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# First record of the dynamics of domoic acid producing *Pseudo-nitzschia* spp. in Indonesian waters as a function of environmental variability

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#### ABSTRACT

Within the past few decades, harmful algal blooms (HABs) have occurred frequently in Indonesian waters, resulting in environmental degradation, economic loss and human health problems. So far, HAB related studies mainly addressed ecological traits and species distribution, yet toxin measurements were virtually absent for Indonesian waters. The aim of the present study was to explore variability of the potentially toxic marine diatom genus Pseudo-nitzschia, as well as its neurotoxin domoic acid as a function of environmental conditions in Ambon Bay, eastern Indonesia. Weekly phytoplankton samples, oceanographic (CTD, nutrients) and meteorological (precipitation, wind) parameters were analyzed at 5 stations in the bay during the dry and wet seasons of 2018. Liquid chromatography - tandem mass spectrometry (LC-MS/MS) was used to detect particulate DA (pDA). Vegetative cells of Pseudo-nitzschia spp. and pDA were found in 98.6% and 51.4% of the samples, respectively. pDA levels were low, yet detected throughout the campaign, implying that Ambon Bay might potentially be subject to amnesic shellfish poisoning. The highest levels of both Pseudo-nitzschia spp. cell abundance and pDA were found in the wet season, showing a strong positive correlation between both parameters, compared to the dry season, (r = 0.87 and r = 0.66 (p < 0.01), respectively). Statistical analyses revealed that temperature and mixed layer depth positively correlated with Pseudo-nitzschia spp. and pDA during the dry season, while ammonium showed positive correlations in both seasons. This study represents the first successful investigation of the presence and variability of Pseudo-nitzschia spp. and its neurotoxin DA in Indonesian waters.

#### 1. Introduction

*Pseudo-nitzschia*, a pennate and chain forming microalga is a cosmopolitan diatom genus that can cause harmful algal blooms (HABs) around the world (Trainer et al., 2000; Pan et al., 2001; Bates and Trainer, 2006). Some species of this genus produce a potent neurotoxin, known as domoic acid (DA). This toxin can be transferred via the food web, eventually causing amnesic shellfish poisoning (ASP) in humans due to the consumption of contaminated shellfish (Bates et al., 1998; Scholin et al., 2000; Bejarano et al., 2008; Mafra Jr. et al., 2010; Zabaglo et al., 2016). Human illness and fatalities due to DA outbreaks were first recorded in Canada in 1987, revealing symptoms such as gastrointestinal distress, confusion, disorientation, memory loss, coma and death (Bates et al., 1989; Lefebvre and Robertson, 2010). DA accumulated in filter feeders can also be fatal to higher marine trophic levels, such as mammals, leading to mortalities of sea lions and whales (Scholin et al., 2000; reviewed by Bejarano et al., 2008; Fire et al., 2010). The total number of described *Pseudo-nitzschia* species in the 2010s was 37 of which 12 were thought to be capable of producing DA (Trainer et al., 2010; reviewed by Trainer et al., 2012). This number recently increased to 52 species, of which 26 are capable of producing DA (Bates et al., 2018).

Environmental properties that promote the proliferation of *Pseudo*nitzschia are very complex and vary between regions and seasons

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(Trainer et al., 2012; Umhau et al., 2018). This complexity and ambiguous causative factors may be related to natural climate variability (Sekula-Wood et al., 2011). Yet, in some cases, *Pseudo-nitzschia* was found to be correlated with nutrient availability. For example, cold upwelled and nutrient-rich waters were associated with *Pseudo-nitzschia* blooms in Californian coastal waters (Anderson et al., 2006; Schnetzer et al., 2007; Sekula-Wood et al., 2011). Eutrophic waters carrying high levels of nitrate derived from local groundwater favoured *Pseudo-nitzschia* to form blooms in the northern Gulf of Mexico (Macintyre et al., 2011).

Regardless of the complexity of the blooms and environmental properties, some factors are likely to be associated with cell specific toxin production. Phosphate and silicate limitation enhanced DA production in *Pseudo-nitzschia* species such as *P. australis, P.pungens* and *P. fraudulenta* (Lelong et al., 2012; Lema et al., 2017). Other factors that might regulate cell specific DA production were found to be pH, CO<sub>2</sub>, irradiance, salinity, bacteria and temperature (Macintyre et al., 2011; Lelong et al., 2012; Zhu et al., 2017). Rue and Bruland (2001) found that *Pseudo-nitzschia* produces DA to bind trace metals in order to increase micronutrient availability (iron) or to reduce intracellular harmful copper. DA was found to accumulate in copepods (Leandro et al., 2010; Tammilehto et al., 2012), indicating grazing on *Pseudo-nitzschia* seriata toxicity increased in the presence of grazers such as copepods, implying that zooplankton may trigger cell specific DA production.

In Indonesian waters, HAB outbreaks and their impacts on marine environments and human wellbeing have been observed since the early 1970s. However, due to a lack of experts, knowledge and awareness of HABs, studies in the country so far focused mainly on general species abundance (Sidharta, 2005). Indonesia has been suffering from paralytic shellfish poisoning (PSP) events caused by *Pyrodinium bahamense* var. *compressum*, which were associated with 427 cases of human illness and 17 deaths (Aditya et al., 2013; Azanza and Taylor, 2001). Other phytoplankton blooms were reported in the country coinciding with massive fish kill events due to dissolved oxygen depletion (Thoha et al., 2007; Wouthuyzen et al., 2007). To our knowledge, no particular studies addressed *Pseudo-nitzschia* spp. and its neurotoxin DA in Indonesian waters, including the eastern part of the country.

