Complex Diseases, Success and Failure

Finding the "right" target → valid targets

which constrains are limiting factors?

Dosage, bioavailability, actual drug concentration in the respective compartment (cell, organelles)

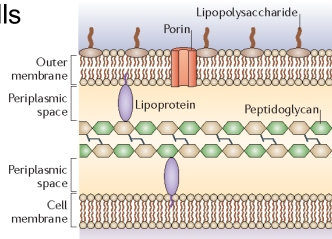
Are we competing against a natural substrate, e.g. ATP? (concentration in the cell: ca. 4 mMol)

Gram-negative bacteria

Biological barriers: e.g. bacterial cell walls

Picture source: N.L.Brown et al. Nature Rev. Biology (2015)

DOI:10.1038/nrmicro3480



Antifungals

Success

Ketoconazole, Fluconazole, Itraconazole, Clotrimazole, ...

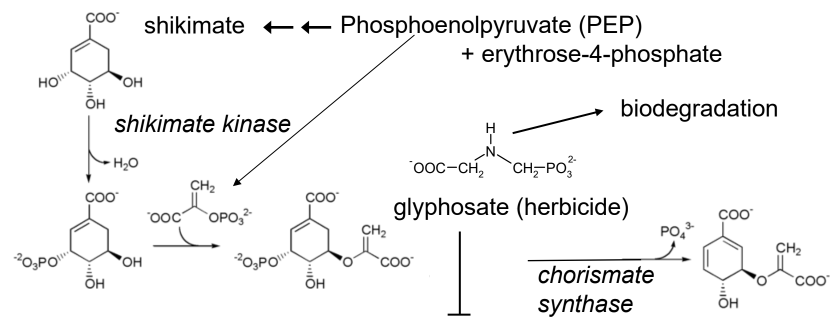
Mechanism of action: Inhibition of $14-\alpha$ -demethylase (CYP51) that is part of the biosynthesis pathway of ergosterol, which is an essential component of the fungal cell membrane

(in mammals: cholesterol).

Cons: Inhibition of Cytochromes causes hepatotoxicity (e.g. ketoconazole). Other conazoles are more specific. Development of resistances (overexpression of efflux proteins).

Failure (so far) (I)

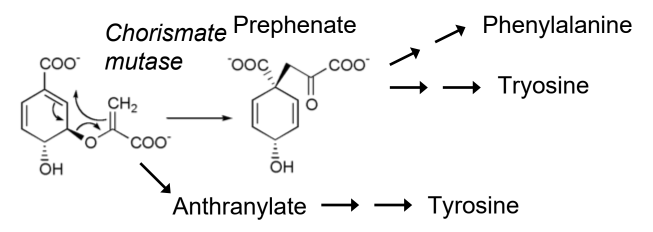
Antibacterial agents targeting enzymes of the Shikimate pathway (responsible for the synthesis of the amino acids Phe, Tyr, and Trp).



5-enolpyruvylshikimate-3-phosphate synthase (EPSPS) chorismate

Pro: Those enzymes are only found in plants, fungi, algae, and bacteria but not in mammals. Thus interference can be ruled out.

Failure (so far) (II)



Pathogens such as *Oxoplasma gondii*, *Plasmodium falciparum*, and *Cryptosporidium parvum* contain the Shikimate pathway and the seven enzymes involved.

Lit. C.W.Roberts et al. J.Infect.Dis. 185 (2002) Suppl.1:S25-36.

Con: Obviously the neccessary inhibitor concentration in the respective compartment could not be achieved.

For comparison:

The cellular level of phosphoenolpyruvate (PEP) is ca. 4 mMol

Complex Diseases

malaria is the tropical disease no.1

300-500 millionen infections per year causing 1-3 million fatalities

clinical symptoms:

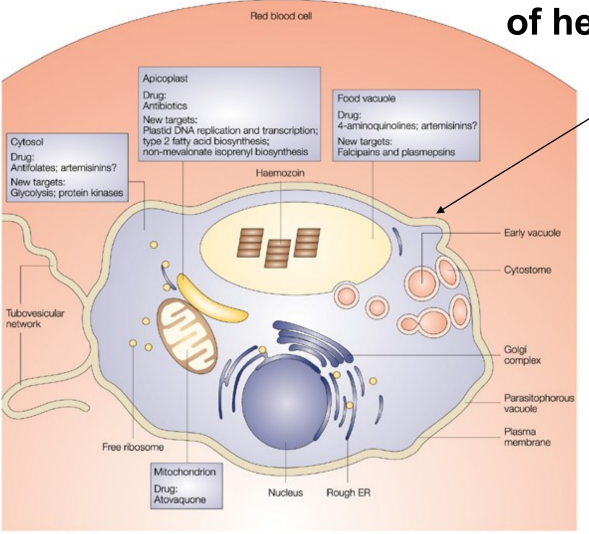
Strong fever, anemia, acidosis, multiple failure of organs



Due to the life cycle of the pathogen *Plasmodium flaciparum*, and the transmission by the *anopheles* fly, there are several starting points for control and therapy.

