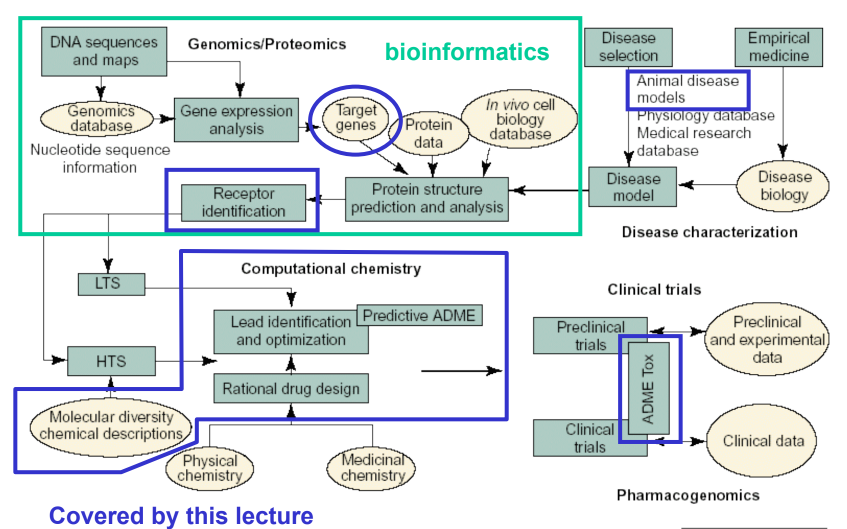
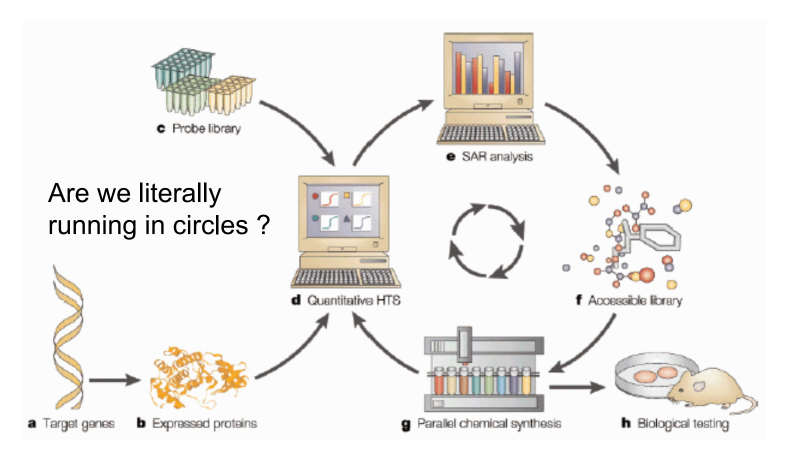
Current Trends



Drug Discovery Today

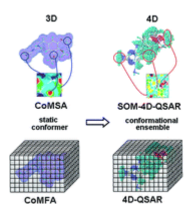
Cycle of optimization in the drug discovery pipeline



Source: D.K. Agrafiotis et al. Nature.Rev.Drug.Discov. 1 (2002) 337.

4D and 5D QSAR

- 3D QSAR: Information from the 3D structure is used
- → 3D descriptors, pharmacophore models
- 4D and 5D QSAR: multiple conformations
- → use of multiple docking results for one compound



Lit: M. Dobler et al. *Helvetica Chim. Acta* **86** (2003) 1554. A. Bak et al. *RCS Adv.* **6** (2016) 76183.

multiple conformations upon docking (I)

The binding pocket of many cytochrome P450 enzymes (esp. CYP3A4 and CYP2D6) is large enough to accomodate the same substrate in different orientations, which leads to different products.

The reactivity of a certain spot of the molecule can be estimated by quantum chemical calculations.

