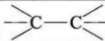
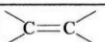
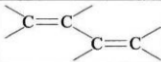

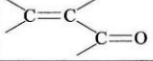




### UV SPECTROSCOPY

Chromophore	Wavelength / nm
	150
C-H	150
	190
	210-230
	195 (280 weak)
	210-250
	254
	270

Chromophore = portion of a molecule that absorbs light

Conjugated structure = 2+ adjacent C=C double bonds

More pi-electrons = more conjugated double bonds

↳ compound will absorb more light and have a longer wavelength and chromophore

### Analytical purposes

When a beam of radiation is passed through a sample,

- some radiation is **absorbed** by the sample
- some radiation is **reflected** or **scattered**
- some radiation **passes** straight through

For analytical purposes, we are interested in the **amount ABSORBED** by the sample and so we want to eliminate reflection and scattering.

This is done by:

- taking  $I_0$  as the intensity of the light passing through a cell when filled with a **blank solution** (everything except substance being measured)
- taking  $I$  as the intensity passing through the cell when filled with the **sample**

### Calculating transmittance

$$\text{Percentage transmittance} = \frac{I}{I_0} \times 100$$

### Equations

• Intensity path length:  $I$  decreases exponentially  $\rightarrow I = I_0 e^{-kcl}$   
 $T = \frac{I}{I_0} = e^{-kcl}$

• Fixed path length:  $I$  decreases in concentration with an increase in concentration  $\rightarrow I = I_0 e^{-k'c}$   
 $T = \frac{I}{I_0} = e^{-k'c}$

• Combining the equations:  $I = I_0 e^{-k'cl}$       $\ln\left(\frac{I}{I_0}\right) = -k'cl$   
 $T = \frac{I}{I_0} = e^{-k'cl}$       $\log\left(\frac{I}{I_0}\right) = \frac{-k'cl}{2.303}$       $\log\left(\frac{I}{I_0}\right) = -\frac{k'cl}{2.303}$

The quantity  $\log\left(\frac{I}{I_0}\right)$  is absorbance (A)  
 • The absorbance is directly proportional to the concentration of the absorbing substance and the path length  $\rightarrow A = \log\left(\frac{I_0}{I}\right) = \frac{k'cl}{2.303}$

### Beer-Lambert Law

Beer-Lambert law = dependence of absorption on concentration and path length

### A=acl

a = absorption coefficient

Represents the absorbance of a solution of unit concentration when measured in a cell of unit path length

### Quantitative analysis

Two methods are used to convert the measured absorbance into a concentration:

1. Calibration curve
2. Using the Beer-Lambert law with a given 'a' value

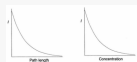
Calibration curve: 5 or more standards are prepared from a sample of pure material to be determined. Their absorbance is measured and their values are used to construct a calibration curve. The sample absorbance is measured and its concentration is read off the curve.

### A pure analyte sample must be available

Beer-Lambert law: Use of **A=acl** will be needed with additional steps to work out the dilution factor



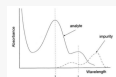
### Relationship between path length and concentration



The value of T (or I) depends on the cell's path length, concentration of absorbing substances and the nature of the substance.

- Increased path length = decreased intensity → light passes MORE through solution and interacts MORE with molecule

### Wavelength selection



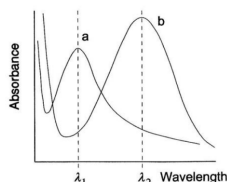
The wavelength for the assay should be chosen so that only the substance of interest absorbs, with no impurities

### Analysis of mixtures

$$A(\lambda) = \sum \epsilon_i c_i l$$

If we assume that the components of the mixture do not react with one another, then the absorbance at some wavelength = the sum of absorbance for each component

### Mixture of 2 substances



At wavelength  $\lambda_1$ ,

$$A(\lambda_1) = \epsilon_a(\lambda_1)c_a l + \epsilon_b(\lambda_1)c_b l$$

At wavelength  $\lambda_2$ ,

$$A(\lambda_2) = \epsilon_a(\lambda_2)c_a l + \epsilon_b(\lambda_2)c_b l$$

For a mixture of 2 substances with different chromophores, each will have different powers of light absorption at some wavelength(s) in the spectrum.

If measurements are made on the mixture at 1 and 2, then a pair of simultaneous equations can be set up to find the unknown concentrations.

### FLUORESCENCE SPECTROSCOPY

What is fluorescence?

When molecules absorb UV-visible radiation, they absorbed energy is converted into kinetic energy due to collisions.

A few excited molecules get rid of excess energy by emitting the absorbed energy as light = **FLUORESCENCE**

Why use fluorescence?