Ambon Bay is a semi-enclosed eutrophic bay in eastern Indonesia (Fig. 1), where proliferations of HAB species have frequently been observed. Human illness and fatalities were reported in the surrounding areas due to frequent occurrences of high densities of the toxic dino-flagellate *Pyrodinium bahamense* var. *compressum* (Likumahua, 2013). Phytoplankton monitoring is conducted in the bay since 2008 revealing other potentially toxic genera, such as *Alexandrium, Gymnodinium, Dinophysis*, and *Pseudo-nitzschia*, which was formerly determined as *Nitzschia* spp (data not published). However, the monitoring program was only focused on the abundance and distribution of phytoplankton with particular attention to dinoflagellates, nor did it include associated toxin analyses.

Given the complete lack of information on *Pseudo-nitzschia* occurrences and potential DA impacts in eastern Indonesian waters, the present study first of all addressed the possible presence and variability of DA producing *Pseudo-nitzschia* in Ambon Bay. Secondly, environmental factors that might regulate the variability of *Pseudo-nitzschia* and DA concentrations were investigated. To this end, a seven-months field campaign was executed in the bay, encompassing both the dry and the wet season, during which weather (wind, precipitation) and physicochemical water column properties (mixed layer depth, stratification, temperature, salinity, irradiance attenuation, nutrients) as well as phytoplankton composition and toxins were analysed.

#### 2. Materials and methods

#### 2.1. Study area and stations

Ambon Island (3°38'17"S and 128°07'02"E) (Fig. 1) is located in Maluku Province, eastern Indonesia, which is resided by roughly 450,000 people. The island has a semi-enclosed estuary, known as Ambon Bay, which opens to the Banda Sea. A narrow and shallow sill (Fig. 2) divides the bay into two parts (Inner and Outer bay), which restricts water circulation and limits the flushing process to the open ocean (Banda Sea). The area has a tropical climate with high rainfall during the wet season, which is associated with the southeast monsoon occurring from March to September. The dry season is associated with the northwest monsoon between October and March, and it is characterized by relatively low precipitation and increased air temperature.

Four sampling stations (B, C, D, and E) were chosen in the inner part of the bay whereas a fifth station (A), with higher oceanic water influence, was set up in the outer bay (Fig. 1). Stations C and D are permanent stations, which have been monitored by the Center for Deep Sea Research Institute (LIPI) since 2008. Those stations are characterized by nearby dense mangrove vegetation and some small river outflows. Two blooms of toxic dinoflagellates had been recorded before at those stations during the wet season (Likumahua, 2013). Stations E and B were chosen close to moderately sized river outflows, which flush massive agricultural waste water into the inner bay during the wet season. These stations had experienced an extensive dinoflagellate bloom during the dry season in March 2013 (data unpublished). Some fish farms are located in the inner bay, close to stations C, D and E. At the LIPI research institute (Fig. 1), a weather station was installed to continuously measure precipitation, air temperature, wind and humidity. Daily average precipitation and wind speed were calculated and calibrated with the data obtained from the local Meteorological and Climate Bureau.

#### 2.2. Field sampling

The five stations were sampled weekly throughout the dry and wet season, between January and July 2018. During the alleged phytoplankton bloom periods, both in the peak of dry (February-March) and wet (June-July) season, samples were collected twice a week. The period between end of March and mid- May was considered as the transition phase from the dry to wet season. Irradiance attenuation was measured using a LiCor Li1400 light meter, at 1, 2, 4, 6, 8, 10, 12, 16 and 20 (m depth). At each station, a Compact Alec CTD Model ASTD 687 was deployed to measure temperature, salinity, turbidity, density, and chlorophyll-a fluorescence.

A 3.5L Nansen bottle was used to collect water samples for nutrient analysis at 2 and 20 m depth, which represented the water above and below the thermocline. Since the water in the Inner bay is often turbid, the water samples were pre-filtered on board through a  $20 \,\mu\text{m}$  filter mesh to eliminate large plankton, other organic materials and sediment particles. The water samples were then transferred to 600 mL bottles and placed in a dark insulation box, containing seawater at ambient temperature. The bottles were pre-cleaned several times using distilled water and rinsed with double distilled water (ddH<sub>2</sub>O) prior to sampling.

At each station, a Hydro-bios plankton net with 40 cm diameter, 100 cm length, and 20  $\mu$ m mesh size was deployed twice vertically to collect plankton samples from 0–20 m depth, with total pulling time ranging between 60 and 70 s, following the manufacturer's guidelines. The difference of approximately 10 s pulling speed (between 0.33 m/s and 0.29 m/s) potentially caused variability of sampled volumes, which was calculated to be less than 12 %. This net tow method has been used to reveal a semi-quantitative comparison of cell abundances and particulate toxin concentrations as described by Fabro et al. (2016) and Almandoz et al. (2017). Plankton samples were transferred to 500 mL bottles and kept in the dark at ambient temperature in the insulation



Fig. 1. Map of Ambon Bay with the location of the five sampling stations (A-E). The red star is LIPI, the Centre of Deep Sea Research in Indonesia. The red triangle is the sill. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).



Fig. 2. Ambon Bay cross-section.

box. The final volume of concentrated plankton samples obtained from two net tows was 400 mL, which was processed in the lab for toxin analysis, microscopic observations and spectrophotometric chlorophylla analysis.

