provided by Electronic Publication Informa

Amphidomataceae

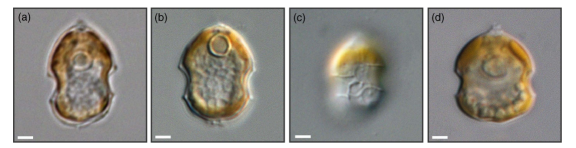


Figure 1 LM micrographs of Azadinium spinosum (a), Az. poporum (b), Az. dexteroporum (c) and Amphidoma languida (d). Scale bars = $2 \mu m$.

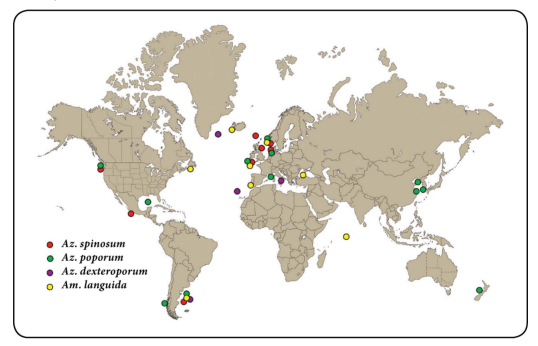


Figure 2 Global records of the four species of Amphidomataceae known to produce azaspiracids (AZA).

AZA producer	no AZA found	not analysed yet
Azadinium spinosum	Azadinium obesum	Azadinium caudatum var. caudatum
Azadinium poporum	Azadinium polongum	Azadinium luciferelloides
Azadinium dexteroporum	Azadinium caudatum var. margalefii	Amphidoma nucula
Amphidoma languida	Azadinium dalianense	Amphidoma acuminata
	Azadinium trinitatum	Amphidoma curtata
	Azadinium cuneatum	Amphidoma depressa
	Azadinium concinnum	Amphidoma elongata
	Azadinium zhuanum	Amphidoma laticincta
	Amphidoma parvula	Amphidoma obtusa
		Amphidoma steinii

576 Harmful Algal Blooms: A Compendium Desk Reference

Amphidomataceae

General: Azaspiraids (AZA) are a group of lipophilic polyether toxins first detected and described in the late 1990s. With the description of *Azadinium spinosum* in 2009, the first source organism has been identified. Currently, there are four out of 22 species of the genera *Azadinium* and *Amphidoma* (merged in the family Amphidomataceae) that have been shown to produce AZA. However, it has to be kept in mind that there is only a limited number of cultured strains available, and it is thus not clear if and to what extent AZA production is a species-specific stable phaenotypic trait.

General Morphology: All AZA-producing species of Azadinium and Amphidoma languida are small (size of about 10-16 µm) and ovoid to elliptical in shape with a hemispherical hyposome. In all these species, the episome is larger than the hyposome, with slightly convex sides ending in a distinctly pointed apex. The cingulum is deep and wide, accounting for roughly 1/5 to 1/4 of the cell length. A central or more posteriorly located large nucleus is visible, which generally is round to elliptical but may become distinctly elongated in shape close to cell division. All species are photosynthetic and possess a presumably single chloroplast, which is parietally arranged, lobed, and normally extends into both the epi- and hyposome. For all of the AZA-producing species, stalked pyrenoid(s) are visible in the light microscope because of a distinct starch cup. Azadinium spp. and Amphidoma languida have delicate thecal plates difficult to detect in light microscopy (LM), so that live cells are sometimes difficult to differentiate from small athecate gymnodinoid species. Plate pattern and thecal plate details are important for determination of the genus and species, but require scanning electron microscopy (SEM). Species of Azadinium are characterised by the Kofoidean plate pattern of Po, cp, X, 3–4', 2–3a, 6", C6, 5S, 6"", 2^{''''}, whereas *Amphidoma languida* has six apical plates and no anterior intercalary plates. A very characteristic feature among the AZA-relevant species is the prominent apical pore complex visible in LM, which is composed of an X-plate and a pore plate with a central round pore covered

