#### Introduction to Flow Cytometry

Stephen De Rosa April 20, 2015

#### Outline

- Instrumentation
  - Fluidics
  - Lasers, light detectors and optics
- Fluorescence
- Spectra
- Multi-parameter flow cytometry
- Flow data display
- Basic compensation

## Flow cytometry

- Microscopy "static" examination
- "Flow" cells flow past the "microscope" or the point of interrogation
- Advantage over microscopy is that large numbers of cells can be examined quickly
- Disadvantage is that cells cannot be examined in situ (must be in suspension) and morphology of cells is not examined

# Examples of applications of flow cytometry

- "Phenotypic" identification of cell type; e.g., the percentage of T cells in blood that express CD4
  - Cannot determine absolute count (unless using counting beads); typically determine count in conjunction with CBC data
- Functional examination of cells cytokines, proliferation
- Multiple other applications

#### Instrumentation

- Analyzers
- Vs.
- Cell sorters (also analyze)



Len Herzenberg

# Cell Sorting: Automated Separation of Mammalian Cells as a Function of Intracellular Fluorescence

Abstract. A system for high-speed sorting of fluorescent cells was able to sort mouse spleen cells from Chinese hamster ovarian cells after development of fluorochromasia. Highly fluorescent fractions separated after similar treatment from mouse spleen cells immunized to sheep erythrocytes were enriched in anti-body-producing cells by factors of 4 to 10.

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## Components of a flow cytometer

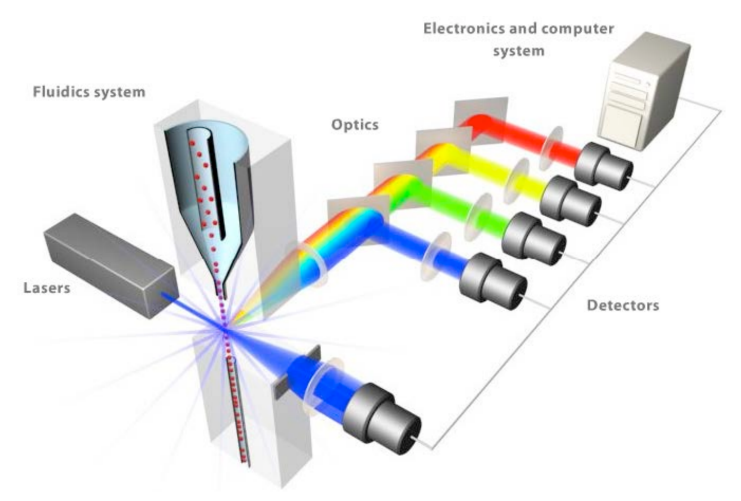
- Fluidics system
- Laser(s)
- Light detectors
- Optical path
- Electronics

(Cell sorting components)

#### **Fluidics**

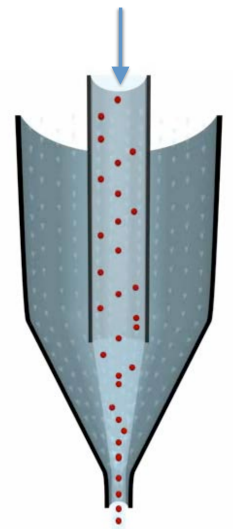
- Fluidics are critical to placing the cell in the correct position to be "viewed" at the point of interrogation
- "Alignment" refers to the focus of the laser beam on the cell in the fluid stream
- Alignment can change either due to laser beam moving or fluid stream moving relative to the laser
  - The latter is more common and can often be corrected by cleaning or priming nozzle

# Flow cytometer 5-parameter, 3-color example



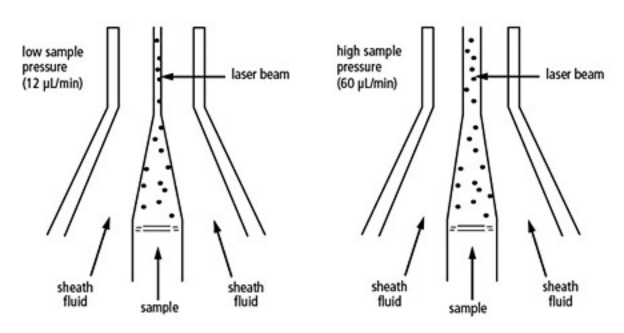
http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

# Hydrodynamic focusing



- Nozzle or flow cell
- Cell stream in center
- Sheath fluid stream surrounds cells and focuses cells
- Single cell per cross section
- Cells pass through laser one cell at a time

#### In some cytometers, cells flow up

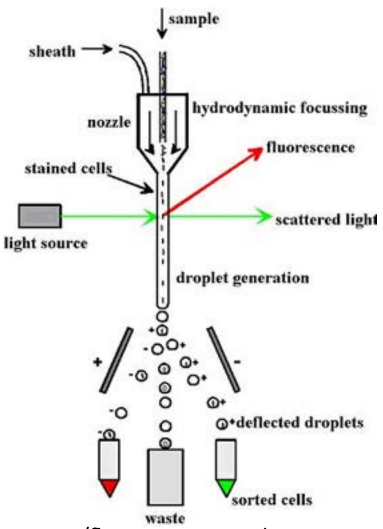


Hydrodynamic focusing of the sample core through the flow cell

## Cell analysis vs. cell sorting

- Some flow cytometers can physically separate or sort cells of interest in addition to analyzing cells
- FACS refers to <u>Fluorescence-Activated Cell</u>
   <u>Sorting</u>
- Each cell to be sorted receives a positive or negative charge and is then deflected from the fluid stream by charged plates
- Same technology as inkjet printers

## Cell sorting

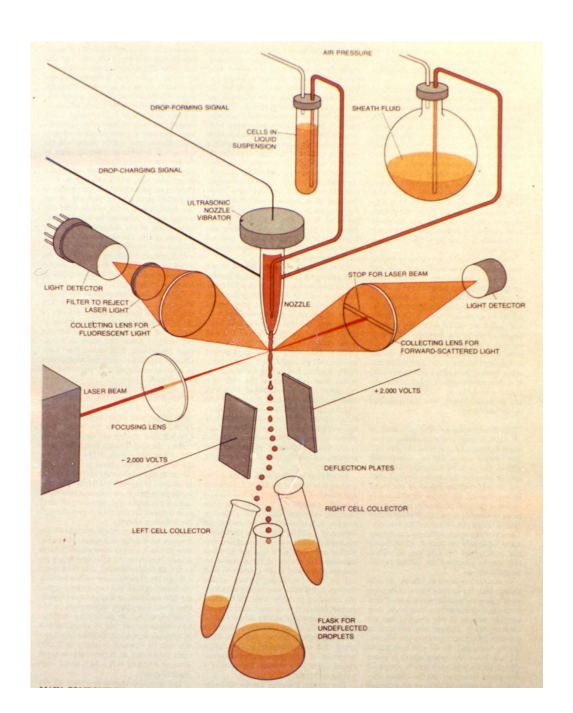


http://www.appliedcytometry.com/flow\_cytometry.php

Leonard A. Herzenberg, Richard G. Sweet and Leonore A. Herzenberg

Scientific American

March 1976



#### Laser light

- <u>Light Amplification by Stimulated Emission of Radiation</u>
- A non-laser light source could be used, but for multi-color flow, light of specific wavelengths is needed
- Also, lasers provide light of sufficient intensity to excite fluorescent dyes to the level necessary to be detectable

