Characterization of multiple isolates from an *Alexandrium ostenfeldii* bloom in The Netherlands

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ARTICLE INFO

Article history:
Received 23 December 2014
Received in revised form 21 August 2015
Accepted 21 August 2015

Keywords:
Harmful algal blooms
PSP toxins
Spiroloide
Gymnodimine
Lytic activity
Allelochemical potency

ABSTRACT

*Alexandrium ostenfeldii* is an emerging harmful algal bloom species forming a global threat to coastal marine ecosystems, with consequences for fisheries and shellfish production. The Oosterschelde estuary is a shallow, macrotidal and mesotrophic estuary in the southwest of The Netherlands with large stocks of mussels, oysters, and cockles. These shellfish stocks were threatened by a recent *A. ostenfeldii* bloom in the Ouwerkerkse Kreek, which is a brackish water creek discharging water into the Oosterschelde. Little is yet known about the characteristics of the *A. ostenfeldii* population in this creek. We therefore isolated 20 clones during an *A. ostenfeldii* bloom in 2013, and characterized these clones on their growth and toxin profile in their exponential growth phase. The cyclic imines were identified by comparison of *A. ostenfeldii* extracts with the retention time and CID spectra of standard solutions, or with published CID spectra. We furthermore assessed the allelochemical potency and phylogeny of a selection of 10–12 clones. Morphology and molecular phylogeny showed that all clones belong to Group 1 of *A. ostenfeldii*. All clones showed comparable growth rates of on average 0.22 ± 0.03 d−1. During exponential growth, they all produced a unique combination of paralytic shellfish poisoning toxins, spirolides and gymnodimines, of which particularly the latter showed a high intra-specific variability, with a 25-fold difference between clones with the lowest and highest cell quota. Furthermore, the selected 12 clones showed high allelopathic potentials with EC50 values based on lysis assays against the cryptophyte *Rhodomonas salina* between 212 and 525 *A. ostenfeldii* cells ml−1. Lytic activities were lower for cell extracts, indicating an important extracellular role of these compounds. A high intra-specific variability may add to the success of genotypically diverse *A. ostenfeldii* blooms, and make populations resilient to changes in environmental and climatic conditions.

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1. Introduction

Harmful algal blooms are a global threat to coastal marine ecosystems, with consequences for fisheries and shellfish production (Anderson et al., 2002; Heisler et al., 2008; Wang, 2008). *Alexandrium* is among the most common bloom forming toxic dinoflagellate genera and is generally held responsible for the outbreak of paralytic shellfish poisoning (PSP; Anderson et al., 2012). Besides PSP toxins, some *Alexandrium* species are known to produce other toxins including different spirolides (SPX) or gymnodimines (Cembella, 2003; Van Wagoner et al., 2011; Borkman et al., 2012; Kremp et al., 2014). These species belong to different phylogenetic groups of *Alexandrium ostenfeldii* as recently defined by Kremp et al. (2014), including *Alexandrium peruvianum*. Most *A. ostenfeldii* strains have been shown to produce SPX, some strains also produce detectable amounts of PSP toxins, while only a few strains were reported to combine PSP toxin, SPX and 12-methylgymnodimine production (Cembella, 2003; Borkman et al., 2012; Kremp et al., 2014).

PSP toxins are a group of neurotoxic compounds, including saxitoxin (STX), neosaxitoxin (NEO), gonyautoxins (GTX), and their N-sulfocarbamoyl variants the B- and C-toxins (Shimizu, 1996; Cembella, 1998). STX is highly toxic with an LD50 value (i.p. mice) of 8 μg kg−1 body weight (Wiberg and Stephenson, 1960). The addition of a sulfate group at the C-11 position forms GTX, thereby reducing the toxicity by up to 40%. A further addition of a sulfonyl...
group at the carbamoyl group forms C-toxins, which exhibit a 99% lower toxicity as compared to STX (Wiese et al., 2010). SPX and
gymnodimines are fast acting highly toxic neurotoxins, with LD_{50}
values (i.e., mice) down to 6.9 and 96 μg kg⁻¹ body weight,
respectively (Munday et al., 2004, 2012). Besides toxicity, various
Alexandrium species were shown to have allelopathic potencies
toward grazers and other phytoplankton species (Tillmann and
John, 2002; Tillmann et al., 2007, 2008; John et al., 2015).

Alexandrium ostenfeldii is globally distributed in brackish and
marine environments, and isolates have been characterized from
locations worldwide (Kriemp et al., 2014). A. ostenfeldii is
historically seen as background bloomer (Cembella et al., 2000;
John et al., 2003), however, dense A. ostenfeldii blooms have been
reported recently in the Narragansett Bay and the New River Estuary in the U.S. east coast (Borkman et al., 2012; Tomas et al.,
2012), the Baltic Sea coast of Finland (Hakanen et al., 2012), along
the Adriatic coast of Italy (Ciminiello et al., 2006), and recently in
a creek of the Oostschelde estuary in the Southwest of The
Netherlands (Burson et al., 2014). This Oostschelde estuary is a
shallow, macrotidal and mesotrophic estuary with large stocks of
mussels, oysters, and cockles (Fig. 1A and B; Troost et al., 2010; van
Broekhoven et al., 2014). Because of potential contamination of
these shellfish with phytoplankton toxins, The Netherlands has a
regular monitoring program for toxic compounds in shellfish, as
well as for the occurrence of harmful algal species in the
Oostschelde estuary (van der Fels-Klerx et al., 2011). A recent
dense A. ostenfeldii bloom in a creek discharging water into the
Oostschelde threatened the shellfish stocks, and was terminated
by the addition of hydrogen peroxide (Burson et al., 2014).
The bloom, however, recurred in 2013 and reached population
densities of up to 4500 cells mL⁻¹ (Fig. 1C). Little is yet known
about the characteristics of the A. ostenfeldii population in the
creek. We therefore isolated a number of A. ostenfeldii clones
during the bloom in 2013, and characterized the bloom population
in terms of growth, morphology, phylogeny, toxin composition,
and lytic activity.