#### 2.3. Sample processing

#### 2.3.1. Nutrients

The pre-filtered water samples were filtered immediately after sampling through a 0.2 µm pore size, 47 mm Ø Fisherbrand nylon membrane filter using a vacuum pump with a maximum pressure of 0.3 bar and stored at -20 °C until analysis. Samples were analyzed fortnightly using a Spectrophotometer (UV-vis Shimadzu 1700) to measure Nitrate (NO<sub>3</sub>), Phosphate (PO<sub>4</sub>) and Silicate (SiO<sub>2</sub>) following Strickland and Parsons (1972), and Ammonium (NH<sub>4</sub>) following APHA (1998).

#### 2.3.2. Chlorophyll-a ( $\geq 20 \mu m$ )

A five-channel filtration set up was used to process the samples from the five stations immediately after arrival in the lab. Sixty mL of net sample was filtered through 25 mm Ø GF/F Whatman filters using a 0.2 bar vacuum pump. Filters were transferred to darkened glass tubes filled with 10 mL of 90% acetone. After mixing, the samples were kept at 4 °C in a refrigerator. After a few hours, the filters were ground to optimize extraction efficiency using a mini glass spatula and subsequently refrigerated overnight. Samples were centrifuged for 8 min at



**Fig. 3.** Weather conditions recorded as average values for 3 h prior to sampling. The grey area refers to the transition period between dry and wet season. The vertical red line separates the two distinct seasons, as determined by statistical PCA analysis of environmental conditions in 2018. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

8,400xg, and measured in a spectrophotometer (UV–vis Shimadzu 1700) at wavelengths of 750, 665, 645 and 630 nm. Chlorophyll-a (Chl-a) was calculated using the equations as described by Strickland and Parsons (1972).

#### 2.3.3. Pseudo-nitzschia spp. abundance

Forty mL of net sample were fixed with acidic lugol iodine solution, and a drop of formalin (4% final concentration) was added to inactivate bacteria. The fixed samples were stored in a 4 °C refrigerator prior to analysis. Samples were settled in an Uthermöhl 10 mL sedimentation chamber and inspected at 200x magnification using an Olympus LH50A inverted microscope. To achieve quantitative cell abundances, a minimum of 400 *Pseudo-nitzschia* spp. cells was counted. The whole chamber was counted when cell densities were low and at higher cell densities a number of randomly chosen fields was inspected until 400 cells were counted. The total abundance of *Pseudo-nitzschia* spp. was calculated in cells per net tow (NT<sup>-1</sup>). Qualitative microscopic information for the dominant *Pseudo-nitzschia* species was obtained by an Olympus IMT-2 inverted microscope at 1000x SPlan APO. The cell size was calibrated and treated using Adobe Light Room and Microscope Image Capture and Measurement (MICAM) software, version 2.4.

#### 2.3.4. Toxin analysis

Pellets were obtained from 300 mL of the net samples, concentrated by centrifugation steps at 12,100 x g for 20 min and transferred to 2 mL Eppendorf tubes. Pellets were stored at -30 °C until further analysis. Particulate domoic acid (pDA) was analysed by liquid chromatography – tandem mass spectrometry (LC–MS/MS). Briefly, cell pellets were transferred to tubes containing ca. 0.9 g lysing matrix D, and 500 µL methanol were added. Extraction was performed at maximum speed (6.5 m/s) for 45 s using a Bio101 FastPrep instrument (Thermo Savant, Illkirch, France). Homogenized samples were centrifuged for 15 min at 16,100xg at 4 °C. Supernatants were transferred to spin filters with a pore size of 0.45 µm (Milipore Ultrafree, Eschborn, Germany) and centrifuged for 30 s at 5700 x g. The filtrate was subsequently transferred to HPLC vials for the pDA analysis, according to Krock et al. (2008). The detection limit of pDA in this study was 0.60 ng NT<sup>-1</sup>.

#### 2.4. Statistical analysis

Three hour averages were calculated for wind speed and the rainfall just prior to the sampling trip and used as environmental variables for the statistical analysis. The light extinction coefficient (kd) was calculated by linear regression of log transformed data. Mixed layer depth (MLD) and stratification index were determined using the threshold method as described by González-Pola et al. (2007) and Somavilla et al. (2017). Tidal elevation data were obtained from the Hydrography and Oceanography Centre, Indonesian Navy. PCA multivariate analysis score plot was employed to distinguish variation among stations and between seasons. The analysis involved all parameters including weather (precipitation and wind speed), physicochemical parameters (the average temperature and salinity (0–20 m), light attenuation coefficient (kd), MLD, stratification, tidal elevation and nutrients at both depths (2 and 20 m)) and biological parameters (Chl-a from both net and CTD, pDA and *Pseudo-nitzschia* spp.). The correlation analysis between *Pseudo-nitzschia* spp. cell abundance, pDA and environmental factors was performed using a non-parametric Spearman's rank, as the data were not normally distributed. Statistical analysis was done using R and the Minitab18 package.

#### 3. Results

#### 3.1. Weather variability

Low average rainfall was found during the dry season, with only six rainy days, ranging between 0.01 and 0.7 mm of total daily precipitation. Precipitation occurred more frequently during the wet season, starting with two high rainfall days (1.8 and 2.0 mm of total daily precipitation), respectively (Fig. 3). High wind speeds were recorded at the beginning of the year, after which a slightly decreasing trend was observed towards the transition period (Fig. 3). An upward trend in wind speed occurred during the wet season, ranging between 1.0 m.s<sup>-1</sup> on day 158 and 4.6 m.s<sup>-1</sup> on day 190.

