Temporal changes in phytoplankton biomass and cellular properties; implications for the IMO Ballast Water Convention

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Abstract

At two locations, coastal waters of the Wadden Sea, the Netherlands and at the station L4 (Western Channel Observatory) in the English Channel, UK, the temporal size class distribution of the phytoplankton community was investigated with respect to the size classes identified by the International Maritime Organization’s Ballast Water Management Convention. As part of this Convention, allowable discharge concentrations of organisms within classes were defined, with the lower size range (10-50 µm) consisting mainly of phytoplankton.

Traditional size fractionation methods that use nylon mesh filtration (10 µm mesh) showed considerable size bias. On average 23.1% of the larger than 10 µm cells were still present in the < 10 µm filtrate but 21.8% of the smaller sized cells were also retained on the mesh. In particular the latter would result in an overestimate of the number of cells per mL by as much as a factor of 5.4.

Flow cytometry was applied to give the precise size classifications of each cell. Temporal measurements, covering an annual cycle, indicated that at both test sites the phytoplankton in the size range 2 to 50 µm was dominated by the smaller sized phytoplankton (<10 µm). In terms of number of cells that fit the >10-≤50 µm size class these were on average only 3.6% and 2% in the Wadden Sea and the L4 sampling site, respectively. In terms of chlorophyll biomass they represent 28.7% and 12%, respectively. This was mainly caused by the cellular increase in chlorophyll concentration which increases in proportion to increasing cell size. In contrast, the mesh filtration method resulted in much higher chlorophyll values for the 10-50 µm size range; 53.7% in the Wadden Sea and 38% at station L4. This overestimation appears
to be caused by cells in 6-10 \( \mu \text{m} \) size range being retained on the mesh rather than passing through.

Present findings are relevant in the context of the size class distribution based on flow cytometry and semi-quantification using chlorophyll as proxy for cell density. - Keywords: Ballast Water Management Convention, Flow cytometry, Fluorometry, Chlorophyll, Phytoplankton

1. Introduction

With respect to particle size distribution in nature there are several universal laws based on allometric distributions. In many cases they are based on size and physiological or metabolic properties [1, 2]. In the oceanic environment, a commonly accepted rule is that the numbers of organisms per unit of volume tend to increase exponentially with decreasing size [3, 4, 5, 6]. Also, within a phyla or class, sizes can vary considerably. Phytoplankton varies by up to 6 orders of magnitude in size and up to 9 orders of magnitude in volume [7]. Associated with these differences maximum cell density [8] and various cellular properties also co-vary similarly to chlorophyll [9] and even the size of the genome [10].

This cell size to number relationship has recently received new interest as a result of the International Maritime Organization’s Ballast Water Management Convention (BWMC) [11]. In order to minimize the spread on non-indigenous organisms through ballast water, this Convention is limiting the number of living organisms in ships’ ballast water discharges. To this end, the Convention has defined specific size range distributions including a size range of \( >10 \) to \( \leq 50 \mu \text{m} \). In nature, this size range tends to be dominated by phytoplankton in terms of numbers while other organisms (e.g. microzooplankton) are far less abundant (less than 5%, unpublished results). However, this only represents a small component of the whole size range compared to the entire range of phytoplankton sizes present in marine or fresh waters. The
smallest known phytoplankton is only 0.7 µm (Prochlorococcus) [12] but other species can reach up to >2 cm in the case of colonies or chains [13]. In the latter case and according to the BWMC, the individual should be measured as it is the smallest unit able to reproduce [14]. The main reason for defining a regulation based on allowed concentrations of organisms in ships’ ballast discharge has been the fact that many of the toxic or otherwise harmful phytoplankton species are found within this size category. However, a significant number of phytoplankton species, including bloom forming harmful algae, are smaller than 10 µm (e.g. Phaeocystis spp., Pfiesteria spp. and Chrysochromulina spp.) [15]. Small sized species also present higher growth rates, which may be an advantage when colonizing a new environment [16, 17].