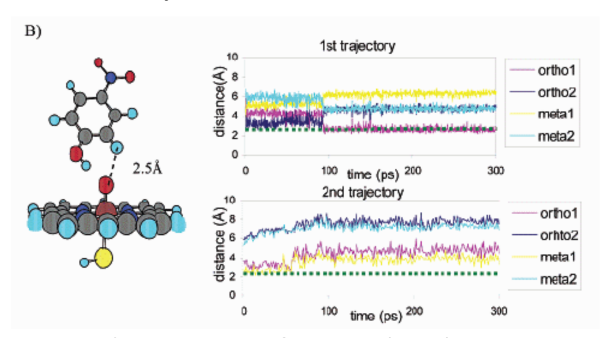
Lit: M. de Groot et al. *J.Med.Chem.* **42** (1999) 4062 S.B. Singh et al. *J.Med.Chem.* **46** (2003) 1330

multiple conformations upon docking (II)

Besides information about the reactivity at a certain spot of the molecule, also the propability of the according binding position in the enzyme is required.

Can be obtained from statistical analysis (clustering) of

- a large number of docking results, or by
- molecular dynamics simulations



Lit: Park & Harris *J.Med.Chem.* **46** (2003) 1645

Drug / Non-Drug Separation (1)

Is it possible to predict the potential suitability of a compound from typical properties of drugs?

approaches:

Reckognition of typical properties in data bases that (almost) exclusively contain drugs

For example:

- World Drug Index (WDI)
- Comprehensive Medicinal Chemistry (CMC)
- MACCS-II Drug Report (MDDR)
- Merck Index (filtering required)
- Drugbank
- ChEMBL (filtering required, e.g. clinical phase)

Drug / Non-Drug Separation (2)

Previous data base analyses:

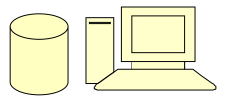
1997 Christopher Lipinski's rule of 5 (Pfizer)

Orally administered drugs typically have



molecular weight < 500 ClogP < 5

less than 5 hydrogen-bond donors (O-H, N-H) less than 10 hydrogen-bond acceptors (N, O, S)



2000 Tudor Oprea (AstraZeneca)

Typical drugs (70% of all) have

less than 3 hydrogen-bond donors between 2 and 9 hydrogen-bond acceptors between 2 and 9 rotatable bonds between 1 and 4 rings

Lipinski's rule of 5 refers to oral bioavailability but not neccessarily drug-likeness!

Drug / Non-Drug Separation (3)

1999 Ghose, Viswanadhan & Wendoloski

Analysis of the Comprehensive Medicinal Chemistry database:

80% of all drugs have

$$-0.4 < logP < 5.6$$

20 < number of atoms < 70

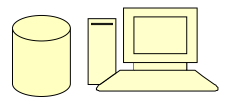
40 < molar refractivity < 130



30 < number of atoms < 55

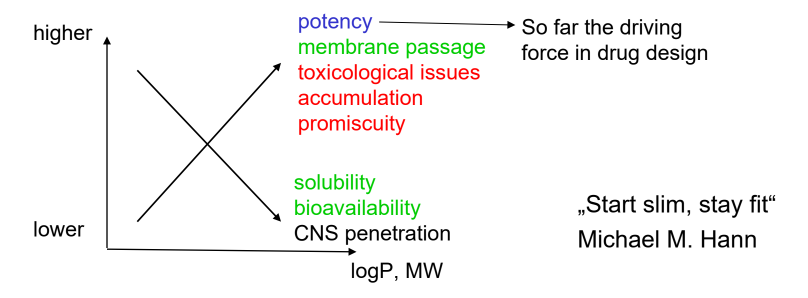
70 < molar refractivity < 110

Lit: A. Ghose et al. J. Comb. Chem. 1 (1999) 55-68.



Drug / Non-Drug Separation (4)

Even tighter restrictions required to avoid adverse effects? Molecular weight < 400 and ClogP < 4 (GSK 4/400 rule)

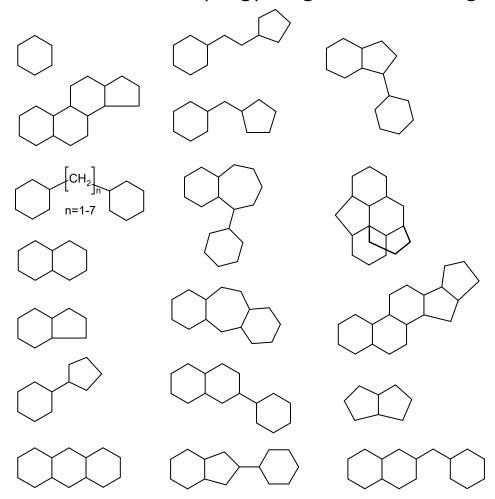


Find smallest crucial parts of molecules → fragments

Lit: M.M. Hann "Molecular Obesity, Potency and Other Addictions in Drug Discovery" *Med.Chem.Commun.* **2** (2011) 349-355.

