

# 流式細胞儀 --- FACSCanto 之基本原理與應用

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## 大綱

- 實驗設計原理
- 流式細胞儀運作原理

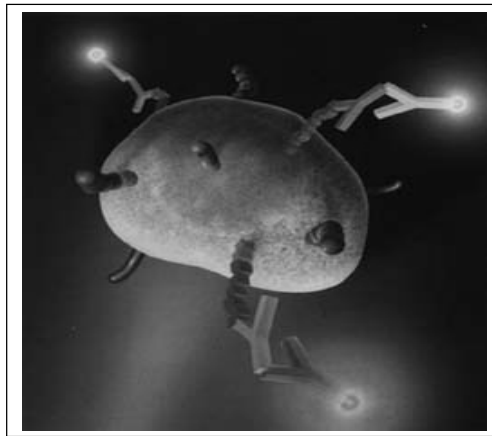


## 實驗設計原理

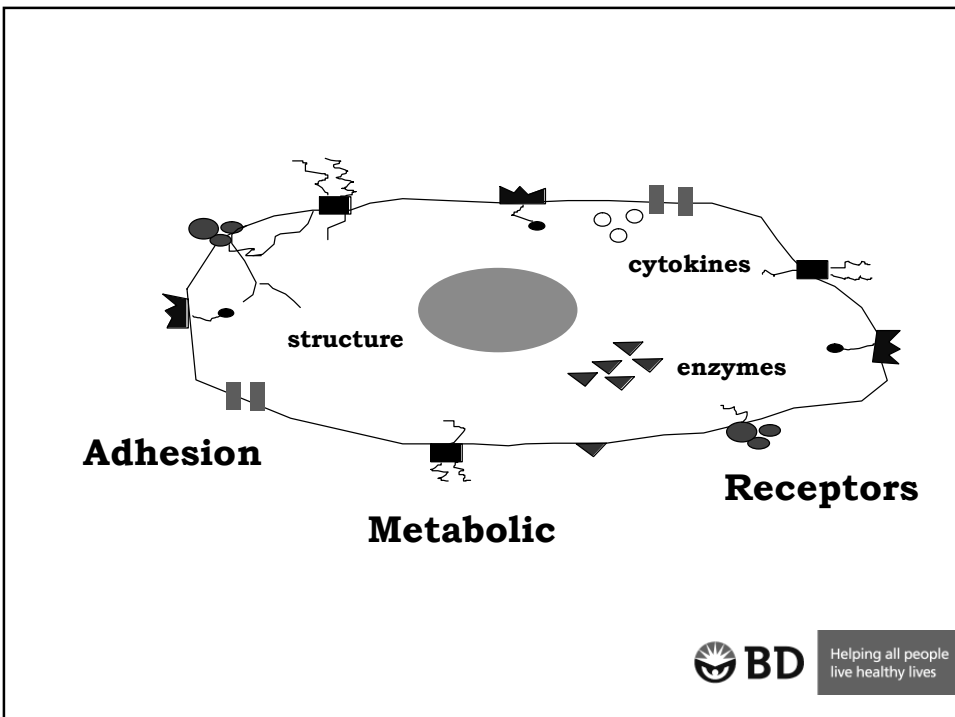
- 藉由螢光抗體標識細胞之抗原特性
- 藉由螢光化合物標識細胞特性
- 藉由螢光染劑標識細胞特性
- Cytometry Beads Array



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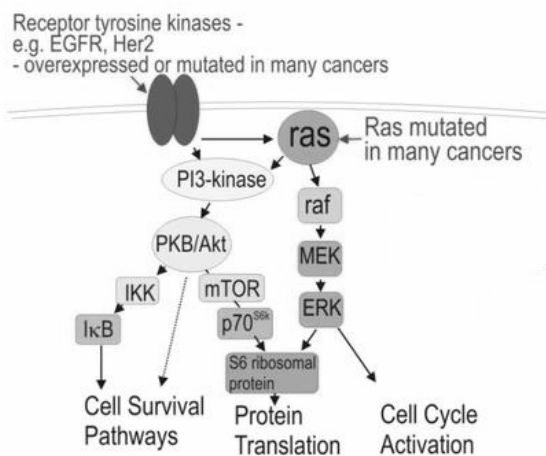


### 淋巴球免疫分型

FITC	PE	使用原理
CD45	CD14	儀器可自動依據 CD45 與 CD14 之免疫螢光染色，測量三群白血球的比例，淋巴球 (CD45 Bright /CD14-/Low FSC/ Low SSC) ，單核球 (CD14+/CD45 Intermediate/High FSC/Intermediate SSC) ，顆粒性球 (CD14 dim/CD45 dim/High FSC/High SSC) ，細胞碎片與紅血球應是 (CD14-/CD45-/Low FSC/Low SSC) 。
IgG1	IgG1	根據陰性對照組之染色程度來劃分陽性/陰性之界線，依此界線在陰性對照的樣品中，假陽性(Quadrant1+2+4)不得超過百分之二。
CD3	CD19	T 細胞必須表達有 CD3+ 抗原，B 細胞表達有 CD19+ 抗原。此兩群細胞與 NK 細胞之總和應約略等於淋巴球的總數，可作為品管的指標。
CD3	CD4	CD4+ T 細胞必須同時表達有 CD3+ 與 CD4+ 抗原。
CD3	CD8	CD8+ T 細胞必須同時表達有 CD3+ 與 CD8+ 抗原。
CD3	CD16+ or CD56+	NK 細胞必須不表達有 CD3 抗原，同時表達有 CD16+ 或 CD56+ 抗原。NK 細胞與 T & B 兩群細胞之總和應約略等於淋巴球的總數，可作為品管的指標。

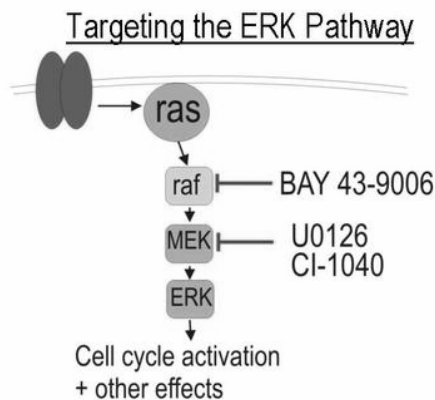
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## 訊息傳導



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## 治癌藥物



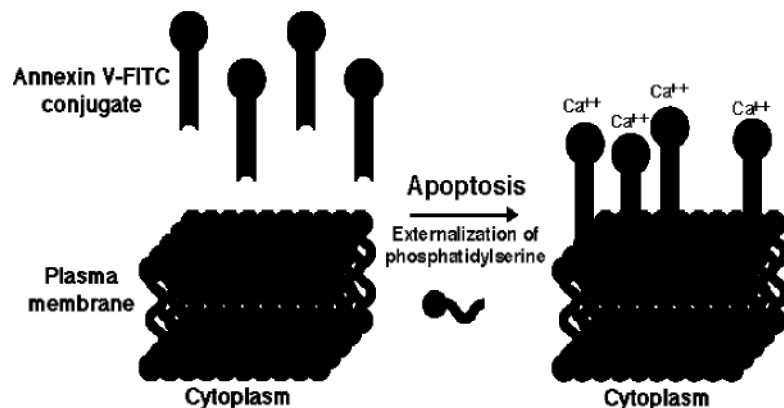
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## 實驗設計原理

- 藉由螢光抗體標識細胞之抗原特性
- 藉由螢光化合物標識細胞特性
- 藉由螢光染劑標識細胞特性
- Cytometry Beads Array



## Annexin V Assay



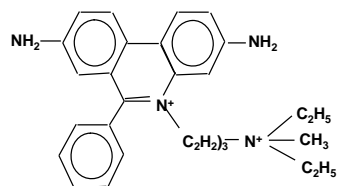
## 實驗設計原理

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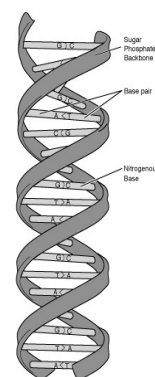
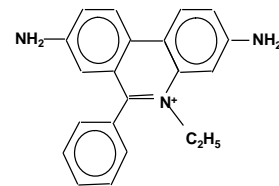


