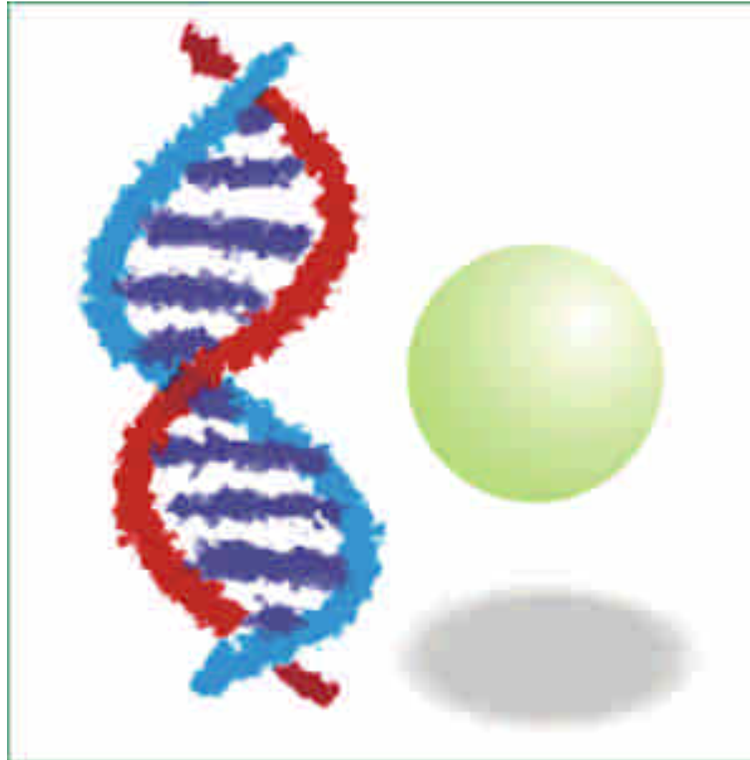


BST
生化科技系

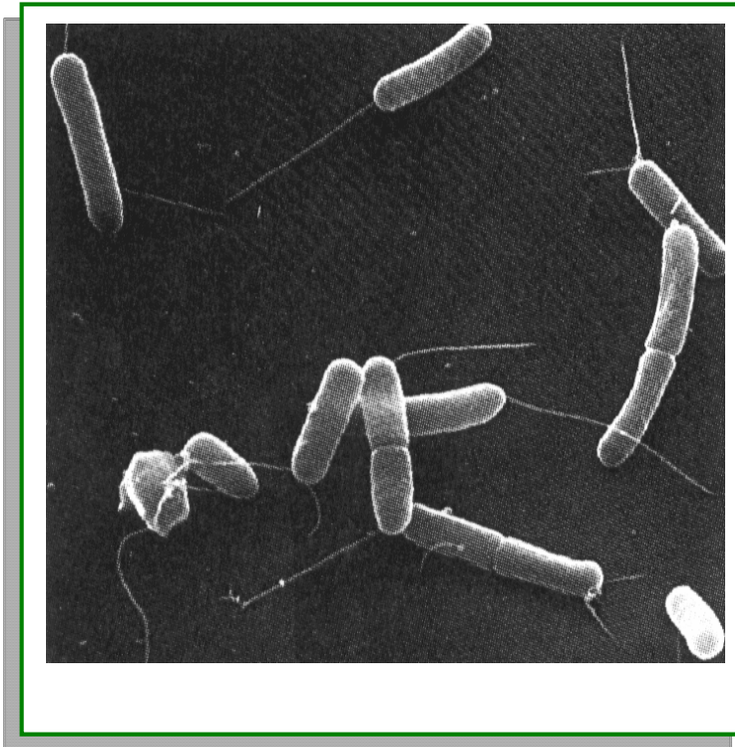
BCX



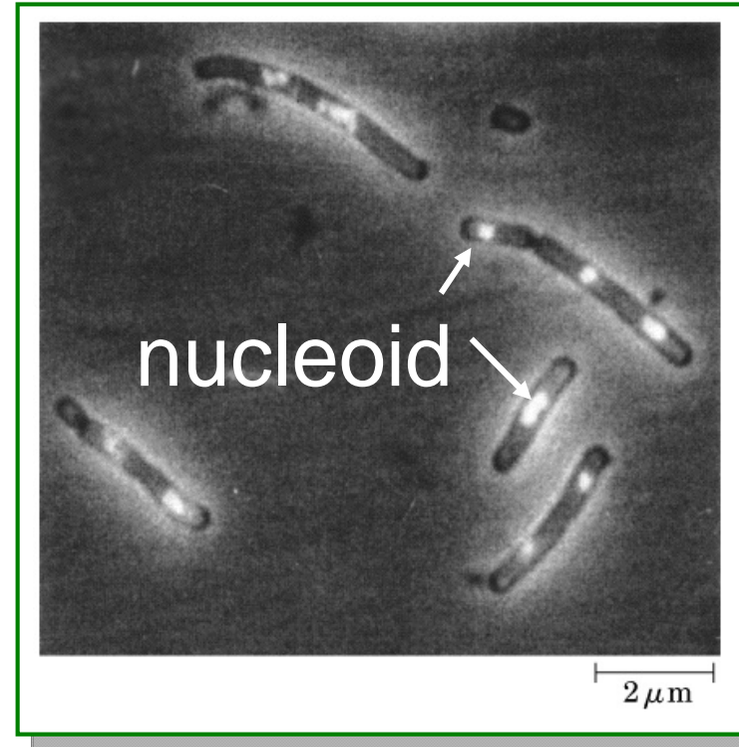
N1

生物化學實驗
染色體 DNA 分離與檢定

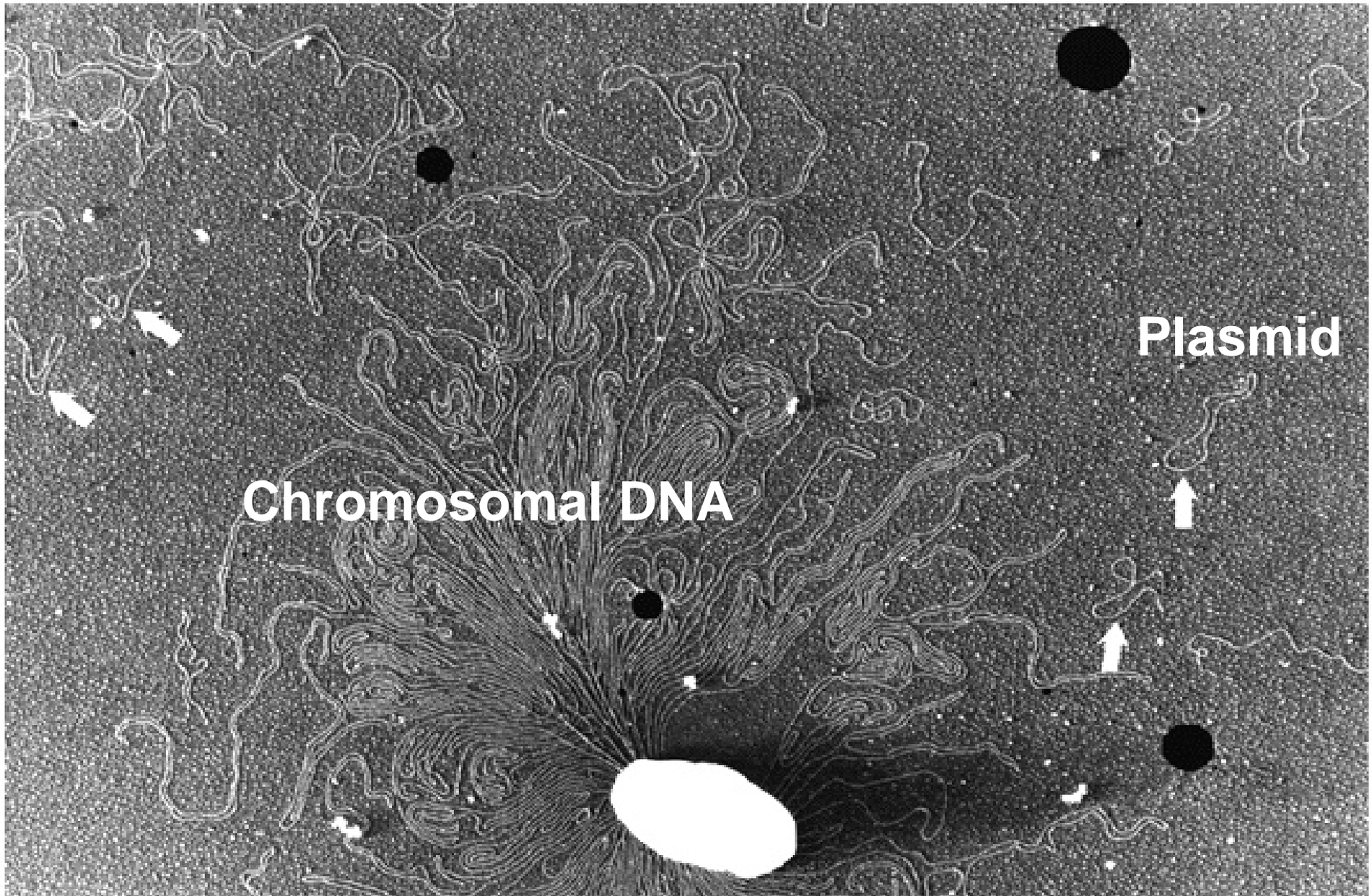
■ 實驗材料：*E. coli*



