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Kód předmětu: C8980

MASARYKOVA UNIVERZITA

Protein expression and purification

VII. Fusion proteins and affinity purification

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Tento projekt je spolufinancován Evropským sociálním fondem a státním rozpočtem České republiky.







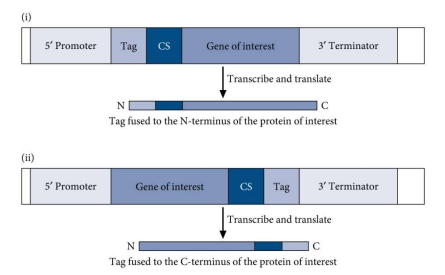


INVESTICE DO ROZVOJE VZDĚLÁVÁNÍ

Fusion proteins (tagged proteins)

Translation fusion of sequences coding a recombinant protein and

- a) short peptides [ex. (His)_n, $(Asp)_n$, $(Arg)_n$...].
- b) protein domains, entire proteins [ex. MBP, GST, thioredoxin ...].



Engineering a tagged protein requires adding the DNA encoding the tag to either the 5' or 3' end of the gene encoding the protein of interest to generate a single, recombinant protein with a tag at the N- or C-terminus. The stretch of amino acids containing a target cleavage sequence (CS) is included to allow selective removal of the tag.

Expression plasmids containing various tags are commercially available.

Purposes of fusion tags

- ➤ Increasing the yield of recombinant proteins Fusion of the N-terminus of the target protein to the C-terminus of a highly expressed fusion partner results in high level expression of the target protein.
- ➤ Enhancing the solubility of recombinant proteins Fusion of the N-terminus of the target protein to the C-terminus of a soluble fusion partner often improves the solubility of the target protein.
- ➤ Improving detection Fusion of the target protein to either terminus of a short peptide (epitope tag) or protein which is recognized by an antibody (Western blot analysis) or by biophysical methods (e.g. GFP by fluorescence) facilitates the detection of the resulting protein during expression or purification.
- ➤ Localization A tag, usually located on the N-terminus of the target protein, which acts as an address for sending a protein to a specific cellular compartment.
- ➤ Facilitating the purification of recombinant proteins Simple purification schemes have been developed for proteins used at either terminus which bind specifically to affinity resins.

No single tag is ideally suited for all of these purposes.

Fusion partner (tag)	Size	Tag placement	Uses
His-tag	6, 8, or 10 aa	N- or C-terminus	Purification, detection
Thioredoxin	109 aa (11.7 kDa)	N- or C-terminus	Purification, solubility enhancement
Calmodulin-binding domain (CBD)	26 aa	N- or C-terminus	Purification
Avidin/streptavidin Strep-tag	8 aa	N- or C-terminus	Purification, secretion
Glutathione S-transferase (GST)	26 kDa	N-terminus	Purification, solubility enhancement
Maltose binding protein (MBP)	396 aa (40 kDa)	N- or C-terminus	Purification, solubility enhancement
Green fluorescent protein (GFP)	220 aa (27 kDa)	N- or C-terminus	Localization, detection, purification
Poly-Arg	5-16 aa	N- or C-terminus	Purification, solubility enhancement
N-utilization substance A (NusA)	495 aa (54.8 kDa)	N-terminus	Solubility enhancement

Combinatorial tagging

➤ No single tag is ideally suited for all purposes. Therefore, combinatorial tagging might be the only way to harness the full potential of tags in a high-throughput setting.

Combinations:

Solubility-enhancing tag + purification tag: MBP + His₆ tag

2x purification tag: IgG-binding domain + streptavidin-binding domain

Detection tag + purification tag: $GFP + His_6 tag$

Detection tag + 2x purification tag + immunodetection: GFP + SBP domain + His₈ tag + c-Myc

^aGST, glutathione S-transferase; MBP, maltose-binding protein; NusA, N-utilization substance A; FLAG, FLAG-tag peptide; BAP, biotin acceptor peptide; His₆, hexahistidine tag; STREP, streptavidin-binding peptide; SET, solubility-enhancing tag; CBP, calmodulin-binding peptide.

Advantages and disadvantages of used fusion tags

➤ Proteins do not naturally lend themselves to high-throughput analysis because of their diverse physiological properties. Affinity tags have become indispensable tools for structural and functional proteomics.



➤ Because affinity tags have the potential to interfere with structural and functional studies, provisions must also be made for removing them.

^bDerivatives of thioredoxin have been engineered to have affinity for immobilized metal ions (His-patch thioredoxin) or avidin/streptavidin [38]. Waugh, 2005

Otázka č. 1:

Jaké jsou důvody pro využívaní tagů/kotev? Vyjmenujte 3.

Increasing the yield of recombinant proteins using fusion technology

Yield enhancing tags are proteins and peptides which can be involved in:

- ➤ Increasing the efficiency of translation initiation (e.g. GST, MBP, NusA...)
 - Advantage of N-terminal tags
 - Providing a reliable context for efficient translation initiation
 - Ribosome efficiently initiates translation at the N-terminal methionin of the tag
 - Deleterious secondary structures are more likely to occur in conjunction with short N-terminal tags because short RNA-RNA interactions tend to be more stable than long-range interactions.

> Protection against proteolytic degradation

- Several studies have shown that the nature of terminal residues in a protein can play a role in recognition and subsequent action by proteases and in some cases affinity tags might improve the yield of recombinant proteins by rendering them more resistant to intracellular proteolysis.
- ➤ Helping to properly fold their partners leading to increased solubility of the target protein (*in vivo* and *in vitro*).

Enhancing the solubility of recombinant proteins

Solubility-enhancing tags

- Advantage of N-terminal tags
- Rather proteins (highly soluble proteins) than peptides
- -Fusion with a soluble fusion partner often helps to properly fold their fusion partners leading to improved solubility (*in vivo and in vitro*) of the target protein.
- -Fusion partners do not perform equally with all target proteins, and each target protein can be differentially affected by several fusion tags (Esposito and Chatterjee, 2006).
- The choice of a fusion partner is still a trial-and-error experience.

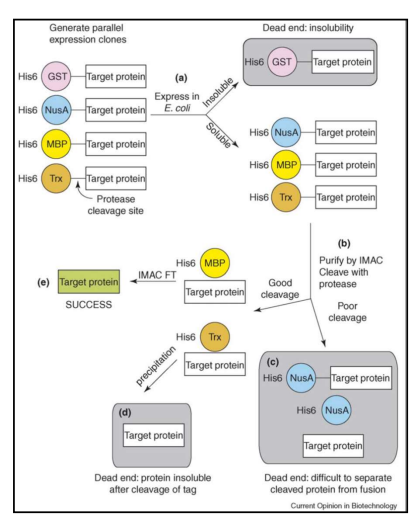
>PROTEINS

Tag	Protein	Source organism
MBP	Maltose-binding protein	Escherichia coli
GST	Glutathione-S-transferase	Schistosoma japonicum
Trx	Thioredoxin	Escherichia coli
NusA	N-Utilization substance	Escherichia coli
SUMO	Small ubiquitin-modifier	Homo sapiens
SET	Solubility-enhancing tag	Synthetic
DsbC	Disulfide bond C	Escherichia coli
Skp	Seventeen kilodalton protein Escherichia	
T7PK	Phage T7 protein kinase Bacteriopha	
GB1	Protein G B1 domain Streptoco	
ZZ	Protein A IgG ZZ repeat domain	Staphylococcus aureus

Adopted from Esposito and Chatterjee, 2006

> PEPTIDES

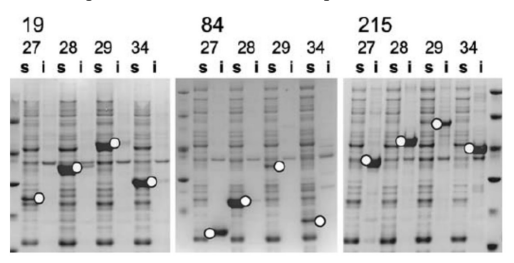
Poly-Arg Poly-Lys



Schematic representation of the pathway from protein expression to purification using solubility tags (Esposito and Chatterjee, 2006).

