

# ***In Vitro* Production of Secondary Metabolites**

**Presented by**

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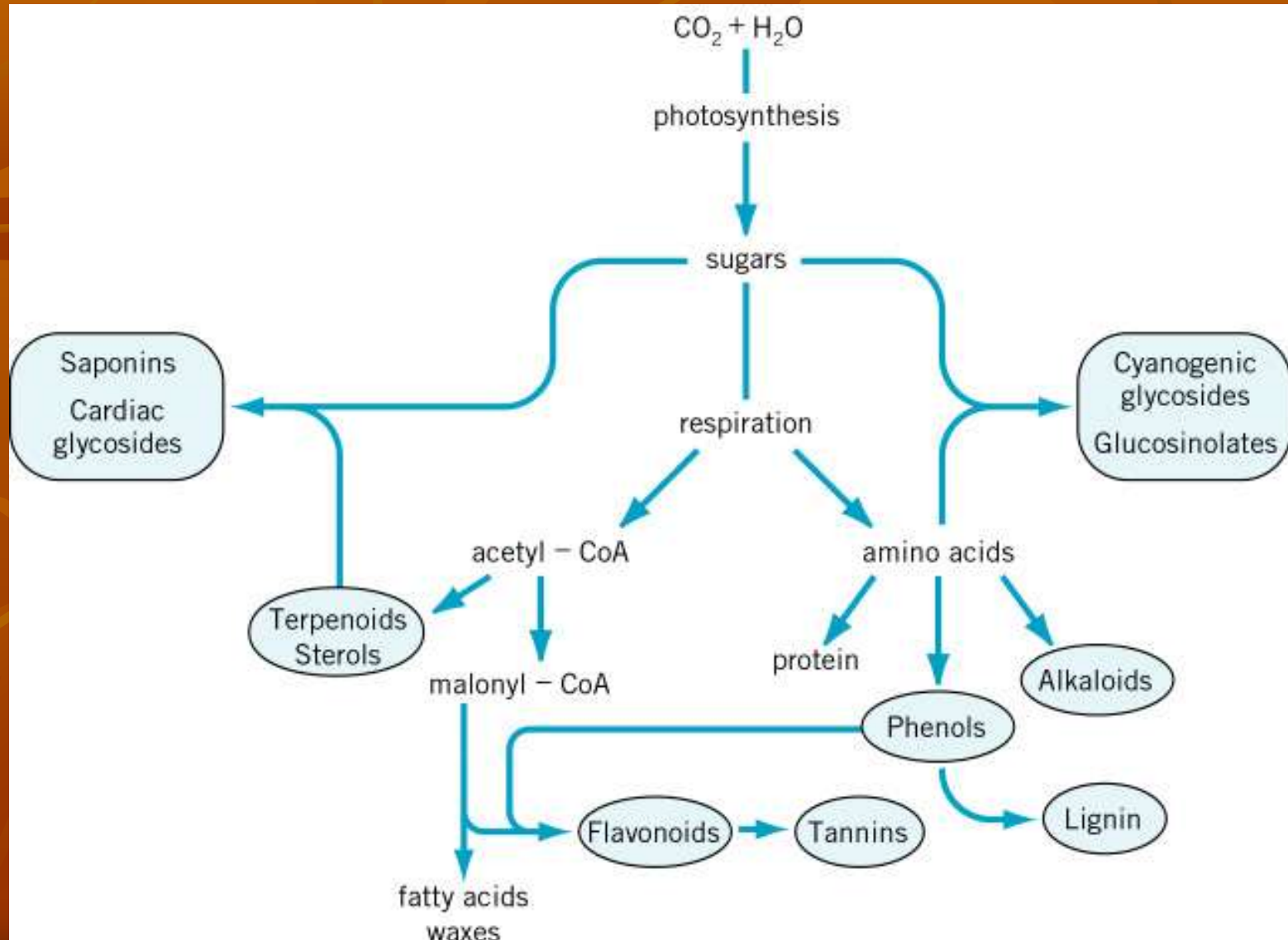
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**TRICHY – 24.**

# Metabolites

- **Primary metabolites:** Molecules that are essential for growth and development of an organism.
- **Secondary metabolites:** molecules that are not essential for growth and development of an organism.