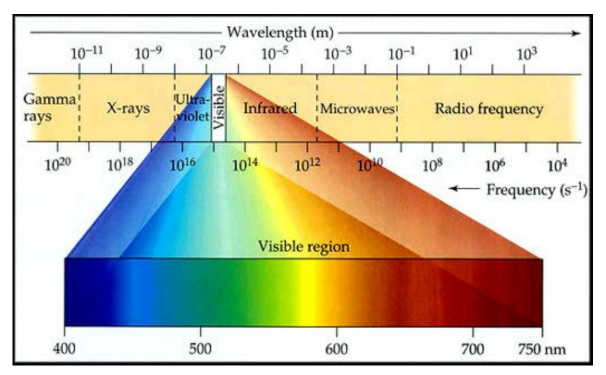
#### Light collector

- Photomultiplier tubes (PMTs)
- Convert light to an electrical signal
- Can be very sensitive, as little as one photon
- Amplifies signal

 Voltage applied to the PMT can be adjusted and determines the signal intensity

#### Brief note on light

- Wavelength determines color
- Higher energy light has shorter wavelengths (blue) and lower energy longer wavelengths (red)



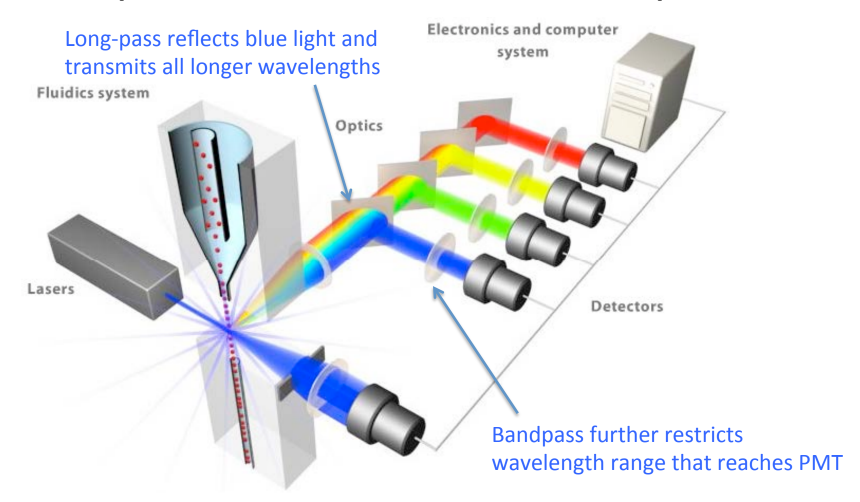
#### Optical path

- Path of laser light directed to the fluid stream
- Path of scattered light from the cell to each PMT
  - Can be quite complex for 20-parameter instruments
- Either can be "direct" and positioned using mirrors or transmitted through optical fibers
- Alignment is critical

#### Light filters

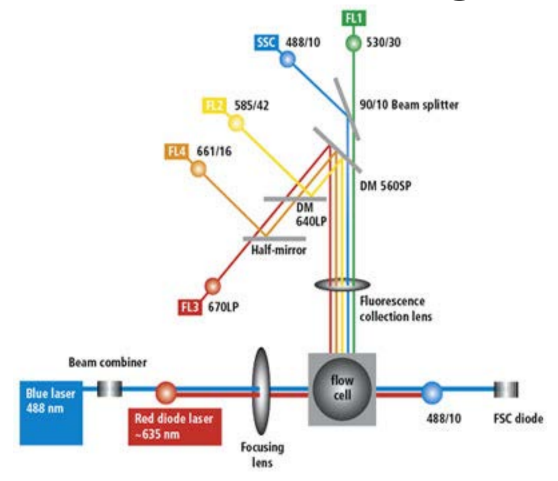
- Restrict light to selected wavelengths
- Each PMT has a bandpass filter so that only light of a selected wavelength range reaches the PMT
- Naming: center/range, e.g., 530/30 passes light of 515 to 545 nm
- Other filters reflect all light above or below a wavelength (long- or short-pass filters)

# Flow cytometer 5-parameter, 3-color example



http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

## **BD FACSCalibur Configuration**



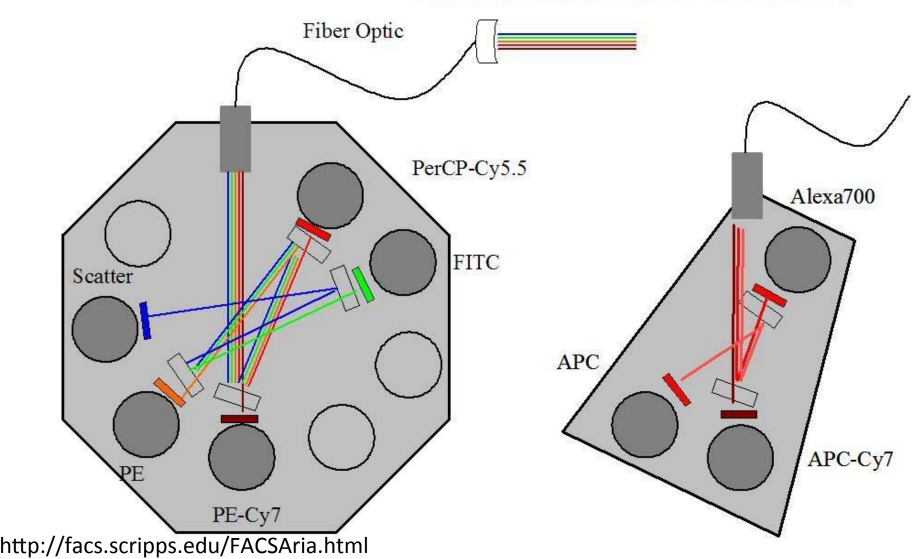
BD FACSCalibur Optical Path Configuration

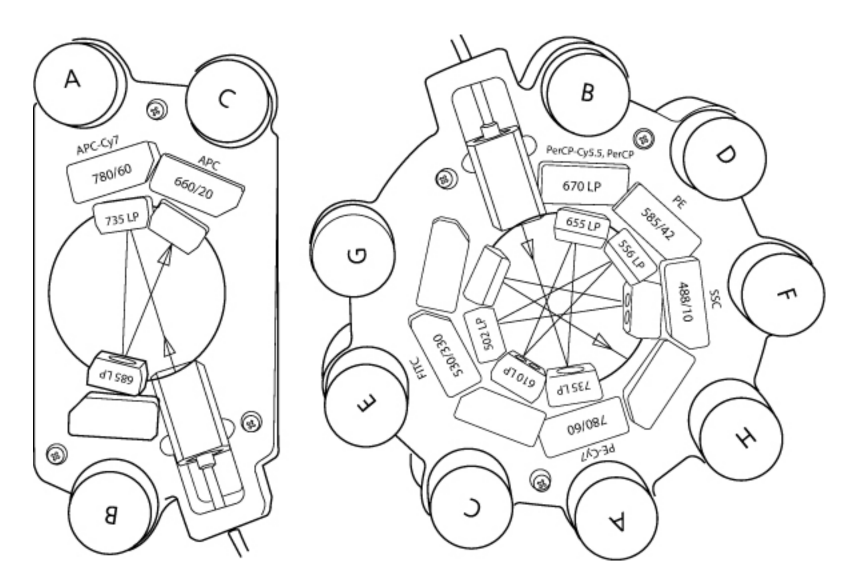
http://www.bdbiosciences.com/instruments/facscalibur/features/index.jsp

