZA-54 (MW = 869 Da)

AZA-54 is a new azaspiracid with $[M+H]^+$ at m/z 870 that represented the major component of the toxin profile of the

Mediterranean A. dexteroporum strain. Compared to AZA-1 (C₄₇H₇₁NO₁₂), its molecular formula (C₄₉H₇₅NO₁₂) indicated an elemental increment of C₂H₄ in the molecule, whereas the ring double bond equivalents (RDB) of the two molecules were identical (Table 2). Apart from that, AZA-54 showed the same characteristic group 2-6 fragments as AZA-1 in both the CID spectrum (Fig. 3A) obtained on the Q-Trap MS and HRMSⁿ spectra obtained on the LTQ-Orbitrap MS (Table 4). This suggested that the structural difference between AZA-54 and AZA-1 was located in the region C1-C9 (Fig. 4). The elemental increment of C₂H₄ could be caused by either the elongation of the side chain of two methylene groups or the presence of one ethyl or two methyl groups or a combination of both. However, the presence of an ion at m/z 808 followed by two water losses in the CID spectrum of AZA-54 (Fig. 3A) pointed to the loss of CO₂ and H₂O from the pseudomolecular ion. This ion was not as abundant as in the case of 3hydroxylated AZAs ([12], revised structures in [15]) but clearly suggested a change in the substitution pattern in the vicinity of the carboxylic acid functionality. The presence of very weak $[M+H-CO_2-nH_2O]^+$ ions (n = 1, 2, 3) was observed also in HR CID/CID MS³ spectra of AZA-54 (Table 4; ESM Fig. S2). Considering that AZA-1, which has a linear side chain, did not eliminate CO₂ neither in CID nor in HR CID/ CID MS³ (Table 3; ESM Fig. S3), the loss of CO₂ from the pseudomolecular ion suggested that the simple elongation of the side chain of AZA-54 with two methylene groups was quite unlikely and that some branching of the alkyl chain was instead occurring (Fig. 4). Interestingly, the intensity of the signal at m/z 808 in both CID (<10 % of the base peak) and HR CID/CID MS³ spectra (<5 % of the base peak) was much lower than intensity of ions due to CO₂ loss in 3-hydroxylated AZAs, likely as a consequence of the different effect on fragmentation of the alkyl group versus the hydroxyl group.

Novel AZA-55 (MW = 867 Da)

This new azaspiracid with $[M+H]^+$ at m/z 868 and molecular formula $C_{49}H_{73}NO_{12}$ was the second most abundant

Table 5 Retention time (Rt) shifts of azaspiracids contained in *A. dexteroporum* acetone extract on Q-Trap MS

	[M+H] ⁺ an original ex	nd Rt of AZAs in the	[M+H] ⁺ and I TMSDM deri	Rt of AZA methyl esters after vatization
Azaspiracid (AZA)	m/z	Rt (min)	m/z	Rt (min)
AZA-54	870	12.8	884	13.3
AZA-55	868	12.5	882	13.9
3-epiAZA-7	858	11.5	872	12.1
AZA-56	884	14.8	898	_
AZA-57	844	11.5	858	12.8
AZA-58	828	12.1	842	_
AZA-35	830	12.2	844	13.6



Fig. 4 Structural hypothesis for the seven azaspiracids contained in the Mediterranean *A. dexteroporum* strain based on the analysis of their fragmentation patterns versus AZA-1. The regions of AZA-1 *circled in red* are those where structural modifications occur

AZA-54: Branching at C2 or C3 with C2H4 (either 2 methyls or 1 ethyl)

AZA-55: 1 double bond in ring H or I

Branching at C2 or C3 with C2H4 (either 2 methyls or 1 ethyl)

3-epiAZA-7: 1 OH at C3

AZA-56: elongation of the side chain with C₃H₆ (3 methylenes)

AZA-57: 1 CH₂ less in the side chain 1 OH more at C2 or C3

AZA-58: 1 CH₂ less in the side chain

AZA-35: 1 CH₂ less in the side chain No double bond in the side chain

component of the Mediterranean *A. dexteroporum* strain. It presented two H less and one unsaturation more than AZA-54 (Table 2). Its fragmentation pattern obtained by CID (Fig. 3B) and HRMSⁿ (Table 4) was almost identical to that of AZA-54 with the only exception that all group 1–6 fragments were downshifted of 2 Da compared to the corresponding fragments of AZA-54. This suggested that AZA-55 contained an additional double bond in ring H or I. Similarly to AZA-54, very weak $[M+H-CO_2-nH_2O]^+$ ions (n = 3, 4) were observed also in CID and HRMS³ spectra, suggesting that AZA-55 also presented an alkyl branching in the vicinity of the carboxylic acid functionality. Thus, AZA-55 is an H/I ring unsaturated derivative of AZA-54 (Fig. 4).