Lit. D.A.Fidock et al. Nature Rev. Drug Disc. 3 (2004) 509

malaria pathogens cause degradation of hemoglobin



Plasmodium falciparum trophozoite

Further pathogens in human:

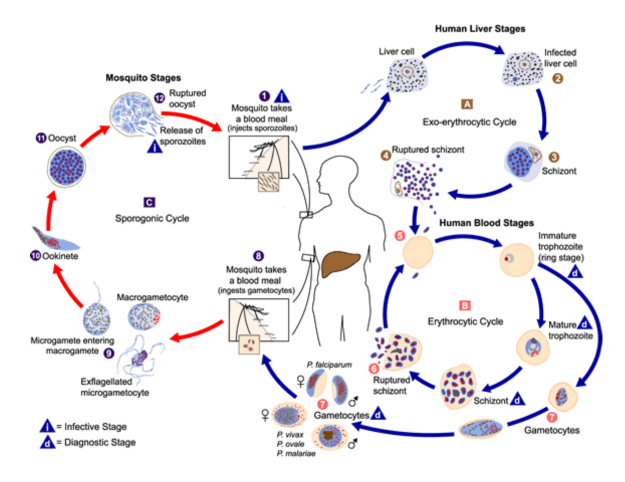
P. vivax

P. malariae

P. ovale

and about 56 more species of *Plasmodium*

Lifecylce of the malaria pathogens



source: http://www.dpd.cdc.gov/.../body_Malaria_page1.htm

Approaches to controlling (I)

1960-1980 exhaustive use of insecticides against the Anopheles fly with very good results by the use of DDT (dichloro-diphenyl-trichloroethane)

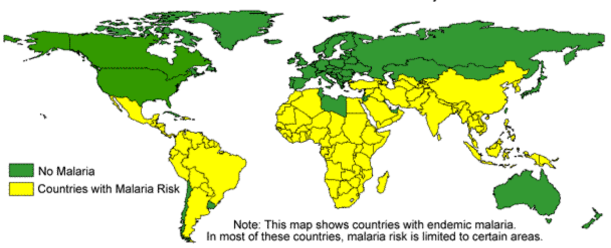
Disadvantages:

- Accumulation of DDT in the adipose tissue [Fettgewebe] of all creatures (mammals, birds, fish)
- DDT is biologically (almost) undegradable
- Metabolismus leads to a neurotransmitter-like substance (acts as contact insecticide!)

Increasing resistance to DDT has been observed

Distribution of Malaria (I)





Areas with risk of malaria

Distribution of malaria (II)

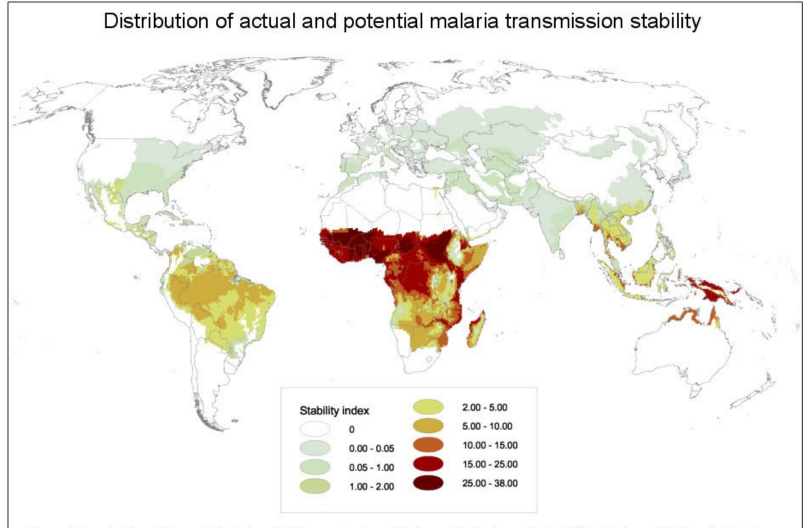
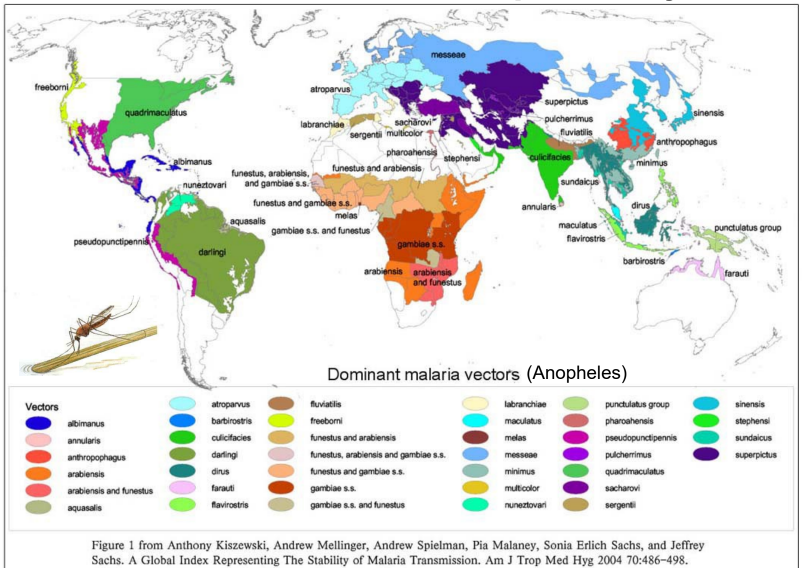


