

INTRODUCTION:

A number of different techniques are routinely used in the biopharmaceutical discovery and development workflow. These include single cell analysis, sorting, imaging and dispensing into individual wells of microtitre plates (MTPs). Traditionally, different instruments would be required for each technique; which is costly, time-consuming and requires extensive lab space. Picodroplet techniques allow for sophisticated and sensitive manipulation of cells at the single cell level. Cyto-Mine® technology is the first integrated device to automatically perform all of these crucial techniques in a single compact system. This high-throughput instrument uses picodroplet technology and microfluidics to process around 10 million heterogeneous mammalian cells in a few hours. Each cell is encapsulated in a picodroplet containing cell culture growth media, which acts as a bioreactor to compartmentalise the cell and let it grow; eventually trapping secreted molecules such as antibodies (Abs). The system also allows selection and deposition of single cells into well of 96- and 384-well MTPs - ensuring high fidelity monoclonality assurance (required for FDA-approval of all monoclonal Abs).

CHALLENGES IN ANTIBODY DISCOVERY:

- Low efficiency of hybridoma technology.
- Single cell screening is needed for deep repertoire interrogation.
- Single cell assays are required to detect surface-bound and/or secreted antibodies.
- Good screening specificity is desirable to reduce downstream costs.
- A large number of cells needs to be screened.
- Cell friendly processing is desirable to maximize transcript recovery.

CHALLENGES IN CELL LINE DEVELOPMENT:

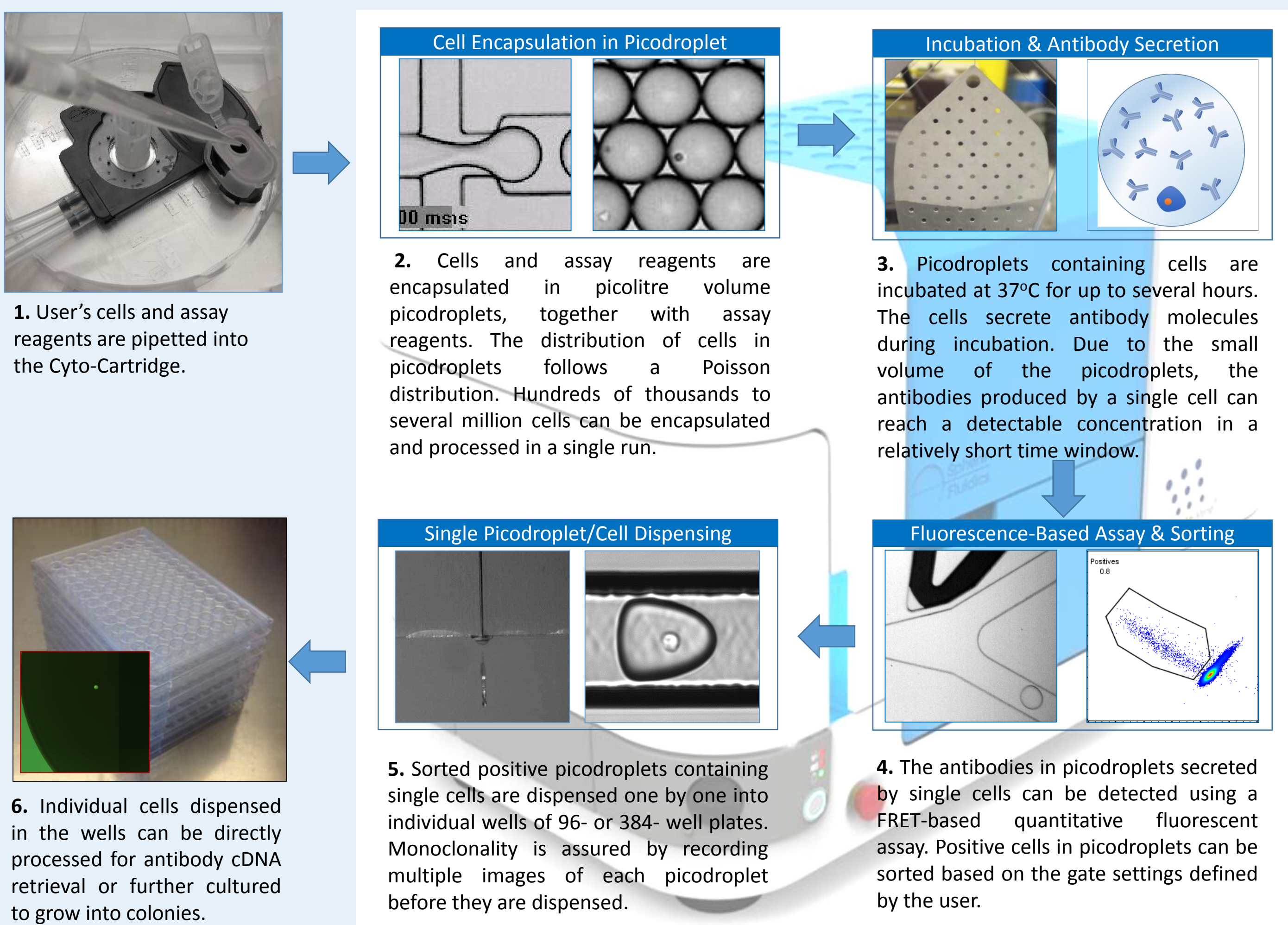
- Larger scale cloning is desirable to maximize the chance of winning.
- Random cloning results in costly post cloning screenings.
- Lack of assays reliably ranking and selection based single cell productivity.
- Cell friendly processing is desirable to allow cell out-growth.
- Evidence of monoclonality is a regulatory requirement.