Novel 3-epi-AZA-7 (MW = 857 Da)

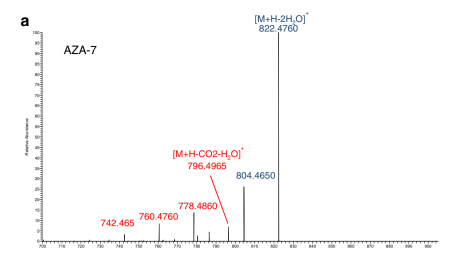
The third most abundant component of the Mediterranean A. dexteroporum toxin profile had been tentatively assigned to AZA-7 by Percopo et al. [13]. This compound with $[M+H]^+$ at m/z 858 had indeed the same molecular formula (C₄₇H₇₁NO₁₃) and RDB value as AZA-7 [5], thus differing from AZA-1 (C₄₇H₇₁NO₁₂) just by the presence of one additional O atom. The fragment ions previously detected by in-source fragmentation on TOF MS [13] were all confirmed by CID analysis on the Q-Trap MS (Fig. 3C) and by HR HCD MS² analysis on the LTQ-Orbitrap FTMS (Table 3; ESM Fig. S1). In more detail, its fragmentation pattern contained the same characteristic group 2-6 fragments as AZA-1 and AZA-7 [5]. This indicated that the compound at m/z 858 shared the B to I ring system with AZA-1 and AZA-7. Thus, compared to AZA-1, both the compound at m/z 858 and AZA-7 have the additional oxygen (likely a hydroxyl group) in the region C1-C9. In contrast to AZA-1, this compound displayed a very complex group 1 fragment cluster characterized by elimination of CO2 and subsequent water losses from the pseudomolecular ion (Fig. 3C, Table 3). These fragments appeared also in the CID MS spectrum of AZA-7 reported by James et al. [5] as minor unassigned fragments at m/z 760 and m/z 742. This clearly suggested that both the compound at m/z 858 and AZA-7 were hydroxylated at C3 (Fig. 4); otherwise, like in most AZAs, the CO₂ elimination would not have been observed ([12], revised structures in [15]). However, a comparative LC-HRMSⁿ analysis of the compound at m/z 858 versus a reference sample of AZA-7 isolated from shellfish allowed to exclude the identity of the two compounds based on retention times and relative ratios of the [M+ H-nH₂O]⁺ and [M+H-CO₂-nH₂O]⁺ ions contained in the group 1 fragments. The compound at m/z 858 eluted 0.55 min later than AZA-7; in addition, group 1 fragments of the former were dominated by the [M+H-CO₂-H₂O]⁺ ion whereas group 1 fragments of AZA-7 were dominated by [M+H-2H₂O]⁺ ion (Fig. 5). Such evidence suggested that the A. dexteroporum compound was not an unrelated compound but rather a stereoisomer of AZA-7 and that the steric difference is probably located at C3 because this is the only position that could explain the different intensities of the CO₂ losses in both compounds. This is consistent with the compound at m/z 858 being the epimer at C3 of AZA-7 (3-epi-AZA-7). Noteworthy, the finding of 3-epi-AZA-7 in the Mediterranean A. dexteroporum represents the second case of de novo biosynthesis of a 3hydroxylated AZA in a dinoflagellate, the first case being A. poporum, which can produce AZA-11 [14]. Until recently, 3-hydroxylated AZA were regarded exclusively as shellfish metabolites of AZA-1 and AZA-2 [9].

