

Application Note

Total cell count (TCC) of bacteria in water



Introduction

The quantification of the bacterial amount is important to control the microbial ecology in different water samples. Sysmex Partec instruments use an improved and standardized application to determine bacterial counts in water: The CyStain™ BacCount reagents.

The old standard - Heterotrophic plate count

Worldwide, the bacterial quality control of water is performed with the 'heterotrophic plate count' (HPC). This method goes back to Robert Koch, who more than a century ago determined the microbiological status of drinking water by counting microbial colonies on agar plates.

Today it is well known that this routine cultivation technique has severe limitations:

 Not all microbes present in water samples grow and form colonies on solid cultivation media. These bacteria are e.g. the so-called viable but nonculturable (VBNC) bacteria.

- 2. The data obtained from colony counting for result interpretation is weak as it represents down to less than 0.1 % of the bacteria present in a water sample.
- 3. Plate counting requires considerable manpower and delivers results with some days of delay, which is caused by the necessary period of cultivation.



Fig. 1: Despite of the many disadvantages, heterotrophic plate count is still the standard methods for the quality control of drinking water.

The delay of the results (1 - 14 days) poses a risk, since the sampling of water and reporting of results are decoupled: By the time the results become available, the drinking water has already reached the consumer and, most likely, has been consumed for some time.

State-of-the-art: Flow cytometry

More and more scientific publications recommend bacterial cell counting (Total cell count, TCC) in drinking and industrial water via flow cytometry. One advantage of this applications is the potential of high throughput automation [1 - 5].

Sysmex Partec standardized and improved bacterial cell count in water samples using the accurate and convenient CyFlow™ Cube 6 V2m Flow Cytometer together with the two CyStain™ BacCount kits. The CyStain™ BacCount Total kit enables the detection of the total amount of bacteria in water samples. The total cell count is based on the labelling of all bacteria with the membrane-permeant nucleic acid-binding dye CyStain™ Green. After excitation the fluorescence signal is counted for each individual microorganism (Fig. 4).

Flow cytometry allows to determine the total bacterial count within less than 15 minutes

This procedure allows to determine the total bacterial cell count (TCC) in a water sample and to differentiate between bacterial populations with a high (HNA bacteria) and a low content of nucleic acid (LNA bacteria) within less



Fig. 2: The CyFlow[™] Cube 6 V2m Flow Cytometer with the CyFlow[™] Robby 6 V2m Autoloading Station.

Tab. 1: Advantages of flow cytometry compared to HPC

	Flow cytometry	НРС
Reproducibility	High	Low
Speed	15 min	Days
Automation	Yes, easy to scale up	No
Bacteria detected	All	Only those growing on agar
Sensitive for changes in cell count	Yes	No
Information	Total-, Living-, Dead-, VBNC-cells; Fingerprint (LNA/ HNA ratio)	Colony forming unit (CFU)

than 15 minutes from the time of sampling. The procedure is simple, reliable, fast and samples can be analysed by the small, compact and robust CyFlow™ Cube 6 V2m Flow Cytometer that is capable of volumetric counting of particles.

In contrast to HPC, this method detects not only those bacteria growing on agar plates but the entire population of bacteria, regardless of their potential to grow in an artificial or natural environment. This explains why HPC results and the total bacterial cell count determined by flow cytometry can differ by a factor of 100 to 10,000. Monitoring the TCC allows drawing conclusions on the present microbiological status of drinking water, but also of its development. The method is suitable for drinking and surface water.

To further discriminate between viable and non-viable bacteria the CyStain[™] **BacCount Viable** kit discriminates between live and dead bacteria (More information in the Application Note "APN_015A_CyStain BacCount Viable").

Make sample automation simple

There are 2 options using the CyStain™ BacCount Total kit·

- Manual measurements using the kit and the CyFlow™
 Cube 6 V2m Flow Cytometer, or
- Automated measurements using the kit, the CyFlow[™]
 Cube 6 V2m Flow Cytometer and the CyFlow[™] Robby
 6 V2m Autoloading Station.

