Basic Principles in Flow Cytometry

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Flow Cytometry

- » Flow Cytometry is the technical process that allows for the individual measurements of cell fluorescence and light scattering. This process is performed at rates of thousands of cells per second.
- » This information can be used to individually sort or separate subpopulations of cells.

History

- Flow cytometry developed from microscopy. Thus Leeuwenhoek is often cited in any discussion regarding it's history.
- F.T. Gucker (1947)build the first apparatus for detecting bacteria in a **LAMINAR SHEATH** stream of air.
- L. Kamentsky (IBM Labs), and M. Fulwyler (Los Alamos Nat. Lab.) experimented with fluidic switching and electrostatic cell sorters respectively. Both described cell sorters in 1965.
- M. Fulwyler utilized Pulse Height Analyzers to accumulate distributions from a Coulter counter. This feature allowed him to apply statistical analysis to samples analyzed by flow.

History

 In 1972 L. Herzenberg (Stanford Univ.), developed a cell sorter that separated cells stained with fluorescent antibodies. The Herzenberg group coined the term <u>Fluorescence Activated Cell Sorter (FACS).</u>



Cellular Parameters Measured by Flow

<u>Intrinsic</u>

- No reagents or probes required (Structural)
 - Cell size(Forward Light Scatter)
 - Cytoplasmic grabularity(90 degree Light Scatter)
 - Photsynthetic pigments

Extrinsic

- Reagents are required.
 - Structural
 - DNA content
 - DNA base ratios
 - RNA content
 - Functional
 - Surface and intracellular receptors(Immunofluorescence).
 - · DNA synthesis
 - DNA degradation (apoptosis)
 - Cytoplasmic Ca++
 - Gene expression

Flow Cytometry Applications

- Immunofluorescence
- Cell Cycle Kinetics
- Cell Kinetics
- Genetics
- Molecular Biology
- Animal Husbandry (and Human as well)
- Microbiology
- Biological Oceanography
- · Parasitology
- Bioterrorism

• Flow cytometry integrates electronics, fluidics, and optics.

Electronics are involved in signal processing, computer display and analysis.

Fluidics are applied to sample processing and cell sorting.

Optics component are involved in fluorescence detection.













Light Absorption

Quantum mechanics requires that molecules absorb energy as quanta (photons)

Absorption of a photon raises the energy of the molecule from a ground state to an excited state

As molecules relax to a lower energy state, light is released

From: J.Paul Robinson, Purdue University

Fluorescence

Chromophores are the components of molecules which absorb light, they are generally aromatic rings.

Fluorescence lifetimes can be measured in femptoseconds

Quantum Yield measures the efficiency between photons absorbed and photons emitted

J.Paul Robinson, Purdue University



Excitation and Emission Spectra











From Fluorescence to Computer Display Individual cell fluorescence quanta is picked up by the various detectors(PMT's). PMT's convert light into electrical pulses. These electrical signals are amplified and digitized using Analog to Digital Converters (ADC's). Each event is designated a channel number (based on the fluorescence intensity as originally detected by the PMT's) on a 1 Parameter Histogram or 2 Parameter Histogram.

• All events are individually correlated for all the parameters collected.











Running Samples

- · Prepare samples.
- One sample should be completely negative. This sample should be analyzed first. Adjust the Forward Light Scatter and Side Scatter **amplification**. This sample is also used for adjusting the Fluorescence PMTs <u>amplification</u> voltage.
- Adjust the PMT Voltage until you can see a population peak in the first decade of your 1 parameter histogram and or your two parameter plot.
- Once the instrument settings are optimized, run samples and collect data.
- If you are analyzing 2 or more fluorescence parameters you have to prepare Single Color samples for each of your fluorochromes.

Sorting

Lord Rayleigh, liquid stream emerging from an orifice becomes unstable, and breaks up into droplets (1800's)

If a vibration is applied to a stream (emerging from an orifice) the droplet formation becomes stable

R. Sweet develops the drop charging and deflection technique for ink-jet printing (1965)

In cell sorters, an electromagnetic tunable transducer is incorporated in the flow chamber. This causes the fluid stream to break-off into individual droplets

The stream behaves like a wavelength, drops are spaced one wavelength apart



The resulting droplet pattern can be described using the wavelength equation :

v=f \

v is the velocity of the stream

The droplet pattern is most stable when the break-off point is closest to the orifice

This is achieved when the wavelength is 4.5 stream diameters

Substituting Λ for 4.5 stream diameter v = f(4.5d)

f is the vibration frequency

 Λ is the wavelength or droplet spacing

From Howard Shapiro, Practical Flow Cytometry, fourth edition

Sorting

Putting it together:

v=f(4.5d)

For a 75uM orifice and a stream velocity of 20m/s:

f=(20m/s)/(75 X 4.5)

f=(2.0e7)/(337.5)

f=59,259cycles /s or hertz

The ideal frequency for a 75um flow chamber nozzle is 59,259Hz

From Howard Shapiro, Practical Flow Cytometry, fourth edition

Sorting

Droplet charging circuitry

When a cell meets a pre-determined criteria to be sorted, a voltage is applied to the stream

The applied charge will travel down the stream to the last attached drop

The droplet charge is delayed to coincide with the arrival of the cell to the precise position in the stream of the last attached drop

Charged droplets are physically deflected as they pass through a set of two deflecting plates with opposite polarities.

From Howard Shapiro, Practical Flow Cytometry, fourth edition