# BD Octagons and Trigons

#### The Octagon/Trigon

Reflection is more efficient than Transmission





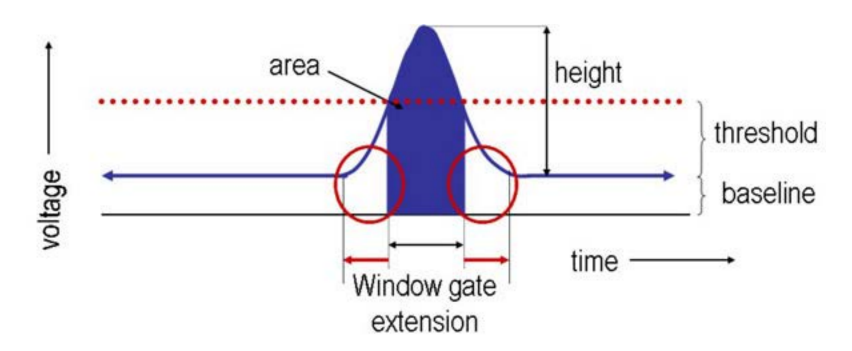
Laser	Laser Power	<u>Channel</u>	Instrument	Bandwidth	Long Pass	Old Detector
		B710	L	710/50	685	
			M	710/50	680	PerCP Cy5.5
			Υ	710/50	685	
Dlive		B515	L	515/20	505	
Blue	488/100mW		M	510/30	505	FITC
			Y	515/20	505	
		SSC	L	488/10	None	
			M	488/10	None	ssc
			L	488/10	None	
Green	532/150mW	G780	M	780/40 780/60	740 725	DE CV7
			Y	780/40	735 740	PE Cy7
			L	None	None	
		G710 G660	М	None	None	PE Cy5.5
			Y	None	None	r L Oys.s
			L	660/40	635	
			M	660/40	635	РЕ Су5
			Y	660/20	640	0,0
			L	610/20	595	
		G610	M	610/20	595	PE Texas Re
			Υ	610/20	600	
			L	575/26	None	
		G575	М	575/26	None	PE
			Υ	575/26	None	
Red	628/200mW	R780	L	780/60	735	АРС Су7
			M	780/60	755	
			Υ	780/60	755	_
		R710	L	710/50	680	
			M	710/50	690	Alx700
			Υ	710/50	685	
		R660	L	660/20	None	
			M	660/20	None	APC
			Υ	660/20	None	
Violet	405/100mW	V780	L	780/60	770	
			M	780/60	770	Qdot800
			Υ	780/60	770	
		V750	L	None	None	
			M	None	None	Not Used
			Y	None	None	
		V710	L	710/40	685	
			M	710/40	685	Qdot705
			Y L	710/40	685	
		V655	M	660/40 660/40	635 635	Odek/ EE
			Y	660/40	635	Qdot655
			L	610/20	585	
		V610	M	610/20	585	Qdot605
			Y	610/20	585	Quotious
		V570	L	575/25	550	
			M	575/25	557	Qdot585
			Y	575/25	557	Guotoco
			L	510/20	495	
		V510	M	510/20	495	Am Cyan
			Υ	510/20	495	
		V450	L	450/50	None	
			М	450/50	None	Pacific Blue
			Υ	450/50	None	
Ultra Violet*	355/20mW -	UV730	L	730/45	685	
			M	730/45	685	
			Υ	730/45	685	
		UV395	L	379/28	None	
			M	379/28	None	
			Υ	379/28	None	
	•					_

# Example configurations for LSR instruments (L, M, Y)

#### What is measured?

# Shape of signal as cell passes through laser

- Typical shape of signal
- Height and/or area is used



#### **Electronics**

- Due to large dynamic range required, logarithmic amplifiers are used on "analog" instruments (data is collected after log transformation)
- Newer "digital" instruments collect data before log transformation

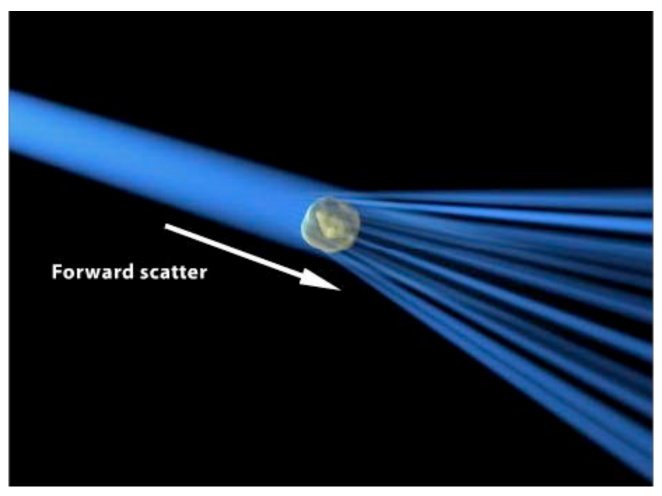
#### What is detected: scattered light

- As cells pass through laser, multiple signals are collected:
  - 1. Scattered light
  - 2. Fluorescence
- Light is scattered by the cell and detectors collect the scattered light
  - 1. Forward scatter
  - 2. Side scatter

#### Forward scatter

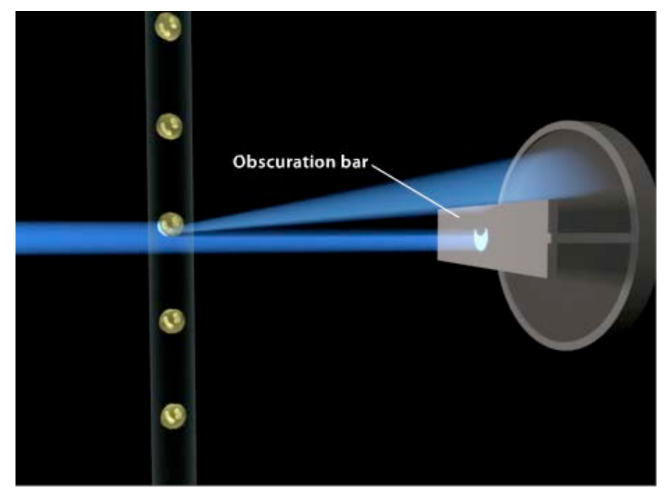
- Often associated with the size of a cell
- E.g., helps to distinguish smaller lymphocytes from larger monocytes