Fluorescence is more selective than absorption of energy

↳ The chances of finding 2 substances which absorb **and emit** at the same wavelength is **LESS** than finding 2 substances that absorb at the same wavelength

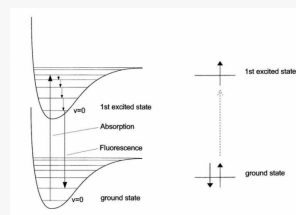
Not all substances fluoresce

↳ Advantage: it makes fluorimetry more selective for compounds that fluoresce

Disadvantage: not all compounds fluoresce

It's more selective than absorption spectrophotometry

### Process of fluorescence



- The emitted radiation is of lower energy than the absorbed radiation
- Molecules in the ground state absorb light and are excited to a vibrational level in the 1st excited state
- Excess vibration is then lost by collisions with other molecules until they are in the V=0 level of the 1st excited state
- The molecules get rid of the remaining energy by emitting it as radiation and dropping to a vibrational level in the ground state

### Types of fluorescence

- Chemiluminescence** When the product molecules are left in an excited state, light is emitted when the molecules return to the ground state
- Bio-luminescence** Biochemical reactions that produce light

### Relationship between structure and fluorescence

- e.g. Fluorescein → fluorescent and rigid structure (prevents loss of energy)
- e.g. Phenolphthalein → non-fluorescent and non-rigid structure (can twist which converts its absorbed energy into rotational and vibrational energy)

### Fluorescence enhancers and inhibitors

Fluorescence enhancers	Fluorescence inhibitors
OH, OCH <sub>3</sub> , NH <sub>2</sub> , NHR, NR <sub>2</sub>	COOH, NO <sub>2</sub> , NO, F, Cl, Br, I
↳ Increase the no. of delocalised electrons	

### Factors affecting fluorescence intensity

$$I = I_0 10^{-\epsilon cl}$$

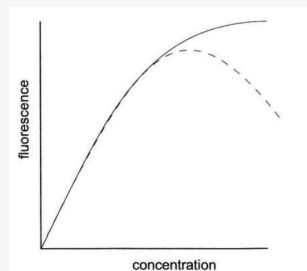
$$F \propto (I_0 - I)$$

More energy absorbed = more energy emitted  
Intensity of fluorescence is **PROPORTIONAL** to amount of light absorbed

Relationship between  $I_0$  and  $I$  given by the Beer-Lambert law

- Intensity of emitted radiation is directly proportional to conc
- Fluorescence is directly proportional to intensity of excitation state
- ↳ brighter lamp = greater fluorescence
- Fluorescence depends on molar absorption coefficient ( $\epsilon$ )
- ↳ stronger absorbance = greater fluorescence

### Fluorescence intensity vs. concentration

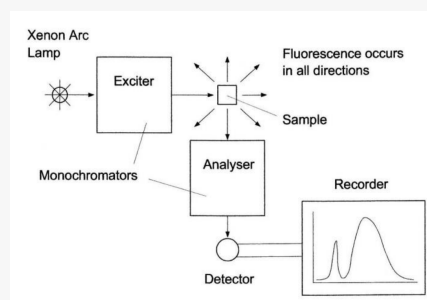


At high concentrations, the fluorescence emission becomes concentrated near the **sides of the sample cell** to which the exciting radiation enters.

As the exciting radiation passes through the sample, its intensity falls and the fluorescence produced also decreases.

**Under these conditions, the equation for fluorescence intensity no longer holds (as  $\epsilon cl$  is no longer small)**

### Experimental measurement of fluorescence



#### 1. Light source

- As intense as possible at wavelength of max absorption
- Continuous source
- Wavelength region of 200-500nm

#### 2. Excitor and detector monochromator

- Used to select excitation/emission wavelengths
- Low diminishing and high light gathering powers
- Filters can be used but are less versatile

#### 3. Sample cell

- Usually silica
- 1x1cm
- 4 clear, optically worked surfaces

#### 4. Detector

- Must be sensitive as possible (varies)
- Photomultiplier tube



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By **eyeeyuu**  
[cheatography.com/eyeeyuu/](https://cheatography.com/eyeeyuu/)

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### Factors affecting fluorescence measurements

- 1. Lower limit of detection**      Lowest concentration of substance that can be determined
- 2. Fluorescent impurities**      Usually from the reagents or from the cell  
Ensure the cell is clean and is always placed in the same way (as the 4 sides have different background intensities)
- 3. Photodecomposition**      The sample can undergo decomposition due to the high intensity of the exciting radiation samples, and standards should NOT be exposed to the radiation longer than necessary
- 4. Quenching**      Substance fluorescence is affected more by its environment than the absorption of the sample  
**Quenching = when fluorescence is decreased by the presence of another sample**



By **eyeeyuu**  
[cheatography.com/eyeeyuu/](https://cheatography.com/eyeeyuu/)

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