## More about QSAR...

QSAR equations form a quantitative connection between chemical structure and (biological) activity.

$$
\log (1 / C)=k_{1} \cdot P_{1}+k_{2} \cdot P_{2}+\ldots+k_{n} \cdot P_{n}
$$

Problems:

- Which and how many descriptors to use?
- How reliable are the predictions (applicability domain)?
- How to test/validate QSAR equations (continued from lecture 5)


## Setting up and testing QSAR equations



## Evaluating QSAR equations (1)

The most important statistical measures to evaluate QSAR equations are (preferred values given in parenthesis):
Correlation coefficient $r$ (in squared from $r^{2}>0.75$ )
Standard deviation se (small as possible, se < 0.4 units)
Fisher value $F$ (level of statistical significance. Also a measure for the portability of the QSAR equation onto another set of data. Should be high, but decreases with increasing number of used variables/descriptors). Therefore only comparable for QSAR equations containing the same number of descriptors
$t$-test to derive the
probability value $p$ of a single variable/descriptor.
Is a measure for coincidental correlation
$p<0.05=95 \%$ significance
$p<0.01=99 \%$
$\mathrm{p}<0.001=99.9 \%$
$\mathrm{p}<0.0001=99.99 \%$

## Evaluating QSAR equations (2)

Example output from OpenStat:


Constant $=0.621$
$\log (1 / C)=-0.517 \cdot$ hbdon $-21.360 \cdot$ dipdens $+0.020 \cdot$ chbba +0.621
Lit: William „Bill" G. Miller, OpenStat Reference Handbook

## Evaluating QSAR equations (3)

A plot tells more than numbers:


Shape of curve indicates non-linear correlation
Source: H. Kubinyi, Lectures of the drug design course http://www.kubinyi.de/index-d.html

## Evaluating QSAR equations (4)

Examples where statistical measures between training set and test set strongly deviate:

$$
\begin{array}{ll}
\text { Training set } & n=15, r^{2}=0.91, s e=0.27 \quad(5 \text { descriptors used) } \\
\text { Test set } & n=5, r^{2}=0.69, s e=0.42
\end{array}
$$

Obvious reason: too many descriptors used in QSAR eq. Therefore the training set becomes overfitted, correlation breaks down for the test set. $\rightarrow$ Limit number of used descriptors in the QSAR equation to three.

Training set $\quad n=26, r^{2}=0.88$, se=$=0.32, F=110.7$ ( 3 descriptors used)
Test set $\quad n=7, r^{2}=0.75$, $\mathrm{se}=0.38, \mathrm{~F}=66.5$
Possible reason: Compounds in the test set are quite different compared to those in the training set.
$\rightarrow$ Check compounds (and descriptor ranges) for similarity, redo compound selection for training and test set e.g. using cluster analysis

## Evaluating QSAR equations (5)

(Simple) $\boldsymbol{k}$-fold cross validation:
Partition your data set that consists of $N$ data points into $k$ subsets ( $k<N$ ).


Generate $k$ QSAR equations using a subset as test set and the remaining $k-1$ subsets as training set respectively. This gives you an average error from the $k$ QSAR equations.

In practise $k=5$ or $k=10$ has shown to be reasonable (refers to 5 -fold or 10 -fold cross validation, respectively)

## Evaluating QSAR equations (6)

## Leave one out cross validation:

Partition your data set that consists of $N$ data points into $k$ subsets ( $k=N$ ).


Disadvantages:

- Computationally expensive
- Partitioning into training and test set is more or less by random, thus the resulting (average) error can be way off in extreme cases.

Solution: (feature) distribution within the training and test sets should be identical or similar

## Evaluating QSAR equations (7)

## Stratified cross validation:

Same as $k$-fold cross validation but each of the $k$ subsets has a similar (feature) distribution as the training set.

$k$ times

The resulting average error is thus more prone against errors due to equal distribution of features between training set and the test sets.

## Evaluating QSAR equations (8)


alternative
Cross-validation and leave one out (LOO) schemes

Leaving out one or more descriptors from the derived equation results in the crossvalidated correlation coefficient $\mathrm{q}^{2}$.
This value is of course lower than the original $r^{2}$. $q^{2}$ being much lower than $r^{2}$ indicates problems...

