

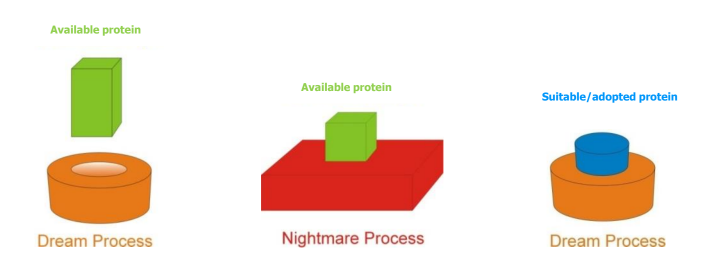
## **Protein Engineering**

- Outline
  - ☐ Limitations of proteins in biotechnology processes
  - ☐ Definition and aim of protein engineering
  - Targeted properties of proteins
  - Basic approaches in protein engineering
    - **DIRECTED EVOLUTION**
    - RATIONAL DESIGN
    - **SEMI-RATIONAL DESIGN**
  - Examples

### Proteins in biotechnology

- key problem -availability of optimal protein for specific process
- ☐ traditional biotechnology adapt process
- modern biotechnology adapt protein

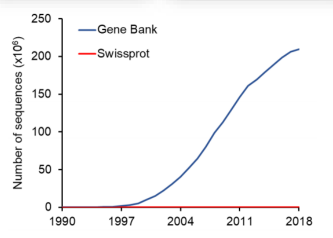
#### **HOW TO OBTAIN OPTIMAL PROTEIN?**



#### Proteins in biotechnology

- classical screening
  - screening culture collections
  - polluted and extreme environment
- environmental gene libraries
  - metagenomic DNA
- data-base mining
  - gene databases
  - (meta)genome sequencing projects
  - numerous uncharacterised proteins





IF SUITABLE PROTEIN DOES NOT EXIST IN NATURE?

PROTEIN ENGINEERING

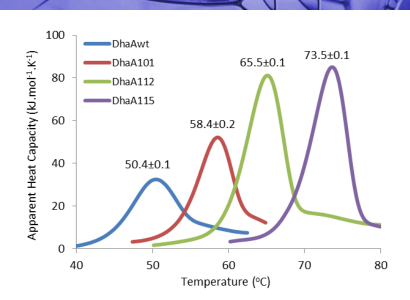
#### Proteins in biotechnology

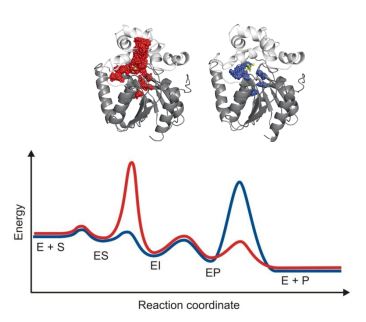
- the process of constructing novel protein molecules by design first principles or altering existing structure "de novo design"
- use of genetic manipulations to alter the coding sequence of a gene and thus **modify the properties of the protein**
- AIMS AND APPLICATIONS
  - technological optimisation of the protein to be suitable in particular technology purpose
  - scientific desire to understand what elements of proteins contribute to folding, stability and function

### Targeted properties of proteins

- structural properties of proteins
  - stability (temperature, solvents)
  - tolerance to pH, salt
  - resistance to oxidative stress

- ☐ functional properties of proteins
  - reaction type
  - substrate specificity and selectivity
  - kinetic properties (e.g.,  $K_{\rm m}$ ,  $k_{\rm cat}$ ,  $K_{\rm i}$ )
  - cofactor selectivity
  - protein-protein or protein-DNA interactions





#### **RATIONAL DESIGN**

1. Computer aided design

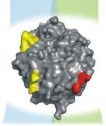


2. Site-directed mutagenesis



Individual mutated gene

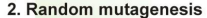
- 3. Transformation
  - 4. Protein expression
    - 5. Protein purification
      - 6. not applied

