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Effects on inactivation of *Tetraselmis suecica* following treatment by KBAL: a UV-based ballast water treatment system with an in-line vacuum drop

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Abstract

The transfer of non-native, possibly invasive species in ship's ballast water is of global concern, and the International Maritime Organization and U.S. Coast Guard have adopted standards to minimize the environmental footprint caused by the maritime industry. In this study, seawater spiked with the phytoplankter *Tetraselmis suecica*, was treated with Knutsen Ballast Water Treatment Technology (KBAL), combining UV irradiation with an in-line vacuum drop. The test water was subsequently incubated in dark tanks, simulating what happens onboard a ship, where ballast water is treated at intake, stored in dark ballast tanks during the voyage, and then treated at discharge. Our results of the test water treated with KBAL and stored 5 days in the dark showed <10 viable *T. suecica* cells ml⁻¹ when assessing reproduction and >10 living cells ml⁻¹ when assessing metabolism. This highlights the challenge UV-based BWTS can encounter when meeting testing regimes assessing different characteristics of life. By comparing the effects caused by KBAL treatment with effects caused by UV irradiation only, we demonstrated that the pressure/vacuum technology seems to improve the disinfection effect. In addition, our investigations point out possible challenges with in situ conditions getting representative ballast water samples.

Keywords Ballast water · KBAL · Pressure/vacuum · UV · Tetraselmis suecica

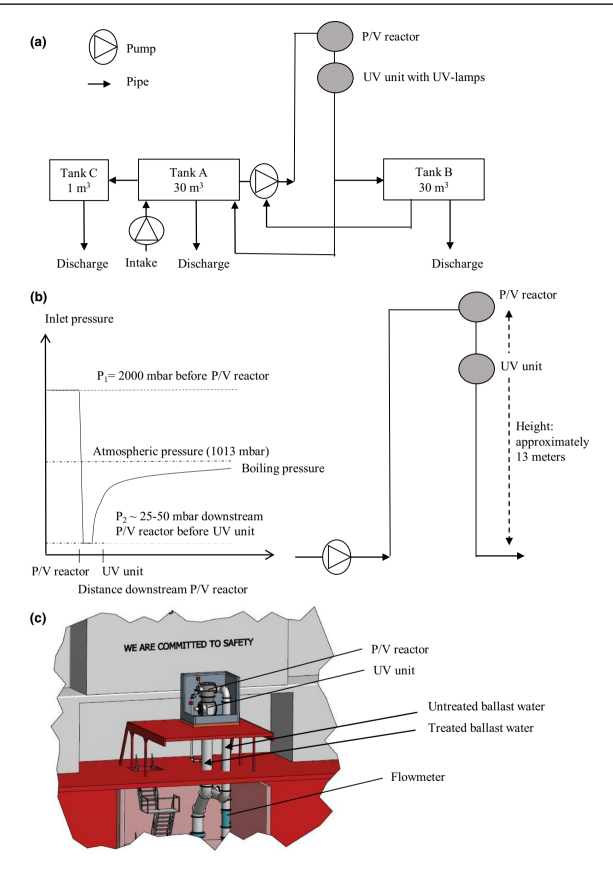
1 Introduction

Non-native, possibly invasive species in ship's ballast water can cause ecological changes and threat the biological diversity and the economy. The transfer of organisms is of global concern, and the International Maritime Organization (IMO) and U.S. Coast Guard (USCG) have adopted standards to minimize the environmental footprint caused by the maritime industry [1, 2]. In response to this, vendors have developed different ballast water treatment systems (BWTS). The commercialization rely on BWTS type approvals, which includes land-based tests and shipboard trials [3, 4]. One of these systems, the Knutsen Ballast Water Treatment Technology (KBAL) developed by Knutsen OAS Shipping AS, combines UV irradiation with an in-line vacuum drop. The developers aimed at avoiding chemicals to fulfill the G8-guidelines of the type approval process. It was also important for them to remove any filtration steps for water purification, due to maintenance concerns onboard ships. KBAL was type approved by IMO in 2016, and the commercialized KBAL system can treat ballast water at intake, during the voyage and/or at discharge. By December 2019, KBAL is the only BWTS on the marked that uses pressure/ vacuum (P/V) and UV irradiation as a treatment technology [5]. Most other BWTS combine UV irradiation with filtration [5]. Only two additional UV-based BWTS sidestep filtration; OceanDoctor BWTS produced by Jiujiang Precision Measuring Technology, instead uses an active substance (OH•) and is type approved according to the G9 guidelines; and BSKYTM BWMS produced by Wuxi Brightsky, which combines UV irradiation with a centrifugal separator [5].

