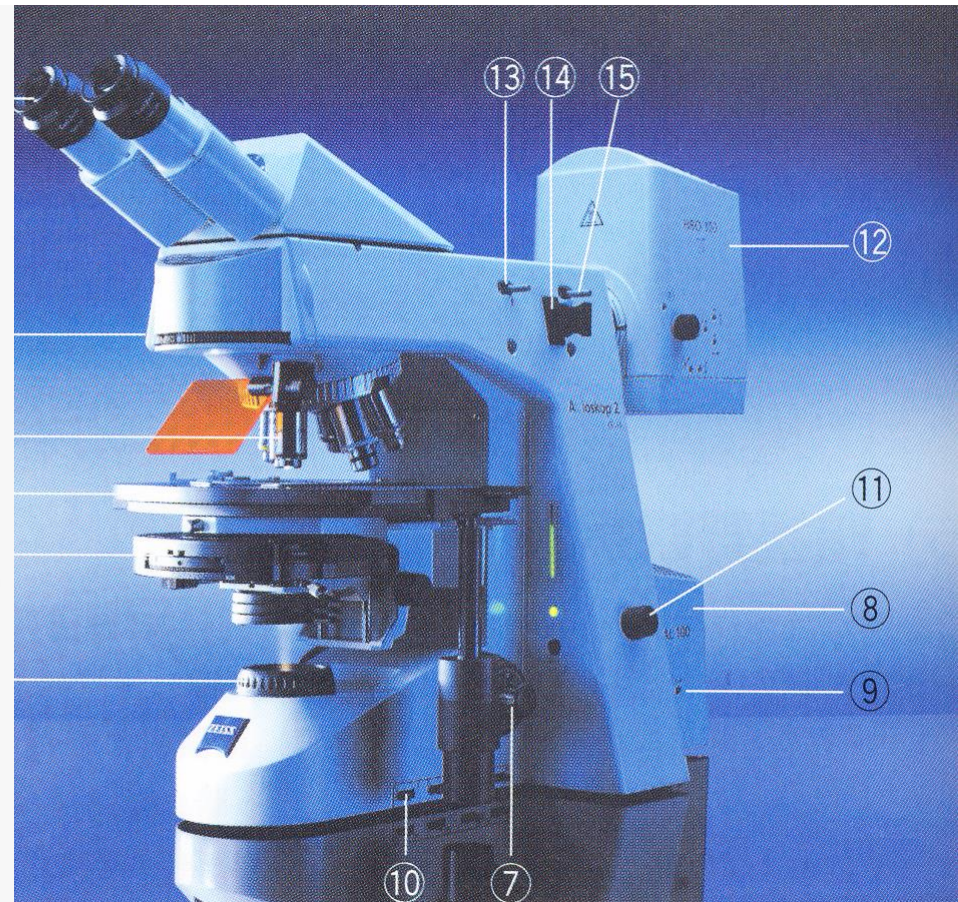
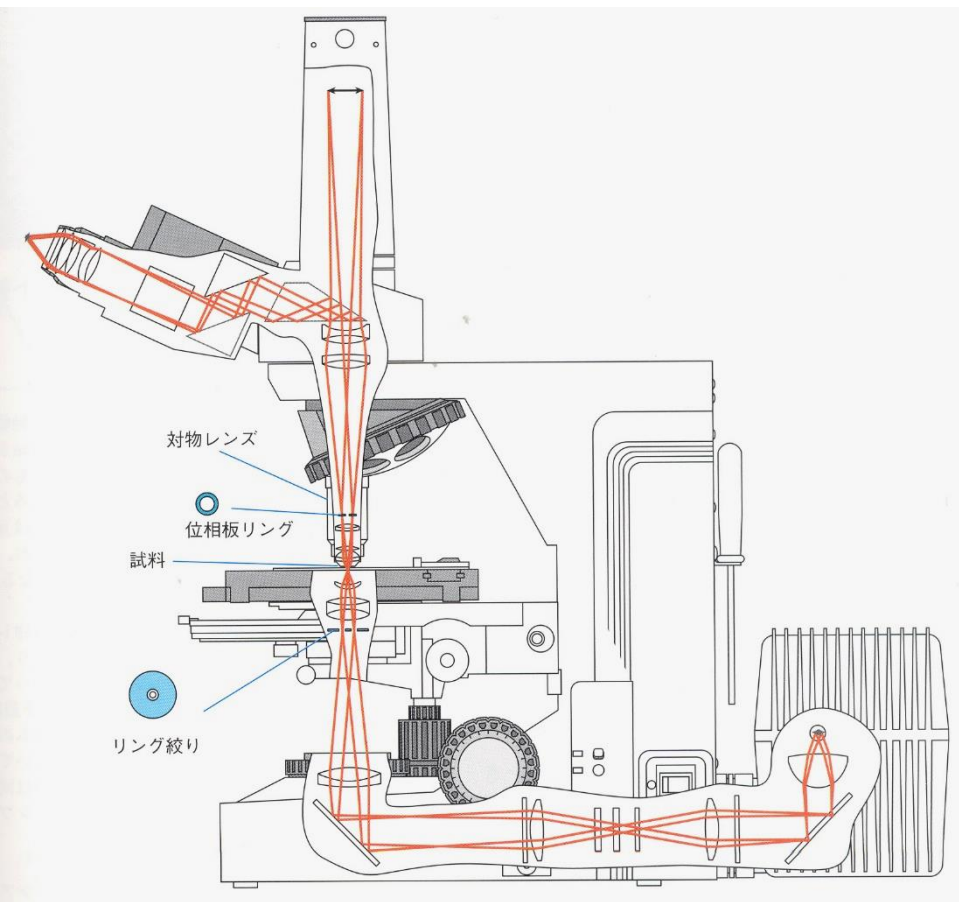


光學顯微鏡 基本原理及運用



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解剖學暨細胞生物學研究所
2025

光學顯微鏡



顯微鏡的分類

正立顯微鏡：觀察組織切片

倒立顯微鏡：觀察培養活細胞

實體顯微鏡：解剖及立體定位

第1章 顯微鏡的分類

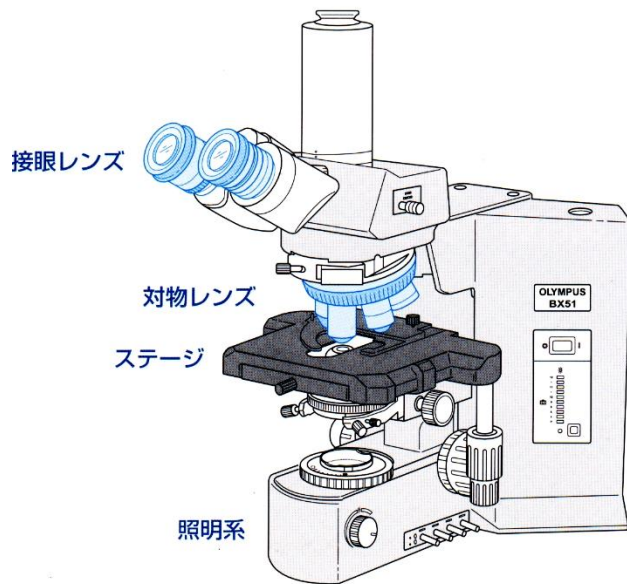


図1 正立型顯微鏡

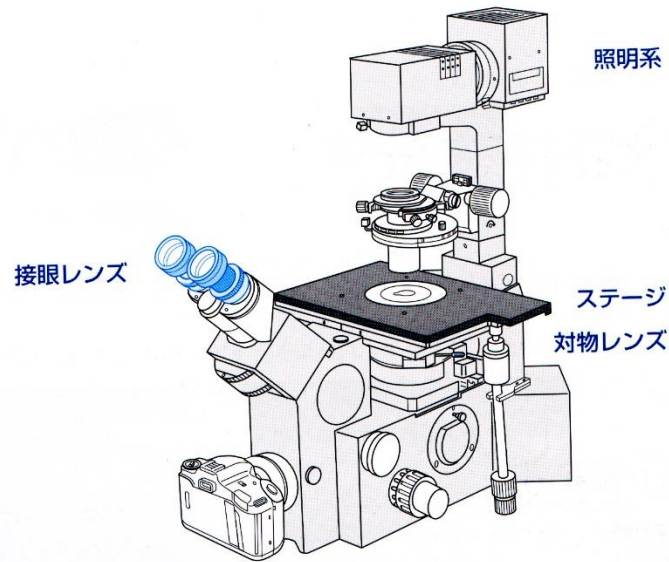
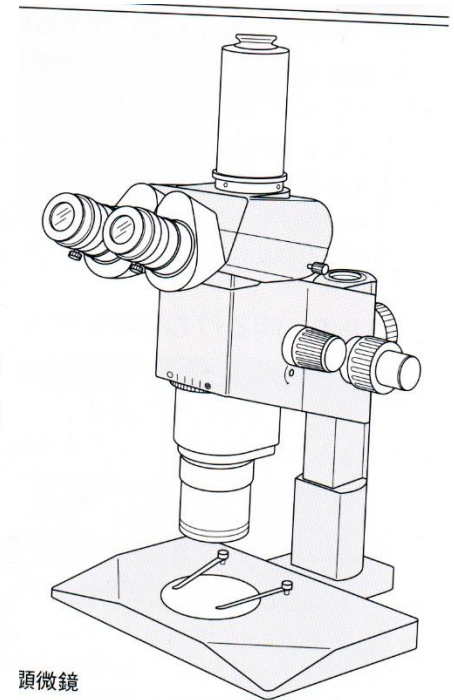


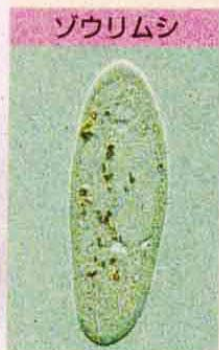
図2 倒立型顯微鏡



D いろいろな顕微鏡と装置

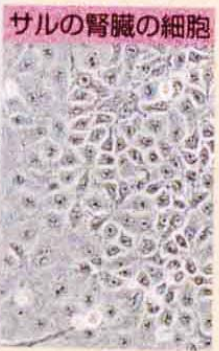
試料をいろいろな面から観察できるような光学顕微鏡と付属装置が開発されている。

●微分干涉顕微鏡



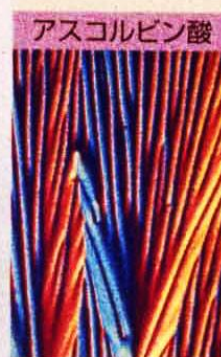
細胞分裂・原形質流動・アメーバ運動・繊毛運動の観察に使用。

●培養倒立顕微鏡



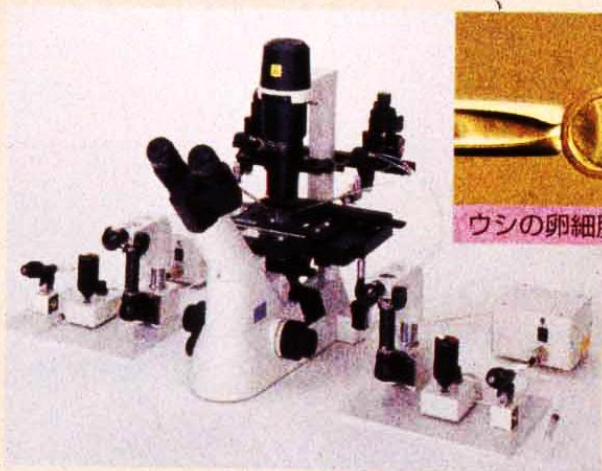
培養容器に入った細胞などの観察に使う。

●偏光顕微鏡



偏光板を利用した顕微鏡で、筋繊維などの微細構造の観察に使用。

●マイクロマニピュレーター



顕微受精などに利用する付属装置。

●蛍光顕微鏡



蛍光色素で抗体などを着色し、試料の発する蛍光を観察する。

●実体顕微鏡

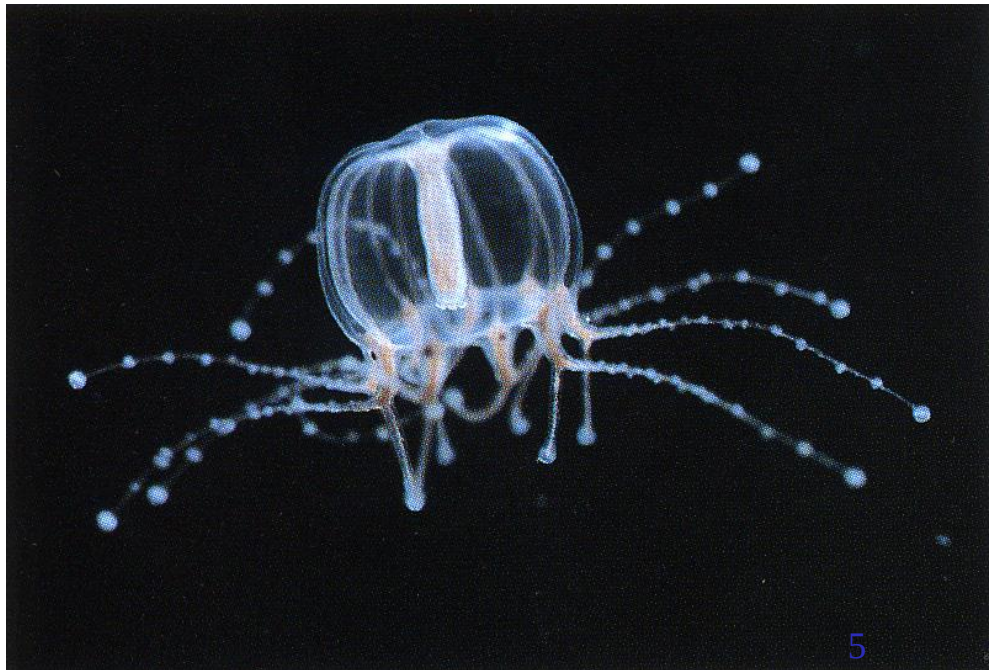
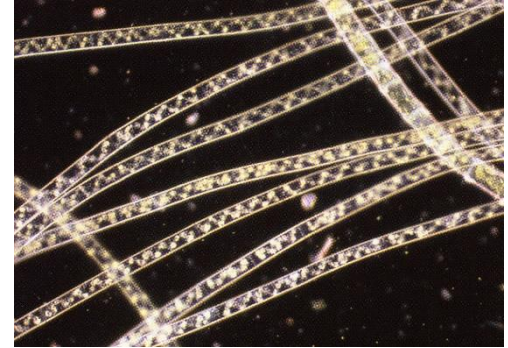
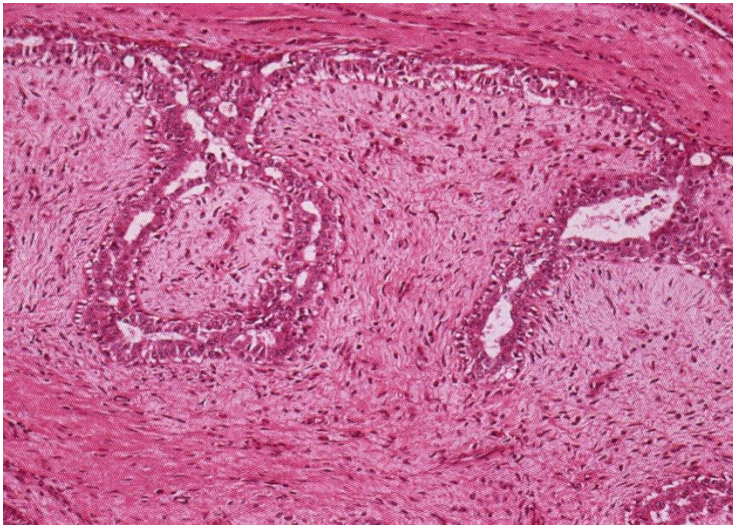
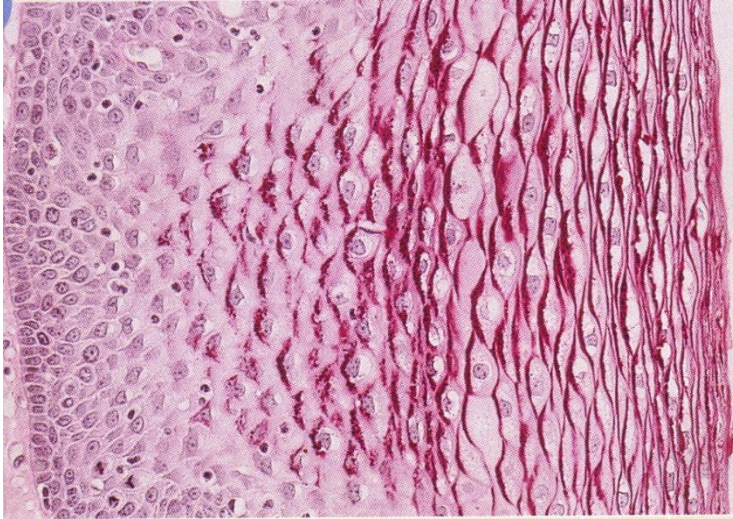