Drug / Non-Drug Separation (5)

The most common (ring) fragments of drugs



Drug / Non-Drug Separation (6)

Rare appearance of certain fragment or side chains does not necessarily mean that it is unsuitable or negligible.

Such fragments can rather

- be difficult to synthesize, or
- be newly introduced, or

possess unsuitable properties

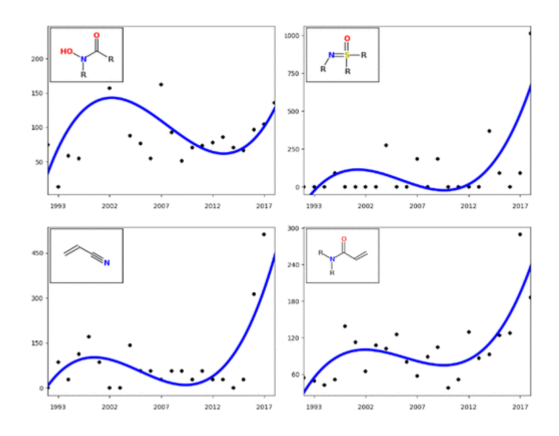
rare ketene fragment

difficult dioxygen bridge

of lactam

Use of fragments over time

While some functional groups seem to indispensable, a number of new and increasingly used fragments have been identified from the ChEMBL database:



Lit. P. Ertl et al. J. Med. Chem. 63 (2020) 8408.

Drug / Non-Drug Separation (7)

Examples of groups that possess wellknown unsuitable properties

Such groups should not be present in clinical drugs, but may be important during synthesis

Lit: D.R.Flower, *J.Mol.Graph.Model.* **16** (1998) 239. M.Hann et al. *J.Chem.Inf.Comput.Sci.* **39** (1999) 897.

Reactive Groups for Irreversible Inhibitors

Covalent binding of inhibitors to enzymes renders these to become unfunctional.

Typical reactive groups are electrophilic, such as acrylamides and α,β -unsaturated carbonyls that bind to noncatalytic cysteine

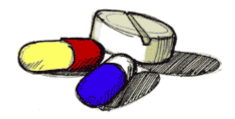
Lit: M.Gehringer & S.A.Laufer Chem. Chem. Biol. 62 (2019) 5673.

Drug / Non-Drug Separation (8)

Further approach:

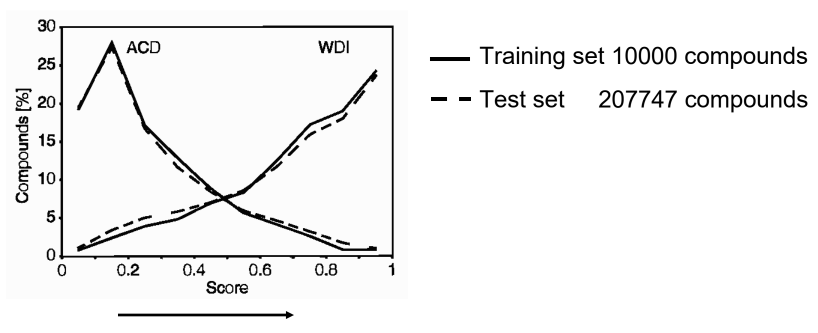
Comparison of compounds in a data base containing solely drugs (World Drug Index) to substances from a data base that predominately consists of non-pharmaceutical compounds (Available Chemical Directory).





Drug / Non-Drug Separation (9)

Classification of compounds according to their atom types using a neural net

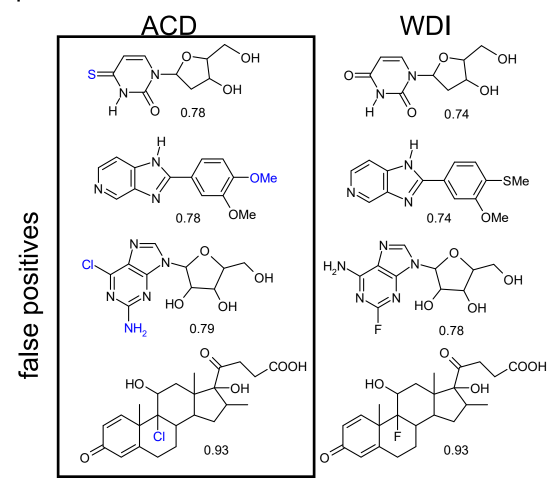


Increasing drug-likeness, but where is a reasonable threshold of the score?