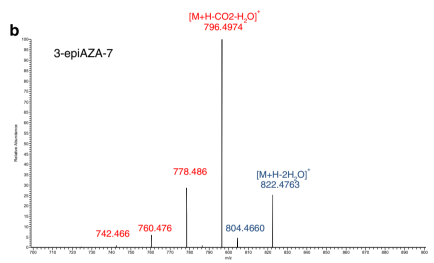
Novel AZA-56 (MW = 883 Da)

AZA-56 is a new azaspiracid with $[M+H]^+$ at m/z 884. Its molecular formula $(C_{50}H_{77}NO_{12})$ showed an elemental increment of C_3H_6 compared to that of AZA-1 and the same RDB value (Table 2). In CID spectrum (Fig. 3D), AZA-56 showed the same characteristic group 2–6 fragments as AZA-1, most of the fragments being confirmed by HRMS³ analyses (Table 4). This suggested that the additional C_3H_6 is located in the region C1–C9 of AZA-1. The absence of other ions deriving from the further fragmentation of the alkyl chain



Fig. 5 HR CID/CID MS³ spectra of **A** the shellfish metabolite AZA-7 and **B** the 3-epiAZA-7 contained in the Mediterranean *A. dexteroporum* extract. Expansion of the mass region *m/z* 700–900 where group 1 fragments appear including [M+H-*n*H₂O]⁺ ions (in *blue*) and [M+H-CO₂-*n*H₂O]⁺ ions (in *red*)





(e.g., the elimination of CO₂ from the pseudomolecular ion and subsequent water losses) suggested that elongation of the side chain of three methylene groups occurs in AZA-56 (Fig. 4).

Novel AZA-57 (MW = 843 Da)

AZA-57 is a new azaspiracid with [M+H]⁺ at m/z 844 and a minor component of the Mediterranean A. dexteroporum toxin profile. Its molecular formula (C₄₆H₆₉NO₁₃) contained one O atom more and one methylene group less than AZA-1 and presented its same RDB value (Table 2). The CID spectrum (Fig. 3E) contained the same group 2–5 fragments as AZA-1, with group 2 and 3 fragments being confirmed also by HRMSⁿ spectra (Table 4). This suggested that the structural modifications in AZA-57 occur in the region C1–C9 of AZA-1. Furthermore, the fragmentation pattern was dominated by the loss of CO₂ and consequent water losses from the pseudomolecular ion (Table 4), which could suggest that AZA-57 is a 3-

hydroxylated derivative of AZA-1. However, a fragment ion at *m/z* 798.4769 was also present, which corresponded to [M+H-HCOOH]⁺ and had never been observed before in any AZA fragmentation spectrum. On the whole, these results suggested that AZA-57 lacks one methylene in the side chain and is hydroxylated in close proximity of the carboxylic group, which could be either C2 or C3, the latter resulting in an enol functionality (Fig. 4). However, the exact position of the hydroxylation cannot be determined by MS alone because this is an unprecedent case.

Another compound contained at trace levels in the extract eluted about 3 min later than AZA-57 (Rt = 13.18 min on system #2) and had the same nominal mass as AZA-57 but a different elemental composition ([M+H]⁺ ion at m/z 844.5202, RDB = 11.5, C₄₇H₇₄NO₁₂, Δ = -0.418 ppm). Due to low abundance, the HRMS³ spectrum contained just the group 1 and 3 fragments suggesting it is an azaspiracid but hampering further structural hypotheses.



Novel AZA-58 (MW = 827 Da)

AZA-58 is a new azaspiracid with $[M+H]^+$ at m/z 828 which had been tentatively assigned to AZA-3 in Percopo et al. [13], having the same molecular formula (C₄₆H₆₉NO₁₂) and exact mass. However, AZA-58 eluted at a different retention time than AZA-3 and had a different structure as revealed by its fragmentation pattern (Fig. 3F, Table 4). Indeed, AZA-58 showed group 2 and 3 fragments identical to AZA-1 (m/z 672 and 462, respectively) and different from AZA-3 (m/z 658 and 448, respectively), whereas the group 4 fragment (m/z 362) was identical for all the three molecules. This fragmentation suggested that AZA-58 differed from AZA-1 just by the lack of a methylene group in the region C1–C9. We can confidently exclude the lack of a methylene group in the ring A since this would lead to group 2 fragments different from AZA-1. Therefore, most likely AZA-58 lacks one methylene in the alkyl chain (either C2 or C3), thus being the normethylene derivative of AZA-1 (Fig. 4).

