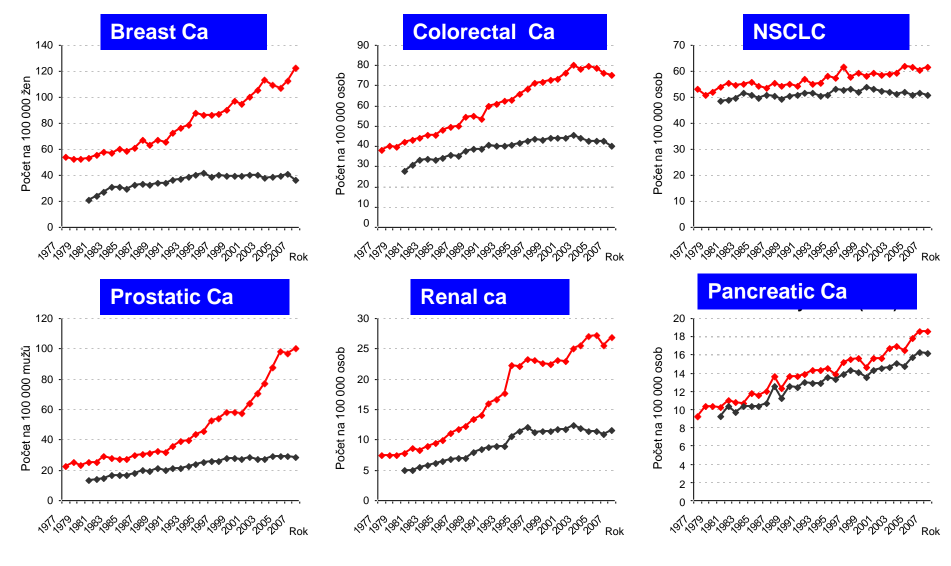
### **Anticancer drugs**



#### Cancer epidemiology - incidence and mortality in the Czech Republic



Czech Cancer Care in Numbers 2015

🛏 incidence

### **Complex Cancer Treatment**

- Surgery
- Radiotherapy

Pharmacoterapy

— Psychotherapy, physiotherapy, nutrition care

 $M \vdash I$ 

### **Pharmacotherapy**

#### -cytostatic agents

-classification according to the mechanism of action

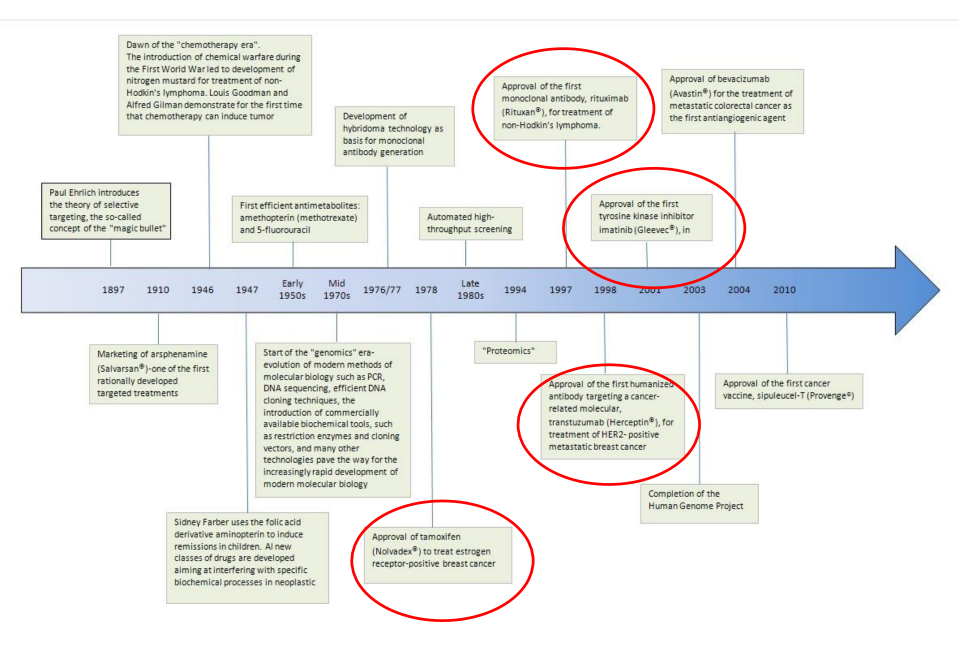
#### -endocrine (hormonal) therapy

#### -targeted therapy

monoclonal antibodies targeting extracellular part of receptors
 tyrosine kinase inhibitors / intracellular signaling cascades inhibitors

#### -pain management, supportive care

MUNI Med



# **Cytostatic drugs**

-route of administration:

—parenterally (i.v. bolus, infusion, intrathecally, intravesically...)
—orally

- —posology: dose in mg/m<sup>2</sup> or mg/kg
- -monotherapy and combination regimens

—repeated administration in cycles pause = patient's recovery, prevention of severe AE + ,,waking" dormant cells in G<sub>o</sub> phase