by a cover plate. Plate details important for species identification include the presence/absence and/or location of a single antapical spine and primarily the position of a ventral pore. Morphology, and in particular the plate tabulation with five different rows of plates, undoubtedly classified the family Amphidomataceae as a member of the dinophycean subclass Peridiniphycidae (Tillmann et al., 2009). The relation to one of the two orders of the subclass (i.e. Gonyaulacales and Peridiniales), however, is less clear as some morphological traits imply affinity to Peridinales and others to Gonyaulacales. Using a concatenated alignment of LSU and SSU, the Amphidomataceae have been placed on the peridinean branch remote from the Gonyaulacales, but the true relation to Peridiniales could not be identified reliably (Tillmann et al., 2014). It thus remains to be determined whether they are part of the Peridiniales or represent a distinct lineage that would deserve the recognition at a higher taxonomic level.

- Known Distribution: Although the first species of *Azadinium* were initially described from the North Sea, there is increasing evidence that AZA-producing species have a wide geographical distribution. Nevertheless, knowledge on the biogeography of the genus or of certain species currently is rather limited and patchy. It is based on the troublesome procedure of isolating, cultivating and fully characterizing local strains; on a very few records of species detected by scanning plankton samples by electron microscopy; or on positive signals using species-specific molecular detection methods.
- **Cysts:** Knowledge on the life cycle of *Azadinium* and/ or *Amphidoma* is quite incomplete. Successful isolation of *Az. poporum* by incubating sediment samples (Potvin *et al.*, 2012; Gu *et al.*, 2013) made the presence of cysts quite likely for that species, and that has been confirmed by Gu *et al.* (2013): in one out of 25 cultured strains, they observed the presence of a few distinct cysts. These cysts are ellipsoid, around 15 μ m long and 10 μ m wide, and are filled with pale granules and a yellow accumulation body. Likewise, the species *Az. polongum* (a non-AZA producer) has been described to produce cysts in culture,

round cells of $10-16 \,\mu\text{m}$ in diameter and with pale white inclusion. No cyst-like cells have been reported for other species, including *Az. spinosum*, *Az. dexteroporum* and *Am. languida*. Clearly, more data and observations are needed to clarify the whole life cycle of Amphidomataceae.

Toxin: Species of Amphidomataceae are the source of azaspiracids (AZA), a class of polyether toxins discovered almost 20 years ago. Azaspiracids are polyketides with a highly hydroxylated carbon chain that is cyclised by ether bridges, and they contain a six-membered cyclic secondary amino ring. To date more than 50 AZA analogs are known. These include about 20 of dinoflagellate origin, and the others are thought to be produced by bioconversion in shellfish (Hess et al., 2014). Azaspiracids are known to be responsible for gastrointestinal disorders with the consumption of AZA-contaminated shellfish, with symptoms quite similar to those of DSP, such as nausea, vomiting, diarrhea and stomach cramps. Preliminary studies of AZA suggested that these compounds are highly toxic with multi-organ damage in mice and teratogenic potential to developing fish, along with a wide array of cellular-level effects, ranging from cytotoxicity to apoptosis and to effects on the hERG potassium channel (reviewed by Twiner et al., 2014). Minimal lethal doses (i.p. mice) for the most dominant AZA in mussels have been determined as 200, 110, and 140 μ g/kg for AZA-1, -2, and -3, respectively (Satake et al., 1998; Ofuji et al., 1999). Consequently, a regulatory limit of 160 µg/kg mussel meat for AZA-1 to AZA-3 was implemented in 2002 into the EU biotoxin legislation. More recent studies vielded similar results for mouse toxicity for AZA-2 and AZA-3, but a distinctly lower dose (higher toxicity) of 74 µg/kg for AZA-1 (Kilcoyne et al., 2014a). Oral mouse studies indicated no additive or synergistic effects when AZA was administered in combination with okadaic acid or yessotoxin (Kilcoyne et al., 2014a).