生試料を正立像で立体的に観察できる。

觀察法的顯微鏡分類

明視野 (bright field)：一般光學組織切片

暗視野 (dark field)：小動物如線蟲及纖毛鞭毛結構

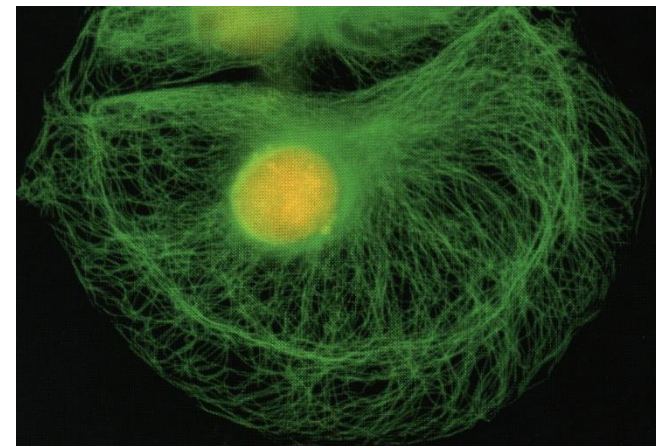
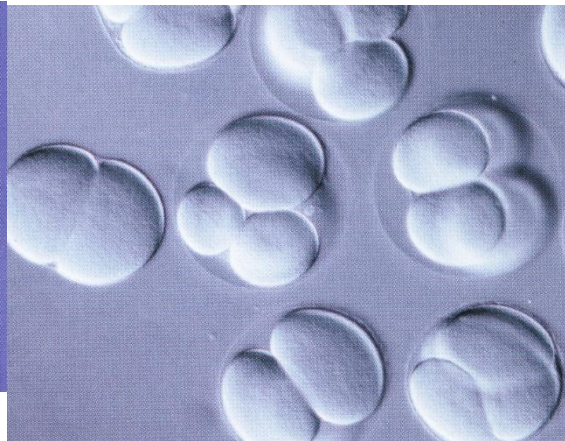
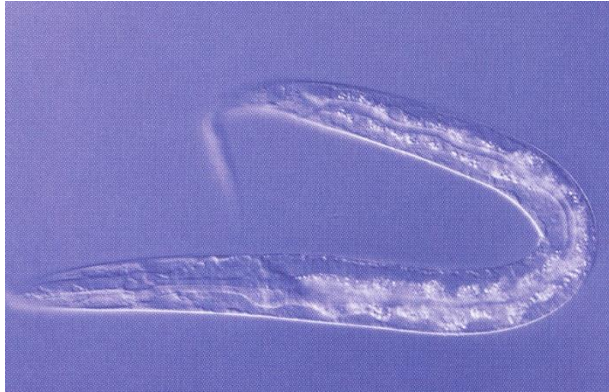
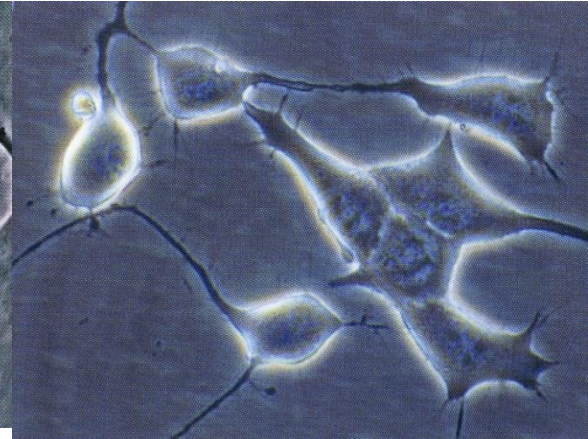
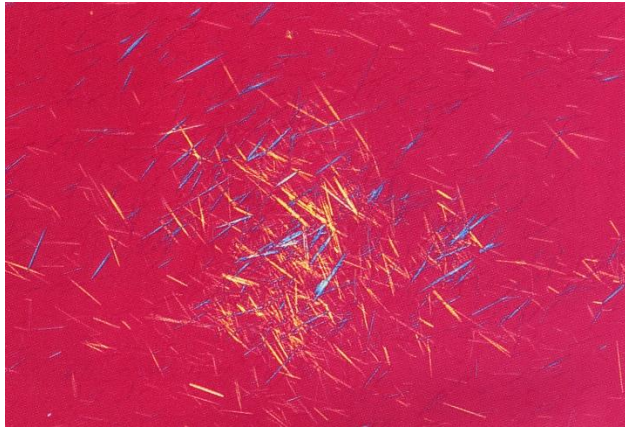


偏光 (polarized light)：堅硬的結晶結構等

位相差 (phase contrast)：一般培養活細胞

微分干涉 (differential interference contrast, DIC)：活體卵細胞及線蟲等

螢光 (fluorescence)：一般螢光染色標示或活體螢光如GFP等



1. **穿透光照明**：一般光學組織切片，薄具透光性；對比與染色
2. **反射照明**：解剖及實體顯微鏡，螢光顯微鏡；表面光源反射

1. **穿透光照明**：一般光學組織切片，薄具透光性；對比與染色
2. **反射照明**：解剖及實體顯微鏡，螢光顯微鏡；表面光源反射

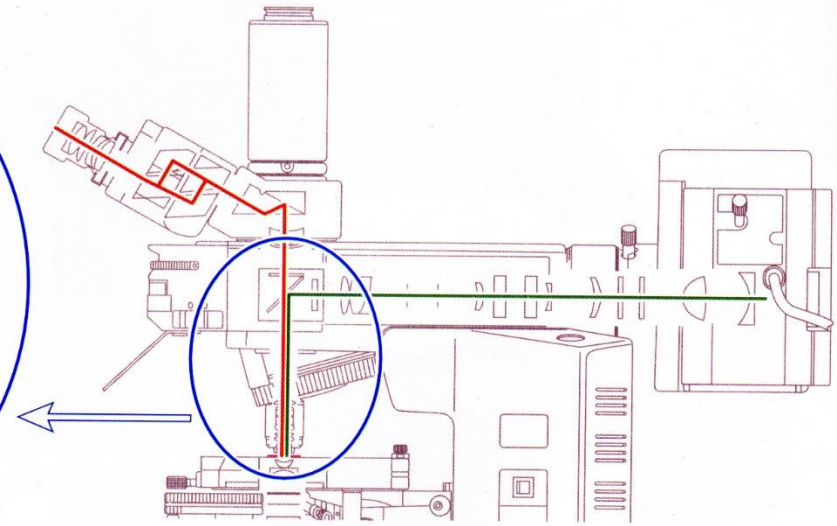
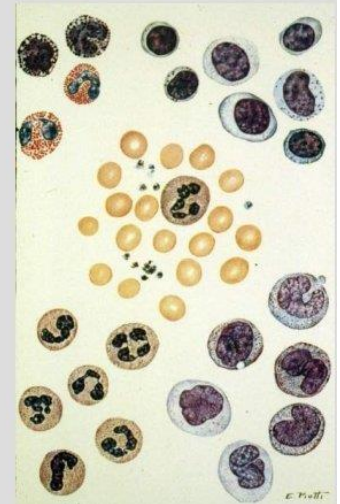


TABLE 1.3. Eye versus Instrument Resolution

	DISTANCE BETWEEN RESOLVABLE POINTS
Human eye	0.2 mm
Bright-field microscope	0.2 μm
SEM	2.5 nm
TEM	
Theoretical	0.05 nm
Tissue section	1.0 nm

SIZE OF HUMAN BLOOD CELLS

CELL/PLATELET	SIZE
ERYTHROCYTES	6.5-8 μm
LEUKOCYTES (WBC)	
% of WBC	
NEUTROPHIL	12-15 μm
60-70%	
LYMPHOCYTE	6-18 μm
25%	
MONOCYTE	12-20 μm
5%	
EOSINOPHIL	12-15 μm
-4%	
BASOPHIL	12-15 μm
-1%	
PLATELETS	2-4 μm



AORTA

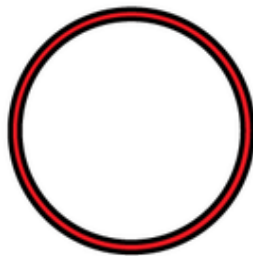
ARTERY

ARTERIOLE

CAPILLARY

VENULE

VEIN



Human

25,000/2,000

4,000/1,000

30/20

8/1

20/2

5,000/500

Mouse

535/55

150/50

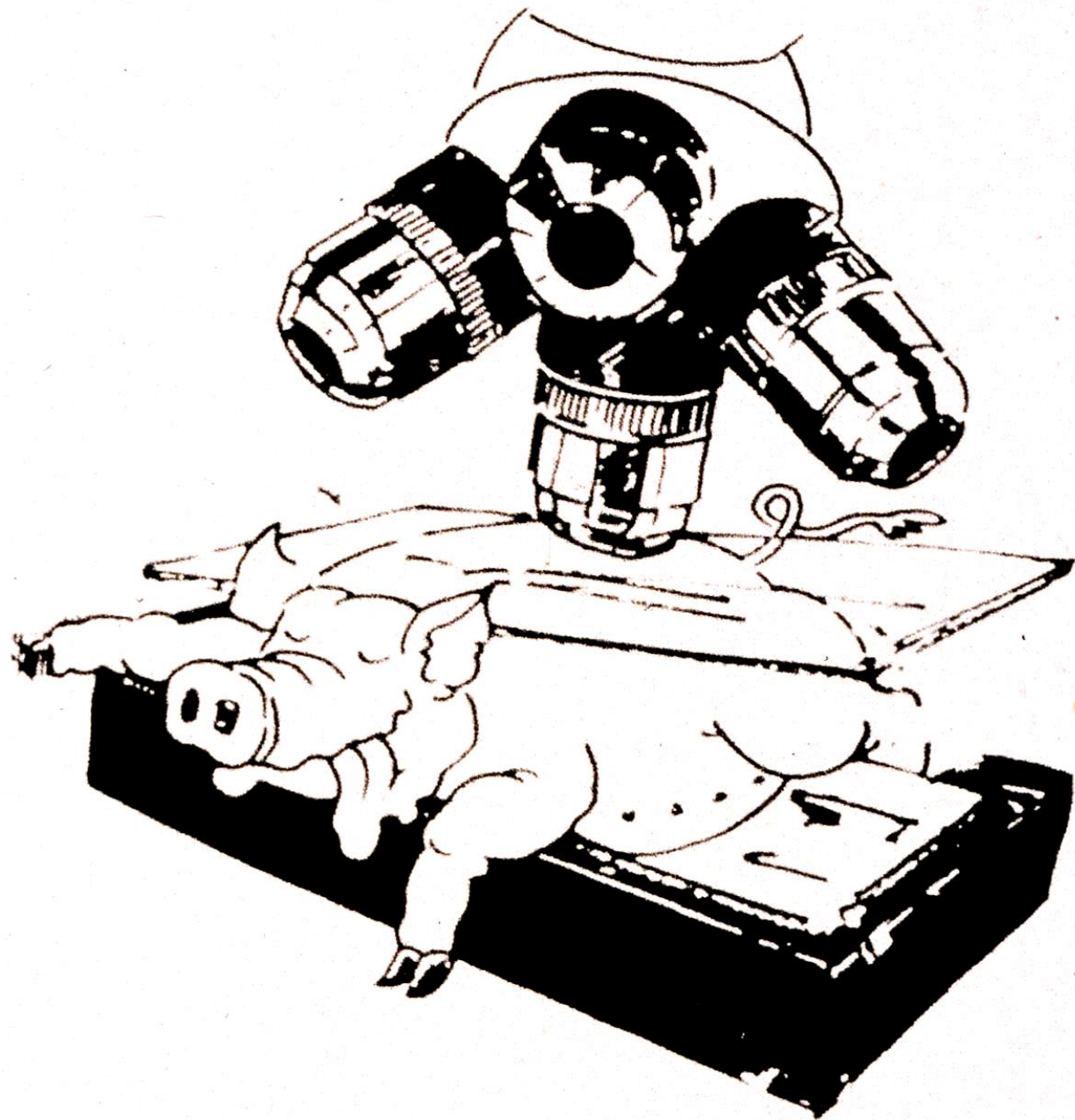
18/4.7

4/0.3

14/1

250/50

The typical diameters/wall thicknesses of blood vessel for humans and mice are given in micrometers. The smallest capillaries have diameters of a very few micrometers and a wall thickness of about one micrometer (μm).



(図：カールツァイス社)

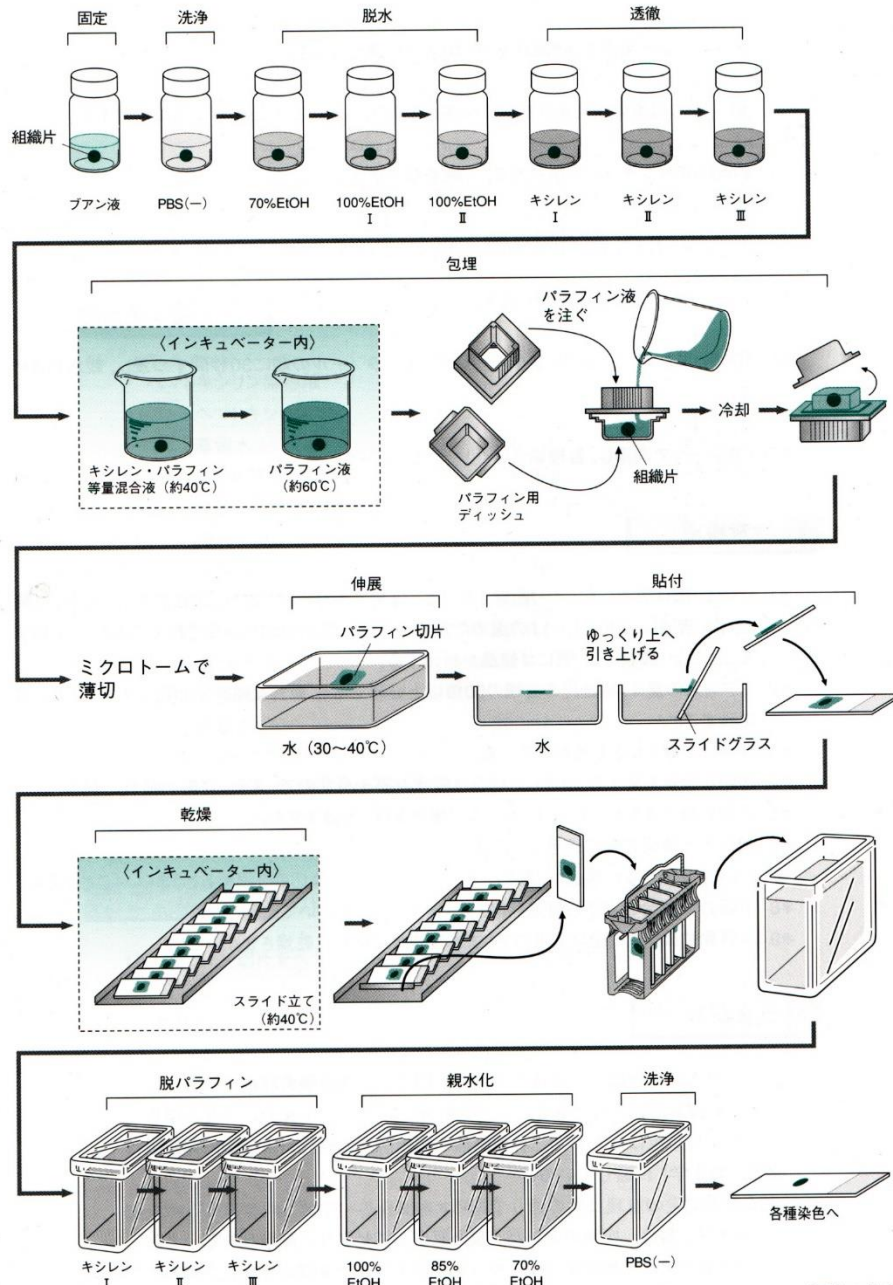


図2-8 パラフィン切片の作製

Cryostat sectioning



物鏡

- 開口值 /NA
- 消色差
- 工作距離
- 色帶
- 蓋玻片厚度



浸液表示色帯

イマージョン
オイル

水

グリセリン

複数



示色帯

0.5×, 1×
0.25×

2×

2.5×

4×, 5×

10×

16×, 20×
25×, 32×

40×, 50×

60×, 63×

100×以上

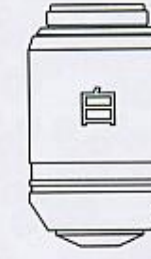
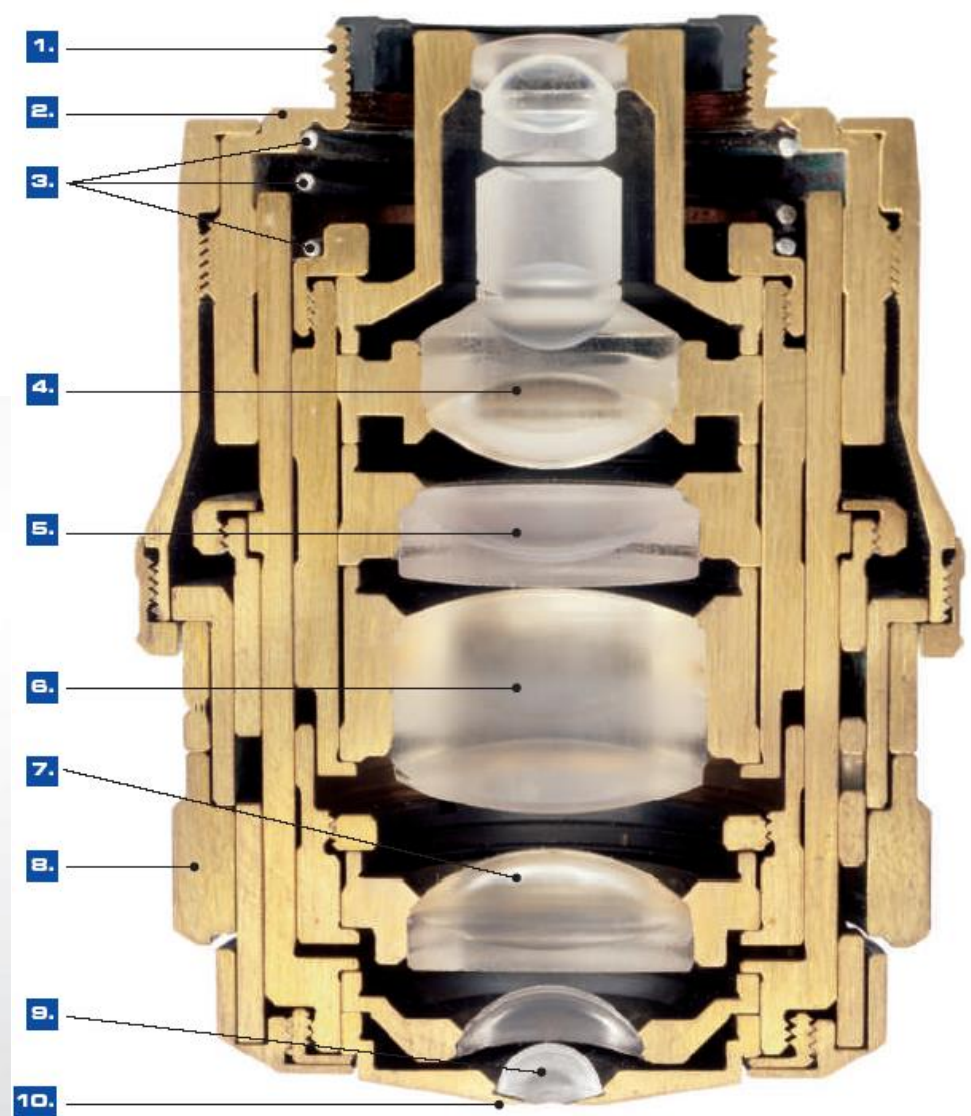