Enhancing the solubility of recombinant proteins

19, 84, 215 – human proteins involved in cancer produced in *E.coli*



Example of SDS PAGE with **soluble (s) and insoluble (i) fractions** following lysis. The results produced from the four different expression vectors (27: His tag only; 28: thioredoxin + His tag; 29: GST + His tag; 34: GB1 + His tag) are shown for three different target proteins (*Hammarstrom et al.*, 2006).

Solubility-enhancing tags - the mechanism of action

-The mechanism by which partners exert their solubilising function is not fully understood and possibly differs between fusion proteins

Examples of possible mechanisms

Maltose binding protein (MBP) has an intrinsic chaperone-like activity. MBP might bind reversibly to exposed hydrophobic regions of nascent target polypeptide, steering the polypeptides towards their native conformation by a chaperone like –mechanism.

N-utilization substance (NusA) decreased translation rates by mediating transtriptional pausing, that might enable critical folding events to occur.

MBP and N-utilization substance (NusA) attract chaperones. The fusion tag drives its partner protein into a chaperone-mediated folding pathway. MBP and NusA interact with GroEL in E. coli (Huang and Chuang, 1999).

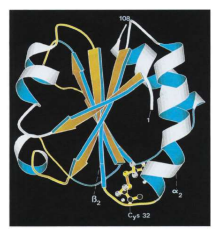
Small ubiquitin related modifier (SUMO) promotes the proper folding and solubility of its target proteins possibly by exerting chaperoning effects in a similar mechanism to the described for its structural homolog Ubiquitin (Ub; Khorasanizadeh et al., 1996).

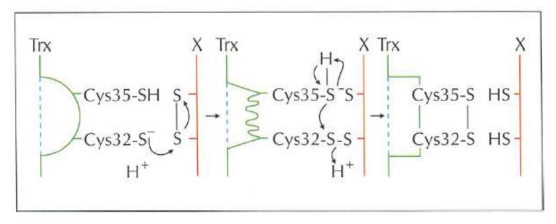
Negative charged tags (highly acidic peptide) inhibit aggregation by increasing electrostatic repulsion between nascent polypepdides (Zhang et. 2004).

Solubility-enhancing tag – mechanism of action

Thioredoxin

- ➤ Its intrinsic oxido-reductase activity is responsible for the reduction of disulfide bonds through thio-disulfide exchange
- Thioredoxin serves as a covalently joined molecular chaperone independently of redox activity. Thioredoxin may, thus, act to prevent the aggregation and precipitation of fused nascent proteins, giving them an extended opportunity to adopt their correct tertiary folds.





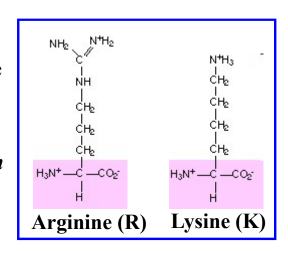
Proposed mechanism of thioredoxin-catalyzed protein disulfide reduction. Reduced thioredoxin [Trx-(SH)2] binds to a target protein via its hydrophobic surface area. Nucleophilic attack by the thiolate of Cys32 results in formation of a transient mixed disulfide, which is followed by nucleophilic attack of the deprotonated Cys35 generating Trx-S2 and the reduced protein. Conformation changes in thioredoxin and the target protein occur during the reaction.

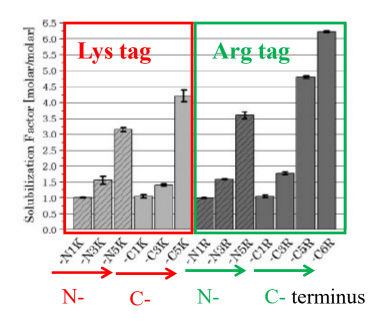
In vitro solubility-enhancing tags

Short peptide tags

Poly-Lys tag, poly-Arg tag = one, three and five lysine or arginine residues fused to the C- or N-terminus of the target protein

Solubility as defined here is the maximum protein concentration of the supernatant after centrifugation of the supersaturated protein sample (in vitro solubility).





BPTI-22 = bovine pancreatic trypsin inhibitor variant containing 22 alanines

The solubilization factor is defined as the molar ratio between the solubility of tagged BPTI-22 variants and that of the reference BPT-22 molecule.

The solubilization effect of poly-Lys tags is lower than that of poly-Arg tags (lysines are less hydrophilic than arginines).

Biochemical properties of poly-Arg and poly-Lys tagged BPTI-22 protein

	Protein Solubility				
Protein	Conc. [mM] (Conc. [mg/ml]) ^a	Solubilization Factor ^b	<i>T</i> _m (°C)	Rel. Trypsin Inhibitor Activity (%) ^c	у
BPTI-22	1.70 (10.00)		38.4	_	
-N1K	1.70 (10.40)	1.00(1.04)	35.2	1.05	
-N3K	2.66 (19.97)	1.56 (2.00)	34.4	1.04	
-N5K	5.37 (35.60)	3.16 (3.56)	34.3	1.05	
-C1K	1.79 (10.95)	1.05 (1.10)	34,6	1.05	The addition of 0.5 M Arg
-C3K	2.41 (15.28)	1.42 (1.53)	36.2	1.05	barely increased its solubility,
-C5K	7.16 (47.47)	4.21 (4.75)	35.0	1.02	and trypsin activity was
-N1R	1.69 (10.34)	0.99 (1.03)	35.5	1.02	inhibited by the high arginine
-N3R	2.70 (17.23)	1.59 (1.72)	35.6	0.99	•
-N5R	6.20 (41.11)	3.65 (4.11)	35.5	0.99	concentration. On the other
-C1R	1.81 (11.07)	1.06(1.11)	35.0	1.05	hand, addition of 50 mM
-C3R	3.02 (19.26)	1.78 (1.93)	34.4	1.05	Arg+Glu was more effective
-C5R	8.23 (54.56)	4.84 (5.46)	34.8	1.08	and increased protein
-C6R	10.59 (73.41)	6.22 (7.34)	32.7	1.1	solubility more than
BPTI-22 ^d	5.63 (33.11)	3.31 (3.31)	NDe	1.09	•
BPTI-22f	2.01 (11.82)	1.18 (1.18)	ND^e	NAg	threefold.

Protein solubility was determined as the maximum supernatant concentration of a supersaturated protein solution at 4°C in 100 mM acetate buffer pH 4.7.

^a Maximum concentrations calculated in milligrams per milliliter are indicated in parenthesis. The Mw of BPTI-22, -N1K and -C1K, -N3K and -C3K, -N5K and -C5K, -N1R and -C1R, -N3R and -C3R, -N5R and -C5R, and -C6R are, respectively: 5880, 6123, 6379, 6636, 6151, 6463, 6776, and 6932 Da.

b Calculated as the ratio between the molar protein solubility of BPTI-22 and that of tagged BPTI-22. Values in parenthesis indicate the ratio calculated in milligrams per milliliters.

^c Relative trypsin inhibitory activity calculated as the ratio between the activity of BPTI-22 and that of tagged BPTI-22. BPTI-22, which lacks R39, an arginine residue involved in two hydrogen bonding interactions with the trypsin residue backbone,³⁴ has a reduced trypsin inhibitor activity corresponding to ~60% of the wt-BPTI and BPTI-[5,55] at stoichiometry and a protein concentration of 280 n.M.¹⁹

^d Solubility in the same buffer as above but with the addition of 50 mM L-Arg + L-Glu.

^e The CD thermal melting curve could not be determined due to the strong absorption of arginine and glutamic acid.

f Protein solubility with 500 mM Arg-HCl added to the above buffer.

⁸ The trypsin activity could not be determined because the high arginine concentration inhibited trypsin activity.