## DNA 特異性染劑

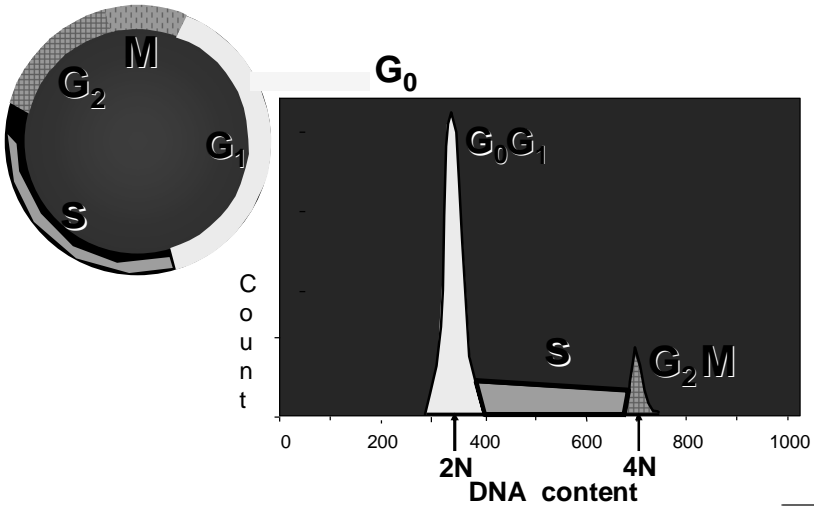
Propidium



Ethidium

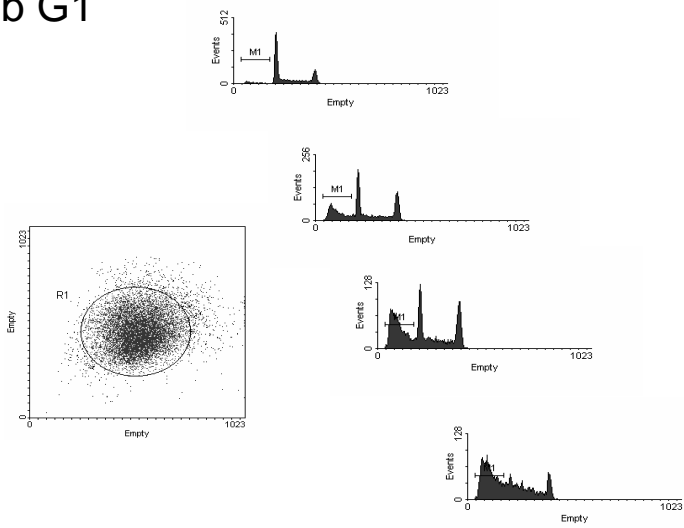


## 細胞周期位相的決定



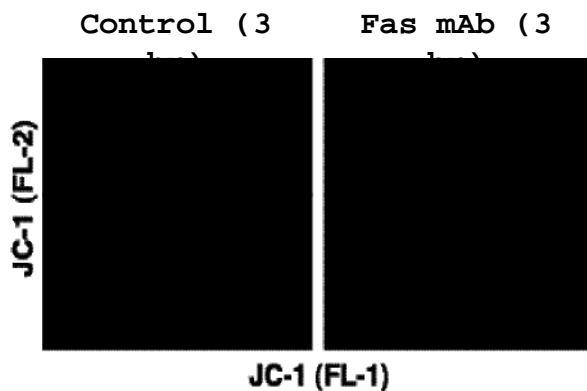
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## Sub G1



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## Mitochondria Membrane Potential



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## 細胞增殖反應

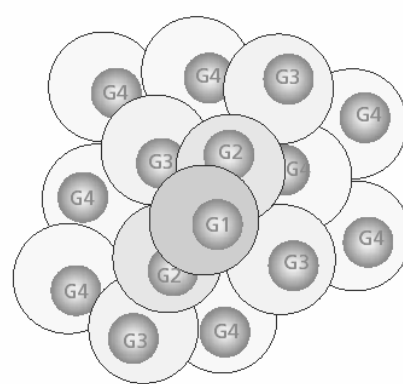


Figure 1. Upon cell division, CFSE is distributed uniformly between daughter cells.



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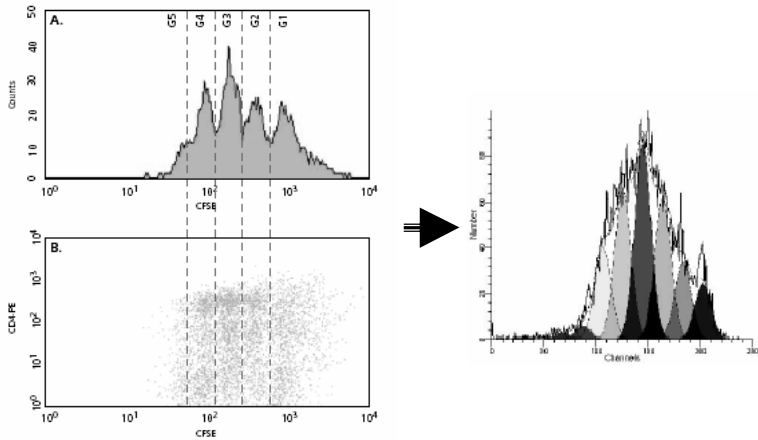


Figure 2. Profile of CSFE-labeled, PHA-stimulated (72 hrs), peripheral blood mononuclear cells (PBMCs) analyzed by flow cytometry.



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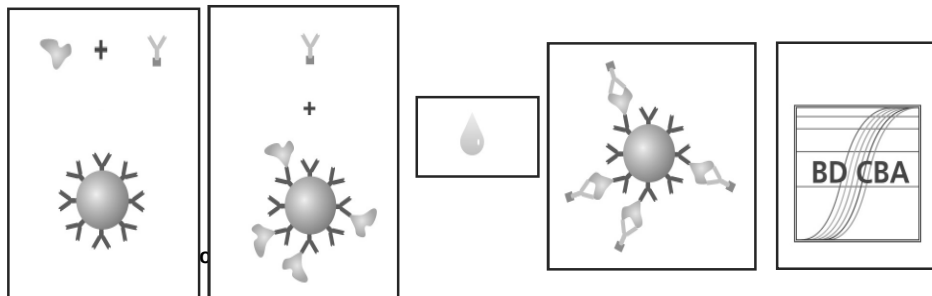
## 實驗設計原理

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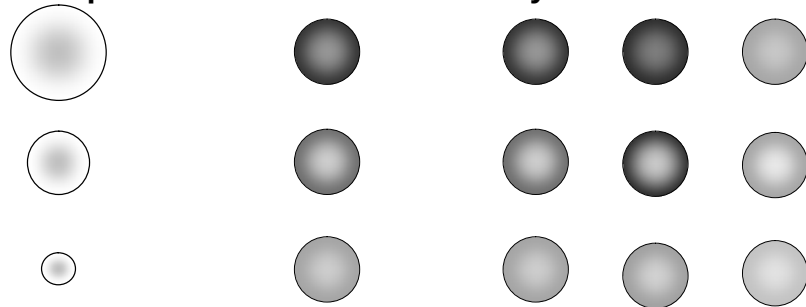
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## Cytometric Beads Array (CBA)



## Beads Provide a Flexible Platform

**Beads provide an expandable assay platform for use with a flow cytometer**



**Multiple sizes**

**Different intensities\***

**Different colors with different intensities**

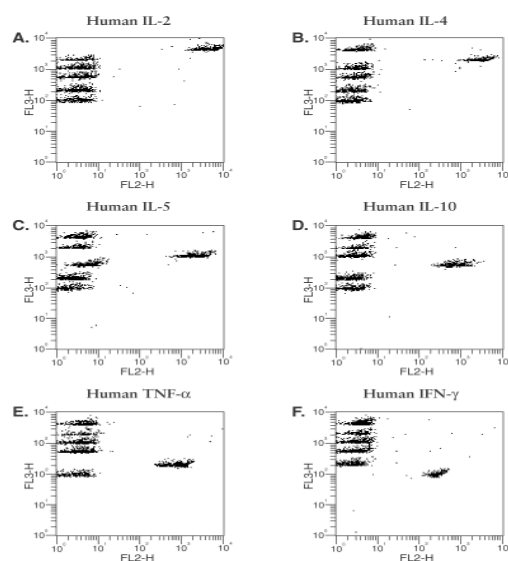


## Advantages of Bead-Based Immunoassays

- Analyze multiple analytes simultaneously
- Reduced sample volume requirements
- Reduced hands-on time by parallel analysis of samples
- Wide dynamic range of fluorescence detection (requires fewer sample dilutions)