2. Material and methods

2.1. Field sampling

The Ouwerkerkse Kreek is a small brackish water creek in the
province of Zeeland, Southwest of The Netherlands (Fig. 1A and B;
See also Burson et al., 2014). The inlet of the creek is connected
with ditches that drain the agricultural lands in the area, and water
from the creek is regularly discharges into the Oostschelde
estuary via a pumping station. The phytoplankton population in
the creek was monitored for Alexandrium ostenfeldii, and weekly
samples were taken for cell counts during the bloom. An integrated
water sample was taken of the upper 1 m of the water column at
location ‘a’ (Fig. 1B), and a 1 L sub sample was fixed with Lugol’s
iodine solution (Lugol) to a final concentration of 1%. A. ostenfeldii
cells were counted in a Sedgewick chamber on an inverted
microscope (Olympus Vanox, Hamburg, Germany). Isolates of A.
ostenfeldii cells were sampled at the onset of the bloom on 16 July
2013 from locations ‘a’ and ‘b’ in the creek (Fig. 1B).

2.2. Isolation

Cells of Alexandrium ostenfeldii were picked from small droplets
in a petridish using a Pasteur pipette or a 10 μL micro-pipette.
Individual cells were cleaned five times in 2 μL droplets of sterile
medium consisting of filtered and diluted North Sea water with a
salinity of about 10, or in sterilized artificial brackish water medium
with a salinity of about 10 (Appendix A). Both media
contained nutrients corresponding to 50% of K-medium (Keller
et al., 1987). A. ostenfeldii cells were subsequently grown in 100 μL
medium mixed in a 1:1 ratio with sterile filtered water from the
creek (0.2 μm membrane filter) in microplate wells. Clones
OKNL-10 were subsequently cultured in artificial brackish water
medium (Appendix A), and clones ONNL11-22 in diluted North Sea
water medium.

2.3. Culturing of clones

Twenty of the successfully isolated clones were grown in
250 mL Erlenmeyer flasks at 15 °C under an incident light intensity
of 100 μmol photons m⁻² s⁻¹ at a light-dark cycle of 16:8. After
acclimation to these growth conditions (i.e. >5 generations), cells
were transferred and growth was monitored by cell counts performed
every second day. At mid-exponential phase (∼6000 cells mL⁻¹) cultures were harvested for analysis of toxins
and for cell size measurements. Cultures were counted again
two days later, confirming that cell sampling two days before was
still during the exponential phase. For toxin sampling, 15 mL samples
were taken for extraction of SPX and gymnodimines, and 50 mL
samples each were taken for PSP toxin and DNA extraction.
Samples were centrifuged at 6800 × g for 15 min (SL16, Thermo
Scientific, Waltham, USA) and after removal of the supernatant the
pellets were stored at −20 °C.

2.4. Cell counts and measurements

Cell densities of Alexandrium ostenfeldii cultures were deter-
mimed by using sedimentation chambers for settling 0.2–1 mL of
culture suspension, and subareas with at least 400 cells were
counted with an inverted microscope (20X, Zeiss Axiosvert 40C).
Observation and documentation of live and fixed cells was carried

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Fig. 1. Overview of the sampling location and the bloom dynamics of Alexandrium ostenfeldii in Rhine-Muese-Scheldt delta, Southwest of The Netherlands (A). With the specific sampling locations ‘a’ and ‘b’ in the Ouwerkerkse Kreek (B), and the A. ostenfeldii population densities during the 2013 bloom sampled from location ‘a’ (C).
out with either an inverted microscope (Axiovert 200M, Zeiss, Germany) or a Zeiss Axioskop 2 (Zeiss, Germany), both equipped with epifluorescence and differential interference contrast optics. Length and width of cells (n > 50 from each clone) were measured on freshly Lugol-fixed cells (1% final concentration) with the program AxioVs40 (v.4.8., Zeiss, Göttingen, Germany) using micrographs taken with a digital camera (Axiocam MRc, Zeiss, Göttingen, Germany).

Growth rates were estimated for each clone by means of an exponential function fitted through all cell counts over time (n = 4), according to \( N_t = N_0 \exp^{\mu t} \), where \( N_t \) refers to the cell concentrations at time \( t \), \( N_0 \) to the cell concentrations at the start of the experiment, and \( \mu \) to the growth rate.

2.5. Cell morphology and morphometry

Pattern, shape and dimension of thecal plates were examined using epifluorescence microscopy of calcifluor-stained cells according to the method of Fritz and Triemer (1985). Morphometric measurements were performed for four randomly selected clones (OKNL11, 12, 14, 19). Cells of exponentially growing cultures were fixed with neutral Lugol (1% final concentration) for 1 h and then collected by centrifugation at 3220 \( \times \) g for 10 min (Eppendorf 5810R, Hamburg, Germany). Cells were resuspended in a drop of filtered seawater and a drop of a 1 mg mL\(^{-1}\) solution of Fluorescence Brightener 28 (Sigma–Aldrich, St. Louis, MO, USA) on a microscope slide. Cells were inspected at 1000 \( \times \) magnification (Zeiss, Axioskop 2, Göttingen, Germany) and photographed with a digital camera (Axiocam MRc, Zeiss, Göttingen). Measurements of size and area of selected diagnostic plates were performed with the program AxioVs40 (v.4.8., Zeiss, Göttingen). Shape of the first apical plate and the anterior sulcal plate was scored into categories given by Kremp et al. (2014).

2.6. Toxins

2.6.1. Toxin extraction

Freeze dried cell pellets were suspended in 500 \( \mu \)L 0.03 M acetic acid for PSP toxin analysis and with methanol for SPX analysis, respectively. The samples were subsequently transferred into a Fast Prep tube containing 0.9 g of lysing matrix D. The samples were homogenized by reciprocal shaking at maximum speed (6.5 m s\(^{-1}\)) for 45 s in a Bio101 Fast Prep instrument (Thermo Savant, Illkirch, France). After homogenization, samples were centrifuged (Eppendorf 5415 R, Hamburg, Germany) at 16,100 \( \times \) g at 4 \(^\circ\)C for 15 min. The supernatants were transferred to spin-filters (pore-size 0.45 \( \mu \)m, Millipore Ultrafree, Eschborn, Germany) and centrifuged for 30 s at 800 \( \times \) g. The filtrates were transferred to HPLC vials and stored at -20 \(^\circ\)C until measurement. Variation in cellular toxin quota and composition was tested by growing four clones in triplicate.