### **Cytostatic drugs**

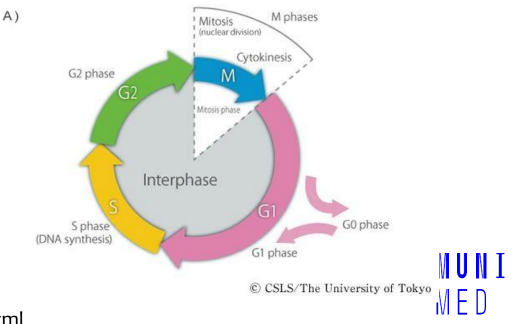
#### Different efficacy according to the cell cycle phase:

#### -Cell cycle non-specific cytostatics (e.g., busulfan)

—Cell cycle specific cytostatics:

-Phase-nonspecific (e.g., some of alkylating agents)

-Phase-specific (e.g., antimetabolites, taxanes)



http://csls-text3.c.u-tokyo.ac.jp/active/13\_01.html

### **Cytostatics according to their MoA**

#### 1. Drugs that damage the structure of DNA

- a) Alkylating agents
- b) Platinum derivatives
- c) Intercalating agents
- d) Bleomycin

#### 2. Drugs that inhibit key enzymes of DNA metabolism

- a) Antimetabolites:
  - i. Purine analogues
  - ii. Pyrimidine analogues
  - iii. Folic acid analogues
  - iv.Hydroxyurea
- b) Topoisomerase inhibitors:
  - i. Inhibitors of topoisomerase I camptothecins
  - ii. Inhibitors of topoisomerase II podophyllotoxins

#### 3. Drugs that alter microtubules

- a) Inhibitors of tubulin polymerization Vinca alkaloids
- b) Inhibitors of tubulin depolymerization taxanes
- 4. Others
  - a) Drugs that inhibit protein synthesis L-asparaginase