Alcamo, I. E. (2000) DNA Technology: The Awesome Skill. 2nd Ed. Academic Press. Fig. 5.9

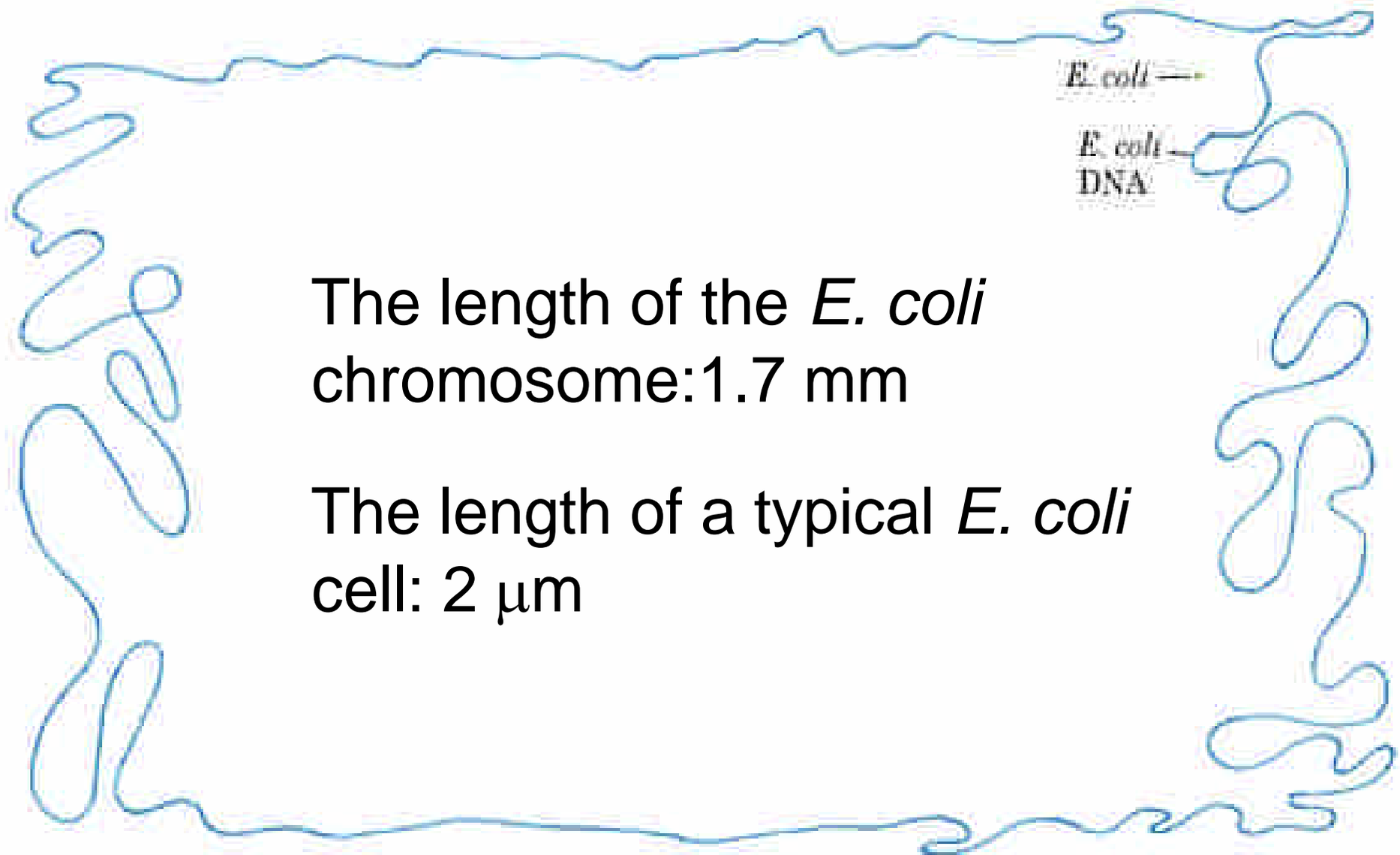


Nelson, D. L. and Cox, M. M. (2000) Lehninger Principles of Biochemistry. 3rd ed., Worth Publishers. Fig. 24-31