2.6.2. Toxin analysis

2.6.2.1. PSP toxins. The aqueous extracts were analyzed for PSP toxins by reverse-phase ion-pair liquid chromatography with fluorescence detection (LC-FLD) and post-column derivatization following minor modifications of previously published methods (Dienert et al., 2006; Krock et al., 2007). The LC-FLD analysis was carried out on a LC1100 series liquid chromatography system consisting of a G1379A degasser, a G1311A quaternary pump, a G1229A autosampler, and a G1321A fluorescence detector (Agilent Technologies, Waldbronn, Germany), equipped with a Phenomenex Luna C18 reversed-phase column (250 mm \( \times \) 4.6 mm id, 5 \( \mu \)m pore size) (Phenomenex, Aschaffenburg, Germany) with a Phenomenex SecuriGuard precolumn. The column was coupled to a PCX 2500 post-column derivatization system (Pickering Laboratories, Mountain View, CA, USA). Eluent A contained 6 mM octanesulfonic acid, 6 mM heptanesulfonic acid, 40 mM ammonium phosphate, adjusted to pH 6.95 with dilute phosphoric acid, and 0.75% tetrahydrofuran. Eluent B contained 13 mM octanesulfonic acid, 50 mM phosphoric acid, adjusted to pH 6.9 with ammonium hydroxide, 15% acetonitrile and 1.5% tetrahydrofuran. The flow rate was 1 mL min\(^{-1}\) with the following gradient: 0–15 min isocratic A, 15–16 min switch to B, 16–35 min isocratic B, 35–36 min switch to A, 36–45 min isocratic A. The injection volume was 20 \( \mu \)L and the autosampler was cooled to 4 \(^\circ\)C. The eluate from the column was oxidized with 10 mM periodic acid in 555 mM ammonium hydroxide before entering the 50 \(^\circ\)C reaction coil, after which it was acidified with 0.75 M nitric acid. Both the oxidizing and acidifying reagents entered the system at a rate of 0.4 mL min\(^{-1}\). The toxins were detected by dual-monochromator fluorescence (\( \lambda_{em} \) 333 nm; \( \lambda_{ex} \) 395 nm). The data were processed with Agilent Chemstation software and calibrated against external standards. Standard solutions of PSP toxins were purchased from the Certified Reference Material Program of the Institute of Marine Biosciences (National Research Council, Halifax, NS, Canada).

2.6.2.2. Lipophilic toxins.

2.6.2.2.1. Selected reaction monitoring (SRM) experiments. Mass spectral experiments for lipophilic toxin detection and quantification were performed on a 4000 Q Trap (AB-SCIEX, Darmstadt, Germany), triple quadrupole mass spectrometer equipped with a TurboSpray\(^{\circledR}\) interface coupled to an Agilent (Waldbrohn, Germany) 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).

After injection of 5 \( \mu \)L of sample, separation of lipophilic toxins was performed by reverse-phase chromatography on a C8 column (50 \( \times \) 2 mm) packed with 3 \( \mu \)m Hypersil BDS 120 A (Phenomenex, Aschaffenburg, Germany) and maintained at 25 \(^\circ\)C. The flow rate was 0.2 mL min\(^{-1}\) and gradient elution was performed with two eluents, where eluent A was water and eluent B was methanol/water (95.5, v/v), both containing 2.0 mM ammonium formate and 50 mM formic acid. Initial conditions were elution with 5% B, followed by a linear gradient to 100% B within 10 min and isocratic elution until 10 min with 100% B. The program was then returned to initial conditions within 1 min followed by 9 min column equilibration (total run time: 30 min).

Mass spectrometric parameters were as follows: curtain gas: 20 psi, CAD gas: medium, ion spray voltage: 5500 V, temperature: 650 \(^\circ\)C, nebulizer gas: 40 psi, auxiliary gas: 70 psi, interface heater: on, declustering potential: 121 V, entrance potential: 10 V, exit potential: 22 V, collision energy: 57 V. SRM experiments were carried out in positive ion mode by selecting the following transitions [precursor ion > fragment ion]: m/z 508 > 490 for gymnodioline A, 522 > 504 for 12-methylgymno- dine, 692 > 164 for 13-desmethyiospirole C (SPX-1), and 694 > 164 for 13-desmethyiospirole D. Dwell times of 40 ms were used for each transition. Standard solutions of SPX-1 and gymnodioline A were purchased from the Certified Reference Material Program of the Institute of Marine Biosciences (National Research Council, Halifax, NS, Canada), and 12-methylgymnodimine was kindly provided by Kiri Harju, VERIFIN, Department of Chemistry, University of Helsinki, Finland.

2.6.2.2.2. Product ion spectra. Product ion spectra were recorded in the Enhanced Product Ion (EPI) mode in the mass range from m/z 120 to 550. Positive ionization and unit resolution mode were used. The mass spectral parameters were as in SRM experiments.
2.7. *Rhodomonas salina* bioassays

Clonal isolates of *Alexandrium ostenfeldii* (OKNL11-22) were grown in batch cultures in 100 mL Erlenmeyer flasks at standard culture conditions described above. Growth was followed by counting Lugol-fixed (1% final concentration) samples every second or third day using an inverted microscope. At late exponential phase, at a population density of 8–12 × 10^3 cells mL⁻¹, cells were harvested for estimating both lytic capacity of whole cell cultures and of cell extracts. A subsample of 20 mL live sample was taken for whole cell lytic activity. A subsample of 50 mL was collected by centrifugation (Eppendorf 5810R, Hamburg, Germany) at 3220 × g for 10 min). The cell pellet was transferred to a 1 mL microtube, centrifuged again (Eppendorf 5415, 16,000 × g, 5 min), and stored frozen (−20 °C) until use.