#### Forward scatter (FS)



http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

# Direct laser light must be removed from forward scatter



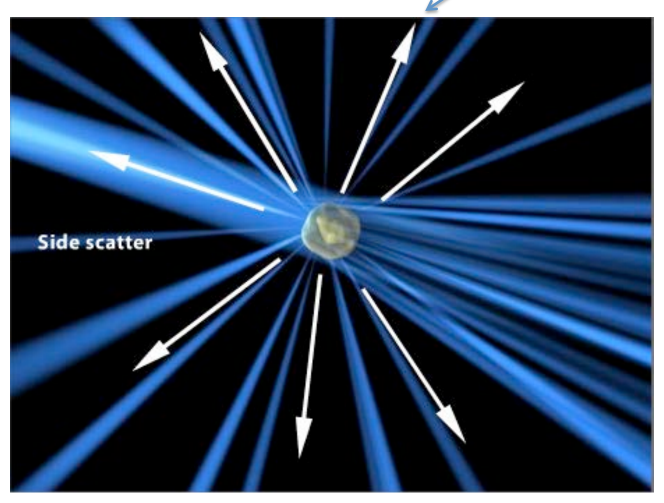
http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

## Side scatter (SS)

- Can be measured at multiple angles, but typically is 90 degrees
- Associated with granularity of cells, e.g., granulocytes are high for SS
- In combination with FS, useful in identifying and excluding dead cells

#### Side scatter

Typically detect at 90 deg

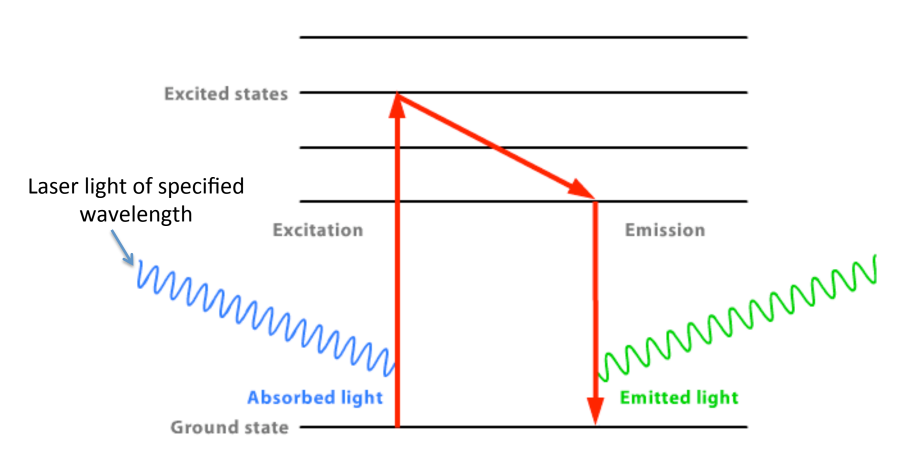


http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

#### What is detected: fluorescence

- As cells pass through laser, fluorescence light from each cell is detected at the appropriate PMT
- Fluorescence from the fluorescent dyes used as tags or markers for cell antigens (but also includes auto-fluorescence)
- The fluorescent light is emitted in all directions, but is only collected and examined from one angle (typically 90 deg)

# Fluorescence: emitted light is of longer wavelength (lower energy)



http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

#### Fluorescent molecules

- Each has excitation and emission characteristics and these can be displayed as spectra
- Commonly used fluorescent molecules have been derived from living organisms or have been created artificially
- E.g., phycobiliproteins are from algae
  - Phycoerythrin, or PE
  - Allophycocyanin, or APC



#### Commonly-used fluorescent dyes

- PE and APC
- Fluorescein or FITC
- Texas red, often as tandem, PE-TR
- Cyanine dyes, often as tandems, PE-Cy7
- Alexa dyes in many colors, e.g., Alx488, Alx655
- Quantum dots
- Pacific Blue
- Propidium iodide (PI)
- "Brilliant" dyes

### Brilliant fluorophores

- Conductive organic polymers
  - Stems from discovery that earned the 2000 Nobel prize in chemistry
- Synthetically tunable network of  $\pi$ -orbitals
  - Many repeat units in a polymer chain
- Extraordinarily high molecular extinction coefficients, similar to PE and APC
- Series of tandem dyes
- Initial series was violet excited, now UV and blue excited

# Fluorescent compound intrinsic properties

- Quantum yield probability of emitting a photon once a photon of light is absorbed (efficiency of energy conversion)
- Molar extinction coefficient probability of absorbing a photon of light at the wavelength of excitation

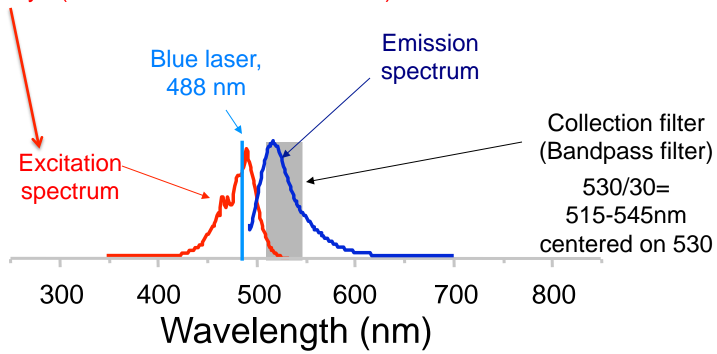
#### Brilliant violet characteristics

Fluorochrome	Quantum Yield	Molar Extinction Coefficienta	Excitation/ Absorbance
Brilliant Violet	0.69	2,500,000	Violet (405 nm)
R-PE	0.82	1,960,000	Blue (496 nm)
APC	0.68	700,000	Red (650 nm)
Quantum Dot 655	≈0.3 <sup>b</sup>	5,700,000	Violet (405 nm)
Quantum Dot 585	≈0.7 <sup>b</sup>	2,200,000	Violet (405 nm)
AlexaFluor 660	0.37	132,000	Red (663 nm)
Fluorescein	0.5	86,000	Blue (488 nm)
AlexaFluor 488	0.92	71,000	Blue (495 nm)
Pacific Blue	0.78	46,000	Violet (405 nm)

<sup>&</sup>lt;sup>a</sup>Measured at the indicated excitation/absorbance wavelength, cm<sup>-1</sup>M<sup>-1</sup> <sup>b</sup>Quantum yields generally increase for larger Quantum Dots

#### **Spectra of Fluorescein**

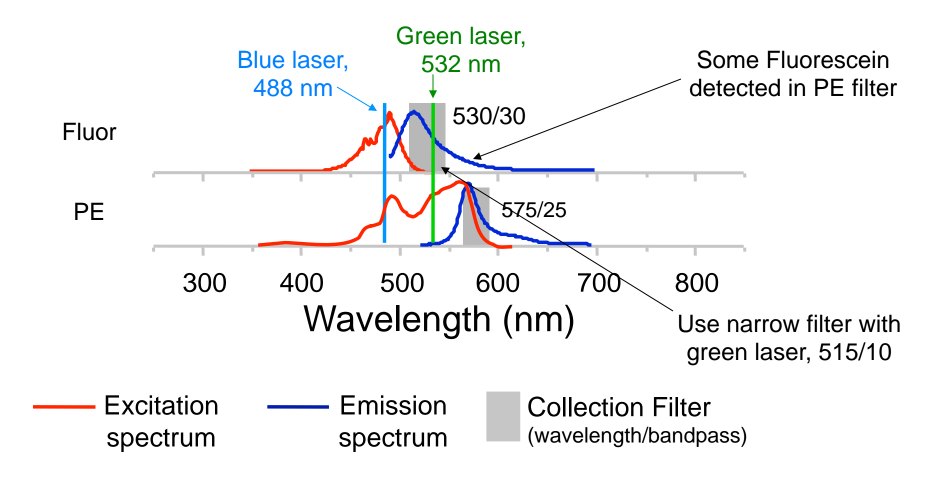
Excitation spectrum shows range of wavelengths that can excite dye (determines which laser to use)