Figure 1 from Anthony Kiszewski, Andrew Mellinger, Andrew Spielman, Pia Malaney, Sonia Erlich Sachs, and Jeffrey Sachs. A Global Index Representing The Stability of Malaria Transmission. Am J Trop Med Hyg 2004 70:486-498.

Distribution of the Anopheles fly



Approaches to controlling (II)

chloroquine: since the late 1940's worldwide application at

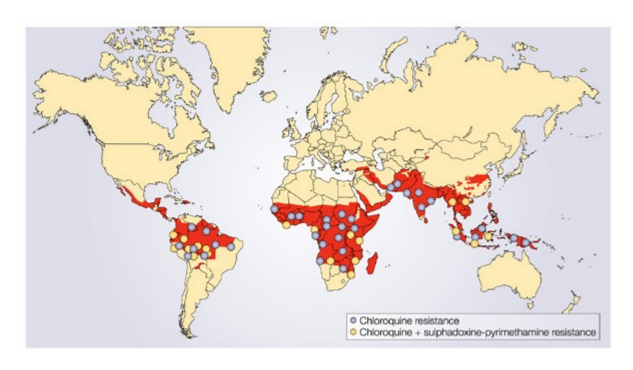
very low costs (0.2 US\$ per dose)

mode of action (still partly unclear): binds to HEM groups inhibition of the glutathion-S-transferase

sulfadoxine antibacterial

pyrimethamine blocks the dihydrofolate reductase respectively the dihydropterate synthetase

Resistance of the Anopheles fly



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red: areas with malaria

Approaches to contolling (III)

Alternatives to chloroquine and sulfadoxine/pyrimethamine amodiaquine respectively chlorproguanil/dapsone

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

Disadvantage: expected build up of resistances due to identical targets

Approaches to contolling (IV)

Profile for new drugs and chemoprophylaxis

- efficient, cheap
- effective against the more rare, but lethal *Plasmodium vivax*
- Avoiding of restistances by the use of combinations drugs (several targets at the same time)

Example for chemoprophylaxis: mefloquine (Lariam®)

Mode of action due to interaction with phospholipids (cell membrane, fatty acid synthesis)

Only very few adverse effects

Approaches to controlling (V)

Example for combination drugs: atovaquone (antiparasitic) together with an antibiotic

Drugs derived from natural compounds: artemisinin → artemether and artesunate (form cytotoxic radicals in the presence of HEM iron)

Disdavantage: rapid metabolization and thus short half life

New malaria targets (I)