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### Proteins Measured

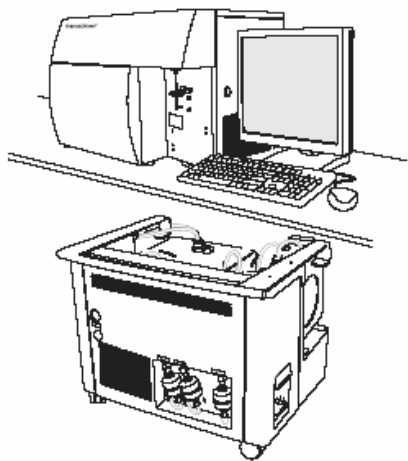
- A. Interleukin (IL)-2
- B. IL-4
- C. IL-5
- D. IL-10
- E. Tumor Necrosis Factor- $\alpha$
- F. Interferon- $\gamma$



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# BD FACSCanto™

Figure 1-1 BD FACSCanto system



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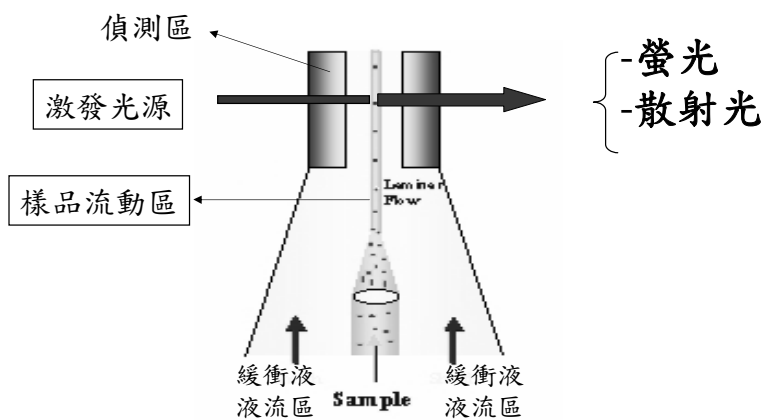


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## 流式細胞儀的工作原理



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## 流式細胞儀能測量：

- 散射光
  - 細胞大小 (前方散射光)
  - 細胞折射率 (側方散射光)
- 各色螢光

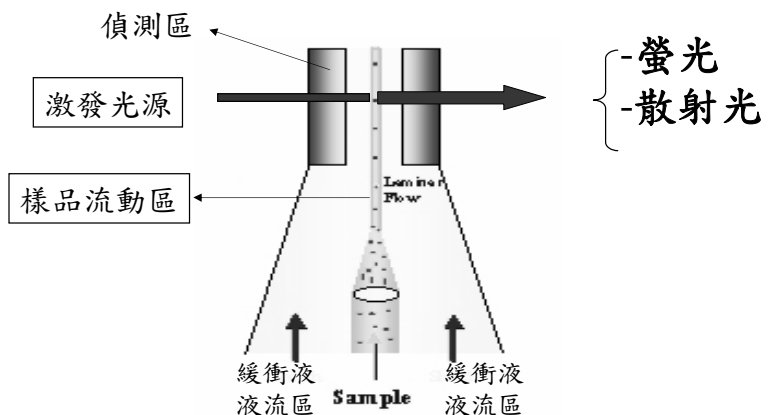


## 綜合三個系統的功能：

- 液流系統:將細胞依序送到測量區受檢。
- 光學系統:產生並收集螢光、光散射等信號。
- 電子系統:
  - 將光學訊號轉換成電子訊號。
  - 分析所輸出的電流訊號，以脈衝高度、寬度、積分面積顯示。
  - 量化訊號並傳至電腦。

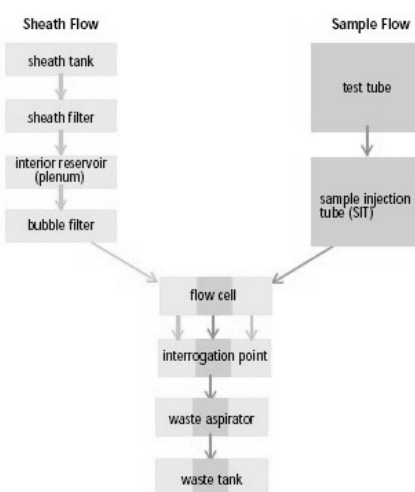


## 液流系統



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## FACSCanto 的液流系統



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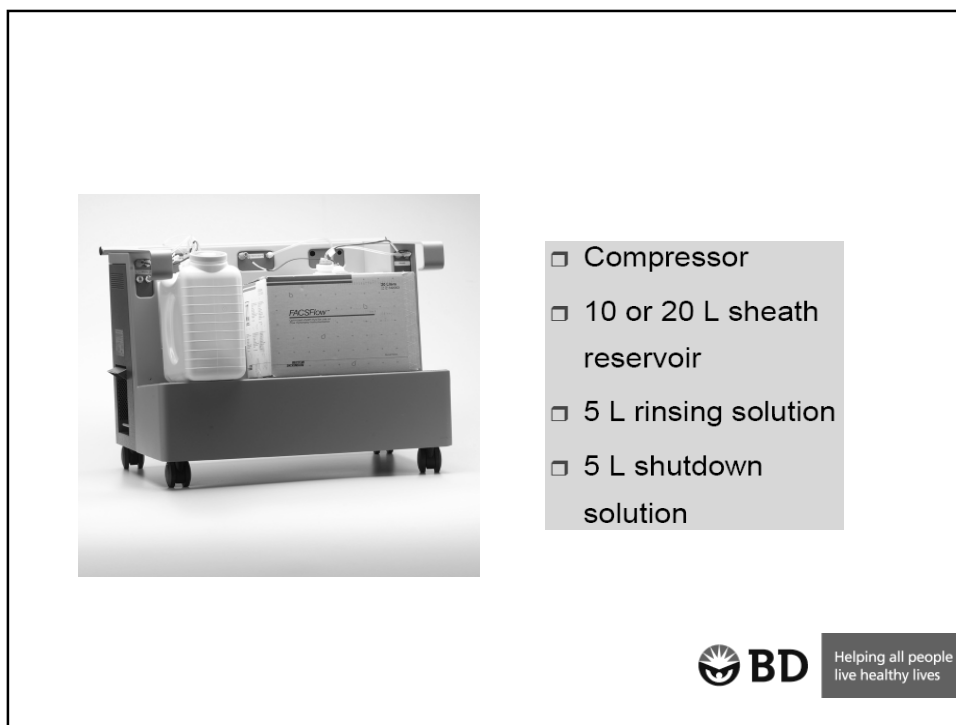
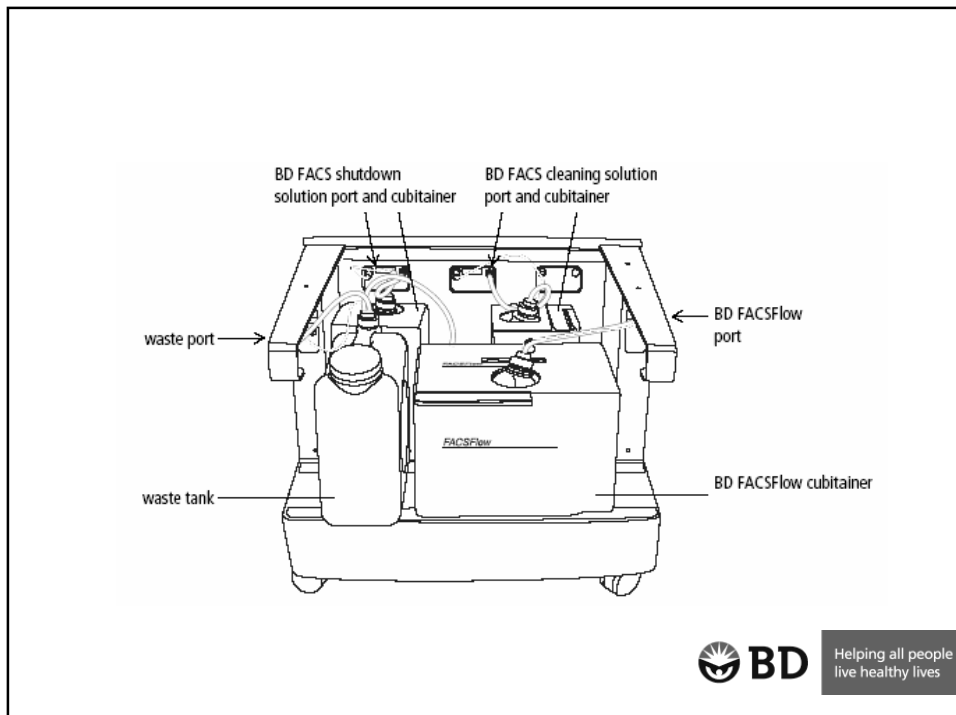
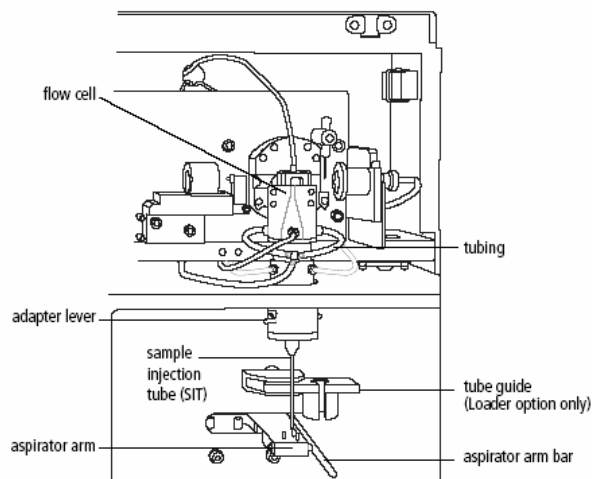