Lytic activity of whole cell culture and cell extract were both estimated by a *Rhodomonas salina* lysis assay (Tillmann et al., 2009). For this bioassay, *R. salina* (strain KAC30, Kalmar culture collection, Sweden) was pre-adapted to and subsequently grown in 10 PSU medium for at least four weeks. For testing the variation in lytic activity, a full dose-response-curve of lytic activity of whole culture and cell extract was recorded for triplicate cultures of clone OKNL19. Here, nine different dilutions ranging from 4000 to 50 cells mL⁻¹ and from 30,000 to 100 cells mL⁻¹ for whole culture and cell extract, respectively, were prepared in triplicate and processed as described below. Lytic activity of all other clones was estimated with a reduced number of dilutions. For whole cell lysis estimates, four different dilutions of the *Alexandrium ostenfeldii* live samples were prepared (undiluted, 3000, 1000, and 250 cells mL⁻¹; final concentration in the assay) in triplicate volumes of 3.9 mL in 6 mL glass vials. Each sample was spiked with 0.1 mL of a pre-adjusted culture of *R. salina* yielding a final concentration of 10 × 10⁶ cells mL⁻¹. The first of two negative controls contained only K-medium, whereas the second negative control was performed with *Alexandrium tamarense*, strain Alex5, a strain which previously was shown to be non-lytic (Tillmann and Hansen, 2009). Vials were then incubated at 15 °C in the dark. After 24 h, samples were fixed with Lugol (2% final concentration), and concentrations of intact cells of both species were determined under an inverted microscope.

For estimating lytic activity of cell extracts, cell pellets were thawed and re-suspended in 1 mL filtered seawater. Cellular extracts were prepared by sonicating the cell suspension with a Sonoplus HD70 disintegrator equipped with a MS73 sonotrode for 1 min on ice using the following settings: 50% pulse cycle, 70% amplitude. Based on the known number of cells in the pellet, the extract was used to prepare four concentrations of “*Alexandrium ostenfeldii* cells” in the 4 mL volume used in the bioassays (15,000, 7000, 3000, and 1500 cells mL⁻¹). The bioassays were then performed as described above except that just intact cells of *Rhodomonas salina* were counted.

All results were reported as final concentration of *Rhodomonas salina* expressed as percent of the seawater control. The data from each clone were plotted as log-transformed cell concentrations of *Alexandrium ostenfeldii* and corresponding percentage survival of *R. salina* (mean of three replicates). Data, even if a limited number of dilutions were performed, followed a sigmoid curve and were fitted as a dose–response curve with Eq. (1) in order to estimate the EC₅₀ of *A. ostenfeldii*, defined as the concentration of *A. ostenfeldii* cells resulting in a mortality of 50% of the *R. salina* population.

\[
Y = \frac{100}{1 + \left(\frac{X}{EC_{50}}\right)^k}
\]

Here, *Y* is the concentration of intact *Rhodomonas salina* cells (as percentage of the control), *X* is log-transformed cell concentration of *Alexandrium ostenfeldii*, and EC₅₀ and *k* are fitted parameters.

For the cell extract of clone OKNL18, EC₅₀ calculation by curve fitting failed as the highest dose (15,000 cells mL⁻¹) yielded just 41% cell lysis. EC₅₀ here was defined as “>15,000 cells mL⁻¹”.

2.8. Phylogeny

Ribosomal DNA sequence data for the phylogenetic analyses were obtained from *Alexandrium ostenfeldii* clones OKNL11-22. Initially, 50 mL samples of exponentially growing cells were collected by centrifugation at 3220 × g for 15 min at room temperature (Eppendorf 5810R, Hamburg, Germany). The cell pellets were frozen at −20 °C for 20 min before extraction of total DNA with the DNeasy Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. The purity and quantity of the DNA was assessed by UV-spectroscopy with a NanoDrop ND-1000 system (Peqlab, Erlangen, Germany) and the integrity of DNA was confirmed using 1% agarose gel electrophoresis where a majority of the extracted genomic DNA exceeded 20 kilobases.

The D1/D2 regions of the 28S large subunit (LSU) ribosomal DNA and the internal transcribed spacer region, including the ITS1, 5.8S subunit, and ITS2 sequences were amplified from total DNA extract by polymerase chain reaction (PCR). The forward and reverse primer for LSU amplification were: D1R-F (5’-ACG TGC TTA ATT TAA GCC TA-3’) and D2C-R (5’-CCT TGG TCC GTT CTT CAA GA-3’), respectively. The forward and reverse primers for ITS1/5.8 amplification were: ITS a (5’-CCA AGC TTC TAG ATG GTA ACA AGG (ACT)TCT AGG T-3’) and ITS b (5’-CCT GCA TGC CAC A(CT)ATG TTA A(AG)T TCA GC(G) GG-3’), respectively. Reaction conditions were as follows: HotMasterTag® (5Prime, Hamburg, Germany) buffer 1X, 0.1 mM of dNTPs, 0.1 mM of each forward and reverse primer and 1.25 units of Taq polymerase was added to 10–30 ng of the extracted genomic DNA in total reaction volumes of 50 μL. For 28S rDNA amplifications, the reactions were subjected to the following thermo cycling conditions: one cycle at 95 °C for 7 min, 35 cycles at 94 °C for 45 s, at 54 °C for 2 min, and at 70 °C for 1.5 min, and a final extension at 70 °C for 5 min. The thermal cycling conditions for the ITS5.8 amplifications were: one cycle at 94 °C for 4 min, 9 cycles at 94 °C for 50 s, at 60 °C for 40 s and at 70 °C for 1 min, and 29 cycles at 94 °C for 45 s, at 50 °C for 45 s, and at 70 °C for 1 min, and a final 5 min extension step at 70 °C. Samples were kept at 10 °C until analysis on 1% agarose gel, in order to ensure the expected amplification products were present. After PCR amplification and subsequent cloning into the vector provided with the TOPO TA Cloning® kit (Invitrogen, Carlsbad, California, USA), 3–8 clones per amplicon were sequenced using the M13 vector primers supplied with the kit. Sequencing was conducted with a standard cycle sequencing chemistry ABI 3.1 (Applied Biosystems, Darmstadt, Germany). Cycle sequencing products were analyzed on an ABI 3130 XL capillary sequencer (Applied Biosystems, Darmstadt, Germany) and the generated sequences were assembled with CLC main workbench version 6.0 (www.CLCBio.com). The resulting sequences were submitted to GenBank (Appendix B).

The LSU and ITS5.8 sequences were then combined with those available in GenBank (Appendix B) and a combined alignments of LSU and ITS sequences was constructed in order to conduct phylogenetic analyses (Appendix C). The data set was aligned with MAFFT using the q-insi option (Katoh et al., 2005). The full ML analysis (GTRGAMMA, 1000 bootstrap replicates) was conducted with RAxML (Stamatakis et al., 2005).