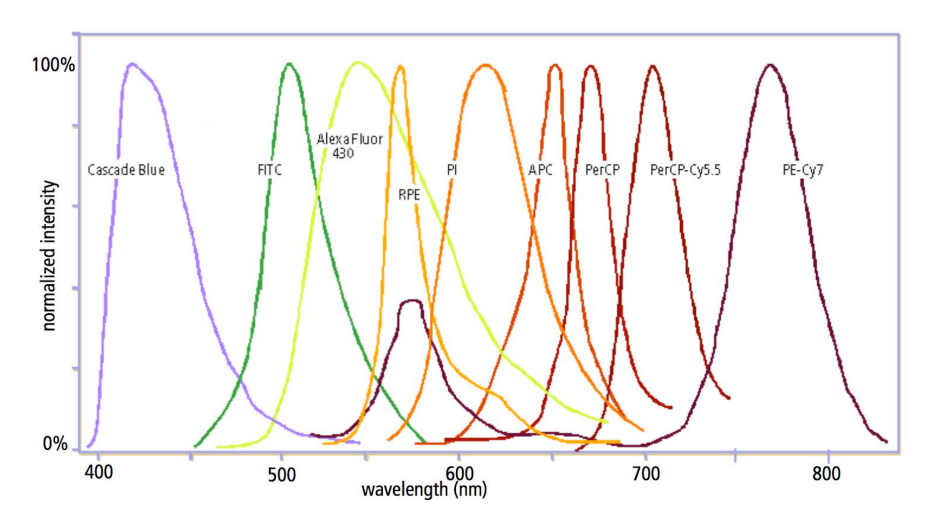
Useful resources = http://www.bdbiosciences.com/spectra/ http://probes.invitrogen.com/resources/spectraviewer/

#### Spectra of dyes: 2-color example

Different emission spectra allow signals to be distinguished



#### Emission spectra of common dyes



http://www.rockefeller.edu/fcrc/pdf/ 642221 BD LSRII Users Guide.pdf

#### Cell identification

### Specific cell labeling

- Flow cytometry was made feasible because of two technologies: lasers and monoclonal antibodies
- Monoclonal antibodies are "specific", i.e., bind to only a single antigenic determinant
- Antibodies are an important component of the immune system, but for flow, they are simply used as a tool
- For human studies, antibodies to human antigens are produced in other species (mouse, rat)

#### Cell antigens

- Antigens refer to "determinants" that can induce an antibody response
- Different cell types often express different antigens
- For the purpose of flow these antigens are simply used as "markers" even though they likely have important functions for that particular cell type

#### Fluorochrome-antibody reagents

- Monoclonal antibodies to the marker of interest provide specificity
- To allow visualization in flow, the antibody must be labeled with a fluorescent dye
- Many different methods for attaching the fluorescent dye to the antibody have been developed (typically through covalent chemical bonds)
- These are often referred to as <u>antibody-dye</u> conjugates or <u>reagents</u>

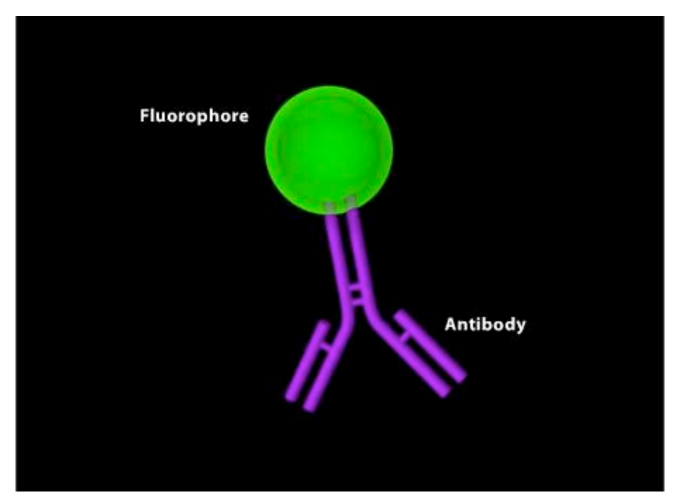
#### Fluorescent labeling of cells

- Specificity is typically provided by monoclonal antibodies specific for selected cell antigens (such as CD4, CD8...)
- Antibodies of different specificities are tagged or labeled with fluorescent molecules of different "colors"
- Cells are labeled by reaction with the antibodydye reagent ("staining"). Any reagent that does not stick to the cell is washed off.

#### Note on nomenclature

- Antigens of interest on cells often have been assigned "CD" numbers (cluster of differentiation)
- These antigens may have been chosen as the marker of interest because of a function they perform, or simply because they have been shown to label a cell of interest

# Example of larger fluorophore attached to antibody

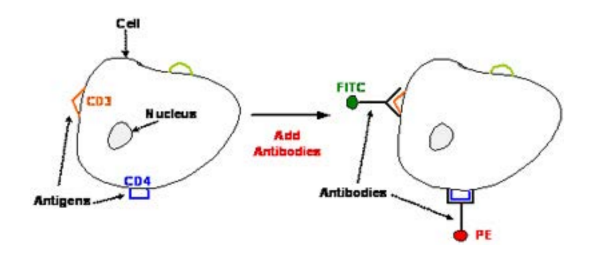


http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

# Multi-parametric flow cytometry

- Multiple measurements on each cell (in addition to forward and side scatter)
- A single marker is often insufficient to identify cell of interest
- A common example is measurement of T cells
  - Three markers typically used
  - CD3 is one component of the T cell receptor and therefore marks all T cells
  - CD4 and CD8 mark the two major types of T cells

# Example of CD3 and CD4 staining



# Multi-parametric flow cytometry requirements

- Dyes of different "colors" used in combination
- Must be able to distinguish the signals from each of these dyes
- Choose dyes that emit light of different colors
- Use different detectors for each dye and only collect light of the appropriate color in each detector

#### Nomenclature

- Each different dye in a multi-parameter flow assay is often referred to as a "color", such as an "8-color" experiment
- On the flow cytometer, each color requires a light path and light detector and this is often referred to as a channel
- The color and/or channel may be named using the name of the fluorescent dye such as the FITC or PE channel or simply numbered (FL1)
  - Generic naming by laser color and center of bandpass filter, e.g., B515, G575