Around 20 AZA analogs are currently described to be of dinoflagellate origin. Among the dominant AZA found in shellfish, AZA-1 and AZA-2 are produced by *Azadinium*, whereas no planktonic source of AZA-3 is

known yet. An increasing number of new AZA are discovered in the Amphidomatacean cultures. Initial mass spectral data (Krock *et al.*, 2012) as well as structural elucidation by nuclear magnetic resonance (NMR) spectroscopy (Kilcoyne *et al.*, 2014b; Krock *et al.*, 2015) showed that some of the new AZA discovered in dinoflagellates are structurally unique from previously reported analogues by having a modification of the nitrogen-containing I-ring of the molecule, which consists of either a missing methyl group at C39 or an additional double bond.

All four described European strains of *Az. spinosum* have the same toxin profile consisting of AZA-1, -2, and -33 (Tillmann *et al.*, 2012b), and a few minor compounds have additionally been found in the Scottish strain (Kilcoyne *et al.*, 2014b). For *Az. poporum*, a larger number of strains from different areas around the globe have been described, and this is reflected by a considerable diversity within this species in terms of toxin profiles. Whereas all three available North Sea strains produce AZA-37, *Az. poporum* from the Asiatic Pacific region produces more complex AZA profiles, including AZA-2, -11, -36, -40, -41 in different combinations, and also strains without any known AZA have been described.

Azaspiracid-2 (AZA-2) is the major AZA produced by *Az. poporum* from Argentina and by a strain from the Mediterranean, whereas strains from the Pacific coast of Chile produce AZA-11. Most recently, the new AZA-59 was identified from *Az. poporum* strains isolated from Puget Sound, WA (Kim *et al.*, 2017). A feature that is shared among some Asian Pacific and Argentinean strains of *Az. poporum* is the production of minor amounts of AZA-related compounds with higher molecular masses. For the Argentinean strains, one of these compounds has been identified as AZA-2 phosphate, which is the first report of a phosphated marine algal toxin (Tillmann *et al.*, 2016).

The presence of AZA has also been unambiguously described for the Mediterranean strain of *Az. dexteroporum* (Percopo *et al.*, 2013), and detailed LC-MS analysis confirmed the presense of six novel AZA and AZA-35 (Rossi *et al.*, 2017). A new strain of *Az. dexteropo*-

rum, isolated from the subarctic Irminger Sea, however, clearly lacked any of these or other known AZA (Tillmann *et al.*, 2015).

The type strain of *Amphidoma languida* isolated from Ireland and a strain originating from the Iceland area produce AZA-38 and -39 (Krock *et al.*, 2012; Tillman *et al.*, 2015). In contrast, *Am. languida* from the Atlantic coast of southern Spain produce AZA-2 and -43 (Tillman *et al.*, 2017).

Cell quotas of AZA were found to be variable within and among strains and species but are typically in the range of 5–20 fg cell⁻¹. A maximum value of 220 fg cell⁻¹ for *Azadinium spinosum* grown at 10 °C was reported (Jauffrais *et al.*, 2013).

In vitro toxicity along with structure elucidation for some of the new AZA detected in *Az. spinosum* (Kilcoyne *et al.*, 2014b) and *Az. poporum* (Krock *et al.*, 2015) have recently been determined, and they showed both lower and higher cytotoxicity compared to AZA-1. For other compounds (e.g. AZA produced by *Am. languida* [AZA-38, -39] and *Az. dexteroporum*), specific toxicity is not known yet.