図3 対物レンズの外と内

A: 60×のプランアポクロマート油浸対物レンズの外表面。外表面に記されている文字は、明視野観察用では
目、位相差観察用では緑、偏光観察用では赤に色分けされている。また、機械的鏡筒長は対物レンズの

Lens:

Most important part of Microscope

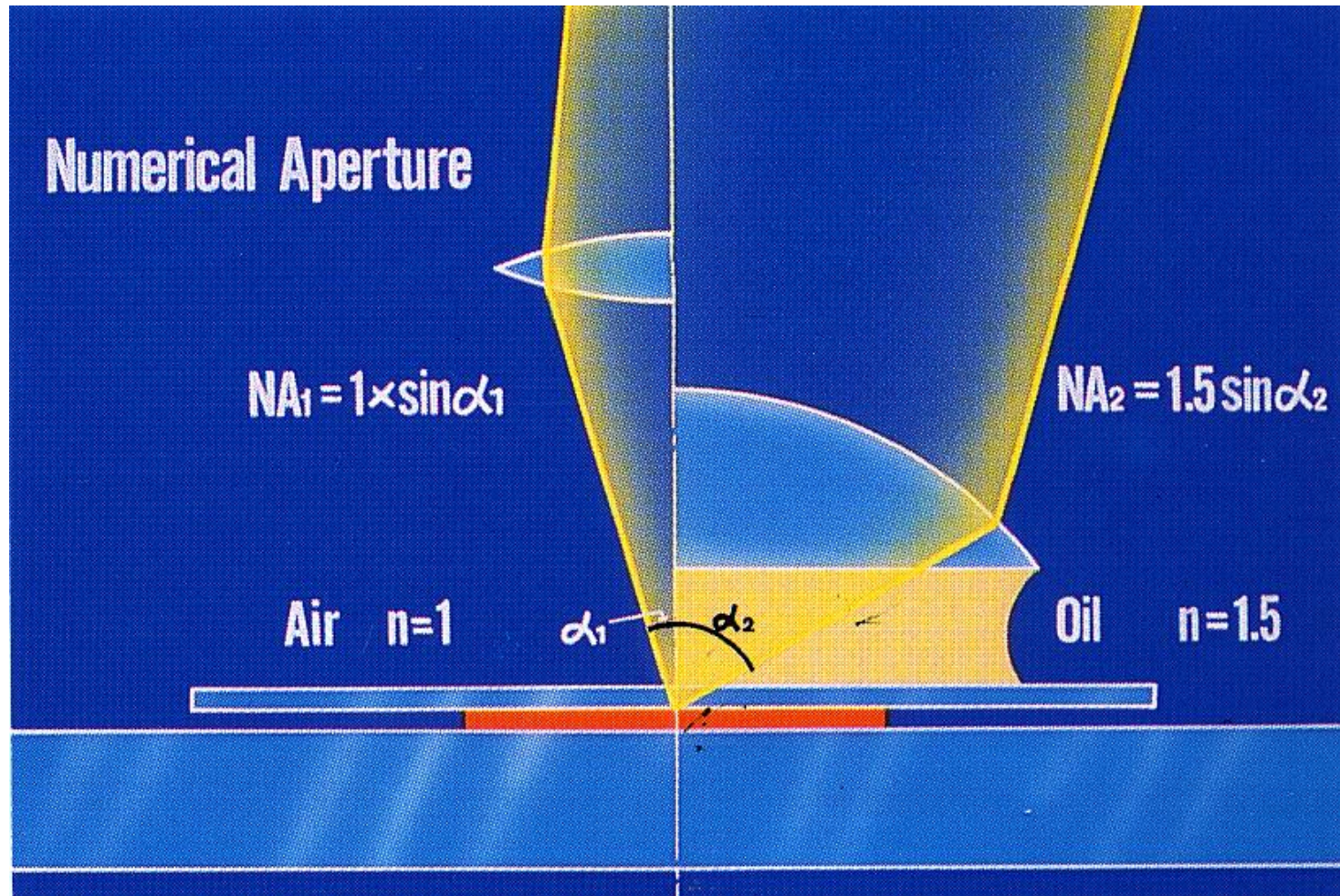


Cross section of an objective

- 1 Objective thread
- 2 Stop face of the objective
- 3 Spring system for the specimen-protection mechanism
- 4 - 7. Lens groups for the correction of image errors
8. Correction collar for adapting to deviating cover glass thicknesses or temperatures
9. Front lens system
10. Front lens holder

開口值 (NA)

- 開口值越大，鏡頭聚光能力越強，解析能力越強。



- Correction efficiency (UV – VIS – IR)

- N. A.

- Type (What's CS Objectives ?) - Quality

PL-APO (Delta, HCS)

PL-Fluotar

N Plan

C Plan

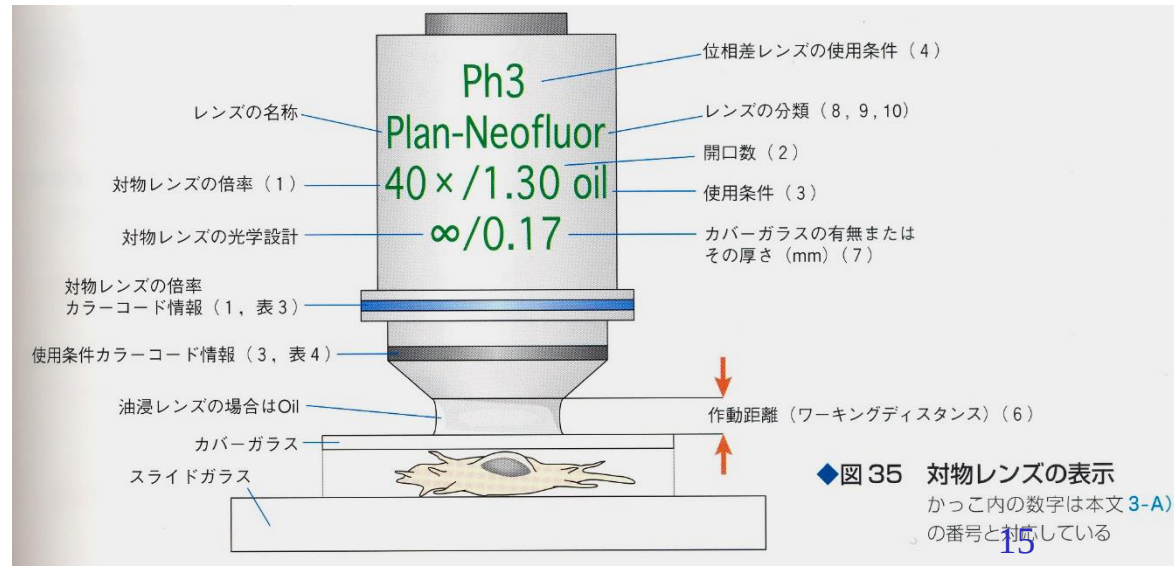
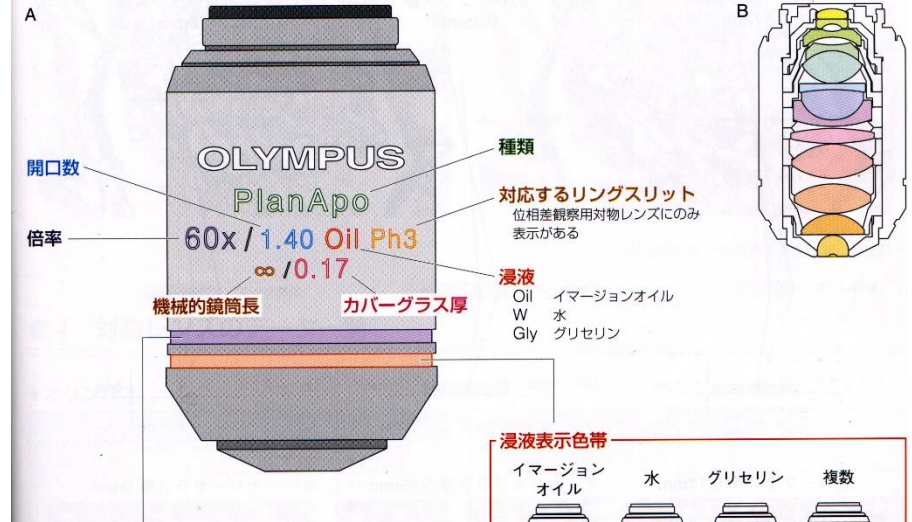
Air, Water, Oil (Imm)

Cover slip thickness
correction ?

Chamber Type ?

2. 対物レンズの外と内

対物レンズの外面には、種類、倍率、開口数、カバーガラス厚などが記されている。また、対物レンズの先端側には倍率と浸液を示す色帯(カラーリング)が付いており、外面の文字が見えなくてもカラーリングの色から倍率や浸液がわかるようになっている(図1A)。対物レンズの内部は、10枚前後のガラスが図1Bのように複雑に組み合わされている。



The Best Lens: Apochromat



Plan-APOCHROMAT: protects sensitive samples
Plan-APOCHROMAT objectives demonstrate top-class optical performance. They make it possible to see structures at the boundary of what is visible. Their outstanding performance features include: excellent correction, extremely high apertures and maximum resolution, color purity, contrast and flatness of field. All this combines to produce brilliant, needle-sharp images for observation, digital documentation and, in particular, fluorescence applications. The i Plan-APOCHROMAT of the 63x objective has been developed specifically for Live Cell Imaging – for optimal focus stability for time-lapse experiments.

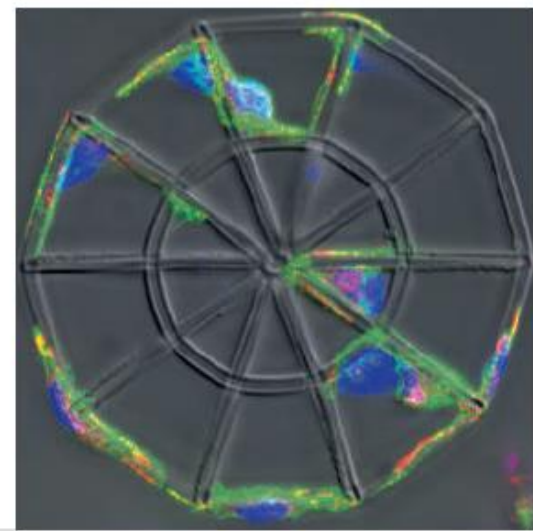


Image courtesy of Martin Bastmeyer und Franziska Klein, University of Karlsruhe, Germany

Resolution table using green light with
 $\lambda = 0.550 \mu\text{m}$:

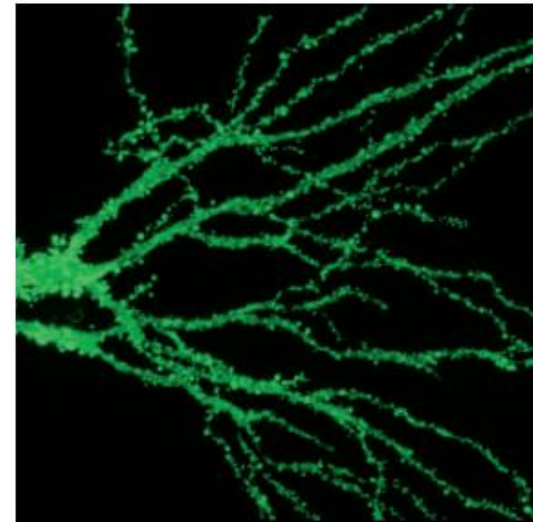
Magnification	/	NA	Resolution (μm)
10x	/	0.30	1.10
40x	/	0.75	0.45
63x	/	1.40 Oil	0.24
100x	/	1.30 Oil	0.26

- Field of view: 25 mm
- Flatness: ★★★★★
- Color correction: ★★★★★

Objectives with optimum correction of flatness of field and color; suitable for Digital Imaging



W Plan-APOCHROMAT: apochromatically correct
The immersion variant of the Plan-APOCHROMAT series – an addition to the water objectives of the ACHROPLAN class – has been specifically designed for electrophysiology. W Plan-APOCHROMAT objectives have apochromatic correction from visible light to the near infrared (VIS - IR) and are intended for use without a cover glass. Typical transmission values are greater than 80% from 450 nm to 1,000 nm and greater than 50% at 365 nm. These are also ideal prerequisites for use in 2-photon microscopy. The front of this slender objective is made of a special inert plastic that was originally developed for food technology.

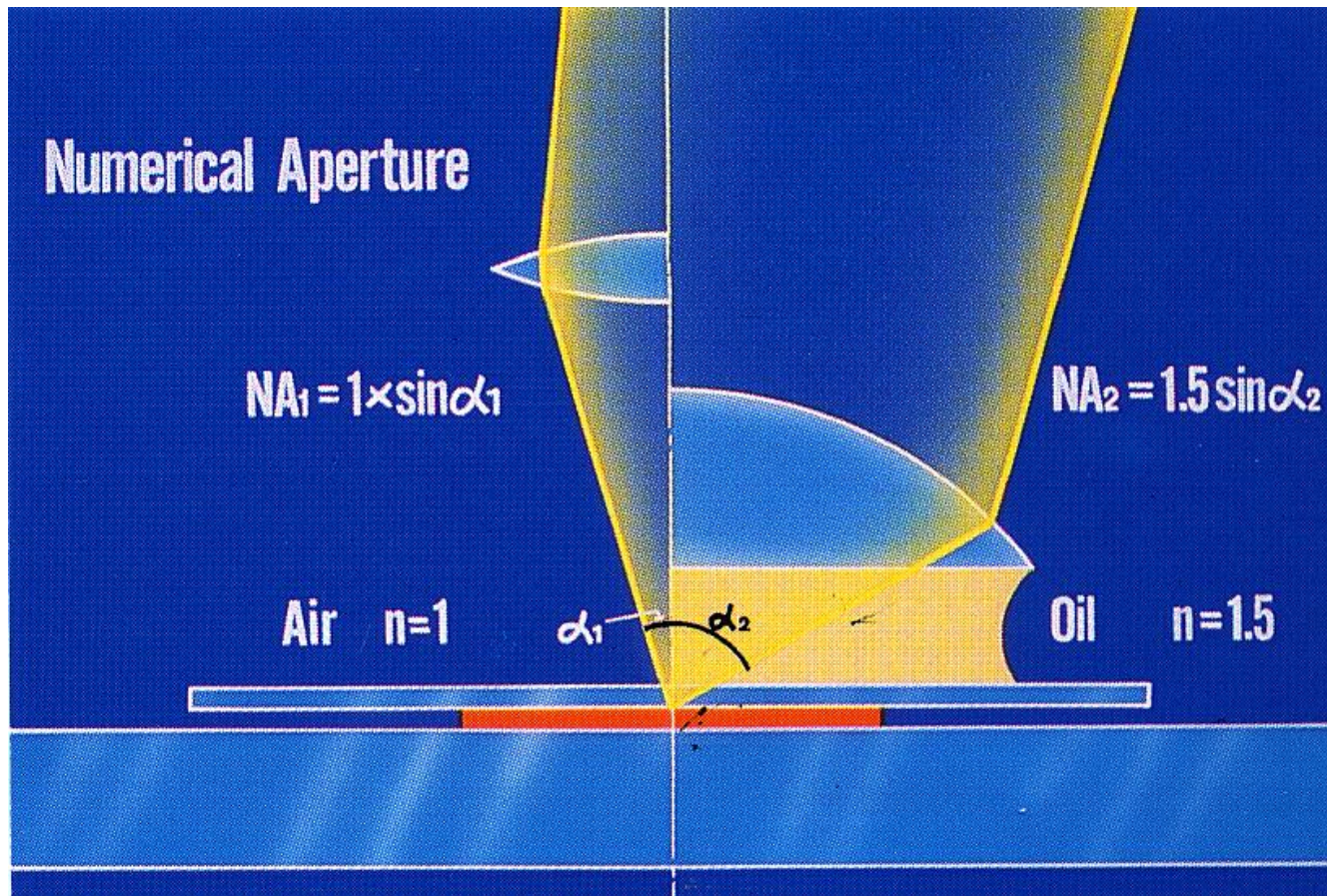


- Field of view: 20 mm
- Flatness: ★★★★★
- Color correction: ★★★★★

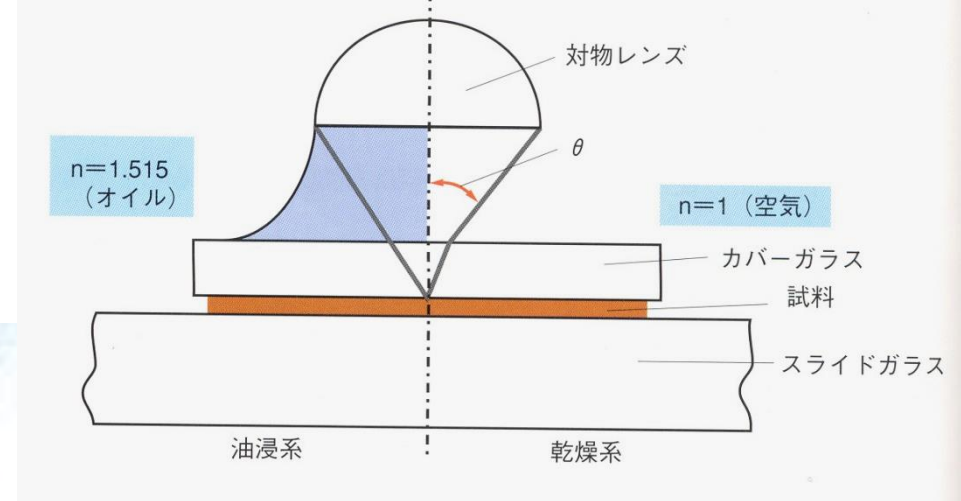
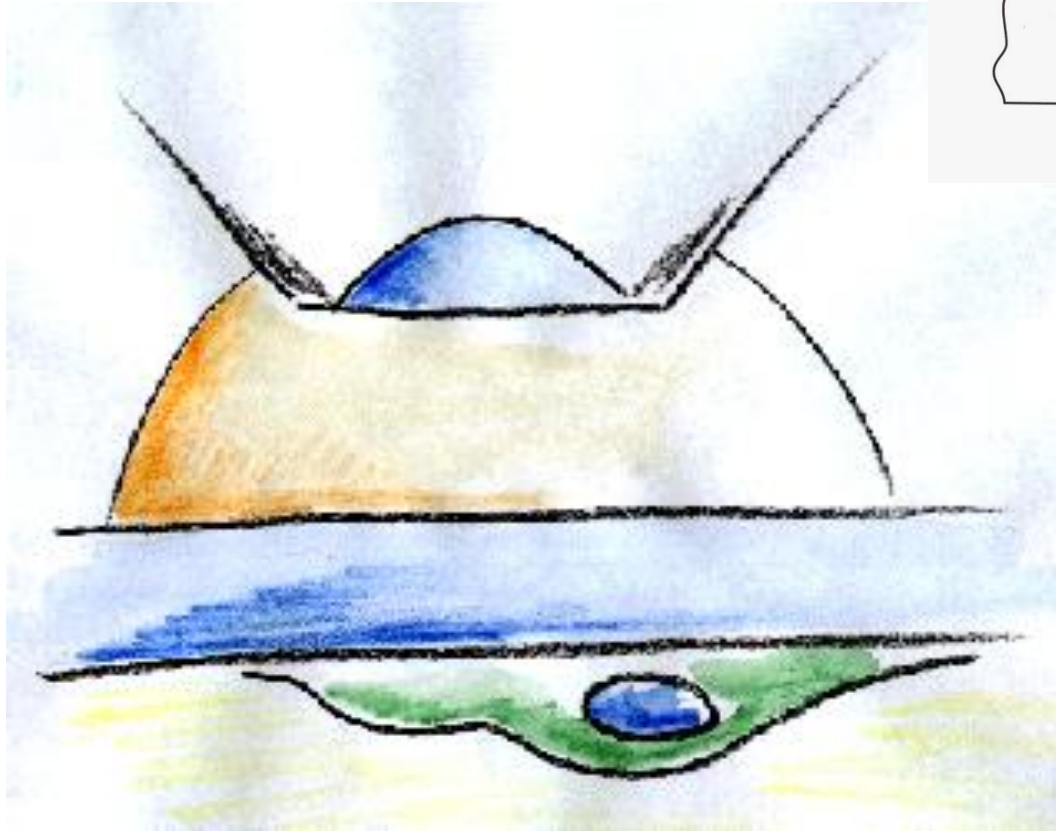
Apochromatically corrected immersion objectives for applications in physiology