Although not very common in BWTS, the use of P/V technology is well-known from other industry processes, for example as a physical pretreatment for extraction of microbial products [6]. The process is called high-pressure homogenization, where a liquid is transported with high pressure through a very narrow nozzle, causing a sudden drop in pressure causing cavitation, which induces destruction of the

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◄Fig. 1 Schematic drawings explaining a how the pumps, pipes and tanks are arranged in the KBAL test rig, where we have performed the experimental part, b the basic principle of the technology, and c how the technology is implemented onboard a ship. The ballast water is pumped into the P/V reactor that works in combination with a vertically drop line followed by a UV unit for irradiation. The water drop line below the P/V reactor will cause a subsequent reduction in pressure, so that the water reaches its boiling point. Figures are used with permission from the inventor of the technology in Knutsen OAS Shipping

cells leading to inactivation [7]. The high-pressure homogenization technology is easy to operate, cost and time effective, and can be used for large volumes. It also has the ability to reduce particle sizes [8], and it has been demonstrated that multiple rounds of high-pressure homogenization gives an additive effect on viability reduction [7].

UV irradiation is a more commonly used ballast water treatment technology, installed in various ship types [5, 9]. UV irradiation affects the genetic material in organisms, possibly inhibiting replication and transcription [10, 11]. The low-pressure (LP) and medium-pressure (MP) mercury UV lamps are emitting monochromatic UV irradiation at 254 nm, or a broader spectrum of wavelengths between 200 and 400 nm, respectively. Studies have shown that MP UV lamps have a higher disinfection capacity compared to LP UV lamps [12–14] and MP UV lamps have higher power output than LP UV lamps. Therefore, the numbers of MP UV lamps needed in a UV-based BWTS is reduced compared to a system based on LP UV lamps. In addition, MP UV lamps are shorter and more compact, reducing the size of the UV reactor. When retrofitting ships with BWTS, ship owners and operators concern about power consumption and footprint parameters, and using MP UV lamps can be beneficial with respect to minimal footprint and low power consumption compared to LP UV lamps. UV treatment has many benefits; it is considered environmentally friendly, no neutralization of chemicals is needed, it has low risk to operators and requires little training and maintenance, and parameters such as pH and temperature does not influence the UV-lamp functionality [15, 16]. Furthermore, microorganisms can employ survival strategies, such as cyst formation, when exposed to periods of unfavorable conditions [9]. Cysts have been recognized in ships' ballast tanks [10], and cysts can be resistant to various disinfectants [11, 12]. However, UV irradiation has been shown to inhibit cyst germination of *Giardia* species and algae [13, 14, 17]. On the other hand, biotic and abiotic particles in seawater can affect the BWTS inactivation, by protecting and shielding microorganisms during UV treatment [18, 19]. In addition, fouling of the quartz sleeves housing the UV lamps may reduce its efficiency by absorbing radiation that would otherwise be available for disinfection [20-22].

In this study, *Tetraselmis suecica* was used as a challenge organism for validation of the KBAL system. *T. suecica* is a representative of the ≥ 10 to $< 50 \ \mu m$ size class of the D-2 discharge standard, and belongs to the prasinophytes [23]. Several studies have previously argued that *Tetraselmis* is well suited as a standard test organism for validation of UV-based systems, since they are more or at least as resistant to LP and MP UV treatments as other naturally occurring seawater organisms, and do not generate a positive or negative bias to any technology [24, 25]. We used a dual staining flow cytometry (FCM) protocol with 5-carboxyfluorescein diacetate acetoxymethyl ester (CFDA-AM) and SYTOX Blue [26], plate count analysis, and a most probable number (MPN) analysis to analyze the effect of the treatments. We aimed at answering the following questions:

- (1) Are the vitality and viability results from the KBAL test in accordance with the IMO and USCG regulations?
- (2) How does the KBAL technology affect the cells?