## Why multiple lasers

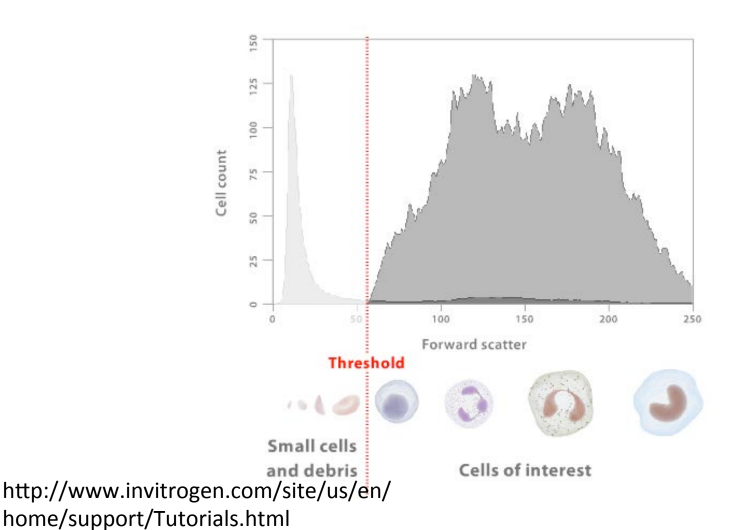
- Each laser type emits light of a specific wavelength(s)
- Multiple lasers allow an increase in the total number of colors or markers that can be examined in each assay (because there is greater choice in dyes)
- Dyes that are excited at different wavelengths can be used in combination (even if they emit at similar wavelengths)

# Flow data display

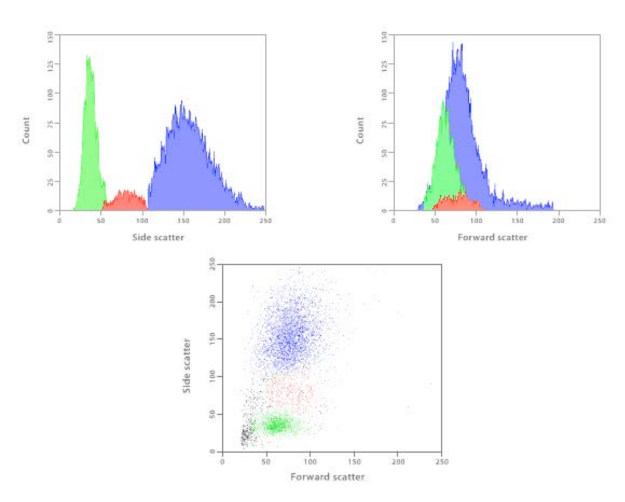
### Flow data display

- Single dimensional histogram
- Two-dimensional
  - Dot plot
  - Contour plot
  - Pseudo-color plot

# Histogram display: distribution of number of cells (y-axis) vs. signal intensity (x-axis) (forward scatter example)

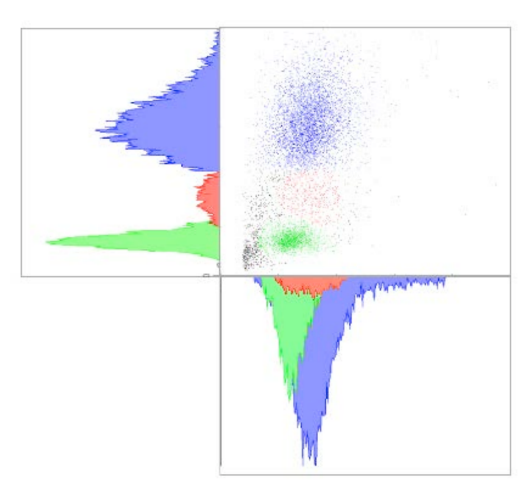


#### Two-dimensional plots vs. histograms



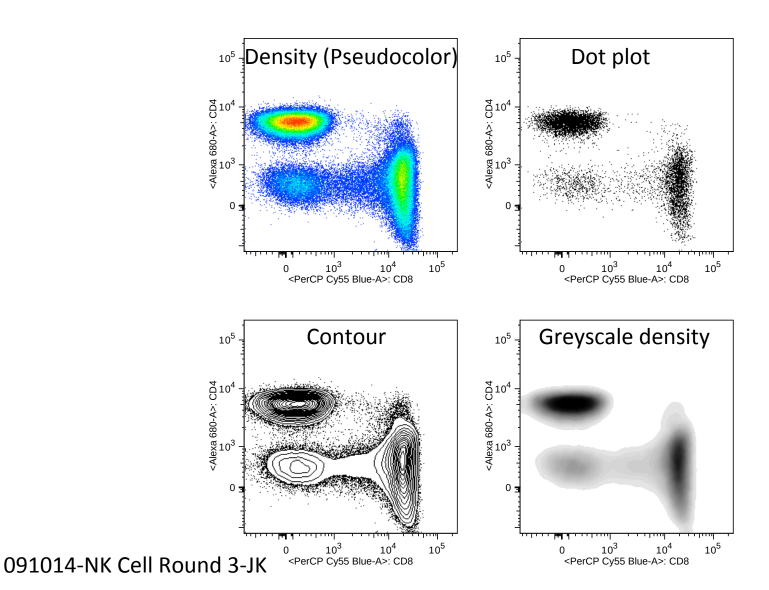
http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

### Visualizing two dimensions



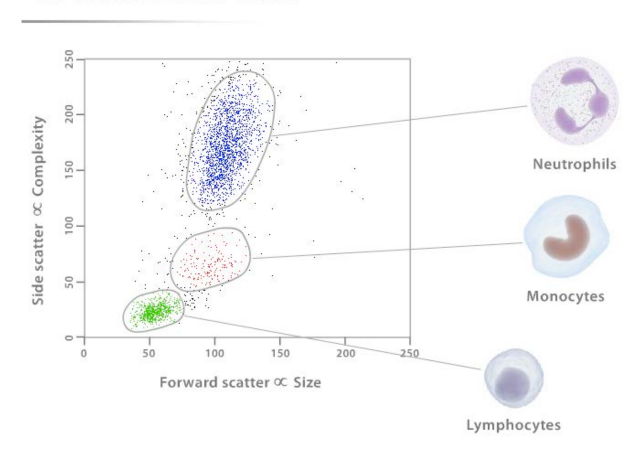
http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

# Display options



# Even scatter alone (without fluorescence signal) can identify cell types

2D Scatter Plot of Blood



http://www.invitrogen.com/site/us/en/home/support/Tutorials.html

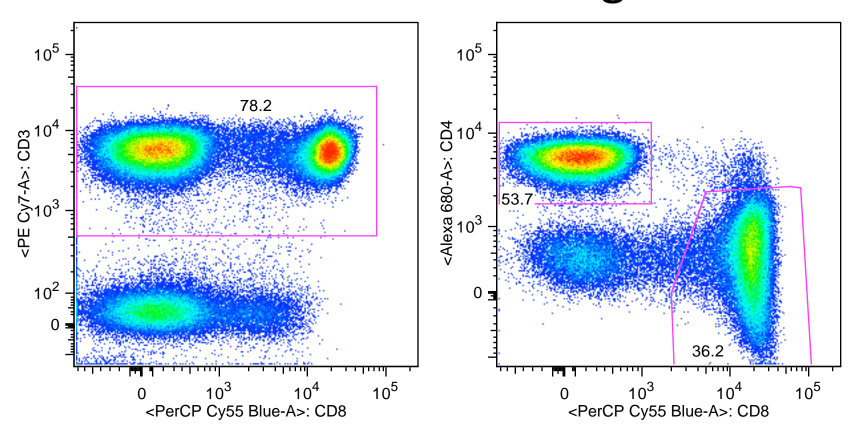
### Flow data analysis

- A "gate" is used to select a group of cells of interest
- It is typically a rectangle or polygon
- Typically, multiple gates are used and often in a hierarchical format (a gating tree) in which each new gate is a subset of cells in the prior gate