CYTO-MINE® KEY FEATURES

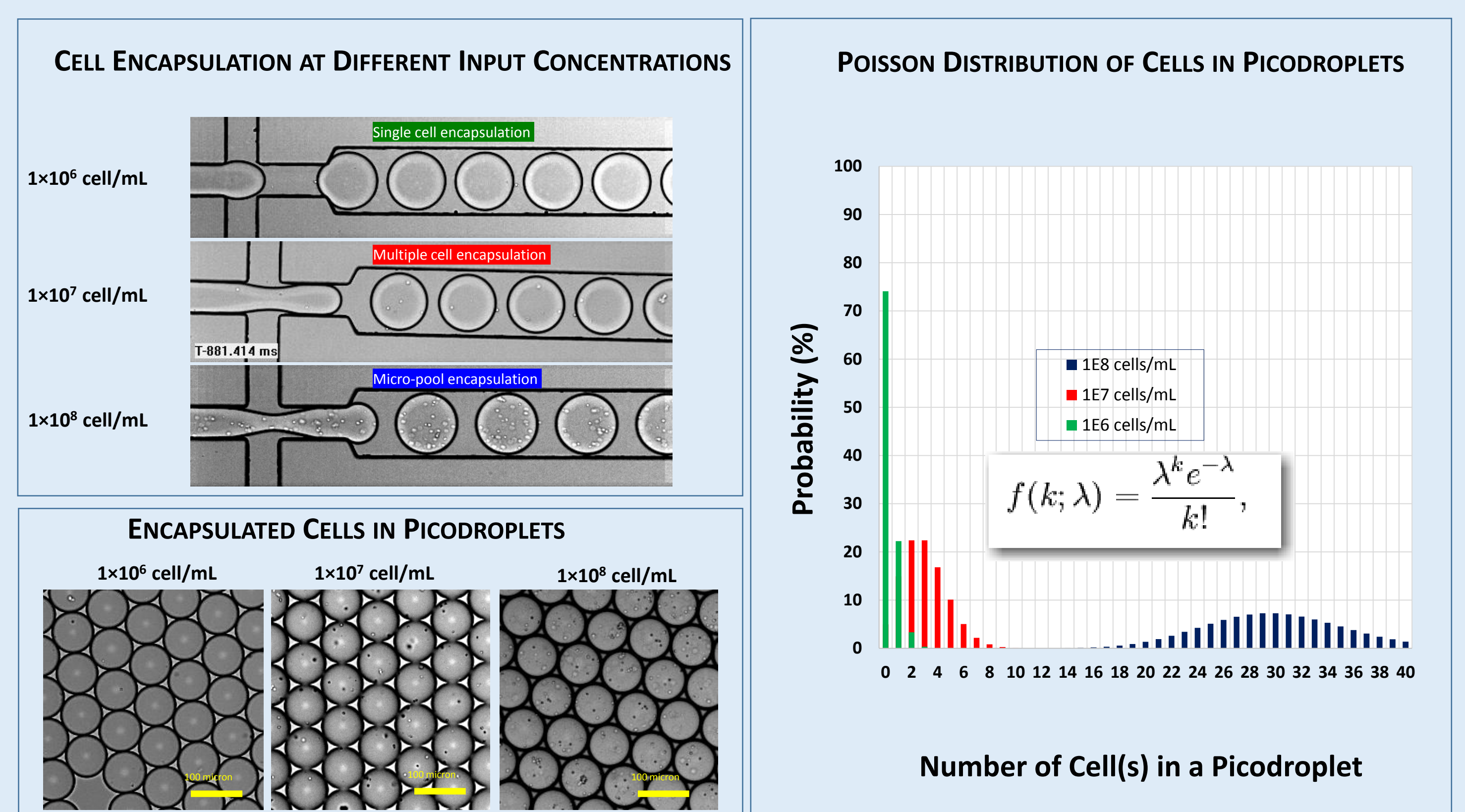
SCREENING	CLONING	OTHERS
<input type="checkbox"/> Quantitative single cell secretion assay	<input type="checkbox"/> Cell friendly dispensing	<input type="checkbox"/> Sterile
<input type="checkbox"/> Customizable assay designs	<input type="checkbox"/> Monoclonality assurance	<input type="checkbox"/> Animal origin free
<input type="checkbox"/> Cell friendly sorting	<input type="checkbox"/> Database compatible	<input type="checkbox"/> User friendly
		<input type="checkbox"/> Automation compatible