# Secondary metabolites are derived from primary metabolites



# Why secondary metabolites?

- Chemical warfare to protect plants from the attacks by predators, pathogens, or competitors
- Attract pollinators or seed dispersal agents
- Important for abiotic stresses
- Medicine
- Industrial additives

# Secondary metabolites

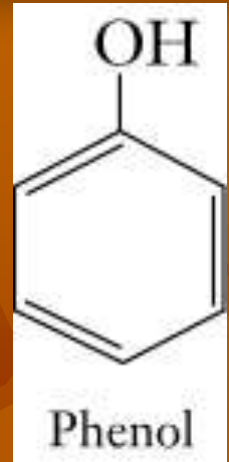
- Possibly over 250,000 secondary metabolites in plants
- Classified based on common biosynthetic pathways where a chemical is derived.
- Four major classes:  
Alkaloids, glycosides, phenolics,  
terpenoids

# Alkaloids

- Most are derived from a few common amino acids (i.e., tyrosine, tryptophan, ornithine or arginine, and lysine)
- Compounds have a ring structure and a nitrogen residue.
- Indole alkaloids is the largest group in this family, derived from tryptophan
- Widely used as medicine

# Phenolics

- Derived from aromatic amino acids, such as phenylalanine, tyrosin, and tryptophan.
- All contain structures derived from phenol
- Some examples:
  - Coumarins: antimicrobial agents, feeding deterrents, and germination inhibitors.
  - Lignin: abundant in secondary cell wall, rigid and resistant to extraction or many degradation reagents.



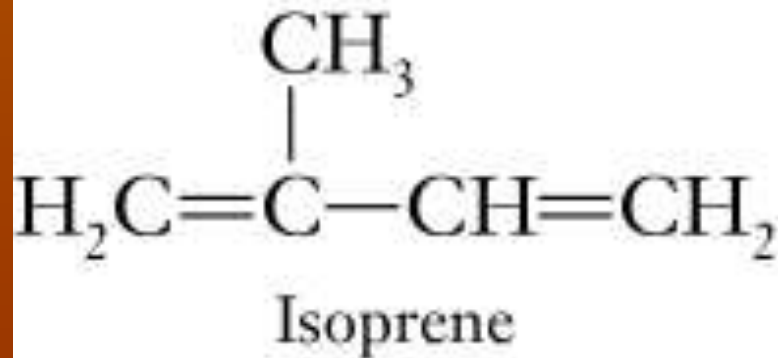
# Glycosides

- Compounds that contain a carbohydrate and a noncarbohydrate
- Glucosinolates: found primarily in the mustard family to give the pungent taste.

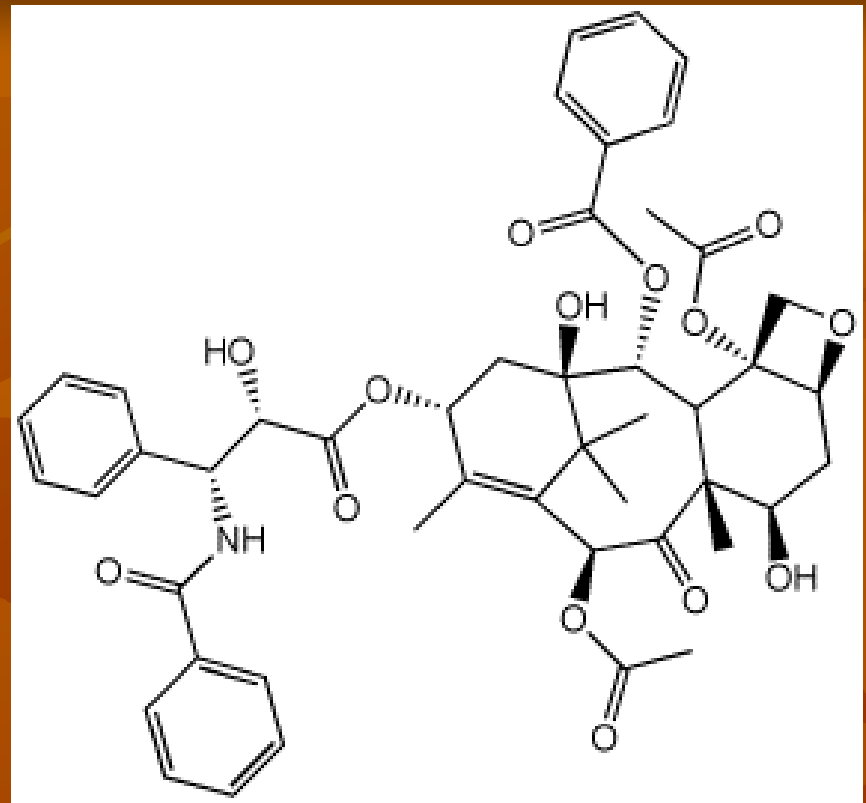


# Terpenoids

- Terpenes are generally polymers of 5-carbon unit called isoprene
- Give scent, flavors, colors, medicine...
- Three plant hormones are derived from the terpenoid pathway.



