

Normalization of Agilent Seahorse™ XF Data with Bright Field-Based Direct Cell Counting

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Introduction

Mitochondrial respiration and glycolysis are two highly important parameters for understanding energy metabolism and cellular functions. Cellular metabolism can also be a key indicator of tumorigenesis, aging, stress, and other diseases.

In the recent years, the Seahorse™ XF Analyzer has been diligently utilized to measure the Oxygen Consumption Rate (OCR) and Extracellular Acidification Rate (ECAR)/ Proton Production Rate (PPR) for assessing mitochondrial respiration and glycolysis, respectively [1].

One of the important steps in performing Seahorse™ XF cell metabolism analysis is normalization of the resultant data to minimize inconsistency and variations from well-to-well due to potential non-uniform cell seeding across the microplate. In general, normalization has been commonly performed using total protein amount in each well such as BCA assays, however cells can be lost during the assay due to the metabolic inhibitors used during the assay. Additionally, normalizing Seahorse™ data using total protein analysis is time-consuming and requires multiple steps such as cell lysing, incubation, and transferring lysed content for analysis. The BCA assay can take up to ~45 min for the entire 96 well plate. Washing and transfer procedure can introduce risk such as cell loss or pipetting error, thus skewing the final normalized results [2].

An alternative method that can improve Seahorse™ XF normalization is performing direct cell count per well prior to the metabolic stress tests. Cell counting is a non-destructive method to generate data for Seahorse™ normalization, which can take account of non-uniform seeding density across the well and plate. Since cell counting is non-destructive, downstream assays are still possible after normalization. Finally, direct cell counting can significantly reduce the amount of time required to perform normalization [1, 2].

In recent years, Celigo Image Cytometer has been used to perform bright field or fluorescence-based direct cell counting for Seahorse™ XF data normalization [3]. Due to its ability to capture whole well images, at high speed; Celigo direct cell counting application has been adapted into the general workflow for Seahorse™ XF96, XFe96, and XF24 Analyzers. The system contains built-in plate profiles for the Seahorse™ plates and will acquire one whole well image of a 96-well plate in ~4-5 seconds, and analysis of the entire plate is ~6 min in bright field, and ~9 min in fluorescence.

Celigo Image Cytometer can be used to perform direct cell

counting of suspension and adherent cells in bright field, as well as fluorescence such as Hoechst 33342 or Live-nuclear dyes staining. Both bright field and fluorescence methods can be used effectively to normalize Seahorse™ XF data. The commonly used DNA minor groove-binder Hoechst 33342 is non-toxic when used at low concentrations, but requires the use of Ultraviolet (UV) excitation light, potentially damaging cells, which may introduce cytotoxic effects. The use of live nuclear dyes (Nexcelom Bioscience) can emit a brighter fluorescence signal, and have little to no cytotoxic & cytotoxic effects. In addition, the cell identification algorithms are highly accurate for segmenting suspension or adherent cells for both bright field and fluorescence imaging. Fluorescence live nuclear dyes can improve accuracy for counting adherent cells that tend to flatten out after attachment.

One major requirement for obtaining reproducible and reliable data with the Seahorse™ XF Analyzers is working with a consistently uniform population of target cells that are evenly distributed and firmly attached to the culture dish. Celigo Image Cytometer provides a fast, effective, and cell stress-free technique to count cells plated onto uncoated or coated culture plates. Furthermore, Celigo has a large dynamic range of counting cells in whole well, i.e. counting 1 to ~120,000 cells per well for suspension cells and 1 to ~90,000 cells for adherent cells in a 96-well plate.

In this application note, we demonstrate the use of Celigo Image Cytometer for direct cell counting in bright field and fluorescence using Hoechst to normalize Seahorse™ XF96 data and compare to protein-based normalization. The results showed significantly improved normalized data, as well as a reduction in assay time, which effectively improved the entire OCR and ECAR measurements.

