Rapid detection method of ship ballast water biomass based on spectrophotometry

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Abstract—The invasive aquatic species carried by ship ballast water has caused serious environmental damage and great economic loss worldwide. The issue of ship ballast water discharge has received extensive attention in recent years. According to the IMO Ballast Water Management Convention, the vast majority of vessels must perform ballast water treatment that complies with a set of parameters according to regulation D-2. Thus, it is urgent to solve the problem of testing the quantity of organisms in ballast water. But few report has been made on the fast microbial counting method that can really support the D-2 standard in ship ballast water and the existing technology methods have obvious disadvantages. In the study, we reported a rapid detection method of ship ballast water biomass based on spectrophotometry. This method can be used for rapid and convenient detection of microalgae cell biomass of 10-50μm in ship ballast water, and has the advantages of portability, speed, simplicity and no need of fluorescence probe.

Index Terms—spectrophotometry; ballast water; long optical;microorganisms.

I. INTRODUCTION

Due to the rapid development of shipping industry in recent years, the invasive aquatic species carried by ship ballast water has caused serious environmental damage and great economic loss worldwide[1-3]. International Maritime Organization (IMO) Ballast water Management convention had been come into force on 8 September 2017. According to the IMO Ballast Water Management Convention, regulation D-2 of the Convention stipulates that ships meeting the requirements of the Convention must discharge:

- less than 10 viable organisms per cubic meter greater than or equal to 50 micrometers in minimum dimension.
- less than 10 viable organisms per millilitre less than 50 micrometers in minimum dimension and greater than or equal to 10 micrometers in minimum dimension.

The convention puts forward new requirements on the performance ability of maritime administrative bodies. The competent maritime authority is about to carry out PSC inspection on ships in port under this convention[4]. For shipowners, the D-2 standard also imposes new requirements on them. Before the international maritime organization marine environmental protection committee, new ships built on or after September 8, 2017 shall comply with D-2 standards from the date of delivery[5]-[6]. Thus, it is urgent to solve the problem of testing the quantity of organisms in ballast water.

But few report has been made on the fast microbial counting method that can really support the D-2 standard in ship ballast water. The method, fluorescent probe staining and microscopic examination, has been widely used presently. However, this method has the disadvantages of high lower limit, time consuming and high detection intensity. The detection result is questionable because a single fluorescence probe is not effective against all living organisms[7]. Thus, many researchers and companies have worked on increasing the number of fluorescent probes and used more efficient and reliable detection instruments, such as Flow Cytometry (FCM)[8]-[10] and Microfluidic Chip[11]-[12].But the current research can only be done in the laboratory, not in practice. In addition, the government regulatory authorities also lack of effective means of on-site monitoring of the discharge of ballast water from ships[13]-[15].

In the present paper, we reported a rapid detection method of ship ballast water biomass based on spectrophotometry. In this method, it is assumed that photosynthetic microalgae are the main living microorganisms in ship ballast water. The number of living microorganisms in ballast water can be estimated roughly by detecting chlorophyll content in ballast water. The device described below can be used for rapid and convenient detection of microalgae cell biomass of 10-50μm in ship ballast water. This method realizes the rapid detection of microalgae cells, and has the advantages of portability, speed, simplicity and no need of fluorescence probe.

II. METHOD AND PROCESS

The long path flow cell (LPC-500CM, optical path 500cm, Wavelength Range 280-730 nm) were purchased from Oceanoptics Co.Ltd. Acetone were purchased from Sinopharm Chemical Reagent Co. Ltd. Deionized water is used in the gas water backwashing device. The detection system is shown in figure 1.

The ballast water sample to be tested is introduced into the sampling chamber, and the sampling valve, the liquid discharging valve and the sampling pump are turned on. The sample is driven by the pump into the concentrator device (the volume of the concentrator device is 0.5 ml, and the filter at the lower end is provided with a pore size of 50 μm. The upper end outlet is provided with a filter with a pore size of 10 μm) which intercepts 10 -50 μm size of organisms, and the remaining liquid is discharged into the waste tank. The final concentration sample is displayed and controlled by a sampling flow meter to be 500 ml. Then, the flush valve is opened, and the gas-water backwash device backwashes the
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The computer connected to the spectrometer performs statistical analysis on the repeatedly measured data, which rejects the individual abnormal data automatically (refers to the measured value of the set of values that deviates from the average value by more than twice the standard deviation) and performs the mean calculation on the remaining data. The content of chlorophyll is calculated by the following calculation formula, and the corresponding biomass concentration in the initial ballast water is converted.

\[
\text{Chlorophyll } \alpha = \frac{(11.64 \times (D663-D750) - 2.16 \times (D645-750) + 0.10 \times (D630-750)) \times V}{V \cdot \delta}
\]

\(V\) —— the volume of the water sample (the volume of the water sample through the sampling flowmeter during the detection process) \(\text{(L)}\)  
\(D\) —— absorbance  
\(V1\) —— the volume after the constant volume of the extract (the actual volume of the concentrator) \(\text{(ml)}\)  
\(\delta\) — optical path of cuvette \(\text{(cm)}\)

The purpose of measuring the absorbance at a wavelength of 750 nm is to avoid interference of suspended matter. Generally, the wavelength used for measuring turbidity in water is 680 nm. To avoid the absorption value of chlorophyll \(\alpha\) at 680 nm, the wavelength of turbidity will be selected above 710 nm.

In the calculation formula, the absorbance values at 750 nm should be subtracted from the absorbance values at 750 nm to deduct the interference of suspended matter, and the values should be less than 5% of the absorbance value of the chlorophyll \(\alpha\) absorption peak \((D750 < 5\% D663; D750 < 5\% D645; D750 < 5\% D630)\). Assuming that the absorbance value of the sample at 663 nm is 0.03, the absorbance at 750 nm should not be greater than 0.0015, which requires high clarity of the test solution. Otherwise, it should be re-filtered and the concentration factor reduced (the concentration factor is reduced to 1/2).

If the detection result after reducing the concentration factor (the concentration factor is reduced to 1/2) is still that the absorbance value at 750 nm is not less than 5% of the absorbance value of chlorophyll \(\alpha\) absorption peak, the enhancement measure of adding cell lysate is adopted.

IV. SUMMARY

Based on the long-path flow cell and the method of spectrophotometric determination of chlorophyll \(\alpha\), we designed a method for detecting 10-50 \(\mu\)m microalgae cells in ship ballast water, and realized the rapid detection of microalgae cells in ship ballast water. Compared with the detection methods currently reported, this method has the advantages of simple device structure, convenient operation, efficient detection process, independent of fluorescent dyes, and no shaking of the ship. It can quickly determine the microalgae cell biomass in the sample and thus provide a means for maritime law enforcement to effectively and quickly detect ballast water. The new method proposed in this paper has the following technical characteristics:

1) Calculating the concentration of chlorophyll \(\alpha\) in ballast
water by spectrophotometry and calculation formula, and then converting the concentration of the microalgae cells biomass.

2) By increasing the optical path, the concentration factor required for the sample is reduced, thereby greatly reducing the sampling amount and saving the time required for concentration. (using a long optical path flow cell is to increase the optical path. The standard cell optical path is 1cm, and this method chooses to use a 500cm long path flow cell).

3) Multiple sets of absorbance data can be obtained by repeated measurement at higher frequencies. By statistical analysis of these data, the detection accuracy can be effectively improved and the invalid samples can be identified, and the reliability of the test results can be improved.

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