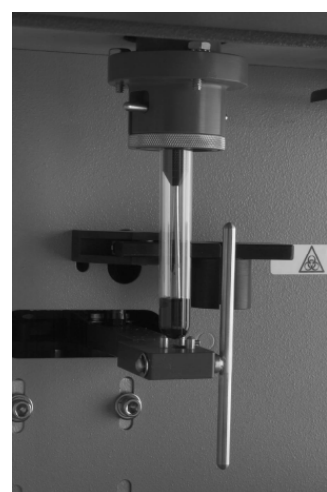
Figure 1-2 Sample injection tube



10,000 events/sec



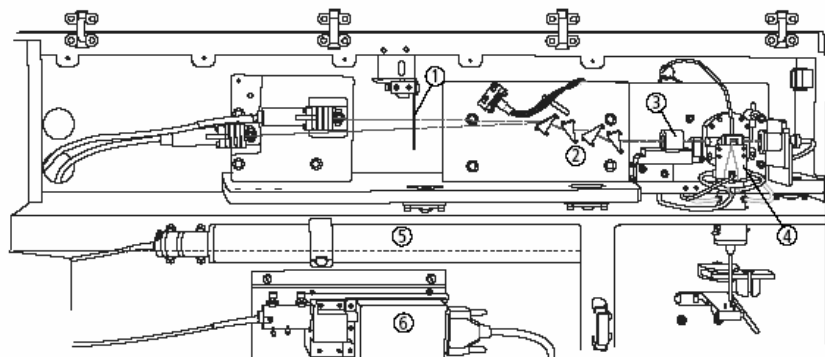
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## 光學系統

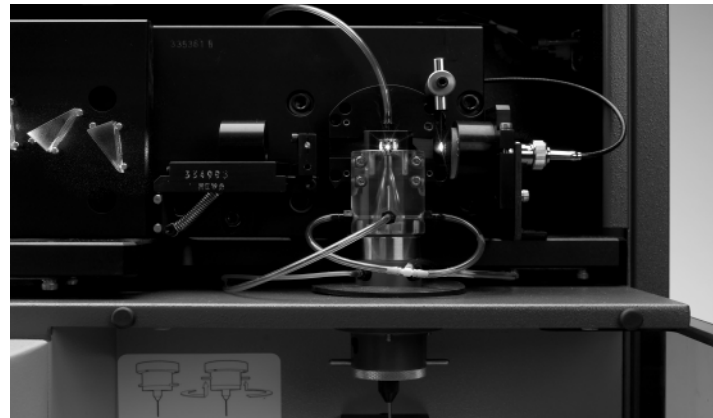
Sensitivity : 100 FITC MESF, 50 PE MESF



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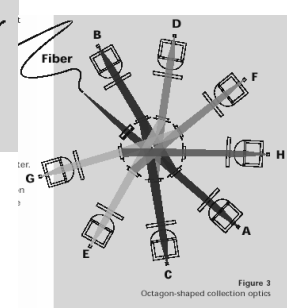
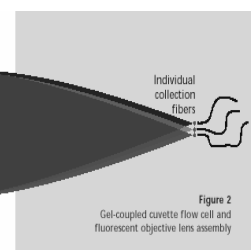


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## Cuvette Flow Cell & Fiber Optics



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## 螢光濾片

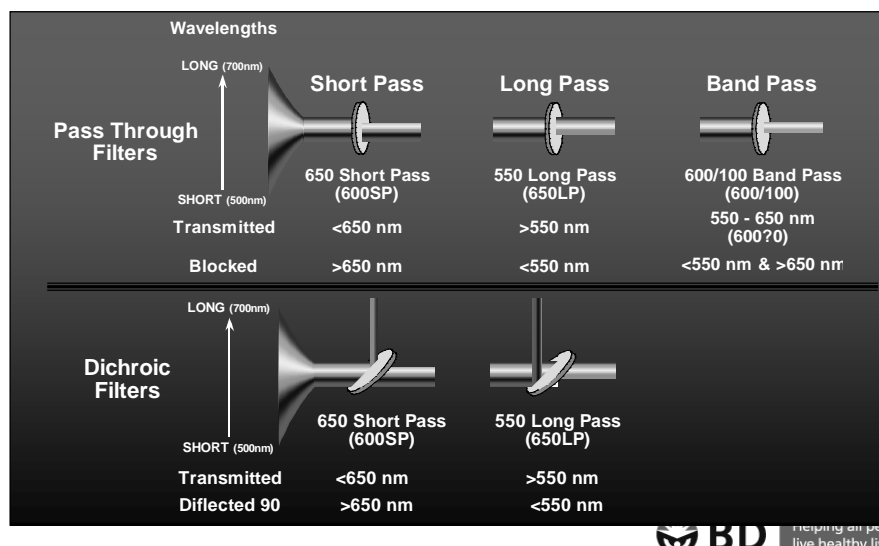
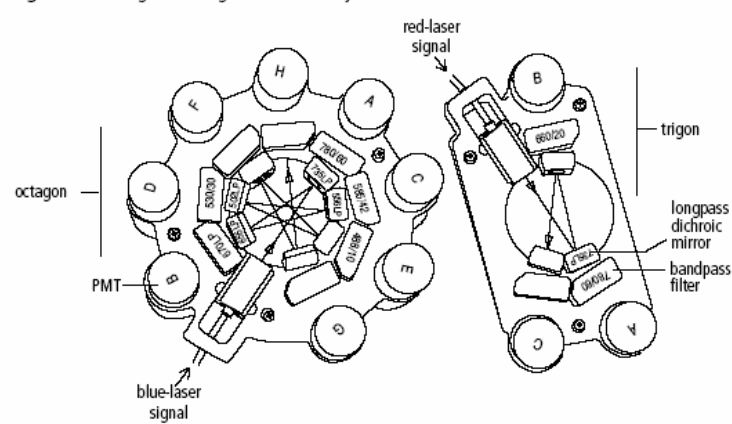


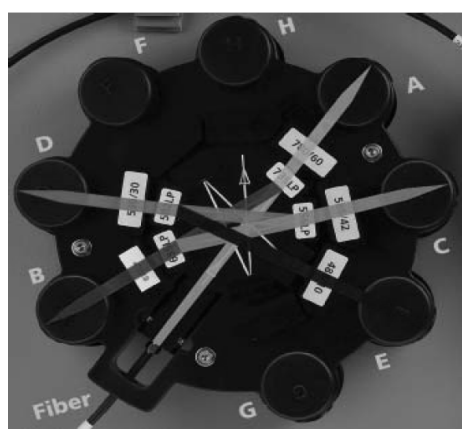
Figure 1-5 Octagon and trigon detector arrays



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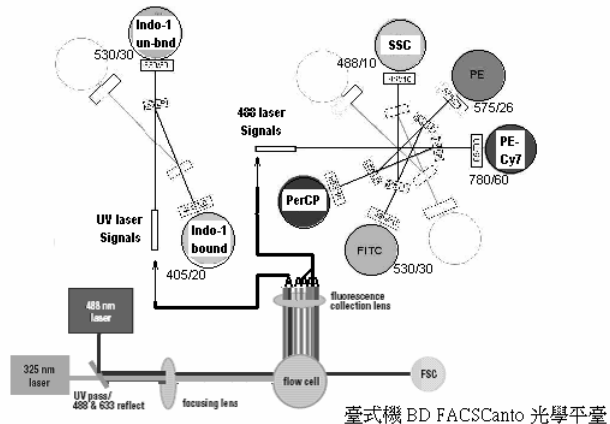