介質的種類

- 乾式物鏡
- 油鏡
- 水鏡



- **Observation and optical setup**



Micro objective

Immersion

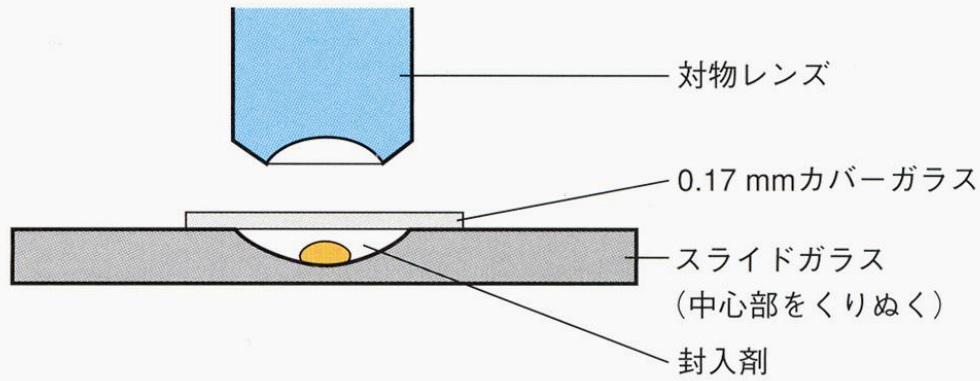
Cover slip

Sample

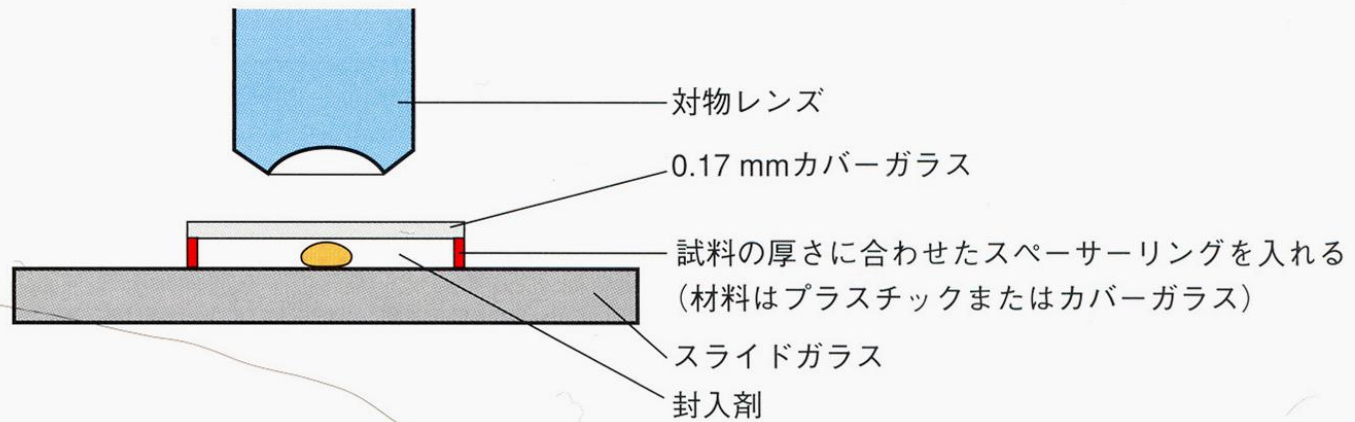
The last 3 optical elements are usually added by the user !

1) 正立型顕微鏡の場合

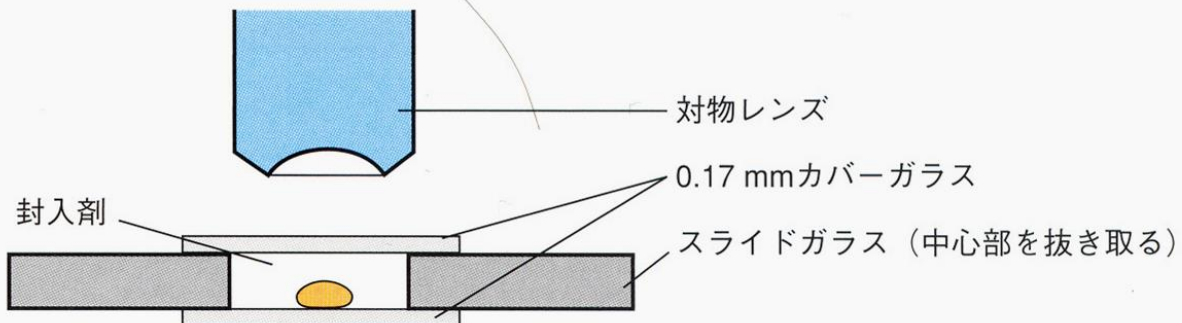
a)



b)

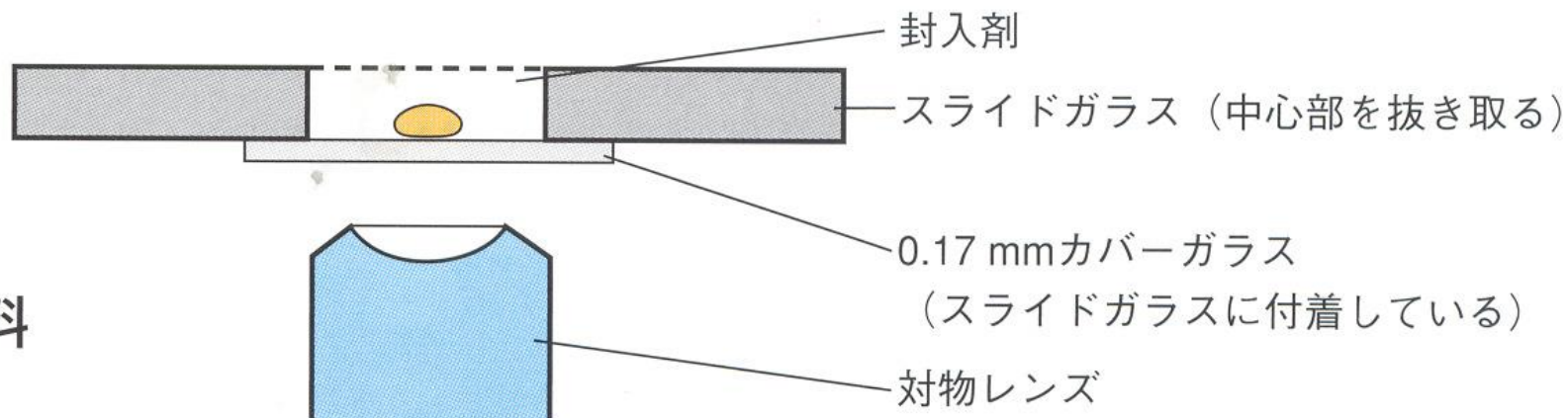


c)