2 Materials and methods

2.1 Culture maintenance

The phytoplankter Tetraselmis suecica (K-0297) was obtained from the Scandinavian Culture Collection of Algae and Protozoa (University of Copenhagen, Denmark), and cultured in 36 g/kg artificial seawater (ASW) (Marine SeaSalt, Germany) supplemented with 0.12% Substral (The Scotts Company (Nordic) A/S, Denmark). The cultures were incubated at room temperature (23-24 °C) under fluorescent light (T5 (25 W), and T8 (18 W) Sylvanian Grolux fluorescent tubes) in two 100 dm³ glass aquariums with aeration, filled with \sim 70 dm³ algal culture each. The amount of motile cells were detected in a counting chamber in an Optiphot-2 microscope (Nikon, Japan) by subtracting the number of non-motile cells from the number of total cells in an equivalent heat-killed sample. In addition, the quantification and viability were tested by growth on 1% BactoTM Agar (Difco Laboratories, Becton, Dickinson and Company, France) in artificial seawater with salinity of 24 ppt (which is normal along the Norwegian coast and fjords) supplemented with 0.12% Substral. A minimum of 1000 motile cells ml⁻¹ were added during tests in the KBAL test rig.

2.2 Treatment at the KBAL test rig

A 50 m³/h KBAL test rig is located in Haugesund, Norway. In the KBAL treatment system (Fig. 1a), water is pumped into a P/V reactor (Knutsen OAS Shipping AS, Norway). The water drop line below the P/V reactor (approximately 13 m) will cause a subsequent reduction in pressure, so that the water reaches its boiling point (Fig. 1b). This almost immediate pressure to vacuum process will have a negative effect on microorganism. In addition, the vacuum process will enhance inactivation by the UV irradiation, by ensuring a more evenly distribution of particles in the water stream and thus reducing the shadowing effects. The UV unit has 4 MP UV lamps (1.1 kW, polychromatic mercury lamp) (BestUV, The Netherlands) (Fig. 1a). A typical ship installation is illustrated in Fig. 1c. The test rig consists of two 30 m³ tanks, one for test water (A), and one for storage of the water after treatment (B) (Fig. 1a). In addition, there was a 1 m³ tank for seawater (C) (Fig. 1a). Before this test, the tanks were washed with a high-pressure hose and the system was flushed with municipal drinking water.

Tank A was filled with approximately 20 m³ seawater obtained from 1 to 2 m depth in Smedasundet, Haugesund, Norway, on the 2. November 2016. The temperature of the seawater was 11 °C at the time. At the same time, 400 g Kaolin (Sigma-Aldrich, product number 18672, CAS number 1332-58-7) and 430 g Maizena (Unilever Norge AS, Knorr, product number 2041499) were added, as well as T. suecica culture to a final concentration of minimum 1000 motile cells ml⁻¹ (test water). To ensure mixing, three extended outboard electric motors (Biltema, product number 25-235) (propeller: two blades, 240 mm in diameter) were used at maximum speed (corresponding to a 15.4 kg thrust) for about 30 min. During the first treatment, the test water was pumped with a flow of 50 m³ h⁻¹. The theoretical UV dose was set to 100 mJ cm⁻² for UV transmittance (UVT) at 60%, though the UVT measured by a UVT10-meter (BestUV, The Netherlands) was 86%. During treatment, the pressure was set to 2000 mbar upstream the P/V reactor and measured to 25-50 mbar downstream the reactor before the UV chamber. The treated test water was stored in tank B for 5 days, and subsequently mixed with the outboard electric motors for about 30 min at maximum speed, followed by a second treatment on the 7. November 2016 with the same conditions as during the first treatment. Untreated test water was stored in tank C for 5 days, and prior to sampling at day 5, the water in this tank was mixed with a paddle.

Samples were collected as follows:

- (1) Seawater.
- (2) Test water (seawater added *T. suecica*, Kaolin and Maizena).
- (3) Test water treated once.
- (4) Untreated test water and subsequently 5 day holding time.
- (5) Test water treated once and subsequently 5 day holding time.
- (6) Test water treated twice, including 5 day holding time between the treatments.

Table 1 summarizes how the seawater and test water was stored and/or treated.

Three parallel bottles of approximately 1 l were collected for each sample, except for sample number 5, where only one bottle was collected. The bottles were stored in the lab at 15 °C in the dark after sampling. Notice that the 5 day holding time for sample 4, 5 and 6 were in dark outdoor tanks of the KBAL test rig, further incubation was at 15 °C in the lab. The air temperature these 5 days of outdoor storage varied between -0.7 °C and 5.1 °C.