Target location	Pathway/mechanism	Target molecule	Examples of therapies		References
			Existing therapies	New compounds	
Cytosol	Folate metabolism Glycolysis Protein synthesis Glutathione metabolism Signal transduction Unknown	Dihydrofolate reductase Dihydropteroate synthase Thymidylate synthase Lactate dehydrogenase Peptide deformylase Heat-shock protein 90 Glutathione reductase Protein kinases Ca ²⁺ -ATPase	Pyrimethamine, proguanil Sulphadoxine, dapsone Artemisinins	Chlorproguanil 5-fluoroorotate Gossypol derivatives Actinonin Geldanamycin Enzyme inhibitors Oxindole derivatives	82,83 84 85 86 87 88 89
Parasite membrane	Phospholipid synthesis Membrane transport	Choline transporter Unique channels Hexose transporter	Quinolines	G25 Dinucleoside dimers Hexose derivatives	71 91 92
Food vacuole	Haem polymerization Haemoglobin hydrolysis Free-radical generation	Haemozoin Plasmepsins Falcipains Unknown	Chloroquine Artemisinins	New quinolines Protease inhibitors Protease inhibitors New peroxides	93,94 95,96 97,98 99,100
Mitochondrion	Electron transport	Cytochrome c oxidoreductase	Atovaquone		101
Apicoplast	Protein synthesis DNA synthesis Transcription Type II fatty acid bio- synthesis Isoprenoid synthesis Protein farnesylation	Apicoplast ribosome DNA gyrase RNA polymerase FabH FabI/PfENR DOXP reductoisomerase Farnesyl transferase	Tetracyclines, clindamycin Quinolones Rifampin	Thiolactomycin Triclosan Fosmidomycin Peptidomimetics	102 29 32,33,103 30 25,104
Extracellular	Erythrocyte invasion	Subtilisin serine proteases		Protease inhibitors	97,105

DOXP, 1-deoxy-p-zylulose 5-phosphate; PfENR, Plasmodium falciparum enoyl-ACP reductase.

Lit. D.A.Fidock et al. Nature Rev. Drug Disc. 3 (2004) 509

New malaria targets (II)

- → Target identification on the gene level homolog enzymes of known diseases
- → Improvment of drugs that are already in use against other (infective) diseases:

dihydrofolate reductase → cancer

cysteine protease → osteoporosis

protein farnesyl transferase → cancer

protein synthesis → other parasites

vaccines: proteins that are expressed on the cell surface

→ sequencing of the *Plasmodium falciparum* genome

The first malaria vaccine Mosquirix (recombinant protein viruslike particle) has been endorsed by the WHO in October 2021.

18

(New) malaria drugs and targets (as of 2018)

Target	Drug
Fe(II)protoporphyrin IX	mefloquine
Fe(II)protoporphyrin IX	primaquine
Ferredoxin-NADPH reductase	tafenoquine (approved)
Dehydroorotate dehydrogenase	atovaquone
Dehydroorotate dehydrogenase	DSM265 (phase II)
Posphatidylinositol-4 kinase	MMV390048 (phase II)
Glutathione S-transferase	artesunate
Glutathione S-transferase	artefenomel (phase II)
Mitochondrial Enlogation Factor G	M7517 (phase I)
Ca ²⁺ transporting P-ATPase 4	cipargamin (phase II)

New malaria targets (III)

Sequencing of Plasmodium falciparum

25 Mb on 14 chromosomes, ca. 5000 genes6 Kb genome of the mitochondrium35 Kb circular DNA of the Apicoplast

Similar dimensions are also to be expected for *P. yoelii* and *P. vivax*.

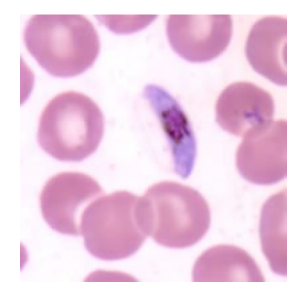
http://www.ncbi.nlm.hih.gov/Malaria/

http://plasmodb.org (annotated Plasmodium genome)



http://sites.huji.ac.il/malaria/ (contains EC numbers)

Lit. S.L.Hoffman et al. Nature 415 (2002) 702



Neglected Tropical Diseases (I)

Infections with pathogens prevalent in developping regions around the tropical belt of Africa, Asia, and America.

ascariasis, trichuriasis, necatoriasis, ancyclostomiasis infection by soil transmitted helmintics (worms)

Schistosomiasis (snail fever, bilharzia)

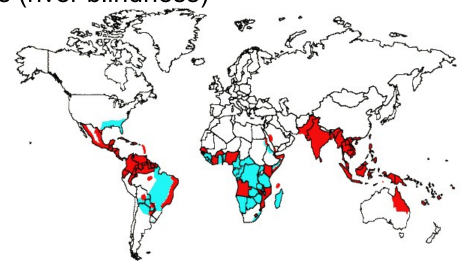
Trachoma and onchoceriasis (river blindness)

Leishmanias

Chagas disease

Leprosy

African Trypanosomnias (sleeping sickness)



The impact of this diseases in numbers is similar to that of malaria and tuberculosis

Neglected Tropical Diseases (II)

The World Health Organisation lists further diseases, such as

Cysticerosis (infection by the pork tapeworm)

Dengue / dengue haemorrhagic fever (virus transmitted by mosquitos)

Rabis [Tollwut] (viral)