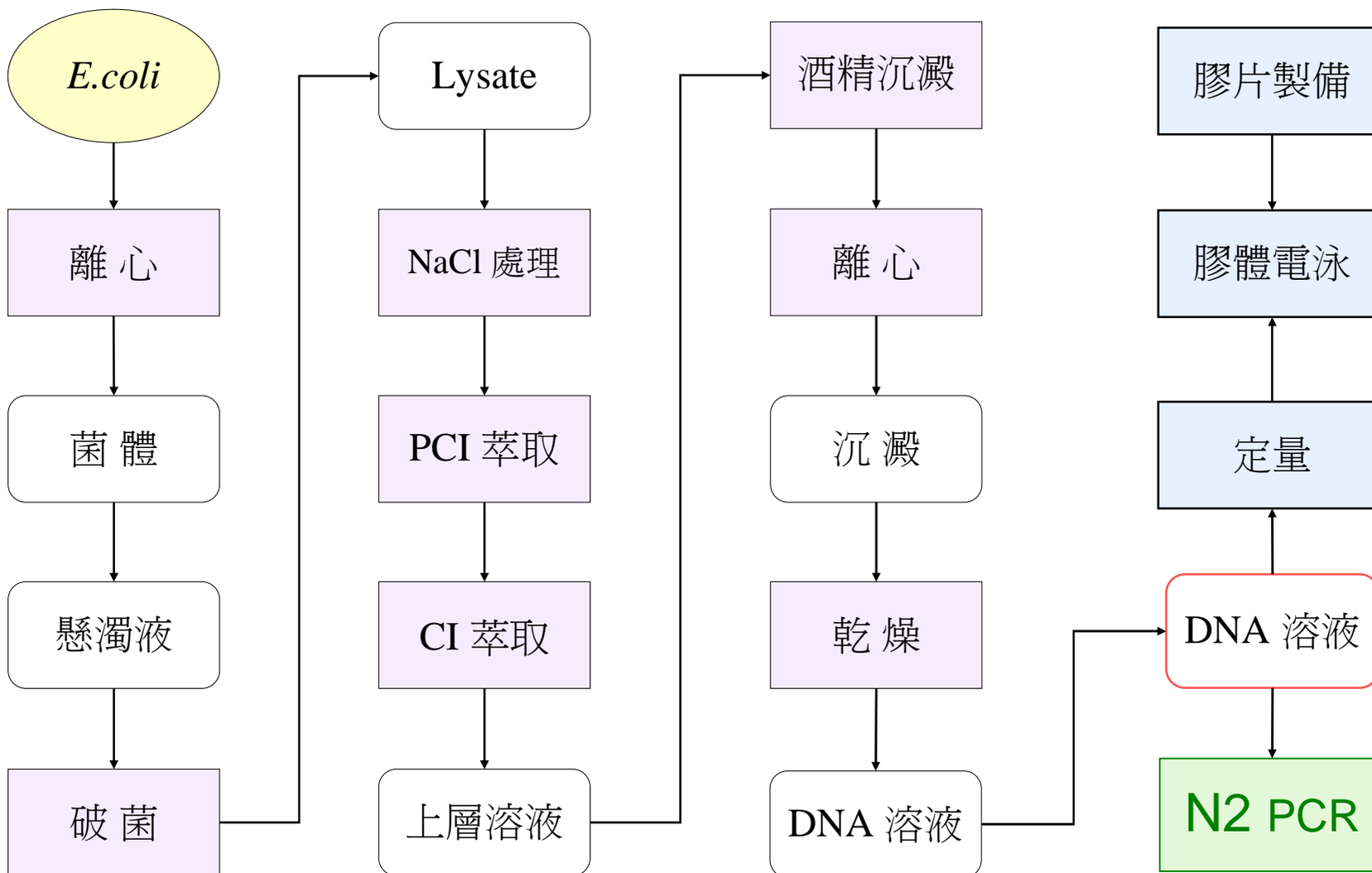
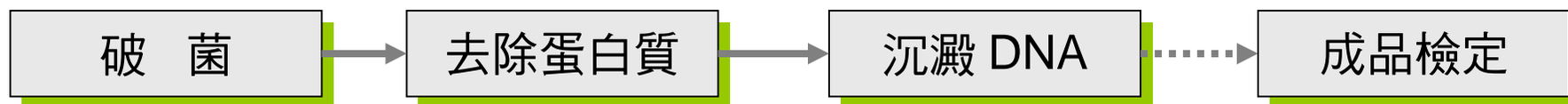
DNA from a lysed *E. coli* cell

圖取材自：Nelson, D. L. and Cox, M. M. (2000) *Lehninger Principles of Biochemistry*. 3rd ed., Worth Publishers. Fig. 24-6



The length of the *E. coli* chromosome: 1.7 mm

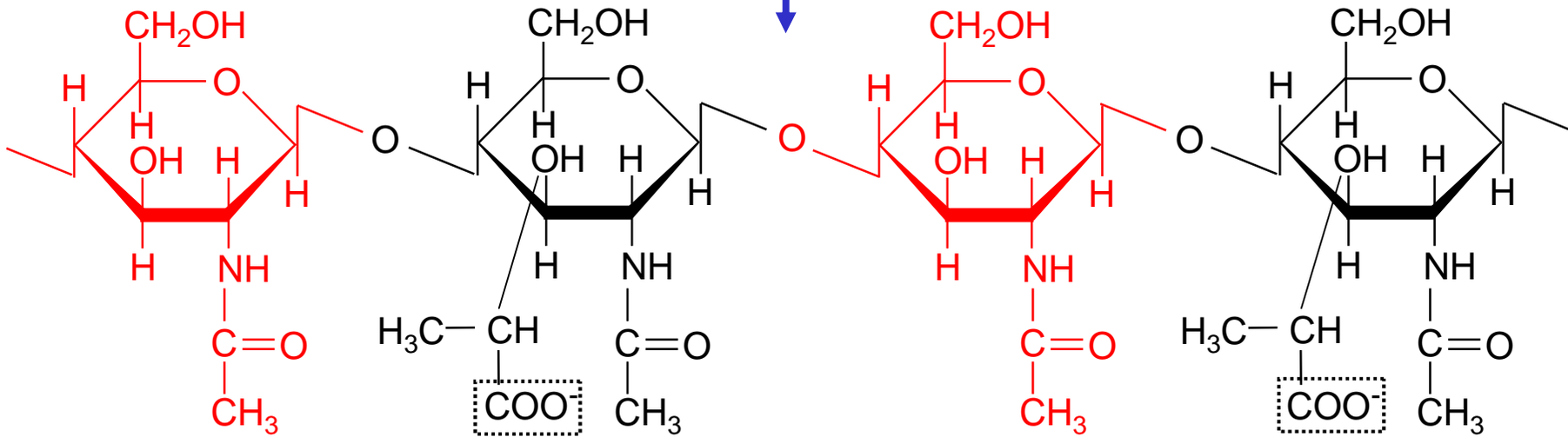
The length of a typical *E. coli* cell: 2 μm