2.3 Ballast water analysis

Prior to analysis with the attune acoustic focusing cytometer (Thermo Fisher Scientific), 30 ml of the samples were filtered through a 100 µm nylon filter (BD Falcon Cell strainer 100 µm nylon, ref. 352360, BD, USA) to prevent large organisms and/or particles from clogging the tubing of the FCM instrument. All samples were dual stained with the esterase substrate CFDA-AM (C1354, Thermo Fisher Scientific, USA) and SYTOX Blue Dead Cell Stain (S34857, Thermo Fisher Scientific, USA), as described previously [26]. 4 ml of each sample was analyzed at a flow rate of 1000 μ l min⁻¹ and the sample dispensed options were set at standard collection mode. The trigger was set at red fluorescence (BL3 detector) and the threshold at 10,000. Samples 1, 2 and 3 were analyzed at day 0 (2 h after sampling), 1, 2, 5, 6, 7, 8, 9, and 12, whereas samples 4, 5, and 6 were analyzed at days 5, 6, 7, 8, 9 and 12.

To determine the numbers of viable cells, plate count analysis was performed and the cells were cultured on 1%

Table 1	Summarizes how the			
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stored a	nd/or treated			

Sample Content number		KBAL treat- ments	Holding time in KBAL tanks	Number of replicates	
1	Sea water	0	0 days	3	
2	Test water	0	0 days	3	
3	Test water	1	0 days	3	
4	Test water	0	5 days in tank C (1 m ³)	3	
5	Test water	1	5 days in tank B (30 m ³)	1	
6	Test water	2	5 days in tank B (between treatments)	3	

BactoTM Agar (Difco Laboratories, Becton, Dickinson and Company, France) in artificial seawater with salinity of 24 ppt supplemented with 0.12% Substral. The agar plates were incubated in light at 36 W m⁻² at 15 °C for 14 days. The number of colony forming units (cfu) for each plate was determined, and the mean values of the three replicates were calculated (except for sample 5, where only one sample was analyzed). In addition, MPN analysis was performed as described previously [27]; however, continuous light was used during incubation. Positive (visible by eye) growth were scored against an MPN table for a three-replicate design [28].

3 Results

3.1 FCM results of dual stained Tetraselmis suecica

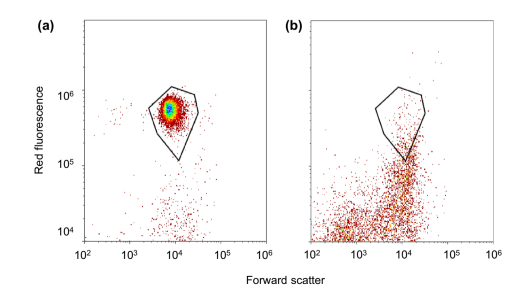
The *T. suecica* cells passed the laser light sources in the FCM instrument, one cell at a time. FCM signals are here presented as dot plots (Figs. 2, 3 and 5), where each dot represents one cell. Figure 2 shows forward scatter (FSC) intensity on the *x*-axis. FSC intensity is proportional to the diameter of the cell due to light diffraction around the cell. The *y*-axis shows the intensity of red autofluorescence. Since phytoplankton contain chlorophyll, the cells emit red autofluorescence when excited with a blue (488 nm) laser. A gate (black frame) defines the *T. suecica* population used for further vitality analysis. Since red autofluorescence from *T. suecica* cells was slightly reduced during the incubation period, the gate was designed to include these cells (Fig. 2a). At the same time, the gate was contracted to exclude other marine organisms and/or particles in the seawater (Fig. 2b).