Yaws (bacterial) a similar treponemal disease is syphillis

Snake bites

Tropical diseases with outbrakes in other areas due to transmission by mosquitos:

West Nile virus

Ross River fever

Complex diseases

obesity [Fettleibigkeit]



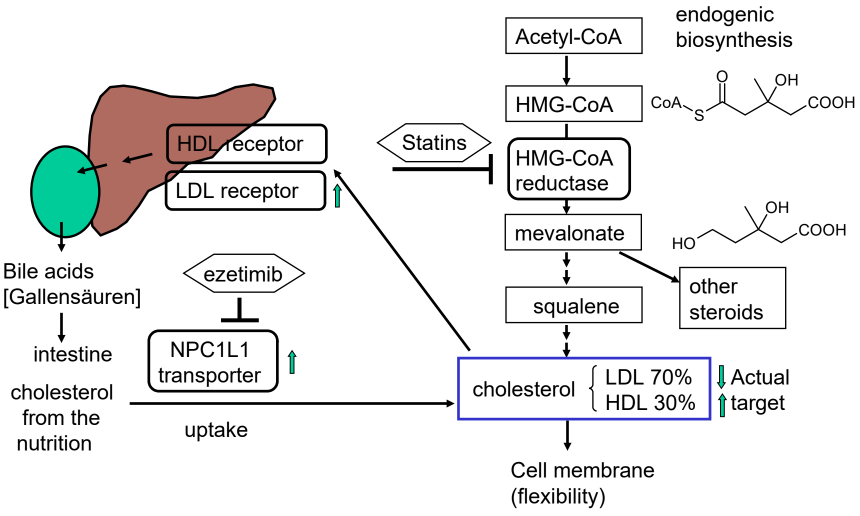
typical symptoms:

- excess weight
- increased levels of chlolesterol
 - → arteriosclerosis
- hypertension

increased cardiovascular risc

The connection to obesity has been established by the genetic lack of cholesterol receptors (hypercholesterolaemia) and especially cholesterol-rich nutrition in animal studies.

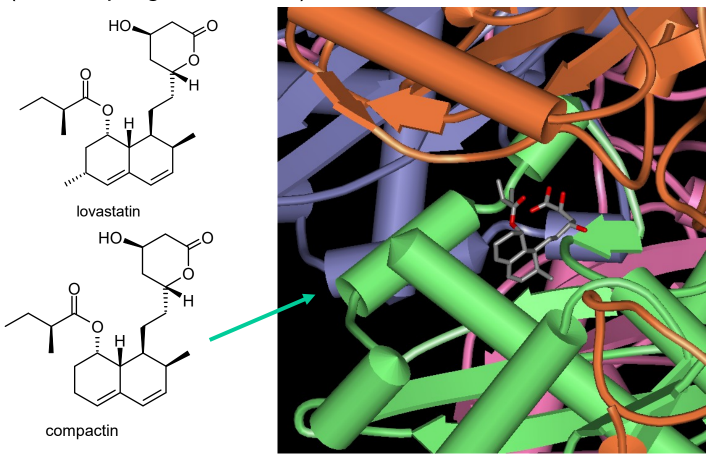
Regulation of the cholesterol pool



Lit. F.Rinninger & H.Greten *Dtsch. Ärztebl.* **102** (2005) A516 J.A.Tobert *Nature Rev. Drug Disc.* **2** (2003) 517

Inhibition of HMG-CoA reductase (I)

compactin (from *Penicillium citrinum*) and mevinolin (=lovastatin) (from *Aspergillus terreus*) were first found as inhibitors.



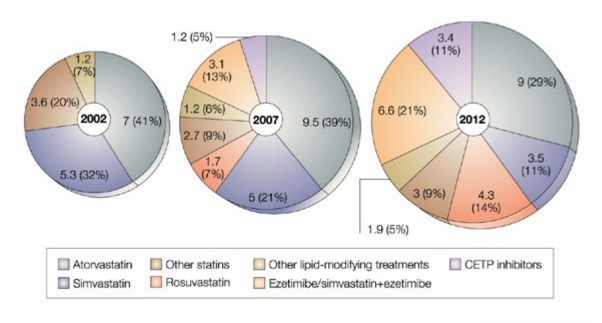
Lit. J.A.Tobert Nature Rev. Drug Disc. 2 (2003) 517

Inhibition of HMG-CoA reductase (II)

The actually effective substance is the metabolite

Sales potential of Statins

Market volume of cholesterol reducing agents



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Turnover in billion US\$ for USA, France, Germany, Italy, Spain, England and Japan, (market volume in %)
CEPT= cholesteryl ester transferase protein