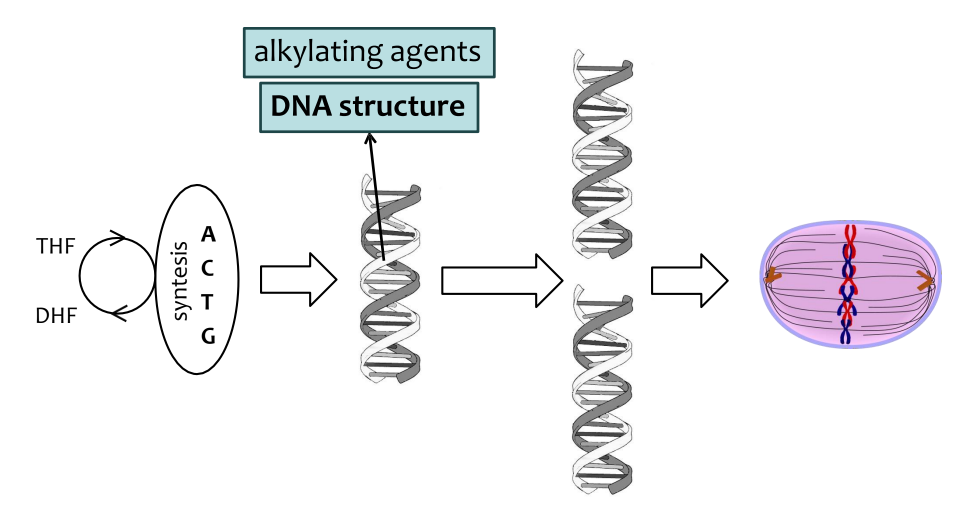
### **Drug groups overview**

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# 1a) Alkylating agents

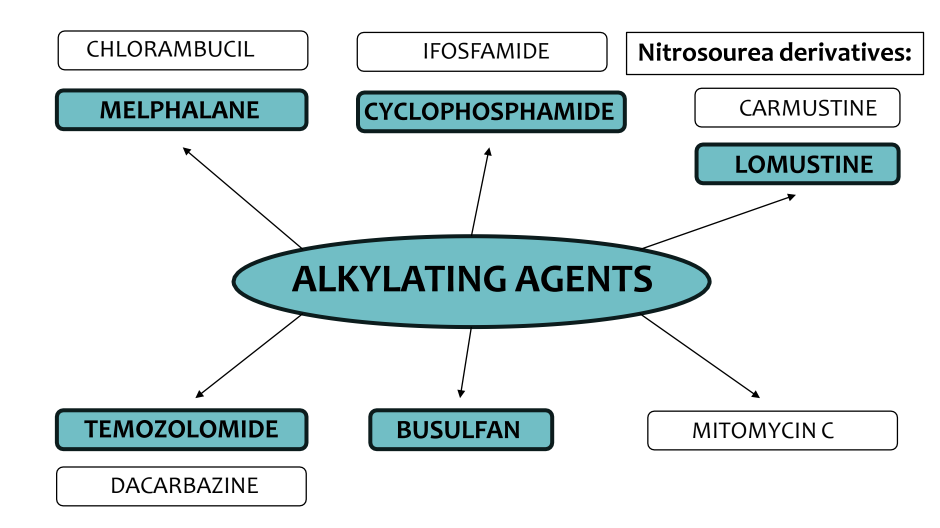


# 1a) Alkylating agents

- MoA: transfer of the alkyl group on nitrogen in nucleobases, covalent bond between two guanines of one or two DNA strands
  - Inhibition of replication, cell cycle arrest
- 50s: first derivatives of sulphur mustard in the clinical practice



# 1a) Alkylating agents – main drugs



# 1a) Alkylating agents

#### Melphalane

- i.v., p.o. administration
- treatment of hematological malignancies and solid tumors

#### Cyclophosphamide

- i.v., p.o. administration
- prodrugs  $\rightarrow$  CYP450  $\rightarrow$  cytotoxic metabolites
- AE: urotoxicity, emetogenity
- low doses immunosuppressant
- hematological malignancies and solid tumors

#### Lomustine

- p.o. administration
- lipophilic, crosses  $BBB \rightarrow$  treatment of brain tumors