Lit: J. Sadowski & H. Kubinyi *J.Med.Chem.* **41** (1998) 3325.

Drug / Non-Drug Separation (10)

Compounds for which a high drug-likeness score was predicted:



Lit: J. Sadowski & H. Kubinyi *J.Med.Chem.* **41** (1998) 3325.

Drug / Non-Drug Separation (11)

Classification of compounds using their ISIS *fingerprint* (set of 73 descriptors which indicate the presence of structural and topological features, and encode chemical properties)

→ Allow comparison of the compounds by their similarity using the Tanimoto index.

These 73 binary descriptors were used as input layer of a neural net, which was trained with compounds from drug data bases and non-drugs from the ACD.

result: about 80% of all compounds were classified correctly.

Lit: Ajay et al. J. Med. Chem. 41 (1998) 3314.

Drug / Non-Drug Separation (12)

Classification of compounds according to atom types that represent so-called *pharmacophoric points*:

Among these functional groups are

preselection: a compound is potentially drug-like, if it contains at least one ring and between 2 and 7 of such functional groups.

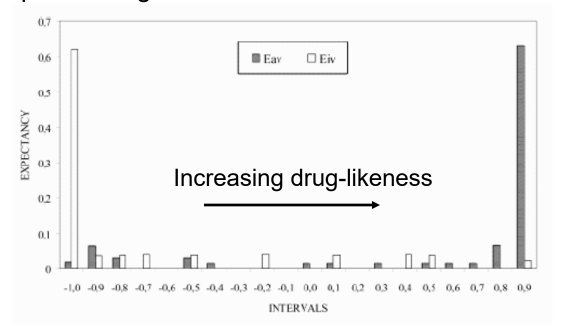
The atoms of the molecule are classified according to the affiliation to one of these *pharamacophoric points* and used as input layer of a neural net.

Here again compounds of the ACD were compared to drug data bases.

Lit: J.Muegge et al. J.Med.Chem. 44 (2001) 1841.

Drug / Non-Drug Separation (13)

Classification of compounds according to topological descriptors using a neural net.



680 compounds of the Merck Index, of which about 76 % were classified correct.

Lit: M.Murcia-Soler et al. J.Chem.Inf.Comput.Sci. 43 (2003) 1688.

Drug / Non-Drug Separation (14)

Classification of compounds using a *decision tree*. Used were atom types that represent certain functional groups.

Advantages of a decision tree compared to a neural net:

The criteria for classification at each branching point can be traced easily and a corresponding error can be assigned.

results:

- ¾ of all compounds can be assigned based on the presence of 6 chemical groups.
- Non-drugs typically contain not enough of these functional groups

Lit: M.Wagner et al. J.Chem.Inf.Comput.Sci. 40 (2001) 280.

Drug / Non-Drug Separation (15)

Preliminary resume:

Neither the presence of atom types, nor that of (sub-) structure fragments or functional groups, allows to classify a substance precisely as drug-like (> 95% accuracy) Seemingly an (even) larger variety of descriptors, e.g. those that account for electronic properties are required.

→ use of quantum chemical descriptors?

Lit: N.Schneider et al. *J.Chem.Inf.Model.* **48** (2008) 613. M.C.Hutter *Curr.Med.Chem.* **16** (2009) 189.

Drug / Non-Drug Separation (16)

Principal component analysis (PCA) of 26 descriptors of compounds from the Maybridge data base yielded the numerical value of the 3rd principal component as most significant separation criteria.