2.9. Statistical analysis

Normality of cell size parameters (length, width, length/width ratio, and volume), toxins (PSP toxins, SPX and gymnodimines) and...
lytic activity was tested using the Shapiro–Wilk test. Significance of
differences between clones was tested using a one-way ANOVA
if the data was normal distributed, and a Kruskal–Wallis test if the
distribution was non-normal. The associations between the
cellular quota of the different toxins were tested using Pearson
product-moment correlation.

3. Results

3.1. Growth, morphology, morphometry, and phylogeny

Growth rates for all isolates were comparable and ranged
between $0.16^{-1}$ and $0.26^{-1}$ with an average of $0.22 \pm 0.03$ d$^{-1}$
($n = 20$). Generally, cells were round to ellipsoid in shape (Fig. 2A–F).
The epitheca was slightly variable in shape with an outline ranging
from sigmoid to dome-shape or round (Fig. 2B–F). The generally
round hypotheca occasionally was slightly asymmetric (Fig. 2K).

Within and among clones there was also a considerable variability in
size (Fig. 2G–K). Cell length of all measured cells ranged from
26.3 to 53.4 μm, the width from 30.6 to 40.5 μm, and the length/
width ratio from 0.9 to 1.2 with significant differences among clones
in mean cell length (Kruskal–Wallis test; $H = 259; df = 19; P < 0.001$),
width (Kruskal–Wallis test; $H = 322; df = 19; P < 0.001$), and length/
width ratios (Kruskal–Wallis test; $H = 175; df = 19; P < 0.001$).

The plate pattern with a pore plate, 4 apical, 6 precingular,
5 postcircular, 2 antapical, 6 cingular, and 8 sulcal plates

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**Fig. 2. Alexandrium ostenfeldii morpho**gy. (A–K) Bright field images of a living cell (A), and Lugol-stained cells (B–K) to show variability in cell shape (A–F) and cell size (G–K). (L–Ad) Epifluorescence images of calcofluor stained cells showing the plate pattern (L–R), and details of the 6" plate (S and T), the 1" plate and ventral pore (U–X), and the s.a plate (Y–Ad). Plate labels according to the Kofoidean system. Sulcal plates abbreviations: s.p. = posterior sulcal plate. s.s.p. = left posterior sulcal plate. s.d.p. = right posterior sulcal plate. s.m.p. = median posterior sulcal plate. s.a.s. = left anterior sulcal plate. s.a.c.a. = anterior accessory sulcal plate. s.m.a. = median anterior sulcal plate. s.a. = anterior sulcal plate.
Table 1
Size measurements of all tested OKNL clones with length (L), width (W), and L/W ratio. Values show mean ± SD with the number of analyzed cells (n).

<table>
<thead>
<tr>
<th>Clone</th>
<th>L (μm)</th>
<th>W (μm)</th>
<th>L/W ratio</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>OKNL</td>
<td>41.7 ± 1.1</td>
<td>38.4 ± 3.4</td>
<td>1.09 ± 0.06</td>
<td>50</td>
</tr>
<tr>
<td>OKNL1</td>
<td>41.1 ± 2.8</td>
<td>38.0 ± 3.2</td>
<td>1.08 ± 0.06</td>
<td>50</td>
</tr>
<tr>
<td>OKNL2</td>
<td>43.1 ± 3.9</td>
<td>39.3 ± 3.1</td>
<td>1.10 ± 0.05</td>
<td>50</td>
</tr>
<tr>
<td>OKNL3</td>
<td>39.9 ± 3.2</td>
<td>37.7 ± 3.4</td>
<td>1.06 ± 0.05</td>
<td>42</td>
</tr>
<tr>
<td>OKNL4</td>
<td>40.9 ± 3.1</td>
<td>37.8 ± 2.9</td>
<td>1.06 ± 0.06</td>
<td>40</td>
</tr>
<tr>
<td>OKNL5</td>
<td>38.6 ± 3.5</td>
<td>33.6 ± 2.7</td>
<td>1.15 ± 0.07</td>
<td>50</td>
</tr>
<tr>
<td>OKNL6</td>
<td>41.9 ± 3.8</td>
<td>37.9 ± 2.5</td>
<td>1.11 ± 0.07</td>
<td>50</td>
</tr>
<tr>
<td>OKNL7</td>
<td>42.1 ± 2.9</td>
<td>40.5 ± 2.3</td>
<td>1.04 ± 0.03</td>
<td>56</td>
</tr>
<tr>
<td>OKNL8</td>
<td>39.1 ± 4.5</td>
<td>37.0 ± 4.3</td>
<td>1.07 ± 0.05</td>
<td>56</td>
</tr>
<tr>
<td>OKNL9</td>
<td>37.6 ± 6.1</td>
<td>35.3 ± 5.4</td>
<td>1.07 ± 0.05</td>
<td>56</td>
</tr>
<tr>
<td>OKNL10</td>
<td>40.6 ± 4.2</td>
<td>38.8 ± 3.7</td>
<td>1.04 ± 0.05</td>
<td>55</td>
</tr>
<tr>
<td>OKNL11</td>
<td>42.7 ± 4.4</td>
<td>40.4 ± 3.5</td>
<td>1.06 ± 0.05</td>
<td>53</td>
</tr>
<tr>
<td>OKNL12</td>
<td>41.3 ± 3.8</td>
<td>38.8 ± 3.1</td>
<td>1.07 ± 0.05</td>
<td>55</td>
</tr>
<tr>
<td>OKNL13</td>
<td>40.2 ± 3.9</td>
<td>38.9 ± 3.1</td>
<td>1.03 ± 0.04</td>
<td>55</td>
</tr>
<tr>
<td>OKNL14</td>
<td>34.4 ± 3.8</td>
<td>32.6 ± 3.4</td>
<td>1.06 ± 0.05</td>
<td>56</td>
</tr>
<tr>
<td>OKNL15</td>
<td>42.3 ± 4.1</td>
<td>40.1 ± 3.2</td>
<td>1.06 ± 0.05</td>
<td>55</td>
</tr>
<tr>
<td>OKNL16</td>
<td>39.8 ± 4.4</td>
<td>37.3 ± 4.2</td>
<td>1.07 ± 0.05</td>
<td>55</td>
</tr>
<tr>
<td>OKNL17</td>
<td>32.5 ± 4.2</td>
<td>30.6 ± 4.1</td>
<td>1.06 ± 0.04</td>
<td>54</td>
</tr>
<tr>
<td>OKNL18</td>
<td>40.8 ± 3.3</td>
<td>39.3 ± 2.6</td>
<td>1.04 ± 0.05</td>
<td>56</td>
</tr>
<tr>
<td>Overall mean</td>
<td>40.1 ± 2.6</td>
<td>37.6 ± 2.6</td>
<td>1.07 ± 0.03</td>
<td>20</td>
</tr>
</tbody>
</table>