# Taxol



- Taxol is a terpenoid
- "the best anti-cancer agent" by National Cancer Institute
- Has remarkable activity against advanced ovarian and breast cancer, and has been approved for clinical use.

# How can we make more usual plant secondary metabolites?

- Plant more trees, extract directly from plants
- Genetic engineering
  - Increase the yield
  - Introduce the biosynthetic pathway to other plants that normally does not produce certain compound
- Tissue culture to produce a compound
- Chemical synthesis
- Semi-chemical synthesis

# Culture Conditions

- ❖ Environment – Culture Medium : Simple nutrient medium – MS, B5, - Mineral Salts Macro and Micro.
- ❖ Carbon Source – Sucrose, glucose, fructose, maltose.
- ❖ GRs- Auxins, Cytokinins, GA<sub>3</sub>
- ❖ Amino acids- glycine, glutamine, proline, phenyl, alanine, arginine
- ❖ Vitamins – nicotinic acid, Pyriodoxine Hcl, thiamine Hcl, biotin, folic acid, cyanocobalamine
- ❖ Organic supplements: Yeast extract, malt extract, casein hydrolysase, coconut water.
- ❖ Gelling agents : Agar – Agar

- ❖ Temp- of incubation 22 -28 ° C
- ❖ Illumination – 0- 5000 lux, photoperiod – 8-16 h
- ❖ Subculture of tissues : 2-8 weeks for Static cultures & 1-2 weeks for cell suspensions.

# Culture initiation

- ✓ Explant – callus induction
- ✓ Growth & Secondary Metabolite Production
- ✓ Growth curve – sigmoid
  - Lag –no growth cells adjust to the new media after subculture
  - Stationary - primary metabolism & cell proliferation come to a halt as nutrients in the medium are exhausted. Dry weight may decline as cells utilize stored reserve material.
  - Exponential – primary metabolism increases & tissue proliferates rapidly with the consumption of medium nutrients