Sample automation is the key requirement for quality and process control of water samples. The CyFlow™ Robby 6 V2m Autoloading Station enables a complete automation solution, which offers simplicity, sustainable reliability and exceptional walk-away convenience. This automated system is the ideal platform to tailor all customers needs in the water industry.

Instrument requirements

All analysis require the CyFlowTM Cube 6 V2m Flow Cytometer (Ref. No. CY-S-3061R). This flow cytometer is equipped with a blue laser (light excitation, 488 nm) and is able to analyse side scatter (SSC), green (536 \pm 40 nm) and red (> 630 nm) fluorescence as detection parameters. For an automated analysis the CyFlowTM Robby 6 V2m Autoloading Station (Ref. No. CY-S-3083) is required.

Material and methods

Kit components

The CyStain™ BacCount Total kit contains the following reagents:

- 5 Aliquots x 40 μL CyStain™ Green
- 29 mL CyStain™ Dilution Buffer



Fig. 3: CyStain[™] BacCount Total kit (Ref. No. 05-5008).

Additional required equipment

- 2 mL Reaction Tubes (Safe-Lock)
- 3.5 mL Sample Tubes (Ref. No. 04-2000)
- Heating block/water bath set to 37 °C ± 0.5 °C
- Vortex mixer

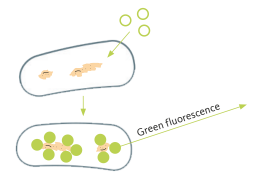


Fig. 4: The membrane-permeant CyStain™ Green dye shows a characteristic fluorescence after intercalating with the bacterial DNA.

Sample preparation

NOTE: There are two options for sample preparation: Manual and automated sample analysis. In this Application Note both options are presented in parallel.

Manual analysis

- 1a. Prepare a 10X working solution by diluting the 1000X stock solution of CyStain™ Green 1:100 with CyStain™ Dilution Buffer. E.g. for 10 samples mix 10 μL CyStain™ Green with 990 μL CyStain™ Dilution Buffer.
- 2a. Mix 100 μL CyStain™ Green working solution with 900 μL water sample in a 2 mL reaction tube.
- 3a. Mix sample with a vortex mixer for 3 seconds.
- 4a. Incubate sample for 13 minutes at 37 °C ± 0.5 °C, protected from light in a heating block or water bath.
- 5a. Mix sample with a vortex mixer for 3 seconds.
- 6a. Pipette 850 μ L of the sample into a sample tube for flow cytometry.

Automated analysis

- 1b. Prepare a 10X working solution by diluting the 1000X stock solution of CyStain™ Green 1:100 with CyStain™ Dilution Buffer.
- 2b. Mix 20 μL CyStain™ Green working solution with 180 μL water sample in a 96 Well Plate (Ref. No. O4-2020).
- 3b. Mix gently by pipetting up and down the solution.
- 4b. Incubate sample for 13 minutes at 37 $^{\circ}$ C \pm 0.5 $^{\circ}$ C, protected from light.

Analysis range

The maximum cell number for an accurate data acquisition is 2 x 10^5 cells/mL. It is recommended to dilute a water sample if the cell number is higher. Dilute the water sample with ultrapure water or 0.1 μ m filtered water before staining with the CyStainTM BacCount Total kit.

Tab. 2: Analysis range of CyStain™ BacCount Total kit

Detection Limit	Reportable Range
200 cells/mL*	1 x 10 ³ - 2 x 10 ⁵ cells/mL

*A specific treatment of the CyFlow™ Cube 6 V2m Flow Cytometer is required if the expected total cell count is lower than 1 x 10⁴ cells/mL. Please refer to section General hints for further information.

Data acquisition

- Switch on the CyFlow[™] Cube 6 V2m Flow Cytometer and use if required the CyFlow[™] Robby 6 V2m Autoloading Station.
- 2. Start the CyViewTM Software after the device has booted
- 3. For manual analysis load the script "CyStain BacCount Total_..._Standalone.cvc85". For automated analysis load the script "CyStain BacCount Total_..._Robby. cvc85".
- 4. For initial cleaning of the device start the Priming procedure and follow the instructions on the display until Priming is completed.
- 5. After completed Priming start the Quality Check (QC) procedure and follow the instructions on the display until the QC is completed.