#### 3.2. Physicochemical characteristics

Average (0-20 m) water temperatures showed a downward trend during the sampling period, ranging between 30.2 °C (day 78 at Station D) and 26.0 °C (day 141 at Station A). During the dry season, the average temperature increased slightly from 29.2 °C to 29.9 °C and started to decrease during the transition period (Fig. 4A). A steady temperature decrease was found during the wet season reaching lowest values at the end of the sampling period. Throughout the campaign, lowest temperatures were found at Station A (Fig. 4A). During the dry and transition seasons, Station D was warmer than the other stations, but was cooler during the subsequent wet season. Similar conditions were observed for the surface water temperature, being slightly warmer than the average at 20 m. Surface temperatures ranged between 31.1 °C (day 106 at Station D) and 26.4 °C (day 200 at Station A).

Average salinity 0–20 m overall showed a slightly upward trend, ranging between 32.5 (day 141 at Station E) and 33.8 (day 197 at Station A). Meanwhile, surface salinity was much lower than the average over 0-20 m, ranging between 27.4 (day 16 at Station D) and 33.6 (day 200 at Station A). Salinity was relatively stable during the dry season, even though there was a slightly downward trend before the transition period (Fig. 4B). During the dry season, the average salinity



**Fig. 4.** Temporal variability of physical properties for the five sampling stations. A: Average water temperature 0-20 m; B: Average salinity 0-20 m; C: Irradiance attenuation kd 0-20 m; D: Mixed Layer Depth (MLD) (m); E: Stratification Index. The grey area refers to the transition period between dry and wet season. The vertical red line separates the two distinct seasons, as determined by statistical PCA analysis of environmental conditions in 2018. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

0-20 m ranged between 33.0 and 33.3, and in the transition period between 33.0 and 33.1. A gradual upward trend occurred during the wet season (Fig. 4B), with the average 0-20 m ranging between 33.0 and 33.5. Highest salinities were recorded at Station A throughout the campaign. Lowest salinities were found for station D during the dry season, whereas station B and E showed lower salinities during the wet season (Fig. 4B). These lower salinities were also observed at the surface.

During the dry season and the transition period irradiance attenuation was stable, ranging between 0.03 (day 71, Station A) and 0.29 (day 134, Station D). An upward trend in kd was observed for all stations during the wet season (Fig. 4C), reaching highest levels towards the end of the campaign (day 207; Station C = 0.37 and Station D = 0.38). Lowest kd values were found for Station A throughout the season, while Station D had the highest kd for both dry and wet seasons.

Mixed layer depth (MLD) was highly variable in space and time. However, a slightly downward trend was found from the dry to the wet season (Fig. 4D). MLD was roughly above 10 m depth during the wet season for the inner bay stations, interrupted by deep mixing events. Deepest MLDs were found at Station A in the outer bay. Stratification index showed an overall increasing trend from the dry to the wet season (Fig. 4E). During the dry season and the transition period, the stratification index was found below 1 kg.m<sup>3</sup>, increasing to 2.9 kg.m<sup>3</sup> at day 190 (Station E) in the wet season. The water column in the inner bay was much more stratified than in the outer bay (Station A:  $\leq$  0.5 kg.m<sup>3</sup>), which showed the lowest stratification index throughout the sampling period. Highest indices were found at stations B and E during the wet season.

Phosphate concentration ranged between not detected and 3  $\mu$ M throughout the sampling period. Surface (2 m) phosphate showed a high variability over time and space, with pronounced peaks in both seasons, yet a gradual decreasing trend was observed for all stations (Fig. 5A). Phosphate concentrations were higher at the inner bay stations compared with station A in the outer bay (Fig. 5A) except for days 16 and 162. Phosphate concentrations at 20 m, showed similar variability and overall trends as surface levels (Fig. 5B). However, concentrations were generally higher at 20 m. Stations C, D and E showed the highest phosphate levels at 20 m, and station A the lowest.

Generally, surface silicate concentrations were higher (mostly  $\geq 10~\mu$ M) than those at 20 m (mostly  $\leq 10~\mu$ M) (Fig. 5 C and D). Both depths showed an upward trend over time. At the end of the wet season, silicate concentrations increased considerably at both depths. Station D showed the highest surface concentrations during the dry season while stations B and E had highest levels during the wet season. Highest silicate peaks occurred at Stations C and D (52.1  $\mu$ M and 54.2  $\mu$ M, respectively) at the end of the season (Fig. 5C). In contrast, station D showed highest silicate levels at 20 m throughout the sampling period (Fig. 5D).

Although highly variable, nitrate concentrations showed an increasing overall trend during the sampling period, ranging between 0.01 and 3.0  $\mu$ M (Fig. 5 E and F). A clear upward trend occurred at the surface, even though there was a drop by the end of the dry season between days 65 and 90. Nitrate concentrations started to increase in the transition period, reaching a maximum level of 2.5 µM (Station D) in the wet season (Fig. 5E). An opposite trend was found for the concentrations at 20 m during the dry season, which exhibited a strong decrease during the course of the dry season (Fig. 5F). The highest nitrate level at 20 m was 3.0 µM, as found during the beginning of the campaign. Concentrations reached lowest levels by the end of the dry season (day 65-90), similar to the observed surface trends. Station D showed the highest nitrate concentrations both at the surface and the 20 m samples. Surface nitrate concentration at Station A was similar to the other stations. In contrast, at 20 m, concentrations were lower than at the other stations.