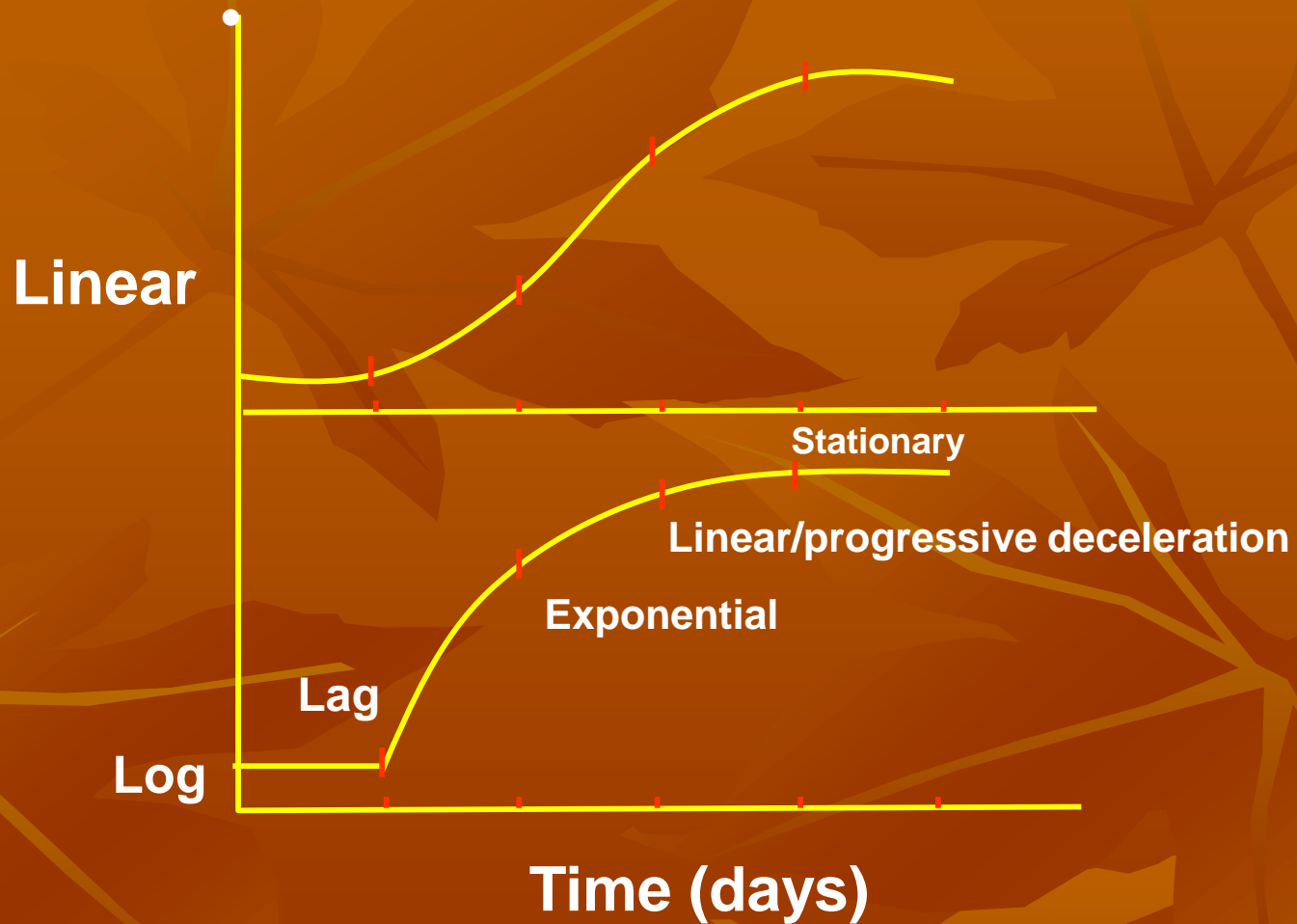
# Growth and Metabolism of Plant in *In Vitro*

## (i) Batch / Solid Culture

1. Lag phase
2. Log / Exponential Phase
3. Stationary Phase
4. Declining Phase

## (ii) Continuous cultures

# Batch Culture Growth Cycle (Passage)





**Factors  
(Stress)**



**Abiotic – Light, Temperature, Nutrients,  
Growth regulators, Precursors**

**Biotic – Elicitors, Biotransformation, Hairy  
root culture, Genetic transformation**

# Temperature

- ❖ Effect on enzymes 25 - 28° C
- ❖ pH – 5-6 effect on anthocyanin, anthraquinones, alkaloids, anthocyanin degrades at higher pH.
- ❖ Aeration – in liquid shake flasks or in bioreactors – secondary metabolite production
- ❖ Anaerobic – direct effect on primary metabolism & consequently production of secondary metabolites or direct effect on secondary metabolites. Anaerobic - ethanol production

## Nutrients

- Carbon – 2-3 %
- Nature & Concentration of 'C'
- Sucrose is better than all other carbohydrate sources for growth and concentrations >3% often enhance the biosynthesis of phytochemicals by heterotrophic plant cells.
- Interactions of carbohydrates & nitrogen is important in cell dry weight, fresh weight, protein content and cell proliferation.

❖ Maximum amount of mineral nitrogen assimilated by cultured cells is used for the biosynthesis of a – acids, proteins (enzymes also) & nucleic acids.

❖ High  $\text{NH}_4^+$  - inhibitors to secondary metabolites

❖ Low or nil – increase to secondary metabolites.

❖ Phosphate

❖ Increased phosphate – alkaloid, anthraquinone

❖ Inorganic phosphate - photosynthesis, respiration, phospholipids synthesis.