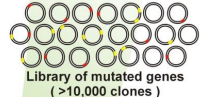


Improved protein

#### **DIRECTED EVOLUTION**

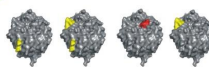
1. not applied





- 3. Transformation
- 4. Protein expression
- 5. not applied
- 6. Screening and selection
  - stability
  - selectivity
  - affinity
  - activity





Selected mutant enzymes



Constructed mutant enzyme

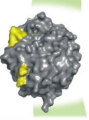
7. Biochemical testing

#### Directed evolution

- directed evolution techniques emerged during mid-1990s
- inspired by natural evolution
- this form of "evolution" does not match what Darwin had envisioned
  - requires outside intelligence, not blind chance
  - does not create brand new species, macroevolution,
     but only improvements of molecules, molecular evolution
  - does not take millions of years, but happens rapidly

#### Directed evolution

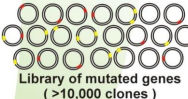
- evolution in test tube comprises two steps
  - random mutagenesisbuilding mutant library (diversity)
  - screening and selection
     identification of desired biocatalyst
- prerequisites for directed evolution
  - gene encoding protein of interest
  - method to create mutant library
  - suitable expression system
  - screening or selection system



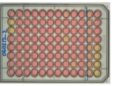
Improved protein

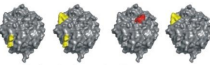
1. not applied

2. Random mutagenesis



- 3. Transformation
- 4. Protein expression
- 5. not applied
- 6. Screening and selection
  - stability
  - selectivity
  - affinity
  - activity





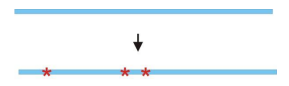
7. Biochemical testing



### Methods to create mutant libraries

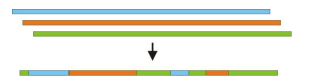
- □ technology to generate large diversity
  - NON-RECOMBINING

one parent gene -> variants with point mutations



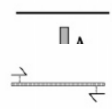
#### RECOMBINING

several parental homologous genes -> chimeras



#### Non-recombining mutagenesis

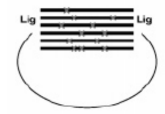
- UV irradiation or chemical mutagens (traditional)
- mutator strains lacks DNA repair mechanism mutations during replication (e.g., Epicurian coli XL1-Red)
- error-prone polymerase chain reaction (ep-PCR)
  - gene amplified in imperfect copying process
     (e.g., unbalanced deoxyribonucleotides concentrations,
     high Mg<sup>2+</sup> concentration, Mn<sup>2+</sup>, low annealing temperatures)
  - 1 to 20 mutation per 1000 base pairs
- saturation mutagenesis
  - randomization of single or multiple codons
  - gene site saturation mutagenesis
- other methods
  - insertion/deletions (InDel)
  - cassette mutagenesis (region mutagenesis)





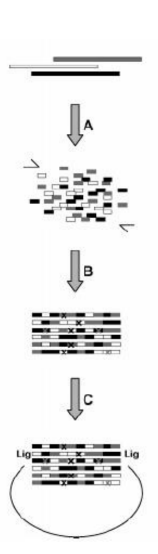






#### Recombining mutagenesis

- □ also refered to as "sexual mutagenesis"
- DNA shuffling
  - fragmentation step
  - random reassembly of segments
- □ StEP staggered extension process
  - simpler then shuffling
  - random reannealing combined with limited primer extension
- other methods
  shuffling of genes with lower homology down to 70%
  (e.g., RACHITT, ITCHY, SCRATCHY)