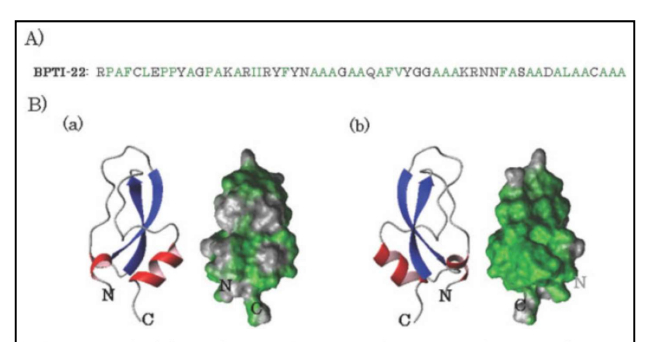


FIGURE 3 Hydrophobic residues in BPTI-22. A: One letter amino acid sequence of BPTI-22 with the hydrophobic residues (A,V,I,L,F,P) shown in green letters. B: Left, BPTI-22 ribbon model with α -helices colored red and β -strands colored blue. Right, surface representation of BPTI-22 with the hydrophobic area determined as low electrostatic potential regions according to MOL-MOL, solored green. The molecule is oriented with the β -sheet pointing to the back in (a) and to the front in (b). The N- and C-termini are labeled "N" and "C," respectively. The C-terminal end is located on the same face as a large hydrophobic patch shown in green, whereas the N-terminal end is on the opposite side of the molecule and is shown with a light gray letter "N" in panel (b).

- > The solubilization factor of all C-terminal tags was slightly higher than that of the respective N-terminal tags.
- ➤ The C-terminus of BPTI-22 is close to a large hydrophobic patch, whereas the N-terminus is located on the opposite side of the molecule, away from the hydrophobic patch.
- > Charged residues seem to act through repulsive electrostatic interaction and thus hamper intermolecular interaction arising from the hydrophobic cluster.

Solubility-enhancing tags – comparison of peptide and protein tags, conclusions

- ➤ Protein tags are inherently large and need to be correctly folded in order to enhance solubility.
- > Protein tags are often natural affinity tags.
- ➤ Peptide tags are small, and, importantly, they do not need to be folded, which provides a significant advantage over protein tags.
- The use of small tags (< 30 amino acids long) does not increase protein size substantially and reduces steric hindrance, which simplifies downstream structural and functional applications without the need to remove the tag.
- The solubilization enhancement effect depends on the size of the target protein. Solubility enhancement of fusion partners such as thioredoxin, GB1 is less pronounced for larger target proteins (above 25 kDa).

MANY TAGS SUFFER FROM THE SAME PROBLEM – THEY DO NOT FUNCTION EQUALLY WELL WITH ALL TARGET PROTEINS.

Otázka 2: Který tag/kotvu by jste využily pro zvýšení rozpustnosti proteinu bohatého na cysteiny?

Removal of fusion tags- the Achilles' heel of the fusion approach

All tags, whether small or large, have the potential to interfere with the biological activity of a protein, impede its crystallization (presumably due to the conformational heterogeneity allowed by the flexible linker region), be too large for NMR analysis, cause a therapeutic protein to become immunogenic or otherwise influence the target protein's behavior.

The fusion tags can be removed by:

- **≻** Chemical cleavage
 - **≻** Self cleavage
- > Enzymatic cleavage

Removal of fusion tags – chemical cleavage

➤ Rarely used.

Cyanogen bromide Met-X

Hydroxylamine Asn-Gly



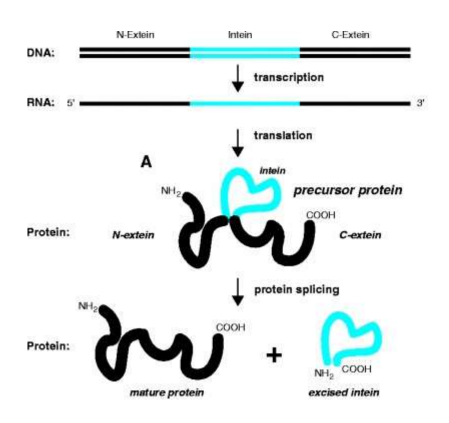
Amino – acid sequence of the *P. falciparum* C-terminal segment of CSP (PfCSP C-ter) fused to a purification tag (*Rais-Beghdadi et al., 1998*).

Chemical cleavage is a harsh method, efficient, but rather non-specific and may lead to unnecesary denaturation or modification of the target protein.

Removal of fusion tags - self - cleaving

➤ Use of self-cleaving fusion tags

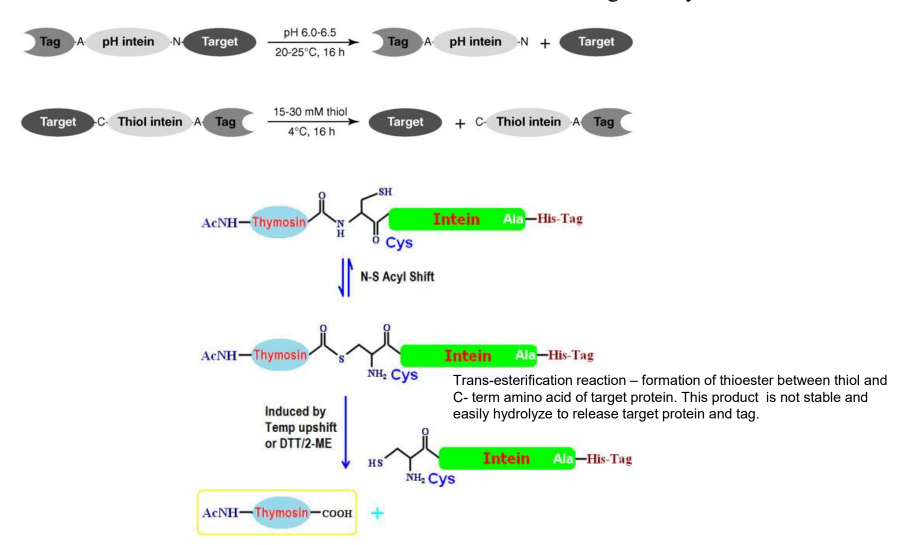
1. Inteins



Inteins (*int*ervening prot*eins*) are protein segments that can excise themselves from protein precursors in which the are inserted and rejoin the flanking regions.

> Self - splicing inteins can be mutated at the N- or C- terminal splice junction to yield self cleaving inteins, which can be used to mediate self cleaving of various tags.

- > Two categories of inteins:
 - inteins with pH-induced C-terminal cleaving activity
 - inteins with thiol-induced N- and C-terminal cleaving activity



Removal of fusion tags - self - cleaving fusion tag

2. HHHHHHH— SrtA60-206 —LPXT G— target protein

System based on the catalytic domain of *Staphylococcus aureus* **sortase A (SrtA).** SrtA cleaves the Thr-Gly bond at the conserved LPXTG motif in the substrates. Cleavage is inducible by adding calcium (cofactor of SrtA).

N(pro) —C X— target protein

N-terminal protease (N^{pro}) is the first protein of the pestivirus polyprotein. It posesses autoproteolytic activity and catalyzes the cleavage by switching from chaotropic to cosmotropic conditions.

FrpC modul (from G+ bacteria *Neisseria meningitides*): FrpC protein undergoes calcium – inducible autocatalytic processing at the peptide bond between residues Asp and Pro. Cleavage reaction is catalyzed by a self processing modul (SPM).

Vibrio cholerae secretes a large multifunctional autoprocessing repeats-in-toxin (MARTX) toxin that undergoes proteolytic cleavage during translocation into host cells. Proteolysis of the toxin is mediated by a conserved internal **cystein protease domain (CPD)**, which is activated upon binding of inositol hexakisphosphate.

Removal of fusion tags - self - cleaving fusion tag

Inteins (1)

- ➤ Uncontrolled in vivo cleavage or in complete in vitro cleavage
- > Target protein modification pH or thiols can modify the target protein
- ➤ Protein compatibility with cleaving conditions pH induced inteins
- ➤ Compared to the traditional protease based method, the intein-based approach requires fewer steps and lower costs.