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## FACSCanto 的光學系統



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Laser	Wavelength (nm)	Min. Power <sup>a</sup> (mW)	Commonly Used Fluorochromes
Coherent® Sapphire™ Solid State	488 (blue)	20	FITC, PE <sup>b</sup> , PE-Texas Red®, PerCP, PerCP-Cy5.5, PE-Cy7, PI
JDS Uniphase™ HeNe Air Cooled	633 (red)	17	APC, APC-Cy7



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## BD FACSCanto – Default dyes

Instrument	Laser	Excitation Laser Line (nm)	Fluorescence Channel	Fluorochromes from BD Biosciences
BD FACSCanto Flow Cytometer	Blue	488	Green Yellow Red Infra Red Red Infra Red	FITC PE PerCP PE-Cy7 APC APC-Cy7
	Red	633		PerCP-Cy5.5

**Table 1**  
Validated fluorochrome combinations with the BD FACSCanto flow cytometer.



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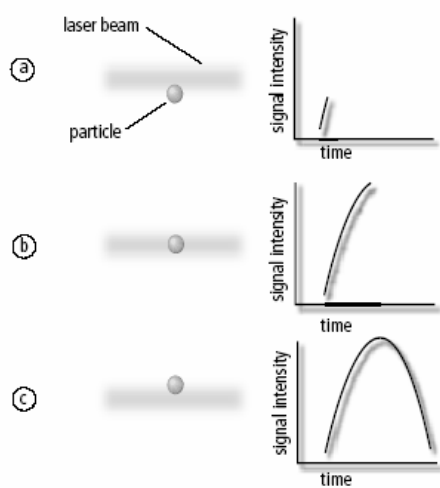
**Table 1-1** Octagon and trigon optical filters

Detector Array (Laser)	PMT Position	LP Mirror	BP Filter or LP Mirror	Intended Dye
Octagon (488-nm blue laser)	A	735	780/60	PE-Cy7
	B	655	670 LP	PerCP-Cy5.5, PerCP
	C	556	585/42	PE
	D	502	530/30	FITC
	E	blank optical holder	488/10	Side scatter (SSC)
	F	blank optical holder	blank optical holder	—
	G	blank optical holder	blank optical holder	—
	H	blank optical holder	blank optical holder	—
Trigon (633-nm red laser)	A	735	780/60	APC-Cy7
	B	blank	660/20	APC



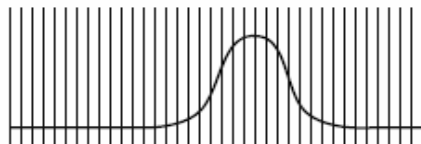
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## 電子系統

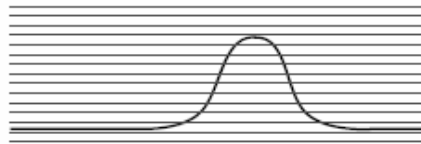


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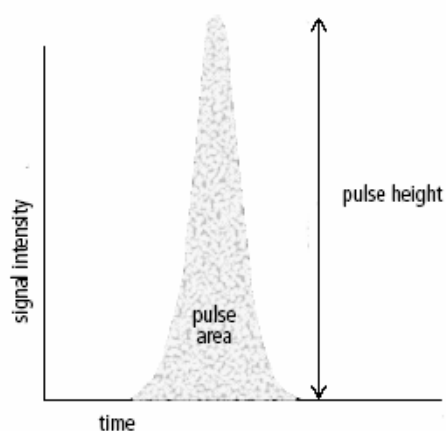
measured 10,000,000 times  
every second



digitized into 16,384 levels

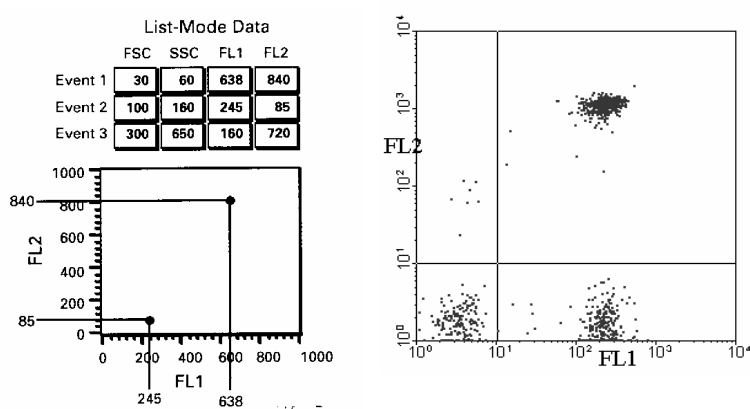


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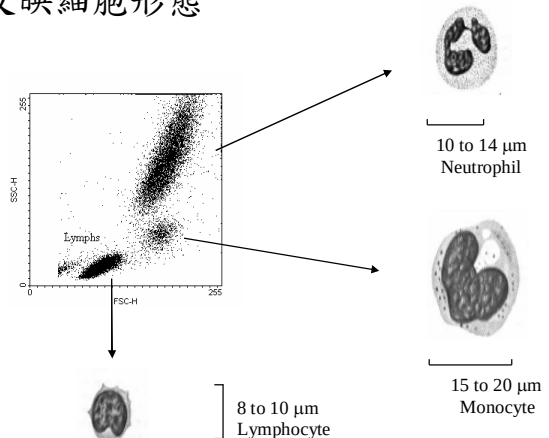
## 實驗數據的呈現



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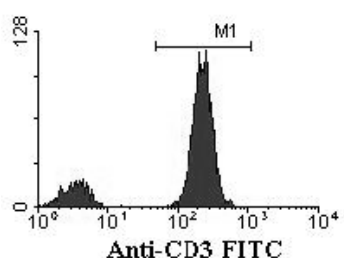
## 常見的數據呈現