# Light

- ❖ Blue Light – anthocyanin in *Haplopappus gracilis*
- ❖ White Light – anthocyanin in *Catharanthus roseus*, *Populus* sps.,
- ❖ Napthoquinone inhibited – *Lithospermum erythrorhizon* in blue or white light
- ❖ Anthocyanin – *Daucus carota* – in white light

*Helianthus tuberosum*

*Linum usitatissimu*

*Vitis vinifera*

- UV (280-320 nm) – stimulate flavone glycoside synthesis in *Petroselinum hortense* cell suspension cultures.
- UV glyceollin in *Glycine max*

### White light

- Coumarins, alkaloids – *Ruta graveolens*
- Indole alkaloids – *Catharanthus roseus*
- Light – direct on development of chlooplasts enzymes of photosynthesis, lipid metabolism, phytochrome

# Precursors

Precursors molecules which are directly incorporated into synthesis of secondary metabolites

<i>Ruta graveolens</i>	4 – OH 2 – Quinoline	Dictamine – 0.6% DW
<i>Chinchona ledgerianer</i>	Tryptophan	Quinoleines – 0.9% DW
<i>Lithospermum erythrorhizon</i>	Phenylalanine	Shikonin 37- 126 µg -1 FW
<i>Ephedra gerardiana</i>	Phenylalanine	Ephedrine 0.17 – 0.5 % DW

Need to optimize the growth and production conditions for each species and strain, and also for each metabolite

# Advantage of callus and cell suspension culture

- ❖ **The production of secondary metabolites from plant cell and tissue culture, which are immediate relevance to the industry.**
- ❖ **Independence from environmental factors.**
- ❖ **The production system is not limited by seasonal consideration.**
- ❖ **The more consistent product quality and yield.**
- ❖ **The product is free from microbes.**
- ❖ **The synthesis of novel natural products, which are not normally produced in normal plants.**
- ❖ **A means of synthesizing novel natural product where the source plant is difficult to grow.**



# Immobilization

## Type of immobilization

Gel entrapment  
Biofilms  
Adsorption  
Foam immobilization  
Membrane

## Advantages of immobilization

1. Facilitate the use of continuous flow process.
2. Facilitate sequential chemical treatment of the cells or organelles and harvest of metabolites from the medium.
3. The biocatalyst can be reused and can be easily separated from the reaction medium.
4. Control aggregation of cells.
5. The cell/ cell contact induced by immobilization can be beneficial for differentiation process and for secondary metabolites production. Since differentiation process is a prerequisite for the production of secondary plant products.
6. Immobilization protects the sensitive plant cells against shear forces.
7. Immobilized cells can be used for much longer periods than free cells.

## Disadvantages

Some of the problems with immobilized cells are the introduction of gradients in the gel beads, the necessity of product excretion to the use of immobilized cells and the fact that some immobilization materials can affect cell viability in a negative way. Besides immobilization introduces an extra cost factor.

# Plant species with immobilized cells employed for the production secondary Metabolite (s)

Plant culture species	Immobilization method	Substrate	Product
<i>Catharanthus roseus</i>	Entrapment in agarose	Cathenamine	Ajmalicine
<i>Digitalis lanata</i>	Entrapment in alginate	Digitoxin	Digoxin
<i>Capsicum frutescens</i>	Entrapment in polyurethane foam	Sucrose	Capsaicin
<i>Catharanthus roseus</i>	Entrapment in alginate, agarose, carrageenin	Sucrose	Ajmalicine
<i>Petunia hybrida</i>	Entrapment in hollow fibres	Sucrose	Phenolics
<i>Morinda citrifolia</i>	Entrapment in alginate	Sucrose	Anthraquinone
<i>Solanum aviculare</i>	Attachment polyphenylene beads	Sucrose	Steroid glycosides
<i>Glycine max</i>	Entrapment in hollow fibres	Sucrose	Phenolics

## Elicitors

- Keen – Coworkers (1972) – elicitation response.
- Fungal cultures – fresh cultures homogenized, autoclaved at 121°C for 20 min, and suitably diluted fungal preparations or chemicals are used to evaluate the elicitation effect.
- Eg. *Pythium*, *Fusarium*, *Phytophthora*, *Alternaria*, *Penicillium* etc.

## *Elicitor-induced secondary metabolites production in plant cell culture*

<b>Elicitor microorganism</b>	<b>Plant cell culture (s)</b>	<b>Secondary metabolite (s)</b>
<i>Aspergillus niger</i>	<i>Cinchona ledgeriana</i> , <i>Rubia tinctoria</i>	Anthraquinones
<i>Pythium aphanidermatum</i> <i>Botrytis</i> sp	<i>Catharanthus roseus</i>  <i>Papaver somniferum</i>	Ajmalicine, Striclosidine, Catharanthine Sanguinarine
<i>Phytophthora megasperma</i>	<i>Glycine max</i>	Isoflavonoids, Gluceollin
<i>Dendryphion</i> sp	 <i>Papaver sominiferum</i>	 Sanguinarine
<i>Alternaria</i> sp	 <i>Phaseolus vulgaris</i>	 Phaseollin
<i>Fusarium</i> sp	 <i>Apium graveolens</i>	 Furanocomarins
<i>Phythium aphanidermatum</i>	 <i>Daucus carota</i>	 Anthocynins
<i>Penicillium expansum</i>	 <i>Sanguinaria canadensis</i>	 Benzophenanthridine Alkaloids