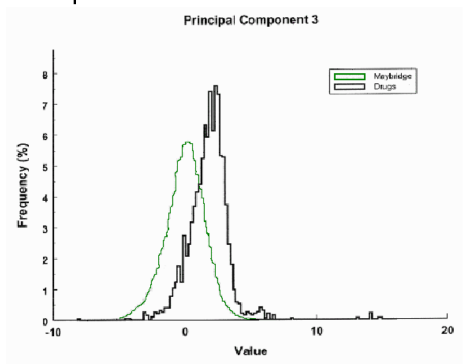
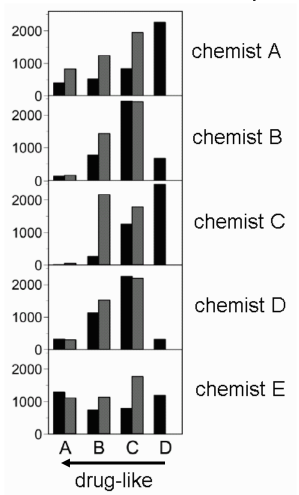


Figure 3. Frequency histogram for PC number 3 (MEP-) for drugs and nondrugs.

Drug / Non-Drug Separation (17)

Classification of compounds based on chemical intuition



3980 compounds were classified by 5 chemists according to their drug-likeness and the according synthesic efford

Table 2. Correlation Coefficient between Scores Assigned by Each Chemist on Drug-Likeness

	chemist A	chemist B	chemist C	chemist D	chemist E
chemist A chemist B chemist C chemist D chemist E	1.00	0.55 1.00	0.63 0.51 1.00	0.57 0.50 0.52 1.00	0.58 0.50 0.54 0.54 1.00

Table 3. Correlation Coefficient between Scores Assigned by Each Chemist on Ease of Synthesis

	chemist A	chemist B	chemist C	chemist D	chemist E
chemist A chemist B chemist C chemist D chemist E	1.00	0.50 1.00	0.40 0.42 1.00	0.40 0.47 0.40 1.00	0.56 0.52 0.48 0.48 1.00

Lit: Y.Takaoka et al. J.Chem.Inf.Comput.Sci. 43 (2003) 1269

Drug / Non-Drug Separation (18)

try yourselves!

Classify these compouds into drug or non-drug

Compare your results to that of the property prediction module at http://www.organic-chemistry.org/prog/peo/index.html

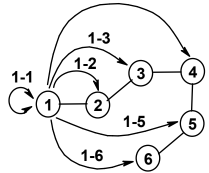
Drug / Non-Drug Separation (19)

Back to the basics:

So far it has been only assumed that there is an unequal feature distribution between drugs and non-drugs. How can we statistically prove this assumption?

Idea: Certain combinations of atom types are found with a different frequency among drugs compared to non-drugs

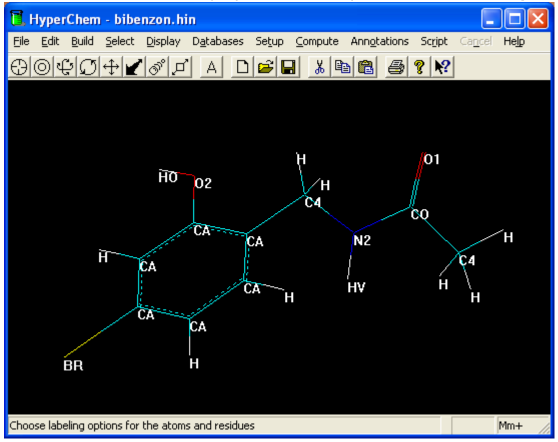
- 1-1 Interaction: the atom itself
- 1-2 Interaction: bond between two atoms 1 and 2
- 1-3 Interaction: angle between atoms 1 and 3
- 1-4 Interaction: dihedral angle between atoms 1 and 4 1-4
- 1-5 Interaction: between atoms 1 and 5



Drug / Non-Drug Separation (20)

What atom types and how many should be used?

Atom types should account for the chemical diversity Thus, elements only (C, N, O,..) are not enough



Here, atom types from the MM+ force field are used (total of 47)

Drug / Non-Drug Separation (21)

When is an atom pair combination i-j statistically overrepresented?

ightarrow If its frequency q_{ij} is higher than that by chance (= relative probability S')