散點圖反映細胞形態



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## 直方圖分析報告



File: IS20924004  
Date: 24-Sep-93  
Total Events 6000  
Gated Events 2785

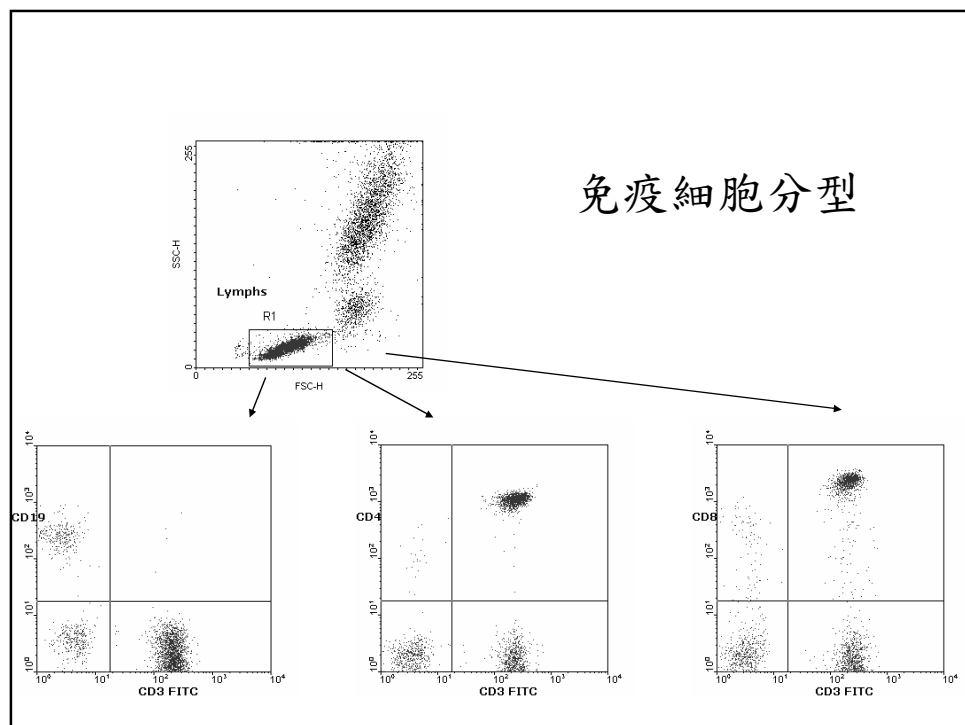
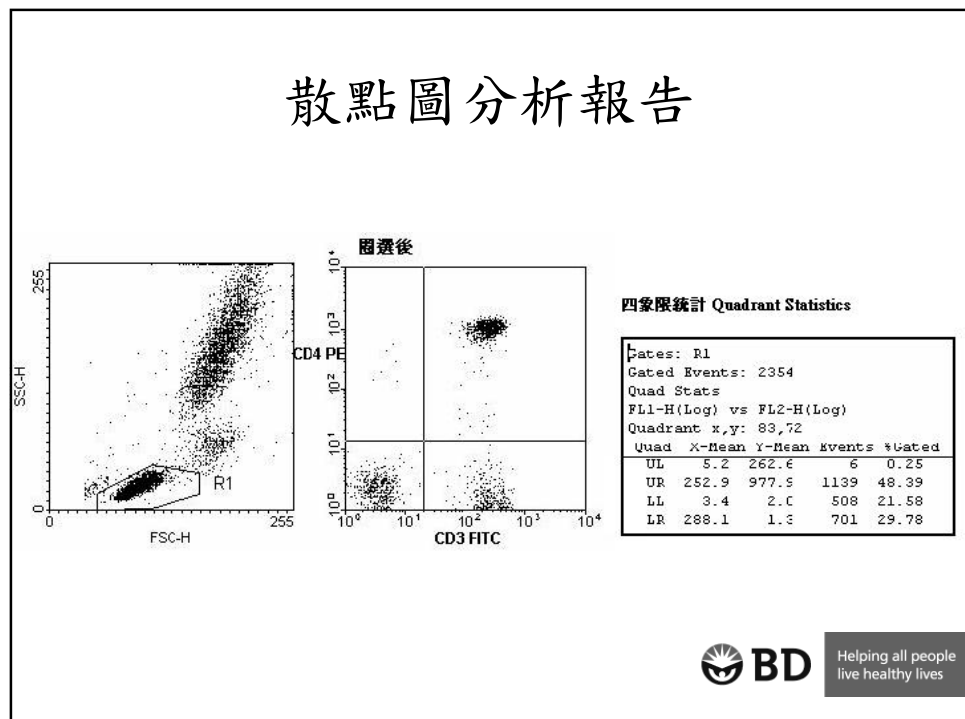
M	Low,High	Events	% Gated	GMean	CV
0	0, 255	2785	100.00	106.55	35.70
1	108, 196	2270	81.51	231.20	6.06

CV=S.D./Mean



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## 散點圖分析報告



# **Running Samples with BD FACSDiva Software**

The following topics are covered in this chapter:

- Instrument Startup on page 54
- Instrument Quality Control on page 57
- Optimization of Instrument Settings on page 68
- Data Recording and Analysis on page 79
- Daily Shutdown on page 90



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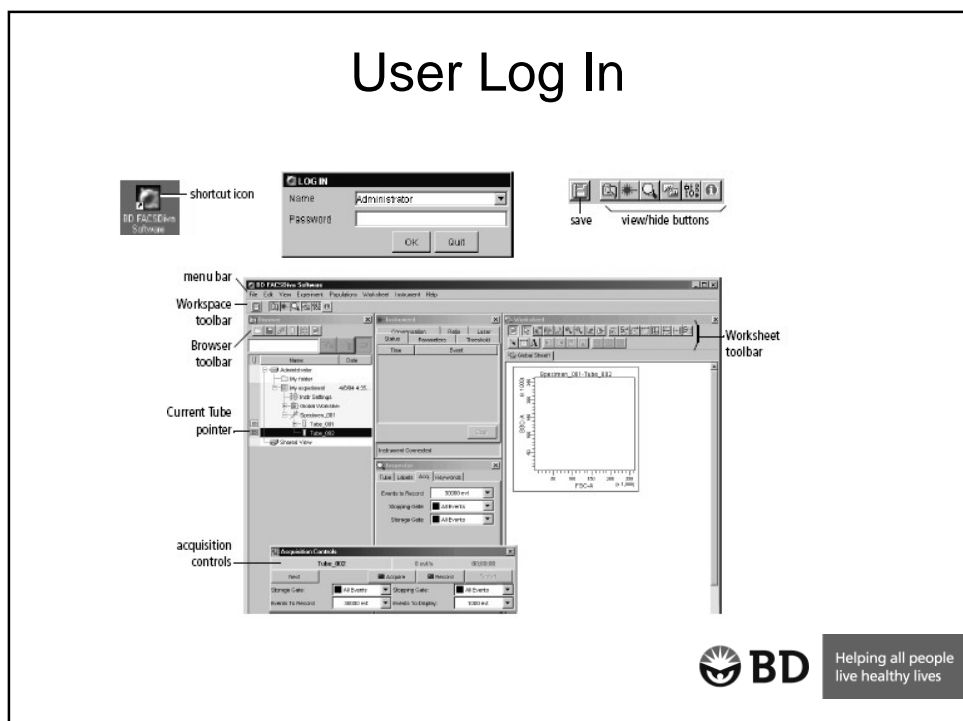
## **儀器之調試**

- 設閾參數
- 閾值
- FSC / SSC 散點圖
- 螢光信號接受器 FL 1-4
- 自動化色差補償



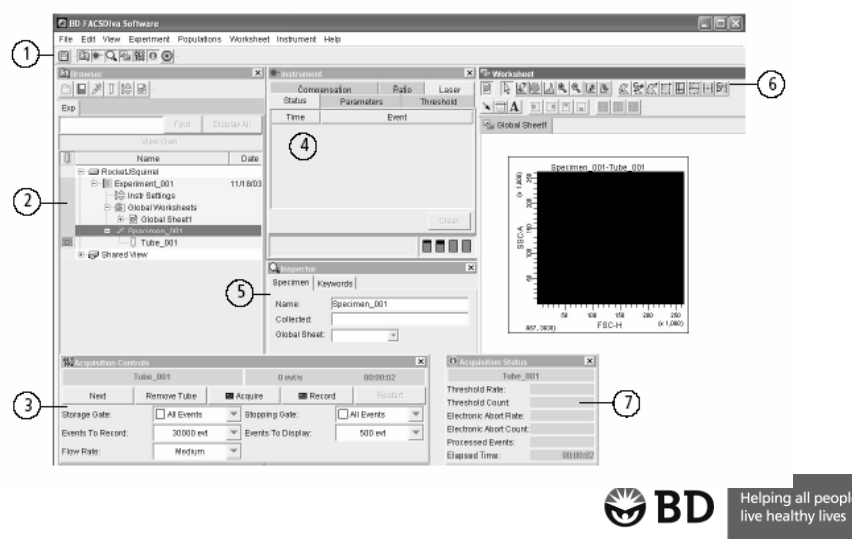
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## User Log In



## FACSDiva Workspace

Figure 2-1 BD FACSDiva workspace



## 常用螢光染劑

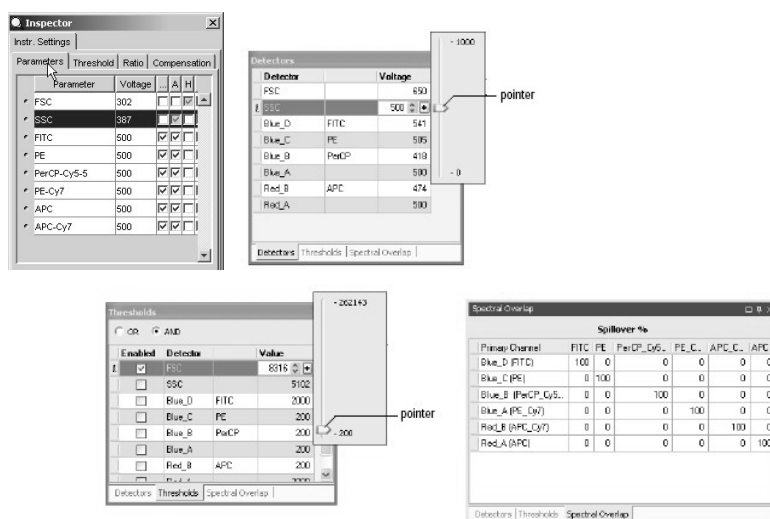
Instrument	Laser	Excitation Laser Line (nm)	Fluorescence Channel	Fluorochromes from BD Biosciences
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	Red	633		PerCP-Cy5.5

Table 1  
Validated fluorochrome combinations with the BD FACSCanto flow cytometer.