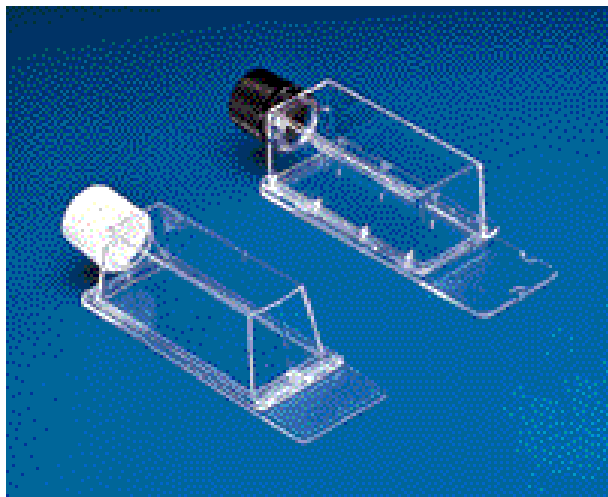


2) 倒立型顕微鏡の場合

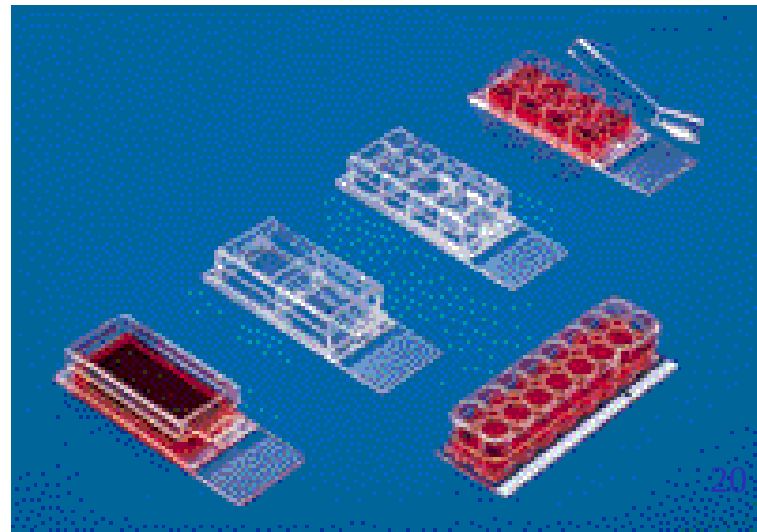
ための試料



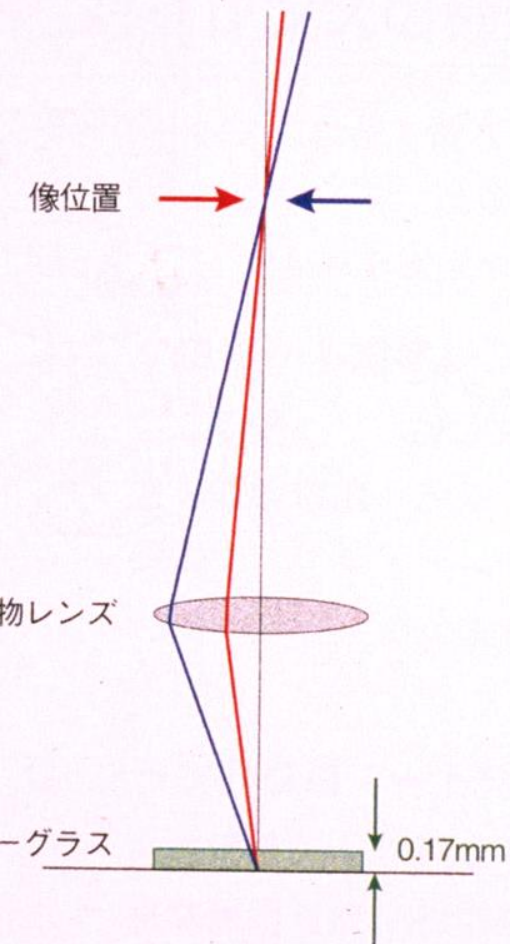
NUNC FLASKETTE® CHAMBER SLIDE/FLASKS



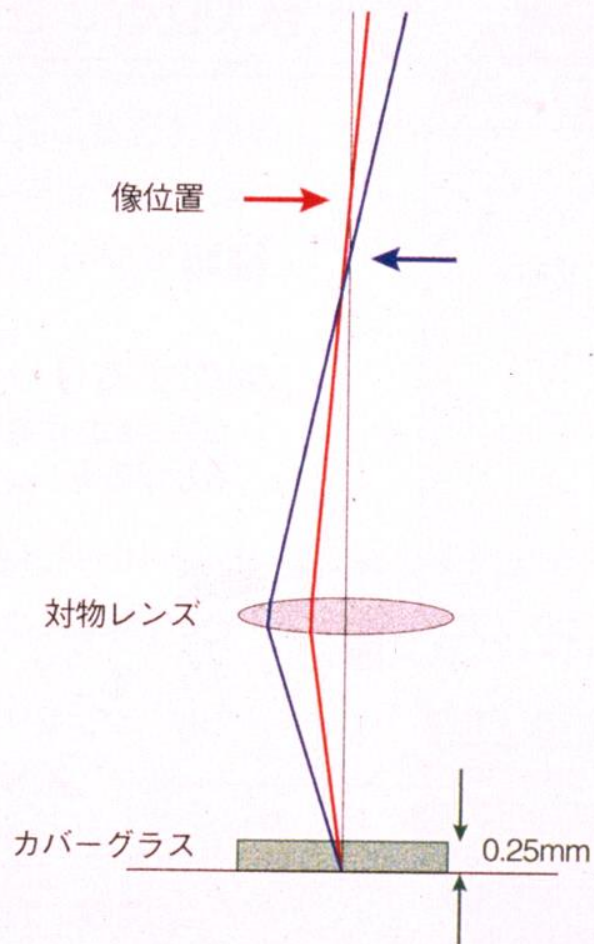
LAB-TEK® II CHAMBERED COVERGLASS



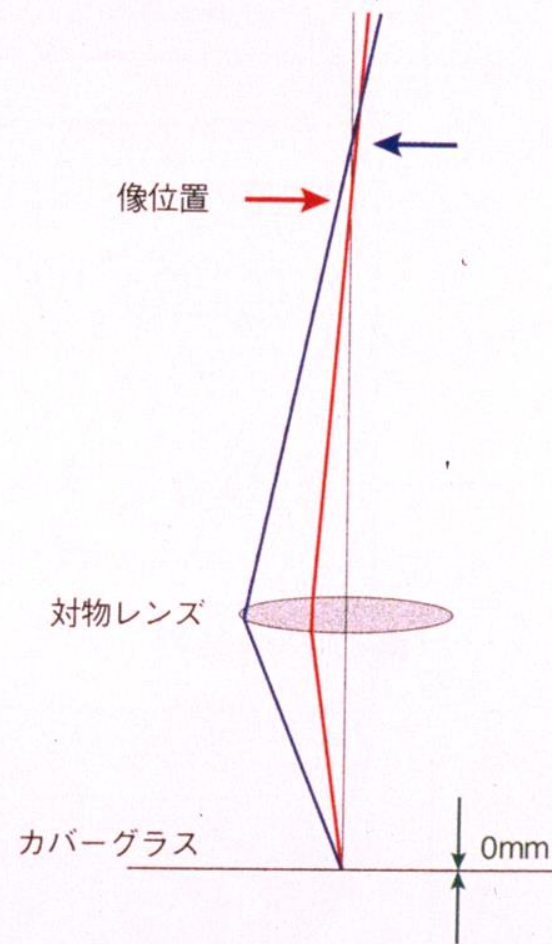
カバーガラス厚
0.17mm



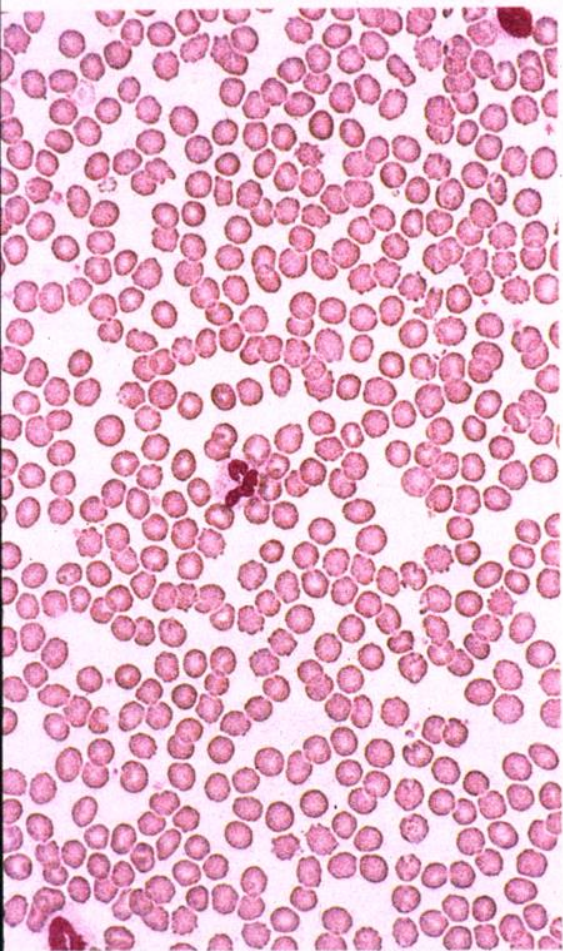
カバーガラス厚
0.25mm



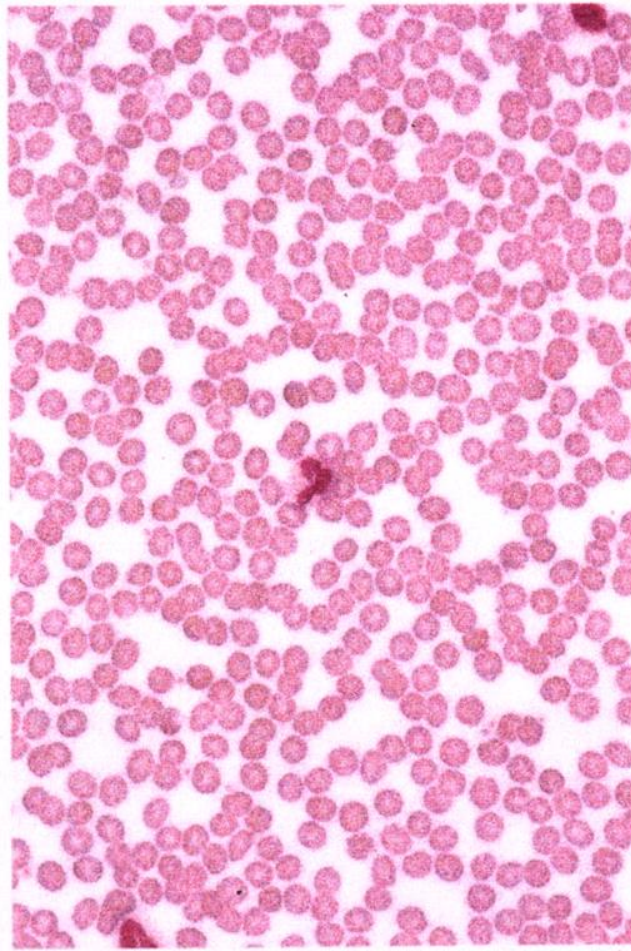
カバーガラス厚
0mm



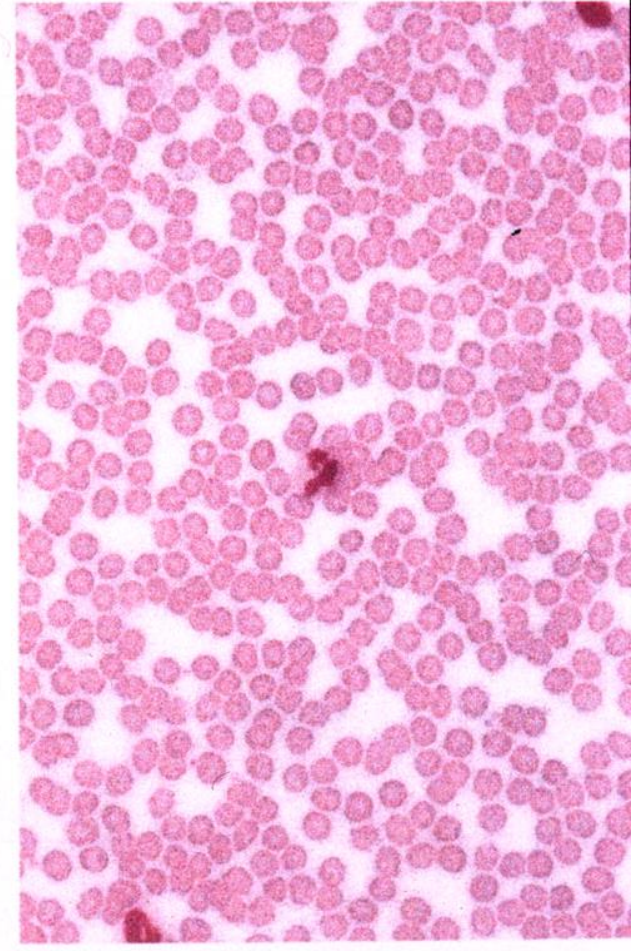
カバーガラス厚 0.17mm



カバーガラス厚 0.25mm



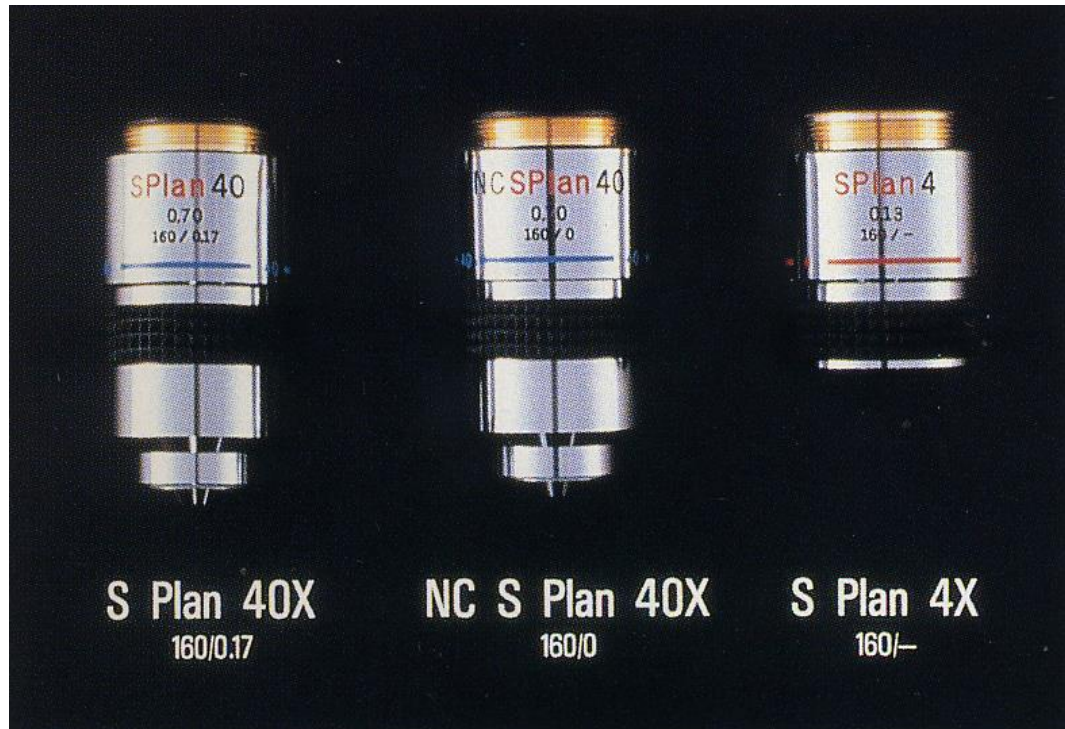
カバーガラス厚 0mm



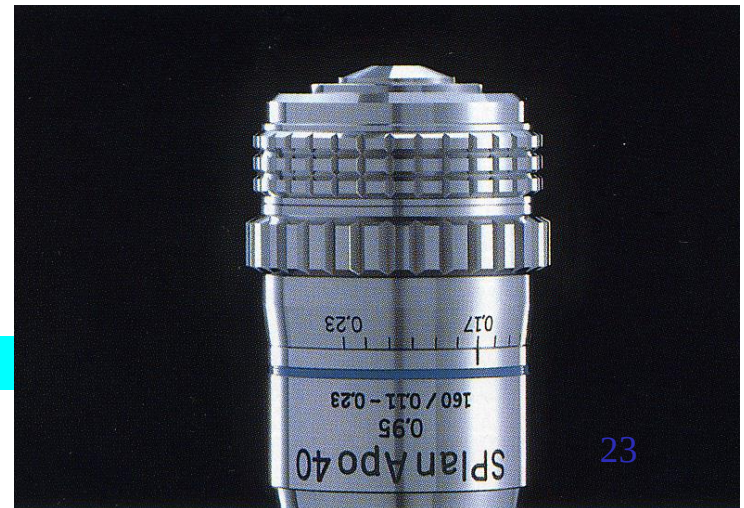
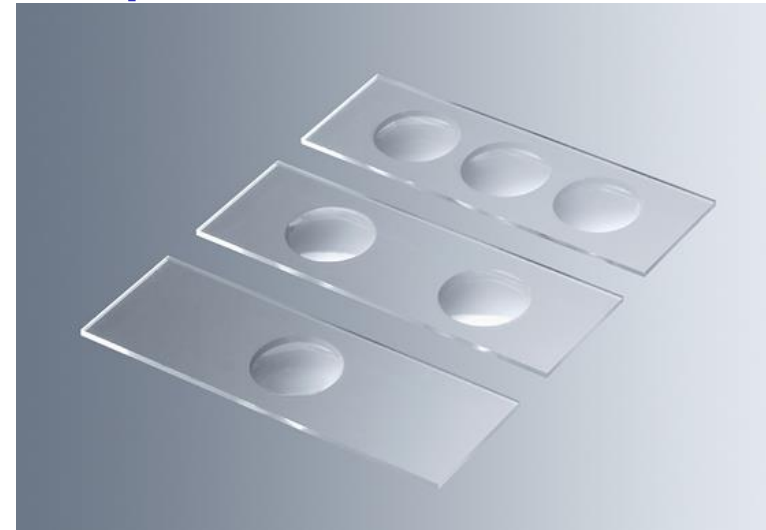
可修正蓋玻片的物鏡

- 高開口值物鏡需要可調整蓋玻片厚度，以取得最佳影像 · (0.11-0.23mm)

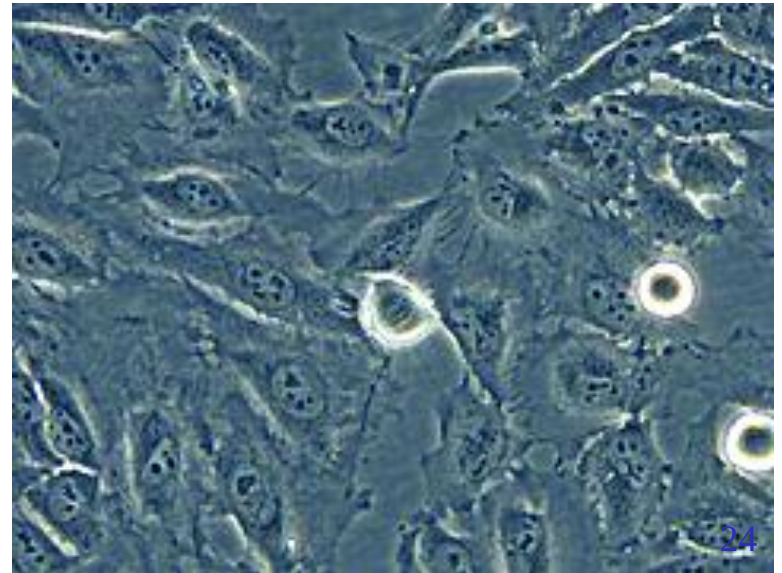
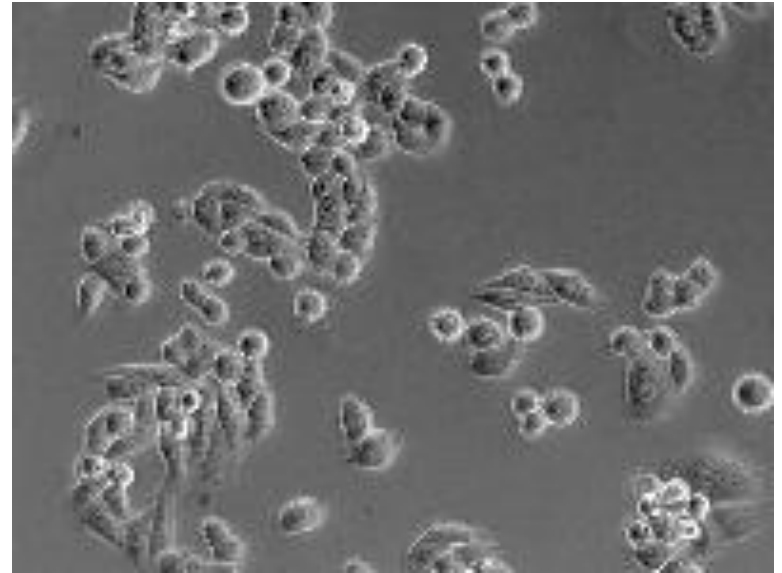
Depression slide



使用0.17mm 蓋玻片 不使用蓋玻片 蓋玻片厚度無關

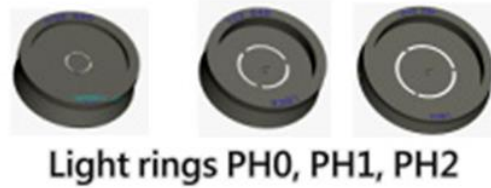


位相差顯微鏡 (Phase Contrast)

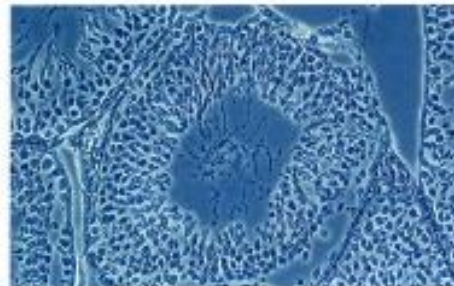
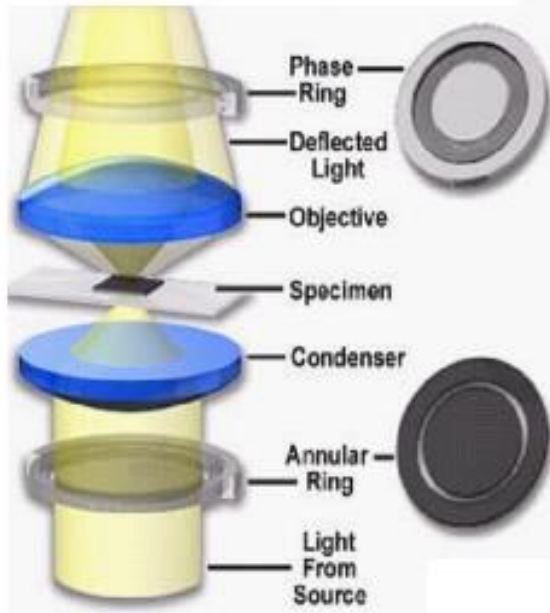


位相差的配件

- 位相差專用物鏡
- 位相差環
- 萬用聚光鏡
- 調整用目鏡



Phase contrast 光路 / 光學組件



Phase contrast



DIC

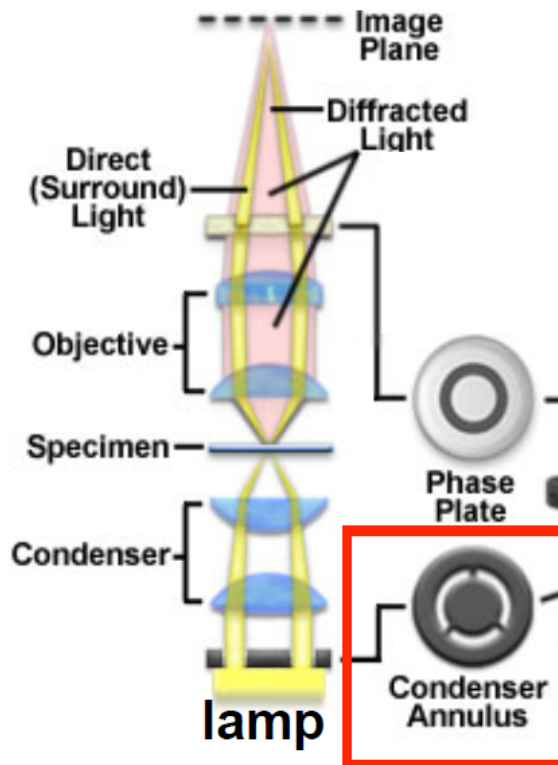
位相差的原理



Objective Apertures and Phase Contrast Optics



Phase Contrast Microscope Configuration



Restricts angles of illumination so diffracted and undiffracted light can be selectively modulated at phase plate

Technical specifications

Mateo TL base configuration

Light source	LED
Contrast methods	Transmitted light (brightfield and phase contrast)
Condenser	S40/0.45 condenser, working distance 50 mm
Phase turret	Encoded 3-position turret (BF, PH0, PH1)
Objective nosepiece	Encoded 4-position nosepiece
Preinstalled objectives	HI PLAN 4x /0.10 PH0 HI PLAN I 10x /0.22 PH1
Stage	Fixed stage (L*W) 262 mm x 212 mm
Focusing	Coarse and fine focusing, travel range 7 mm, min. adjustment 2 µm
Camera	Integrated CMOS, 6 megapixel color camera, sensor size 1 / 1.8", pixel dimensions 2.4 µm x 2.4 µm, sensor format H: 3072 px; V: 2048 px
Software	Stand-alone operating system. PC is not required.
Image output	JPEG and TIFF
Display	Color monitor with adjustable tilt, 15.6 inch, 1920 pixels x 1080 pixels
Onboard storage	Total storage 16 GB (Approximately 9 GB for the operating system and 7 GB for storing images and logfiles)
USB ports	1x USB 3.0 and 2x USB 2.0
Data transfer	USB or WIFI connection to smart device (optional)
Dimension (depth x width x height)	Monitor in display position: 310 mm x 376 mm x 530 mm Monitor in folded position: 310 mm x 376 mm x 385 mm
Weight	11 kg
Transportation aid	Supporting handle on the back of microscope
Other items included in the base configuration	Wireless mouse, mouse pad, dust cover, power adaptor, power cord, user manual and quick start guide

Optional accessories

Objectives	HI PLAN I 20x/0.30 PH1 HI PLAN I 40x/0.50 PH1
Object guide kit	Including > one attachable object guide > holding frame #1 for petri dishes: 35 mm & 60 mm, glass slides, chambered slides and chambered coverglasses > holding frame #2 for multi-well plates
Software module	Confluency module
WIFI dongle	WIFI Dongle 2.4 GHZ for data transfer

Mateo TL

Digital transmitted light inverted microscope for cell culture



Mateo TL makes the process of cell culture checks effortless for every team member, with intuitive setup and use.

It improves reproducibility for your experiments through objective confluency evaluation across individuals or experiments.