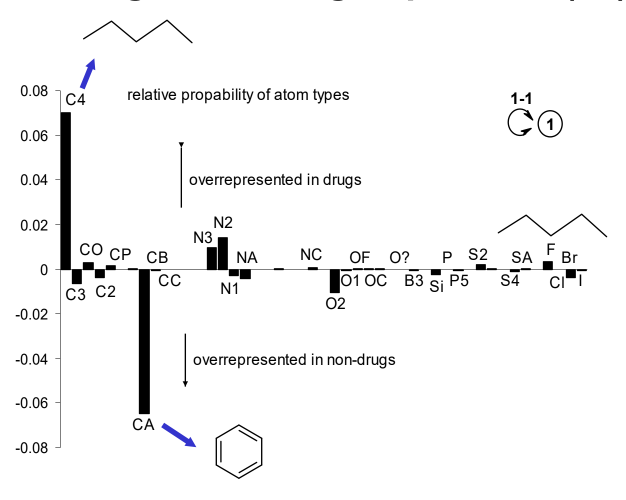
$$S'_{ij} = \frac{q_{ij}}{p_i \cdot p_j}$$

where p_i is the individual frequency of an atom of type i

For better handling we use the logarithmic value = log odds score

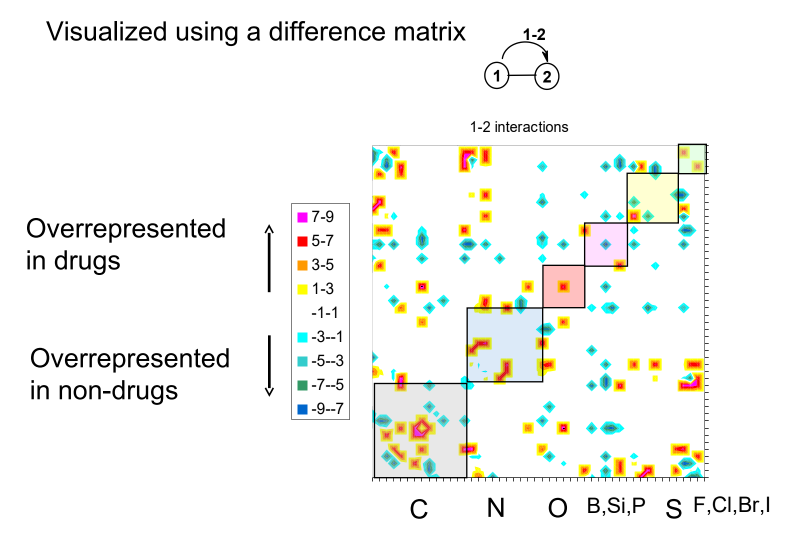
$$S_{ij} = \ln \frac{q_{ij}}{p_i \cdot p_j}$$
 $\begin{cases} >0 \text{ overrepresented} \\ <0 \text{ underrepresented} \end{cases}$

Drug / Non-Drug Separation (22)



Distribution of atom types (1-1 interaction) alone is not sufficient

Drug / Non-Drug Separation (23)



Similar to amino acid exchange matrix!

Drug / Non-Drug Separation (24)

But how to calculated the drug-*likeliness* from the atom type distribution?

Simply add up corresponding matrix entries and divide by the number of occuring atom pairs in the molecule:

Drug-likeliness score L

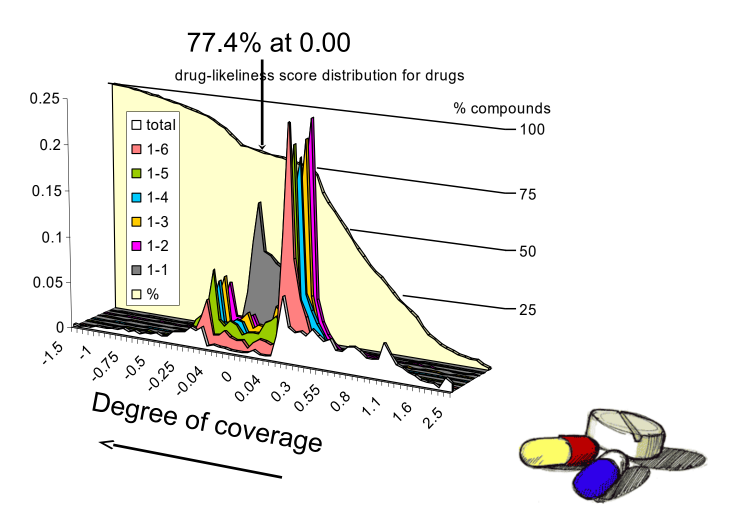
$$L = \sum_{1}^{6} \frac{\sum S_{ij}}{M}$$
 { >0 drug-like <0 non-drug-like