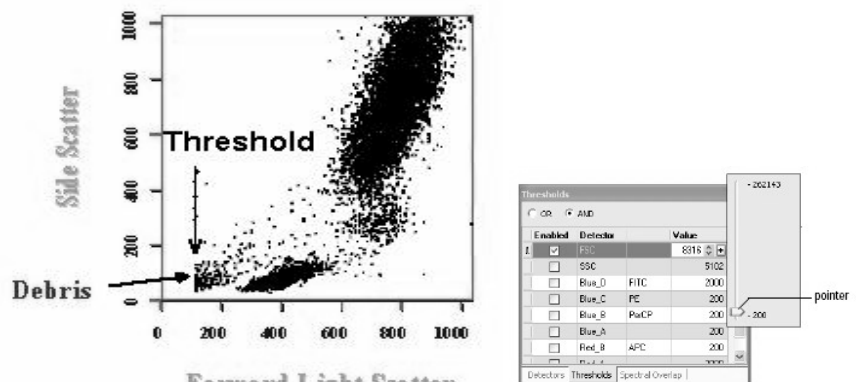
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## Instrument Set Up

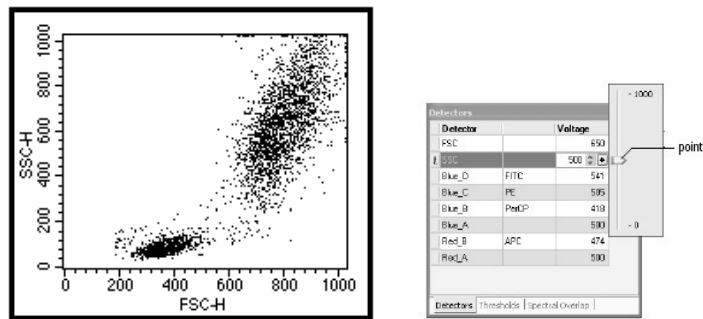


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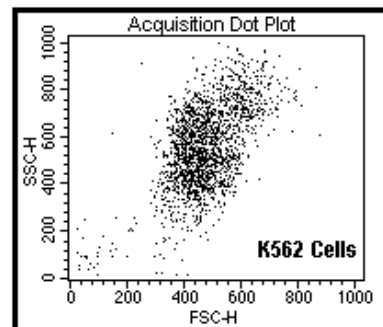
## 閥值的調整



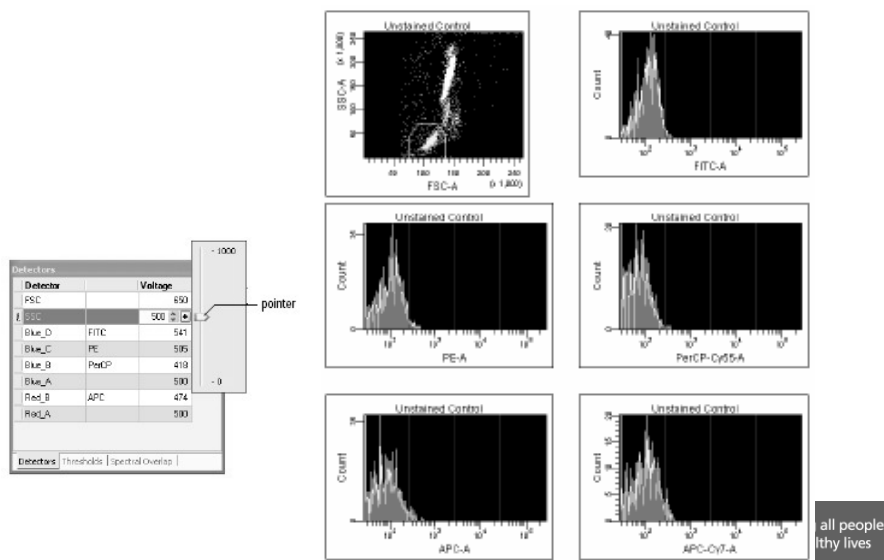
## 散射光信號的調整



## 散射光信號的調整 Cell Line

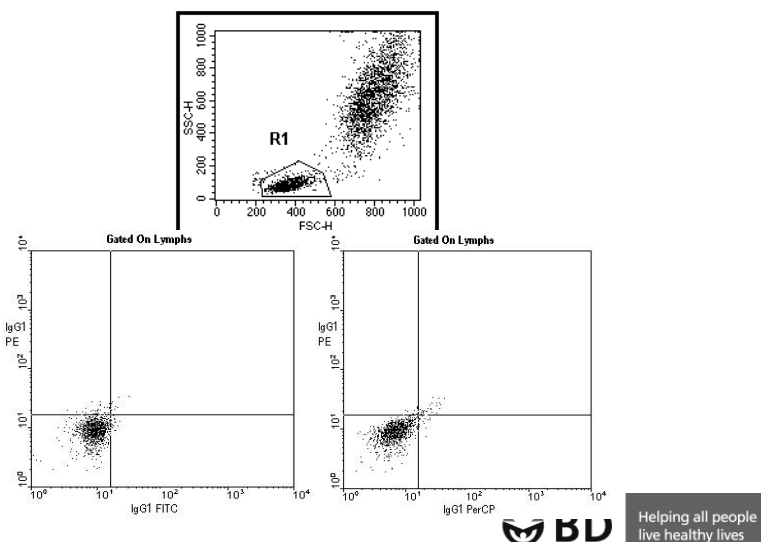


## 自體螢光信號的調整

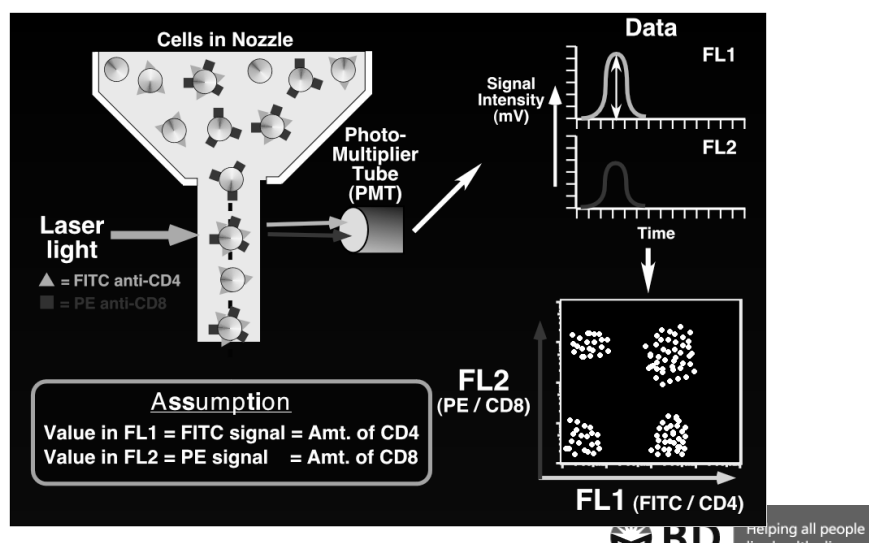


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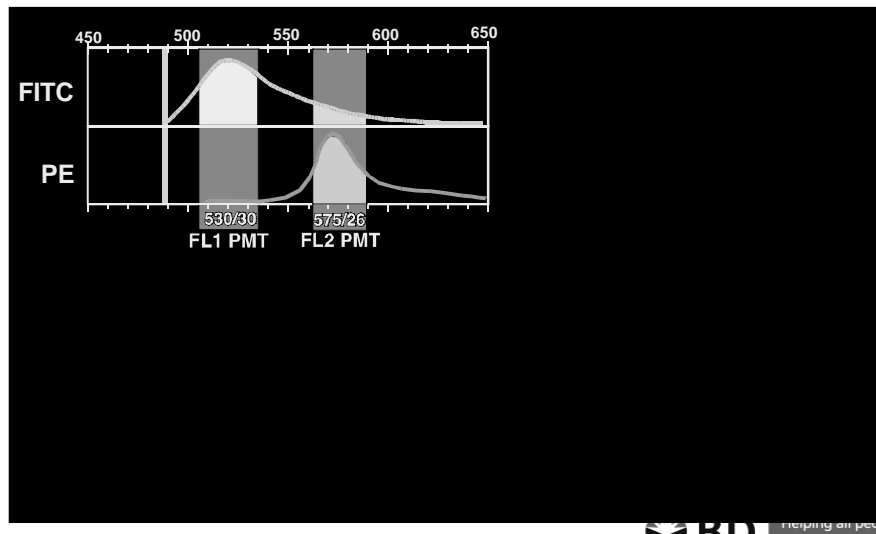
## 自體螢光信號的調整 2-3C