Phytoplankton (or specific sub-populations of) biomass and dynamics are generally studied as a whole so the establishment of a fixed size range imposes new criteria on studies. With the exception of phytoplankton blooms, the defined size range has a relatively low numerical abundance relative to smaller sized cells [18]. On the other hand, larger cells possess much higher concentrations of cellular chlorophyll, a cell component commonly used to estimate biomass or even cell density. As chlorophyll concentrations vary hugely with cell size, errors on cell density estimates based on chlorophyll concentrations will be significant. Even within a relatively small size range of 10-50 µm the diameter of the cell varies by a factor of 5 and, therefore, the volume of the cells (assuming they are spherical) will vary by a factor of 125.

The present study was conducted in order to examine the application of flow cytometry and fluorometry in characterizing natural phytoplankton communities with special attention to cell size. In addition, the annual variability of cellular properties like cell size and chlorophyll fluorescence combined with the actual size distribution of the cells was also investigated. The study covers a whole year at 2 different locations the Western Wadden Sea in the Netherlands and the Western English Channel in the UK.
2. Material and Methods

2.1. Area of study

2.1.1. Den Oever Harbour (the Netherlands)

Water samples (ca. 1 L) at the test site in Den Oever (Western Wadden Sea, the Netherlands, 52°56.07’N; 05°02.19’E – Fig.1) were collected weekly during a full year (2016). The harbour is in the inner part of the Wadden Sea, a shallow estuary repeatedly influenced by fresh water input from a nearby lake (Lake IJssel). During the year the temperature varied from 1 to 22 °C, and nutrients (PO₄, NO₃ and silicate) were depleted from May until the end of September. Whole samples and samples gently filtered over a 10 µm mesh filter were analysed within 30 min of collection.

2.1.2. L4 station (Western English Channel, UK)

Samples were collected from the coastal station, L4, of the Western Channel Observatory (WCO) in the English Channel, about 13 km off Plymouth, in waters of approximately 50 metres deep (coordinates 50°15.0’N; 04°13.0’W – Fig. 1) [19]. Relatively open sea characteristics may be found at the L4 site as well as features resulting from the influence of land with the inflow of water with higher concentrations of nutrients coming from rivers [20, 21, 22].

L4 samples are collected on a weekly basis, weather permitting, for ongoing research projects conducted by the Plymouth Marine Laboratory (PML) and the Marine Biological Association (MBA). These are some of the longest time-series in the world for phytoplankton and zooplankton. In the present study L4 samples were collected from June 2016 to May 2017.
Water samples from L4 were collected from the surface using a bucket and were analysed immediately or, in a few exceptional instances, samples were kept in a constant temperature room (held at L4 seawater temperature) and were analysed within 18 hours after collection.

Fig. 1: Study area showing Den Oever harbour in the Wadden Sea, The Netherlands (52°56.07’N; 05°02.19’E) and the L4 sampling site in the Western English Channel, UK (50°15.0’N; 04°13.0’W).

2.2. Methods

2.2.1. Flow cytometry

Flow cytometry (FCM) is widely applied in biological research including plant cells, yeast, phytoplankton bacteria and viruses [23, 24, 25]. In short, a set of bio-optical parameters is analysed from particles passing a narrowly focussed laser beam. While passing the laser a variety of cell properties related to size (Forward Light Scatter: FS) optical density (Side Scatter: SS) of auto or induced fluorescence are generated by each individual cell. This is done as peak or integrated values, varying with instrumentation. This information can afterwards be analysed semi-quantitatively and allow selective visual clustering of cells with matching values.

Flow cytometric analyses were conducted using a Beckman Coulter (BC) EPICS-XL-MCL in Den Oever and with a Bekton Dickinson (BD) FACSort™ at PML. 2 mL samples were analysed in triplicate, with single values or averages of the triplicates being used for further analysis according to Veldhuis & Kraay [26].

The settings of the instruments were adapted to display phytoplankton cells in the size range from 2 to 50 µm. The size was measured as the scattered light in the forward direction (FS), the measurement best related to size [25]. The red fluorescence from the phytoplankton
chlorophyll was measured after excitation with blue laser light (488 nm) as autofluorescence of the chlorophyll pigment (emission > 630 nm).

Standard spherical beads with known diameters (9.7 and 50 μm, Polysciences) were used as an internal standard for instrument calibration. These beads are uniform in size with known coefficients of variation (C.V. <2%) and measurements should possess the same spread for size and fluorescence.