Lysozyme : 作用於 peptidoglycan 中的糖苷鍵



Lysozyme



NAG

NAM

NAG

NAM

N-acetylmuramic acid

N-acetylglucosamine

■ 去除核酸中的蛋白質：



DNA or RNA solution



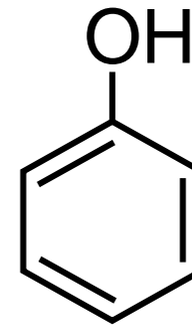
Aqueous phase



Aqueous phase



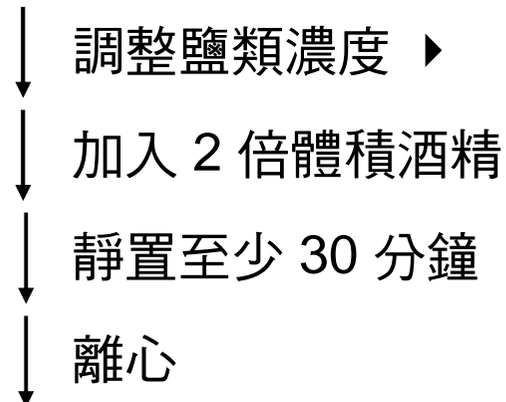
Deproteinized DNA or RNA



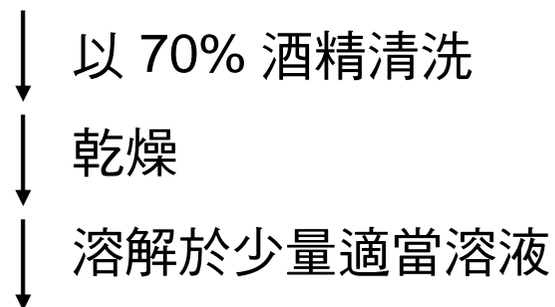
Phenol

酒精沈澱：濃縮 DNA 的方法

DNA 溶液



DNA 沈澱



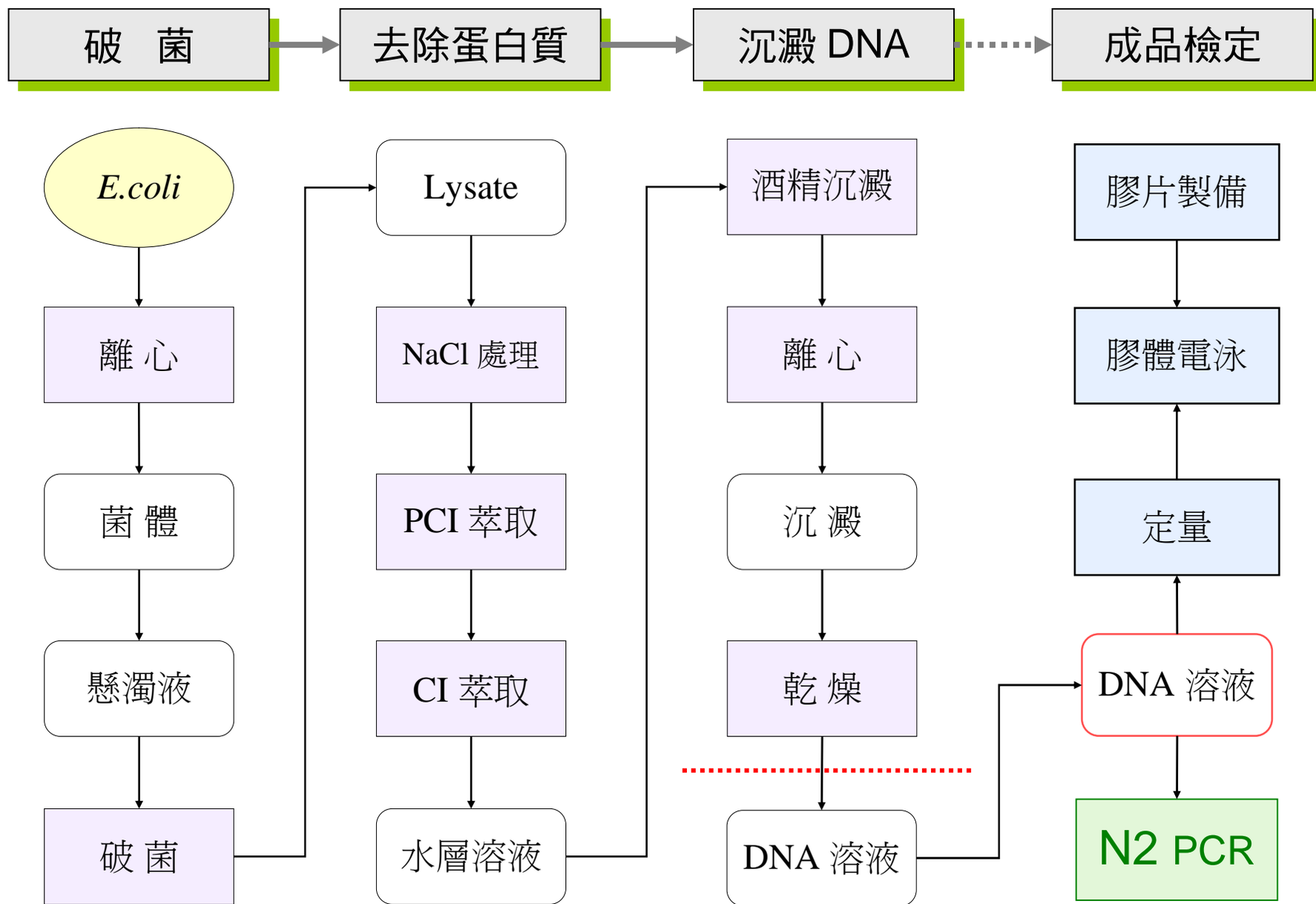
濃縮 DNA 溶液

▶ 鹽類及最終濃度：

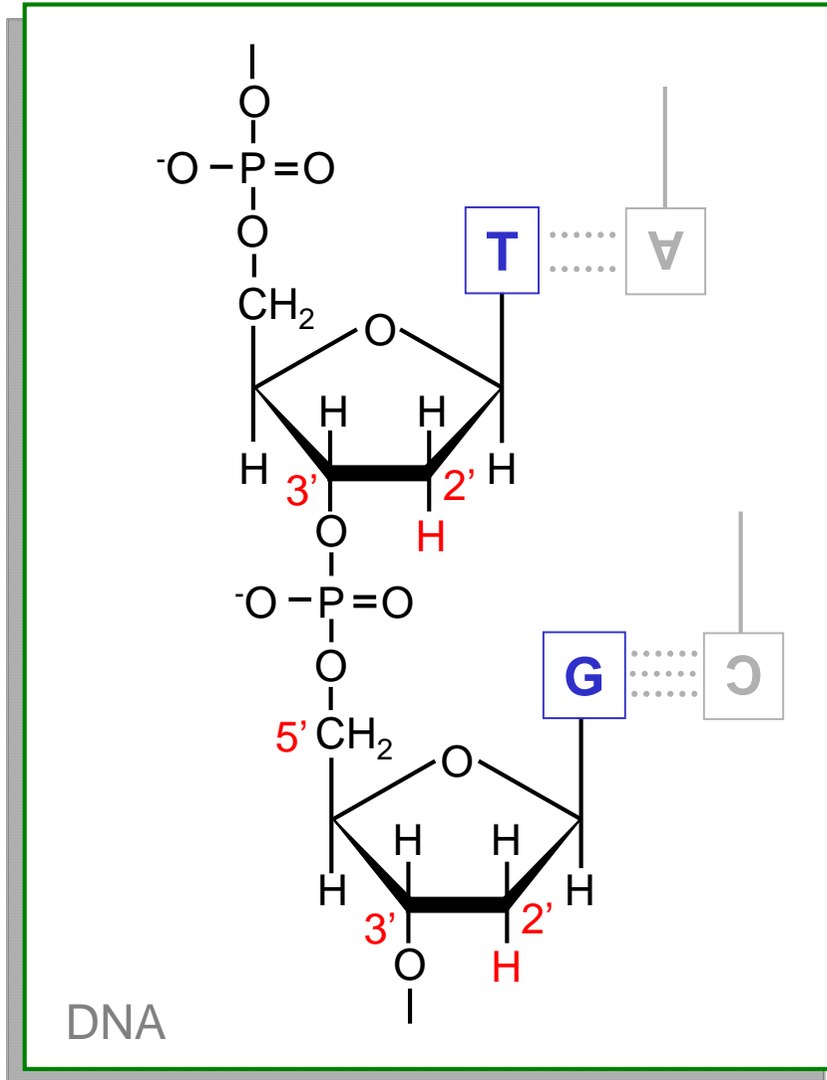