NOTE: For further information see CyStain™ BacCount reagents – Quality Check Manual [6]

- Two FCS Express[™] reports will be automatically loaded to verify the QC. Enter QC material information for LOT N° and expiry date.
- 7. The analysis of stained water samples can be started after a "valid" QC procedure.

NOTE: Make sure that your device is qualified to perform measurements with the CyStainTM BacCount Total kit. If tests fail, the device is not qualified to perform measurements. Please see section Troubleshooting.

- 8. For manual analysis connect the sample tube from step 6a and start measurement. For automated analysis insert 96 Well Plate from step 4b into the CyFlowTM Robby Autoloading Station and start measurement.
- 9. The measurement automatically stops at the end of analysis.
- 10. FCS Express[™] automatically starts and opens analysis report.
- 11. All further samples are analysed as stated before.
- 12. Start the Shutdown procedure after the sample analysis is completed and follow the instructions on the display.

Exemplary data

Quality Check

The QC procedure controls background, laser power, optical alignment, gate positioning and counting precision. After the QC Part 1 and 2 are completed the results will be analysed automatically by FCS Express™. An exemplary report can be found in the CyStain™ BacCount reagents − Quality Check Manual (section 4.2.2) [6].

Water samples

For this application Note Evian mineral water (commercially available bottled water) samples with different cell numbers were chosen. Evian mineral water is used due to its natural content of high nucleic acid (HNA) and low nucleic acid (LNA) bacteria.

Sample 1: Evian mineral water

Sample 2: Evian mineral water, 0.1 µm filtered

Results

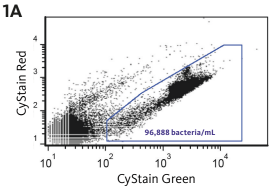
96,888 bacteria/mL were found in Evian mineral water. The 0.1 μ m filter successfully filter almost all bacteria in the water sample: Only 6 bacteria/mL were detected.

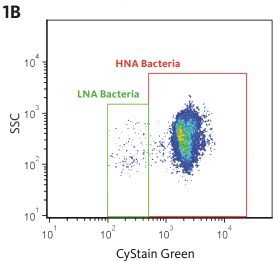
Tab. 3: Amount of detected bacteria in the 2 water samples

Water sample	Total bacteria
Evian mineral water	96,888/mL
Evian mineral water, 0.1 µm filtered	6/mL

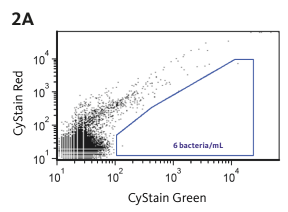
NOTE: The lower detection limit of the CyStainTM BacCount Total kit is 200 cells/mL. Therefore, a detected cell count of less than 200 cells/mL represent a negative result.

Sample 1: Evian mineral water





Sample 2: Evian mineral water, 0.1 µm filtered



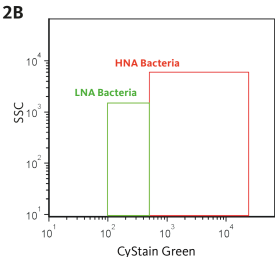


Fig. 5: Different samples of Evian mineral water stained with CyStain™ BacCount Total kit. (1) Evian mineral water, (2) Evian mineral water, 0.1 μm filtered (A): Dot plot showing CyStainTM Green vs. CyStainTM Red. All bacteria are located and automatically counted within the predefined blue gate. (B): Dot plot showing the side scatter (SSC) vs. CyStain™ Green which allows to determine the LNA and HNA numbers - the "Fingerprint" of a water sample.

General hints

Cleaning steps between different samples

Depending on the cell count of the water sample, as well as the analysis method cleaning steps between individual measurements are recommended to reduce the possibility of cross contamination and background signals.