Lit. J.Quirk et al. Nature Rev. Drug Disc. 2 (2003) 769

Further statins

cerivastatin (Bayer)

fluvastatin (Sandoz)

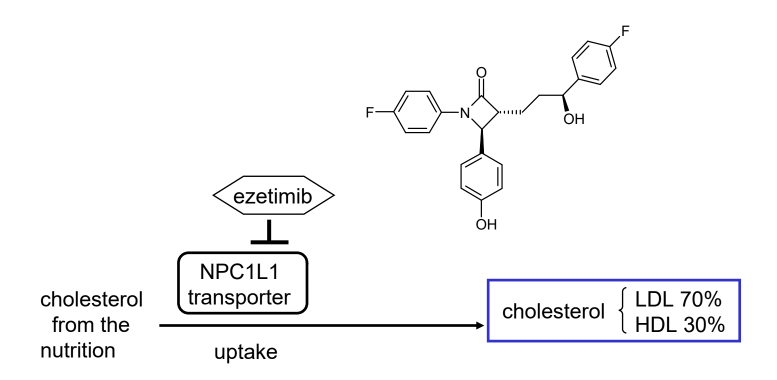
rosuvastatin (Astra-Zeneca)

atorvastatin (Warner-Lambert)

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Further lipid lowering agents (I)

ezetimib inhibits the cholesterol transporter



Lit. Van Heek *Brit.J.Pharmacol.* **129** (2000) 1748.

Further lipid lowering agents (II)

avasimibe inhibits the acetyl-coenzyme-A-cholesterol-acetyltransferase (ACAT-inhibitor)

Further lipid lowering agents (III)

competitive cholesterol analogs

cholesterol

Further lipid lowering agents (IV)

Bile acid sequestrants

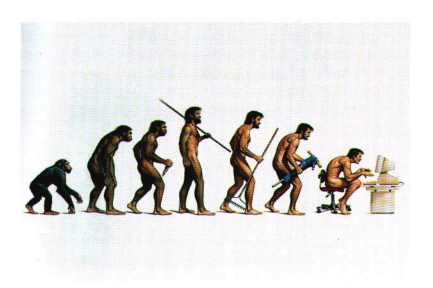
Polymers that are not absorbed from the intestine

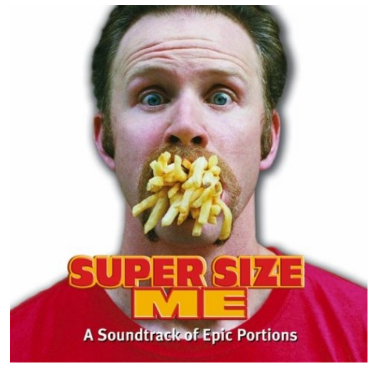
absorb cholesterol and bile acid and therefore prevent uptake of cholesterol

Opinion drugs vs. life style modification

"obesity is a form of depression in which the eating

is an antidepressant"

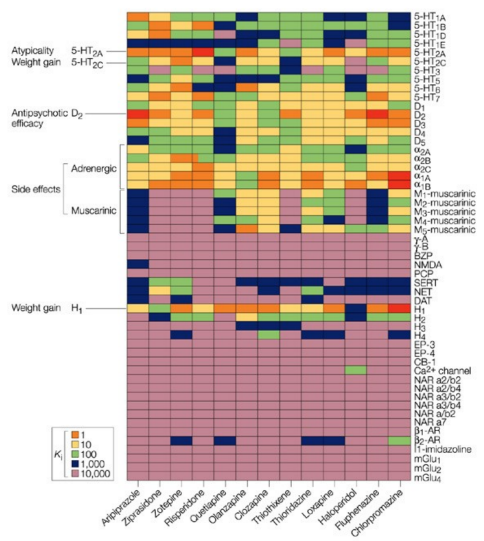




Fat storage is most efficient to preserve energy

Anorexic drugs (I)

Due to their complex affinity profile regarding a whole series of receptors ("dirty drugs") psychoactive drugs also modify the eating behaviour



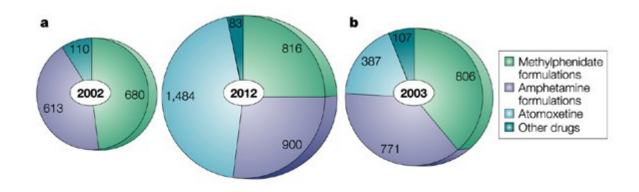
Lit. B.L.Roth et al.

Nature Rev. Drug Disc. 3 (2004) 353.