CYTO-CARTRIDGE®	CYTO-MINE® INSTRUMENT	CYTO-SURF®
<ul style="list-style-type: none"> Fully integrated microfluidic functional modules. Mass manufactured. Sterilized. Single-use. Animal origin free. 	<ul style="list-style-type: none"> Fully integrated control modules. User-friendly GUI. Compact design. Single use consumables. Animal origin free. Robotic arm accessible. 	<ul style="list-style-type: none"> Patented formulation. Biocompatible. Sterilize. Animal origin free.
Dimensions L x H x W = 860 x 566 x 463 mm		
Weight 85 Kg		
Detection mode Laser induced fluorescence		
Excitation Laser wavelength λ _{ex} =488 nm		
Detection colours Two		
Picodroplet size 300 pL		
Screening capacity Up to 2,000,000 picodroplets/run		
Screening throughput 10,000-10,000,000 cells in 2-7 hours		
Speed of dispensing 1 picodroplet per second		
Max. no. of dispensings 10,000		
Consumables Cyto-Cartridge® Cyto-Surf®		
Compatible cell types CHO, B Cells, Hybridoma		

CYTO-MINE® WORKFLOW:

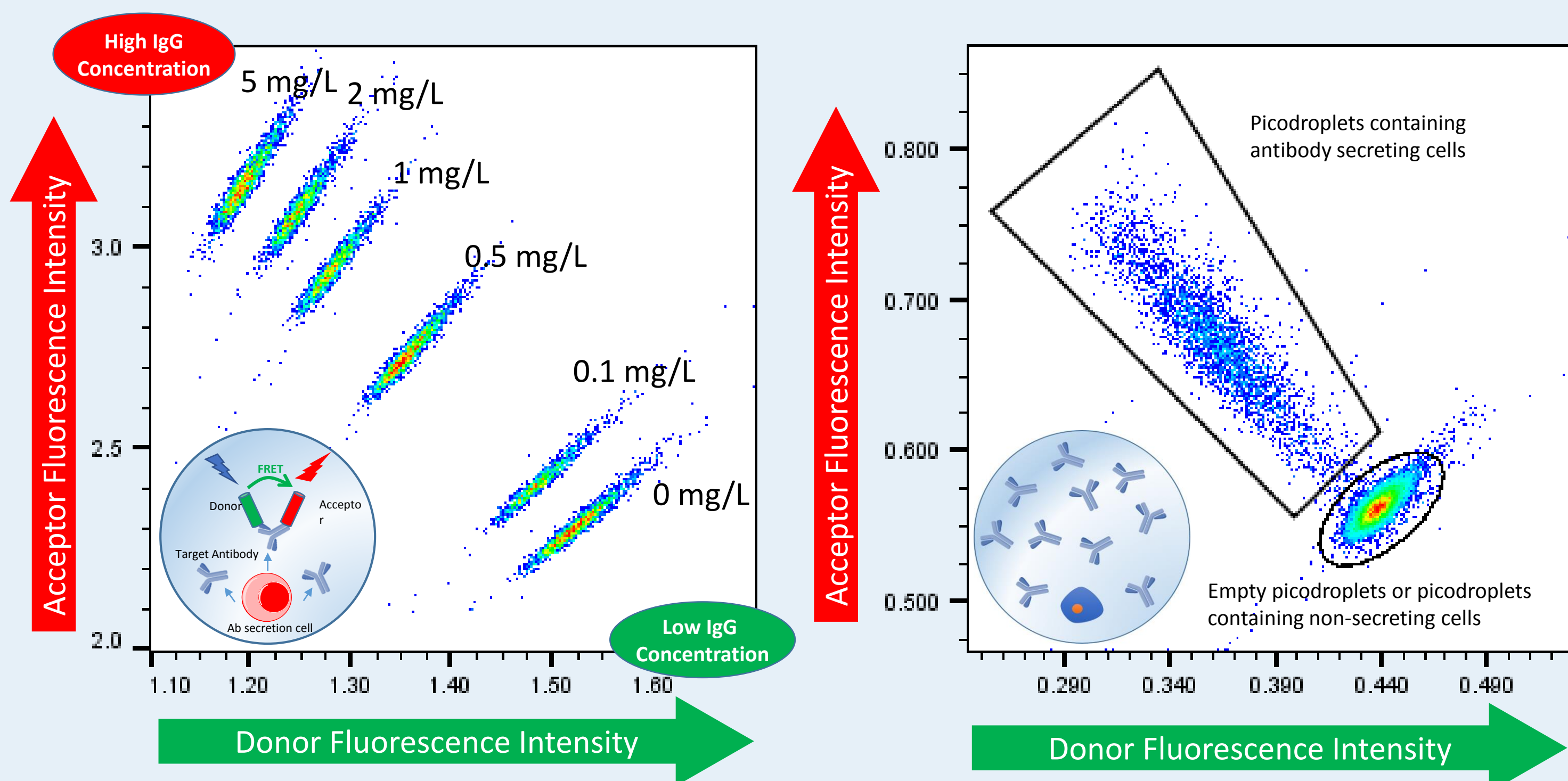


CELL ENCAPSULATION :



The cell number distribution in picodroplets is a factor which can be controlled and optimized by the user. High-quality cell encapsulation is achieved by: 1) accurately counting and adjusting the cell concentration in the medium and 2) minimizing cell sedimentation during the encapsulation process. In an ideal case scenario, the cell number in picodroplets will follow a Poisson distribution.

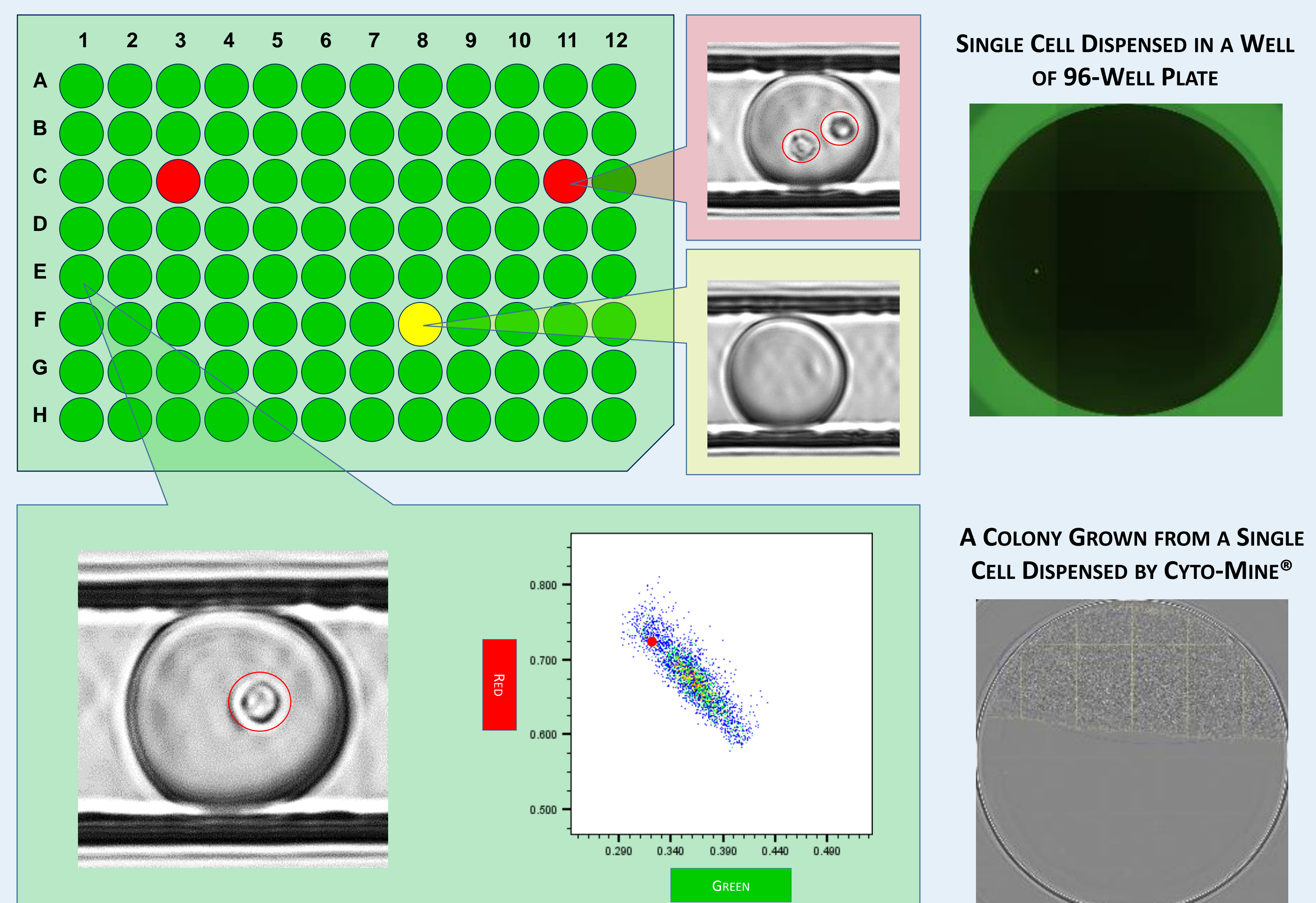
ASSAY AND SCREENING:



Encapsulating single cells in picodroplets enables detection of molecules secreted by the cells (rather than cell surface-bound molecules). Sphere Fluidics has developed a panel of homogeneous FRET assays which allows detection of IgG from various species inside picodroplets. The homogeneous assays can provide information on the endpoint concentration of the antibodies in the picodroplets, essentially, enabling single cell sorting and cloning based on ranking their productivities. The above scatter plot shows results of detection of a library of picodroplets containing different concentrations of target IgG. Picodroplets containing assay reagents and 0, 0.1, 0.5, 1, 2, 5 mg/mL target IgG were generated separately, then mixed at equal ratio.

The size of the picodroplets generated in Cyto-Mine® is only several hundreds of pico-litres in volume, about 5-6 orders of magnitude lower than the volumes in conventional assays. This means, given the sample incubation time, the concentration of secreted antibodies from a single cell in picodroplets is 5-6 orders of magnitude higher than in conventional assay. In Cyto-Mine®, it only takes 0.5-4 hours incubation time before the system can detect antibody secretion from each of the encapsulated cells. In a Cyto-Mine® instrument run, the user can gate and sort a (sub-)population of picodroplets by manually drawing a region of interest on the scatter plot. From 10,000 up to 2,000,000 picodroplets can be analysed in a single Cyto-Mine® experiment run.

SINGLE CELL DISPENSING:



Cyto-Mine® uses a patented technology to dispense each picodroplet containing a cell into 96 or 384-well MTPs, after the screening step. Just prior to the picodroplet being dispensed, the system takes several images of the picodroplet and uses an object recognition algorithm in the software to identify the number of cells inside this picodroplet. A second fluorescent reading on each picodroplet is also carried out at dispensing step. The movement of the MTP is synchronized with each dispensing event and the location of the dispensed picodroplet/cell in the well will also be recorded by the system. After each experimental run, the system will provide a data pack of: 1) a map of monoclonality in the wells, 2) images of each picodroplet prior to dispensing and 3) fluorescence intensity reading for that picodroplet.

The Cyto-Mine® work flow is bio-friendly. No reduction in cell viability is observed after the Cyto-Mine® process, the CHO cells dispensed into wells are still viable and can proliferate into colonies.

CONCLUSIONS:

By encapsulating single cells in picolitre volume water-in-oil droplets, Cyto-Mine® enables rapid detection of molecules (e.g. IgG) secreted by individual cells, followed by selective cloning of single cells based on their productivity. The system also enables monoclonality assurance thus providing a comprehensive solution for accelerated biopharmaceutical discovery and development. The entire systems is also animal origin free, ISO 9001 and GLP-compliant.