# 1a) Alkylating agents

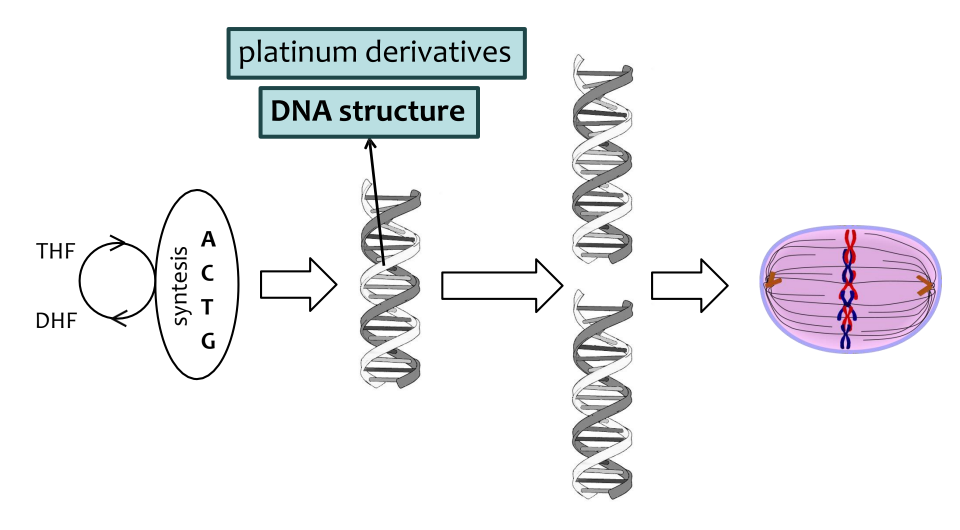
#### Temozolomide

- 100% bioavailability after oral administration
- crosses BBB  $\rightarrow$  treatment of brain tumors

#### Busulfan

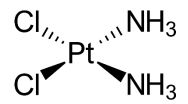
- i.v., p.o. administration
- bone marrow transplantation
- treatment of hematological malignancies

# 1b) platinum derivates



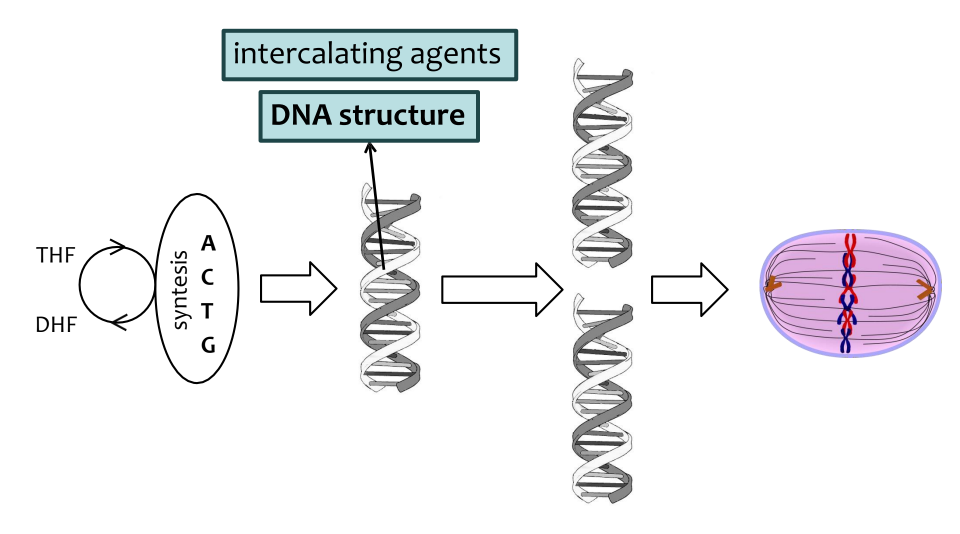
# 1b) platinum derivates

- **MoA:** binding on DNA, cross-linking of DNA strands, inhibition of topoisomerases
- AE most important: emetogenity, nephrotoxicity
  - AE are dose-dependent
  - prevention of nephrotoxicity: i.v. hydration, forced diuresis
- **cisplatin** high nephrotoxicity
  - treatment of solid tumors



- others:
  - carboplatin
  - oxaliplatin typical neurotoxicity

# 1c) intercalating agents



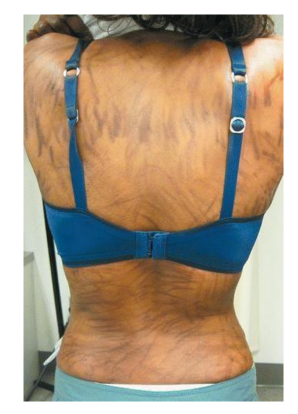
# 1c) intercalating agents

#### **Anthracyclines**

- MoA: intercalation = insertion between base pairs, binding of DNA strands
- AE typical toxicity: acute and chronic cardiotoxicity
- cardioprotective cumulative dose = restraint of therapy (e.g., doxorubicin 550 mg/m<sup>2</sup>)
- i.v., intravesical administration
- doxorubicin
  - treatment of hematological malignancies and solid tumors
  - modern dosage form (PEGylated liposomes) higher cumulative dose (860 mg/m<sup>2</sup>)
- others: epirubicin...