# Biotransformations

- Biotransformations is of two types
  - (i) Transformation of low cost precursors into valuable product or conservation of racemic /inactive compounds into active forms

Eg. Conversion of D- menthol to L – menthol

- (ii) Transformation with the help of Agrobacterium

# Biotransformation by plant cell culture

Plant cell culture	Substrate	Product
<i>Digitalis lanata</i>	Digitoxin	Digoxin
<i>Papaver somniferum</i>	Codeinone	Codeine
<i>Nicotiana tobacum</i>	Carvoxine	Cavaxone
<i>Daucus carota</i>	Digitoxigenin	Periplogenin
<i>Mucuna pruriens</i>	L – Tyrosine	L- Dihydroxy phenylalanine (L-DOPA)
<i>Mentha sp</i>	(-)- Methone	(+) – Neomenthol
<i>Coffea arabica</i>	Vanillin	Vanillin –D – glucoside
<i>Solanum tuberosum</i>	Solavetivone	Hydroxylated derivatives
<i>Galium mollugo</i>	2-Succinyl benzoate	Anthra quinones
<i>Datura sp</i>	Hydroquinone	Arbutin
<i>Citrus sp</i>	Valencene	Nootkatone
<i>Chiisya ternata</i>	Ellipticine	5-Formyl ellipticine
<i>Digitalis purpurea /</i> <i>Stvia rebandiana</i>	Steviol	Steviocide, Steviobiocide



# Hairy root – *Agrobacterium rhizogenes*

Eg. *Atropa belladonna* – Atropine

- *Datura stramonium* – Hyoscamine
- *Hyscyamus multicus* – Hyoscamine
- *Cathyranthus roseus* – Ajmaline, Serpentine, Catharnthine
- *Lithospermum, Erythrorhizon* – Shikonin
- *Chinchona ledgeriana*

❑ The formation of transformed roots following infection of plants with *A. rhizogenes* is limited to dicotyledonous species only.

❑ The use of transformed root cultures for the production of secondary metabolites may also be restricted to species in which the products are synthesized in roots of intact plants.

## Plant species used in hairy root cultures for the production of secondary metabolite (s)

Plant species	Secondary metabolite (s)
<i>Nicotiana tabacum</i>	Nicotine, anatabine
<i>Atropa belladonna</i>	Atropine
<i>Datura stramonium</i>	Hyoscyamine
<i>Lithospermum erythrorhizon</i>	Shikonin
<i>Catharanthus roseus</i>	Ajmalicine, serpentine
<i>Cinchona ledgeriana</i>	Quinine alkaloids
<i>Mentha vulgaris</i>	Monoterpenes
<i>Solanum laciniatum</i>	Steroid alkaloids





# Production

# Bioreactors

- Optimization of secondary metabolite production in plant cells.
- 1984- *Lithospermum erythrorhizon* cells in Japan – 750 bioreactor for shikonin (a dye & chemical compound).
- Sanguinarine – *Papaver somniferum* cells. USA
- Vanilla flavour USA
- Germany Co., Taxol, *Taxus* cell cultures 75 m<sup>3</sup>.

**Growth period – bacteria - < 1hr doubling time  
plant cells – 24 – 72 h**

## **Batch cultivation**

**Phase I - lag period – adaptations to the new environment of the bioreactor**

**II - exponential growth – growth required nutrients are present in excess (O<sub>2</sub> also)**

**III - nutrient- limited rate – (C,N,P)**

**IV Stationary (Sec. metabolites) – growth ceases & the cell density reaches maximum, cells remain metabolically active.**