Methods for Toxin Identification: In 2011, the EU replaced the mouse bioassay with LC-MS/MS as the primary monitoring method for the analysis of AZA (and other lipophilic toxins) in shellfish. A number of validated LC-MS/MS methods for detection and quantification of AZA in shellfish have been described (Hess *et al.*, 2014). Work on alternative detection methods for AZA has been limited. An antibody-based ELISA assay, as a rapid analytical technique using inexpensive instrumentation, has recently been described as a suitable tool for shellfish toxin analysis (Samdahl *et al.*, 2015).

Ecological Observations: As species of Amphidomataceae have only recently been detected and identified, knowledge on their biology and ecology is rather limited. A first set of growth experiments indicated that *Az. spinosum* was fairly easy to grow with a number of standard culture media (indicating no special nutritional requirement) and at a wide range of different salinities, temperatures, and light conditions. Quantitative abundance data of toxic Amphidomataceae are hardly available, but dense blooms (> 10^6 cells L⁻¹) from a species of Azadinium from the Argentinean shelf have been observed (Akselman and Negri, 2012). Pathway and transfer kinetics of AZA into bivalve molluscs are just getting started to be explored. Azaspiracid accumulation in mussels following direct feeding on Az. spinosum has been proven experimentally, but Az. spinosum also had a significant negative effect on mussel feeding behavior and slightly increased mussel mortality compared to a control food (Jauffrais et al., 2012). Azaspiracids have been detected in a number of micrograzers (e.g., Protoperidinium crassipes, Favella ehrenbergii), so that a role of plankton vectors for mussel intoxication needs to be explored.

General Notes: With their small size, their distinctive and species-specific morphological characteristics that are hardly or not at all visible at the LM level, and with the close resemblance of toxigenic and non-toxigenic species, the AZAproducing Amphidomataceae are a good example for the necessity of applying molecular detection methods in monitoring and early warning systems. Molecular probes have been developed for the first three described species, Az. spinosum, Az. poporum, and Az. obesum (Toebe et al., 2013), but specific probes for other AZA-producing species (Az. dexteroporum and Am. languida) are still missing. In addition, it has to be kept in mind that there probably are more AZA-producing species that are not yet identified. Am. languida, for example, is the only species of the genus Amphidoma known so far for AZA production, and there are eight more species described, for which AZA production cannot be excluded. A general probe recently developed to detect a broad range of Amphidomataceae will be helpful to screen field samples and to aid in the detection, isolation and characterisation of AZA-producing species (Smith et al., 2016).

Azadinium spinosum Elbrächter et Tillmann (Tillmann et al., 2009)

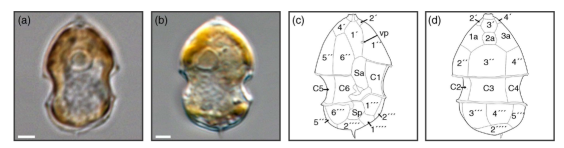


Figure 3 Az. spinosum LM micrographs (a, b) and schematic drawings (c, d) including the thecal plates in Kofoidean notation (vp = ventral pore). Scale bars = $2 \mu m$.

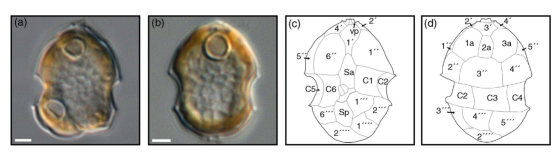
Synonyms: None.

Morphology: *Azadinium spinosum* is a small $(12-16 \,\mu\text{m} \text{ length} \text{ and } 7-11 \,\mu\text{m} \text{ width})$, slender (length-width ratio = 1.6), and slightly dorsoventrally compressed thecate, photosynthetic dinoflagellate. The conical episome with convex sides ends with a conspicuous apical pore complex (APC) and is larger than the hemispherical hyposome. It has a wide and descending cingulum, which is displaced by about half its width. In the light microscope, one large pyrenoid visible by its starch sheath is located in the episome. Eponymous for the species is the presence of a single small antapical spine located slightly asymmetrically at the right side of the cell.