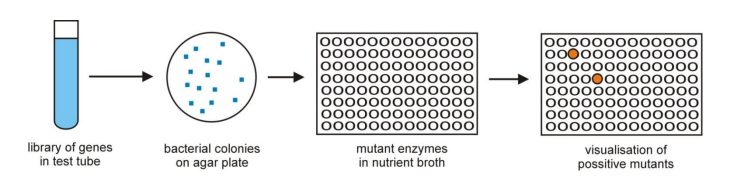


#### Screening and selection

- most critical step of direct evolution
- ☐ isolation of positive mutants hiding in library
  - HIGH THROUGHPUT SCREENING
     individual assays of variants one by one



DIRECT SELECTION
 display techniques (link between genotype and phenotype)



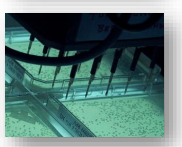


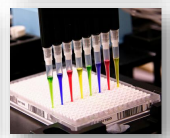
### (Utra) High throughput screening

- common methods not applicable
- agar plate (pre)screening
- microtiter plates screening
  - 96-, 384- or 1536-well formate
  - robot assistance (colony picker, liquid handler)
  - 10<sup>4</sup> libraries
  - volume 10 100 uL
- microfluidic systems (Lesson 5)
  - water in oil emulsions (up to 10 kHz)
  - FACS sorting (10<sup>8</sup> events/hour)
  - 10<sup>9</sup> libraries
  - volume 1 10 pL

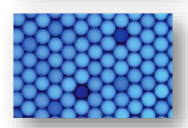


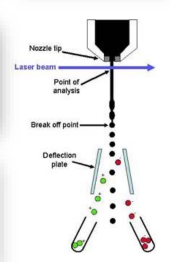








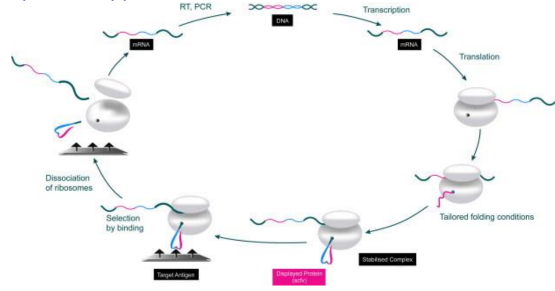


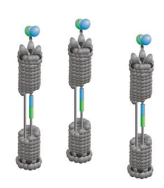


#### Direct selection

- □ not generally applicable (mutant libraries >10 variants)
- ☐ link between genotype and phenotype
- display technologies
  - ribosome display
  - phage display
- life-or-death assay
  - auxotrophic strain
  - toxicity based selection

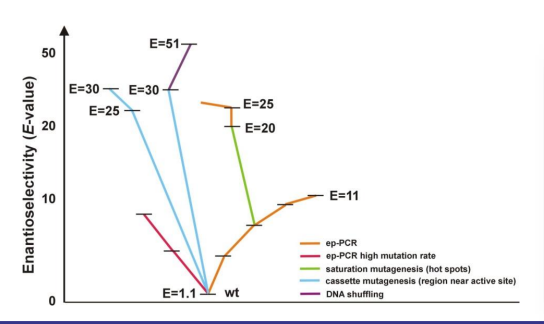


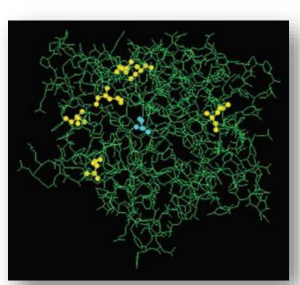




#### Example of Directed evolution

- directed evolution of enantioselectivity
  - lipase from P. aeruginosa (E-value improved from 1.1 into 51)
  - spectrophotometric screening of (R)- and (S)-nitrophenyl esters
  - 40 000 variants screened
  - the best mutant contains six amino acid substitutions





#### **RATIONAL DESIGN**

1. Computer aided design



2. Site-directed mutagenesis



Individual mutated gene

- 3. Transformation
  - 4. Protein expression
    - 5. Protein purification
      - 6. not applied