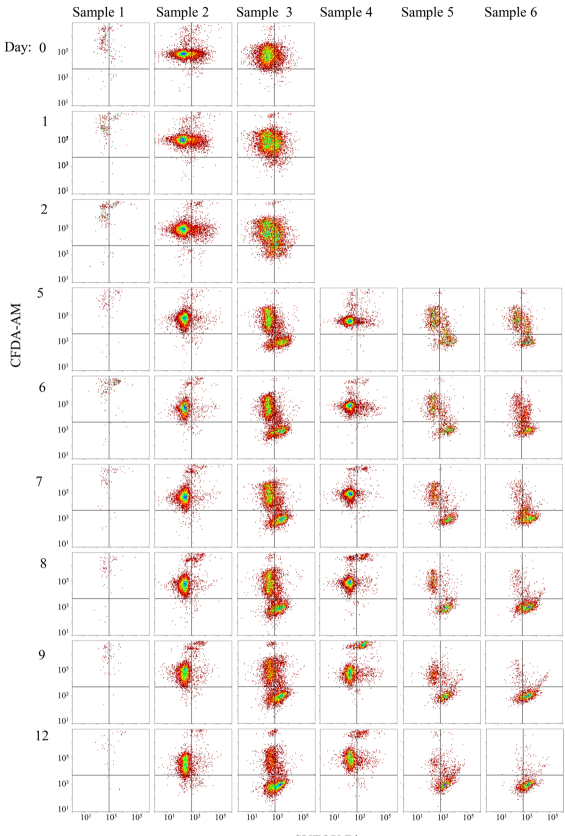
The cells inside the gate (Fig. 2a) was subsequently presented in a dot plot with blue fluorescence on the x-axis and green fluorescence on the *y*-axis (Fig. 3), displaying membrane permeability and esterase activity, respectively. These dot plots were separated into 4 quadrants (Q) by a vertical and a horizontal line based on the green and blue fluorescence intensity from the cells, as described previously [26]. The quadrants reflect the physiological cellular characteristics of esterase activity and membrane permeability, and thus vitality. The cells in Q1 are live (high esterase activity and low membrane permeability/intact membrane), severely damaged in Q2 (high esterase activity and high membrane permeability), and dead in Q3 and Q4 (low esterase activity with either low (Q3) or high (Q4) membrane permeability). The low blue fluorescence in Q3 can be explained by degradation of DNA/RNA resulting in fewer or no binding sites for the SYTOX Blue stain [29].

As expected, the FCM analysis detected organisms with esterase activity (Q1), 44 ± 15 org. ml⁻¹, in sample 1 (seawater) (Fig. 3a). These signals were most likely not caused by *T. suecica* cells, since cultivation methods did not show growth of this phytoplankter (Tables 2 and 3). The numbers of signals decreased during incubation.

For sample 2 (untreated test water, incubated in the lab), most cells were live (Q1) during the entire incubation period (Fig. 3). An FCM population (= single cells grouped together based on their fluorescence signals) appeared in Q2 (severely damaged cells) at days 0, 1 and 2; however, this FCM population disappeared at day 5 and was absent during the rest of the analysis period. FCM results for sample 2 and sample 4 (untreated test water, stored in 1 m³ outdoor KBAL tank for 5 days) (Fig. 3) match perfectly from day 5, where most cells were detected as live (Q1) during the entire storage period. The numbers of live cells in sample 2 and 4 varied for the three replicates at some of the analysis days, as seen in the standard deviations (SD) for the three replicates analyzed (Fig. 4).

Fig. 2 FCM dot plots of the gate (black frame) used to define the *Tetraselmis suecica* population used for vitality analysis (Fig. 3). **a** Analysis of the lab-culture before added to the KBAL test rig, 10,000 cells were analyzed. **b** Analysis of 1 ml seawater sample (dual stained) shows that most other organisms and/or particles were excluded from the gate





SYTOX Blue

◄Fig. 3 FCM dot plots of *Tetraselmis suecica* stained with SYTOX Blue (*x*-axis) and CFDA-AM (*y*-axis). The vertical columns show results for sample 1–6. For an overview of the various samples, see Table 1. The horizontal rows show the number of days of incubation. Notice that samples 4–6 were analyzed only from day 5, due to the holding time in outdoor KBAL tanks prior to sampling. Intersections in the dot plots separate signals into 4 quadrants (Q1–Q4)

The FCM signals for sample 3 (test water treated once, incubated in the lab) followed a pattern as previously described [26]. The cells were in Q1 (live) at day 0 with some cells in Q2 (severely damaged), and the population in Q2 increased during days 1 and 2 (Fig. 3). When comparing sample 3 with sample 5 (test water treated once, stored in 30 m³ outdoor KBAL tank for 5 days) (Fig. 3), the results from days 5-12 show the same reduction of cells in Q1 (live), an increasing population of damaged cell in Q2, and also an increase of dead cells in Q3/Q4 over time. Our results show that there were fewer cells in the sample with 5 day holding time in the outdoor KBAL tank (sample 5), only ~ 50 cells ml⁻¹ live cells at day 12, compared to ~ 400 cells ml⁻¹ for sample 3. When comparing the FCM results of sample 2 (untreated test water) with sample 3 (test water treated once) (Fig. 3), both incubated under similar conditions in the lab, less esterase activity and more variance in the fluorescence intensity was observed in sample 2 immediately after treatment.