## Evaluating QSAR equations (9)

Problems associated with $q^{2}$ and leave one out (LOO)
$\rightarrow$ There is no correlation between $\mathrm{q}^{2}$ and test set predictivity, $q^{2}$ is related to $r^{2}$ of the training set



Kubinyi's paradoxon: Most $r^{2}$ of test sets are higher than $q^{2}$ of the corresponding training sets [due to manual selection?]
Lit: A.M.Doweyko J.Comput.-Aided Mol.Des. 22 (2008) 81-89.

## Evaluating QSAR equations (10)

One of most reliable ways to test the performance of a QSAR equation is to apply an external test set.
$\rightarrow$ partition your complete set of data into training set (2/3) and test set ( $1 / 3$ of all compounds, idealy)
Compounds of the test set should be representative (confers to a 1 -fold stratified cross validation)
$\rightarrow$ Cluster analysis using the descriptor values of each compound plus their activities.
$\rightarrow$ Use cluster centroids as test set and the remaining
compounds for the training set (these account for the diversity)


## Evaluating QSAR equations (11)



Compounds of the test set must cover the same activity range as those of the training set

- Training set
- Test set


## Evaluating QSAR equations (12)

Estimating the error (range) of predicted values is difficult. Approaches to give a confidence range (as in statistics) or determining the applicability domain of the model:

- Distance based: similar to $k$-nearest neighbor; where is the predicted compound located in the descriptor space?
Close to one group or rather in between clusters?
Large training sets can be split into a further calibration set that is used for estimating the error of unseen data based on their similarity.
- Are there consistent outliers in the data set?
$\rightarrow$ These are either too dissimilar or are likely to contain experimental errors.

Lit: K. Roy et al. Chemomet. Intell. Lab. Sys. 145 (2015) 22-29.

## Interpretation of QSAR equations

The kind of applied variables/descriptors should enable us to

- draw conclusions about the underlying physico-chemical processes
- derive guidelines for the design of new molecules by interpolation

$$
\log \left(1 / K_{i}\right)=\oplus 1.049 \cdot n_{\text {fluorine }} \in @ .843 \cdot n_{O H}+5.768
$$

Higher affinity requires more fluorine, less OH groups
Some descriptors give information about the biological mode of action:

- A dependence of $(\log P)^{2}$ indicates a transport process of the drug to its receptor.
- Dependence from $\mathrm{E}_{\text {Luмо }}$ or $\mathrm{E}_{\text {номо }}$ indicates a chemical reaction


## Evaluating QSAR equations (13)

Reduce the number of available descriptors before performing a regression analysis:

- More descriptors cause longer run times
- More descriptors raise the likelihood of accidental correlation (see also slides further below)
- Descriptors might be correlated to each other and thus do not provide more information
- Can you interpret what your descriptors mean?
$\rightarrow$ get rid of the "garbage"



## Correlation of descriptors

Other approaches to handle correlated descriptors and/or a wealth of descriptors:

Transforming descriptors to uncorrelated variables by

- principal component analysis (PCA)
- partial least square (PLS)
for example applied in comparative molecular field analysis (CoMFA), see below

Methods that intrinsically handle correlated variables

- neural networks, especially deep learning networks


## Partial least square (I)

The idea is to construct a small set of latent variables $t_{i}$ (that are orthogonal to each other and therefore uncorrelated) from the pool of inter-correlated descriptors $x_{i}$.


In this case $t_{1}$ and $t_{2}$ result as the normal modes of $x_{1}$ and $x_{2}$ where $t_{1}$ shows the larger variance.

## Partial least square (II)

The predicted term $y$ is then a QSAR equation using the latent variables $t_{i}$

$$
y=b_{1} t_{1}+b_{2} t_{2}+b_{3} t_{3}+\ldots+b_{m} t_{m}
$$

where

$$
\begin{aligned}
& t_{1}=c_{11} x_{1}+c_{12} x_{2}+\ldots+c_{1 n} x_{n} \\
& t_{2}=c_{21} x_{1}+c_{22} x_{2}+\ldots+c_{2 n} x_{n} \\
& \cdot \cdot \cdot \cdot \cdot \\
& t_{m}=c_{m 1} x_{1}+c_{m 2} x_{2}+\ldots+c_{m n} x_{n}
\end{aligned}
$$

The number of latent variables $t_{i}$ is chosen to be (much) smaller than that of the original descriptors $x_{i}$.
But, how many latent variables are reasonable?
$\rightarrow$ plot $r^{2}$, se, $q^{2}$ and their fluctations against the number of latent variables and identify the minimal number of latent variables.