Plate pattern and thecal plate details are important for determination of the genus

and species, but require SEM. The Kofoidean thecal tabulation of *Az. spinosum* is Po, cp, X, 4', 3a, 6'', 6C, 5S, 6''', 2''''. *Az. spinosum* has a distinct ventral pore located on the left side of the first apical plate.

Distribution: *Azadinium spinosum*, the type of the genus, has been isolated off the Scottish coast, the coast off Denmark, the Shetland Islands, the Norwegian coast, and from coastal Atlantic waters in Ireland. A species of *Azadinium* most likely *Az. spinosum* has been recorded in SEM samples from coastal Pacific waters off Mexico. *Az. spinosum* has also been identified in SEM field samples from the Argentinean shelf (South Atlantic). Recently *Az. spinosum* was detected by qPCR from Puget Sound, WA, U.S.



Azadininium poporum Tillmann et Elbrächter (Tillmann et al., 2011)

Figure 4 Az. poporum LM micrographs (a, b) and schematic drawings (c, d) including the thecal plates in Kofoidean notation (vp = ventral pore). Scale bars = $2 \mu m$.

Synonyms: None.

Morphology: *Azadininium poporum* is small (11–16 μ m length, 8–12 μ m width), ovoid (length-width ratio = 1.3), slightly dorsoventrally compressed, with a broad and slightly descending cingulum, and with a hyposome slightly smaller than the episome ending in a

conspicuous APC. In *Az. poporum*, there may be several (up to four) pyrenoids with a starch sheath visible in LM located in both the epiand hyposome. The most distinctive morphological feature of *Az. poporum* requires SEM; it is the characteristic position of the ventral

pore, which is located anterior at the cell's left side of the pore plate at the junction with the first two apical plates.

Distribution: *Azadinium poporum* was described based on strains from the North Sea off Denmark and has also been recorded in Ireland and along the Norwegian coast. A number of strains have been obtained from outside Europe. *Az. poporum* obviously is quite widely distributed in the Asian Pacific. As a first record of *Azadinium* in Pacific waters, *Az. poporum* has been isolated from Shiwha Bay in Korea, and subsequently, 25 different strains of *Az. poporum* originating from China covering the Bohai Sea and the East and South China Seas were established. Most recently, *Az. poporum* was detected in New Zealand both by qPCR and by establishing a culture. Likewise, *Az. poporum* cultures were obtained from samples from the South Atlantic (Argentina), the South Pacific (Chile), and the Gulf of Mexico. Most recently *Az. poporum* was identified by qPCR and isolated strains from Puget Sound, Washington.

Azadinium dexteroporum Percopo et Zingone (Percopo et al., 2013)

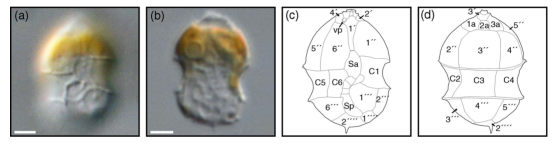


Figure 5 Az. dexteroporum LM micrographs (a, b) and schematic drawings (c, d) including the thecal plates in Kofoidean notation (vp = ventral pore). Scale bars = $2 \mu m$.

Synonyms: None.

Morphology: *Azadinium dexteroporum* is the smallest species of *Azadinium* (7.0–10.0 μ m in length and 5.0–8.0 μ m in width). Cells are slightly elongated (length-width ratio = 1.4) and dorso-ventrally compressed, with the episome longer and slightly larger than the hyposome. The hyposome is slightly asymmetrical, with a small spine located in its posterior right side. The cingulum is deeply excavated and notably wide. One pyrenoid visible by its starch cup is present in the episome. Species-specific morphological details visible at the SEM level include the characteristic arrangement of the ventral pore, which is located at the right posterior end of the

markedly asymmetric pore plate. A pronounced concavity of the median intercalary plate 2a has been highlighted as a peculiar feature of the Mediterranean type material, but this plate was plain for a subarctic strain originating from the Irminger Sea.