Other system (2-5)

> Tested on limited number of cases

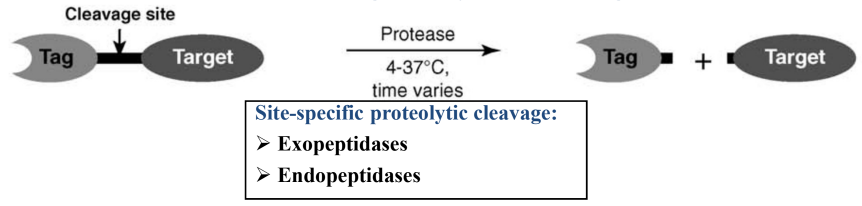
Table 3 General features of the five self-cleavage fusion systems discussed in the text

Self- cleaving tag	MW ^a (kDa)	Purification tag	Cleavage condition	Advantages	Disadvantages
Intein	51; 22; 17; 15 ^b	CBD, CBM, phasin, ELP	Thiols; pH and/or temperature shift	Flexible fusion and cleavage options; allowing generation of target protein with native sequence	Lack of solubility-enhancing capacity; in vivo cleavage; incomplete cleavage; miscleavage
SrtA	17	His-tag, biotin	5 mM Ca ²⁺	Potential of enhancing target protein expression and solubility	In vivo cleavage; incomplete cleavage; introduction of an extra Gly residue to the <i>N</i> -terminus of the target protein
N ^{pro}	19	His-tag	Kosmotropic conditions	Allowing generation of target protein with native sequence	Limited to proteins capable of refolding; in vivo cleavage; incomplete cleavage; long cleavage time
FrpC	26	His-tag, CBD	10 mM Ca ²⁺	Efficient and tightly controlled cleavage; insensitive to protease inhibitors	Lack of solubility-enhancing capacity; introduction of an extra Asp residue to the <i>C</i> -terminus of the target protein; single <i>C</i> -terminal fusion option
CPD	23	His-tag	50–100 μM InsP6	Potential of enhancing target protein expression and solubility; efficient and tightly controlled cleavage; insensitive to protease inhibitors	Introduction of up to four non-native residues to the <i>C</i> -terminus of the target protein; single <i>C</i> -terminal fusion option

a Molecular weight of the self-cleaving tag

b Inteins with different sizes are available

Removal of fusion tags – enzymatic cleavage



Exopeptidases (aminopeptidases and carboxypeptidases):

DAPase (TAGZyme)	Exo(di)peptidase	Cleaves N-terminal. His-tag (C-terminal) for purification and removal
Aeromonas aminopeptidase	Exopeptidase	Cleaves N-terminal, effective on M, L. Requires Zn
Aminopeptidase M	Exopeptidase	Cleaves N-terminal, does not cleave X-P
Carboxypeptidase A	Exopeptidase	Cleaves C-terminal. No cleavage at X-R, P
Carboxypeptidase B	Exopeptidase	Cleaves C-terminal R, K

APM, CPA and CPB release sequentially a single amino-acid from the N- or C- terminus of a protein until

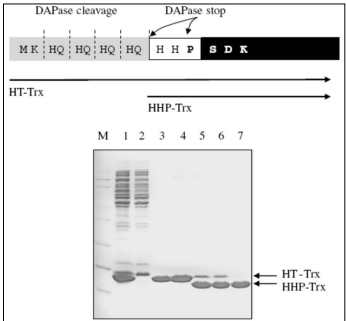
the stop site is reached.

TAGZyme system (Qiagen):

➤ DAPase (dipeptidyl aminopeptidase I)

TAGZyme stop points

Amino acid	DAPase stop point (♣) sequence*
Lysine (Lys, K)	Xaa-XaaXaa-Xaa ↓ <mark>lys-Xaa</mark>
Arginine (Arg., R)	Xaa-XaaXaa-Xaa ↓ Arg-Xaa
Proline (Pro, P)	Xaa-XaaXaa-Xaa ↓ Xaa-Xaa Pro-Xaa
Proline (Pro, P)	Xaa-XaaXaa-Xaa ↓ Xaa-Pro Xaa-Xaa
Glutamine (Gln, Q)1	Xaa-XaaXaa-Xaa ↓ Gln-Xaa



Arnau et al., 2006

Removal of fusion tags - enzymatic cleavage

Endopeptidases

> The enzymatic cleavage site has to be placed between the fusion tag and the target protein.

Enzyme	Cleavage site	Comments	, Protease site
Enterokinase	DDDDK*	Secondary sites at other basic aa	His _e V
Factor Xa	IDGR*	Secondary sites at GR	- ° (MBP) Target
Thrombin	LVPR*GS	Secondary sites. Biotin labeled for removal of the protease	protein
PreScission	LEVLFQ*GP	GST tag for removal of the protease	19
TEV protease	EQLYFQ*G	His-tag for removal of the protease	
3C protease	ETLFQ*GP	GST tag for removal of the protease	
Sortase A	LPET*G	Ca2+-induction of cleavage, requires an additional affinity tag	
		(e.g., his-tag) for on column tag removal	
Granzyme B	D^*X, N^*X, M^*N, S^*X	Serine protease. Risk for unspecific cleavage	

Enterokinase Asp-Asp-Asp-Lys/X

sequenceGSDYKDDDDK-X ₁ -ADQLTEEQIA of a GST-cal- modulin fusion protein was tested using 5 mg protein digested with 0.2 Uof enterokinase for 16 h at 37 °C		
Amino acid in position X ₁	Cleavage of enterokinase (%)	
Alanine	88	
Methionine	86	
Lysine	85	
Leucine	85	
Asparagine	85	
Phenylalanine	85	
Isoleucine	84	
Aspartic acid	84	
Glutamic acid	80	
Glutamine	79	
Valine	79	
Arginine	78	
Threonine	78	
Tyrosine	78	
Histidine	76	
Serine	76	
Cysteine	74	
Glycine	74	
Tryptophan	67	
Proline	61	

Table 4 Cleavage (%) of enterokinase through densitometry (Hosfield and Lu 1999) based on the amino acid residue X_1 . The

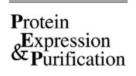
Removal of fusion tags - enzymatic cleavage

A critical review of the methods for cleavage of fusion proteins with thrombin and factor Xa

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The purpose of this review was to demostrate that both thrombin and factor Xa can hydrolyze a variety of peptide bonds within the fused proteins of interest.

Sequences cleaved by thrombin in polypeptide hormones

Polypeptide homones ^a	Sequence cleaved	
Secretin	ELSLSRL R DSA	
Secretin	ELSLSRLR (much slower than above)	
Vasoactive intestine polypeptide	DNYTRLRK	
Vasoactive intestine polypeptide	YTRLRKQM	
Choleocystokinin	APSGRVSM	
Choleocystokinin	VSMIKNLQ	
Dynorphin A	RIRPKLKW	
Somatostatin-28	AMAPRERK	
Somatostatin-28	NFFWKTFT	
Gastrin releasing peptide	KMYP R GNH	
Salmon calcitonin	QTYPRTNT	

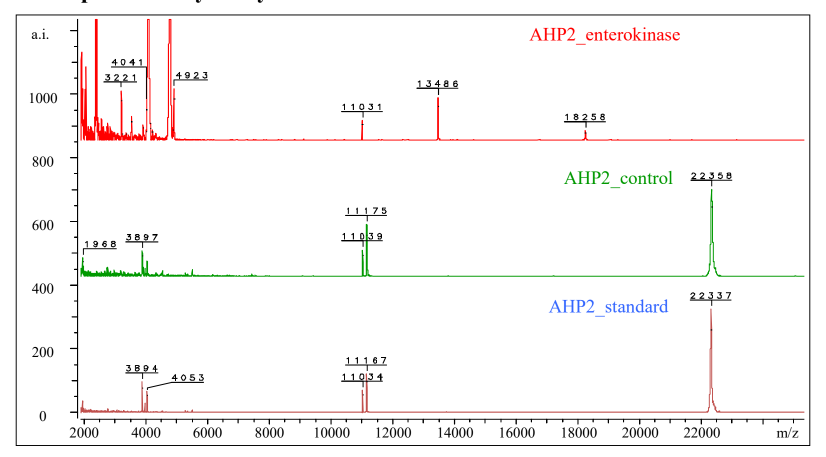
 $^{^{}a}$ The reaction mixtures contained 0.5 NIH units thrombin and 1.0 nmol peptide in 20 μ L of 50 mM NH₄CO₃, pH 8.0, at 25 $^{\circ}$ C. The conditions were designed to obtain an enzyme/substrate ratio of 1:60 (w/w).

Accuracy of cleavage has to be precisely verified!