Data analysis was based on clustering (sub) populations with identical size and chlorophyll fluorescence properties and considering the IMO size classifications, the fixed size defined implies that size rather than a specific population of cells was selected. Since phytoplankton populations usually have a broad size range, even within a species, the implication of this selection may be that only a part of the groups meets the size requirement. Next to cluster analysis resulting in grouped average values of cell size and chlorophyll fluorescence a frequency distribution of the cell size of the entire phytoplankton population was also made. This was done by reducing the standard 1024 channels, covering 4 decades of variation in size, into a 256 channel logarithmic mode, i.e. increasing bin size at the larger size ranges.

2.2.2. Fluorometry

Samples collected in Den Oever were analysed for phytoplankton biomass, in terms of chlorophyll fluorescence and photosynthetic efficiency, after dark adaptation using a WALZ-Water-PAM fluorometer, equipped with a blue excitation LED according to Schreiber [27]. The instrument was calibrated for background fluorescence using 0.2 μm filtered water.

This analysis provides an estimate of the chlorophyll-α concentration of the total and <10 μm phytoplankton (F₀ and F₀<10). The difference between both values was used to calculate the chlorophyll-α fluorescence of the >10 μm (F₀>10) fraction.
L4 samples were analysed using the Ballast Check 2 PAM fluorometer. This uses two measuring LEDs with multiple turnover to determine organisms’ photosynthetic activity. The equipment includes a filtration step (10 µm mesh filter) and based on the measured variable fluorescence it provides an estimated abundance for cells >10 µm based on the conversion of a fluorescence value divided by a set constant value of fluorescence per cell. To estimate the total number of cells we used a 0.2 µm filter. Therefore the calculated size fraction in this case is for cells smaller than 10 µm.

A dark adaptation period of at least 15 minutes was always observed before analyses.

2.2.3. Size range determination

Internal standard beads (9.7 µm, Polysciences) were used to distinguish between two size classes of phytoplankton (sub) populations combined with a series of size fractionation experiments. These were conducted to establish the relationship between the arbitrary estimates of size, determined as the forward light scatter (FS), and size based on selective filtration.

A suspension of mono algal cultures and samples collected from the field, the latter with clearly distinguishable subpopulations, were gently filtered over a series of filters ranging from 20 to 0.2 µm. The 20 and 10 µm filters were nylon mesh filters with nucleopore filters (8, 5, 3, 2, 1 and 0.6 µm) being used for the subsequent filtration steps. During the sequential filtration steps great care was taken that some sample fluid remained on top of the filter and that the filter was not run dry thereby avoiding damage to cells. Three to five replicates of samples were analysed and the number of cells passing through each filter were counted using flow cytometry. Using a logistic (sigmoidal) fit the size, as estimated spherical diameter (ESD), of the cells was determined as the number relating to 50% retention on the filter according to equation 1 (equation 1) using SigmaPlot (version 12.5).
Equation 1: \[ f = \frac{a}{1 + \left( \frac{x}{x_0} \right)^b} \]

- Where \( x_0 \) = infinitive pore size
- \( x \) = pore size of filter applied
- \( a \) and \( b \) computed constants
- \( f \) = fraction of cells passing filter

Using this sigmoidal curve fit for each phytoplankton population the average size, as an ESD, was determined using a level of 50% of the population present. All cultures and field samples used had a length to width ratio of a factor of less than 3.

Fig. 2: fraction of initial cell number of 4 different phytoplankton species remaining present in filtrate as a function of applied filter pore size. Lines are calculated fit of logistic function. Arrows are associated cell size based on 50% of cells present.

In total 21 samples, sampled throughout the year in Den Oever, were fully analysed using flow cytometry and the estimated spherical diameter of the phytoplankton subpopulation was compared with the corresponding forward light scatter signal, as a proxy for cell size (Fig. 3). No linear relationship was found but, based on the curve regression fit, the ESD of subpopulation or individual cells can be determined based on the FS measured.