(Fig. 2L–R) was typical for the genus *Alexandrium* and the typical shape of the first apical plate and the large ventral pore (Fig. 2L–N) clearly identified all isolates as *Alexandrium ostenfeldii*. Detailed morphometric analyses of four isolates indicated some variability in shape of the diagnostic plates 1’, 6’ and s.a. (Table 2). For most of the cells the right anterior margin of the first apical plate 1’ was straight (Fig. 2L and M), however for all four clones there were cells (up to 27% for clone OKNL11) with curved or irregular margins (Fig. 2U and V). A significant number of cells ranging from 22% to 43% had an anteriorly extended 1’ plate (Fig. 2W and X), which can explain some of the variability of the 1’ area (Table 2). The mean ventral pore area ranged from 2.3 to 4.6 μm² and was different between the four analyzed clones (Table 2). The shape of the anterior sulcal plate s.a. was found to be variable within all four analyzed clones. For the majority of cells the shape was classified as “A-shaped” (Fig. 2A and Ab), but about 20% and 30% of the cells of each isolate had a “door-latch” shaped (Fig. 2Y and Z) and a “horse-shoe” shaped (Fig. 2Ac and Ad) s.a. plate, respectively (Table 2). The s.a. plate was more wide than high with the width/height ratio of about 1.3 being quite similar for the four clones (Table 2). The terminal precingular plate 6’ was found to be quite variable in its width/height ratio (Fig. 2S and T), with a mean being even below one for clone OKNL12 (Table 2). Based on the ITS/LSU phylogenetic analyses (Fig. 3), the tested clones belonged to Group 1 of *A. ostenfeldii*.

3.2. Toxins and lytic activity

All 20 isolates of *Alexandrium ostenfeldii* produced PSP toxins with cellular quota ranging between 9.5 and 51 pg cell⁻¹ (Fig. 4A). The total PSP content was significantly different (one-way ANOVA; F₃,₁₇ = 21.3; P < 0.001) between the four replicated clones (OKNL11, 12, 15 and 19), with an average relative standard deviation (i.e. the standard deviation relative to the average PSP content of each clone) of 10 ± 5%. The GTX and C1/C2 toxins were the predominant analogs in all clones, the latter comprising around 84 ± 5% of the total PSP content (Fig. 4B). The relative contribution of GTX2/3 was also largely comparable between all clones contributing to around 9.6 ± 2% of the total PSP content, while STX showed a much larger variability contributing between 2.4% and 18% of the total PSP content (Fig. 4B). Among the cyclic imine toxin group, 13-desmethysprodigiosin C (SPX-1) and gymnodiurm A (data not shown), and 12-methylgymnodimide could be identified by comparison of retention time (Fig. 5) and CID spectra (Fig. 6) from *A. ostenfeldii* extracts with those from standard solutions. 13-desmethysprodigiosin D was identified by comparison with a previously published CID spectrum (Sienko et al., 2004).

The total gymnodiurm content (i.e. gymnodiurm A and 12-methylgymnodimide) showed a large variation between clones with a 25-fold difference between the lowest and highest values (Fig. 7A). The total gymnodiurm content differed significantly between the four tested clones (one-way ANOVA; F₃,₁₇ = 98.2; P < 0.001), with an average relative standard deviation of 7 ± 3%. All clones predominantly produced gymnodiurm A, while most clones also produced 12-methylgymnodimide with relative contributions ranging between 0.2% and 37% (Fig. 7A). The cellular quota of total SPX (expressed as SPX-1 equivalents) generally ranged between 2.4 and 4.4 pg cell⁻¹, with three clones containing less than 1.8 pg cell⁻¹ (Fig. 7B). The SPX included 13-desmethysprodigiosin C (76% to 86%) and 13-desmethysprodigiosin D (14% to 24%, Fig. 7B). All clones also produced trace levels of several unknown SPX-like compounds (data not shown). The total SPX content significantly differed between the four tested clones (one-way ANOVA; F₃,₁₇ = 37.1; P < 0.001), with an average relative standard deviation of 8 ± 2%. We observed a significant correlation between the cellular quota of total PSP toxins and total SPX (r = 0.568; n = 20; P = 0.009), total PSP toxins and total gymnodiurms (r = 0.472; n = 20; P = 0.031), and total SPX and total gymnodiurms (r = 0.526; n = 20; P = 0.017).

All 12 selected clones induced lysis of *Rhodomonas salina*, with EC₅₀ values ranging between 212 and 525 *Alexandrium ostenfeldii* cells ml⁻¹ (Fig. 8A and B). The lytic response toward cellular extract was less pronounced and corresponded to 1222 to 5839 *A. ostenfeldii* cells ml⁻¹, and was even beyond the highest tested cell concentration of 15,000 cell ml⁻¹ for strain OKNL18 (Fig. 8C). A quantitative comparison of cell lysis between whole cell culture and cell free supernatant of strain OKNL21 showed that about 70% of the total lytic activity is present without the cells (data not shown). We did not observe a significant correlation between the EC₅₀ values and the different cellular toxin quota.

4. Discussion

The morphological characteristics show a high variability within the population, supporting earlier descriptions of members

Table 2
Diagnostic plate features of four OKNL clones. Values show mean, and mean ± SD, with the number of analyzed cells (n).