NaOAc (pH 5.2), 0.3 M

NH₄OAc, 2.0-2.5 M

NaCl, 0.2 M



核酸抽取要點：



(1) 避免分解

(2) 避免斷裂

(3) 避免污染

(4) 保持清潔

■ 常用核酸電泳系統：



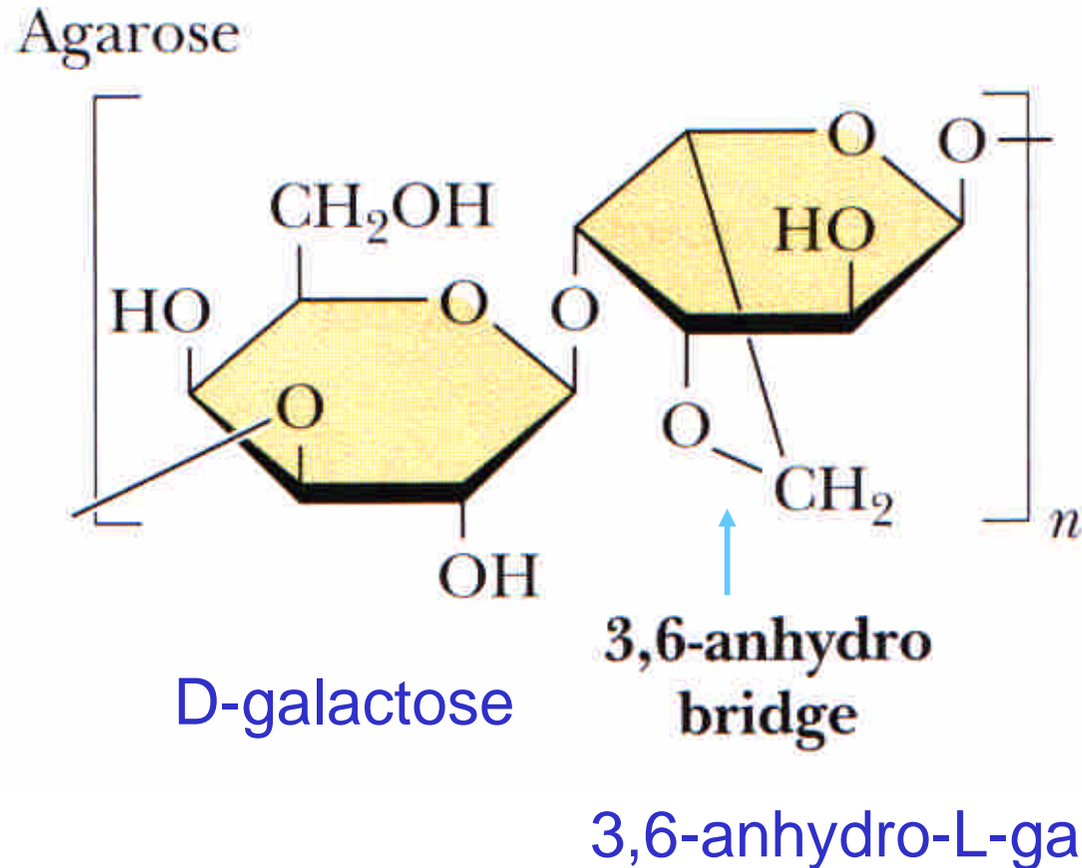
□ Agarose gel electrophoresis

- ▶ Non-denaturing gel
- ▶ Denaturing gel

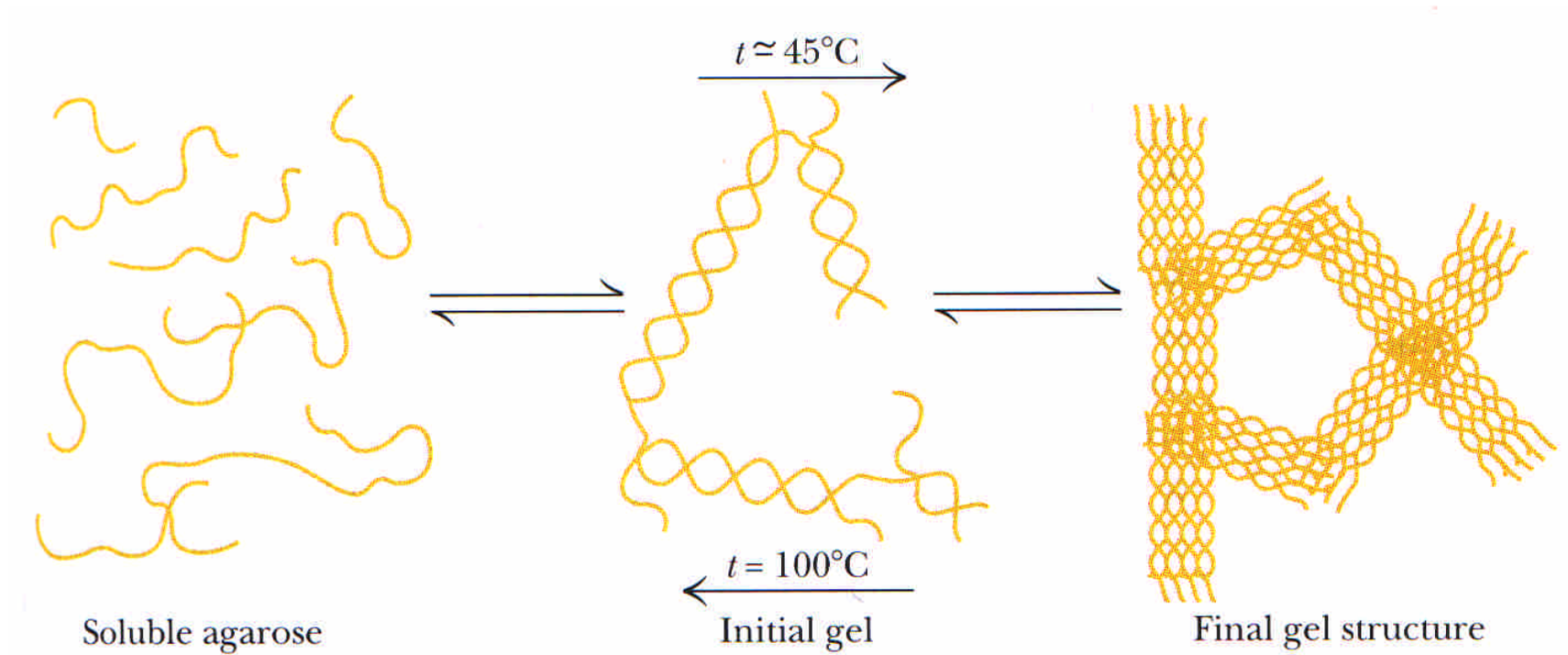
□ Polyacrylamide gel electrophoresis

- ▶ Non-denaturing gel
- ▶ Denaturing gel

■ Agarose 基本組成：

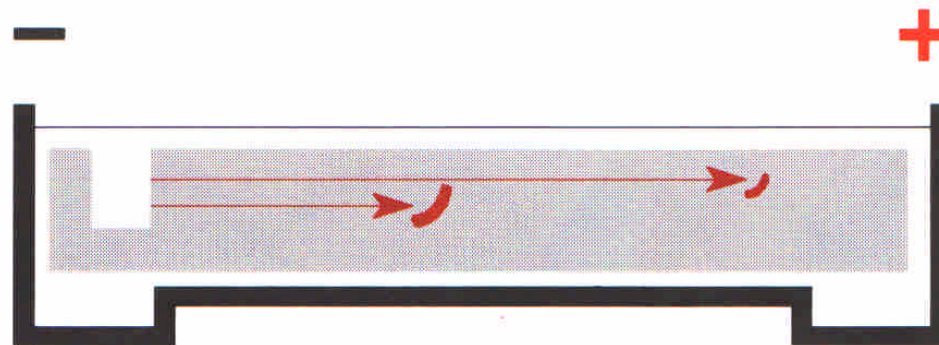
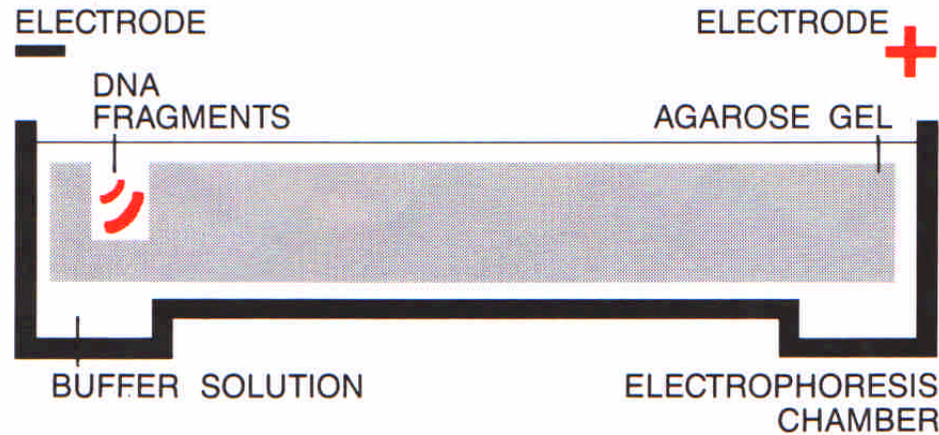