Distribution: Azadinium dexteroporum was initially described from the Mediterranean (Naples), but a new strain representing the species was recently obtained from the Subarctic (Irminger Sea). Az. dexteroporum was also identified in SEM preparation of spring bloom samples from the South Atlantic (Argentinean shelf) and is on a species list (as Az. cf. dexteroporum) of Madeira (North Atlantic off Morocco).

Amphidoma languida Tillmann, Salas et Elbrächter (Tillmann et al., 2011)

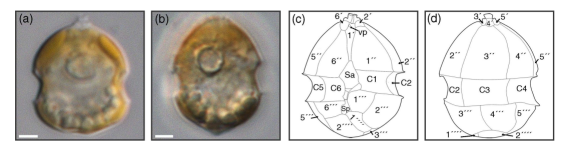


Figure 6 Am. languida LM micrographs (a, b) and schematic drawings (c, d) including the thecal plates in Kofoidean notation (vp = ventral pore). Scale bars = $2 \mu m$.

Synonyms: None.

- Morphology: Cells of Amphidoma languida are ovoid to slightly elliptical (length-width ratio = 1.3), with a conical episome and a distinctly pointed APC. Cells are small $(12.9-15.5 \,\mu\text{m} \text{ in length and } 9.7-14.1 \,\mu\text{m})$ in width). The episome is slightly larger than the spherical hyposome, which ends in a pointed antapex. At the light microscope level Am. languida is very similar to small species of Azadinium. Electron microscopy, however, reveal major differences in plate pattern, with a Kofoidean plate pattern of Po, cp, X, 6', 0a, 6", 6C, 5S, 6"", 2"". Am. languida, as other species of the genus Amphidoma, has thus 6 apical and no anterior intercalary plates, whereas Azadinium has 3-4 apical plates and 2-3 anterior intercalary plates. Other specific details visible with SEM are the presence of a large antapical pore (which in fact is a field of a number of small pores) and the location of a ventral pore on the anterior right side of the first apical plate.
- Distribution: The AZA-producing species Amphidoma languida has first been isolated from a bay in Ireland, but definitely has a much wider distribution. Sequence data from plankton samples of the Skagerrak area and strains of this species from the Norwegian coast and from Iceland indicate the presence of Am. languida in the North Sea and the North Atlantic as well. More recently, it has been observed in SEM from a seawater sample collected at Saint-Pierre and Miquelon in 2012 and at several sampling locations along the southern coast of the Black Sea in 2014. In contrast to the shallow coasts of Ireland and the Subarctic near Iceland, cells most likely determinable as Am. languida have been observed in SEM from a sample collected at the open West Indian Ocean as well. Moreover, Am. languida was present in a 1991 bloom sample from the Argentinean shelf. Finally, a culture of Am. languida has been established from water off the Atlantic coast of southern Spain.

References

- Akselman, R., and A. Negri. 2012. Blooms of *Azadinium* cf. *spinosum* Elbrächter et Tillmann (Dinophyceae) in northern shelf waters of Argentina, Southwestern Atlantic. *Harmful Algae*, **19**: 30–38.
- Gu, H., Z. Luo, B. Krock, M. Witt, and U. Tillmann. 2013. Morphology, phylogeny and azaspiracid profile of *Azadinium poporum* (Dinophyceae) from the China Sea. *Harmful Algae*, **21–22**: 64–75.
- Hess, P., P. McCarron, B. Krock, J. Kilkoyne, and C.O. Miles. 2014. Azaspiracids: chemistry, biosynthesis, metabolism, and detection. In: *Seafood and Freshwater Toxins*. L.M. Botana (Ed.). CRC Press, Boca Raton: p. 799–821.
- Jauffrais, T., A. Contreras, C. Herrenknecht, P. Truquet, V. Séchet, U. Tillmann, and P. Hess. 2012. Effect of *Azadinium spinosum* on the feeding behaviour and azaspiracid accumulation of *Mytilus edulis. Aquatic Toxicology*, **124–125**: 179–187.
- Jauffrais, T., V. Séchet, C. Herrenknecht, P. Truquet, S. Veronique, U. Tillmann, and P. Hess. 2013. Effect of environmental and nutritional factors on growth and azaspiracid production of the dinoflagellate *Azadinium spinosum. Harmful Algae*, 27: 138–148.
- Kilcoyne, J., T. Jauffrais, M. Twiner, G. Doucette, J.A. Aasen Bunæs, S. Sosa, B. Krock, V. Séchet, C. Nulty, R. Salas, D. Clark, J. Geraghty, C.