Benefits:

- > Simple to use, also for non-experts
- > Go from setup to first image in under 1 minute
- > One common criterion for consistent confluency measurements
- > Enhance your team's comfort



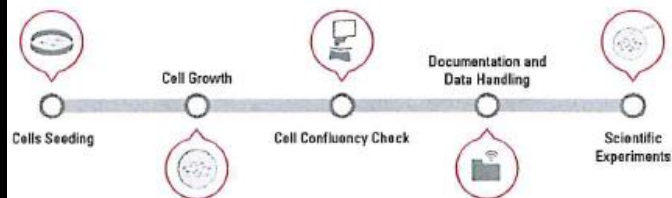
Enhance your team's comfort

Minimize fatigue by eliminating the need to use oculars

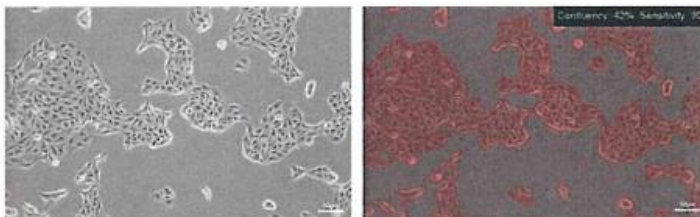
- > Mateo TL can be placed inside a laminar flow hood of compatible dimension for convenient cell culture work inside the hood, which also helps to minimize cell contamination.

Confluency Check Workflow with Mateo TL

In life science research, such as cancer, stem cells, or regenerative medicine, experiments often require cells in specific growth conditions. These conditions include cell morphology and confluency which are checked regularly.



For researchers who need consistent experimental outcomes, Mateo TL is an intuitive-to-use digital transmitted light inverted microscope that enables all lab members to check and document cell growth status conveniently and comfortably, and measure confluency consistently, thereby increasing the confidence in the success of their downstream experiments.



Phase-contrast image of a MDCK-cell culture and its respective confluency measured by Mateo TL.

MATEO FL

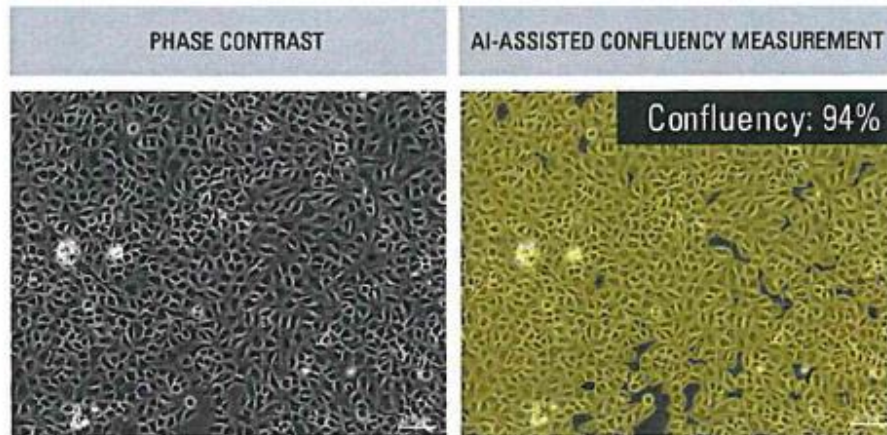
Digital fluorescence inverted microscope for advanced cell culture experiments



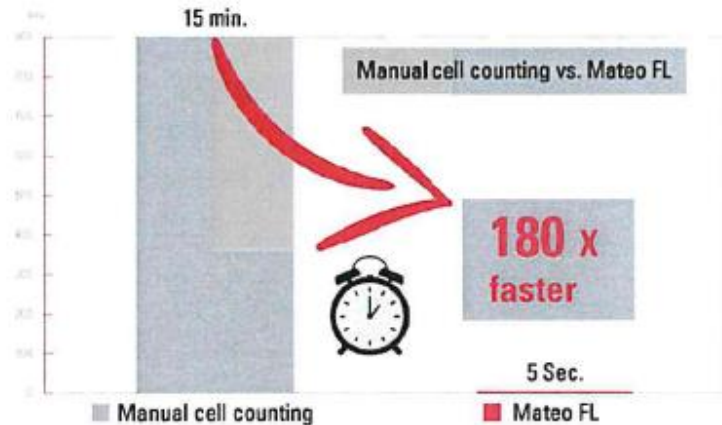
■ Make your experiments easier with AI-powered workflows

Boost reliability and consistency in your cell culture experiments with AI-powered algorithms and analysis.

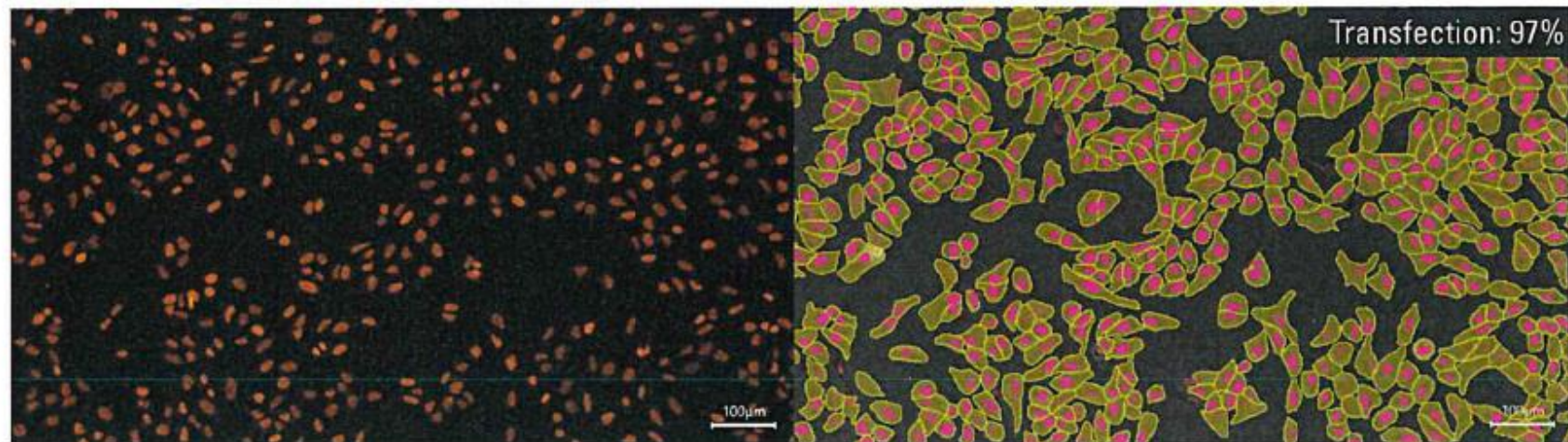
> Eliminate guesswork with AI-assisted confluency measurements and increase confidence in your downstream experiments.



> Get an accurate, automated cell count in 5 seconds, saving you an average of 15 minutes of manual cell counting.



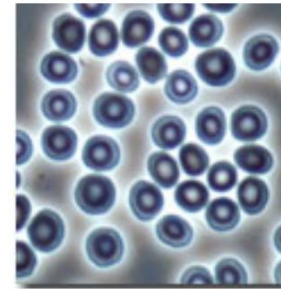
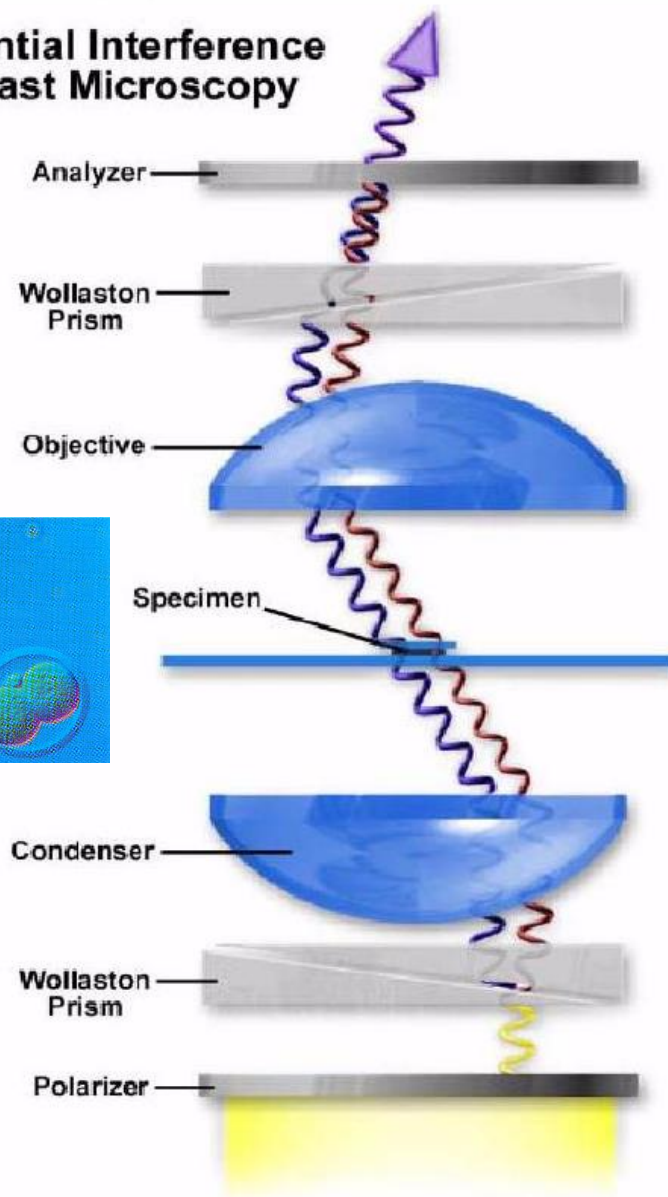
> Optimize your transfection process with smart insights.



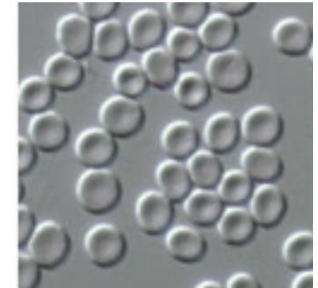
微分干涉顯微鏡

DIC: an alternative technique for enhancing contrast

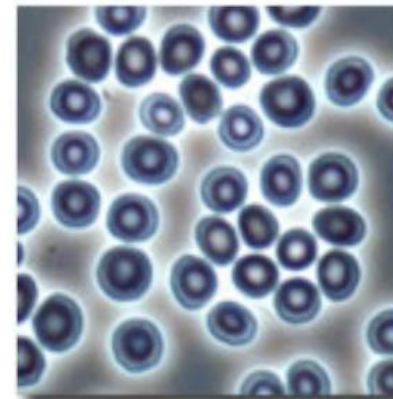
Differential Interference Contrast Microscopy



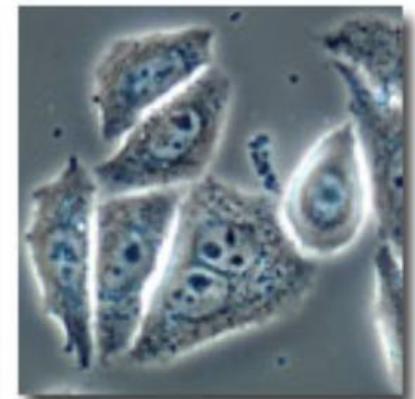
Phase



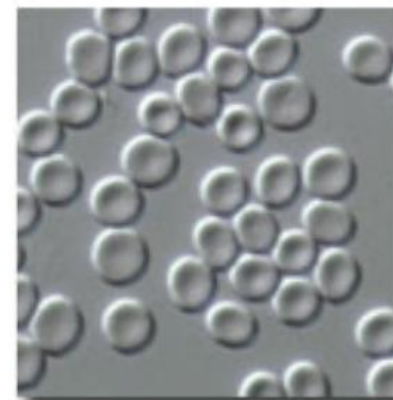
DIC (Differential Interference Contrast)



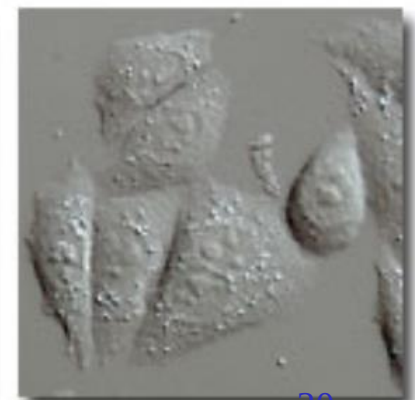
(a)



(c)



(b)



(d)

微分干涉顯微鏡的配件

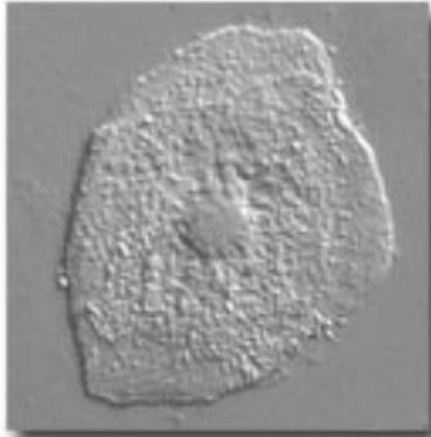
- 萬物聚光鏡
- 微分干涉鍍鏡
- 起偏鏡
- 檢偏鏡



微分干涉 位相差 的比較

	位相差	微分干涉
對比方式	以標本厚度轉換成對比	以厚度變化轉換成對比
對比的調整	無法調整	以鍍鏡調整
影像的特性	以明暗度表現	以立體感及明暗表現
解析度	比微分干涉差	高
合適的標本	細微構造 標本厚度10 μm	細微構造，外觀・ 標本厚度數百 μm
容器	可使用塑膠類	不可使用塑膠類

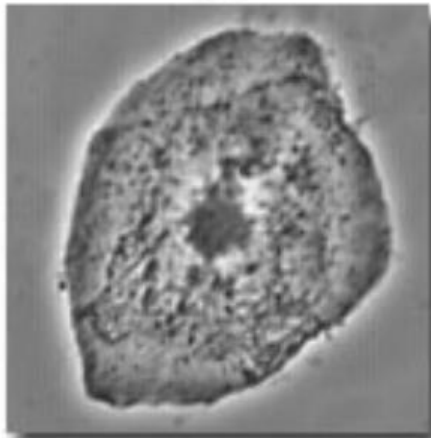
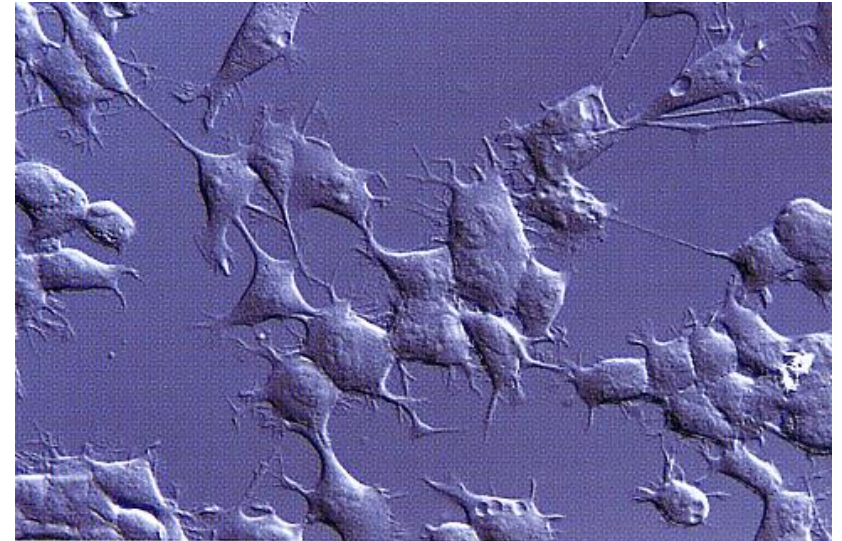
微分干涉及位相差的實例



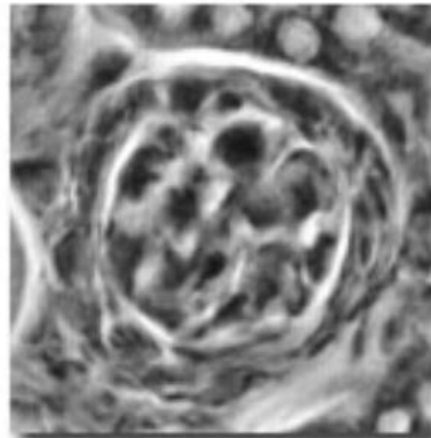
(a)



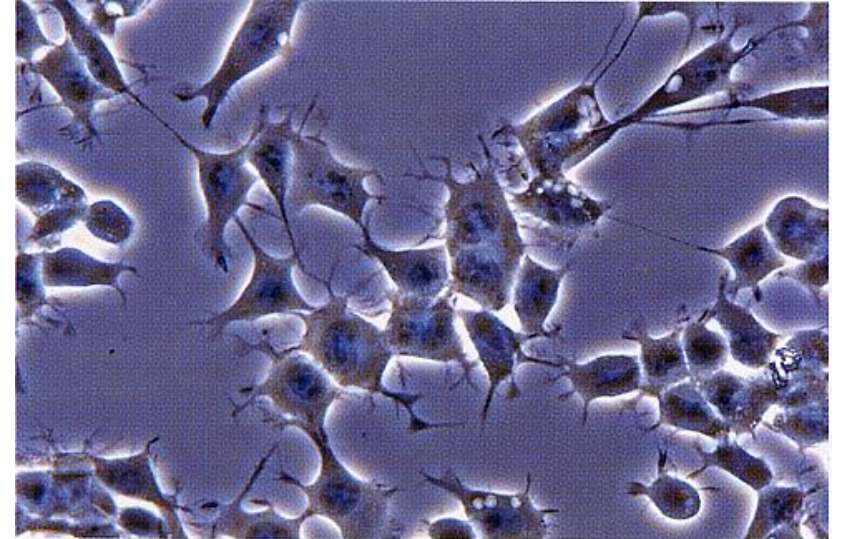
(c)



(b)