Surface and 20 m ammonium concentrations increased during the dry season, followed by a decrease during the transition period and

subsequent wet season (Fig. 5 G and H). Yet, during the wet season, peak ammonia levels were recorded at both depths, (14.8  $\mu$ M, day 200 at Station C). During the wet season, stations A and C showed the highest ammonium concentrations, respectively at the surface (14.3  $\mu$ M and 13.0  $\mu$ M) and at 20 m (15.1  $\mu$ M and 14.7  $\mu$ M).

#### 3.3. Chl-a, pDA and Pseudo-nitzschia spp. abundance

Chlorophyll-a (Chl-a) data was obtained from the net hauls (reflecting the  $\geq 20 \,\mu\text{m}$  phytoplankton fraction) and the CTD measurements, showing a strong positive correlation (r = 0.68 and r = 0.78 (p = 0.00), dry and wet season respectively). On average, net Chl-a level was  $\pm 12$  times lower than the CTD Chl-a measurement (results not shown). During the campaign, Chl-a  $\geq 20 \,\mu\text{m}$  showed an upward trend reaching peak levels at the end of the wet season. Chl-a  $\geq 20 \,\mu\text{m}$  levels ranged between 0.01 (day 46, Station A) and 1.8 mg m<sup>3</sup> (day 207, Station C), (Fig. 6A). Levels were similar for all stations during the dry season. During the wet season the inner bay stations, in particular B, C and D showed the highest Chl-a  $\geq 20 \,\mu\text{m}$  levels while values remained low at Station A in the outer bay.

Particulate domoic acid (pDA) was detected in all samples, yet, only 51.4% was found above the detection limit ( $0.60 \text{ ng NT}^{-1}$ ). Values of pDA were highly variable over time, showing peaks both in the dry and wet season, mainly at the inner bay stations. Yet, pDA levels showed an upward trend from the dry to the wet season, during which peaks were higher and more frequent, with peak levels up to 3,716 ng NT<sup>-1</sup> (day 162, Station D) (Fig. 6B). In the dry season, two pDA peaks were found, on day 53 (1,218 ng NT<sup>-1</sup>) and day 85 (321 ng NT<sup>-1</sup>) both at station E; in the transition period one small peak was found on day 127 at station D (297 ng NT<sup>-1</sup>). During the wet season peaks were found at station D on day 148 (1,819 ng NT<sup>-1</sup>) and day 162 (3,716 ng NT<sup>-1</sup>), and at station C on day 200 (465 ng NT<sup>-1</sup>) and at station E on day 204 (463 ng NT<sup>-1</sup>).

During the campaign, Pseudo-nitzschia spp. cells were observed in 207 out of 210 samples (98.6%), and the abundance variability showed a similar pattern as compared with pDA levels. The cell abundance observed during the campaign remained relatively low and no Pseudonitzschia blooms occurred. Maximal cell densities were observed during both seasons and showed an overall increasing trend (Fig. 6C). Cell abundances ranged between no cells (day 46, Station C) and  $4,456\times10^4$  cells  $NT^{\text{-}1}$  (day 162, Station D). In the dry season, the Pseudo-nitzschia spp. peaked between day 50 and 60, ranging from  $40 \times 10^4$  to 2,651 × 10<sup>4</sup> cells NT<sup>-1</sup>. A small peak in *Pseudo-nitzschia* spp. cell abundance was found at Station D on day 148 (374  $\times$  10<sup>4</sup> cells NT<sup>-1</sup>), which was followed by a peak density in the wet season between day 158 and 169 (4,456  $\times$  10<sup>4</sup> cells NT<sup>-1</sup>). During the dry season, peaks in cell abundance were found at Station E and at Station D during the wet season. Qualitative microscopic inspection revealed the presence of two potentially toxic Pseudo-nitzschia species, of which P. pungens was by far the most abundant. The other species was determined as P. cf delicatissima, which occurred in relatively low numbers compared with P. pungens (data not shown). Species identification was based on average cell size (valve length and width). The cell length of P. pungens in this study ranged between 101 and 123 µm, its width between 3.0 and 4.5 µm, whereas the number of valve striae per 10 µm ranged between 10 and 14 (Fig. 7).

Besides *Pseudo-nitzschia* spp., the net haul phytoplankton communities were dominated by other diatoms including *Chaetoceros* spp., *Thalassionema nitzschioides* and *Rhizosolenia* spp. In addition, dinoflagellate species were observed throughout, including *Gymnodinium catenatum* and *Dinophysis miles*, whereas zooplankton (Copepoda and Meroplankton groups), and fecal pellets were found in high abundance in the net samples.



**Fig. 5.** Temporal variability of major nutrients for the five sampling stations at two water depths. A, C, E, G = 2 m: B, D, F, H = 20 m. The grey area refers to the transition period between dry and wet season. The vertical red line separates the two distinct seasons, as determined by statistical PCA analysis of environmental conditions in 2018. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

## 3.4. Relationship between Pseudo-nitzschia spp., pDA and environmental factors

Station A was set up as a reference station, located in the outer bay, which was characterized by a higher water depth and oceanic influence. The space resolved PCA score plot (n = 210), when based on all data (stations, weather, physicochemical and biological parameters), indeed showed a clear separation of station A samples from samples of the other stations (Fig. 8A), with only one inner bay data point (station B, day 67) grouping with station A. Apart from station A, no clear space resolved clustering was observed, e.g. for the stations in the inner bay (n = 168). The time resolved PCA score plot (Fig. 8B) showed highly similar characteristics for the dry and the transition period but a clear separation from the wet season (n = 210). Four data points of the transition period, belonging to the day where the wet season had just started, revealed the same traits with the wet season. As a result, those points were grouped in the wet season.