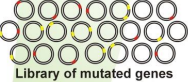


Improved protein

#### **DIRECTED EVOLUTION**

1. not applied

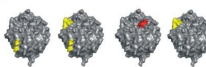
2. Random mutagenesis



(>10,000 clones)

- 3. Transformation
- 4. Protein expression
- 5. not applied
- 6. Screening and selection
  - stability
  - selectivity
  - affinity
  - activity





Selected mutant enzymes



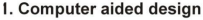
Constructed mutant enzyme

7. Biochemical testing

### Rational design

- emerged around 1980s as the original protein engineering approach
- knowledge based combining theory and experiment
- protein engineering cycle:
  - "structure-theory-design-mutation-purification-analysis"
- ☐ difficulty in prediction of mutation effects on protein property
- de novo design most challenging

#### Principal of rational design



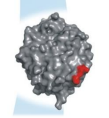


2. Site-directed mutagenesis



Individual mutated gene

- 3. Transformation
  - 4. Protein expression
    - 5. Protein purification
      - 6. not applied



**Improved** protein



Constructed mutant enzyme

7. Biochemical testing

- rational design comprises:
  - design understanding of protein functionality
  - experiment construction and testing of mutants
  - prerequisites for rational design:
    - gene encoding protein of interest
    - 3D structure (e.g., X-ray, NMR) or sequence alignment
    - structure-function relationship
    - computational methods and capacity
    - side directed mutagenesis techniques
    - efficient expression system
    - biochemical tests

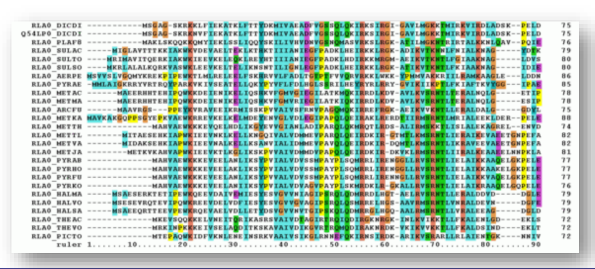
### Design

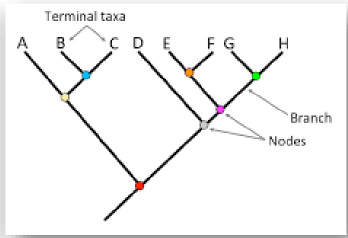


- homologous wild-type sequences alignment
- identifying amino acid residues responsible for differences
- design combination of possitive mutation from all parental proteins

#### ■ ANCESTRAL RECONSTRUCTION

- construction of phylogenetic tree
- design nods prediction by consensus approach

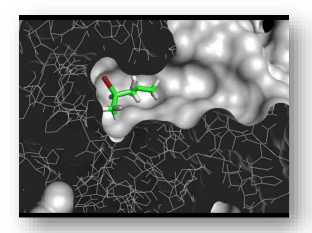


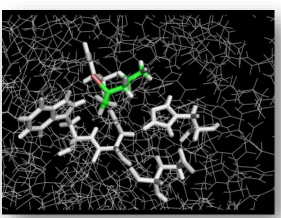


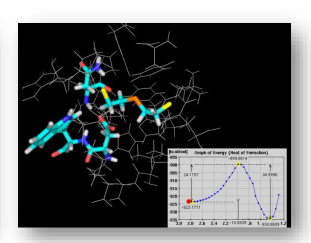
### Design



- prediction of enzyme function from structure alone is challenging
- protein structure (X-ray crystallography, NMR, homology models!)
- molecular modelling
  - molecular docking
  - molecular dynamics
  - quantum mechanics/molecular mechanics (QM/MM)