AZA-35 (MW = 829 Da)

The compound with $[M+H]^+$ at m/z 830 represents a very minor component of the Mediterranean A. dexteroporum toxin profile. The molecular formula (C₄₆H₇₁NO₁₂) contained just two H atoms more than that of AZA-58 and is the same as that of AZA-38 from Amphidoma languida (compound 3 reported by Krock et al. [12]). Based on the fragmentation pattern contained in its CID spectrum (Fig. 3G) and HRMSⁿ (n = 2– 3) spectra (Table 3), we excluded that this compound was AZA-38 since it lacked the characteristic group 2–5 fragments at m/z 686, 448, 348, and 248 of the latter. On the contrary, this compound presented the same group 2–5 fragments as AZA-1 and AZA-58, which suggested that, similarly to AZA-58, it presented the same B to I ring systems as AZA-1 and thus lacked a methylene group in the region C1–C9. The structural modification must be located in the methylene chain (C2–C3) for the same reasons discussed for AZA-58. In addition, this compound also presented group 2-5 fragments identical to AZA-58 and AZA-1 but a lower RDB value (11.5 versus 12.5 of AZA-1 and AZA-58), which suggested that in this molecule the double bond either in the side chain or in the A ring was saturated. The latter hypothesis could be excluded. Indeed AZA-37, which has a saturated A ring ([12], revised structures in [15]), produces group 2 fragments due to elimination of a C7 part structure, which results from a simultaneous cleavage between C6 and its adjacent oxygen and between C7 and C8, and not between C9 and C10 as occurring in AZA-1 (Fig. 1), in AZA-58, as well as in AZA-35. The whole of these observations suggested that this compound, compared to AZA-1, presented one methylene less and one double bond less in the side chain, thus being the saturated form of AZA-58 (Fig. 4). Recently, Kilkoyne et al. [29] have reported AZA-35 from *A. spinosum* having the same molecular formula and fragmentation behavior as the compound we detected. Availability of an *A. spinosum* extract containing AZA-35 allowed us to confirm the identity of this compound as AZA-35 based on retention time (system #2) and HR CID/CID MS³ data, suggesting that AZA-35 is the nor-methylene 4.5-dehydroAZA-1.

Quantitative analyses

The relative abundances of the AZAs detected in the Mediterranean strain of A. dexteroporum were determined in decreasing order as follows: 65.3 % of AZA-54, 18.0 % of AZA-55, 14.9 % of 3-epiAZA-7, 0.6 % of AZA-58, 0.5 % of AZA-35, 0.3 % of AZA-57, and 0.2 % of AZA-56 (Fig. 6). The quantification was based on the SRM mass transitions between the pseudomolecular mass and the first water loss of each AZA (Table 1) on the reasonable assumption that the ratios of these transitions are similar among all AZAs. Nevertheless, it has to be taken into account that this quantification is only a good approximation since the exact molecular responses of the individual AZAs of this strain are still unknown. However, it is clear that the AZA profile of the Mediterranean strain of A. dexteroporum is dominated by AZA-54 making up approximately two thirds of total AZAs. The second most abundant component is AZA-55, which differs from AZA-54 only by an additional unsaturation in the ring H/I. This means that the AZA profile of A. dexteroporum is dominated for over 80 % by compounds with an alkyl branching (AZA-54 and AZA-55), a structural feature completely absent in other AZA-producing species. Also the third most abundant 3-epi-AZA-7 has never been observed in dinoflagellates. The other four AZAs of the Mediterranean A. dexteroporum together only made up less than 2 % and accordingly can be regarded as minor compounds.

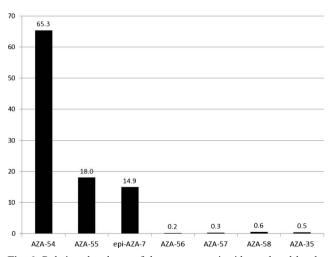


Fig. 6 Relative abundance of the seven azaspiracids produced by the Mediterranean strain of *A. dexteroporum* expressed in percent