# 1d) bleomycin

- MoA: intercalation between base pairs
   + inhibition of thymine incorporation
   → breaks → DNA fragmentation
   ("radiomimetic" effect)
- i.v. administration
- treatment of solid tumors
- typical AE: fever,
   hyperkeratosis and
   hyperpigmentation of skin
   (flagellate = whip-like)
- risk of anaphylactic reaction





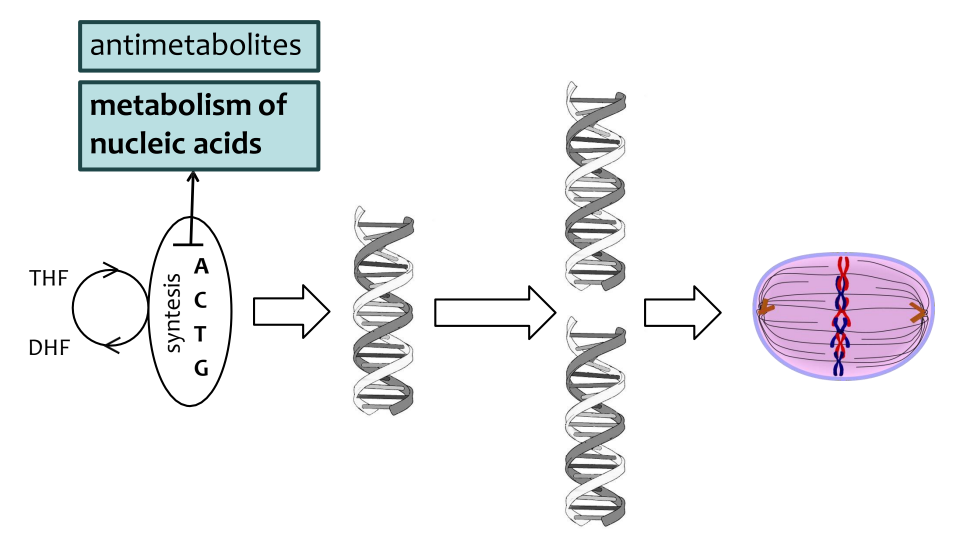
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### 2a. antimetabolites



### 2a. antimetabolites

- MoA: false substrates = affinity to target structure, loss of endogenous effect  $\rightarrow$  blockade of nucleic acid synthesis, inhibition of nucleotides metabolism enzymes, production of *non-sense* DNA sequences
- prodrugs: intracellular activation mostly by phosphorylation
- *a) purine analogues* 6-mercaptopurine, azathioprine, fludarabine...
- **b)** pyrimidine analogues fluorouracil, capecitabine, gemcitabine...
- c) folic acid analogues methotrexate, pemetrexed...

### 2a. Antimetabolites – i/ purine analogs

#### 6-Mercaptopurin

- MoA: inhibition of purine nucleobases biosynthesis *de* novo, inhibition of mutual conversion of purine nucleotides
- thiopurin methyltransferase (TPMT): MP  $\rightarrow$  MeMP
  - genetic polymorphism <sup>t</sup>oxicity / Jefficacy
  - available pharmacogenetic testing of TPMT
- p.o. administration, treatment of hematologic malignancies
- azathioprine prodrug of MP, immunosuppressant