Anorexic drugs (II)

Prominent examples of psychoactive drugs with mit appetite suppressant (side-) effect:

methylphenidate (Ritalin®) ADHD atomexetine (Strattera®) [Aufmerksamkeitsdefizitsyndrome] fluoxetin (Prozac®)



Nature Reviews | Drug Discovery

Market volume of ADHD pharmaca in million US \$

Lit. M.Garland, P.Kirkpatrick Nature Rev. Drug Disc. 3 (2004) 385.

Prodrugs

Actually effective substance is the main metabolite of the drug

Example: ester cleavage

Irreversible inhibitor of cycloxygenase (COX)

Statins as HMG-CoA Reductase Inhibitors

The prodrug is a lactone whereas its metabolite is effective

Antiviral Nucleoside Analogs

Nucleosides missing the 3'-OH group cause disruption of the synthesis of a new DNA strain

Multi level prodrugs

Active uptake of α -Methyldopa-Phe by the dipeptide transporter

first pass metabolism
$$\alpha\text{-Methyldopa-Phe}$$

$$\alpha\text{-Methyldopa}$$

$$\alpha\text$$

 α -Methylnoreprinephrine

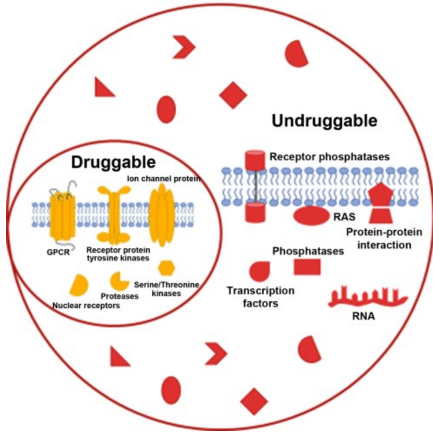
 α -Methylnoreprinephrine is an α_2 agonist

(false neurotransmitter)
ure Modern Methods in Drug Discovery WS21/22

Difficult and Undruggable Targets (1)

If there is no distinct binding pocket for typical small molecules, such targets are hard to inhibit:

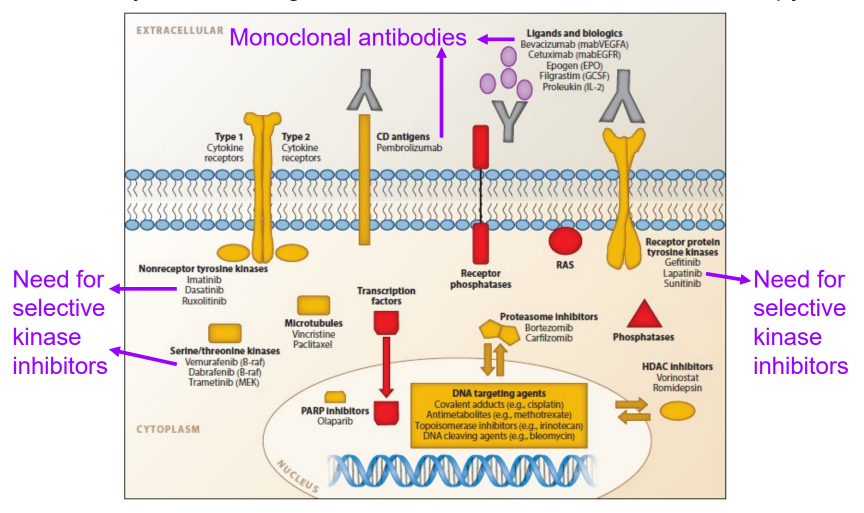
Transcription Factors
Receptor Phosphatases
(soluble) Phosphatases
(K)RAS
RNA
Protein-Protein-Interaction



Lit: J.Wang et al. Chin. J. Chem. 37 (2019) 501.

Difficult and Undruggable Targets (2)

Many of those targets are, however, crucial in cancer therapy



Lit: J.S.Lazo & E.R.Sharlow Annu.Rev.Pharmacol.Toxicol. Chem. 56 (2016) 23.

Lifestyle vs. Disease

The great challenges

- Virostatics
- Antibiotics (Zn-β-lactamases, malaria)
- Anticancer drugs
- Neurodegenerative diseases (Antidementia, Alzheimer)
- Diabetes type 2
- civilization diseases (obesity, ADHD)?