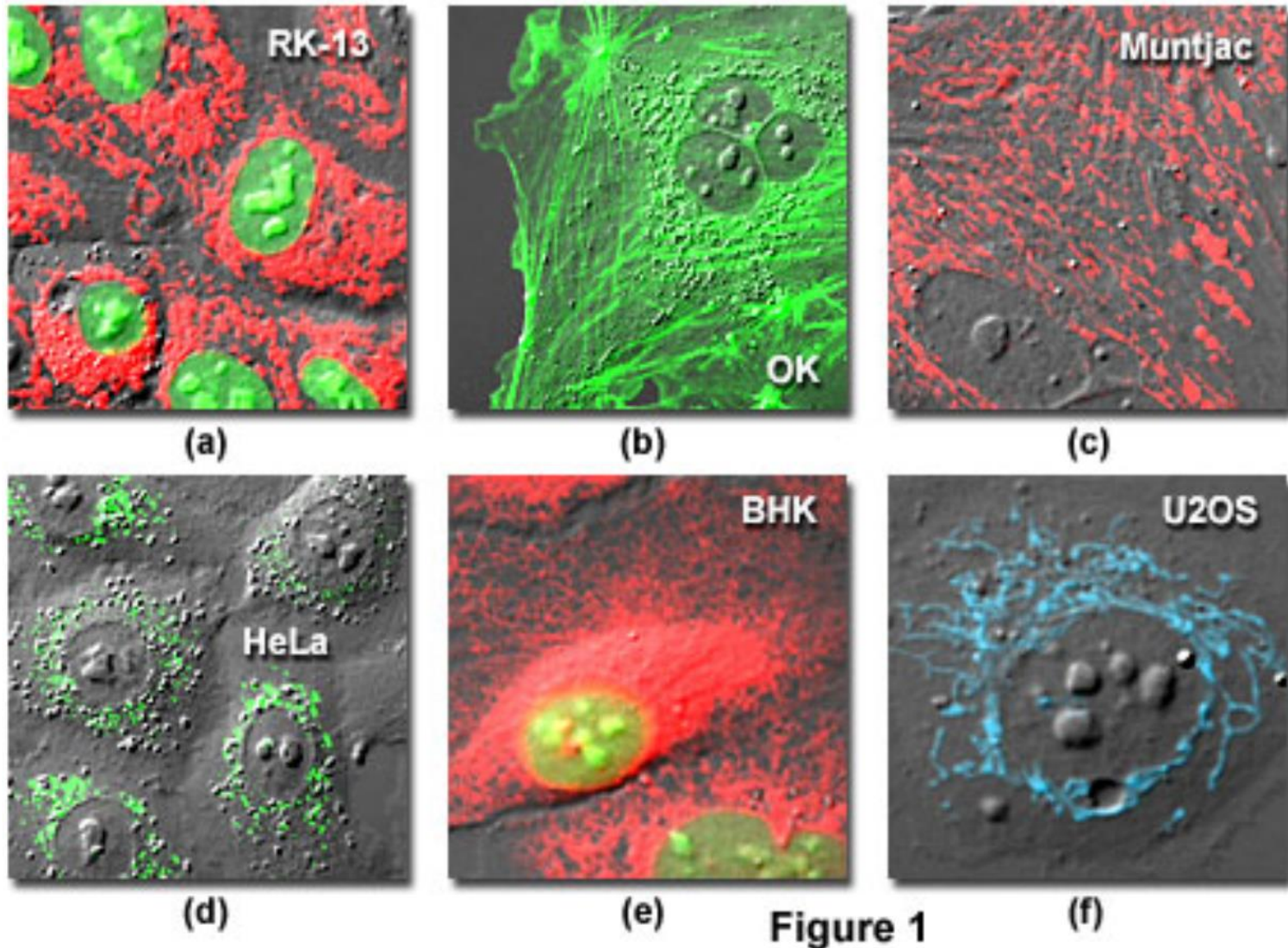


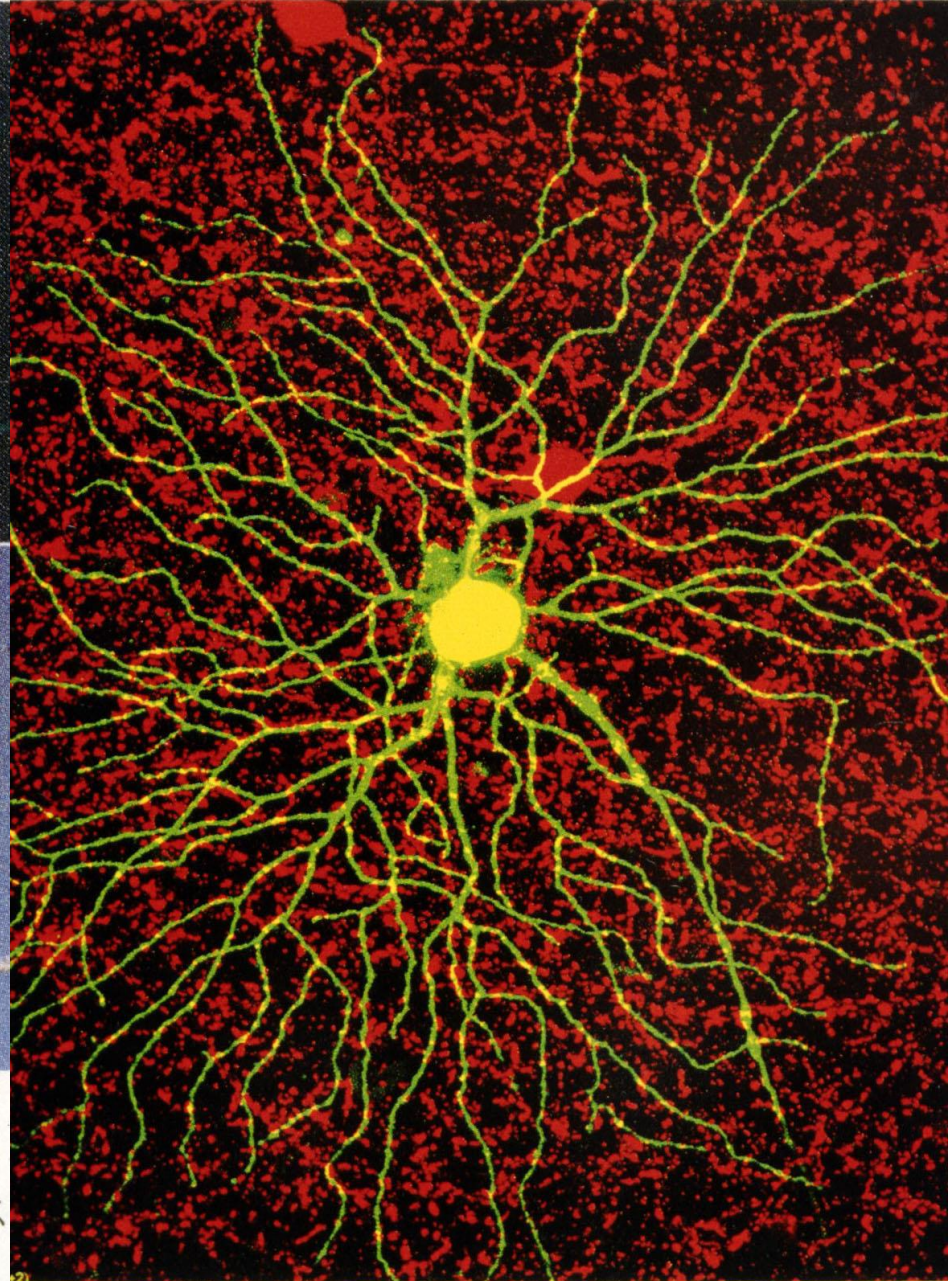
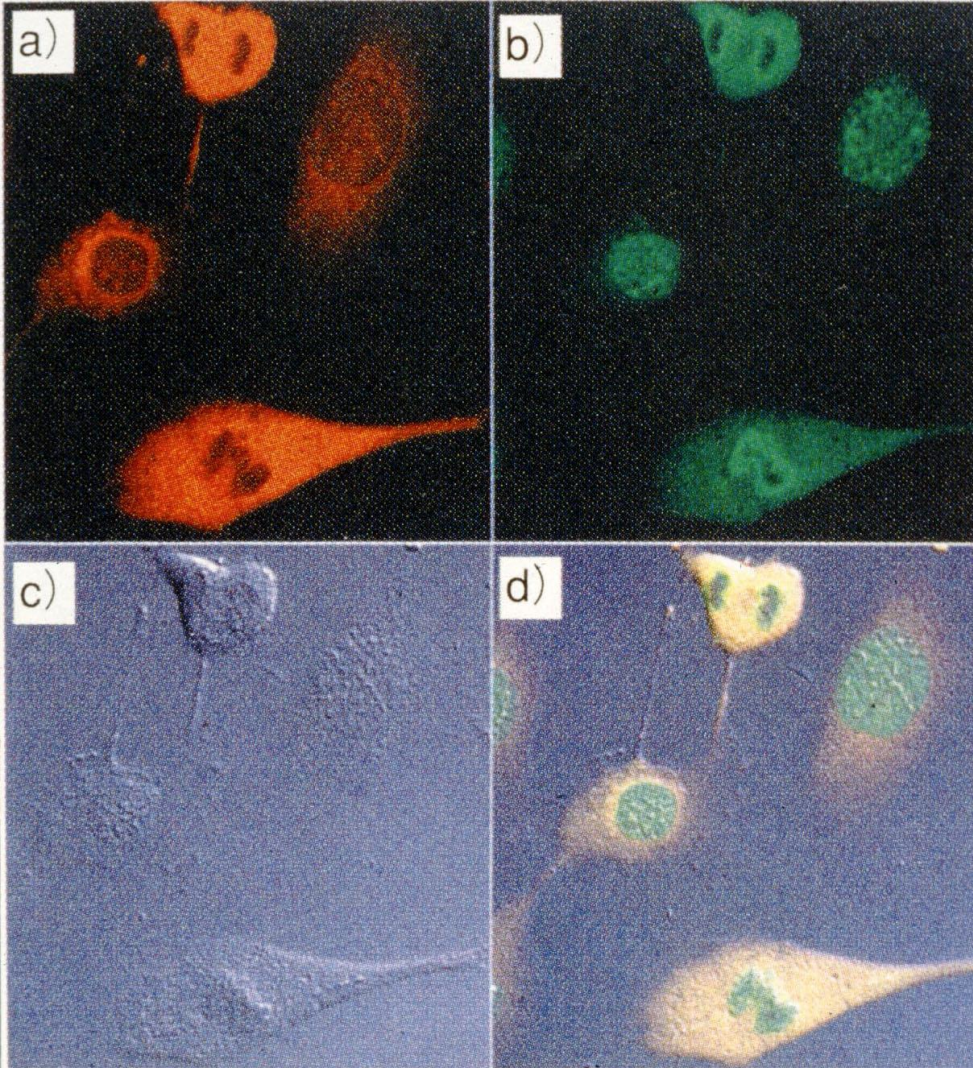
(d)



微分干涉及位相差的實例

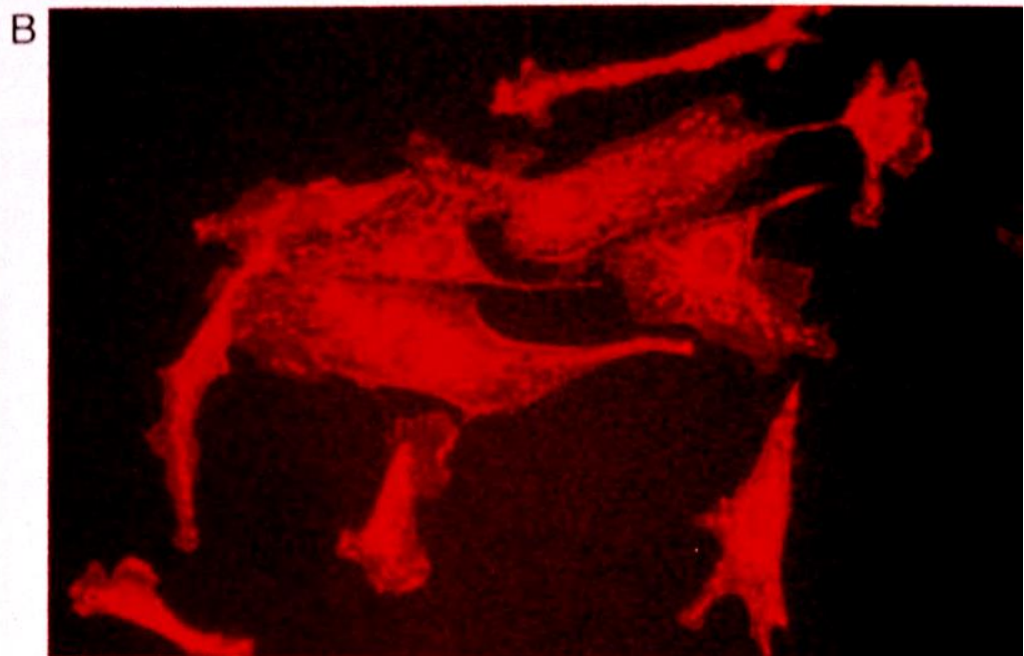
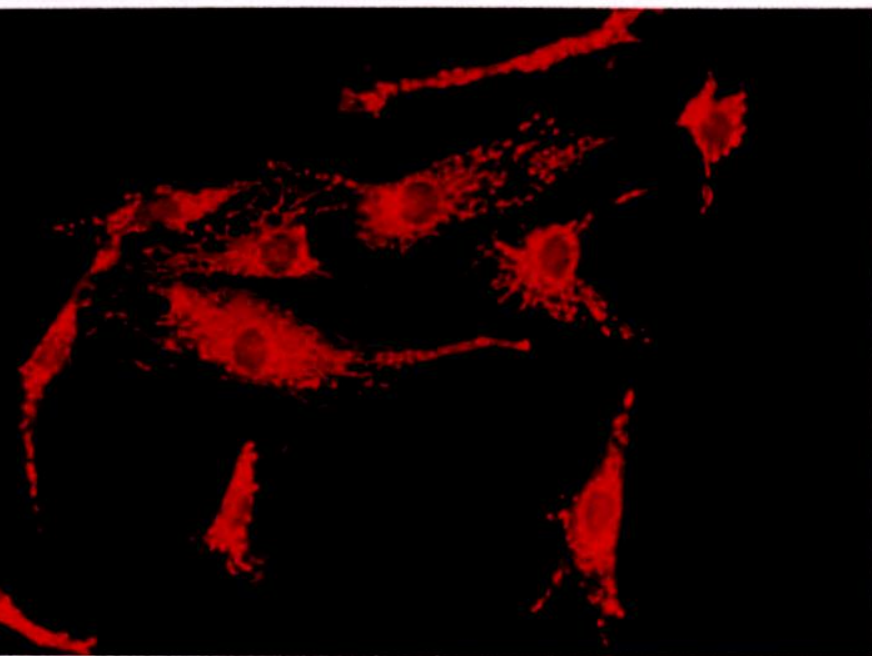
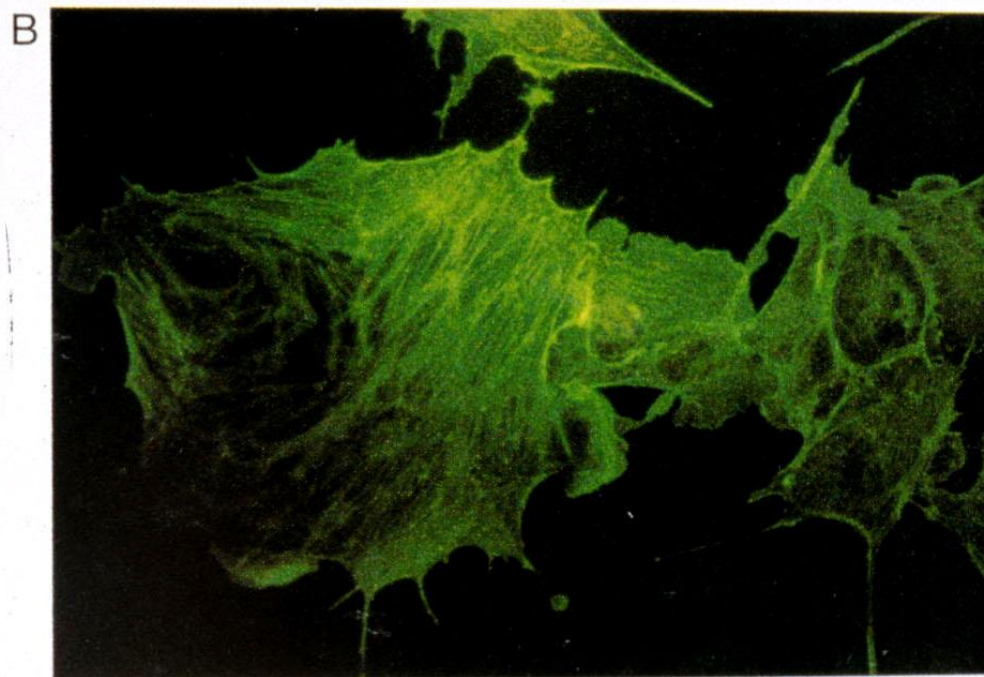
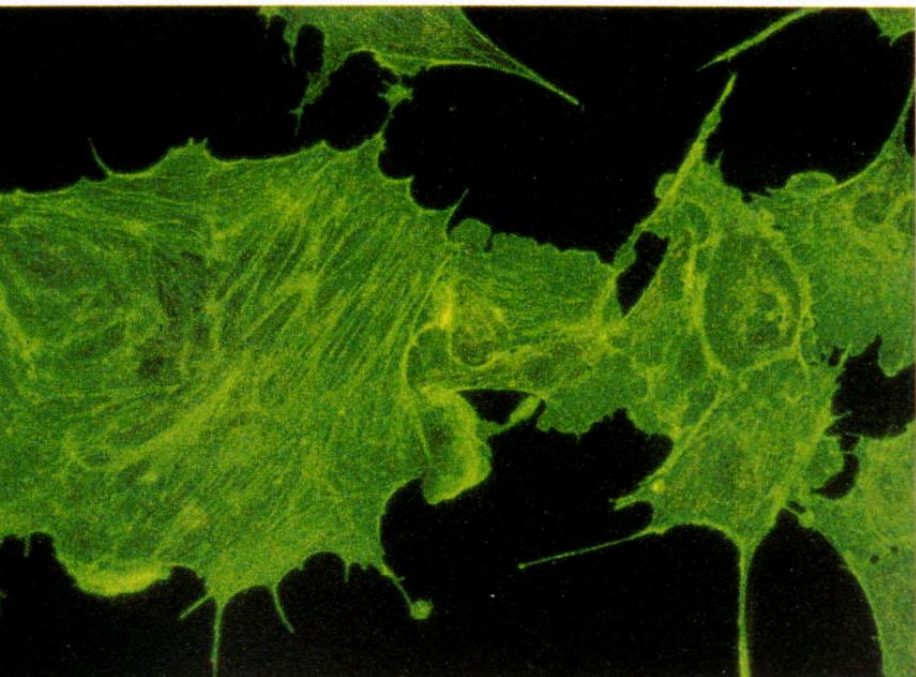
Live-Cell Imaging with Fluorescent Proteins and DIC