**V Cell lysis – generally not in bioreactors**

**Plant cell process systems is divided into three areas**

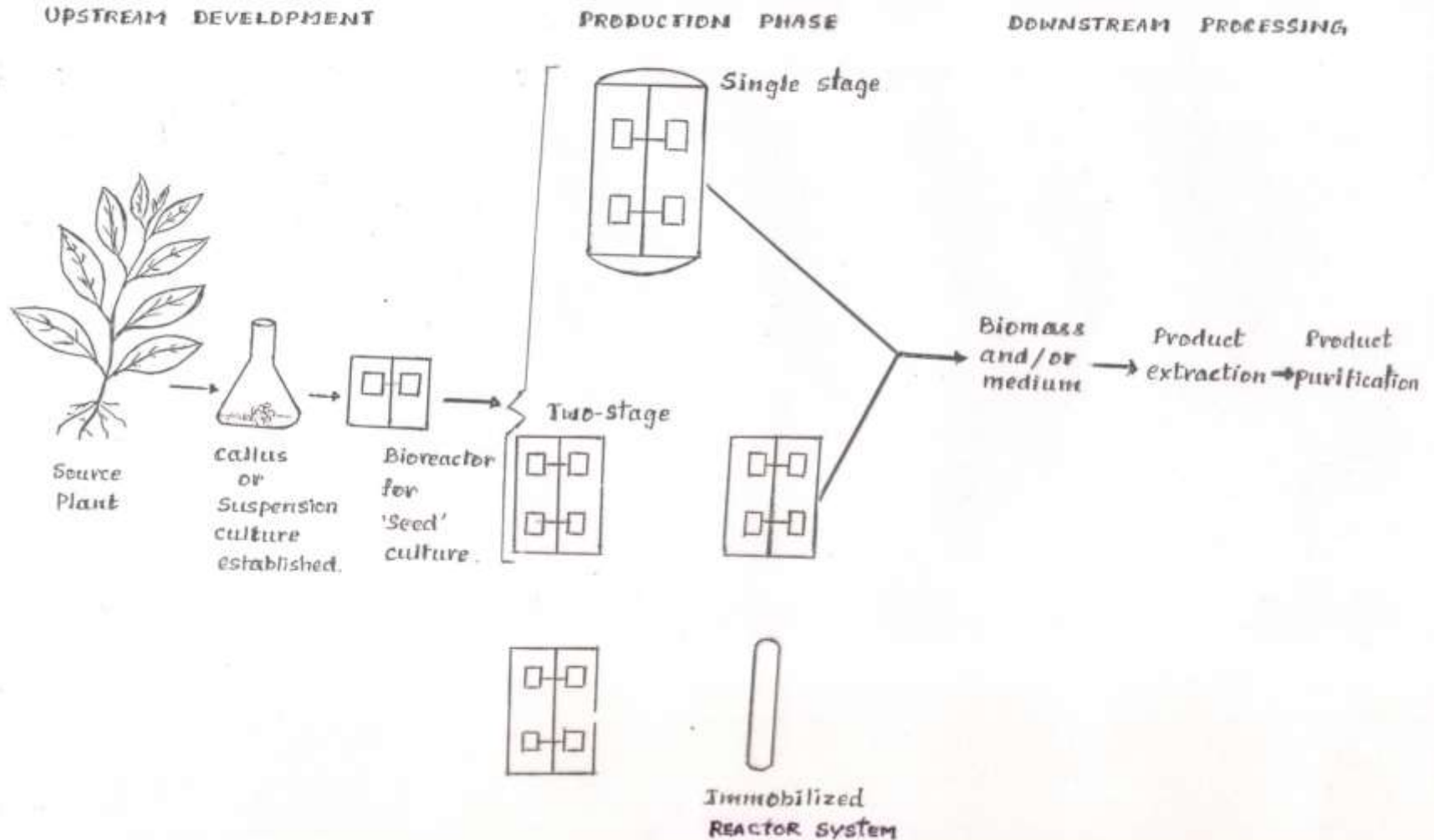
- Upstream processing**
- the production or reactor phase**
- downstream processing**

**First phase – batch/ continuous system**

**Second phase – batch cultures / immobilized cell technology**

# Steps of large scale secondary metabolites production

## STAGES IN THE DEVELOPMENT AND OPERATION OF A PLANT BIOTECHNOLOGY PROCESS



**List of some culture secondary products  
obtained through in vitro culture**

<b>Species</b>	<b>Products/Compounds</b>	<b>Culture type</b>
<i>Anethum graveolens</i>	Scopoletin	Root
<i>Atropa belladonna</i>	Hyoscyamine, Scopolamine	Root
<i>Calystegia sepium</i>	Hyoscyamine, Scopolamine	Root
<i>Catharanthus roseus</i>	Ajmalicine, Catharanthine Cephaeline	Root
<i>Cephaelis ipecacuanha</i>	Valepotriates	Root
<i>Centranthus macrosiphon</i>	Valepotriates	Root
<i>Centranthus ruber</i>	Quinine, Quinidine	Root
<i>Cinchona ledgeriana</i>	Hyoscyamine, Scopolamine	Root