<table>
<thead>
<tr>
<th>Clone</th>
<th>Shape 1’ plate</th>
<th>1’ extension</th>
<th>Shape s.a.</th>
<th>Area vp (μm²)</th>
<th>Area 1’ (μm²)</th>
<th>6’ ratio w/h</th>
<th>s/a ratio w/h</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>OKNL1</td>
<td>72.6 (62)</td>
<td>21.9 (64)</td>
<td>16.1</td>
<td>55.4</td>
<td>28.6</td>
<td>1.04 ± 0.05</td>
<td>1.03 ± 0.05</td>
<td>50</td>
</tr>
<tr>
<td>OKNL2</td>
<td>86.5 (37)</td>
<td>22.2 (45)</td>
<td>24.4</td>
<td>40.0</td>
<td>35.6</td>
<td>1.06 ± 0.04</td>
<td>1.02 ± 0.01</td>
<td>45</td>
</tr>
<tr>
<td>OKNL3</td>
<td>92.1 (63)</td>
<td>42.9 (63)</td>
<td>17.9</td>
<td>53.8</td>
<td>28.2</td>
<td>1.03 ± 0.04</td>
<td>1.02 ± 0.01</td>
<td>50</td>
</tr>
<tr>
<td>OKNL4</td>
<td>93.8 (96)</td>
<td>33.3 (99)</td>
<td>14.4</td>
<td>58.9</td>
<td>26.7</td>
<td>1.03 ± 0.04</td>
<td>1.02 ± 0.01</td>
<td>45</td>
</tr>
</tbody>
</table>

The total ratio.
from Group 1 of Alexandrium ostenfeldii. The high variability in cell size within each clone, in combination with a relatively consistent length/width ratio of 1.1, is in line with earlier findings for clones belonging to Group 1 (Kremp et al., 2014). About 30% of the investigated cells showed an anteriorly extended 1’ plate, more common for Group 2. Previously considered as an important diagnostic plate, our analysis shows a variety of shapes for the s.a. plate within the clones, dominated by A-shaped plates generally accounting for 52%, but also with up to 36% of the cells containing a horseshoe-shaped s.a. plate, previously ascribed to Alexandrium peruvianum. We thus confirm the conclusions by Kremp et al. (2014) that the diagnostic characters originally defined to separate A. ostenfeldii and A. peruvianum are more variable than previously assumed and can show considerable intra- and inter-strain variability. Furthermore, Kremp et al. (2014) showed that A. ostenfeldii and A. peruvianum morphotypes often had nearly identical rDNA sequences indicating that they represent extreme ends in a continuum of A. ostenfeldii morphotypes (Kremp et al., 2014). Nevertheless, Kremp et al. (2014) also showed that selected plate features, when tested statistically, as well as toxin profiles, differed significantly among phylogenetic clusters. Our morphometric analysis thus indicate a designation of the Dutch A. ostenfeldii population as members of the Group 1 (see also Kremp et al., 2014), a conclusion clearly supported by both the molecular data and the presence of PSP toxins.

The phylogenetic analysis reveals a close relationship of the Dutch Alexandrium ostenfeldii population with isolates covering a wide geographical distribution, ranging from the coastal areas in the Baltic Sea to embayments in the North-East coast of the US (Fig. 3). This geographically wide distribution may indicate recent anthropogenic-driven dispersal, or that the phylogenetic clade comprises globally distributed populations with a broad trait range similar to the Groups (I-V) of the former Alexandrium tamarensis species complex (John et al., 2014). The emergence of A. ostenfeldii in the Ouwerkerkse Kreek seems to have occurred only recently, although some sporadic observations of Alexandrium sp. in the area were made in the last decade during monitoring programs (unpublished data). However, only in depth population genetics will help to evaluate a common origin and/or the potential dispersal routes of the A. ostenfeldii Group 1 populations.

Interestingly, within Alexandrium ostenfeldii Group 1, toxin profiles of the isolates from The Netherlands are most comparable to two isolates from North America, which are the only representatives which also produce PSP toxins, SPX, and gynomodimines (Van Wagoner et al., 2011; Borkman et al., 2012; Tomas et al., 2012). The Dutch isolates, however, differ in their gynomodimine composition. Specifically, the isolates from North America were reported to produce 12-methylgymnodimine, while the OKNI isolates produce both 12-methylgymnodimine as well as gynomodimine A, a compound previously known from Karenia only (Seki et al., 1995; MacKenzie et al., 1996; Miles et al., 2000). It thus seems that the clones we report here are the first record of gynomodimine A for A. ostenfeldii, and possess a unique combination of toxins. Whether the isolates from North America are not producing gynomodimine A, or if the levels are below detection limit requires further analyses. Generally, taking into account that advancing analytical techniques will lower the limit of detection for a multitude of toxins, future studies may show that more complex toxin profiles might generally be common for A. ostenfeldii populations.
The cellular quota of PSP toxins fell within the range of earlier reported *Alexandrium ostenfeldii* and *Alexandrium tamarense* clones (Tillmann et al., 2009; Suikkanen et al., 2013). The predominance of C1/C2 has been earlier reported for *A. ostenfeldii* clones, though the PSP toxin composition differs between isolates (Kremp et al., 2014). B1 toxins were not found in clones OKNL4, 12, 13, and 14, which may indicate that these clones lack the ability to produce B1, or that the values were below the level of detection (<0.1 pg cell⁻¹). Both the cellular quota of SPX and gymnodimines are relatively low as compared to an earlier reported *A. ostenfeldii* clone (then designated to *Alexandrium peruvianum*) from the US East Coast (Tatters et al., 2012), while SPX quota fell within the range of several *A. peruvianum* clones from the NE Atlantic (Suikkanen et al., 2013). SPX consisted mainly of 13-desmethylspirolid A, which was largely consistent with other clones in Group 1 and 2 of the *A. ostenfeldii* species complex (Kremp et al., 2014), though all OKNL clones also contained some 13-desmethylspirolid D. Clones OKNL13 and OKNL22 did not produce 12-methylgymnodimine, or only very low amounts that fell below the limit of detection (<0.1 pg cell⁻¹). We furthermore observed differences between clones with the lowest and highest detected toxin quota of about 5-fold for PSP toxins, 4-fold for SPX and 25-fold for gymnodimines. Thus, our results demonstrate a high intraspecific variability in toxin quota, which is in line with earlier reports on *Alexandrium* sp. populations (Gribble et al., 2005; Tillmann et al., 2009; Alpermann et al., 2010; Tillmann et al., 2014).
This variability cannot be explained by differences in growth conditions or growth phase, as all cultures were grown under the same conditions and harvested during the mid-exponential growth phase. The observed variability in toxin quota thus seems the result of genetic differences between clones.