Materials and Methods

Bright field-based direct cell counting method for Seahorse™ XF normalization

In this experiment, the glycolytic function (ECAR) was assessed for DCIS and HCT116 cell lines for approximately 160 min. The Seahorse™ XF data were normalized to both protein amount and bright field cell counting using the Celigo Image Cytometer. The normalized time-dependent ECAR data were then compared, which showed improvement in cell line differentiation using the cell count normalization.

1. DCIS and HCT116 cells were seeded at 12,500 cells/well and 10,000 cells/well, respectively, in the Seahorse™ XF 96-well plate and incubated overnight

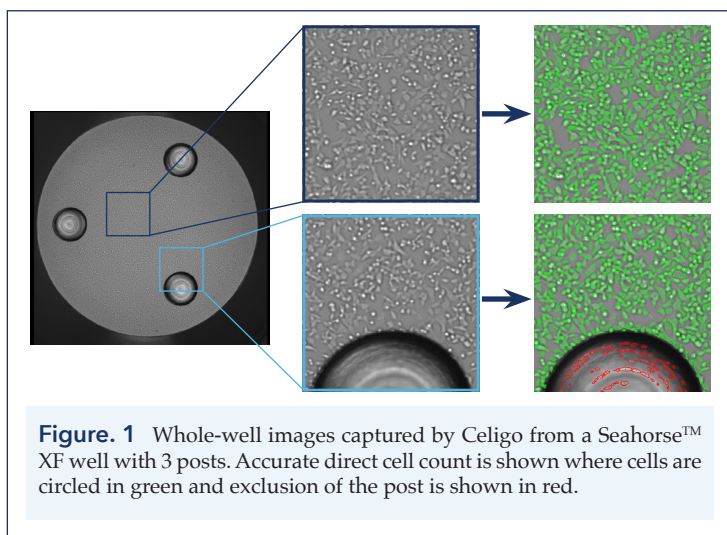
2. The plate was first visualized on microscope for approximately 95% confluence
3. The confluence was then confirmed at 98% using the Celigo, as well as QC for cell health, morphology, seeding uniformity and purity (no contamination) prior to washing
4. Next, washed with the appropriate assay media
5. Subsequently, the plate was imaged and analyzed on the Celigo using direct cell counting application
6. Next, the plate is placed in a 37° C incubator without CO2 for 45 minutes
7. The plate was then analyzed using Seahorse™ XFe96 to measure ECAR for 160 min
8. Finally, the plate was used to perform protein analysis for normalization

Fluorescence-based direct cell counting method for Seahorse™ XF normalization

In this experiment, both mitochondrial respiration (OCR) and glycolytic function (ECAR) were assessed for T cells for approximately 60 min. The Seahorse™ XF data were normalized to both protein amount and fluorescent cell counting using the Celigo Image Cytometer. The normalized end point ECAR data were then compared, which showed improvement in drug response differentiation using the cell count normalization.

1. T cells were seeded at 100,000 cells/well in the Seahorse™ XF 96-well plate
2. The cells were allowed to settle to the bottom of the plate
3. The plate was then placed into the Seahorse™ XF Analyzer and allowed delivery of Hoechst 33342, and subsequently analyzed for 60 min
4. The plate was then counted on the Celigo for total Hoechst positive cells