Species	Products/Compounds	Culture type
<i>Citrus paradisi</i>	Naringin	Shoot
<i>Cinchona ledgeriana</i>	Quinine, Quinidine	Shoot
<i>Cinchona succirubra</i>	Cinchonidine	Shoot
<i>Chrysanthemum cinerariaefolium</i>	Pyrethrine, Cinerin	Shoot
<i>Chrysanthemum americanum</i>	O-methylated flavanol glucosides	Shoot
<i>Datura innoxia</i>	Hyoscyamine, Scopolamine	Shoot
<i>Digitalis purpurea</i>	Digitoxin	Shoot
<i>Digitalis lanata</i>	Digitoxin	Shoot
<i>Digitalis lutea</i>	Digitoxin	Shoot
<i>Digitalis mertonensis</i>	Digitoxin	Shoot
<i>Digitalis ferruginea</i>	Digitoxin	Shoot
<i>Digitalis ambigua</i>	Digitoxin	Shoot
<i>Dioscorea composita</i>	Diosgenin	Shoot
<i>Foeniculum vulgare</i>	Anethole	Shoot
<i>Heimia salicifolia</i>	Vertine, Lyfoline	Shoot
<i>Lavandula angustifolia</i>	Vertine, Lyfoline	Shoot
<i>Origanum vulgare</i>	Monoterpenes	Shoot
<i>Papaver bracteatum</i>	Thebaine	Shoot
<i>Papaver somniferum</i>	Thebaine	Shoot
<i>Pelargonium fragrans</i>	Pinene	Shoot
<i>Pelargonium graveolens</i>	Geraniol, citronellol	Shoot
<i>Pelargonium tomentosum</i>	Methone, isomenthone	Shoot



Species	Products/Compounds	Culture type
<i>Atropa belladonna</i>	Artopine	Hairy root
<i>BetaBeta</i>	Betanin	Hairy root
<i>Bidens</i>	Thipenes	Hairy root
<i>Catharantthus roseus</i>	Ajrnalicine,Catharanthine	Hairy root
<i>Cinchona ledgeriana</i>	Quinine, Quinidine	Hairy root
<i>Coreopsis</i>	Thiophenes	Hairy root
<i>Datura stramonium</i>	Hyoscyamin, Scopolamine	Hairy root
<i>Dubosia hopwoodii</i>	Hyoscyamin, Scopolamine	Hairy root
<i>Digitalis lanata</i>	Digoxin	Liquid culture
<i>Digitalis purpurea</i>	Digitoxin	Liquid culture
<i>Dioscorea deltoidea</i>	Diosgenin	Suspension
<i>Papaver somniferum</i>	Codeine	Suspension
<i>Papaver somniferum</i>	Morphine	Suspension
<i>Rauwolfia serpentina</i>	Reserpine	Suspension
<i>Matricaria chamomilla</i>	Sesquiterpenes	Suspension
<i>Pimpinella anisum</i>	Anethole	Suspension
<i>Coleus blumei</i>	Rosemarinic acid	Suspension
<i>Catharanthus roseus</i>	Serpentine	Suspension
<i>Catharanthus roseus</i>	Ajmalicine	Suspension
<i>Nicotiana tabacum</i>	Glutathione	Suspension



Species	Products/Compounds	Culture type
<i>Papaver bracteatum</i>	Thebaine	Callus
<i>P. somniferum</i>	Thebaine	Callus
<i>Mentha piperita</i>	Geraniol, linolol	Callus
<i>Mentha piperita</i>	Menthone, menthol	Callus
<i>Matricaria chamomilla</i>	Sesquiterpenes, a-bisabolol	Callus
<i>Pelargonium spp.</i>	Monoterpenes	Callus
<i>Cassia tora</i>	Anthraquinones	Callus
<i>Panax ginseng</i>	Ginsenosides	Callus
<i>Lithospennum erythrorhizon</i>	Shikonin	Callus

Sources: Charlwood et al. (1990), Petersen (1993) Starford (1991), Rhodes et al 1990)