## Principal Component Analysis PCA (I)

Problem: Which are the (decisive) significant descriptors?
Principal component analysis determines the normal modes from a set of descriptors/variables.
This is achieved by a coordinate transformation resulting in new axes. The first principal component then shows the largest variance of the data. The second and further normal components are orthogonal to each other.


## Principal Component Analysis PCA (II)

The first component ( pc 1 ) shows the largest variance, the second component the second largest variance, and so on.


Lit: E.C. Pielou: The Interpretation of Ecological Data, Wiley, New York, 1984

## Principal Component Analysis PCA (III)

The significant principal components usually have an Eigen value >1 (Kaiser-Guttman criterion). Frequently there is also a kink that separates the less relevant components (Scree test)


## Principal Component Analysis PCA (IV)

The obtained principal components should account for more than $80 \%$ of the total variance.



## Principal Component Analysis (V)

Example: What descriptors determine the logP?

| property | pc1 | pc2 | pc3 |
| :--- | ---: | ---: | ---: |
| dipole moment | 0.353 |  |  |
| polarizability |  | 0.504 |  |
| mean of +ESP | 0.397 | -0.175 | 0.151 |
| mean of -ESP | -0.389 | 0.104 | 0.160 |
| variance of ESP 0.403  -0.244 <br> minimum ESP -0.239 -0.149 0.548 <br> maximum ESP <br> molecular volume 0.422  0.170 <br> surface <br> fraction of total <br> variance 0.519 0.506 0.115$\quad 28 \%$ | $22 \%$ | $10 \%$ |  |

Lit: T.Clark et al. J.Mol.Model. 3 (1997) 142


## Comparative Molecular Field Analysis (I)

The molecules are placed into a 3D grid and at each grid point the steric and electronic interaction with a probe atom is calculated (force field parameters)


For this purpose the GRID program can be used:
P.J. Goodford J.Med.Chem. 28 (1985) 849.

Problems: „active conformation" of the molecules needed All molecule must be superimposed (aligned according to their common scaffold)

Lit: R.D. Cramer et al. J.Am.Chem.Soc. 110 (1988) 5959.

## Comparative Molecular Field Analysis (II)

The resulting coefficients for the matrix $S(N$ grid points, $P$ probe atoms) have to determined using a PLS analysis.

| compound | $\log$ (1/C) | S1 | S2 | S3 | $\ldots$ | P1 | P2 | P3 | $\ldots$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| steroid1 | 4.15 |  |  |  |  |  |  |  |  |
| steroid2 | 5.74 |  |  |  |  |  |  |  |  |
| steroid3 | 8.83 |  |  |  |  |  |  |  |  |
| steroid4 | 7.6 |  |  |  |  |  |  |  |  |
| $\ldots$ |  |  |  |  |  |  |  |  |  |
|  |  |  |  | $\nu^{N}$ |  |  |  |  |  |

## Comparative Molecular Field Analysis (III)












Lit: R.D. Cramer et al. J.Am.Chem.Soc. 110 (1988) 5959.

## Comparative Molecular Field Analysis (IV)

Analog to QSAR descriptors, the CoMFA variables can be interpreted. Here (color coded) contour maps are helpful

yellow: regions of unfavorable steric interaction blue: regions of favorable steric interaction

Lit: R.D. Cramer et al. J.Am.Chem.Soc. 110 (1988) 5959

## CoMFA (V) 3-D Database online:


„A 3-D QSAR Models Database for Virtual Screening"

Compounds can be screened against a large set of precalculated models

Jmol

| Maps Table |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| \% | 0 | 10 | 20 | 30 | 40 | 50 | 60 | 70 | 80 | 90 | 100 |
| PLS_Coeff | © | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ |
| ComFA_Maps | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ | $\bigcirc$ |

Rino Ragno et al. Università di Roma (Italy)

## Comparative Molecular Similarity Indices Analysis (CoMSIA)

CoMFA based on similarity indices at the grid points


Lit: G.Klebe et al. J.Med.Chem. 37 (1994) 4130.

## Neural Networks (I)

Neural networks can be regarded as a common implementation of artificial intelligence. The name is derived from the network-like connection between the switches (neurons) within the system. Thus they can also handle inter-correlated descriptors.


modeling of a (regression) function

From the many types of neural networks, backpropagation and unsupervised maps are the most frequently used.

## Neural Networks (II)

A typical backpropagation net consists of neurons organized as the input layer, one or more hidden layers, and the output layer


Furthermore, the actual kind of signal transduction between the neurons can be different:

bipolar
hard limiter

sigmoidal transfer logic

## Recursive Partitioning

Instead of quantitative values often there is only qualitative information available, e.g. substrates versus non-substrates
Thus we need classification methods such as

- decision trees, naïve Bayes classification
- support vector machines
- boosting
- (neural networks): partition at what score value ?