■ Agarose 溶解與凝膠的結構：



Garrett, R. H. and Grisham, C. M. (1999) Biochemistry. 2nd Ed. Saunders College Publishing. Fig. 7.32

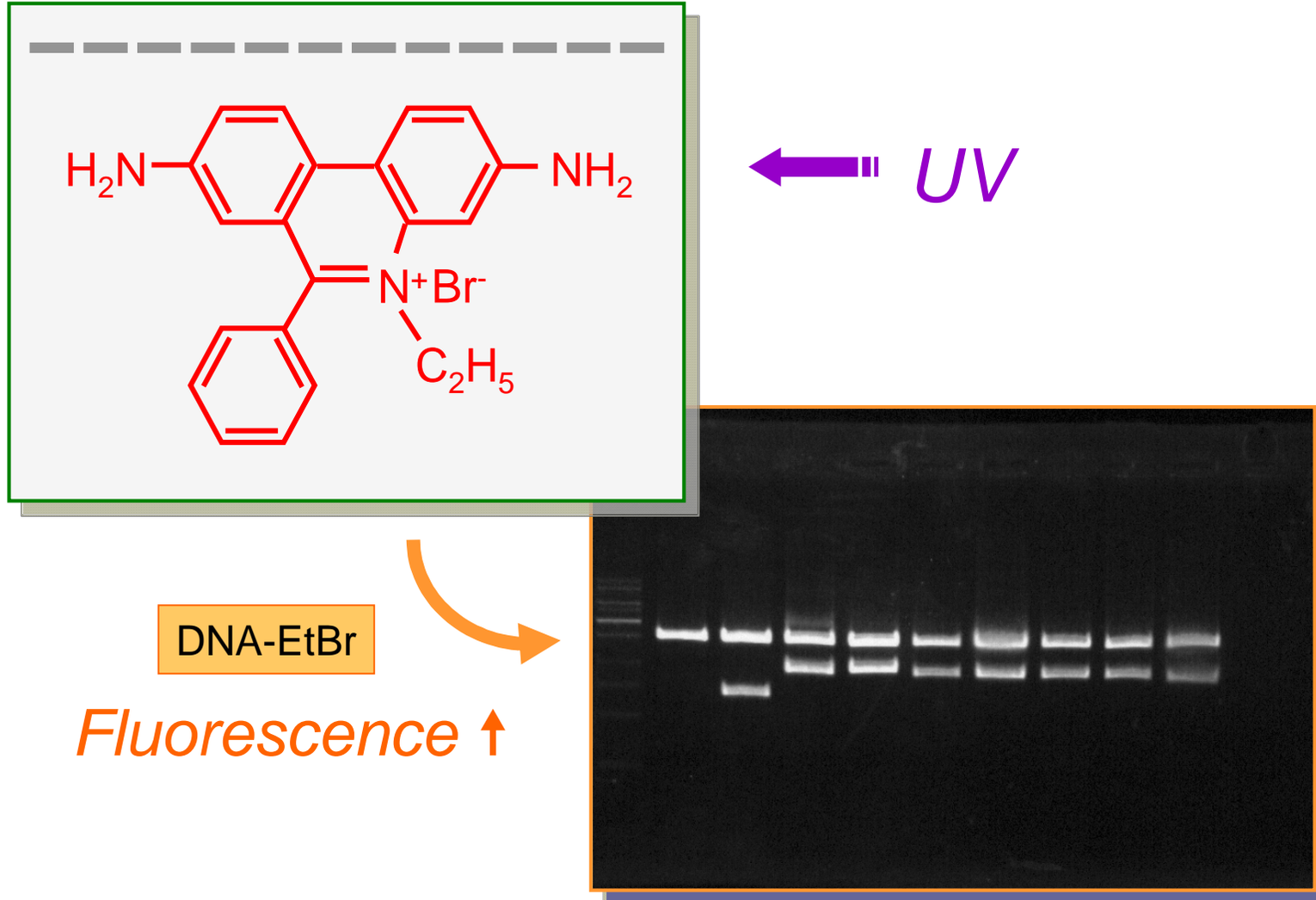
以水平電泳槽進行 agarose 膠體電泳：



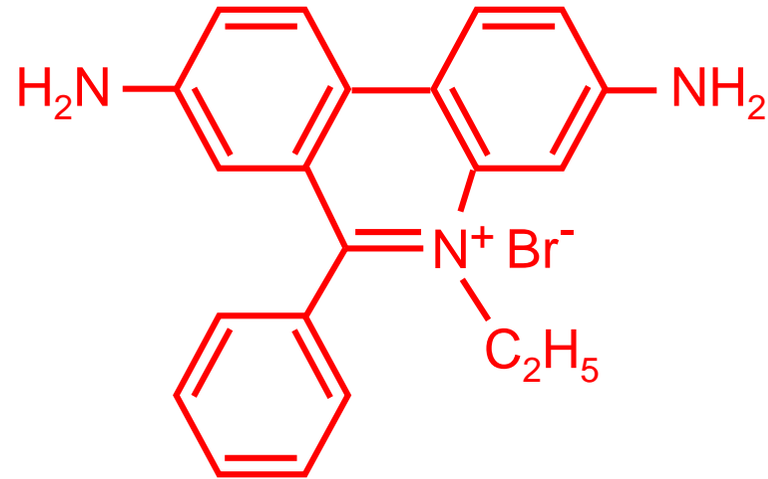
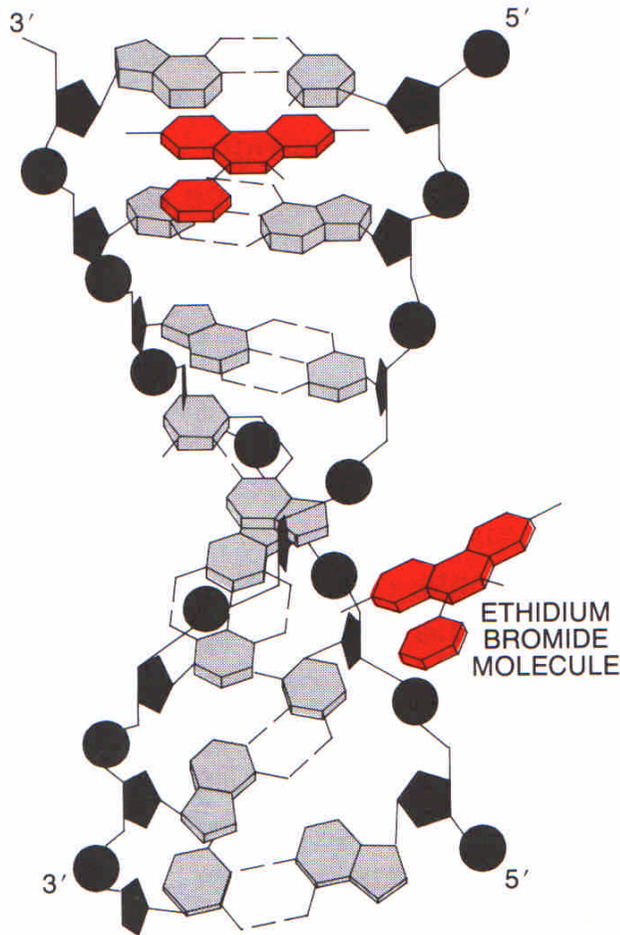
Small DNA fragment moves further through gel than large fragment