Fig. 3: forward light scatter versus size fractionated estimated spherical cell diameter

3. Results

3.1. Temporal distribution
Figure 4a shows the annual distribution of phytoplankton numbers in Den Oever with a typical spring and autumn bloom and low cell density in the winter season. The phytoplankton community was throughout the year dominated by smaller sized (<10 µm) phytoplankton using the 9.7 µm reference beads as a selection criteria for size. In terms of cell density, the number of phytoplankton cells larger than 10 µm varied between 26 and 2662 cells per mL (annual average 982 cells/mL, table 1). Compared to the total number of phytoplankton cells measured, this size class was only a minor fraction of the total, ranging from 0.1 to 14 % (mean value of 3.6 %).

The L4 site (Figure 4b) showed a similar trend with a distinct spring/summer and autumn blooms and lower numbers during the winter (October to March). Previous studies described the spring and autumn blooms composed mainly by diatoms whilst dinoflagellates are dominant during the summer [28]. The total number of cells per mL found for the whole period was 12590 in average, however cells larger than 10 µm corresponded in average to only 201 cells/mL (CV%= ± 87, table 1).

Fig. 4: annual number of total phytoplankton and fraction <10 micron (bottom graph). Number and percentage of phytoplankton cells in fraction >10 micron (top graph): Den Oever (a) and L4 (b).

Cell density, cell size of each individual cell and the chlorophyll auto-fluorescence ($F_{0_{icm}}$) of each cell were measured concurrently. The collective values of these cellular $F_{0_{icm}}$ values also provide an estimate of chlorophyll biomass (Fig. 5a, Table 1). The percentage of chlorophyll associated with the larger cell sizes (> 10 µm) varied considerably throughout the year in Den Oever, ranging from 0.8 to 80% of the total, but the annual mean value of 28.7% was higher than the value based on cell number.
chlorophyll biomass results measured using flow cytometry for L4 samples also showed a
similar trend to the pattern found for number of cells (Fig. 5b). And, as experienced in Den
Oever, results from the fraction larger than 10 µm were higher because larger cells have
higher chlorophyll content, showing an average of 12% (CV% = ± 108, Table 1), with values
ranging from 3 to 30% of the total.

Fig. 5: annual flow cytometric integrated chlorophyll concentration of total phytoplankton and
fraction <10 µm (bottom graph). Integrated chlorophyll concentration and percentage of
chlorophyll in fraction >10 µm (top graph): Den Oever (a) and L4 (b).

The annual analysis of the cellular characteristics of the phytoplankton in Den Oever showed
alongside to a variation in terms of numbers also changes in the cellular properties of size (FS)
and chlorophyll autofluorescence (F0fcm) (Fig. 6). For the total phytoplankton community
these average values varied by as much as a factor of 4 for both size and chlorophyll
throughout the year. Using the conversion of equation 1 the corresponding average sizes
would range from 5 to 15 µm (Fig. 3).

Identical results were measured at station L4 where the minimum average value represented
c. one quarter of the average values found for cell size and chlorophyll content. While the
ratio between maximum and minimum single results varied by a factor of ca. 15 for both
cellular properties.

On a more detailed level, and based on a clearly visible subpopulation, flow cytometrically
derived values of size and chlorophyll also co-varied indicating a clear relationship between
size and chlorophyll content (Fig. 7). This relationship was found for all size classes covering
the entire size range of phytoplankton cells in both sampling sites.
Fig. 6: annual variation in cell size and cellular chlorophyll autofluorescence of total phytoplankton community (2 - 50 micron) and size class > 10 micron: Den Oever (a) and L4 (b).

Fig. 7: Covariation between cell size (measured as forward light scatter) and cellular chlorophyll autofluorescence of total phytoplankton community (2 - 50 μm) and different subpopulations 2 – 4 μm, 4 to 7 μm, 7 to 10 μm and > 10 μm (Den Oever data - a). Same covariation for the total number of cells (2 to 50 μm) and for organisms between 2 and 10 and from 10 to 50 μm (L4 data - b).

Table 1 near here

3.1. Detailed size classification

Because flow cytometry generates values of size and chlorophyll fluorescence data for each individual cell, it is possible to generate a complete frequency distribution of the size distribution of each sample analysed (Fig. 8).