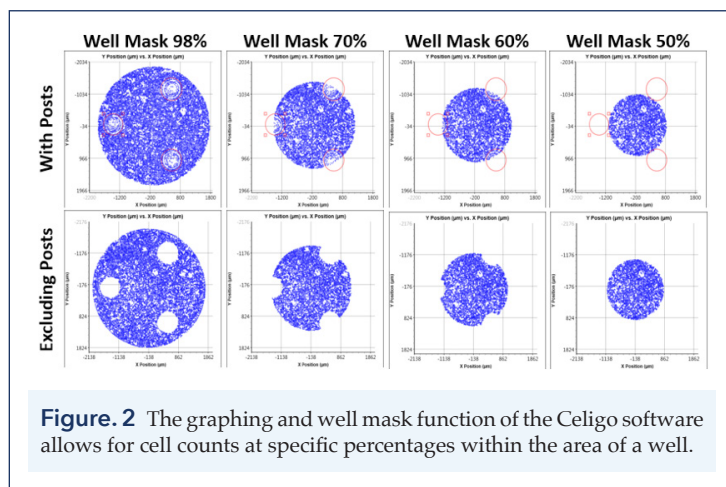
Results



Bright field-based direct cell counting method for Seahorse™ XF normalization

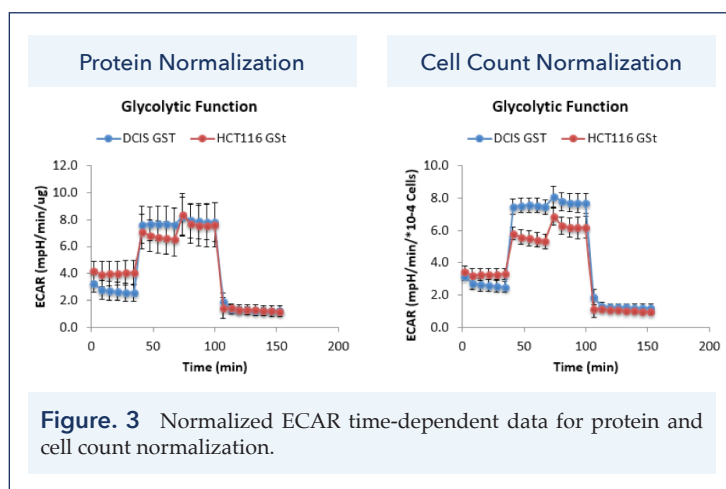
The Celigo captured bright field images are displayed in Figure 1, which showed a whole well image of a Seahorse™ XF well with the 3 posts. In addition, the zoomed in images showed accurate direct cell counting of the HC116 cells (green outline), as well as exclusion of the post area (red outline).

By utilizing the graphing function and well mask feature in the Celigo software, researchers can count cells at specific percentages



of area within the well, which may allow better normalization for the Seahorse™ XF data (Figure 2).

The normalized ECAR time-dependent data for protein and cell count normalization are shown in Figure 3. It is obvious that measurements with protein normalization cannot distinguish between the glycolytic function of DCIS and HCT116 cell lines. In contrast, the cell count normalization method showed a reduction in standard deviation as well as a clear separation between the ECAR values of DCIS and HCT116 cell lines.



Fluorescence-based direct cell counting method for Seahorse™ XF normalization

The Celigo captured bright field and Hoechst fluorescence overlay images are shown in Figure 4. The counted image showed yellow outlines indicating counted Hoechst positive cells. Similarly,

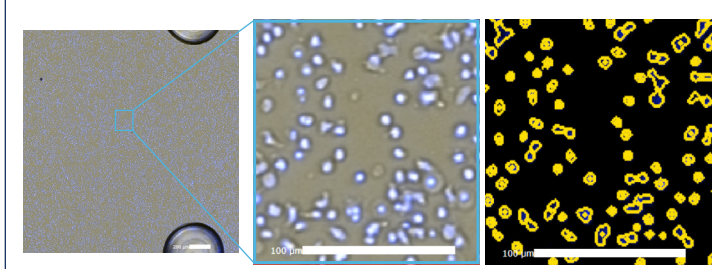


Figure. 4 Celigo captured bright field image (left), zoomed merged bright field and Hoechst image (center), and yellow ROI counted Hoechst stained cells (right).