Duffy, B. Foley, M.A. Quilliam, P. McCarron, C.O. Miles, J. Silke, A. Cembella, U. Tillmann, and P. Hess. 2014a. Azaspiracids: toxicological evaluation, test methods and identification of the source organisms (ASTOX 2). Galway, Ireland.

Kilcoyne, J., C. Nulty, T. Jauffrais, P. McCarron, F. Herve, B. Foley, F. Rise, S. Crain, A.L. Wilkins, M.J. Twiner, P. Hess, and C.O. Miles. 2014b. Isolation, structure elucidation, relative LC-MS response, and in vitro toxicity of azaspiracids from the dinoflagellate *Azadinium spinosum. Journal of Natural Products*, 77: 2465–2474.

Kim, J.W., U. Tillmann, N.G. Adams, B. Krock, W.L. Stutts, J.R. Deeds, M.S. Han, and V.L. Trainer. 2017. Identification of *Azadinium* species and a new azaspiracid from *Azadinium poporum* in Puget Sound, Washington State, USA. *Harmful Algae*, 68: 152–167.

Krock, B., U. Tillmann, D. Voß, B.P. Koch, R. Salas, M. Witt, E. Potvin, and H.J. Jeong. 2012. New azaspiracids in Amphidomataceae (Dinophyceae): proposed structures. *Toxicon*, **60**: 830–839.

Krock, B., U. Tillmann, E. Potvin, H.J. Jeong, W. Drebing, J. Kilcoyne, A. Al-Jorani, M.J. Twiner, Q. Göthel, and M. Köck. 2015. Structure elucidation and in vitro toxicity of new azaspiracids isolated from the marine dinoflagellate *Azadinium poporum. Marine Drugs*, **13**: 6687–6702.

Ofuji, K., M. Satake, T. McMahon, J. Silke, K.J. James, H. Naoki, Y. Oshima, and T. Yasumoto. 1999. Two analogs of Azaspiracid isolated from mussels, *Mytilus edulis*, involved in human intoxication in Ireland. *Natural Toxins*, 7: 99–102.

Percopo, I., R. Siano, R. Rossi, V., Soprano, D. Sarno, and A. Zingone. 2013. A new potentially toxic *Azadinium* species (Dinophyceae) from the Mediterranean Sea, *A. dexteroporum* sp. nov. *Journal of Phycology*, **49**: 950–966.

Potvin, E., H.J. Jeong, N.S.T. Kang, U. Tillmann, and B. Krock. 2012. First report of the photosynthetic dinoflagellate genus *Azadinium* in the Pacific Ocean: morphology and molecular characterization of *Azadinium cf. poporum*. *Journal of Eukaryotic Microbiology*, **59**: 145–156.

Rossi, R., C. Dell'Aversano, B. Krock, P. Ciminiello, I. Percopo, U. Tillmann, V. Soprano, and A. Zingone. 2017. Mediterranean *Azadinium dexteroporum* (Dinophyceae) produces AZA-35 and six novel azaspiracids: a structural study by a multi-platform mass spectrometry approach. *Analytical and Bioanalytical Chemistry*, 409: 1121–1134.