Spearman rho regression and analyses was done to investigate possible environmental impacts on *Pseudo-nitzschia* spp. abundances and pDA levels. Given its strong deviation from the other stations, the outer bay station A was not included further. Moreover, based on the strong seasonal clustering, analyses were done for the dry and wet season separately, whereas data points from the transition period were included in the dry season following the first PCA analysis (Fig. 8B). In this way, the clearest visible regulation of environmental factors on biological properties could be obtained. Spearman rho analysis in the dry season revealed strong positive correlations among *Pseudo-nitzschia* spp., pDA, MLD and Temperature (Table 1). Phosphate, nitrate and silicate at 20 m as well as stratification on the other hand negatively correlated with *Pseudo-nitzschia* spp. and pDA (Table 1). In the wet season, *Pseudo-nitzschia* spp. and pDA positively correlated with ammonium and negatively correlated with MLD (Table 1).

Spearman rho analysis revealed a strong positive correlation between *Pseudo-nitzschia* spp. and pDA during both seasons. The correlation in the wet season was stronger than in the dry season, r = 0.87(p = 0.00) and r = 0.66 (p = 0.00), respectively (Table 1). *Pseudonitzschia* spp. and pDA showed a strong correlation when based on linear regression in the dry and wet season ( $R^2 = 0.72$  and 0.63, respectively, Fig. 9).

#### 4. Discussion

This study provides the first evidence of domoic acid (DA) associated with *Pseudo-nitzschia* spp. in Indonesian waters. Although overall low, pDA and *Pseudo-nitzschia* spp. were found to be persistent throughout the dry and the wet season in the semi-enclosed Ambon



**Fig. 6.** Temporal variability of biological properties for five stations. A: Chl-a from net (cells  $\geq 20 \,\mu$ m); B: particulate domoic acid (pDA) levels; C: *Pseudo-nitzschia* spp. cell abundance. The grey area refers to the transition period between dry and wet season. The vertical red line separates the two distinct seasons, as determined by statistical PCA analysis of environmental conditions in 2018. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

Bay. This implies that the area might potentially be subject to amnesic shellfish poisoning (ASP).

In our study, *Pseudo-nitzschia* spp. abundances and pDA concentrations were derived from net samples with 20  $\mu$ m mesh size. Although most *Pseudo-nitzschia* cells occurred in chains, given the average cell width of 3.3  $\mu$ m, we cannot exclude that an unknown fraction of cells passed the net. Hence, total *Pseudo-nitzschia* cells as well as pDA levels might be underestimated in our study. Furthermore, the towing speed of the net ranged between 0.33 m/s and 0.29 m/s, which potentially might have created a 12% variability in total sampled volume per net tow. Clearly, the choice for using nets is not ideal, yet, when investigating particulate toxin dynamics under non-bloom conditions (like aimed for here), water sample processing (by a.o. Niskins) would not reveal quantitative results sufficiently above the toxins detection limits.

Both cell determination and quantification of *Pseudo-nitzschia* were performed at the genus level, leading to the total *Pseudo-nitzschia* spp. abundance. In other words, for practical reasons the most abundant *P. pungens* as well as *P.* cf *delicatissima* cells were pooled. This implies that if environmental conditions would act in a different way on these two species, this could not be revealed in our study.

The tropical climate of Ambon Island is characterized by seasonal monsoons. During the dry season, high water temperatures are associated with the northwest monsoon, while lower temperatures and high precipitation during the wet season associates with the southeast monsoon. These typical monsoonal characteristics massively influence water physicochemical properties, including Ambon Bay, related with upwelling (southeast monsoon) and sinking (northwest monsoon)



**Fig. 7.** Light microscope photographs of *Pseudo-nitzschia pungens* cells, found in Ambon Bay in 2018. (A) A single cell (scale bar  $= 10 \,\mu$ m). (B) Valve view (scale bar  $= 10 \,\mu$ m). (Cw) Cell width ( $\mu$ m). (Cs) 13–14 striae per 10  $\mu$ m.

processes in the Banda Sea (Wyrtki, 1961; Boëly et al., 1990; Zijlstra et al., 1990). In our study, the decrease in water temperature and increase in salinity (Fig. 4 A and B) as observed during the wet season indicated the onset of upwelling in the Banda Sea, affecting all stations in Ambon Bay, but most of all Station A. Similar events had been shown earlier by Saputra and Lekalette (2016), where cold-saline water at 20 m or lower showed enhanced intrusion of the inner bay during the southeast monsoon as the result of tidal force upwelling in the Banda Sea. In addition, more than 70% of upwelled water from the outer bay had been found to replace local water masses in the inner bay (Riantika, 2011 (data not published); Corvianawatie et al., 2014).

Other monsoon related factors controlling physical water properties include wind speed and precipitation, which might have opposing effects on mixed layer depth and stratification. Yet, as found in the present study, enhanced wind speeds in the wet season did not lead to deeper mixed layers in the inner bay (Figs. 3 and 4D). Instead, the increase in precipitation on average caused elevated irradiance attenuation due to enhanced runoff as well as water column stratification, although occasional strong winds caused regular stratification breakdown (Figs. 3, 4C and E).

