

From textbook: (Trease and Evans Pharmacognosy, 16th Ed.)
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Alkaloids

- Alkaloids are extremely difficult to define for they do not represent a homogeneous group of compounds either from the chemical, biochemical, or physiologic viewpoint. Consequently, Alkaloids are all organic nitrogenous compounds.
- Plants have been a rich source of alkaloids, but some are found in animals, fungi, and bacteria. Most possess basic properties, due to the presence of an amino nitrogen, and many possess marked physiologic activity.
- In spite of the difficulties attending a precise definition, the term is an extremely useful one, commonly applied to basic nitrogenous compounds of plant origin which are physiologically active.

Distribution

- 1- Angiosperms: Leguminosae, Papaveraceae, Ranunculaceae, Rubiaceae, Solanaceae, and Berberidaceae. (but not the Labiatae and Rosaceae).
- 2- The gymnosperms only (Taxaceae).
- 3- The monocotyledons: Amaryllidaceae and Liliaceae are two of the most promising families in which to search for alkaloid-yielding plants.

- Alkaloids may occur in various parts of the plant:
- In seeds (nux vomica, areca),
- In fruits (black pepper, conium),
- In leaves (belladonna leaf, hyoscyamus),
- In underground stems (sanguinaria, corydalis).
- In roots (aconite, belladonna root),
- In rhizomes and roots (ipecac, hydrastis),
- In barks (cinchona, pomegranate).
- They are also found in the fungi (ergot, *Amanita citrina*).

The names of the alkaloids are obtained in various ways:

- (1) From the generic name of the plant yielding them (hydrastine, atropine).
- (2) From the specific name of the plant yielding them (cocaine, belladonnine).
- (3) From the common name of the drug yielding them (ergotamine).
- (4) From their physiologic activity (emetine, morphine).
- (5) from the discoverer (pelletierine).
- By agreement, chemical rules designate that the names of all alkaloids should end in "ine."

Toxicity to Humans and Other Vertebrates

- Animal intoxication by alkaloids is mostly caused by accidental ingestion of food contaminated with alkaloid-containing plants.
- Clearly, the amount of ingested alkaloid and the sensitivity of the target animal are key factors leading to intoxication.
- Some alkaloids can be extremely harmful to mammals, which is the case of the steroidal alkaloid cyclopamine in lambs, identified as the compound in *Veratrum californicum* (Liliaceae) responsible for teratogen effects.

The possible function of alkaloids in plants as:

- (1) Poisonous agents protecting the plant against insects and herbivores.
- (2) End products of detoxification reactions representing a metabolic locking-up of compounds otherwise harmful to the plant.
- (3) Regulatory growth factors.
- (4) Reserve substances capable of supplying nitrogen or other necessary elements to the plant's economy.

PROPERTIES:

- Most alkaloids are well-defined crystalline substances which unite with acids to form salts.
- In the plant they may exist in the free state, as salts or as N-oxides.
- [N-oxides, also referred to as amine oxides, are organic compounds that contain the functional group N+-O-. Amine oxides are weak bases and highly polar molecules. Small amine oxides are found to be hydrophilic in nature and hence possess excellent water solubility].
- In addition to the elements carbon, hydrogen and nitrogen, most alkaloids contain oxygen, and an additional few, coniine, nicotine, and sparteine, which lack oxygen in their molecules, are liquids.
- Although colored alkaloids are relatively rare, they are not unknown; berberine for example is yellow and the salts of sangularine are copperted.

- Alkaloids usually contain one nitrogen atom, although some like ergotamine may contain up to five. The nitrogen may exist as a primary amine (RNH2), as a secondary amine (R2NH), or as a tertiary amine (R3N).
- Since the nitrogen atom bears an unshared pair of electrons, such compounds are basic and resemble ammonia in chemical properties. The degree of basicity varies greatly depending upon the structure of the molecule and the presence and location of other functional groups.
- Like ammonia, the alkaloids are converted into their salts by aqueous mineral acids, and when the salt of an alkaloid is treated with hydroxide ion, nitrogen gives up a hydrogen ion and the free amine is liberated.

- Quaternary ammonium compounds [R4N+ X-] such as tubocurarine chloride or muscarine chloride have 4 four organic groups covalently bonded to nitrogen, and the positive charge of this ion is balanced by some negative ion.
- The quaternary ammonium ion, having no proton to give up, is not affected by hydroxide ion; consequently, quaternary ammonium compounds have chemical properties quite different from those of the amines.

- For the most part the alkaloids are insoluble or sparingly soluable in water but the salts formed upon reacting with acids are usually freely soluble.
- The free alkaloids are usually soluble in ether or chloroform or other relatively nonpolar, immiscible solvents in which, however, the alkaloidal salts are insoluble.
- This permits a ready means for the isolation and purification of the alkaloids as well as for their quantitative estimation.