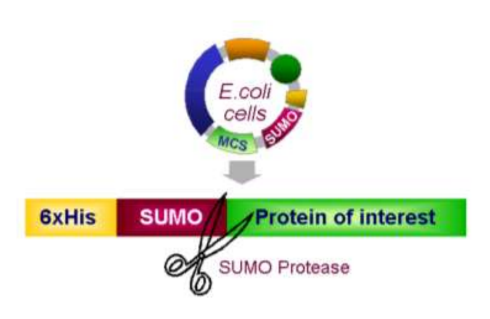
pRSETB::AHP2 Enterokinase cleavage site

N'MRGSHHHHHHGMASMTGGQQMGRDLYDDDDKDPSSRSAAGTMEFMDALIA......GIVPQVDIN C'
Theoretically: 3,4 kDa

Intact mass spectrometry analysis



SUMO gene fusion system



SUMO protease recognizes the tertiary structure of SUMO rather than an amino acid sequence.

As a result, SUMO protease never cleaves within the fused protein of interest!

Removal of fusion tags - enzymatic cleavage

- **Optimization of protein cleavage conditions** (mainly enzyme-to-substrate ratio, temperature, pH, salt concentration, length of exposure).
- Cleavage efficiency (Optimization is needed. The efficiency varies with each fusion protein in an unpredictable manner, probably due to aggregation or steric issues; the problem can be solved by introducing short linkers between the protease site and the fusion tag).
- ➤ Unspecific cleavage (SOLUTION: optimization of protein cleavage conditions or using re-engineered proteases with increased specificity such as ProTEV and AcTEV proteases).
- Precipitation of the target protein when the fusion partner is removed (so-called soluble aggregates; SOLUTION: another approach for protein solubilization has to be found).
- > Failure to recover active or structurally intact protein
- ➤ Target protein modification (some proteases like thrombin, TEV, Precision leave one or two amino-acids on the target protein near the cleavage site).
- **Re-purification step** is needed to separate the protease from target protein.
- High cost of proteases

The alternative is to leave the tag in place for structural analysis:

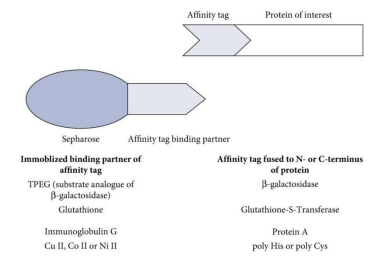


The small tags are a better choice in structural and functional analysis of proteins.

Otázka 3: Jaký je rozdíl mezi inteinem a samo-vyštěpujícím tagem odvozeným od inteinu?

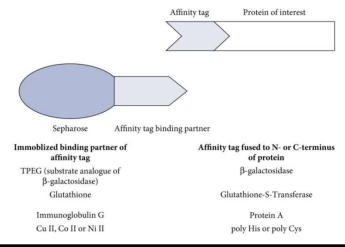
Affinity chromatography (AC)

A type of adsorption chromatography, in which the molecule to be purified is specifically and reversibly adsorbed to a complementary binding substance (ligand, L) immobilized on an insoluble support (matrix, M).



- > AC has a concentrating effect, the high selectivity of separations derived from the natural specificities of the interacting molecules.
- AC can be used (1) to purify substances from complex biological mixtures, (2) to separate native forms from denatured forms of the same substance, and (3) to remove small amounts of biological material from large amounts of contaminating substances, (4) and to isolate protein complexes from the native source.
- ➤ the first application was in 1910 (adsorption of amylase onto insoluble starch) but it developed during the 1960s and 1970s.

Affinity tags and affinity purification



Tag	Residues	Sequence	Size (kDa)
Poly-Arg	5–6	RRRRR	0.80
	(usually 5)		
Poly-His	2-10	НННННН	0.84
	(usually 6)		
FLAG	8	DYKDDDDK	1.01
Strep-tag II	8	WSHPQFEK	1.06
c-myc	11	EQKLISEEDL	1.20
S-	15	KETAAAKFEROHMDS	1.75
HAT-	19	KDHLIHNVHKEFHAHAHNK	2.31
3x FLAG	22	DYKDHDGDYKDHDIDYKDDDDK	2.73
Calmodulin-binding peptide	26	KRRWKKNFIAVSAANRFKKISSSGAL	2.96
Cellulose-binding domains	27-189	Domains	3.00
			20.00
SBP	38	MDEKTTGWRGGHVVEGLAGELEOLRARLEHHPOGOREP	4.03
Chitin-binding domain	51	TNPGVSAWOVNTAYTAGOLVTYNGKTYKCLOPHTSLAGWEPSNVPALWOLO	
Glutathione S-transferase	211	Protein	26.00
Maltose-binding protein	396	Protein	40.00

A tag is fused to the N- or C-terminus of the protein of interest to facilitate purification, which relies on a specific interaction between the affinity tag and its immobilized binding partner. Genetically engineered fusion tags allow the purification of virtually any protein without any prior knowledge of its biochemical properties.

Purification tags

Affinity tags

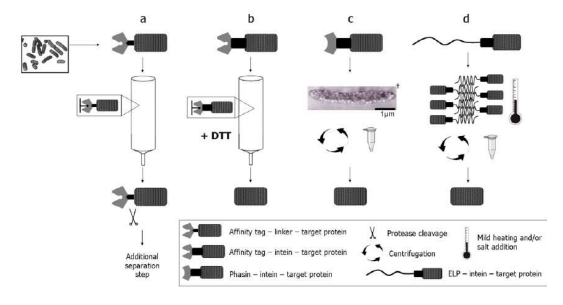
Affinity tag	Matrix
Poly-Arg	Cation-exchange resin
Poly-His	Ni ²⁺ -NTA, Co ²⁺ -CMA (Talon)
FLAG	Anti-FLAG monoclonal antibody
Strep-tag II	Strep-Tactin (modified streptavidin)
c-myc	Monoclonal antibody
S	S-fragment of RNaseA
HAT (natural histidine affinity tag)	Co ²⁺ -CMA (Talon)
Calmodulin-binding peptide	Calmodulin
Cellulose-binding domain	Cellulose
SBP	Streptavidin
Chitin-binding domain	Chitin
Glutathione S-transferase	Glutathione
Maltose-binding protein	Cross-linked amylose

- > Traditional purification tags
- ➤ The tag binds strongly and selectively to an immobilized ligand on a solid support.
- ➤ After optimization one could achieve > 90% purity.

Non - chromatographic tags

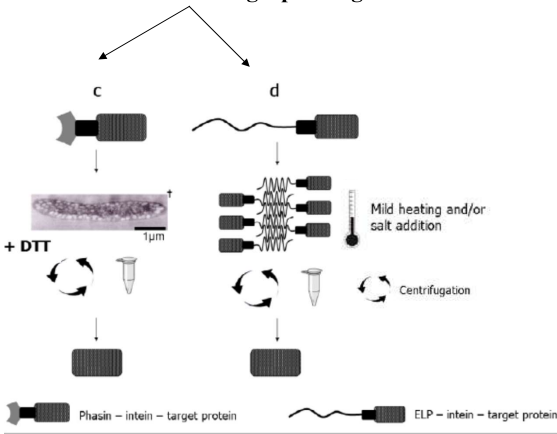
<u> 1ag</u>	Matrix
ELP	None
PHB	Intracellular PHA granules
annexin B1	None

- ➤ These tags can eliminate affinity resin. Proteins are isolated by other nonchromatographic methods (centrifugation, filtration)
- > typically combined with self-cleaving tags
- > 75 % 95 % purity



Purification tags

Non - chromatographic tags



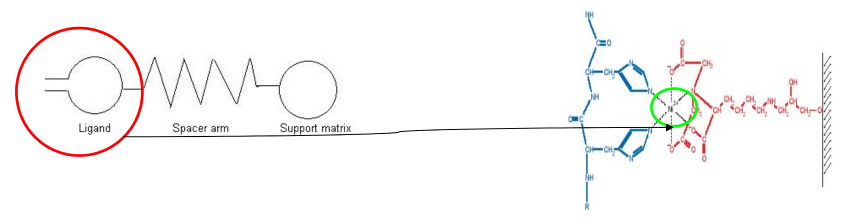
The PHB system (c):

- ➤ PHB (polyhydroxybutarate): subclass of biodegradable polymers produced in various organisms, use as storing excess carbon.
- ➤ The system includes *in vivo* production of PHB small granules (from the plasmid carrying PHB-synthesis genes).
- ➤ Target protein in fusion to self cleaving phasin tag.
- ➤ Tagged protein binds to the PHB particles via phasin tag, which allows the granules and the tagged protein to be co-purified via centrifugation.
- ➤ DTT induced cleaving activity of intein and thus elution of the target protein.