This was done based on a logarithmic distribution of the bin-size varying from 2.7 FS units at the lower size range to 340 FS units per bin of the largest bin. Throughout the year the size distribution of the phytoplankton community remained rather constant despite changes in absolute numbers. Only during typical bloom events, in spring or autumn, a relative increase in certain size ranges (6 to 8 and 10 to 15 μm) was observed. These were usually related to the episodic occurrence of blooms of mono-specific phytoplankton species. On the basis of the annually averaged values, the highest numbers of phytoplankton fall within the FS size range of 10 to 200 μm, these values correspond with an ESD ranging from 2 to 20 μm (Equation 1) (Fig. 8a).

A frequency histogram of the observed FS values at station L4 is shown in Fig. 8b. At this station the values of FS for total phytoplankton varied from 7.2 to 112.3 and were
concentrated between 10 and 100 on a logarithmic distribution. Also from L4 data we can see a constant size distribution throughout the year regardless changes in absolute number.

Fig. 8: frequency distribution of cell size of each sampling day in Den Oever (average of 3 replicates, black lines) and annual average ± 1 sd (right scale) (a) and frequency distribution of cell size for total phytoplankton at station L4 considering all samples/replicates in the period (N=114, FS= 28.2 CV%= ± 63) (b).

3.1.1. Size fractionation 10 μm mesh

As mentioned, the IMO’s ballast water performance standard (Regulation D-2) provided in the BWM Convention is defined on a size class basis. Having this in mind, the following section describes the results of the samples filtered over 10 μm mesh, the commonly applied method to separate size classes, compared to entire sample. Measurements of total phytoplankton biomass (PAM fluorescence, F₀), cell density and FCM-integrated chlorophyll values based on the mesh separation method resulted in distinct differences when compared to those based on flow cytometric values for size (Figure 5, Table 2). On average, the values based on the filter screening were substantially higher for the >10 μm size fraction at both test sites.

Applying a standard fluorescent measurement showed that 53.7% of the total phytoplankton chlorophyll fluorescence (PAM-F₀) was associated with phytoplankton retained on the 10 μm mesh in Den Oever. Therefore the theoretical concentration of cells in the >10 μm size fraction would be 6148 cells/mL, or 22.7% of the total. This is 6.2 times higher than measured using flow cytometric size selection (982 cells/mL). The FCM-integrated chlorophyll measurements also showed that 47.3% of the chlorophyll was retained on the 10 μm mesh. This percentage is close to the value based on the bulk chlorophyll fluorescence (53.7%).
In addition to the whole water sample, a detailed flow cytometric analysis of size and cellular chlorophyll fluorescence was conducted on the fraction of phytoplankton passing the 10 µm mesh (Table 2). Analysis showed that both cells with a flow cytometric determined size of > 10 µm were passing the 10 µm mesh but also that smaller sized cells were retained on the filter.

On average 227 cells/mL were measured that were >10 µm on basis of their size (FS) in the 10 µm mesh filtered water samples. Compared to the total number of phytoplankton cells this was only 1.1% but as much as 23.1% of the number in the same size range of cells in the unfiltered sample.

Alternatively, an average of 5392 cells/mL were retained on the nylon mesh corresponding to 20.6% of the total phytoplankton number classified on basis of the FS-size <10 µm.

Applying the same procedure for the L4 data, resulting F₀ measurements from the BC 2 fluorometer showed 38% of the total phytoplankton associated with the fraction over 10 µm, which would mean an average of 1812 cells/mL. This is 9 times higher than the 201 cells/mL average number detected with FCM and 14.4% of the total cells in average.

Table 2 near here

The effect of the 10 µm mesh filtration on the size distribution of the phytoplankton was also analysed on the level of each individual cell for samples collected in Den Oever, similar to that shown in Fig. 8 for the unfiltered sample. For this analysis the frequency distribution of the cell density for the entire year was integrated instead of using the annual average (Fig. 9). The top graph of Fig. 9 shows the size distribution of a cell culture of *Tetraselmis* sp. (average cell diameter of 12 µm). The detailed cell size analysis showed that, as commonly observed for phytoplankton, the population of *Tetraselmis* was far from uniform in size distribution and
varied by as much as a factor 3. The size of 95% of the cells varied between values for FS of 80 and 240.

