Six different phenomena form the basics of Optical spectroscopic methods:

1. **Absorption** – causes the promotion of a particle from its normal room temperature state to one or more higher energy excited states.
   - The process of absorption of quantized; i.e., for absorption to occur, the excitation energy must exactly match the energy difference between the ground state and one of the excited states.
   - For molecules, obtaining a radiant energy that match a transition is easy as there are several states (electronic, vibration, rotation). For atoms, this is a challenge because they contain only electronic states.
   - For molecules, there are several relaxation means by which the excited species can get rid of the excess energy (i) non-radiative, (ii) fluorescence, and (iii) phosphorescence.
   - For molecular absorption, we need materials that allow only a part of the electromagnetic radiation to be exposed to the analyte depending on the information needed.

2. **Fluorescence** -

3. **Phosphorescence** -

4. **Scattering** - only interested in the small fraction of the scattered photons (approximately 1 in 10 million) that are scattered by an excitation, with the scattered photons having a frequency different from that of the incident photons.

5. **Emission** – release of radiant light by a substance resulting from heat.

6. **Luminescence** - emission of light by a substance not resulting from heat.
   - E.g., Chemiluminescence: a result of a chemical reaction,
   - Photoluminescence: a result of absorption of photons.
Although the required instrument components for measuring each phenomenon somewhat differ in configuration, most of their basic components are remarkably similar.

Typical spectroscopic instrument contain five components:

1. Stable source for radiant energy
2. Transparent container for holding the sample
3. Device that isolates restricted region of the spectrum for measurement
4. A radiant detector that converts radiant energy to useable electrical signal
5. Signal processor and readout
1. Absorption

2. Fluorescence and Phosphorescence

3. Emission or Luminescence

- Emission spectroscopy and Chemiluminescence spectroscopy differ from the others in that no external radiation source is required.
- The sample itself is the emitter
- Emission: sample container is a plasma, or flame that contains the analyte, and also causes it to emit radiant light
- Chemiluminescence: radiation source is a solution of the analyte plus reagents held in a transparent sample holder
**Sources of Radiation**

A suitable source is the one that is (1) stable and (2) generates sufficient radiant power for easy detection.

**Continuum sources:**
1. Emit radiation that changes only slowly as function of wavelength.
2. Widely used in absorption and fluorescence spec.

**Line sources:**
1. Emit discrete of limited number of lines or bands of radiation, each of which spans a limited range of wavelengths.
2. Widely used in atomic absorption spectroscopy, atomic and molecular fluorescence spectroscopy, and Raman spectroscopy.

<table>
<thead>
<tr>
<th>Spectral region</th>
<th>VAC</th>
<th>UV</th>
<th>Visible</th>
<th>Near IR</th>
<th>IR</th>
<th>Far IR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sources</td>
<td>Ar lamp</td>
<td>Xe lamp</td>
<td>H$_2$ or D$_2$ lamp</td>
<td>Tungsten lamp</td>
<td>Nernst glower (ZrO$_2$ + Y$_2$O$_3$)</td>
<td>Nichrome wire (Ni + Cr)</td>
</tr>
<tr>
<td>Continuum</td>
<td>Hollow cathode lamps</td>
<td>Lasers</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Line</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Sources of Radiation

Most common continuum sources:
1. UV region – deuterium lamps

\[ D_2 + E_e \rightarrow D_2^* \rightarrow D' + D'' + \nu \text{ (ultraviolet photon)} \]

- \( E_e \) = electrical energy
- \( D' \) = excited deuterium molecule
- \( E_e = E_{D2*} \approx E_{D'} + E_{D''} + \nu \)

- \( E_{D2*} \) = fixed quantized energy of \( D_2^* \)
- \( E_{D'} \) and \( E_{D''} \) = kinetic energies of the two deuterium atoms, the sum of varies from 0 to \( E_{D2*} \)
  i.e., when \( E_{D'} + E_{D''} = E_{D2*} \), then \( \nu = 0 \)
  and when \( E_{D'} + E_{D''} = 0 \), \( \nu = E_{D2*} \)

- Since these conditions are determined by chance, the consequence is a true continuum
Sources of Radiation

Most common continuum sources:

2. Visible Region – tungsten filament lamp or quartz-halogen lamps
   - Temperature dependent
   - Bulk of the radiation is emitted in the IR region during heating
   - Absorption of radiation by glass case that houses the filament absorbs the IR radiations, and imposes the lower limit
   - Quartz case allows higher temperatures (3500 K) to be used, which yields higher radiant intensities, and expands the range of the lamp to UV regions
   - Tungsten-halogen lamp: contain iodine, which sublimes to react with tungsten vapor to form a volatile compound WI₂. When molecules of this compound strikes the filament, decomposition occurs, which re-deposits tungsten and thus extending the lifetime of the lamp