The FCM signals for sample 6 (test water treated twice) also follow the Q1–Q2–Q3/Q4 pathway, similar to the samples treated once (samples 3 and 5) (Fig. 3). When comparing the second treatment with the first treatment (Fig. 3), both with a 5 day holding time in the outdoor KBAL tank, the cells died more rapidly when treated twice, meaning that cells were detected earlier in Q3/Q4. In addition, the numbers of cells in Q1 decreased more rapidly due to the second treatment (Fig. 4).

To get an indication on effects caused by the P/V technology, the results of *T. suecica* cells treated at the KBAL test rig were compared with previous results from *T. suecica* cells treated with UV irradiation only. Dot plot analyses from days 0, 1, 2 and 7 of seawater (sample 1) and test water treated once at the KBAL test rig (sample 3) were compared to a sample irradiated with an MP UV lamp at 100 mJ cm⁻¹ [26] (Fig. 5). The cells exposed to UV only, first lost their membrane permeability followed by a loss of esterase activity. On the other hand, cells treated with KBAL lost both membrane permeability and esterase activity simultaneously.

3.2 Plate count and MPN

Parallel to the FCM analysis, the samples were analyzed by the cultivation methods plate count (Table 2) and MPN (Table 3) to examine the numbers of viable cells (i.e., reproductive). No viable *T. suecica* cells were detected by plate count (Table 2) in the seawater (sample 1) during the analysis period. The plate count results showed a reduction of viable cells over time in samples 2–6. The numbers of viable cells for both the untreated test water samples (sample 2 and sample 4), was reduced during the analysis period to ~50% (> 500 org. ml⁻¹) at day 12. Sample 3 (test water treated once, incubated in the lab) showed 6 ± 5 viable cells ml⁻¹ after 2 days, and <3 viable cells ml⁻¹ 5–12 days after treatment. Sample 5 (test water treated once, stored in KBAL tank for 5 days) (one sample only), showed <10 viable cells ml⁻¹ when analyzed at days 5–12. Sample 6 (test water treated twice) showed <3 viable cells ml⁻¹ immediately after the second treatment and throughout the incubation period.

The MPN results (Table 3) also showed a reduction of viable cells over time in all samples analyzed. An exception was results for sample 2 (untreated test water) at day 7, as well as sample 4 (untreated test water with 5 day holding time in 1 m³ outdoor KBAL tank) at day 12, which both had higher numbers of viable cells compared to previous days. Since only one bottle was collected from the first treatment stored 5 days in the 1 m³ outdoor KBAL tank (sample 5), no MPN results were available for this sample.

4 Discussion

Both IMO and USCG have similar test water requirements when it comes to minimum concentration of organisms needed during land-based testing. For the ≥ 10 to $<50 \ \mu\text{m}$ size class, $\geq 1000 \ \text{org. ml}^{-1}$ is required at intake, $> 100 \ \text{org. ml}^{-1}$ in the control discharge, and $< 10 \ \text{org. ml}^{-1}$ in the treated discharge according to the D-2 discharge standard [1, 3, 4]. We added a minimum of 1000 motile cells ml⁻¹ to the test water, and during our 12 day analysis period, $> 500 \ T. \ suecica$ cells ml⁻¹ remained viable and living in the control samples, verifying that the test conditions were according to the requirements. Our land-based test mimics what happens onboard a ship: the ballast water is treated at intake (1st treatment), then stored in dark ballast tanks during the voyage (the 5 day holding time), and then treated again at discharge (2nd treatment).