取材自 Micklos, D. A. and Freyer, G. A. (1990) DNA Science: A First Course in Recombinant DNA Technology. Cold Spring Harbor Laboratory Press. (p.48)

■ Ethidium bromide 可染出 DNA 色帶：



Ethidium bromide 可嵌入雙股 DNA



Ethidium bromide

突變劑！請小心使用！

左圖取材自 Micklos, D. A. and Freyer, G. A. (1990) DNA Science: A First Course in Recombinant DNA Technology. Cold Spring Harbor Laboratory Press. (p.49)

■ 影響 DNA 泳動的主要因素：

- ▶ Agarose 濃度
- ▶ DNA 構形與大小
- ▶ 電壓
- ▶ 電泳緩衝液
- ▶ Intercalating dye 存在與否

選擇適當膠體濃度



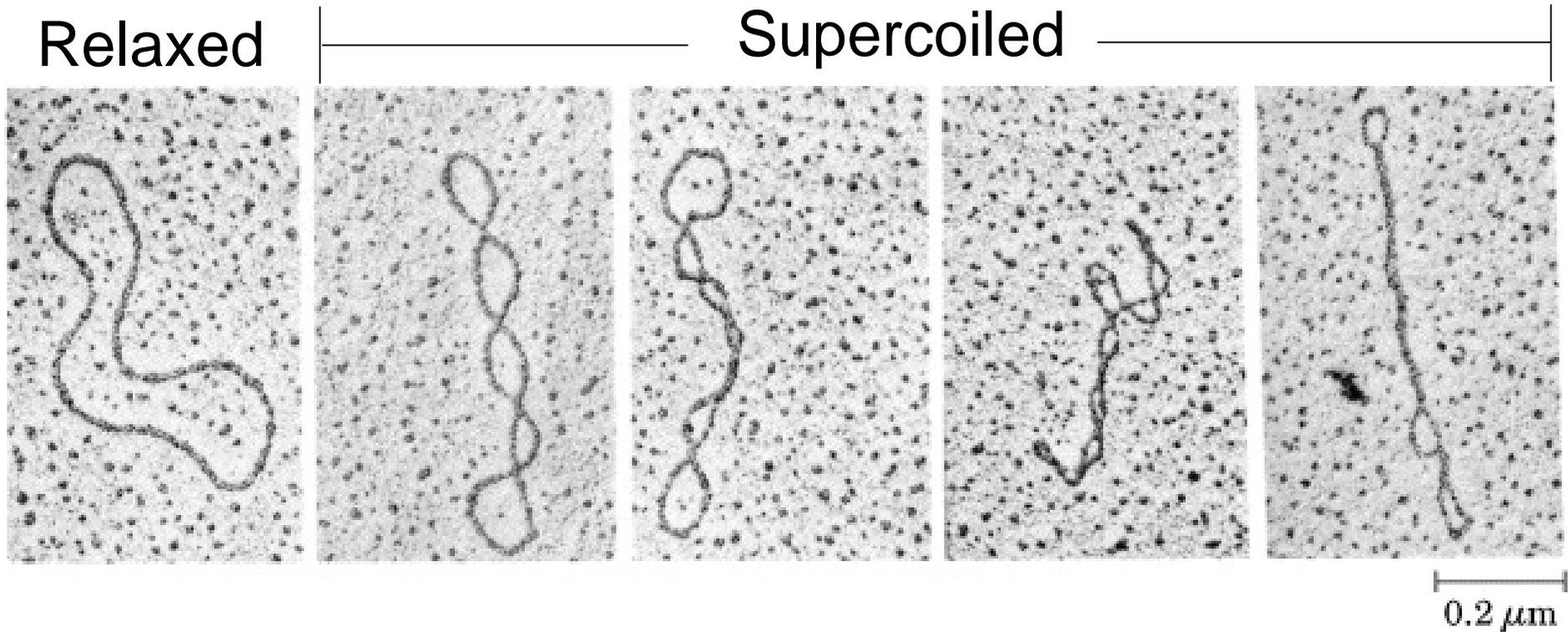
Agarose concentration in gel (% [W/V])	Range of separation of linear DNA molecules (kb)
0.3	5-60
0.6	1-20
0.7	0.8-10
0.9	0.5-7
1.2	0.4-6
1.5	0.2-3
2.0	0.1-2

資料取材自 Sambrook, J., Russell, D. W. (2001) Molecular Cloning: A Laboratory Manual. 3rd ed., Cold Spring Harbor Laboratory Press. Table 5.5

■ 影響 DNA 泳動的主要因素：

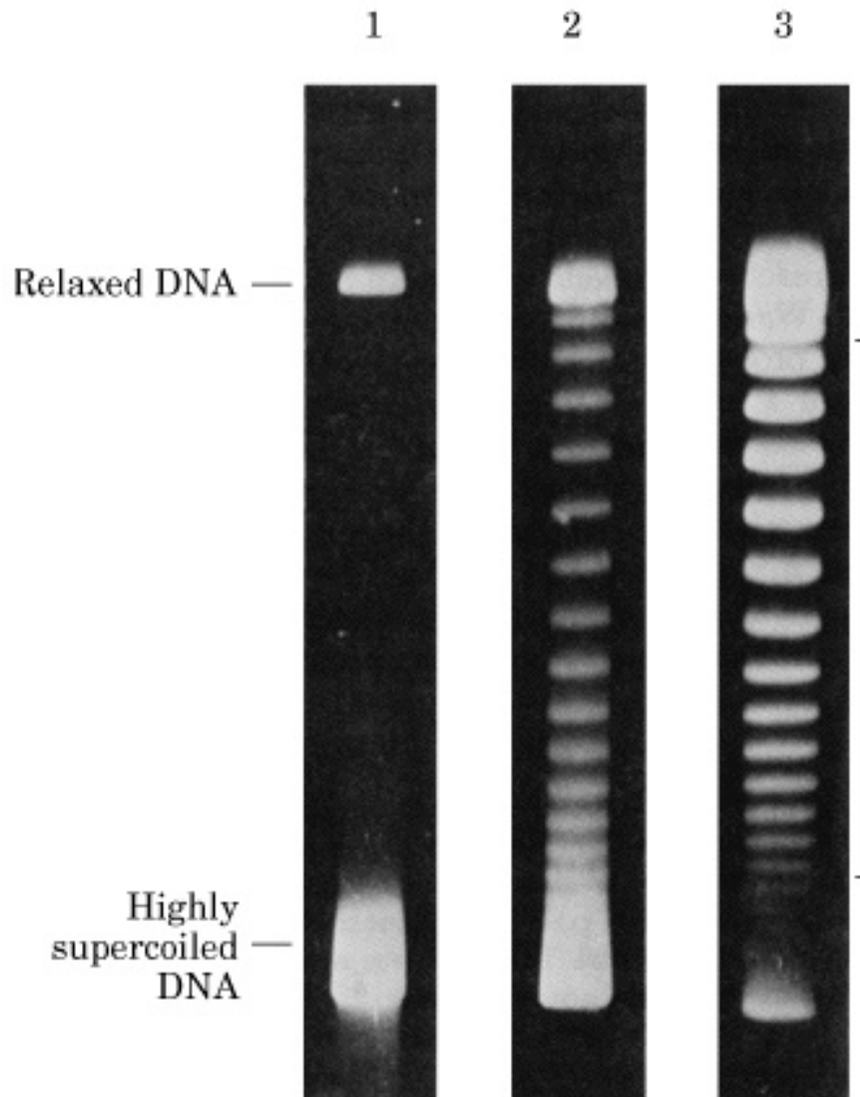
- ▶ Agarose 濃度
- ▶ DNA 構形與大小
- ▶ 電壓
- ▶ 電泳緩衝液
- ▶ Intercalating dye 存在與否