# Gating tree example in Flow Jo

$\psi \ \overline{\mathbb{T}}$	00.0.00		218184	NonComp	SAC 214404	DMSO
₩	<b></b> <i>s</i>	96.4	210294			
₩	€ Live	99.6	209549			
₩	€ L	70.4	147586			
₩	€ 3+	77.6	114523			
₩	€ 4+	30.6	35088			
	€ 107	0.023	8			
	IFNg+	0.046	16			
	<b></b> IL2+	0.054	19			
	TNF+	0.063	22			
₩	€ 8+	56.1	64267			
	<b>€</b> 107		98			
	IFNg+	0.025	16			
		0.011	7			
	TNF+	0.067	43			
$\psi \ \overline{1}$	374094.fcs		212640	NonComp	SAC 200087	ENV
₩	<b>ⓒ</b> <i>S</i>	94.6	201253			
▼		91.2	183454			
₩	€ L	65	119265			
₩	€ 3+	76.2	90916			
₩	<b>€</b> 4+	51.3	46623			
	€ 107	0.015	7			
	IFNg+	0.015	7			
	€ IL2+	0.056	26			
	TNF+	0.06	28			
₩	€ 8+	37.5	34096			
	<b>€</b> 107		16			
	IFNg+	0.018	6			
	€ IL2+	0.026	9			
	₹ TNF+					

# Gating example: T cells identified as CD3+, then CD4+ and CD8+ cells gated



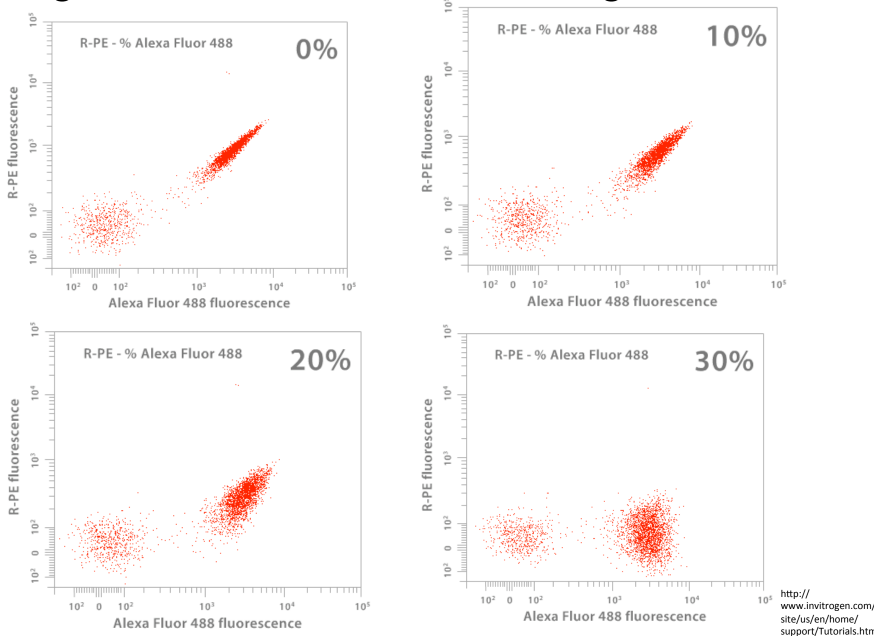
#### Compensation

- Fluorescent signals from different dyes overlap so that the signal from dye 1 is detected in the signal from dye 2, and/or viceversa
- Mathematically, this can be corrected by subtracting out the part of the signal in the dye 2 channel that is due to dye 1
- Single-stained controls are used to determine compensation percentages

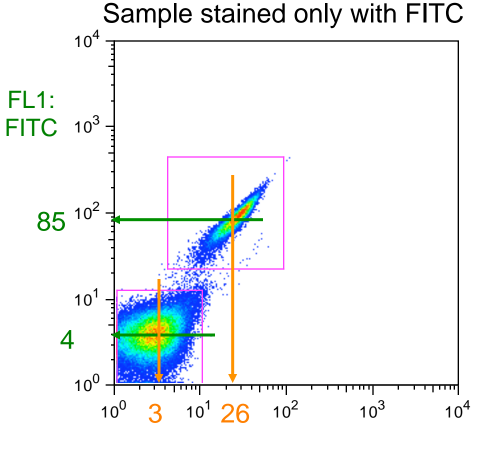
#### Compensation

- Spectral overlap between dyes results in the detection of the fluorescence from one dye in one or more other detectors (primary vs. spillover fluorescence)
- The spillover fluorescence must be subtracted from the total fluorescence detected in the secondary detector
- This spillover fluorescence is proportional to the level of the primary fluorescence
- For each cell, a specified percentage of the primary fluorescence is subtracted from the total fluorescence detected in the secondary detector

Single stain Alx488: subtract signal in PE



#### FITC into PE



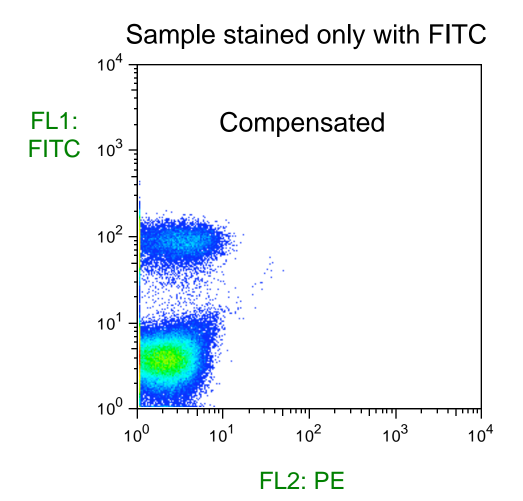
# % spillover of FITC into PE=

$$\frac{\text{MFI}_{\text{FL2}}(\text{pos}) - \text{MFI}_{\text{FL2}}(\text{neg})}{\text{MFI}_{\text{FL1}}(\text{pos}) - \text{MFI}_{\text{FL1}}(\text{neg})} \times 100$$

$$\frac{26-3}{85-4}$$
 x100 = 28%

FL2: PE

#### FITC into PE



# % spillover of FITC into PE=

$$\frac{\text{MFI}_{\text{FL2}}(\text{pos}) - \text{MFI}_{\text{FL2}}(\text{neg})}{\text{MFI}_{\text{FL1}}(\text{pos}) - \text{MFI}_{\text{FL1}}(\text{neg})} \times 100$$

$$\frac{26 - 3}{85 - 4} \quad x100 = 28\%$$

# Advanced topics in Fluorescence Spillover Compensation

Mario Roederer, Cytometry 45:194-205 (2001)

Spectral Compensation for Flow Cytometry: Visualization Artifacts, Limitations, and Caveats