A lesson learned from sampling is that the number of live *T. suecica* cells can easily be biased under variable conditions. For example, we experienced that cell concentration most likely was underestimated after the 5 day holding time, due to difficulties with mixing in the storage tanks. A previous study with known cell density of *T. suecica* culture in a storage tank, showed that the cell concentration was underestimated at discharge [30]. *Tetraselmis* species have three life stages, including flagellated, vegetative non-motile, and cyst stages [23]. *Tetraselmis* species in the cyst stage lose their flagella [23], and vegetative non-motile cells and

Table 2 Plate count results for samples 1–6 at different analysis days. The first day of analysis for samples 4–6 were at day 5, since these samples were stored in outdoor KBAL tanks for 5 days prior to sampling. Data are means (± 1 SD) of 3 replicates, except for sample number 5, where only 1 bottle was collected

Analysis day	Sample						
	1	2	3	4	5	6	
0	< 3	1058 ± 124	488±64				
1	< 3	988 ± 117	133 ± 5				
2	< 3	809 ± 158	6 ± 5				
5	< 3	621 ± 76	< 3	836 ± 55	< 10	< 3	
6	< 3	688 ± 252	< 3	839 ± 127	< 10	< 3	
7	< 3	788±37	< 3	697 ± 77	< 10	< 3	
8	< 3	709 ± 105	< 3	803 ± 50	< 10	< 3	
9	n.d.	591 ± 57	< 3	612 ± 5	< 10	< 3	
12	n.d.	536 ± 64	< 3	548 ± 50	< 10	< 3	

Table 3 MPN results for samples 1–4 and 6. 95% confidence intervals are in brackets. The first day of analysis for samples 4 and 6 were at day 5, since these samples were stored in outdoor KBAL tanks for

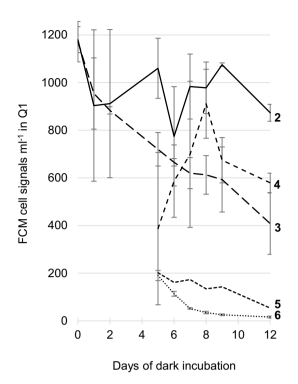
5 days prior to sampling. There are no MPN results for sample number 5, as only 1 bottle was collected

Analysis day	Samples					
	1	2	3	4	6	
1	< 3 (0–9.5)	930 (180–4200)	93 (18–420)		·	
2	< 3 (0–9.5)	930 (180-4200)	14 (36–420)			
5	0.3 (0.15–9.6)	430 (90–1800)	0.92 (0.14-3.8)	750 (170-2000)	0.36 (0.09-0.94)	
6	< 3 (0–9.5)	430 (90-1800)	2.3 (0.46–9.4)	750 (170-2000)	< 0.3 (0–9.5)	
7	< 3 (0–9.5)	2400 (420-20,000)	< 3 (0–9.5)	430 (90–1800)	< 0.3 (0–9.5)	
8	< 3 (0–9.5)	750 (170-2000)	0.36 (0.09-0.94)	930 (180-4200)	0.92 (0.14-3.8)	
9	< 3 (0–9.5)	240 (42-2000)	< 3 (0–9.5)	430 (90–1800)	< 0.3 (0–9.5)	
12	< 3 (0–9.5)	750 (170-2000)	< 3 (0–9.5)	4600 (900-20,000)	< 0.3 (0–9.5)	

cysts formed in the dark outdoor tanks might settle down in tanks during the holding time. In addition, the exposure to UV irradiation can influence the swimming capacity of algae [31]. The numbers of live cells also varied between replicates in some of the samples, which can be explained by patchiness of organisms in the tanks at sampling. Tests have showed that the concentration of organisms in different size groups can vary in different areas inside the tanks, with high concentration of large organisms and low concentration of small organisms, and vice versa [32], explained by predator–prey interactions [33]. This highlights possible challenges with in situ conditions, and the importance of getting representative ballast water samples.

Results from all analysis methods to assess the effectiveness of KBAL, showed a reduction in the concentration of live and viable *T. suecica* cells after treatment. This is consistent with previous in vitro studies of *T. suecica* irradiated with an MP UV lamp [26, 27, 34, 35], and also other UV-irradiated microalgae [36, 37]. The cultivation based methods (plate count and MPN) revealed similar results, with a decrease in the numbers of viable cells during dark incubation (except for two outstanding high MPN results for the untreated control sample). The results show that there were < 10 viable *T. suecica* cell ml⁻¹ in the test water after the first treatment and the 5 day holding time. Consequently, the results confirm compliance with the IMOs discharge standard assessing reproduction already at this stage, and additional effects caused by the second treatment was, therefore, not possible to detect. However, the same samples did not meet the discharge standards of IMO and the USCG when assessing metabolism, not even after the second treatment. The different results for metabolic (living) and reproductive (viable) *T. suecica* cells after KBAL treatments, clearly highlights a challenge UV-based BWTS can encounter when meeting different testing regimes, which previously has been highlighted by several authors [38, 39].