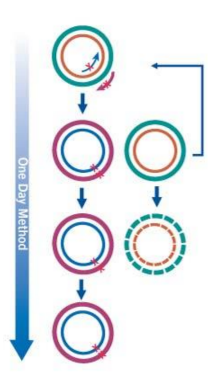






#### Construction

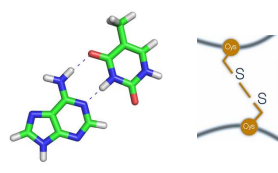
- ☐ site-directed mutagenesis
  - introducing point mutations
- multi site-directed mutagenesis
- gene synthesis
  - commercial service
  - codone optimisation

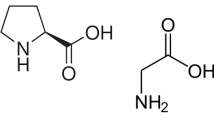


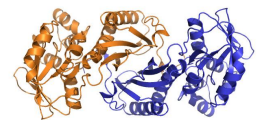




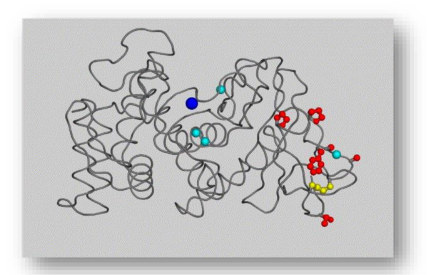
- rational design of protein stability
  - stability to high temperature, extreme pH, proteases etc.
  - stabilizing mutations increase strength of weak interactions
    - o salt bridges and H-bonds Eijsink et al., Biochem. J. 285: 625-628, 1992
    - S-S bonds
       Matsumura et al., Nature 342: 291-293, 1989
    - o addition of prolines
      Watanabe et al., Eur. J. Biochem. 226: 277-283, 1994
    - o **less glycines** *Margarit et al., Protein Eng. 5: 543-550, 1992*
    - o oligomerisation
      Dalhus et al., J. Mol. Biol. 318: 707-721, 2002







- engineering protein to resist boiling
  - reduced rotational freedom Ser65Pro, Ala96Pro
  - introduction of disulfide bridge Gly8Cys + Asn60Cys
  - improved internal hydrogen bond Ala4Thr
  - filling cavity Tyr63Phe



Half-lifes (min.)	80°C	100°C
wild type	17.5	>0.5
mutant	stable	170

#### RATIONAL DESIGN

1. Computer aided design



2. Site-directed mutagenesis



Individual mutated gene

- 3. Transformation
  - 4. Protein expression
    - 5. Protein purification
      - 6. not applied

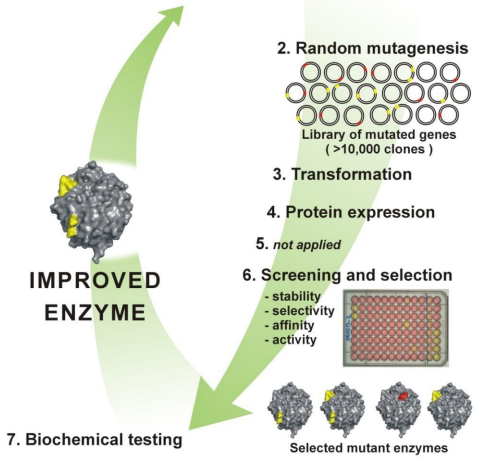




7. Biochemical testing

#### DIRECTED EVOLUTION

1. not applied



#### **RATIONAL DESIGN**

1. Computer aided design

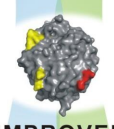


2. Site-directed mutagenesis



Individual mutated gene

- 3. Transformation
  - 4. Protein expression
    - 5. Protein purification
      - 6. not applied



IMPROVED ENZYME

#### DIRECTED EVOLUTION

## SEMIRATIONAL DESIGN

2. Random mutagenesis



(>10,000 clones)

- 3. Transformation
- 4. Protein expression
- 5. not applied
- 6. Screening and selection
  - stability
  - selectivity
  - affinity
  - activity