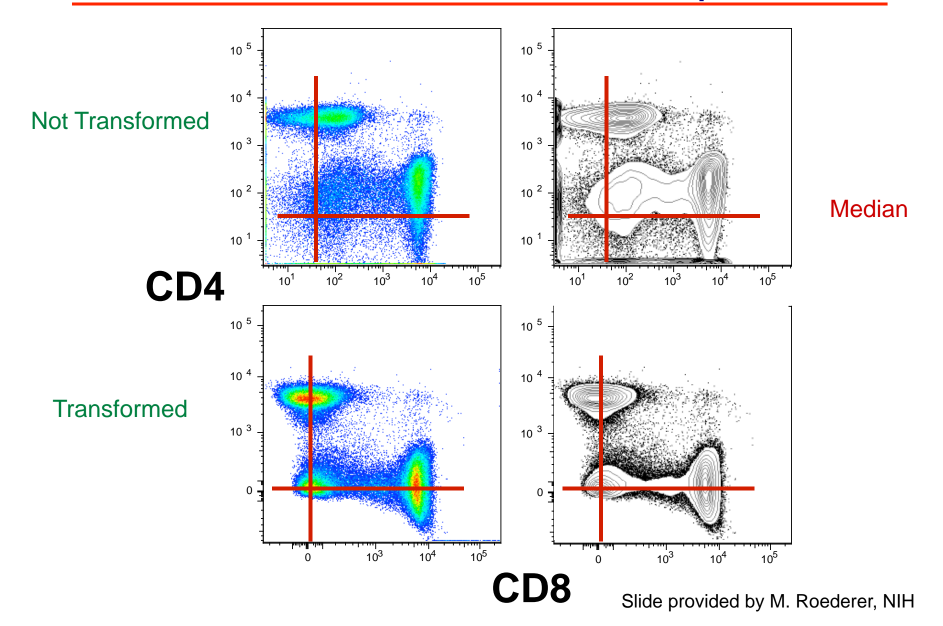
Nicole Baumgarth and Mario Roederer Journal of Immunological Methods 243:77-97 (2000)

A practical approach to multicolor flow cytometry for immunophenotyping

#### Bi-exponential or logicle transformation

- Transforms the log scale to display values below zero
- This allows better visualization of populations centered around zero
- This display feature is now available in most FACS analysis software

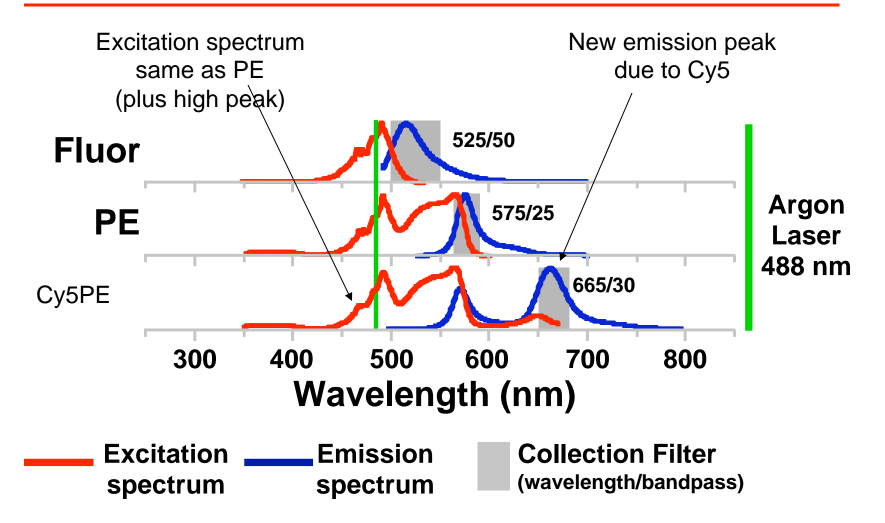
#### Transformation Confirms Compensation



#### **Tandem dyes**

- Dyes covalently linked so that energy from one dye is transferred to the other (resonance energy transfer)
- Ideally, when the dye is excited, it emits mainly at the wavelength characteristic of the second dye (with little emission from the first dye)
- Examples: PE-Cy5, PE-TR (ECD), PerCP-Cy5.5, PE-Cy7, APC-Cy7

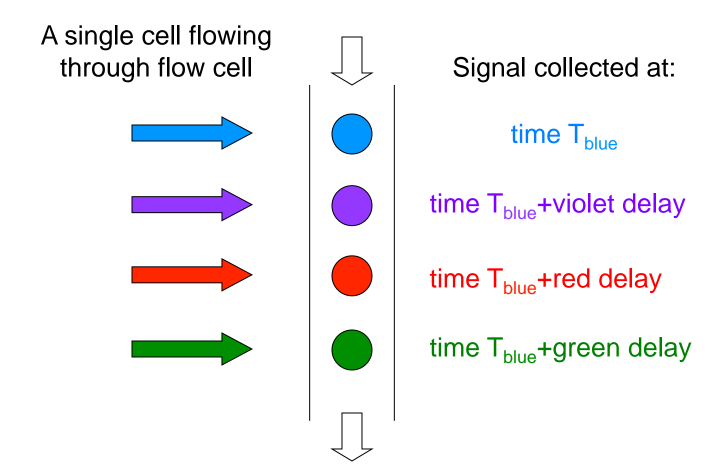
#### Spectra: 3-color example (one laser)



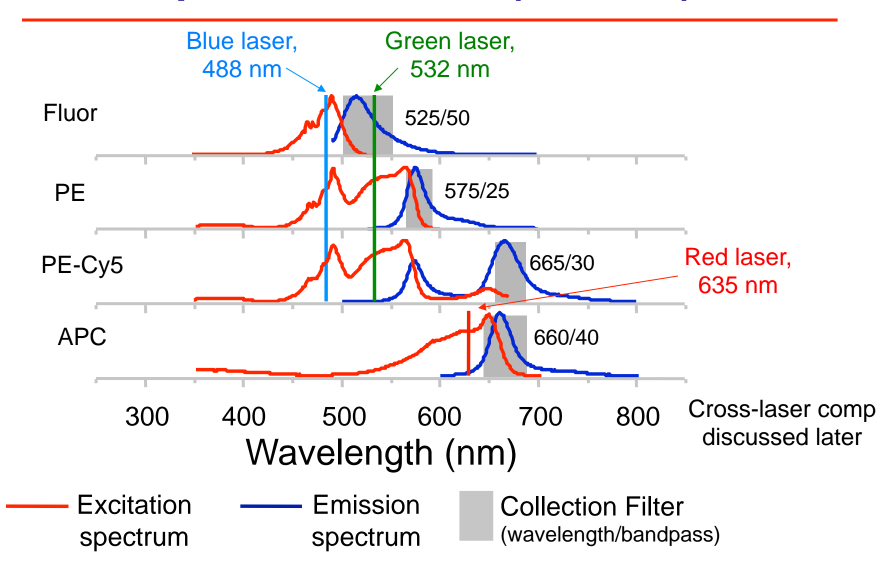
#### **Additional lasers**

- The addition of a second (or third or fourth) laser allows for use of dyes excited at another wavelength(s)
- There are time delays for different lasers, so even if dyes have similar emission spectra, the signals can be distinguished by time
  - •But, if a dye is excited by more than one laser, then "cross-laser" compensation is required

#### Laser Delay



#### Spectra: 4-colors (2 lasers)



#### Learning points

- What are the parts of a flow cytometer?
- What is a flow cytometer used for?
- How are fluorescent dyes and monoclonal antibodies used in flow cytometry?
- How are flow cytometric data displayed?
- What limits the number of colors or parameters measured?