During the southeast monsoon (wet season) nutrient variability showed upward trends at both water depths, likely caused by elevated precipitation and associated runoff. However, our data did not show strong positive correlations between precipitation and nutrient concentrations during the wet season. This might be due to rapid flushing processes in the bay, promoted by the deep water intrusion. The residence time in the inner bay has been calculated to be less than 14 days (Pello et al., 2014). Meanwhile, high nutrient concentrations at the beginning of the year might be due to mixing events caused by the observed high wind speeds during January and February (Fig. 3). Similar results were found between 2012 and 2015, where high nutrient concentrations were detected at the start of the northwest monsoon, declining during the transition period, and subsequently increasing during the southeast monsoon (LIPI monitoring program, not published). Ikhsani et al. (2016) reported that high nutrient levels during the southeast monsoon (wet season) in the inner Ambon Bay could be regulated by runoffs as well as cold-nutrient rich upwelled water entering from the outer bay, implying that intermonsoonal and -annual variability in precipitation may partly be responsible for the observed nutrient dynamics in our study.

Throughout the dry and wet season, Pseudo-nitzschia spp. was found to be persistent in the bay. Abundance maxima found in this study were relatively low compared to other locations (Klein et al., 2010; Bowers et al., 2018; Liefer et al., 2013; Vilas et al., 2014) including Asia (Teng et al., 2013). Interestingly, Pseudo-nitzschia spp. abundance strongly correlated with pDA levels throughout the campaign (Fig. 9). The average toxin cell quota in this study varied among seasons, which was higher during the wet season (on average 0.4 pg DA cell<sup>-1</sup>; ranging between 0.01 and 4.1 pg DA cell<sup>-1</sup>) than during the dry season (on average 0.1 pg DA cell<sup>-1</sup>; ranging between 0 and 0.6 pg DA cell<sup>-1</sup>). Previous field studies had shown weak correlations between Pseudonitzschia and DA level (Bates et al., 1998; Klein et al., 2010; reviewed by Lelong et al., 2012). Delegrange et al. (2018) found Pseudo-nitzschia and DA to be correlated in spring (0.02 pg DA cell<sup>-1</sup>) but not during the rest of season in the southern North Sea. In contrast, DA was found to correspond to Pseudo-nitzschia abundance in the Southern California Bight during two years of monitoring (2013 (5.7  $\pm$  10.6 pg DA cell<sup>-1</sup>); 2014 (0.3  $\pm$  0.9 pg DA cell<sup>-1</sup>)) (Smith et al., 2017). Moreover, Pseudonitzschia blooms were found to be associated with high DA levels in the eddy area of Juan de Fuca (Trainer et al., 2002, reviewed by Trainer et al., 2012). Umhau et al. (2018) found a significant correlation between particulate DA and Pseudo-nitzschia abundance in an offshore area of Santa Barbara Basin, California (37  $\pm$  124 pg DA cell<sup>-1</sup>). In the Bay of Seine, France, high DA concentrations were correlated with Pseudo-nitzschia australis, P. pungens and P. fraudulenta, but no correlation was found with a bloom of *P. delicatissima* (Thorel et al., 2017). The observed high variability in DA content may be related to species or strain specific differences, apart from cell specific DA variability related with environmental conditions such as nutrients, temperature or biotic influences (Trainer et al., 2012; Lelong et al., 2012).

The persistent occurrence of pDA and strong correlations with *Pseudo-nitzschia* spp. abundance as found in the present study imply that this species thrives and survives in Ambon Bay throughout the seasons. Even though detailed taxonomic identification based on electron microscope could not be done, the morphology of the most dominant *Pseudo-nitzschia* cells showed highly similar traits to *P. pungens* (Fig. 7). Based on the distribution of *Pseudo-nitzschia* species around the world, *P. pungens* is ubiquitously distributed in both tropical and temperate waters (Naz et al., 2012; Lim et al., 2014; Kim et al., 2015). Furthermore, *P. pungens* is regularly observed in Indonesian waters, yet toxin analyses were not reported before (Trainer et al., 2012; Lelong et al., 2012).

Several environmental factors stimulated both Pseudo-nitzschia spp. abundance and pDA levels in the bay, but these factors differed between seasons (Table 1). Spearman Rho Analysis showed no clear single factor that strongly governed Pseudo-nitzschia spp. abundance and pDA. Yet, ammonium was found as the main factor that correlated with Pseudonitzschia spp. abundance and pDA throughout the seasons. High constant ammonium availability, exceeding that of nitrate (Fig. 5E-H), might benefit persistent Pseudo-nitzschia growth and pDA production in the bay. Wastewater is discharged continuously in Ambon Bay supporting nitrification of ammonium as well as direct uptake, providing a substantial nitrogen source for the primary producers (McLaughlin et al., 2017), including Pseudo-nitzschia and its toxin (Cochlan et al., 2008; Auro and Cochlan, 2013; Martin-Jézéquel et al., 2015). By utilizing multiple nitrogen resources such as ammonium, urea and nitrate, Pseudo-nitzschia had demonstrated a persistent growth and enhanced toxin production (Thessen et al., 2009). The presence of fish farms in Ambon Bay, especially near stations C, D and E might contribute further to nitrogen input, however, no clear spatial differences in Pseudo-



Fig. 8. PCA analysis of weather, environmental and biological data represent station (A) and season (B) groupings based on score plots (n = 210).

#### Table 1

Spearman rho correlations between *Pseudo-nitzschia* spp., particulate domoic acid (pDA) and environmental properties (Abbreviations; MLD (Mixed Layer Depth), P0 (Phosphate 2 m), N0 (Nitrate at surface 2 m), S0 (Silicate 2 m), A0 (Ammonium 2 m), P20 (Phosphate 20 m), N20 (Nitrate 20 m), S20 (Silicate 20 m), A20 (Ammonium 20 m), Chl-a net (Chlorophyll-a  $\geq$  20 µm from the net)).