環狀質體的不同構形



Closed circular plasmid DNA

不同構形具不同泳動率



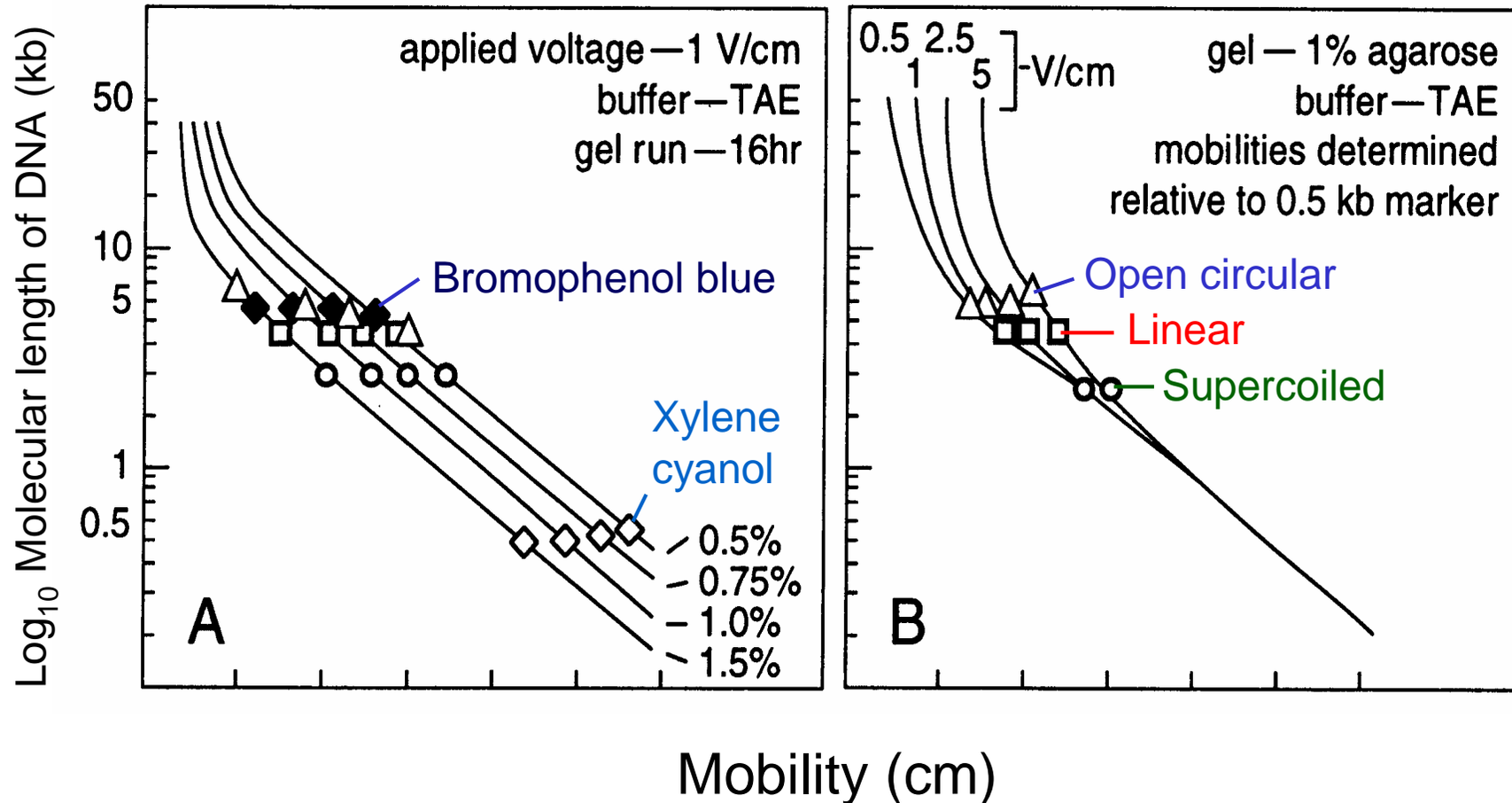
* Lane 2 & 3:
DNA treated with a
type I topoisomerase

圖取材自：Nelson, D. L. and Cox, M. M. (2000)
Lehninger Principles of Biochemistry. 3rd ed.,
Worth Publishers. Fig. 24-19

■ 影響 DNA 泳動的主要因素：

- ▶ Agarose 濃度
- ▶ DNA 構形與大小
- ▶ 電壓
- ▶ 電泳緩衝液
- ▶ Intercalating dye 存在與否

膠體濃度及電壓的影響



Ausubel, F. M., Brent, R., Kingston, R. E., Moore, D. D., Seidman, J. G., Smith, J. A. and Struhl, K. (1987) Current protocols in molecular Biology. Fig. 2.5A.2.

■ 常用定量 DNA 的方法：



- ▶ 測定 260 nm 吸光值
- ▶ Hoechst 33258 定量法
- ▶ Ethidium bromide 法

測定 260 nm 吸光值


$$\text{Beer-Lambert Law: } A = \epsilon bC$$

$$\epsilon_{260\text{nm}} \text{ of ss DNA} = 0.027 (\mu\text{g/ml})^{-1}\text{cm}^{-1}$$

$$\epsilon_{260\text{nm}} \text{ of ds DNA} = 0.020 (\mu\text{g/ml})^{-1}\text{cm}^{-1}$$

$$\epsilon_{260\text{nm}} \text{ of ss RNA} = 0.025 (\mu\text{g/ml})^{-1}\text{cm}^{-1}$$

註：此處之 $\epsilon_{260\text{nm}}$ 為平均值，且不適用於 oligonucleotides

■ 常用定量 DNA 的方法：靈敏度比較

Table A.3D.3 Properties of Absorbance and Fluorescence Spectrophotometric Assays for DNA and RNA

Property	Absorbance (A_{260})	Fluorescence	
		H33258	EtBr
Sensitivity ($\mu\text{g/ml}$)			
DNA	1-50	0.01-15	0.1-10
RNA	1-40	n.a.	0.2-10
Ratio of signal (DNA/RNA)	0.8	400	2.2

Ausubel, F. M., Brent, R., Kingston, R. E., Moore, D. D., Seidman, J. G., Smith, J. A. and Struhl, K. (1994) Current protocols in molecular Biology. John Wiley & Sons, Inc. Table A.3D.3.