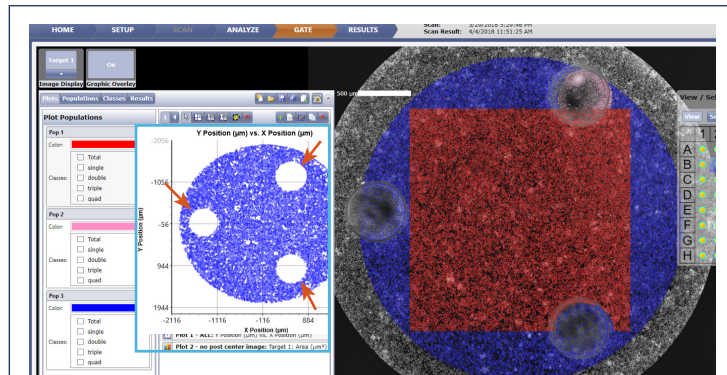


Figure. 5 Using the Celigo gating function, the location graph is generated to gate out the 3 posts (indicated by red arrows above) in the well using fluorescence. In addition, if only the area between the 3 posts are analyzed (shown with red square), scanning and counting require only 2 minutes per plate.

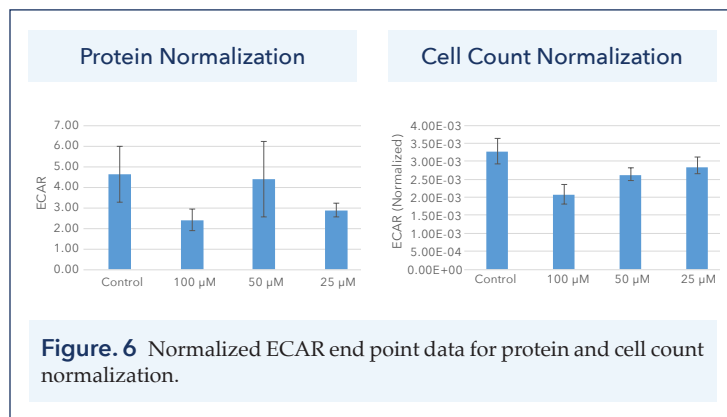


Figure. 6 Normalized ECAR end point data for protein and cell count normalization.

under the gating function, the location graph can be generated to gate out the 3 posts in the well using fluorescence (Figure 5).

The normalized ECAR end point data for protein and cell count normalization are shown in Figure 6. Similarly, the protein and cell count normalization method showed distinct results. Using protein normalization, large standard deviation for each sample can be observed, as well as inexplicable drug treatment trends. On the other hand, the cell count normalization showed a reduction in standard deviation, as well as showing expected drug treatment results with a decrease in ECAR as drug concentration increased.

Conclusions

The ability to employ high-throughput and high-speed direct cell counting method for Seahorse™ XF normalization enables improvement in final OCR and ECAR results as well as a reduction in time required from the BCA protein analysis. Below we will discuss the 4 key advantages of using Celigo Image Cytometer for Seahorse™ XF data normalization.

Speed to Results

The Celigo Image Cytometer can scan whole well images and directly count cells in less than 6 and 9 minutes per 96-well plate (less than 4 and 5 seconds per well) using bright field and fluorescence imaging, respectively (Figure 7). If only the area between the 3 posts are captured, the entire process requires only 2 min. Researchers investigating mitochondrial respiration and glycolytic functions using Seahorse™ XF Analyzer can rapidly

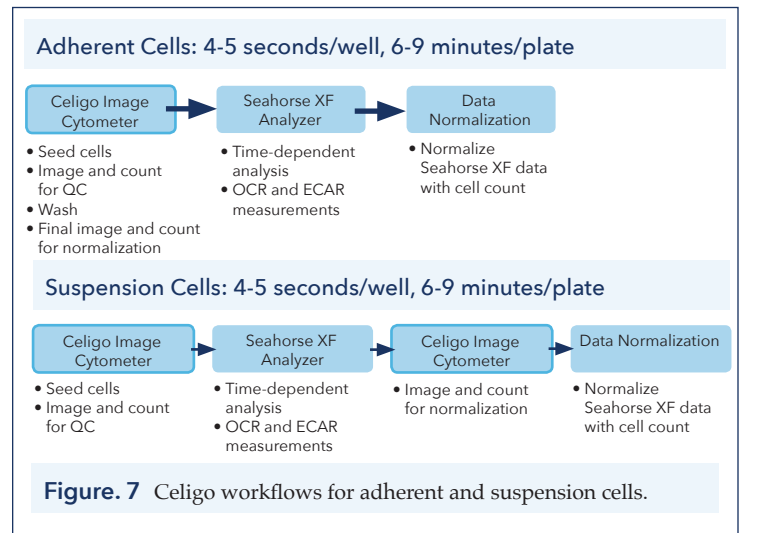


Figure. 7 Celigo workflows for adherent and suspension cells.

and effectively normalize the OCR and ECAR measurements without additional procedures like protein analysis.