Environmental factors	Dry season ( $n = 96$ )		Wet season $(n = 72)$	
	Pseudo- nitzschia spp.	pDA	Pseudo- nitzschia spp.	pDA
Pseudo-nitzschia spp.	-	r = 0.66***	-	r = 0.87***
MLD	r = 0.35***	r = 0.32***	r = -0.28*	r = -0.28*
Stratification	$r = -0.28^{**}$	r = -0.24*	nc	nc
Temperature	nc	r = 0.5***	nc	nc
Salinity	nc	nc	nc	nc
Precipitation	r = -0.26*	r = -0.24*	nc	nc
PO	nc	r = -0.24*	nc	nc
N0	nc	nc	nc	nc
S0	nc	nc	nc	nc
A0	r = 0.23*	r = 0.24*	r = 0.24*	nc
P20	$r = -0.4^{***}$	$r = -0.5^{***}$	nc	nc
N20	r = -0.21*	$r = -0.4^{***}$	nc	nc
S20	nc	r = -0.23*	nc	nc
A20	nc	r = 0.3***	r = 0.25*	r = 0.26*
Chl-a net	$r = 0.6^{***}$	r = 0.57***	$r = 0.84^{***}$	r = 0.77***

Remarks: \*\*\*p = 0.00; \*\* $p \le 0.01$ ; \* $p \le 0.05$ ; nc (not correlated).

nitzschia spp. as well as pDA level were found. The other nutrients (phosphate, nitrate and silicate) inversely correlated with both Pseudonitzschia spp. and pDA in the dry season. Similar negative correlations were observed in the San Pedro Channel and Los Angeles harbor in Southern California (Schnetzer et al., 2007) and in the Western English Channel (UK) (Downes-Tettmar et al., 2013). Meanwhile, in the period of higher nutrient availability during the wet season, both particulate DA levels (Fig. 6B) and DA cell quotas were found to be higher. In both seasons, peaks of pDA were found when phosphate and silicate decreased dramatically, implying that the lack of these nutrients increased toxin production. In accordance with this result, previous studies both in the field and lab experiments had shown that phosphate and silicate limitation triggered high DA production in Pseudo-nitzschia (Fehling et al., 2004; Lelong et al., 2012; Trainer et al., 2012; Tatters et al., 2012; Parson et al., 2013; Terseleer et al., 2013; Lema et al., 2017; Bowers et al., 2018).

In our study, *Pseudo-nitzschia* spp. and pDA were positively correlated with mixed layer depth in the dry season but negatively correlated in the wet season. In addition, a strong positive correlation between pDA and temperature occurred during the dry season. Strong and warm winds blow in Ambon Island during the northwest monsoon provoking upper water mixing in the inner bay, enriching the water column with nutrient rich bottom water. During the wet season however, nutrient rich freshwater input enhances water column stratification despite elevated wind speeds. Coupled with the runoff discharges, the upwelling onset with high nutrient concentration seemed to fuel *Pseudonitzschia* spp. growth and more pDA production during the southeast



◎ Dry 🛛 🛆 Wet — — - Linear (Dry) ••••••• Linear (Wet)

Fig. 9. Linear regression model of Pseudo-nitzschia spp. cell abundance versus pDA during the dry and wet season.

monsoon, even though cell abundances were moderate and no blooms were found. In Monterey Bay, increases of *Pseudo-nitzschia* cell density coincided with upwelled nutrient-rich water (Bowers et al., 2018). Meanwhile, high DA levels were associated with relatively low sea surface temperature and high salinity in the Santa Barbara Channel (Anderson et al., 2009). Temperature dependent DA and *Pseudo-nitzschia* responses vary between regions, seasons and species as has been shown by both field and laboratory studies (Amato et al., 2010; Klein et al., 2010; Ljubešić et al., 2011; Marić et al., 2011; Hansen et al., 2011; Sahraoui et al., 2012; Thorel et al., 2014; Kim et al., 2015; Bresnan et al., 2017). In the subtropical Gulf of Mexico, the average temperature during the *Pseudo-nitzschia* spp. bloom period was  $28 \pm 1.3$  °C (Liefer et al., 2013), which was similar to sea water temperatures in Ambon Bay during both seasons.

#### 5. Conclusions

In conclusion, several environmental conditions seem to govern Pseudo-nitzschia spp. and pDA in Ambon Bay, of which ammonium variability seems the most pronounced. Thus, based on the present data, the prediction of future Pseudo-nitzschia spp. outbreaks is problematic, although factors like human population growth affecting waste water discharge, and elevated precipitation as well as land discharges through massive runoffs during the wet season might stimulate Pseudo-nitzschia spp. through enhanced stratification and nutrient (ammonium) input. Furthermore, possible influences of biotic factors including grazers (Leandro et al., 2010; Tammilehto et al., 2012; Selander et al., 2019), allelopathy (Van Meerssche et al., 2018) and bacteria (Lelong et al., 2012; Sison-Mangus et al., 2014) were not determined and therefore need to be addressed in future studies. pDA and Pseudo-nitzschia spp. were observed throughout the year during both the dry and wet season, implying that Ambon Bay might be potentially subject to amnesia shellfish poisoning. Thus, future research needs to reveal if the observed low but persistent DA levels accumulate in local shellfish, thereby causing a potential threat for the environment as well as human health.

#### Author declaration

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- Each author has participated sufficiently in the work to take public responsibility for all the content.

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Submission of an article implies that the work described has not

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#### S. Likumahua, et al.

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