### 2a. Antimetabolites – ii/ pyrimidines analogs 5-Fluorouracil

- MoA: incorporation to RNA + inhibition of thymidylate synthetase
- combined chemotherapeutic regimens of solid cancers (i.v.)
- **AE typical toxicity:** GIT toxicity (mucositis)
- biochemical modulation of effect: leucovorin (folinic acid) enhances binding on thymidylate synthetase, i.v. administered before FU
  - "FUFA" regimen = colorectal carcinoma
- capecitabine prodrug

#### 2a. Antimetabolites – iii/ folic acid analogs Methotrexate

- **MoA:** inhibition of dihydrofolate reductase, thymidylate synthetase and other enzymes
- i.v., intrathecal administration, p.o.
- leucovorin (folinic acid) "rescue therapy", antidote forces free MTX out of healthy cells ; in cancer cells, polyglutamylation is more intensive → more MTXPG → MTXPG cannot be forced out
   TDM – calculation of time interval from MTX administration, frequently in pediatric patients, less frequent in adults
- AE typical toxicity:

**nephrotoxicity** – precipitation (acute renal failure) prevention: hydration, urine alkalinization (pH 7–7,5) pneumotoxicity

- low-dose MTX = immunosuppressant (p.o.)
- high-dose MTX = hematological malignancies

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# **2b. Topoisomerase inhibitors** Α THF syntesis C Т G DHF **DNA replication** topoisomerase inhibitors

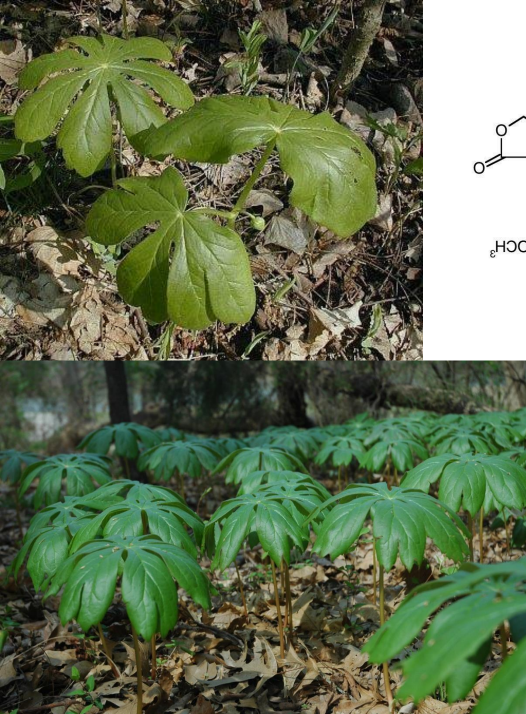
# **2b. Topoisomerase inhibitors**

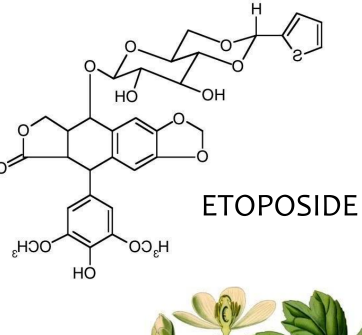
#### **Topoisomerase I inhibitors – camptothecins**

- plant-derived drugs identification in bark of the tree *Camptotheca acuminata*
- derivatives: irinotecan, topotecan
  - treatment of solid tumors

#### **Topoisomerase II inhibitors – podophyllotoxins**

- plant-derived drugs identification in *Podophyllum peltatum*
- derivatives: etoposide, teniposide
  - treatment of solid tumors (etoposide) and hematological malignancies (teniposid)







