

Reduce FACS Sorting Speed by 10x with Superior DTC Cleanup

Introduction

Resection of tumor tissue for in-depth study is a staple of cancer research. Dissociation of these tissues into a single cell suspension via enzymatic and/or mechanical techniques yields a cellular milieu that provides the closest representation of the clinical scenario by retaining the characteristics and cell type fractions present in the tumor. Access to these samples is critical to understanding the innate biology of the tumor and the host response. The fields of personalized medicine, drug/biomarker discovery, and preclinical/translational research rely heavily on the ability to study dissociated tumor cells (DTCs) from a wide range of cancer types using several different techniques. Central to the ability to study DTCs using any of these technologies is sample quality post-cryopreservation, particularly cell viability and purity (absence of debris).

Common Challenges With DTC's

The freeze-thaw process that these samples undergo inherently damages some portion of the total sample, typically manifested through cell loss due to routine handling and removal of the cryopreservative via washing. Cell viability may also be negatively impacted by the freeze-thaw process. Further, the dissociation process itself can vary widely depending on the tumor and the host organ. The quality of the DTC suspension plays a significant role in the ultimate quality and usability of the sample post-cryopreservation.

In addition to the negative effects of tissue dissociation and cryopreservation on total cells and overall viability, these samples are often plagued by the presence of cellular debris. A common use of these samples is to characterize them using routine cell sorting or analysis, but this can be impeded by the amount and characteristics of the cellular debris present.

Excessive debris and dead cells can confound the ability to efficiently analyze the sample via cell sorting or analysis. Small bits of debris and dead cells can cloud the analysis while larger debris can clog the

KEY HIGHLIGHTS

Levitation Enabling Gentler Cell Processing

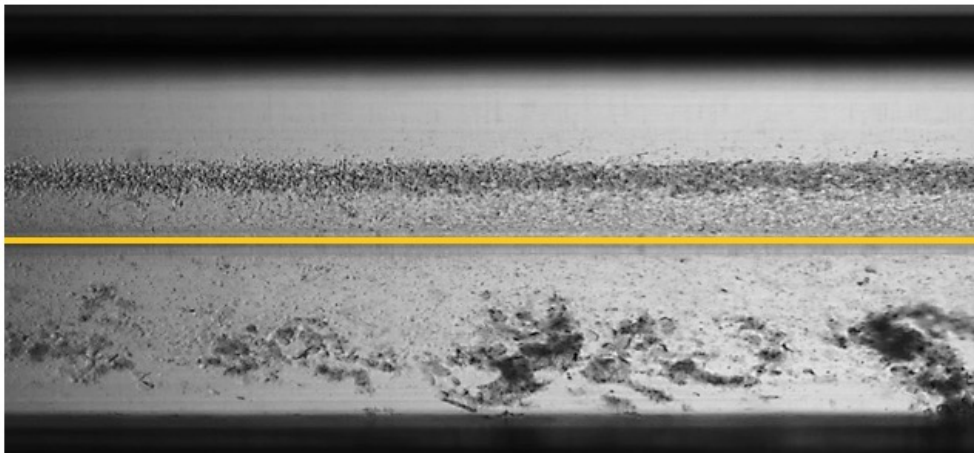
- ✓ Increased number of viable cells from initial sample
- ✓ Removal of all debris for the most efficient cell sorting or analysis
- ✓ Lower FC nozzle pressure to protect sensitive cell types

instrument, resulting in extra time spent cleaning the instrument while the cells sit idle, further degrading the quality of the sample.

During a typical flow cytometry run, if there is a lot of debris and dead cells, high nozzle pressure is often used to process enough events to ensure the viable cells of interest are observed. This higher pressure in turn adds undue mechanical stress on the cells. Fragile cells may not tolerate these conditions, and if the cells of interest are particularly fragile, there is a risk of losing those cells altogether due to artificial cell death.

Viable Cell Enrichment and Debris Removal Using LeviCell System

Optimizing the condition of these precious samples by magnetic levitation on the **LeviCell system** enriches the most viable cells and removes both small and large debris, as well as dead cells from the suspension (Figure 1). There are two primary benefits of sample optimization by levitation upstream of the flow cytometry workflow. First, enriched viable samples generate more meaningful events in the final analysis. Fewer dead cells and small debris events are present, allowing more confidence in the analysis and gating strategy. Second, if the sample contains a higher relative percentage of viable cells, then lower nozzle pressure can be used which allows for more fragile



Viable cells
Split line = 0
Dead / dying cells
Debris

Figure 1. Viable cells levitate above the split line (midpoint between the magnets), dead cells are immediately below the split line, and debris is observed as larger objects and clumps near the bottom of the levitation chamber.

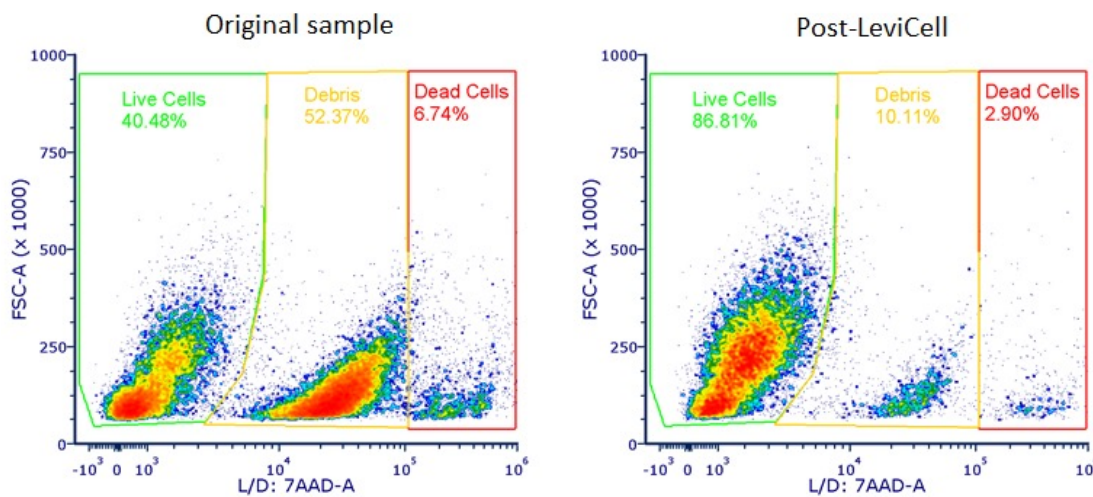


Figure 2. Flow cytometry analysis of a human lung DTC sample before (left) and after (right) viable cell enrichment and dead cell/debris removal with the LeviCell system. After levitation, debris decreased more than 5-fold, and viable cells increased almost 2-fold from <50% of events to nearly 90% of all events.

cells to potentially withstand the mechanical stresses of flow cytometry.

As seen in Figure 2, this human lung DTC was characterized before and after magnetic levitation to enrich viable cells and remove dead cells and debris with the LeviCell system. 7-Aminoactinomycin D (7AAD) was used to distinguish between intact dead cells (7AAD^{Pos}) and viable cells (7AAD^{Neg}). The population of events lying between the dead and

viable cells were defined as debris.

Conclusion

Fast and gentle magnetic levitation on the LeviCell system can significantly clean-up poor quality samples like DTC's, generating viable high quality cells prior to flow analysis. A higher viability, debris-free sample promotes more efficient flow cytometry analysis while enabling lower nozzle pressures, leading to better survival of particularly sensitive cell types.

For more information, visit levitasbio.com, or contact sales@levitasbio.com.

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