The ELP system (d):

➤ ELP (elastin-like polypeptide) selectively and reversibly precipitates in response to changes in temperature and buffer salts. This allows soluble and insoluble contaminants to be removed by filtration or centrifugation.

Components of a matrix for affinity chromatography

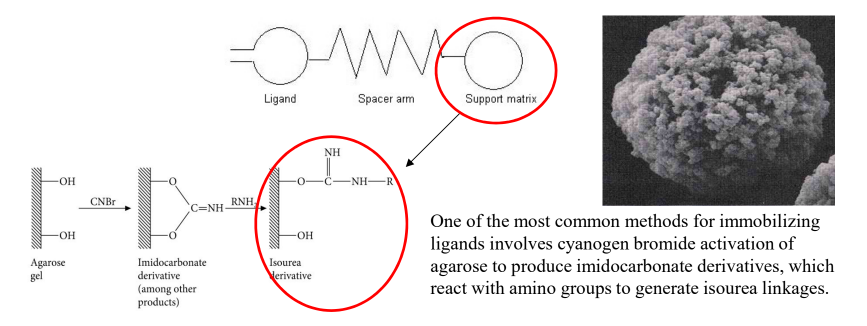


Ni²⁺ NTA sepharose

A ligand

- \triangleright The dissociation constant (Kd) for the ligand target complex should ideally be in the range 10^{-4} to 10^{-8} M in free solution to allow efficient elution under conditions which will maintain protein stability.
- ➤ A ligand has to be attached to the matrix with a suitable chemically reactive group. The mode of attachment must not compromise the reversible interaction between the ligand and protein.

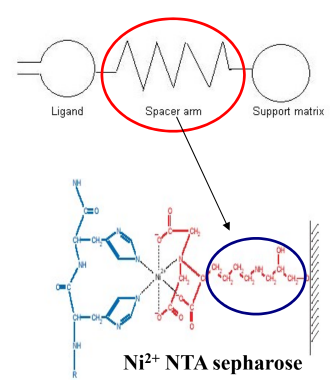
Components of a matrix for affinity chromatography



A matrix

- ➤ Typically, a macroporous polysaccharide bead such as agarose, that provides a porous structure so that there is an increased surface area to which the target molecule can bind.
- A matrix has a suitable attachment site for the ligand. Typically matrices are chemically activated to permit the coupling of the ligand. A number of activation methods are available which depend on the nature of the matrix and the availability of compatible reactive groups on the ligand.

Components of a matrix for affinity chromatography



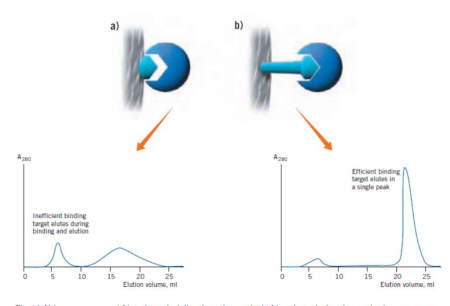
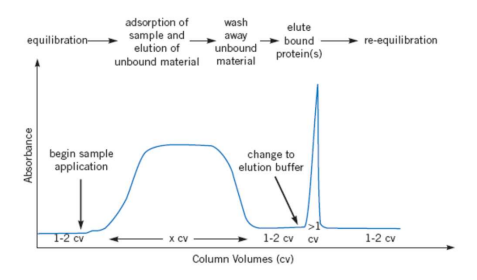


Fig. 56. Using spacer arms. a) Ligand attached directly to the matrix. b) Ligand attached to the matrix via a spacer arm.

Spacer arm

A spacer arm will be required in cases where direct coupling of the ligand to the matrix results in steric hindrance and subsequently the target protein will fail to bind to the immobilized ligand efficiently. The introduction of a spacer arm between the ligand and the matrix minimizes this steric effect and promotes optimal adsorption of the target protein to the immobilized ligand.

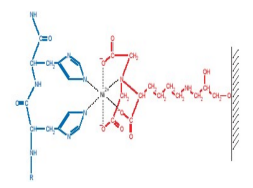
Typical affinity purification steps



- In the equilibration phase, buffer conditions are optimized to ensure that the target molecules interact effectively with the ligand and are retained by the affinity medium as all other molecules wash through the column.
- During the washing step, buffer conditions are created that wash unbound substances from the column without eluting the target molecules or that re-equilibrate the column back to the starting conditions (in most cases the binding buffer is used as a wash buffer).
- ➤ In the elution step, buffer conditions are changed to reverse (weaken) the interaction between the target molecules and the ligand so that the target molecules can be eluted from the column.

Affinity chromatography - Immobilized metal ion affinity chromatography (IMAC)

- > The most common purification tag is typically composed of six consecutive histidine residues.
- ➤ Histidine, cysteine, and tryptophan residues are known to interact specifically with divalent transient metal ions such as Ni²⁺, Cu²⁺, Co²⁺, and Zn²⁺.
- ➤ Histidine is the amino acid that exhibits the strongest interaction with immobilized metal ion matrices as the electron donor groups on the histidine imidazole ring readily form coordination bonds with an immobilized transition metal.



M (kDa)
170 —
116 —
86 —
56 —

27 —
20 —

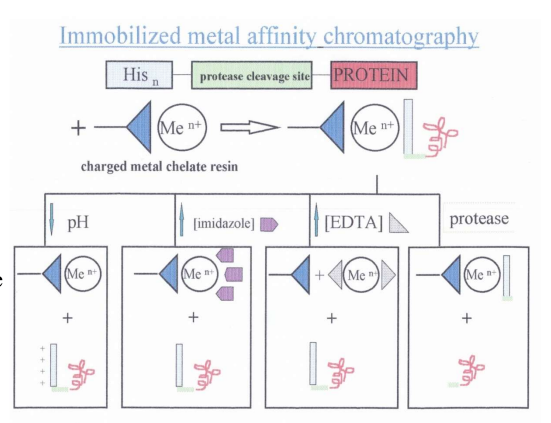
Binding strength of His tag to metal ions:

$$Cu^{2+} > Ni^{2+} > Zn^{2+} \sim Co^{2+}$$

Zn²⁺ Ni²⁺ Co²⁺ Cu²⁺
(Zouhar et al., 1999)

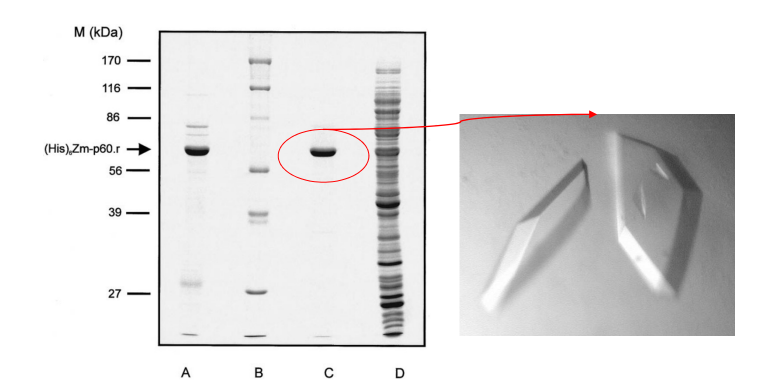
- > IMAC can be used under native and/or denatured conditions.
- A highly purified protein can often be obtained in one or, at most, two purification steps.

- ➤ Optimal binding of recombinant protein with metal ion is achieved at pH 7–8.
- ➤ Buffers with a high salt concentration (0.5–1 M NaCl) reduce nonspecific electrostatic interaction.
- ➤ Nonionic detergents or glycerol reduce nonspecific hydrophobic interactions.
- ➤ Elution of contaminating proteins can be achieved by lowering the pH or using low concentrations of imidazole.
- ➤ Elution of tagged protein is achieved at high imidazole concentrations (0–0.5 M), by strongly decreasing the pH, or by using EDTA.