Samdal, I., K.E. Lovberg, L.R. Briggs, J. Kilkoyne, J. Xu, C.J. Forsyth, and C.O. Miles. 2015. Development of an ELISA for the detection of Azaspiracids. *Journal of Agricultural and Food Chemistry*, 63: 7855–7861.

Satake, M., K. Ofuji, K. James, A. Furey, and T. Yasumoto. 1998. New toxic events caused by Irish mussels. In: *Harmful Algae*. B. Reguera, J. Blanco, M.L. Fernandez, and T. Wyatt (Eds.). Xunta de Galicia and International Oceanographic Commission of UNESCO, Santiago de Compostela: p. 468–469.

Smith, K.F., L. Rhodes, D.T. Harwood, J. Adamson, C. Moisan, R. Munday, and U. Tillmann. 2016. Detection of *Azadinium poporum* in New Zealand: the use of molecular tools to assist with species isolations. *Journal* of *Applied Phycology*, 28: 1125–1132.

Tillmann, U., M. Borel, F. Barrera, R. Lara, B. Krock, G. Almandoz, and N. Trefault. 2016. *Azadinium poporum* (Dinophyceae) from the South Atlantic off the Argentinean coast produce AZA-2. *Harmful Algae*, **51**: 40–55.

Tillmann, U., M. Elbrächter, U. John, and B. Krock. 2011. A new non-toxic species in the dinoflagellate genus Azadinium: A. poporum sp. nov. European Journal of Phycology, 46: 74–87.

Tillmann, U., M. Elbrächter, B. Krock, U. John, and A. Cembella. 2009. *Azadinium spinosum* gen. et sp. nov. (Dinophyceae) identified as a primary producer of azaspiracid toxins. *European Journal of Phycology*, 44: 63–79.

Tillmann, U., M. Gottschling, E. Nézan, and B. Krock. 2015. First record of *Azadinium dexteroporum* and *Amphidoma languida* (Amphidomataceae, Dinophyceae) from the Irminger Sea off Iceland. *Marine Biodiversity Records*, **8**: 1–11.

Tillmann, U., M. Gottschling, E. Nézan, B. Krock, and G. Bilien. 2014. Morphological and molecular characterization of three new *Azadinium* species (Amphidomataceae, Dinophyceae) from the Irminger Sea. *Protist*, **165**: 417–444.

Tillmann, U., D. Jaen, L. Fernandez, M. Gottschling, M. Witt, J. Blanco, and B. Krock. 2017. *Amphidoma languida* (Amphidomataceae, Dinophyceae) with a novel azaspiracid toxin profile identified as the cause of molluscan contamination at the Atlantic coast of southern Spain. *Harmful Algae*, 62: 113–126.

Tillmann, U., R. Salas, M. Gottschling, B. Krock, D. O'Driscoll, and M. Elbrächter. 2012a. *Amphidoma languida* sp. nov. (Dinophyceae) reveals a close relationship between *Amphidoma* and *Azadinium*. *Protist*, **163**: 701–719.

Tillmann, U., S. Söhner, E. Nézan, and B. Krock. 2012b. First record of *Azadinium* from the Shetland Islands including the description of *A. polongum* sp. nov. *Harmful Algae*, 20: 142–155.

Toebe, K., A.R. Joshi, P. Messtorff, U. Tillmann, A. Cembella, and U. John. 2013. Molecular discrimination of taxa within the dinoflagellate genus *Azadinium*, the source of azaspiracid toxins. *Journal of Plankton Research*, **35**: 225–230.

Twiner, M., P. Hess, and G.J. Doucette. 2014. Azaspiracids: toxicology, pharmacology, and risk assessment. In: *Seafood and Freshwater Toxins*. L.M. Botana (Ed.). CRC Press, Boca Raton: p. 823–855.