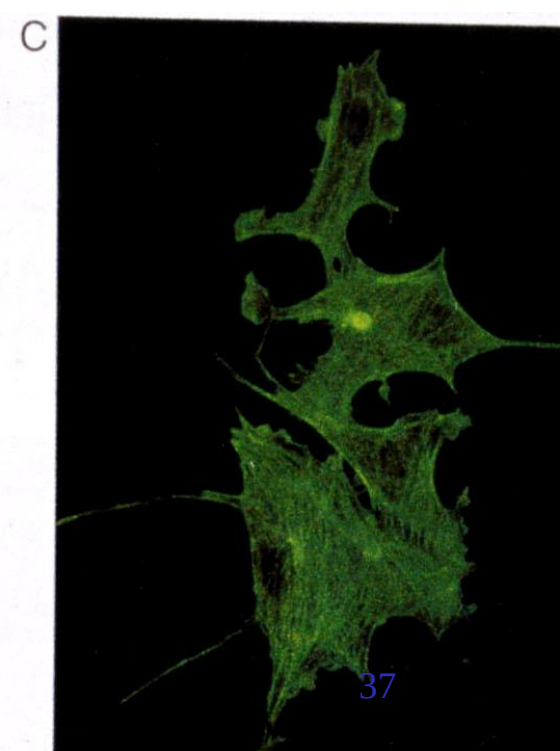
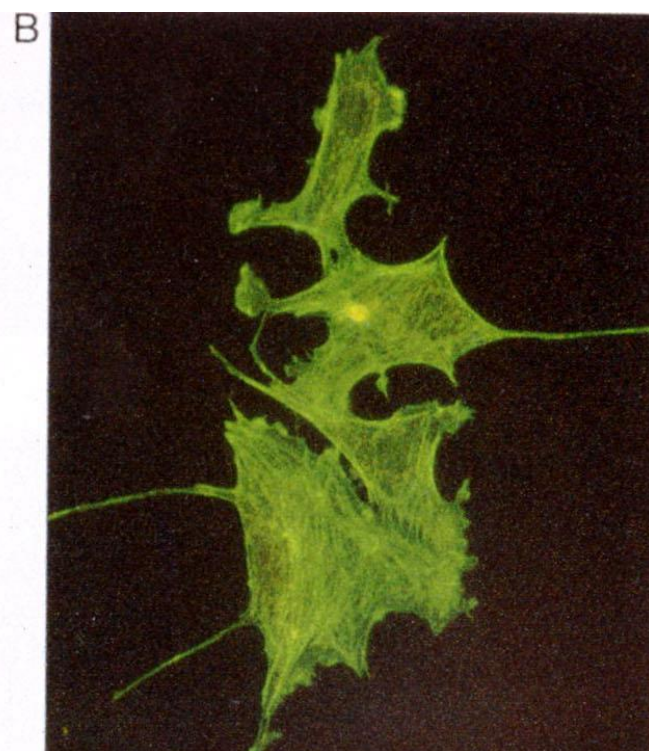
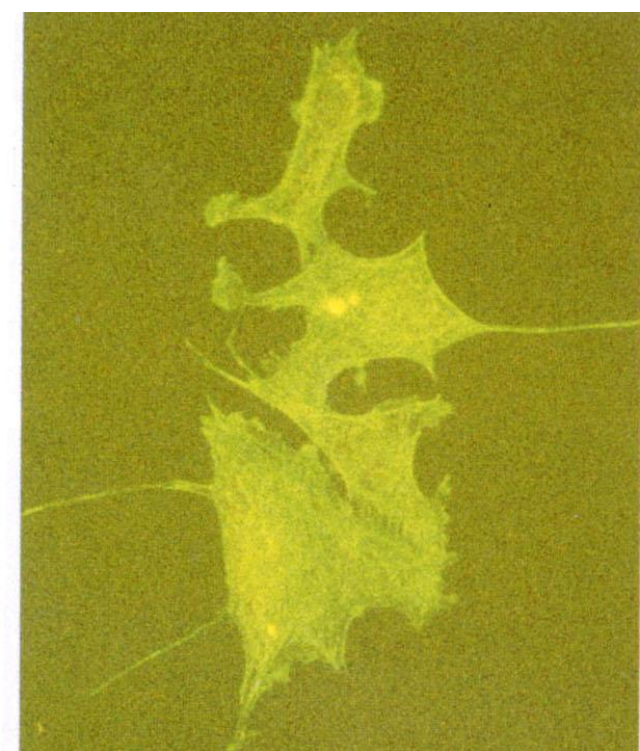
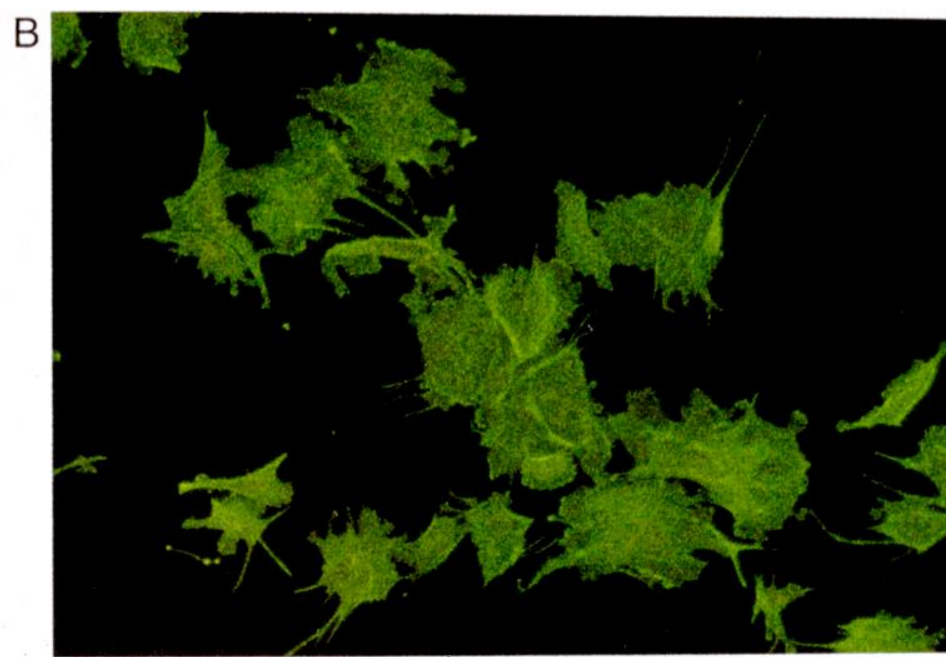
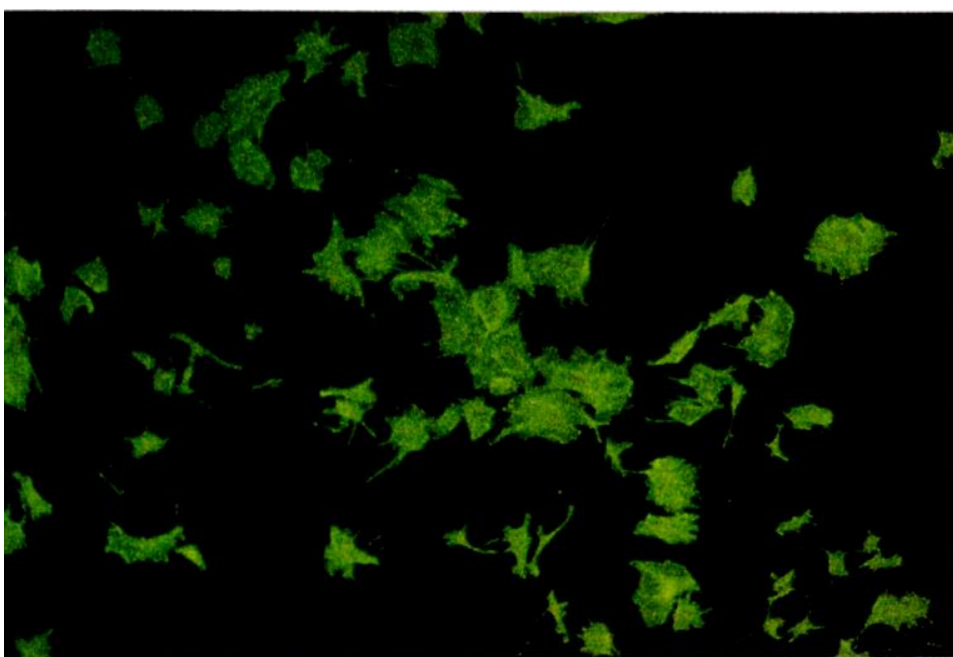


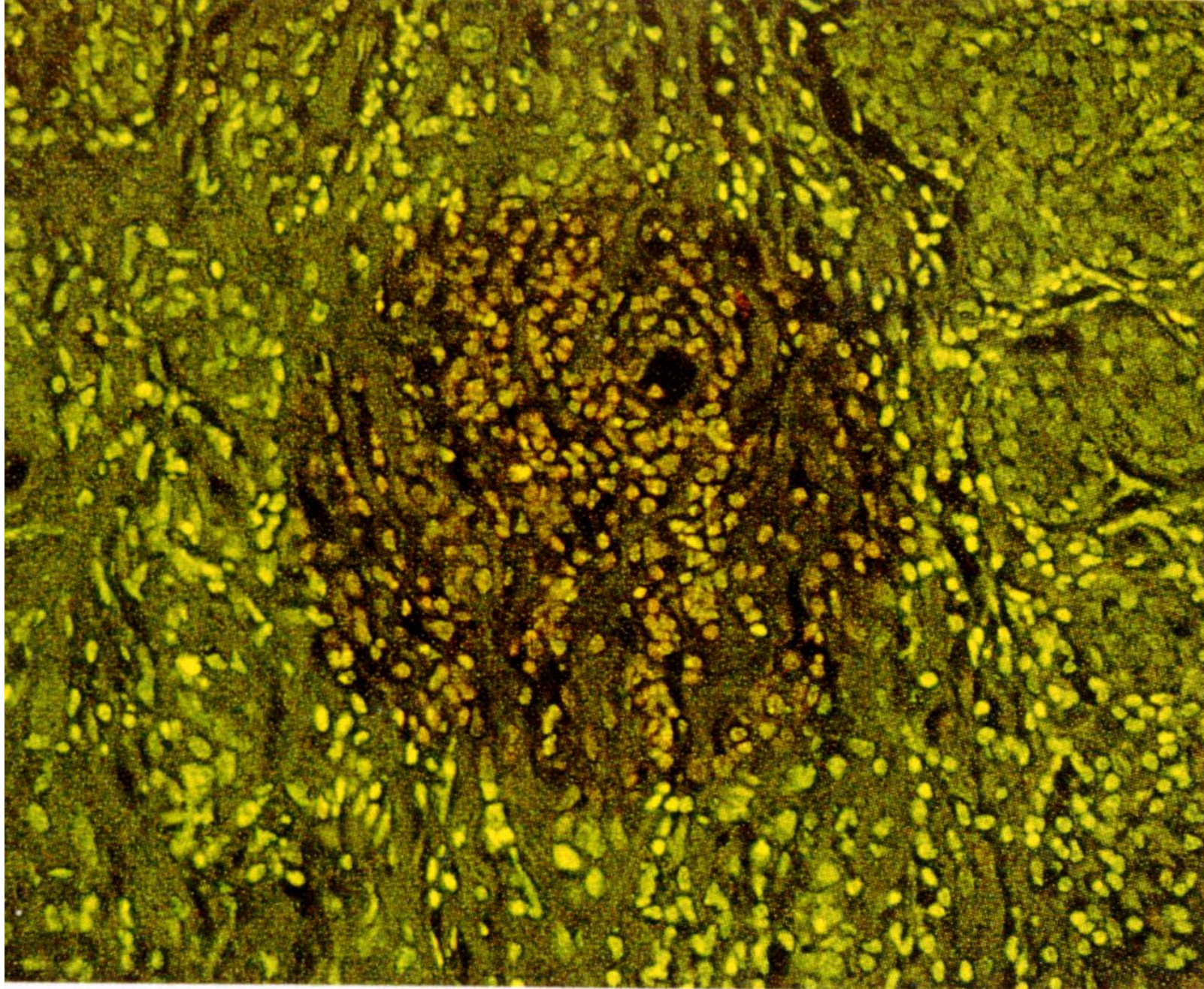


12 FITC, Rhodamine, 微分干渉および合成像
 a) F-アクチン/Rhodamine 染色像, b) クリ
 スン/FITC 染色像, c) 微分干渉像, d) 合成像.
 ラット由来の筋芽細胞 (L-6) の観察像.
 東京大学大学院総合文化研究科広域科学専攻生
 環境科学系跡見順子先生より提供

細胞とドーパミン作動性アマクリン細胞のレーザー顕微鏡エクステ
 像. Lucifer Yellowで網膜神経節細胞, Texas Redでドーパミン
 を染色.







褪色してしまった例

常見的抗退劑

- P-phenylenediamine
- DABCO
- N-propylgallate
- 2-mercaptoethylamine

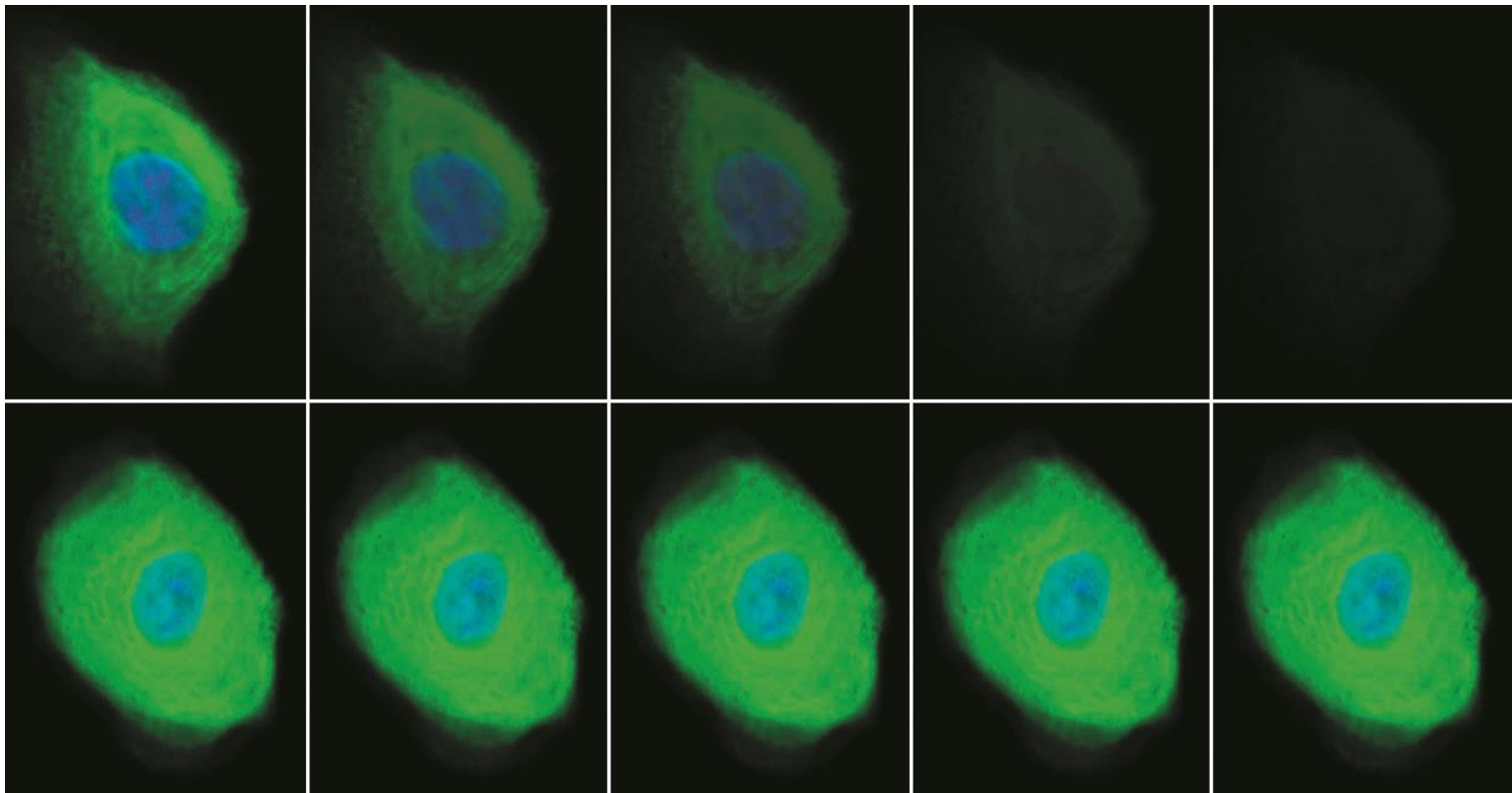
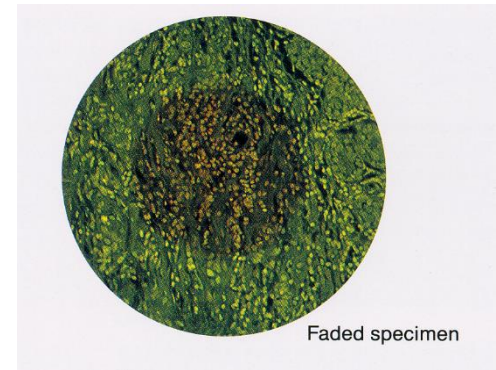


Figure 1. Immunofluorescence imaging of a human keratinocyte stained with FITC-labeled antibodies without (top) and with (bottom) PromoFluor Antifade Reagent at different excitations intervals (5, 10, 15, 20, 25 seconds). **PromoFluor Antifade Reagent**

Thank you for your attention