One-step purification of maize β -glucosidase

- > Perfusion matrix: POROS MC/M
- ➤ Functional group: iminodiacetate, metal ion Zn²⁺
- > Removing contaminated proteins: linear gradient of imidazole (0–50 mM) and pH (pH 7-6.1)
- > Protein elution: 0.1 M EDTA
- ➤ 80% recovery, 95 fold purification
- > Common production and isolation of the wild type protein and soluble mutant form for enzymatic measurements and crystallization.

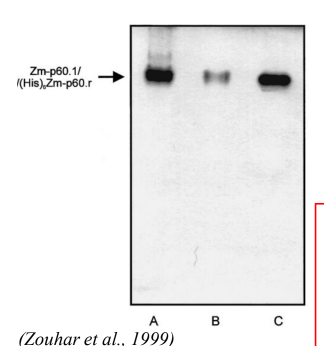


(Zouhar et al., 1999)

- Purification of proteins expressed in inclusion bodies.
- Purification in a high concentration of urea or guanidine chloride.
- Result is a pure protein, but in a denatured form (sufficient for immunization).

Recovery of native conformers (necessary for functional and structural analysis):

- ➤ Binding to the column under strong denaturing conditions (8 M urea)
- > Two possibilities of renaturation:
- 1. The protein is eluted from the column and renatured by dialysis or rapid dilution in renaturing buffers.
- 2. Renaturation of the protein bounded to the column (matrix assisted refolding procedure): gradient from denatured to renatured buffers or pulsion renaturation (8-0M urea).



Identification of properly refolded (His)₆Zm-p60.1 (maize β-glucosidase) using 10% native PAGE, followed by activity in gel staining:

A = crude protein extract prepared from maize seedlings containing the native enzyme

 $B = (His)_6 Zm-p60.1$, renatured product (matrix assisted refolding procedure – 23 renaturing cycles)

 $C = (His)_6 Zm-p60.1$ purified by native IMAC

 K_M (His)₆Zm-p60.1 purified by native IMAC: **0.64 ± 0.06 mM**

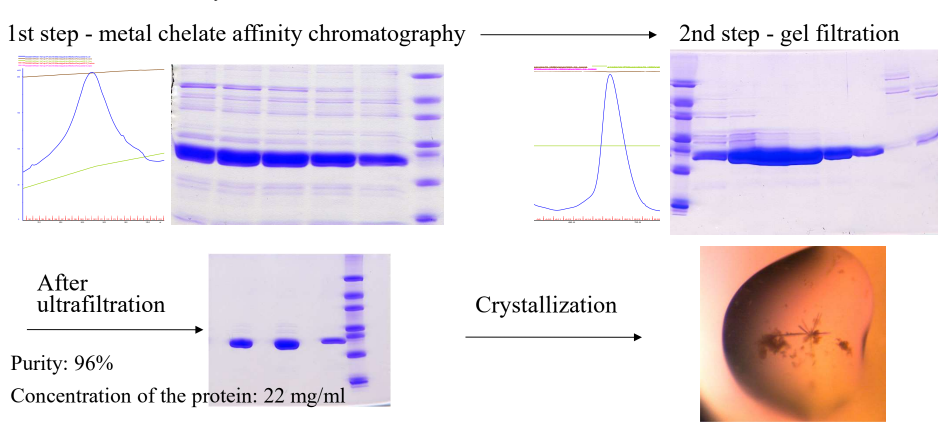
 K_M (His)₆Zm-p60.1 renatured product: **0.6 ± 0.08 mM**

Determination of v_{max} and k_{cat} was hampered by the fact that the refolding process yielded a number of improperly folded polypeptides.

Two-step purification of Arabidopsis histidine phosphotransfer protein 5

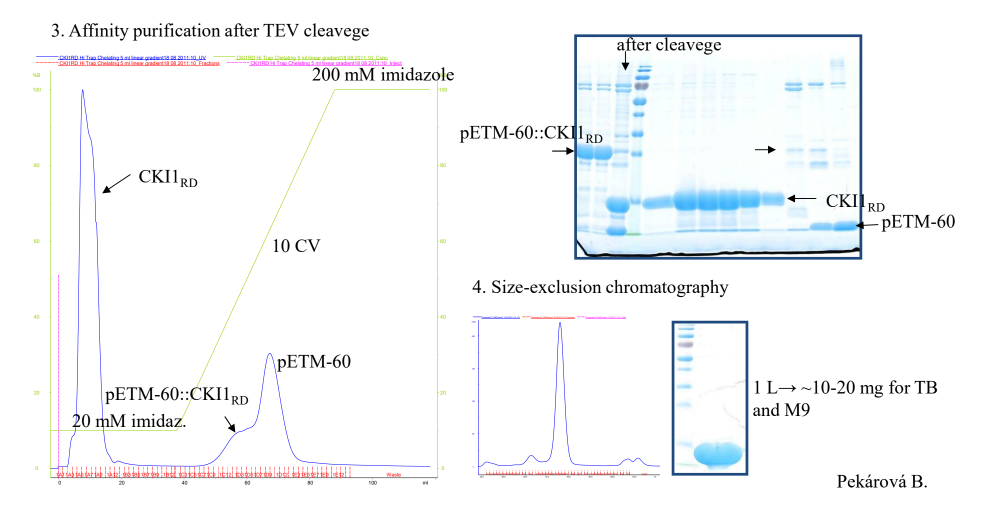
(His)₆AHP5

- ➤ IMAC matrix: highly cross-linked spherical agarose
- Functional group: nitrilotriacetic acid, metal ion Ni²⁺
- > Removing contaminated proteins: linear gradient of imidazole (20–500 mM)
- > Protein elution: 130 mM imidazol
- ➤ Common production and isolation of the wild type protein for protein-protein interaction measurements and crystallization.



Four-step purification of *Arabidopsis* CKI1_{RD}

- 1. Affinity purification (IMAC)
- 2. Tag removal (TEV protease)
- Ub-SGSG-HisTag-SA-TEV-AME-CKI1
- 3. Affinity purification (IMAC)
- 4. Size exclusion chromatography



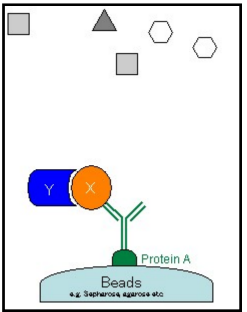
Otázka č.4: Jakými metodami se izolují proteiny fúzované s nechromatografickými tagy/kotvami?

Affinity purification for studying protein-protein interaction

- ➤ Affinity purification provides a high-efficiency method for isolation of interacting proteins and protein complexes:
 - > Co-immunoprecipitation
 - ➤ GST (or His) pull-down
 - > Tandem affinity purification
 - > Testing known protein-protein interaction.
 - > Identification of novel protein-protein interactions.

Co-immunoprecipitation (Co-IP)

- The principle: If protein X is immunoprecipitated with an antibody of X, then protein Y, which is stably associated with X in vivo, may also be precipitated. This precipitation of protein Y, based on a physical interaction with X, is referred to as co-immunoprecipitation.
- An obvious advantage is that complexes are isolated in the state closest to the physiological condition.
- ➤ When a good quality antibody of X is available, Co-IP is a fast method and there is no need to clone and express the component(s) of the complex.

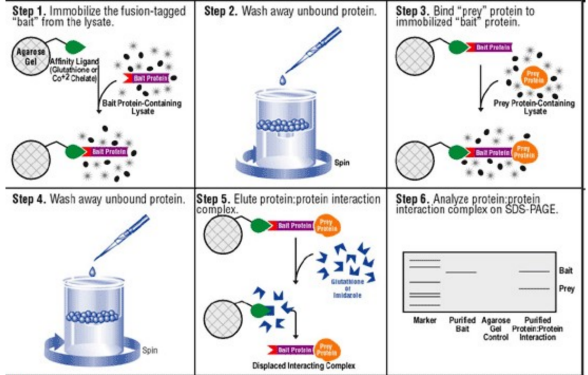


- 1. Cell lysis under mild conditions that do not disrupt protein-protein interactions (using low salt concentrations, non-ionic detergents, protease inhibitors, phosphatase inhibitors).
- 2. The protein of interest (X) is specifically immunoprecipitated from the cell extracts (using an antibody specific to the protein of interest or to its fusion tag).
- 3. The antibody-protein(s) complex is then pelleted usually using protein-A or G sepharose, which binds most antibodies.
- 4. Eluted immunoprecipitates are then fractionated by SDS-PAGE.
- 5. A protein of known identity is most commonly detected by performing a western blot. Identification of novel interaction is carried out by mass spectrometry analysis.