The cell size distribution of the 10 μm mesh filtered sample showed values matching those of the unfiltered sample in the lower range of cell sizes up to an FS value of 40. With increasing cell size, the discrepancy in numbers between total and mesh filtered increased even at values of FS below the value corresponding with a cell size of 10 μm. Above an FS value of 50 the numerical difference between the total and mesh filtered water declined again with increasing cell size. In terms of percentage of reduction in cell density due to the filtration a different trend was observed. At the lowest size ranges the difference was in the order of a few percent increasing to as much as 35% at the value of FS corresponding to a size of 10 μm. At the higher cell sizes this percentage increased rapidly. A near 100% reduction was only measured when the value of FS was higher than a value of 500.

Fig. 9: Flow cytometric data of frequency distribution of phytoplankton cell size (2 - > 50 μm) of total phytoplankton (top graph), cells passing 10 μm mesh, numerical difference and percentage of difference between both data sets. Values are based on annually integrated numbers (below). Top graph is total phytoplankton and *Tetraselmis* sp. as a reference phytoplankton species. Dashed line indicates FS value corresponding with ESD of 10 μm.

**4. Discussion**

This study shows that flow cytometry is a useful, fast, accurate and reproducible tool for the size analysis of phytoplankton cells. Size distribution can be done based on the whole community, subpopulation and even as in this case down to the level of the individual cell.
Although the data for size are usually based on arbitrary units of forward light scatter [10] [25] they can be converted into more realistic values of cell size using simple conversion factors. Jennings & Parslow [29] defined equivalent spherical diameter as the diameter of a sphere that would perform in the same way as the non-spherical particles presented in the sample; the authors highlight that the resultant dimension is always less than the true major dimension though. These conversions not only rely on the shape/dimensions and their conversion into a forward light scattering signal but also on instrumental differences in how the particle’s cross-sectional area is determined [30] therefore indicating the need of proper calibration. The measurements also indicate that even within a single phytoplankton species (e.g. Tetraselmis) the variation in size can be considerable, as microscopic analysis confirmed. For many species of phytoplankton analysed the coefficient of variation of size ranged typically between 40 and 60%. The variability in the dimension of size is therefore natural but also explains the overlap when multiple species are present as in the current samples.

Infrequently and during blooms of selective species higher numbers of certain size classes are seen e.g. Phaeocystis or diatoms in the spring in the Wadden Sea [31, 32]. On an annual time scale these blooms are of minor effect on the general pattern of size versus cell number distributions (Fig. 8). At station L4, Tarran & Bruun [33] described periods of higher abundance for pico- and nanoplankton during the spring / summer in the top 20 meters. The summer peak is probably encouraged by the summer thermocline breakdown allowing the mix of nutrients in the surface layers [34, 33]. Samples from 07 April, 2017 showed the first signs of spring bloom arriving earlier than in recent years, confirmed in subsequent sampling to be dominated by Guinardia delicatula (Dr. Claire Widdicombe - personal communication). L4 results in early May showed very low numbers of cells per mL, probably as a result of being deprived of the nutrients that were consumed by the phytoplankton during the early spring
bloom (L4 buoy data – PML Western Channel Observatory Blog - http://www.westernchannelobservatory.org.uk/blog/?p=870). (Fig. 4 and 5).

These data also confirm the general trend that phytoplankton populations show an inverse relationship between numerical abundance and cell size as has previously been reported in the literature [35, 36, 37].

On an annual basis, smaller sized cells were dominant at all periods not only at the more off-shore station (of the two in this study) L4 (13 Km off Plymouth with influence of the North Atlantic Ocean) but also nearshore in an estuary (Table 1). This dominance is not only restricted to certain periods like the summer when nutrients are normally low [22] but throughout the whole year. Applying the size classification based on the conversion of flow cytometric derived values of size into ESD shows that on average, the relative contribution of cells with a dimension of 10 μm or lager to the total cell number was low, only 3.6% or 2% for the Wadden Sea and station L4, respectively. Even when taking into account the higher chlorophyll a concentration of larger sized cells only 28.7% (Wadden Sea) or 12% (L4) of the chlorophyll is associated with the larger size fraction.