Our results show that the number of live and viable cells were similar for the untreated control sample (sample 4). Some of the untreated cells appeared as damaged (Q2) in the start of the experiment (days 0-2); however, they were detected as live (Q1) later in the incubation period. Physical or chemical stress applied to cells can cause a temporary



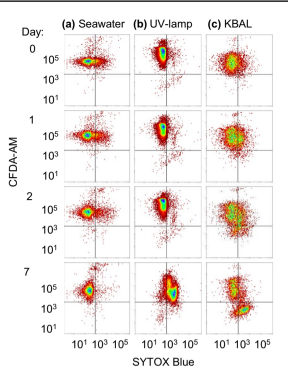


Fig. 4 Numbers of FCM cell signals in Q1 (live) during incubation for samples 2–6 (see Table 1). Sample 1, seawater, was not included in the figure, as cultivation methods did not show growth of *T. suecica.* The first day of analysis for samples 4–6 were at day 5, since these samples were stored in outdoor KBAL tanks for 5 days prior to sampling. Error bars indicate ± 1 SD

membrane permeability and recovery after a few days [40–42]. Even though our *T. suecica* cells were cultured under controllable laboratory conditions, they were diluted in natural seawater to obtain test water quality, which may have caused stress to the cells. The temperature of the seawater was lower compared to the laboratory conditions; however, many marine phytoplankton are able to tolerate high daily and seasonal temperature variations and growth can vary over a range of temperatures [6, 7]. For samples treated once with KBAL (sample 3 and 5), a decrease in esterase activity was seen immediately after the treatment and during the dark hold period, but the number of live cells were greater compared to the number of viable cells. This is consistent with previous studies of UV-irradiated T. suecica [27, 34, 35], but also to bacteria, and yeast [42]. For samples irradiated at low UV doses, no immediate shutdown of the cellular functions occur even when the reproductive capacity is lost (i.e., live but non-viable cells) [29, 38, 42, 43]. Non-viable cells may suffer from UV-induced DNA damages, which can be counteracted by DNA repair mechanisms [44]. Photorepair have been observed in *T. suecica* cells [45], but dark repair is not reported for T. suecica [46]. In addition, seawater turbidity can decrease the effectiveness of UV

Fig. 5 FCM dot plots of *Tetraselmis suecica* stained with SYTOX Blue (*x*-axis) and CFDA-AM (*y*-axis). The vertical columns show results for **a** seawater, **b** *T*. *suecica* cells irradiated with an MP UV lamp at 100 mJ cm⁻¹ [26], and **c** test water treated once at KBAL. The horizontal rows show the number of days of incubation

inactivation [46], and seawater particles and particles added to the seawater in the KBAL tank may have shielded some organisms from UV exposure. Dark incubated cells may also have low metabolic activity level [47, 48], and particle shielding during UV irradiation and low metabolic activity may have caused a delayed/no effect on the esterase activity.

Several studies have examined the use of UV irradiation in ballast water treatment, but knowledge about P/V technology used for ballast water purification is scarce. We, therefore, compared results of T. suecica cells treated at the KBAL test rig, with previous results from T. suecica cells treated with UV irradiation only. This comparison can give indications of effects caused by the P/V technology, even though results from tests with P/V treatment alone is lacking. Comparable UV doses were used for the KBAL testrig experiment and in UV only experiments (100 mJ cm^{-2}), and the T. suecica population were followed over time after treatment. The main observation is that the P/V technology seems to improve the disinfection effect, since T. suecica cells die faster after treatment with KBAL compared to UV irradiation alone (Fig. 5). This can be either an additive or a synergistic effect. The cells exposed to UV only, first lost their membrane permeability followed by a loss of esterase activity. On the other hand, cells treated with P/V and UV

irradiation, simultaneously lost both membrane permeability and esterase activity.

For future research, it would be interesting to study the effects caused by the P/V treatment on turbidity. It will also be important to know more details on how the P/V technology effects a broader range of organisms, since the focus here have been on vitality and viability of the indicator organism *T. suecica* only.

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