Pull-down assay

- ➤ Pull-down assays are a common variation of co-immunoprecipitation and are used in the same way, but pull down does not involve using an antibody specific to the target protein being studied.
- They are used for purification of multiprotein complexes *in vitro*.
- ➤ The target protein is expressed in *E. coli* as GST fusion and immobilized on glutathione-sepharose beads (GST alone is often used as a control).
- ➤ Cellular lysate is applied to the beads or column, and the target protein competes with the endogenous protein for interacting proteins, forming complexes in vitro.
- > Centrifugation is used to collect the GST fusion probe protein and adhering proteins.

The complexes are washed to remove nonspecifically adhering proteins.



- Free glutathione is used to elute the complexes from the beads, or alternatively the beads with attached complexes are boiled directly in an SDS-PAGE sample buffer.
- The proteins are resolved on SDS-PAGE and processed for further analysis.

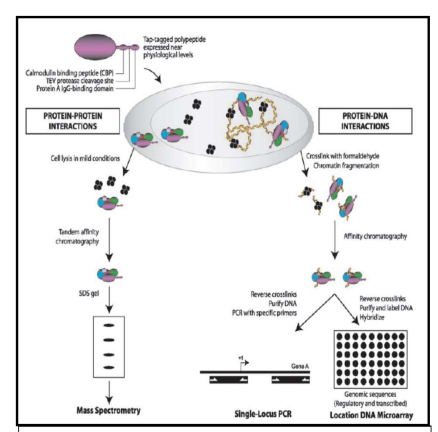
Tandem affinity purification (TAP)

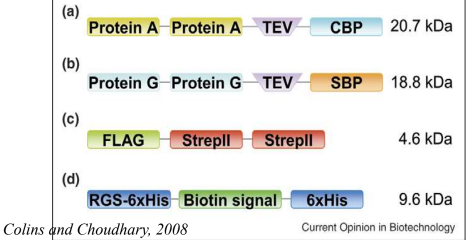
Two-step purification strategy in order to achieve higher purity of isolated multiprotein complexes under near physiological conditions.

This method was originally developed for use in yeast and quickly adapted to higher eukaryotes such as insect cells, human cells and plant cells.

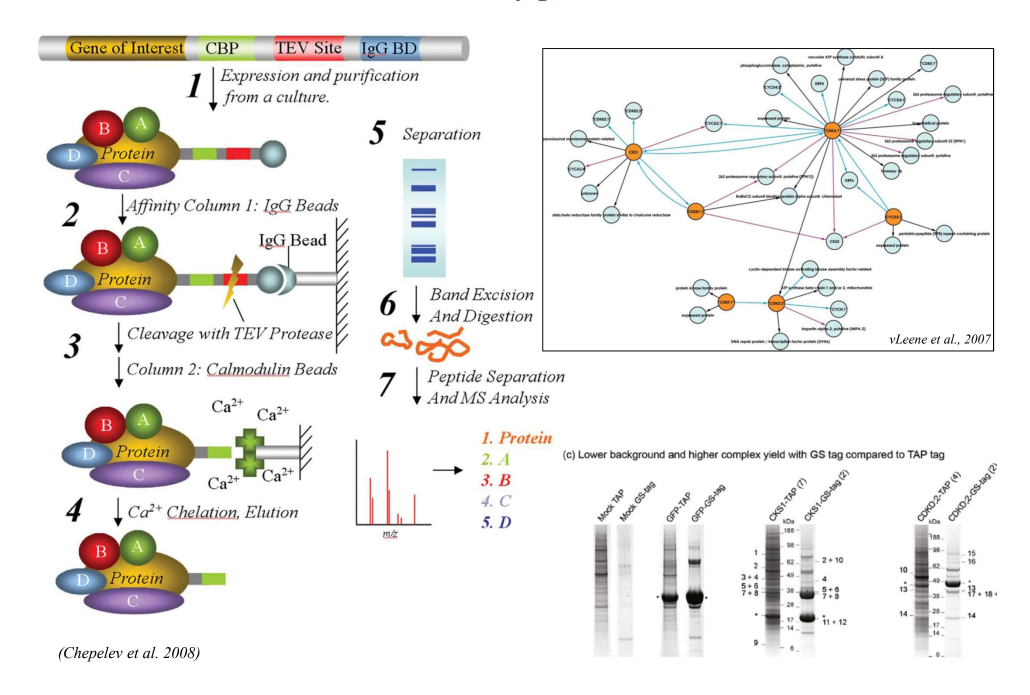
Examples of TAP (tandem affinity peptides) tags

TAP tag: a double affinity tag (highly specific) which is fused to a protein of interest as an efficient tool for purification of native protein complexes.





Tandem affinity purification

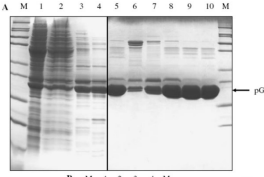


Affinity purification for studying protein-protein interaction

- ➤ An affinity tags can influence protein-protein interactions (testing N- and C-terminal fusions).
- > Loss of weak or transient protein-protein interactions.
- ➤ Non-specificity: controls, affinity tags with higher specificity
- ➤ Verification of newly identified interactors by other methods and biologically relevant mutants.

Comparison of a standard purification process with affinity purification

- ➤ Generally, the yield and efficiency of any specific purification procedure depends on the level of optimization developed for individual proteins and the method. It is therefore recommended to use the data presented in different comparisons as indicative rather than definitive, which it is not e.g., identical elution conditions are optimal for different proteins.
- ➤ Standard chromatographic methods include several steps to obtain a relative pure protein. This results in a time-consuming procedure and a relatively low yield of recovery (typically 50 % of the starting material for optimized processes).
- ➤ The yields obtained in purification of proteins using affinity chromatography can be over 90 % and include a reduced number of steps.



Comparison of purification strategies for recombinant pGAP

Purification step	Total volume (ml)	Activity (U/ml)	Total activity (U)	Yield (%)
Standard purification (untagged	dpGAP)			
Cell extract	750	2.0	1463	100
Phenyl-Sepharose HS	400	2.6	1040	71
Phenyl-Sepharose LS	160	5.6	888	61
Q Sepharose HP	57	10.3	587	40
IMAC purification (his-tag pG.	AP)			
Cell extract	120	19.0	2280	100
Ni-NTA Sepharose	84	26.0	2184	96

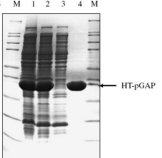


Fig. 2. Comparison of purification strategies for recombinant pGAP. A *B. amyloliquefaciens* pyroglutamyl aminopeptidase (pGAP) was produced in *E. coli* with and without an N-terminal his-tag (HT-pGAP, tag sequence: MEP(H)₆L). For untagged pGAP, purification included ammonium sulfate precipitation and two consecutive separation steps using phenyl-Sepharose. Subsequently, a desalting step using a Sephadex G-25 F column and a final step using Q Sepharose HP were performed. For HT-pGAP, purification was performed with a single IMAC step. (A) Standard purification of pGAP. Lane M: MWM (Novex); lane 1: cell extract; lane 2: supernatant fraction of cell extract; lane 3: pool from first phenyl-Sepharose step; lane 4: pool from the second phenyl-Sepharose step; lane 5: pool after desalting; lanes 6–10: several fractions from Q Sepharose HP containing pGAP. (B) IMAC purification of HT-pGAP. Lane M: MWM; lane 1: cell extract; lane 2: supernatant fraction of the cell extract; lane 3: flow through fraction from the IMAC; lane 4: eluted HT-pGAP. See Table 2 for process yields.