Tests for alkaloids

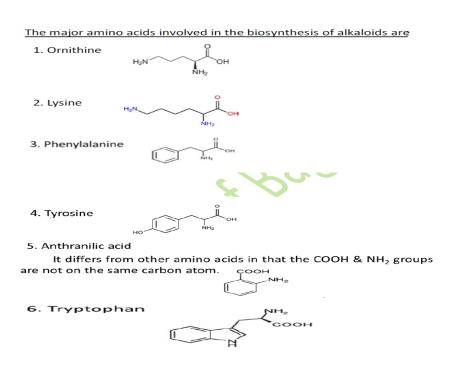
Most alkaloids are precipitated from neutral or slightly acid solution by :

- 1- Mayer's reagent (potassium mercuric iodide solution).
- 2- Wagner's reagent (solution of iodine in potassium iodide).
- 3- Solution of tannic acid.
- 4- Hager's reagent (a saturated solution of picric acid).
- 5- **Dragendorff's reagent** (solution of potassium bismuth iodide).

These precipitates may be amorphous or crystalline and are of various colors: @@@m (Mayer¹s), yellow (Hager¹s), reddishbrown (Wagner's and Dragandorff's).

- Care must be taken in the application of these alkaloidal tests:
- 1- The reagents also give precipitates with proteins. During the extraction of alkaloids from the plant and sub-sequent evaporation. Some proteins will not be extracted and others will be made insoluble (denatured) by the evaporation process and may be filtered out.
- 2- Caffeine, and other purine derivatives, does not precipitate like most alkaloids. It is usually detected by mixing with a very small amount of potassium chlorate and a drop of hydrochloric acid, evaporating to dryness and exposing the residue to ammonia vapor.

A purple color is produced ,this is known as the murexide test.



Classification:

Various schemes for the classification of alkaloids have been suggested.

1. Due to Hegnauer:

True (Typical) alkaloids that are derived from amino acids and have nitrogen in a heterocyclic ring. e.g Atropine

Protoalkaloids that are derived from amino acids and do not have nitrogen in a heterocyclic ring. e.g Ephedrine H_{CH_3}

Pseudo alkaloids that are not derived from amino acids but have nitrogen in a heterocyclic ring.

e.g Caffeine

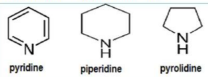
False alkaloids are non alkaloids give false positive reaction with alkaloidal reagents. e.g. homatropine.

2 .Alkaloids that classified according to the nature of the basic chemical structures from which they derive:

- Arecoline, pelletierine, lobeline, coniine, and nicotine are derivatives of pyridine and piperidine.
- atropine, hyoscyamine, and hyoscine are derived from tropane, a condensation product of pyrrolidine and piperidine.



Tropane



3. The Cinchona alkaloids; quinine, quinidine, cinchonine, and cinchonidine contain quinoline as the principal nucleus; hydrastine, d-tubocurarine, emetine, and certain of the Opium alkaloids are characterized by the isoquinoline nucleus.



Quinoline



4. ergonovine, reserpine, and strychnine which derive from the indole ring.



5. pilocarpine having the imidazole ring.

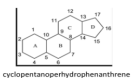


6. caffeine and theobromine which are purine bases.



7. morphine and codeine possessing the phenanthrene ring, and aconitine and protoveratrine which contains a steroidal structure.





Extraction of alkaloids

- The extraction by fractional extraction (From less Polar to more Polar).
- 1-Defatting by non-polar solvent (Petroleum Ether, benzene, alkane,....) To get rid of Chlorophyll, Wax, Volatile oil, Fixed oil.
- 2-Filtration, for marc use methanol or ethanol 95% Evaporate by rotary evaporator (to Concentrate).
- 3-Add Tartaric acid 2% and Ethyl acetate will separate into two layers:
- Organic layer (For week or neutral alkaloids)
- -Aqueous layer (acidic layer, Tartaric acid) which have alkaloidal salt.
- 4- To break the salt, add NH3 or Sodium bicarbonate. then add ethyl acetate again so will it separate into two layers again:
- - Aqueous layer (Quaternary alkaloids 4º)
- - Organic layer (For basic alkaloid 10,2°,3°).

Purification of the extract

- Direct crystallization: The simplest procedure, but it seldom success when a crude mixture is involved.
- The extract is evaporated to dryness & the residue is dissolved in a crystallizing solvent, which may be either as in a single organic solvent or a mixture of two solvents.
- In a general order of increasing solubility of most alkaloids is as follows: Hexane, benzene, ether, ethylacetate, methanol, ac etone, chloroform&dioxane.

Steam distillation:

It is used in rare cases, for liquid alkaloids, e.g. coniine, nicotine & sparteine.

. Crystallization of sparingly soluble salts:

The choice of acid is unlimited, but HCl, HBr, oxalic, picric, perchloric, sulfuric, maleic, tartaric acids are among the widely used acids.

The general procedure for hydro halides involves dissolving the crude base in methanol or acetone & adding an ethereal solution of the acid.

Oxalates, picrates & perchlorates are usually formed by mixing methanolic solutions of the base & the acid.

Distribution between immiscible solvents:

- The alkaloids are taken up in a dilute acid solution.
- From this it may be possible to recover the alkaloids by the addition of ammonium hydroxide solution & extract with water immiscible organic solvent.
- The choice of organic solvent for this method is usually limited to benzene, chloroform or ether.

THANK YOU