Conclusions

This study confirmed the Mediterranean A. dexteroporum as an azaspiracid-producing species and provided a better characterization of the three molecules initially detected [13] in the strain, along with the demonstration of the presence of at least seven azaspiracids. The three compounds at m/z 828, 830, and 858 initially hypothesized by Percopo et al [13] to be AZA-3 [7], compound 3 [12], and AZA-7 [5], respectively, were proven to be different compounds, two of which were so far undescribed. Based on retention time, exact masses, and fragmentation patterns, six out of the seven AZAs produced by the Mediterranean A. dexteroporum strain proved to be new molecules (3-epi-AZA-7, AZA-54 to AZA-58) and one was a previously reported compound (AZA-35). This is the first time that an epimer of the shellfish metabolite AZA-7 is reported in a dinoflagellate extract and that AZA-35 is reported in an Azadinium species other than A. spinosum [29]. Overall, the results obtained in this study show that MS and fragmentation behavior of AZAs on different LC-MS systems may produce complementary data on these molecules, which allow to characterize them reliably.

All the molecules studied in this work, except AZA-55, seem to have the same A to I-ring system as AZA-1, suggesting that the differences among these molecules are all located in the carboxylic side chain. Recently, four new compounds were discovered in Amphidomataceae, all showing the same CID fragmentation patterns with a conserved E-I ring system in common [12]. All this evidence suggests that Amphidomataceae have similar biosynthetic pathways that produce molecules with a relatively conserved polyketide skeleton [30] and a more variable part towards the carboxylic end. Branching of the side chain observed in AZA-54 and AZA-55 has never been observed in any known azaspiracid and proved to cause low abundance of CO₂ loss (<10 %) in contrast to 3-hydroxylation which results in a highly abundant CO₂ loss.

It is worth noting that 3-hydroxylated AZAs have been long thought to be shellfish metabolites of AZA-1 and AZA-2 resulting from enzymatic hydroxylation in the shellfish at C3 [5, 9]. However, recently, Krock et al. [15] found the 3-hydroxylated derivatives AZA-36 and AZA-37 to be de novo synthesized in *A. poporum* strains from the North Sea and Korea, respectively. In addition, some strains of *A. poporum* from the Northwest Pacific region only produced AZA-11, the 3-hydroxylated derivative of AZA-2, without producing AZA-2 itself [14]. Interestingly, also the Mediterranean *A. dexteroporum* produces 3-epi-AZA-7, the 3-hydroxylated derivative of AZA-1 without producing AZA-1 itself. So, our results show that the Mediterranean *A. dexteroporum* is another

species that can produce de novo 3-hydroxylated AZAs which further supports the hypothesis that in some Amphidomataceae an enzymatic pathway leading to 3-hydroxylation exists.

The molecules found in the Mediterranean A. dexteroporum strain further add to the number of so far known AZAs directly produced by microalgae, whereas the majority of the known AZAs is still considered to be transformation products within shellfish. Compared to other Amphidomataceae species, for which only a few azaspiracids have been reported, the diversity of AZAs in the Mediterranean A. dexteroporum strain seems to be exceptional. However, a high diversity of AZAs is now also known from A. spinosum [17, 29] and future studies using the multi-platform approach or computational approaches such as molecular networking [31] might show this for other species/strains as well. The presence of so many AZAs is even more surprising if compared to the complete absence of AZAs in the Arctic strain of the same species [27], although cases of intraspecific differences in toxin profiles are widespread [14, 32].

Recent studies on the toxicity of some purified AZA analogues relative to AZA-1 showed that all analyzed AZAs were toxic and that AZA-2, AZA-3, and AZA-6 were more toxic than AZA-1 by 8.3, 4.5, and 7 times, respectively [10], 37-epi-AZA-1 was 5 times more toxic [11], AZA-33 and AZA-34 were 5 times less toxic and 5 times more toxic, respectively [17], and AZA-36 and AZA-37 were 6 and 3 times less toxic, respectively [15]. As major differences in toxicity can be related to slightly different structures of AZAs, the actual toxicity of the different compounds produced by the Mediterranean A. dexteroporum is worthy of consideration in future toxicological studies. In the meantime, for a correct evaluation of the AZA-related risk in the Mediterranean area, the presence of the azaspiracids produced by A. dexteroporum should be taken into account in the monitoring programs of these marine biotoxins. This seems even more important considering that none of the AZAs currently legislated (AZA-1, AZA-2, and AZA-3) and hence regularly monitored in shellfish is actually produced by this strain.

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Compliance with ethical standard This paper does not raise any concern regarding human and animal rights and does not involve animals or humans.

Conflict of interests There is no conflict of interest pending on this paper



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