3. Infrared region – inert solids heated to 1500 to 2000 K
Sources of Radiation

Most common line sources:

1. Hollo Cathode Lamps

- High field between anode and cathode ionizes inert gas buffer inside the glass
- Ionized inert gases are accelerated to the cathode (negatively charged), which sputters atoms from the cathode
- Sputtered atoms are excited through collision with other atoms in the glass enclosure
- Photons are emitted as the excited atoms decay to the ground state. These photons are of discrete energies, and thus producing line spectrum
Sample containers must be transparent to radiation in the spectra region of interest. Common materials and their useful regions are shown below:

<table>
<thead>
<tr>
<th>Wavelength, nm</th>
<th>100</th>
<th>200</th>
<th>400</th>
<th>700</th>
<th>1000</th>
<th>2000</th>
<th>4000</th>
<th>7000</th>
<th>10,000</th>
<th>20,000</th>
<th>40,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spectral region</td>
<td>VAC</td>
<td>UV</td>
<td>Visible</td>
<td>Near IR</td>
<td>IR</td>
<td>Far IR</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Materials for cells, windows, lenses, and prisms</td>
<td>Lithium fluoride</td>
<td>Fused silica or quartz</td>
<td>Correx glass</td>
<td>Silicate glass</td>
<td>NaCl</td>
<td>KBr</td>
<td>TlBr or TlI</td>
<td>ZnSe</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Two types of Wavelength Selectors:

1. Filters (allows a selected portion of radiation to pass)
   a) Absorption Filters (only for visible region)
   b) Interference Filters (for UV, Visible and IR regions)

2. Monochromators (for scanning)
   a) Prism Monochromators (for UV, Visible and IR regions)
      i. Cornu type (60°)
      ii. Littrow type (30°)
   b) Grating Monochromators (for UV, Visible and IR regions)
      i. Echellette Grating
      ii. Concave Gratings
      iii. Holographic Gratings
Wavelength Selectors – Absorption Filters

Less expensive compared with interference filters
Widely used for band selection in visible region
Bandwidth range: 30 – 250 nm available commercially

Types:
Colored glass → thermally stable
Dye suspension in gelatin and sandwiched between glass
Cutoff filter → 100% transmittance over some portion, and then rapidly falls to zero over the remainder
Combine cutoff filter with a second filter to achieve a narrow spectra band

[Graph showing transmittance of absorption filters, green filter, orange cutoff filter, and combination of two filters over wavelength range]
Wavelength Selectors – Interference Filters

A diagram illustrating the output of a typical filter-based wavelength selector. White radiation is input at the top, passing through a glass plate, metal film, and dielectric layer. This results in a narrow band of radiation as output. A graph showing the percent transmittance with respect to wavelength is also included, with a peak at nominal wavelength and effective bandwidth indicated."
Wavelength Selectors – Interference Filters

Decreased effective bandwidth comes at a cost of reduced percent transmittance.

Effective bandwidth for three types of filters:
- Interference filter: Effective bandwidth ~10 nm
- Absorption filter: Effective bandwidth ~50 nm
- Peak height

Half peak height
Wavelength Selectors – Monochromators

1. Prism Monochromator

Components of Monochromators
1. Entrance slit that provides a triangular optical image
2. Collimating lens that produces a parallel beam of radiation
3. A prism of grating that disperses the radiation into its component wavelengths
4. Focusing element that reform the image of the entrance slit, and focuses it on a focal plane
5. An exit slit in the focal plane that isolates the desired spectral band

Scan wavelength by rotating prism or grating

2. Grating Monochromator
Dispersion of light of two wavelengths by a prism of refractive index $\eta$, apex $\alpha$, and baselength $b$. Collimated rays of wavelength $\lambda_1$ (red) and $\lambda_2$ (blue) are refracted upon entering the prism material and upon exiting it according to Snell’s law. Normal prism materials show higher refractive indices at shorter wavelengths. Hence blue light of wavelength $\lambda_2$ is more highly refracted than red light ($\lambda_1$).
Wavelength Selectors – Prism Monochromators

Angular Dispersion (DA):
From Snell's law, the angle of refraction of light in a prism depends on the refractive index of the prism material. But, since that refractive index varies with wavelength, it follows that the angle that the light is refracted by will also vary with wavelength, causing an angular separation of the colors.

\[ DA = \frac{d\theta}{d\lambda} = \frac{dn}{d\lambda} \times \frac{d\theta}{dn} \]

- Determined by prism material
- Determined by prism design (e.g., \( \alpha \))

\[ n = \text{refractive index of the prism material} \]
\[ \theta = \text{angle of refraction} \]
\[ \lambda = \text{wavelength of refracted ray} \]
Wavelength Selectors – Grating Monochromators

Grating Monochromator

Wavelength selection device based on the constructive interference of light rays that have traveled different distances to reach the same point. Work via two mechanisms:

1. Transmission,
2. Reflection (most common)

Beam 2 travels a greater distance than beam 1, the extra distance = BC + DB

For constructive interference, this extra distance must be a multiple of the wavelength (n\lambda) of the reflected beam

n\lambda = BC + DB

Since angle at CAB = i, and angle at DAB = r,

CB = dsin i, and BD = dsin r

n\lambda = dsin i + dsin r = d(sin i + sin r)
Wavelength Selectors – Grating Monochromators

Performance Characteristics
1. Spectra Purity
2. Dispersion
3. Resolving Power
4. Light-gathering Power

1. Spectra Purity:
The exiting beam is always observed to be contaminated with small amounts of wavelengths far from that of the instrumental setting. This is mainly due to the following reasons:

a) Scattered radiation – caused by the presence of dust particulates inside the monochromator as well as on various optical surfaces. This drawback can be overcome by sealing the monochromator entrance and exit slits by suitable windows.

b) Stray radiation – radiation that exits the monochromator without passing through the dispersion element. This problem, including those related to spurious radiation, can be largely eliminated by painting the internal walls of the monochromator by a black paint.

c) Imperfections of monochromator components – e.g., broken or uneven blazes, uneven lens or mirror surfaces, etc., would lead to important problems regarding the quality of obtained wavelengths.

2. Dispersion:
Angular dispersion ($dr/d\lambda$): change in angle of reflection ($r$) with respect to changes in wavelength ($\lambda$)

Differentiate $n\lambda = d(sin i + sin r)$ at constant incidence angle $\lambda$

$$\frac{dr}{d\lambda} = \frac{n}{d\cos r}$$

Linear dispersion ($D$) is the most appropriate, which represents variations in wavelength as a function of some distance ($y$) on the focal plane.

$$D = \frac{dy}{d\lambda}$$

If the focal length of the collimating mirror is $F$, then

$$dy = Fdr; \text{ hence } D = \frac{Fdr}{d\lambda}$$

Reciprocal linear dispersion ($D^{-1}$) =

$$\frac{nF}{d}$$

At small reflection angles ($<20^\circ$), $\cos r \approx 1$

Hence, $D^{-1} = \frac{d}{nF}$ and $D = \frac{nF}{d}$
Implications for small angle of reflection usually used for grating operation

1. Dispersion of a grating monochromator is linear and independent on wavelength
2. Dispersion of a grating monochromator increases as the line spacing, $d$, is decreased
3. Dispersion of gratings is linear meaning that all wavelengths are dispersed to the same extent, something that simplifies instrumental designs

$$D = \frac{nF}{d}$$
3. Resolving Power ($R$):

Ability of a grating monochromators to separate adjacent wavelengths, with very small difference

$$R = \frac{\lambda}{\Delta \lambda}$$

$\Delta \lambda$ = difference between the two adjacent wavelengths ($\lambda_2 - \lambda_1$)

$\lambda$ = average of the two wavelengths ($\frac{\lambda_1 + \lambda_2}{2}$)

The resolving power can also be defined as:

$$R = \frac{nN}{n\lambda}$$

$n$ = diffraction order

$N$ = number of illuminated blazes

Hence, better resolving powers can be obtained with:

a. Longer gratings
b. Higher blaze density
c. Higher order of diffraction (obtained using echelle gratings)
4. **Light-gathering Power:**

   The ability of a grating monochromator to collect incident radiation from the entrance slit is very important as only some of this radiation will reach the detector.

   \[
   \text{f/number (or f-number or speed) is a measure of the ability of the monochromator to collect incident radiation}
   \]

   \[
   \text{f/number} = \frac{F}{d}
   \]

   \(F = \text{focal length of the collimating mirror or lens}\)

   \(d = \text{diameter of mirror}\)

   The light-gathering power of a grating monochromator increases as the inverse square of the \(f\)-number.

   Thus: \(f/2\) mirror gathers 4 times more light than an \(f/4\)

   The \(f\)-number for most monochromators ranges from 1 to 10.
## General Instrumentation for Spectrochemical Method

<table>
<thead>
<tr>
<th>Wavelength, nm</th>
<th>100</th>
<th>200</th>
<th>400</th>
<th>700</th>
<th>1000</th>
<th>2000</th>
<th>4000</th>
<th>7000</th>
<th>10,000</th>
<th>20,000</th>
<th>40,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>(b) Detectors</td>
<td>Photographic plate</td>
<td>Photomultiplier tube</td>
<td>Phototube</td>
<td>Photocell</td>
<td>Silicon diode</td>
<td>Charge-transfer detector</td>
<td>Photoconductor</td>
<td>Thermocouple (voltage) or bolometer (resistance)</td>
<td>Gelay pneumatic cell</td>
<td>Pyroelectric cell (capacitance)</td>
<td></td>
</tr>
</tbody>
</table>