Constructed mutant enzyme

7. Biochemical testing

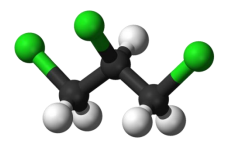






Selected mutant enzymes

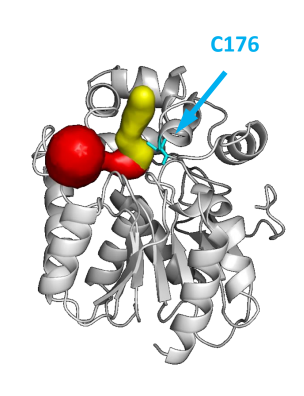
conversion of 1,2,3-trichloropropane
by DhaA from Rhodococcus erythropolis Y2

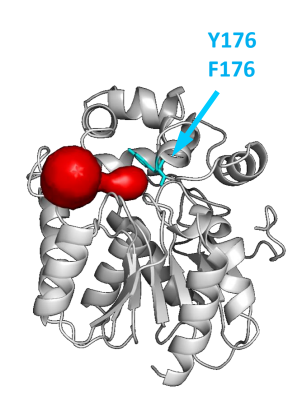




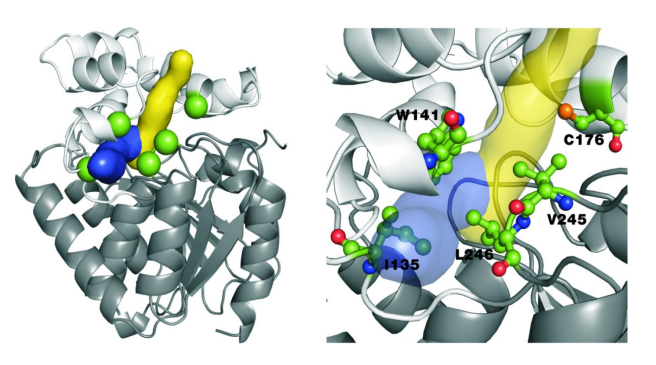


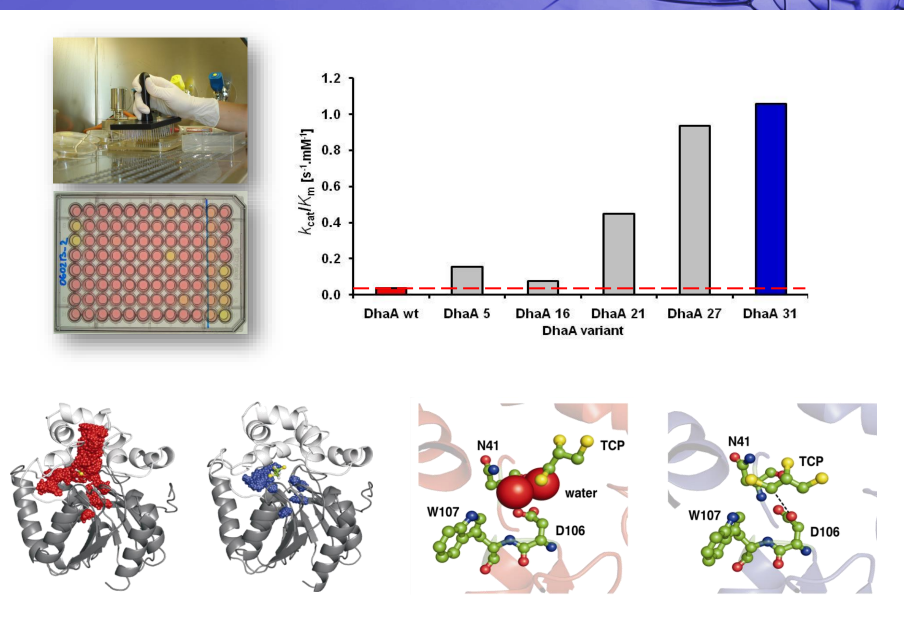
- conversion of 1,2,3-trichloropropane
  by DhaA from Rhodococcus erythropolis Y2
- □ **DIRECTED EVOLUTION** importance of access pathways