In addition to the production of PSP toxins, spirolides, and gymnodimines, all isolates produced allelochemicals with the capacity to lyse cells of the target species Rhodomonas salina. Lytic activity of extracellular secondary metabolites is rather widespread in the genus Alexandrium (Tillmann and John, 2002) and has been shown for various Alexandrium ostenfeldii clones (Hansen et al., 1992; Tillmann et al., 2007; Tomas et al., 2012; Hakanen et al., 2014; Tillmann et al., 2014). Our correlation analyses confirm earlier comparative approaches that have shown that lytic activity is unrelated to PSP toxin (Tillmann and John, 2002) and spirolide content (Tillmann et al., 2007), and also a comparison of gymnodimine cell quota and lytic activity gave no obvious evidence that gymnodimines might contribute to the observed cell lysis of R. salina.

With EC50 values from 0.18 to 0.5 × 10³ cells mL⁻¹ the Dutch population seem to possess a comparable lytic activity as other Alexandrium ostenfeldii isolates of other phylogenetic groups. Their EC50 values have been shown to range from 0.2 to 1.9 × 10³ cells mL⁻¹ (Tillmann et al., 2007; Hakanen et al., 2014)
suggested comparable lytic activities within the species *A. ostenfeldii*. EC50 values for hemolysis of *A. ostenfeldii* Group I from the USA coast given by Tomas et al. (2012) and Tatters et al. (2012) for the same isolate (14 × 10⁸ and 7 × 10⁹ cells, respectively; values given as “cells” and not as a concentration) are more difficult to compare and refer to cell extracts. By using an erythrocyte lysis assay, Tomas et al. (2012) found only limited lysis in cell free supernatant of the *A. ostenfeldii* clones from coastal North Carolina, whereas particularly cell pellets had the highest lytic activity. The authors thus concluded that the lytic agents were most likely intracellular or membrane bound (Tomas et al., 2012). However, our data clearly show that lytic activity of cellular extracts was, with a mean factor of 12 (range: 4–30), remarkably lower compared to the lytic activity of whole cell cultures. The observed 30% loss of lytic activity when removing the cells might indicate a continuous release of lytic compounds during the bioassay incubation, but might also be due to loss of lytic compounds during the centrifugation step, as such compounds have been shown to adsorb to plastic material (Ma et al., 2009, 2010). In any case, our data show that the majority of lytic activity of *A. ostenfeldii* is excreted and acting extracellular, underlining a rather unspecific role in grazer impairment and competition (Cembella, 2003; Legrand et al., 2003). Nevertheless, lytic compounds produced by *A. ostenfeldii* may also be involved in cell-to-cell interactions, e.g. in prey capture and resource acquisition via mixotrophy.

Whole cell lytic activity varied roughly two-fold among clones. The intraspecific variability in lytic activity thus seems to be smaller as compared to multi-clone comparisons of *Alexandrium tamarense* (Alpermann et al., 2010) and *Alexandrium ostenfeldii* (Haken et al., 2014; Tillmann et al., 2014), and might indicate that high lytic activity was a trait positively selected for in the sampled bloom population. During the bloom period in the Ouwerkerkse Kreek, cell concentrations of *A. ostenfeldii* were well above 1000 cell mL⁻¹ (Fig. 1C) and thus far above the concentration that was shown to cause significant cell lysis of *Rhodomonas salina* in our bioassays. Although the role of lytic compounds for bloom initiation, at much lower cell concentrations, is more difficult to evaluate, deleterious effects of lytic extracellular compounds on both competitors and grazers likely play an important role in the success of *Alexandrium* blooms (John et al., 2015).

From a human health perspective, the toxicity of an *Alexandrium* bloom is determined by the population densities, and the amounts and relative toxicities of toxins that are produced. The *Alexandrium ostenfeldii* population in the Ouwerkerkse Kreek reached very high cell densities of up to 4500 cells mL⁻¹ (Fig. 1C), and will be of particular risk if dominated by the clones with high cellular toxin quota. However, the toxin composition in all clones consisted predominantly of the least toxic C1/C2. These C-toxins are hardly found in molluscs as they are readily desulphonated by enzymatic activity (Krock et al., 2007). This will convert C-toxins into the more toxic GTX which may potentially lead to high toxicities of exposed shellfish. The population of *A. ostenfeldii* for Ouwerkerkse Kreek produces two different toxin classes at the same time, and interactive effects among both toxin classes cannot be ruled out as, to the best of our knowledge, no studies have been conducted to investigate synergistic effects between different classes of neurotoxins, such as PSP toxins and cyclic imine toxins. Risks of blooms may furthermore be affected by environmental driven changes in the cellular toxin quota, for instance induced by changes in nutrient availabilities, pCO2 and temperature (Krempp et al., 2012; Tatters et al., 2013; Van de Waal et al., 2013, 2014).

In conclusion, our study shows that the dense bloom in The Netherlands was formed by Group 1 of the *Alexandrium ostenfeldii* species complex and thus highlights a new example of the bloom forming potency of this group around the world. The population contained a highly diverse cocktail of various neurotoxins, which underlines the threat of these blooms for important shellfish production areas in The Netherlands. Growth of isolates from this bloom under fixed laboratory settings was relatively slow, suggesting that allelopathy and presumably a reduced population loss by grazing, played a substantial role in bloom formation and persistence. Our data also demonstrate a high intra-specific variability with respect to toxin quota in the Dutch *A. ostenfeldii* population, as was earlier also reported for other *Alexandrium* populations (Alpermann et al., 2010). Such a high phenotypic variability may add to the success of genotypically highly diverse *A. ostenfeldii* blooms, and makes populations resilient to changes in environmental and climatic conditions.

**Acknowledgements**

The authors thank Pia Tahvanainen from the University of Helsinki for sharing *Alexandrium ostenfeldii* sequences prior to their availability in GenBank. We thank Annegret Müller for analyses of PSP toxins and Nancy Kühne for technical support with sequencing. We are grateful to Cynthia de Kruiff-Van der Voom from Waterschap Scheldestromen, and Suzanne Wieser, Dennis Waas, and Sven Teurlincx from the Netherlands Institute of Ecology (NIOO-KNAW) for their technical support. Kirsí Harju from the University of Helsinki is acknowledged for providing a standard solution of 12-methylglycinnimide.[SS]

**Appendix A. Supplementary data**

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.hal.2015.08.002.

**References**