Tab. 4: Cleaning steps between different samples

Set of water samples		Cell content		
		similar	different	unknown
clooping	manual analysis	not required	intermediate cleaning with Sheath Fluid	intermediate cleaning with Sheath Fluid
cleaning step	automated analysis	cleaning mode none	cleaning mode normal* or intensive**	cleaning mode normal* or intensive**

In case of changing the application from total cell count to viable cell count determination on the same day no additional QC procedure or a Shutdown of the device is necessary. However, a Priming procedure is required.

Intensive cleaning procedure

The analysis of water samples with an expected total cell count lower than 1 x 10⁴ cells/mL requires a special treatment of the CyFlow™ Cube 6 V2m Flow Cytometer. To maintain ultra-clean conditions the following manual cleaning procedure is recommended:

NOTE: In addition to the cleaning procedure a filtration (0.22 µm filter) of Sheath Fluid is recommended.

- Empty the Sheath Fluid bottle and rinse twice with Hypochlorite Solution (Ref. No. 04-4012_R).
- 2. Remove the used Inline Filter (Ref. No. 04-004-1000) of the Sheath fluid bottle.
- 3. Fill the Sheath Fluid bottle with 200 mL Hypochlorite Solution.
- 4. Fill a sample tube with 3 mL Hypochlorite Solution.
- Start a measurement at a speed of 0.2 µl/sec until it stops automatically.
- 6. Wait 90 min and measure again.
- Stop the measurement after 10 min.

^{*}Cleaning with Sheath Fluid (Ref. No. 04-4007_R)

**Cleaning with additional Cleaning Solution (Ref. No. 04-4009_R)

- 8. Exchange the Hypochlorite Solution in the sheath fluid bottle with Sheath Fluid. Rinse twice with Sheath Fluid before filling up the bottle.
- 9. Attach a new Inline Filter to the Tubing of the Sheath Fluid bottle.
- 10. Fill a sample tube with 3 mL Sheath Fluid.
- 11. Start a measurement with Sheath Fluid at a speed of $0.2 \mu l/sec$.
- 12. Stop the measurement after 30 min.

Troubleshooting

If the error you are experiencing is not described or the remedy could not solve your problem, please contact your local Sysmex representative.

Error #1: QC procedure "invalid"

Reason	Remedy
QC material poorly mixed	Shake the QC material vigorously (e.g. by vortexing) and repeat QC procedure
	Check the date of expiry of the QC material
	Perform a priming procedure and repeat QC procedure
	Perform an intensive cleaning procedure and repeat QC procedure
High concentration of Calibration Beads 0.5 μm (Ref. No. 05- 4005)	Perform an intensive cleaning procedure, prepare a new dilution of the QC material and repeat QC procedure
LOT NO° mismatch of QC material	Make sure to use a matching FCS Express™ template and QC material LOT NO°

Error #2: Population(s) of bacteria cells outside of predefined gate

Reason	Remedy
Air bubbles inside the device	Perform a priming procedure, repeat sample preparation and measurement
	Perform an intensive cleaning procedure, repeat sample preparation and measurement

Error #3: No separation between bacteria cells and background signal

Reason	Remedy
Cell number higher than 2 x 10 ⁵ cells/mL	Dilute water sample with filtered (0.22 µm Filter) or ultrapure water, repeat sample preparation and measurement

References

- 1. Van Nevel S., et al., 2017. Flow cytometric bacterial cell counts challenge conventional heterotrophic plate counts for routine microbiological drinking water monitoring. Water Res. 113: p. 191-206.
- 2. Wang, Y., et al., 2010. Past, present and future applications of flow cytometry in aquatic microbiology. Trends Biotechnol. 28, 416-424.
- 3. Van Nevel S., et al., 2013. Routine bacterial analysis with automated flow cytometry. J Microbiol Methods. 94, 73-76.
- 4. Hammes, F., et al., 2008. Flow-cytometric total bacterial cell counts as a descriptive microbiological parameter for drinking water treatment processes. Water Research. 42, 269-277.
- 5. Safford HR., et al., 2018. Flow cytometry applications in water treatment, distribution, and reuse: A review. Water Research. 151, 110-133.
- CyStain[™] BacCount reagents Quality Check Manual.