## 螢光補償調整 (The Problem)



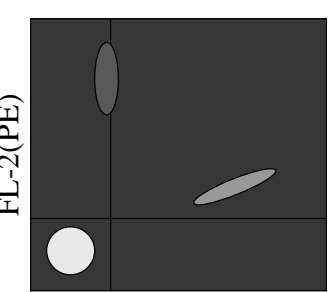
## 螢光補償調整 (The Problem)



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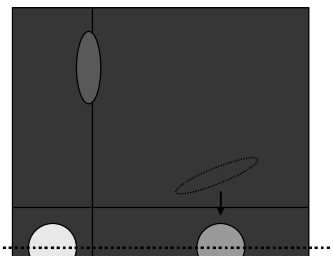
## 螢光補償調節

FL-2(PE)



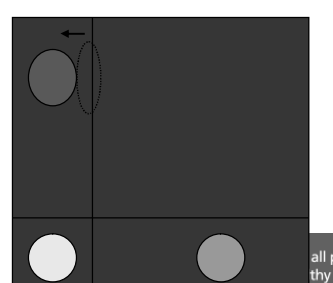
$$\frac{\text{FL2} - \% \text{FL1}}{(18 - 30\%)}$$

$$\xrightarrow{\hspace{1cm}}$$
  
$$\text{FL1} - \% \text{FL2} \\ (0 - 1\%)$$

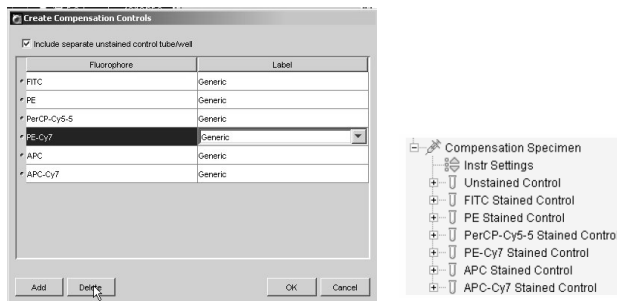


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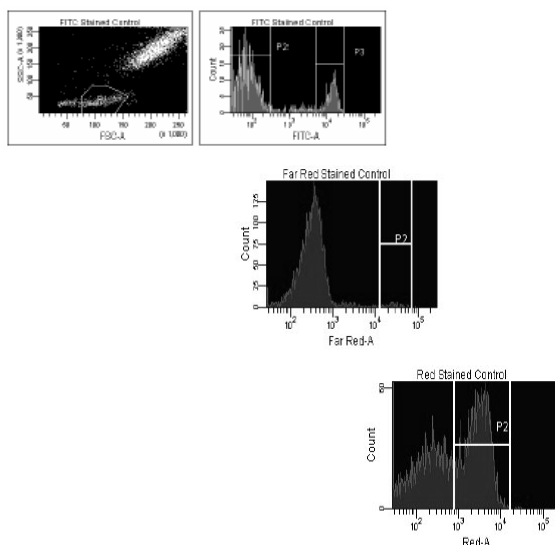
FL-1(FITC)



## Create Compensation Controls

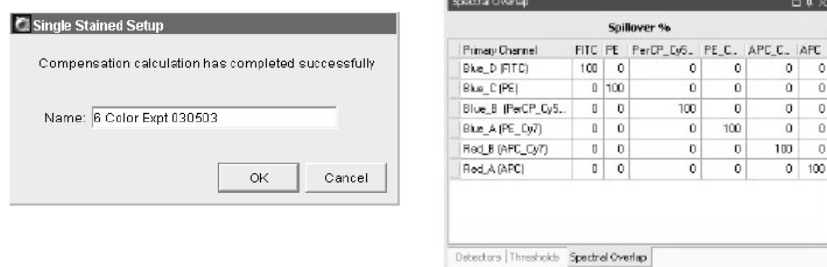


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選擇“Instrument>Instrument Setup>Calculate Compensation”命令。



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## FACSCanto 各部構造



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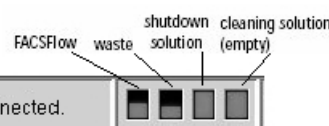
## 正確開機程序

1. 開啟主電源。
2. 開啟電腦。在密碼登錄框中，輸入BDIS，按Enter。
3. 開啟 BD FACSDiva 軟體。
4. 確定軟體與細胞儀完成連線。
5. 完成連線後，檢視各項試劑液面是否正常。
6. 等待細胞儀自動完成液流啟動程式。
7. 當液流啟動程式完成後，請點選OK。
8. 檢視雷射是否暖機完成。

整個程序約需5~10分鐘。



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## 上機檢查程序

1. 檢品濃度調至 $1\times 10^6$  cells/ml ? 一般只需0.5 ml。
2. 是否已小心地去除檢品中之細胞團塊。
3. 是否已將檢品放至FALCON 2052 試管中？試管是否有裂痕？
4. 是否有足量專用鞘液？是否已將廢液倒掉？
5. 是否已執行Auto Clean ?
6. 系統氣壓讀數55~65 Psi ?
7. 是否已將液體過濾器中之氣泡排空？
8. 請填寫使用登記表。



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## 建議使用之鞘液液體

1. FACSFlow (BDIS)
2. 經0.22um過濾之自備含0.05% NaN<sub>3</sub>之 PBS

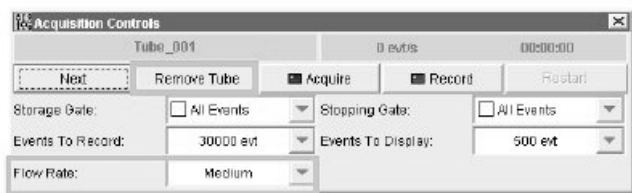
不鼓勵使用下列溶液作為鞘液液體之用：

- Fisher Hematology Diluent
- Isoton III
- Isolac D
- DI water



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Figure 2-7 Acquisition controls unique to the BD FACSCanto



Low: 樣品流速=  $10 \mu\text{L}/\text{min}$

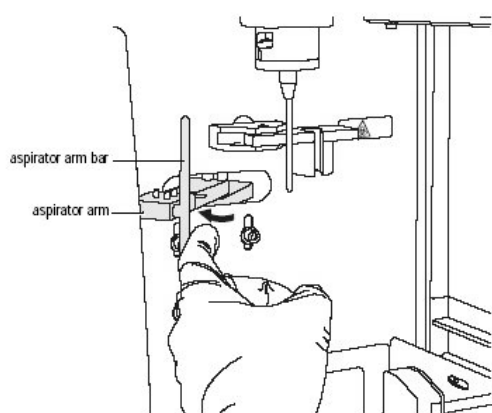
Medium: 樣品流速=  $60 \mu\text{L}/\text{min}$

High: 樣品流速=  $120 \mu\text{L}/\text{min}$



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## 檢體吸取區(SIT)



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## Remove Tube正確步驟

步驟要確實：〔to minimize carry over between samples〕

1. 在Acquisition Controls視窗中，點Remove tube；此時，會出現一個進程顯示對話框。
2. 右手握著小管，左手將Aspirator arm向左搬到底。
3. 取下樣品管。
4. 放開左手，使Aspirator arm回到中央；此時，SIT會自動清洗；清洗完成後，進程顯示對話框會自動消失，在清洗動作完成前，請勿放下一管。



## 儀器清洗與關機正確步驟

清洗步驟要確實：〔尤其是使用PI之後〕

1. 取3 ml FACSClean (10% Bleach) 上樣品。
2. 在Acquisition control上，按Acquire。
3. 讓儀器在Medium Flow Rate模式下，Acquire 5分鐘，再按Acquire以中止收取。取下樣品管。
4. 換上3 ml dH<sub>2</sub>O上樣品。
5. 在Acquisition control上，按Acquire。
6. 讓儀器在Medium Flow Rate模式下，Acquire 5分鐘，再按Acquire以中止收取。取下樣品管。
7. 點選Instrument > Fluidics Shutdown。液流系統關閉中，會出現下方進度報告。
8. 在允許關閉系統的對話框中，點選OK。
9. 關閉電腦，關閉細胞儀。



## 公共儀器的維護



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