Application note No. 3007. Rev. 1.3

NucleoCounter® NC-3000™

Count of PI Stained Cells – Total Count of Mammalian Cells

Product description
The NucleoCounter® NC-3000™ system enables the user to perform automated cell counting and analyses of a broad range of eukaryotic cells.

Application
The PI-Cassette™ is used together with Reagent A100 and B and analyzed using the NucleoCounter® NC-3000™. Thereby it facilitates the determination of cell concentration even if the cell type investigated exhibit a very aggregating phenotype and the cell concentration cannot accurately be determined using the Via1-Cassette™.

Introduction
Propidium iodide (PI) is immobilized inside the PI-Cassette™ and has the ability to stain DNA of non-viable cells. PI enters non-viable cells, as their plasma membrane is permeable. In order to measure the total concentration of cells the plasma membranes of all cells in the sample must be disrupted to render all nuclei susceptible to staining with PI. The disruption is achieved by treatment with a lysis buffer (Reagent A100) followed by a stabilizing buffer (Reagent B). After treating the cells with the Reagent A100 and Reagent B the suspension is loaded into the PI-Cassette™. Once inside the PI-Cassette™ the nuclei are stained with PI. The PI-Cassette™ is placed in the NucleoCounter® NC-3000™ where the cell concentration is determined.

Procedures
If the cell line to be investigated is adherent or semi-adherent, then start by getting all cells into suspension using the preferred method of your laboratory (e.g. trypsin/EDTA treatment).

Materials needed
- Cells to be counted
- PI-Cassette™
- Reagent A100 (Lysis buffer)
- Reagent B (Stabilizing buffer)

1. The original cell suspension is mixed to obtain a homogenous suspension. Pipette a representative cell sample from the cell suspension into a microcentrifuge tube (e.g. 100 µl).
2. Add 1 volume of Reagent A100 to the microcentrifuge tube with the cell sample. E.g., if the volume of the cell sample is 100 µl then add 100 µl of Reagent A100. Mix by pipetting.
3. Add 1 volume of Reagent B to the mixture of cell suspension and Reagent A100. E.g. to 200 µl of the mixture of cell suspension and Reagent A100 add 100 µl Reagent B. Mix by pipetting.
4. Draw the diluted cell suspension into a PI-Cassette™ by inserting the tip of the cassette into the cell suspension and press the piston.
5. Immediately place the loaded PI-Cassette™ in the NucleoCounter® NC-3000™ sample tray, select the protocol “Count of PI Stained Cells” with the sample unit “PI-Cassette™” and press RUN.

After approximately 45 seconds the total cell concentration (cells/ml) is presented in the bottom right result. The presented cell concentration of the total cell count has been compensated for the dilution caused by adding of Reagent A100 and Reagent B. If the sample has been further diluted and the user has entered the volumes or dilution factor into the user interface, the dilution factor has also been taken into account and the cell concentration given is for the original cell sample.
Note
To assure reliable results, it is recommended that the total cell concentration of the cell suspension should be in the range of $5 \times 10^4$ cells/ml to $5 \times 10^6$ cells/ml. If the concentration of cells is below $5 \times 10^4$ cells/ml then the cell concentration may be increased by centrifugation followed by resuspension of the pellet using growth media or PBS. The resuspended cell sample is then treated as described above.

If the total cell concentration is above $5 \times 10^6$ cells/ml, the cell suspension should be diluted with growth media or PBS to achieve the desired concentration. The diluted cell sample is then treated as described in the procedure.

Cell count determination of MCF7 cells. The cells were harvested and the sample treated with Reagent A100 and B was loaded into a PI-Cassette™ and analyzed using the Count of PI stained Cells Assay. The result is presented at the bottom right. An insert shows a close up of a part of the image.

Troubleshooting

Inaccurate and imprecise counting:
When setting up a new cell line it is important to inspect that the cell line is counted correctly. The cells included in the total count can be marked by clicking on the overlay button in the bottom right corner of the image. Visual inspect the image to evaluate in the vast majority of the cells has been counted correctly. If this is not the case right click on the image file in question and choose “Show Raw Data”. Inspect the gates displayed in the Plot Manager. If the gating is inappropriate right click on the image file in question again and choose “Start Protocol Adaptation Wizard”. Adapt the gate(s) to cover the cell population (do not include debris and very large objects) and save the changes to a new protocol. Note that the user is responsible for defining appropriate gating of the particular cell line.
Handling and storage
For handling and storage of ChemoMetec instruments and reagents, cassettes and NC-Slides refer to the corresponding product documentation. For other reagents refer to the material data sheet from the manufacturer of the reagents and chemicals.

Warnings and precautions
For safe handling and disposal of the ChemoMetec reagents, cassettes and NC-slides refer to the corresponding product documentation and the NucleoCounter® NC-3000™ user’s guide. For other reagents refer to the safety data sheet from the manufacturer of the reagents and chemicals required for this protocol. Wear suitable eye protection and protective clothes and gloves when handling biologically active materials.

Limitations
The NucleoCounter® NC-3000™ system is FOR RESEARCH USE ONLY. NOT FOR DIAGNOSTIC OR THERAPEUTIC USE. The results presented by the NucleoCounter® NC-3000™ system depend on correct use of the reagents, NC-slide and the NucleoCounter® NC-3000™ instrument and might depend on the type of cells being analyzed. Refer to the NucleoCounter® NC-3000™ user’s guide for instructions and limitations.

Liability disclaimer
This application note is for RESEARCH PURPOSES ONLY. It is not intended for food, drug, household, or cosmetic use. Its use must be supervised by a technically qualified individual experienced in handling potentially hazardous chemicals. The above information is correct to the best of our knowledge. Users should make independent decisions regarding completeness of the information based on all sources available. ChemoMetec A/S shall not be held liable for any damage resulting from handling or contact with the above product.

Product disclaimer
ChemoMetec A/S reserves the right to introduce changes in the product to incorporate new technology. This application note is subject to change without notice.

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