Whole Well Image and Image Quality

Image quality is a critical advantage for the Celigo Image Cytometer. Because of components of F-Theta lens and

cancer, cell aging, stress, and stem cell research.

References

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2. "Normalization of Agilent Seahorse XF Data by In-situ Cell Counting Using a BioTek Cytation 5", Application, Seahorse Bioscience
3. Riddle et al., "Expansion capacity of human muscle progenitor cells differs by age, sex, and metabolic fuel preference," Am J Physiol Cell Physiol, doi: 10.1152/ajpcell.00135, 2018

*Seahorse™ XF Analyzer is a product of Agilent
<https://www.agilent.com/en/products/cell-analysis/seahorse-analyzers>

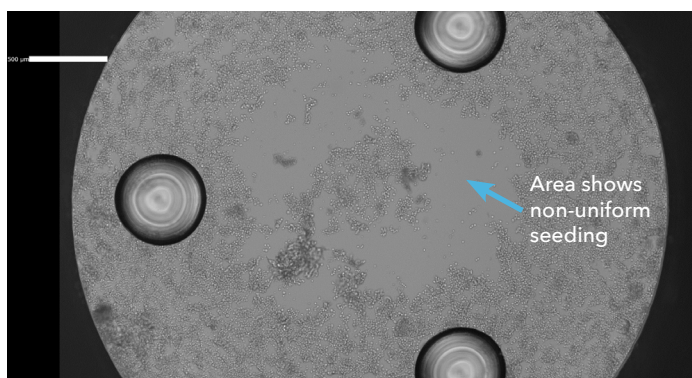


Figure 8 Celigo whole-well bright field image shows non-uniformly seeded cells.

Galvanometric Mirror, Celigo generates highly uniform and rapid whole well imaging in both bright field and fluorescence for the entire plate. Therefore, it is unnecessary to extrapolate from one single image in the middle of the well, which may create uncertainty and inaccurate counting in the well if the cells are not uniformly seeded (Figure 8). These images can also help to identify issues with cells or incorrect seeding if assay results are not as expected.

Sophisticated and Accurate Cell Counting Algorithm

Since cells come in all shapes and sizes, researchers also require a flexible and sophisticated cell counting platform to ensure accurate results for normalization. Both adherent and suspension cells can utilize a variety but specific counting algorithm to count the target cells in both bright field and fluorescent images. In addition, location plots are used to specifically gate out the 3 post areas to further improve cell counting accuracy (Figure 2).

Perform label-free direct cell counting

The Celigo Image Cytometer can be used to directly count both adherent and suspension cells without fluorescent labels, which can eliminate the potential cytotoxic effect on the cells, as well as reducing the time for staining. However, since Celigo has one bright field and 4 fluorescent channels (Blue, Green, Red, and Far Red), fluorescence-based direct cell counting can also be easily performed.

In conclusion, the advantages to using a Seahorse™ Extracellular Flux Analyzer to measure metabolic phenotypes of cells, as opposed to other more traditional metabolism assays, are considerable. Incorporation of the Celigo applications to provide direct cell counts provides better insight into Seahorse™ data analyses and ensures that healthy cultures are presented to the Seahorse™ XF analyzer. Using the Celigo Image Cytometer for direct cell counting can eliminate the time and accuracy issues raised from normalizing data using the protein-based assays. Celigo image cytometry applications can significantly improve assay results for researchers investigating cellular metabolism for