Picture: J. Sadowski \& H. Kubinyi J.Med.Chem. 41 (1998) 3325.

## Decision Trees

Iterative classification


Lit: J.R. Quinlan Machine Learning 1 (1986) 81.

## Support Vector Machines

Support vector machines generate a hyperplane in the multidimensional space of the descriptors that separates the data points.


Advantages: accuracy, a minimum of descriptors (= support vectors) used
Disadvantage: Interpretation of results, design of new compounds with desired properties, which descriptors for input

## Property prediction: So what?

Classical QSAR equations: small data sets, few descriptors that are (hopefully) easy to understand

CoMFA: small data sets, lots of descriptors

easy visual interpretation of resulting interaction regions

Partial least square: small data sets, many descriptors
black box
methods

Neural nets: large data sets, some, preselected descriptors $\}$ often difficult

Support vector machines: large data sets, many descriptors

## Interpretation of QSAR equations (1)

Caution is required when extrapolating beyond the underlying data range. Outside this range no reliable predicitions can be made



Beyond the black stump ...

Kimberley, Western Australia


## Interpretation of QSAR equations (2)

There should be a reasonable connection between the used descriptors and the predicted quantity.
Example: H. Sies Nature 332 (1988) 495.
Scientific proof that babies are delivered by storks


## Interpretation of QSAR equations (3)

Another striking correlation
"QSAR has evolved into a perfectly practiced art of logical fallacy"

S.R. Johnson J.Chem.Inf.Model. 48 (2008) 25.
$\rightarrow$ the more descriptors are available, the higher is the chance of finding some that show a chance correlation

## Interpretation of QSAR equations (4)

The scientific proof that chocolate makes you smarter....


Figure 1. Correlation between Countries' Annual Per Capita Chocolate Consumption and the Number of Nobel Laureates per 10 Million Population.
F.H. Messerli New England J. Med. Oct.10, 2012 DOI:10.1056/NEJMon1211064

6th lecture
Modern Methods in Drug Discovery WS23/24

## Interpretation of QSAR equations (6)

Predictivity of QSAR equations in between data points.
The hypersurface is not smooth: activity islands vs. activity cliffs:
Even small changes in structure can change the activity strongly.
In other words, what is in between the present molecules?

$$
r^{2}=0.99 \mathrm{se}=0.27
$$




Bryce Canyon National Park, Utah

Lit: G.M. Maggiora J.Chem.Inf.Model. 46 (2006) 1535.
S.R. Johnson J.Chem.Inf.Model. 48 (2008) 25.

## Interpretation of QSAR equations (6)

Which QSAR performance is realistic and where does overfitting start?

- standard deviation (se) of 0.2-0.3 log units corresponds to a typical 2-fold error in experiments („soft data"). This gives rise to an upper limit of
- $r^{2}$ between 0.77-0.88 (for biological systems) $\rightarrow$ obtained correlations above 0.90 are highly likely to be accidental or due to overfitting (except for physico-chemical properties that show small errors, e.g. boiling points, logP, NMR ${ }^{13} \mathrm{C}$ shifts)


But: even random correlations can sometimes be as high as 0.84
Lit: A.M.Doweyko J.Comput.-Aided Mol.Des. 22 (2008) 81-89.

## Interpretation of QSAR equations (7)

Accidental correlation of a single descriptor (1000 random descriptors)

$\rightarrow$ Dismiss unsuitable variables from the pool of descriptors.
Lit: M.C.Hutter J.Chem.Inf.Model. (2011) DOI: 10.1021/ci200403j

## Interpretation of QSAR equations (8)

Low correlations...


I DON'T TRUST LINEAR REGRESSIONS WHEN ITS HARDER TO GUESS THE DIRECTION OF THE CORRELATION FROM THE SCATTER PLOT THAN TO FIND NEW CONSTELLATIONS ON IT.

Source: https://xkcd.com

## Interpretation of QSAR equations (9)

According to statistics more people die after being hit by a donkey than from the consequences of an airplane crash.

„An unsophisticated forecaster uses statistics as a drunken man uses lamp-posts - for support rather than for illumination" Andrew Lang (1844-1912)
further literature: R.Guha J.Comput.-Aided Mol.Des. 22 (2008) 857-871.