Timing:

Less than 5 minutes computing the difference matrices and scores for 4083 compounds

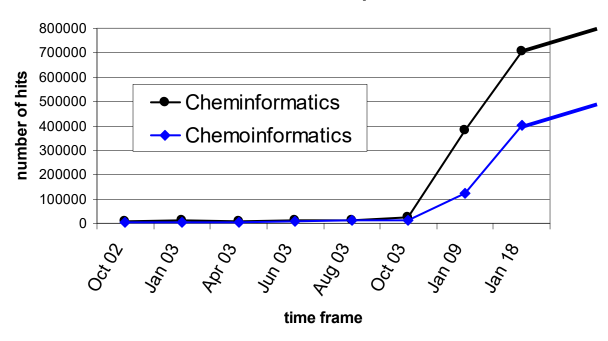
Drug / Non-Drug Separation (25)



Lit: M.C.Hutter *J.Chem.Inf.Model.* **47** (2007) 186-194.

Cheminformatics or Chemoinformatics?

Which term is more accepted?



Data source: http://www.molinspiration.com

http://www.google.de

Doping (I)

Illicit use of substances to achieve an increase in performance (in sport)

→ A definition is difficult, since there must be a causative link between cause and action, similar to drugs



According substances are put together in doping lists by national and international sport committees (e.g. international olympic committee IOC) based on medical knowledge.

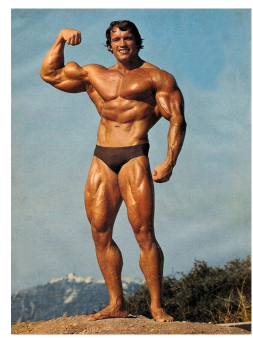
Doping list (I)

Illicit substance groups

 anabolic steroids (anabolics) lead to an increased building up of muscles

naturally in the body occurring steroids such as testosterone, as well as totally artificial steroids e.g. tetrahydrogestrinone (THG) Partly not even allowed for fattening of porks!

- antiestrogenic compounds aromatase inhibitors tamoxifen, etc.
- hormons and related drugs erythropoietin (EPO): increased production of red blood cells
- insulin and insulin-like growth factors
- → substances that increase the oxygen transport capacity of the blood



Doping list (II)

Banned substance groups

- Stimulants increase the short term motivation amphetamines (cardiovasuclar and addiction risks) caffeine (until 2004 with a limit), due to newer results no limits any more
- narcotics and β-blocker show a calming down effect (pain reducing)
 (boxing, archery [Sportschießen])



Doping list (III)

- glucocorticoides (heart and circulation function)
- cannaboids
 hashish, marihuana

Mascing substances

- diuretica (increased elimination, reduction of body weight)
- inhibitors of the steroid- α -reductase (finasterid)
- plasmaexpanders (albumin, dextran) cause reduced drug concentration in the serum



Doping list (IV)

Substances with limits in certain sports

- alcohol (billard, tighter limits e.g. in racing)
- β-blocker (sports that require increased concentration)





 gene doping modification on the genetic level to increase performance (nuclear receptors, mRNA, gene silencing) feasibility, analytical proof?

Doping (V)

Anti-Doping-Kontrolle

Anti-Doping-Kontrolle

Contrôle Antidopage

Controle Antidopage

Doping lists are not comprehensive, which means that all similar compounds and those with a similar effect are included implicitly.

→ possibly not formulated precise enough for legal actions

Doping tests

Mainly urine samples, blood samples less frequent problems: limits for metabolites of naturally occuring compounds, e.g. of testosterone and hematocrite → synthetic steroids show a different ¹²C/¹³C ratio traceability of certain compounds (EPO) new and formerly unknown compounds (e.g. THG) → give rise to unusual patterns in Mass-spectroscopy

Doping (VI)

Why doping tests? fairness, (self-)protection of the athletes

Risks of doping

- anabolic steroids: liver damage
- stimulants: addiction, lethal exhaustion
- known common adverse effects

Many drugs that are included in doping lists can administered with exception permits.

E.g. steroidal anti-inflammatories, anti-asthmatics