Figure 8 shows that throughout the year and irrespective of the location, coastal or more open ocean, the flow cytometric determination of cell size results in a uniform and continuing pattern of size distribution of the phytoplankton covering the entire range from 2 to > 50 μm in diameter. As a result, classification of populations in terms of size classes will therefore be a rather arbitrary exercise. In this study up to 44% of the Tetraselmis population must be classified as cells with a diameter of less than 10 μm (Fig. 9). The rather strict definition used by the BWMC [11] of ‘minimum cell dimension’ would imply that even within a single species, individual cells would not meet the criteria and exact sizing of all cells would be required. While flow cytometry provides a full-scale analysis of cell size in a time span of
several minutes, more detailed microscopic analysis of a large number of cells would take many hours.

Our data also show that the commonly applied method of size selection by means of mesh filtration resulted in significantly different results. Reanalysis of the filtered fraction indicated that as much as 23.1% of the cells or 14.9% of the chlorophyll (F₀_{FCM}) of phytoplankton cells larger than 10 μm passed through the mesh filter. In contrast, 21.8% of the cells and 43.0% of the chlorophyll of phytoplankton cells with estimated cell size <10 μm were retained on the mesh filter. In particular, the bias towards smaller sized cells by the filter resulted in an overestimation of the actual numbers by as much as a factor of 5.4. Also in terms of chlorophyll biomass the difference between both size selection methods was considerable. The direct measurement of chlorophyll biomass, applying PAM-fluorescence analysis, resulted in 53.7% and 38% of the chlorophyll associated with the larger sized cells in Den Oever and L4 respectively. These values were comparable with the flow cytometric data of differences in chlorophyll (47.3% and 29% in Den Oever and L4 respectively). In reality the actual number of cells was much lower, on average only 982 per mL (Den Oever) and 201 per mL (L4); and therefore also their chlorophyll concentration (F₀_{FCM} of 28.7 and 12% in Den Oever and L4 respectively).

Finally, we return to the initial questioning on what would be the effect of cell size on the conversion factors used by fluorometers to convert fluorescence into cell density. In theory, any fluorescence parameter may be converted into number of cells/mL by means of an internal coefficient. However, variations in cells’ size and therefore in the content of chlorophyll within the cell can affect the fluorescence signal measured, which means that a simple conversion value based on chlorophyll fluorescence might not be completely reliable [10, 38, 39]. Another aspect to be considered is the device’s behaviour to a large number of
smaller cells (<10-μm), would their fluorescence signal influence fluorometer’s numeric results? Since many fluorometers used in the quantification of the IMO relevant size class of 10 – 50 μm include a filtration step, the bias due to smaller size cells retained on the filter may be significant.

On the other hand, based on an uniform distribution of size and numbers, an average cell size and accompanying chlorophyll fluorescence can be calculated. For the Den Oever test site the average cell size, as ESD, determined was 20.8 μm (CV ± 44%, ranging from 11.6 to 30.0 μm) and corresponding chlorophyll fluorescence of 329.4 F0_{FCM}/cell (CV ± 85%, ranging from 49.4 to 608.7 F0_{FCM}/cell). But even for the given coefficient of variation there would be a 12-fold variation in cellular chlorophyll, and with a fixed conversion factor an equal variation in corresponding number of cells.

5. Conclusion

This study attempts to add value to the discussion on the possible implications resulting from cellular properties and biomass changes on the IMO’s Ballast Water Convention implementation notably on the ballast water performance standard (Regulation D-2). Additionally, it intends to raise the issue of potential sources of error for further refinement of the instruments regarding a relatively unknown area where portable tools developed for verifying ships’ compliance to the BWMC may produce dubious / false results [40].

In an ocean threatened by increasing CO2 and many other natural and anthropogenic stressors, cell size composition will be affected and therefore the phytoplankton community structure [7]. This will pose additional challenges for indicative tools developed to measure abundance of photoautotrophic cells in the water. Accordingly, conversion factors based on photosynthetic activity will need to be robust enough to face the challenges of a changing ocean.
Sampling and analysis of ballast water samples is supposed to be a relatively rare procedure according to the tiered regulatory enforcement approach agreed at IMO. However, considering the challenges, there remains a feeling that there is not enough discussion and research to provide the needed confidence that is required of ballast water monitoring techniques.

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