Lifestyle vs. Disease (II)

The top selling drugs during recent times (selection):

adalimumab* (arthritis)

apixaban factor Xa-inhibitor (anti-coagulant)

pregabalin calcium channels (epilepsy)

lenalidomide antitumor/apoptisis

nivolumab* (oncology, various cancers)

pembrolizumab* (cancer immunotherapy)

etancercept* (rheumatoid arthritis)

trastuzumab* (breast cancer) bevacizumab* (colon cancer)

rituximab* (autoimmune diseases, cancer)

sofosbuvir antiviral nucleoside

fluticasone anti-inflammatory/corticosteroid

rosuvastatin HMG-CoA reductase

Source: wikipedia (2019)

^{*} monoclonal antibody

Lifestyle vs. Disease (III)

Most "blockbuster" drugs were not predicted by analysts of the marketing departements:

indication

tamoxifen breast cancer

captopril hypertension

cimetidine gastric ulcers [Geschwulstbildung

im Magen]

fluoxetine (Prozac™) depression

atorvastatin (Lipitor™) hyperlipidaemia, obesity

Lit: J.Knowles & G.Gromo Nat.Rev.Drug.Discov. 2 (2003) 63.

Lifestyle vs. Disease (IV)

Innovative new drugs that have emerged (source: Hugo Kubinyi)

2006	Deferasirox	iron chelator (thalassemia)
2003	Roflumilast	PDE-4 inhibitor (asthma)
2002	Ezetimib	cholesterol uptake inhibitor
2001	Imatinib	leucemia (tyrosine kinase inhibitor)
2001	Fondaparinux	thrombosis (antagonist)
1999	Zanamivir	influenza (viral neuraminase inhibitor)
1999	Amprenavir	HIV (protease inhibitor)
1999	Celecoxib	arthritis (COX-2 inhibitor)
1998	Sildenafil	erectile dysfunction (PDE-5 inhibitor)
1998	Orlistat	obesity (pancreas pipase inhibitor)
1997	Sibutramine	obesity (GPCR inhibitor)
1997	Finasteride	prostata (steroidreductase inhibitor)
1997	Nelfinavir	HIV (protease inhibitor)
1996	Indinavir	HIV (protease inhibitor)
1996	Nevirapin	HIV (reverse transcriptase inhibitor)
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Lifestyle vs. Disease (V)

Innovative new drugs from 1982-1996: (source: Hugo Kubinyi)

1996	Meloxicam	arthritis (COX-2 inhibitor)
1995	Dorzolamine	glaucoma (carboanhydrase inhibitor)
1995	Losartan	hypertension (GPCR antagonist)
1994	Famciclovir	herpes (DNA polymerase inhibitor)
1993	Risperidon	psychose (D ₂ / 5HT ₂ antagonist)
1991	Sumatriptan	migraine (5HT ₁ rezeptor antagonist)
1990	Ondansetron	antiemetic (5HT ₃ antagonist)
1988	Omeprazole	gastric ulcers (proton pump inhibitor)
1987	Lovastatin	cholesterol (biosynthesis inhibitor)
1986	Artemisinin	anti-malarial (natural compound)
1985	Fluoxetine	depression (5HT inhibitor)
1985	Mefloquine	anti-malarial
1984	Enalapril	hypertension (ACE inhibitor)
1983	Cyclosporin A	immunosupressant
1982	Ranitidine	gastric ulcers (H ₂ antagonist)
11th lecture	Modern Methods in Drug Discovery WS21/22	

Lifestyle vs. Disease (VI)

How are innovative drugs defined?

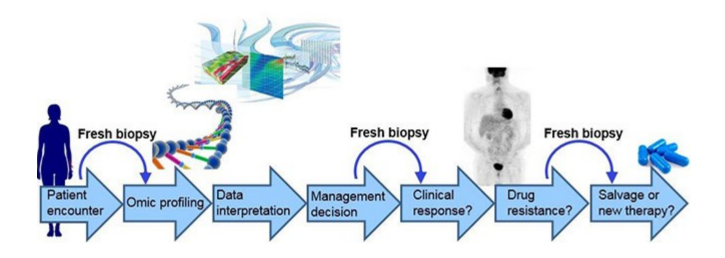
- improved mode of action (selectivity)
- improved ADMET profile
- Improved administration (e.g. oral instead of intravenous)
- pro-drugs
- new targets

personalized medicine

Variable metabolic content and predisposition (Genotyping)

Avoiding rare, complicated adverse effects (in part already used in the clinic)

Will the necessary financial effort of screening and of clinical studies limit the genetic pool to inhibitants of wealthy nations?



picture source: www.dana-farber.org

Resume

The available knowledge on the human genome and the present SNPs in it allow two approaches:

- 1. Finding new targets (either on the genome, the mRNA, or the protein level)
- 2. pharmacogenomic methods will lead to personalized medicine (which drug and at what dosage), esp. for long term application of certain drugs (hypertension, analgesics, anti-psychotics) and those that possess a narrow therapeutic band width (cardiotonics, antineoplastics)