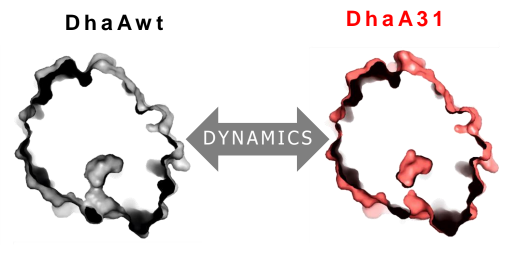


- conversion of 1,2,3-trichloropropane
  by DhaA from Rhodococcus erythropolis Y2
- □ DIRECTED EVOLUTION importance of access pathways
- SEMI-RATIONAL DESIGN hot spots in access tunels
- ☐ library of **5,300 clones** screened

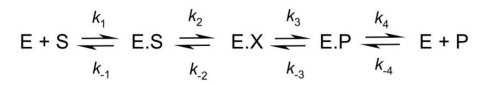


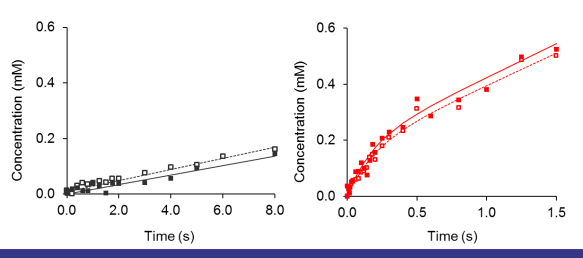


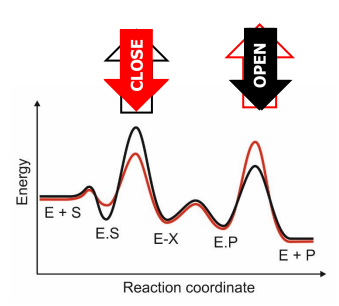
#### Engineering activity of DhaA



	DhaA wt	DhaA 31
К <sub>т</sub> (mM)	0.98 ± 0.17	1.19 ± 0.15
$k_{\rm cat}$ (s <sup>-1</sup> )	0.035 ± 0.002	1.26 ± 0.05
$k_{\rm cat}/K_{\rm m}  ({\rm mM}^{-1}.{\rm s}^{-1})$	0.04	1.06

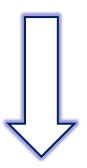






#### STANDARD DESIGN

- random mutagenesis (2-3 positions)
- library of 10<sup>4</sup> clones



#### ADVANCED DESIGN

- random mutagenesis (5-7 positions)
- library of >10<sup>6</sup> clones

volume:  $100^{\circ} \mu L$  assays/day:  $10^{3}$ 





volume: 10' pL assays/day: 10<sup>7</sup>



#### Software tools





loschmidt.chemi.muni.cz/caver



□ CaverDock

(coming soon)



Hotspot Wizard 2.0

loschmidt.chemi.muni.cz/hotspotwizard



☐ FireProt 1.0

loschmidt.chemi.muni.cz/fireprot



🖵 Cavetta

(coming soon)

#### Reading

- Lutz, S. 2010: **Beyond directed evolution semi-rational protein engineering and design**. *Curr Opin Biotechnol*. 21(6): 734-743
- ☐ Computational enzyme redesign and Computational de novo enzyme design (page 5-7)



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# Beyond directed evolution - semi-rational protein engineering and design

#### Stefan Lutz

Department of Chemistry, Emory University, 1515 Dickey Drive, Atlanta, GA, 30322

#### **Abstract**

Over the last two decades, directed evolution has transformed the field of protein engineering. The advances in understanding protein structure and function, in no insignificant part a result of directed evolution studies, are increasingly empowering scientists and engineers to device more effective methods for manipulating and tailoring biocatalysts. Abandoning large combinatorial libraries, the