Podophyllum peltatum L. Image processed by Thomas Schoepke www.plant-pictures.de

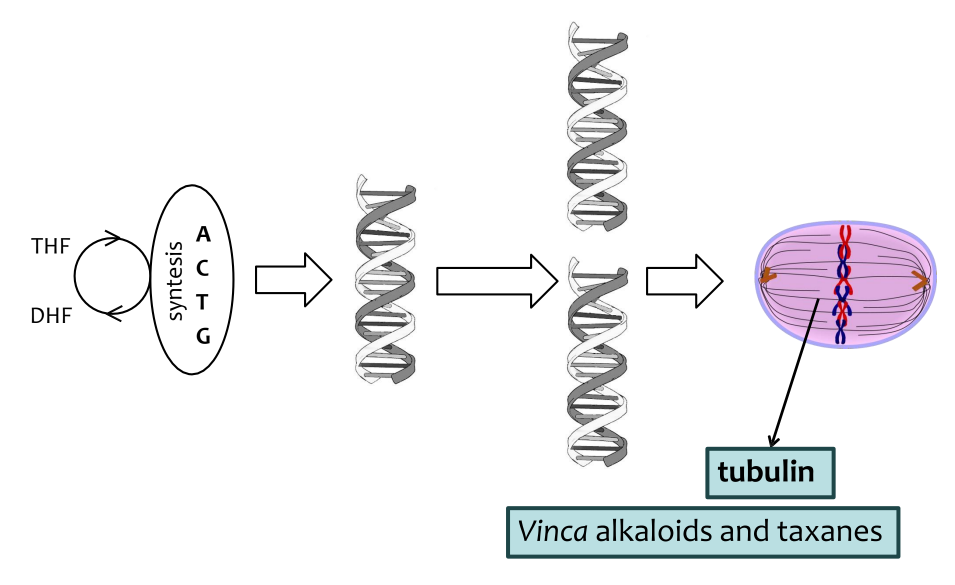
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### 3. Tubulin alterating cytostatics



### **3a. Vinca alkaloids**

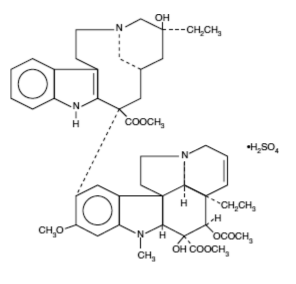
- ant-derived drugs
- MoA: inhibition of tubuline dimers polymerization inhibition of mitotic spindle formation, depolymerization prevails
- i.v. administration, some for p.o. (vinorelbine)
- treatment of hematological malignancies and solid tumors
- AE typical toxicity: peripheral neuropathy
- original alkaloids: vincristine, vinblastine
- semisynthetic derivatives: vinorelbine, vindesin, vinflunine increased affinity to mitotic spindle tubulin, JAE



### Vinca alkaloids

- identification:
   lesser periwinkle
   (Vinca minor)
- isolation:
   Cataranthus roseus





VINBLASTINE

### **3b. taxanes**

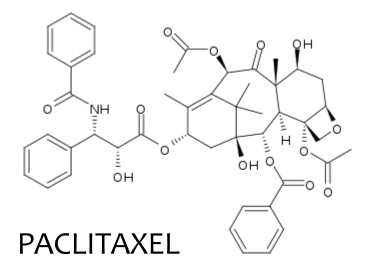
- plant-based drugs
- MoA: inhibition of tubulin depolymerization
- i.v. administration treatment of solid tumors
- AE typical toxicity: neurotoxicity
- paclitaxel, docetaxel, cabazitaxel

- modern dosage form: paclitaxel conjugated with albumine nanoparticles

- transporter protein for albumine in cancer cells = better distribution from circulation into the tissues
- ↓toxicity, ↑efficacy

### **Taxanes**

 identification and isolation: *Taxus brevifolia* (Pacific yew) a *Taxus baccata* (European yew)





# **Combination of cytostatics**

- monotherapy

- combination regimens – examples:

FUFA FOLFOX ABVD BEACOPP

fluorouracil, folinic acid folinic acid, fluorouracil, oxaliplatin doxorubicin, bleomycin, vinblastine, dacarbazine bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristin, procarbazine, prednisone

#### "Targeted therapy"in oncology

# **Targeted therapy in oncology (biological therapy)**

- "target" should be on
- A/ tumor cells
- -Cell membrane receptor extracellular part or/ intracellular signalling pathway
- B/ immune system (specific T-cells) cancer immunotherapy
- -Immune check-point inhibitors (anti-CTLA-4 or anti-PD(L)1)